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# Methods for the Estimation of the NICE Cost Effectiveness Threshold 

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# Methods for the Estimation of the NICE Cost Effectiveness Threshold 

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## Summary

## 1 Introduction

1.1 NICE's comparison of the incremental cost effectiveness ratio (ICER) of a new technology, which is more costly than existing alternatives, with the cost-effectiveness threshold is important in assessing whether the health expected to be gained from its use exceeds the health expected to be forgone elsewhere as other NHS activities are displaced (i.e. whether the new technology is cost effective).
1.2 When NICE issues positive guidance for a new intervention which imposes additional costs on the NHS, the resources required to deliver it must be found by disinvesting from other interventions and services elsewhere. This displacement will inevitably result in health decrements for other types of individual. Thus the threshold represents the additional cost that has to be imposed on the system to forgo 1 QALY of health through displacement.
1.3 Currently NICE uses a threshold range of $£ 20,000$ to $£ 30,000$ QALY gained. There have been a number of calls for further research on the value of the threshold.
1.4 This report details a 2-year project, funded by the NIHR and MRC Methodology Research Programme, to develop methods to estimate the NICE cost effectiveness threshold.
1.5 Given NICE's remit, empirical research on the threshold requires some key characteristics:

- Reflect the expected health effects (in terms of length and quality of life) of NICE guidance through the displacement decisions taken across the NHS rather than what specific services are (or could have been) displaced.
- Facilitate regular updates, based on routinely available data, to reflect NHS changes such as real overall expenditure and ppoductivity. This would encourage accountability through scrutiny by stakeholders and provide predictability for technology manufacturers' investment decisions.
- The nature of service displacement and the magnitude of the health forgone will depend on the scale of the budget impact which should, ideally, be reflected in the value of the threshold.
- Methods should recognise the inevitable uncertainty relating to the evidence currently available for the threshold and reflect its implications for policy.


## 2 Study methods

2.1 The@im was to develop methods to estimate the NICE cost-effectiveness threshold making use of routinely available data. Objectives were:

Informed by relevant literature, to provide a conceptual framework to define the threshold and the basis of its estimation.
ii. Using programme budgeting data for the English NHS, to estimate the relationship between changes in overall NHS expenditure and changes in mortality.
iii. Extend the measure of benefit in the threshold to QALYs by estimating the quality of life (QoL) associated with additional years of life and the direct impact of health services on QoL.
iv. Present the best estimate of the cost effectiveness threshold for policy purposes.
2.2 Earlier econometric analysis estimated the relationship between differences in primary care trust (PCT) spending and associated disease-specific mortality. Expenditure came from programme
budgeting data which allocates the entire volume of health care expenditure to broad programme budget categories (PBCs) according to primary diagnosis.
2.3 This research extended this in several ways including estimating the impact of marginal increases or decreases in overall NHS expenditure on spending in each of the 23 PBCs. These were linked to changes in mortality outcomes by PBC across 11 PBCs.
2.4 The results of the econometric analysis were translated into broader effects in terms of QALYs. The first stage linked estimated effects on mortality to life years taking into account the 'counterfactual' deaths that would have occurred if the population in a given PBC faced the same mortality riss as the general population. The second stage accounted for the health (QALY) effects of changes in mortality due to changes in expenditure reflecting how QoL differs by age and gender. The third stage incorporated those effects on health not directly associated with mortality and life year effects (i.e., the 'pure' QoL effects) to estimate an overall cost per QALY threshold. This effectively used the estimates of mortality and life year effects as 'surrogate outcomes' for a more complete measure of the health effects of a change in expenditure.

## 3 Central or 'best' estimate of the threshold

3.1 The most relevant threshold is estimated using the latest available data (2008 expenditure, 2008-10 mortality). The central or 'best' threshold is estimated to be $£ 18,317$ per QALY.

## 4 Which PBCs have the greatest influence on the overall threshold?

4.1 Although the 11 PBCs where a mortality effect of changes in expenditure could be estimated only account for $36 \%$ of the change in overall expenditure, they account for $80 \%$ of the overall health effects. The other 12 PBCs, where mortality effects could not be estimated, account for the greater part of a change in overall expenditure ( $64 \%$ ) but only $20 \%$ of the overall health effects, i.e., the cost per QALY estimates associated with a change in expenditure in these PBCs are, in general, much higher.
4.2 Insofar as investment and disinvestment opportunities in these PBCs might have been more valuable (offered greater improvement in QoL) than suggested by the implied PBC thresholds, the overall QALY effects will tend to be underestimated and the overall cost per QALY threshold will be overestimated.
4.3 The overall threshold of $£ 18,317$ may be especially conservative (i.e., likely to be overestimated) with respect to health effects in PBC5 (Mental Health Disorders), which accounts for a large proportion of the change in overall expenditure ( $25 \%$ ) and contributes most to the overall health effects ( $9 \%$ ) compared to these other PBCs. The cost per QALY associated with this PBC is based on an extrapolation rather than observations of the direct impact of changes in expenditure on QoL. Available evidence suggests that the investment and disinvestment opportunities in mental health are likely to have been much more valuable than its implied cost per QALY.

## 5 How uncertain are the estimates and what are the implications?

5.1 Simulation methods were used to reflect the combined uncertainty in the various estimates from the econometric analysis. This indicated that the probability that the overall threshold is less than $£ 20,000$ per QALY is 0.64 and the probability that it is less than $£ 30,000$ is 0.92 .
5.2 As the consequences of overestimating the threshold are more serious than underestimating it in terms of population health, a policy threshold will be lower than the mean of the cost per QALY threshold (i.e., lower than $£ 18,317$ ) to compensate for the more serious consequences of overestimating the 'true' value.
5.3 There were other ('structural') sources of uncertainty associated with the estimated threshold, specifically relating to the choice of econometric models and identification of causal effects. Although all the models passed the relevant tests of validity, there remained some uncertainty about the validity of the instruments. This structural uncertainty constituted a greater part of the overall uncertainty associated with the mortality effects of changes in expenditure, but the central estimate of the cost per QALY threshold was robust to this uncertainty.
5.4 The method of analysis used to link the effects of changes in expenditure on mortality to a fuller measure of health expressed in QALYs was also subject to uncertainty. A preferred analysis (or scenario) was identified as making the best use of available information, with assumptions appearing more reasonable than the available alternatives and providing a more complete picture of the likely health effects of a change in expenditure.
5.5 A critical issue is whether, on balance, the central or best estimate is likely to be an underestimate or overestimate of the cost per QALY threshold. Although other assumptions and judgments are possible that retain some level of plausibility, they do not all favour a higher threshold. Indeed, when considered together, they suggest that, on balance, the central or best estimate of $£, 18,317$ is, if anything, likely to be an overestimate.

## 6 The impact of investment, disinvestment and non marginal effects

6.1 The central estimate of the cost per QALY threshold is based on estimates of the health effects of changes in expenditure across all 152 PCTs, some of which will be making investments (where expenditure is increasing) and others making disinvestments (where expenditure is reduced or growing more slowly).
6.2 The threshold is, however, likely to differ across these different types of PCT. It would be expected that, other things equal, more expenditure would increase health but at a diminishing rate. Therefore, the amount of health displaced by disinvestment would be expected to be greater, and the associated threshold lower than the central estimate. Conversely, the health gained from investment would be expected to be lower, and the associated threshold higher.
6.3 This was examined by re-estimating the outcome and expenditure effects separately for those PCTs where their actual budget is under the target allocation from the Department of Health resource allocation formula (i.e., those under greater financial pressure and more likely to be disinvesting than investing), and those that are over target (under less financial pressure and more likely to be investing than disinvesting).
6.4 The results confirm these expectations: the health effects of changes in expenditure are greater when PCTs are under more financial pressure and are more likely to be disinvesting then investing. The analysis suggests that budget impact not only displaces more valuable activities within each PBC but that overall expenditure tends to be reallocated to PBCs which can generate more health. Although further research might enable a quantitative assessment of how the relevant threshold should be
adjusted for the scale of budget impacts, the qualitative assessment seems clear: the central estimate of the threshold is likely to be an overestimate for all technologies which impose net costs on the NHS (almost all technologies appraised by NICE); and the appropriate threshold to apply should be lower for technologies which have a greater impact on NHS costs.

## 7 How does the threshold change with overall expenditure?

7.1 The same methods are used to consider how the cost per QALY threshold is likely to have changed from 2007 to 2008 as overall expenditure has increased. This provides some insights into how the threshold might be expected to change over time as, for example, overall expenditure and NHS productivity changes.
7.2 This has implications for a judgement about the appropriate frequency of periodic reassessment of the cost per QALY threshold. Other things equal, the threshold would be expected to increase following a rise in overall expenditure, although this will depend on whether there is discretion over how additional resources can be spent. However, insofar as the productivity of those activities that are valuable to the NHS also improves through innovation, the threshold will tend to fall. So the net impact of these two countervailing effects on the threshold cannot be determined a priori.
7.3 Differences in the estimated thresholds between 2007 and 2008 are assessed. Although overall expenditure increased by $6 \%$ between 2007 and 2008 which represented real growth of $2 \%$ in 2007 prices, the overall threshold for all 23 PBCs fell by $2 \%$ (in nominal terms and by $5 \%$ in real terms.
7.4 The reasons are complex but reflect changes in productivity, which differ across PBCs, but also a general reallocation of a change in overall expenditure towards those PBCs that appear more valuable in 2008. Given the uncertainty in estimation, subtle differences between 2007 and 2008 should not be over-interpreted. This analysis does suggest, however, that the overall threshold will not necessary increase with growth in the real or even nominal NHS budget. This suggests that the threshold is more likely to fall at a time when real budget growth is flat or falling and PCTs find themselves under increasing financial pressure.

8 What type of health is forgone by approval of a new technology?
8.1 The methods of analysis can identify not only how many QALYs are likely to be forgone across the NHS as a consequence of approving a technology which imposes additional costs on the NHS, but also where those QALYs are likely to be forgone and how they are made up, i.e., the additional deaths, life years lost and the QoL impacts on those with disease.
8.2 As an example, based on the 2008 central estimate of the cost per QALY threshold ( $£ 18,317$ ), the approval of Ranibizumab for the treatment of diabetic macular oedema (prior to the patient access scheme agreement) would have imposed additional annual costs of up to $£ 80 \mathrm{~m}$ on the NHS each year and been likely to displace 4,367 QALYs elsewhere in the NHS. This forgone health is likely to be made up of 295 additional deaths and 1,337 life years forgone, most of which are likely to occur in Circulatory, Respiratory, Gastro-intestinal and Cancer PBCs. However, much of the total health effect of these additional costs (3,509 QALYs) is associated with QoL forgone during disease which is most likely to occur in Respiratory, Neurological and Mental Health PBCs.

## 9 Conclusions

9.1 The methods of analysis presented here go some way to providing an empirically-based and explicit quantification of the scale of opportunity costs the NHS faces when considering whether the health benefits associated with new technologies are expected to offset the health that is likely to be forgone elsewhere in the NHS.
9.2 The study also starts to make the other NHS patients, who ultimately bear the opportunity costs of such decisions, less abstract and more 'known' in social decisions. Since who happens to be known or unknown is only a matter of perspective, time and ignorance, ethical and coherent social decisions require that both should be treated in the same way. These methods contribute to removing some of the 'ignorance' and making the unknown more real.
9.3 This work has implications for the Government's proposals to move to a system of value-based pricing for new prescription pharmaceuticals. These proposals include a widening of the measure of benefit to be used in evaluating new products including wider social benefits such as carer time and a consideration of the unmet need. Implementation of an augmented measure of benefit will also need to be reflected in terms of benefits forgone through service displacement. In principle, this will be feasible given the methods used in this study.


## Chapter 1: Introduction

### 1.1 Policy context

A comparison of the incremental cost effectiveness ratio (ICER) of a new technology with a costeffectiveness threshold is not the only consideration when the National Institute for Health and Clinical Excellence (NICE) and its advisory committees issues guidance. But it is an important one as it allows an assessment of whether the health expected to be gained from the use of a technology exceeds the health expected to be forgone elsewhere as other NHS activities are displaced. For this reason a comparison of the ICER of a technology to a threshold range is a critical part of the reference case in the NICE Guide to Methods of Appraisal[1] and is often taken to be the starting point for deliberations about other considerations including judgements of social value. Therefore, the value of the threshold is critical to the assessment of whether technologies can be regarded as cost-effective. This is also true for other NHS resource allocation decisions which potentially impose additional costs on local NHS commissioners.

From 2014 the Government plans to introduce a new approach to determining the appropriate price of prescription pharmaceuticals. Under value-based pricing, the price the NHS pays for a new product will be directly linked to its cost-effectiveness.[2,3] Therefore, the value of the threshold will be even more important as it will have a major impact on the prices that the NHS pays for pharmaceuticals, the access that NHS patients will have to new drugs and the return that manufacturers ean expect from future research and development.[4, 5]

### 1.2 Estimating the cost-effectiveness threshold

A key part of NICE's remit is to make decisions which are consistent with the efficient use of NHS resources. In the context of the NHS budget constraint, a consideration of efficiency has to reflect the implications of imposing additional costs on the system which will displace existing services thus leading to health decrements for patients other than those benefiting from the new technology being appraised. The cost-effectiveness threshold is an estimate of health forgone as other NHS activities are displaced to accommodate the additional costs of new technologies. A national decision-making body like NICE needs an estimate of what is likely to be forgone across the NHS as we currently find it.[6] Of course, this will change as circumstances and the NHS change; tending to rise with increases in budget and health care costs but tending to fall with increases in the productivity of health technologies and the efficiency of the NHS in general - including better local commissioning decisions.[7] A body like NICE cannot and does not necessarily need to know what specific services and treatments will be displaced in particular localities or who will actually forgo health.

What is required, therefore, is an accountable and empirically-based assessment of the health that is likely to be forgone on average across the NHS. Currently NICE uses a threshold range of $£ 20,000$ to $£ 30,000$ per quality adjusted life year (QALY) gained, where additional considerations are required towards the upper bound.[11 The empirical basis of this range of values is very limited and there have been calls for further research in this area.[8] Explicit scientific methods are required which will provide accountability so that estimates can be scrutinised by a range of stakeholders. Since estimates of the threshold will need to be periodically revised, methods which make best use of routinely available NHS data are needed. As wellas accountability, this will also provide more predictability in likely changes to the threshold for the investment decisions of technology manufacturers.

### 1.3 Aims and objectives

The aim of this research is to develop and to demonstrate methods to estimate the cost-effectiveness threshold for the NHS which makes best use of routinely available data. Methods are required which can capture the impact of a change in expenditure on length and quality of life ( QoL ), indicate how estimates of the threshold have changed over time, reflect uncertainty in any estimates and assess its implications, and indicate the impact of increases or decreases in spending. The project also aims to discuss options for developing data sources in the UK to estimate the threshold more precisely over time. The research has four main objectives:
i. Informed by relevant literature, to provide a conceptual framework to define the threshold and the basis of its estimation.
ii. Using programme budgeting data for the English NHS, to estimate the cost per life year gained on average across the NHS, for marginal changes in budget.
iii. To extend the measure of the health effects of changes in expenditure by estimating the QoL associated with additional years of life and the direct impact of health services on QoL.
iv. To synthesise this work to bring evidence on life-years and QALYs together, to present the best estimate of the cost-effectiveness threshold given existing data, to show the implications of the uncertainty in the current evidence and to provide recommendations for future data collection and analysis.

### 1.4 Report structure

The main report is set out as a series of chapters, most of which are linked to more detailed analysis in separate appendices. Chapter 2 provides a policy context for the research and a conceptual framework for the subsequent empirical work. Chapter 3 outlines a simple theoretical model and associated econometric analysis of programme budgeting data to estimate the link between changes in overall NHS expenditure and mortality. Chapter 4 considers a range of analyses to extend the measure of health effect from mortality to life-years gained and to QALYs. Chapter 5 draws out the main conclusions and insights from the research.

## Chapter 2: Policy Context and Conceptual Framework

### 2.1. Introduction

The purpose of this chapter is to provide the foundation for the empirical chapters that follow. It addresses a series of questions regarding the nature of the cost-effectiveness threshold that NICE use to guide its decisions, and the principles of how it should be estimated.

The chapter uses the results of a systematic literature search relating to these questions. Details of the methods and results of that search, together with a summary of the papers identified, are provided in Appendix A. In brief, the search uses a 'pearl growing' method to identify relevant papers. This identifies a number of initial key articles ('pearls') on the basis of expert advice, and 'grows' these pearls in a series of steps: extraction of citations and references from the initial pearls; identification of further pearls from cited and referenced papers; repetition of citation and reference searches; and manual search of references. This process is repeated until no further papers of relevance are identified. Onthis basis, 76 relevant papers were identified and are referred to, when relevant, in this chapter.

This chapter is organised as follows. The next section considers, at a conceptual level, what the costeffectiveness threshold to inform NHS decisions, such as those made by NICE's adyisory committees, should represent. Section 2.3 considers alternative routes to generating an empirical estimate of such a threshold. The final section provides a brief overview of the methods used in the study.

### 2.2. What should the NICE threshold represent?

### 2.2.1 The threshold as a measure of opportunity cost

NICE uses cost-effectiveness analysis (CEA) to inform the decisions underlying most types of guidance that it publishes. The use of CEA is most prominent in appraisals relating to new medicines, [1] but is also a key input into diagnostics appraisals as well as clinical guidelines and public health guidance.[1, 9] For those interventions and programmes which impose additional costs on the NHS budget, their incremental cost-effectiveness ratios (ICERs) indicate the incremental cost per additional quality-adjusted life-year (QALY) achieved relative to appropriate comparators. Although the ICER is one of a number of evidential inputs into NICE committees' decisions, is has been shown to be the most important, at least for technology appraisals.[10]

Interpreting whether a given ICER is acceptable requires the use of a cost-effectiveness threshold. Given that NICE has no influence on the level of the NHS budget, its decisions need to consider that budget a fixed constraint.[6] Therefore, the threshold should reflect the opportunity costs, in terms of health forgone, resulting from the imposition of additional costs on the NHS. When NICE issues positive guidance for a new intervention which imposes additional costs on the system, the resources required to deliver it must be found by disinvesting from other interventions and services elsewhere.[11] This displacement of existing services will result in health decrements for other types of individual.[12] Thus the threshold represents the additional cost that has to be imposed on the system to forgo 1 QALY worth of health through displacement.

As Figure 2.1 illustrates, CEA effectively becomes an analysis of net health benefits: does the health gain from the new intervention outweigh the health decrement associated with the displacement of existing services necessary to fund it? Figure 2.1 shows the incremental costs and QALYs associated with a new intervention relative to a comparator (the latter being shown at the origin). The new intervention generates 2 additional QALYs per patient and, at price P1, imposes an additional $£ 20,000$ per patient; the ICER is, therefore, $£ 10,000$ per QALY gained. At a threshold of $£ 20,000$ per QALY, the additional cost of $£ 20,000$ per patient translates into a decrement of 1 QALY (the distance between the $y$-axis and the threshold). Therefore, at that price, there is a net health gain of 1 QALY per patient (2 gained from the new intervention and 1 forgone through displacement). At a price of P 2 , the additional cost per patient of the new intervention is $£ 40,000$ and the net health gain is zero: the 2 additional QALYs from the new intervention are the same as the QALYs forgone through displacement. At the highest price of P3, the
adoption of the new intervention would actually result in a net health decrement of 1 QALY as it generates fewer QALYs (2) than are forgone (3).


Figure 2.1: graph showing illustration of the NICE threshold as a basis for assessing net health benefit. Adapted from Claxton et al[4].

The use of the threshold to facilitate this net health benefit (NHB) analysis can be expressed as in Equation 2.1:
$N H B=\Delta h-\frac{\Delta C_{h}}{k}$
Equation 2.1
where $\Delta h$ is the change in health generated by the new intervention, $\Delta C_{h}$ is the additional health care cost imposed on the NHS, and k is the cost effectiveness threshold. The net health gain from adopting the new intervention is, therefore, the health gained, $\Delta h$, minus the health forgone,$\frac{\Delta C_{h}}{k}$.

Understanding the NICE cost-effectiveness threshold as representing opportunity costs in terms of health is explicit in NICE documentation (for example, the methods guide for technology appraisal[1]). It is also clear in reports published by the Department of Health, such as the consultation report on value-based pricing. [8, 13, 14] This conceptualisation of the principles of the NICE threshold is also described in the broader literature. [6, 7] Formally, the threshold can be seen as the shadow price of the budget constraint. [6, 7, 12, 15-17] Although this project focussed on the use and estimation of a costeffectiveness threshold for NICE decisions, the methods and estimates relate to any resource allocation decision within the NHS where the opportunity cost could fall anywhere in the system. Hence it could apply, for example, to Department of Health targets or to Commissioning Board directives, as well as NICE guidance.

### 2.2.2 The threshold as the consumption value of health

Another view of what the threshold used in CEA should represent exists in the literature, however. In general terms, this is based on the rate at which individuals are willing to forgo other forms of consumption to achieve health improvement (sometimes referred to as 'willingness to pay').[18-37] Although this consumption value of health can provide information on the value of health improvement and may guide decisions such as the level of the overall NHS budget, it does not inform decisions regarding how to allocate a fixed budget within the health care system.

The reason for this is that the consumption value of health applies equally to health gained as well as to health forgone. This is shown in Equation 2.2 where the consumption value of health, v , is added to the definition of NHB in Equation 2.1. This simply involves valuing both health gained and health forgone by the same consumption value of a unit of health, $v$. Therefore, the use of the consumption value is irrelevant: a treatment considered cost effective in Equation 2.1 (i.e. to have a positive NHB) will inevitably be considered cost-effective in Equation 2.2, and an intervention with negative NHB (i.e. not cost effective) will remain as such in Equation 2.2. ${ }^{1}$ Therefore, the magnitude of the threshold, $k$, is not a value judgment but an empirical question which can, in principle, be estimated.
$N H B=v . \Delta h-\frac{v}{k} \Delta C_{h}$
Equation 2.2

### 2.3. Estimating the threshold

### 2.3.1 NICE's threshold range

NICE has been reluctant to specify a single cost effectiveness threshold used in its decision making.[10] It has also consistently emphasised that factors other than CEA are taken into consideration by the various advisory committees. [1, 9, 10, 38-40] Therefore, it has preferred to indicate the range within which its threshold value lies - $£ 20,000$ to $£ 30,000$ per QALY gained.[1, 9] Alongside this, it has provided an indication of the role other factors play in determining which point of threshold range is relevant. The latest guide[1] suggests that an ICER below $£ 20,000$ is likely to lead to recommendation unless the evidence is considered highly uncertain; an ICER between $£ 20,000$ and $£ 30,000$ will lead to recommendation if the committee is also happy with the levels of uncertainty in the evidence and/or the QALY does not capture all aspects of benefit; and an ICER above $£ 30,000$ would only be recommended if issues related to levels of evidential uncertainty and a failure to capture all benefits in the QALY are particularly compelling.

In the following year, NICE issued further supplementary guidance relating to the appraisal of interventions for patients with short life expectancy, although this can be considered to relate more to the measure of benefit than factors to be considered outside of cost effectiveness.[41] In 2012 NICE issued


[^1]a draft update of its methods guide which added that, if a new technology has an ICER above $£ 20,000$ per QALY, the committee's deliberations would also consider 'aspects that relate to non-health objectives of the NHS' (e.g. wider social considerations and/or costs that fall outside of the NHS budget).[42]

Although NICE has carefully argued the case for why its decisions are not driven entirely by a comparison of the ICER with its threshold range, it has not provided any empirical evidence for why the threshold range takes the value it does. Indeed it has been widely argued than an empirical basis for these values should be generated.[8, 43-47] For example, the House of Commons Health Select Committee in 2008 argued:
> "The affordability of NICE guidance and the threshold it uses to decide whether a treatment is cost-effective is of serious concern. The threshold is not based on empirical research and is not directly related to the budget, it seems to be bigher than the threshold used by PCTs for treatments not assessed by NICE. Some witnesses, including patient organisations and pharmaceutical companies, thought that NICE should be more generous in the cost per $Q A L Y$ threshold it uses, and should approve more products. On the other hand, some PCTs struggle to implement NICE guidance at the current threshold and other witnesses argued that a lower threshold should be used. We recommend that the threshold used by NICE in its full assessments be reviewed; further research comparing thresholds used by PCTs and those used by NICE should be undertaken...." ([8], page 6).

### 2.3.2 The basis for empirical work

Although there is acceptance of the need for empirical work on the NICE cost-effectiveness threshold, a set of issues exists regarding the starting point for such analysis. One aspect of this is the view that the nature of the services that are displaced in response to additional costs being imposed by NICE guidance, and hence the magnitude of the health forgone for other patients, will depend on the productivity of the NHS and its overall (inflation adjusted) budget, both of which have increased since NICE initially defined its threshold range. [48, 49] In principle an increase in the (real) NHS budget would allow it to introduce interventions which were previously not cost effective which might be expected to increase the threshold if these interventions were the marginal ones displaced in response to the budget impacts of NICE recommendations. However, any increase in the NHS budget may be allocated to non-discretionary expenditure. This would include, for example, expenditure relating to national initiatives such as new contracts for consultants and activities to meet waiting list targets as well as, of course, the implementation of NICE guidance. The non-discretionary nature of such expenditure means that these types of activities cannot easily be disinvested from given a need to release resources to fund NICE guidance. Therefore, if an increase in the NHS budget is largely devoted to these types of nondiscretionary expenditure, there will be a limited impact on the threshold.

Gains in productivity may come through doing worthwhile activities more cost effectively, including for those marginal interventions displaced by NICE recommendations, suggesting a reduction in the threshold. Altematively, productivity gains might come through discontinuing activities which are not worth doing (i.e. that produce no health improvement), freeing resources for additional cost effective interventions which may be the marginal services displaced by NICE guidance - this can have the result of increasing the threshold.

The net effect of these changes on the threshold could not be determined a priori and would depend how any additional (real) budget were allocated and how the gains in productivity where achieved. This does emphasise the fact that the threshold may change over time in response to these and other broader developments, and this would have to be considered as part of any regular updating of the empirical analysis of the threshold.

A second issue to be considered relates to how decisions are taken locally about any displacement following NICE guidance. The principles of CEA suggest that such displacement should relate to interventions which are the least cost effective of those currently covered by the budget.[15] The basis for how local commissioners and providers make their disinvestment decisions is not clear, however, and there have been calls for greater transparency and guidance in this area.[48] It would be entirely unrealistic to assume that displacement only takes place in those existing services which are the least cost
effective. The reality is that numerous criteria are likely to be used by commissioners in implementing disinvestment, and that significant variation will exist between local decision makers.[12] Therefore, NICE needs to know what is likely to happen on average across the NHS given the reality of local decisions. If local decision making changes over time, this may affect the estimate of the threshold.

### 2.3.3 Studying displacement locally

A reasonable conclusion from a consideration of these issues is, therefore, that local decisions about disinvestment are likely to be an important determinant of the NICE threshold.[50-55] Appleby et al sought to assess whether it was possible to study local decisions about service investment and disinvestment to infer the cost effectiveness thresholds being used (implicitly) locally and to draw conclusions about the appropriate level of the NICE threshold.[56] They identified six primary care trusts (PCTs) and undertook structured interviews with each of the directors of public health. They also administrated questionnaires to an opportunistic sample of finance directors from NHS trusts On this basis they developed a list of new services as well as those that had been deferred or discontinued. An attempt was made to estimate the implicit local ICER relating to these decisions by using any cost effectiveness evidence used to inform the decisions together with relevant evidence on cost effectiveness from the published literature.

The study found it quite straightforward to identify specific services that had been introduced, discontinued or deferred, but concluded that these decisions were typically based on clinical and other non-economic factors. A number of 'decisions at the margin' were identified but none of these was based on cost effectiveness analysis. Instead, the basis for changes in services was a 'business case', or overall cost impact. It was possible to impute cost effectiveness formost of the services affected, but the study concluded that, even with a larger sample of commissioners and providers, it would be very difficult to estimate an implied cost effectiveness threshold locally. This would be because, firstly, most PCT decisions were service reconfigurations including demand management and waiting list initiatives. By their nature, teasing out the incremental cost and health effects, potentially across numerous types of patients, would be an enormous challenge. Secondly, there would be difficulty in identifying all local decisions as many options for investment, deferment or discontinuation are rejected before they are made more explicit in documentation. A third problem would be the finding that a range of criteria is used to make local decisions, with relatively little concern for cost effectiveness, making a local threshold estimated in this way hard to interpret. A final challenge would be that it would be very difficult to establish a causal link between a change in local NHS budget and specific local investment and disinvestment decisions. The Appleby et al study highlights the problems that exist in deriving a cost effectiveness threshold from a bespoke study of specific local resource allocation decisions.

### 2.3.4 What evidence is needed?

Given the challenges of studying local decisions as a means of establishing the NICE threshold, a series of important chatacteristics that estimation methods should have from a practical and principled perspective would seem appropriate:

They should reflect the effect of NICE guidance on the average of the displacement decisions taken across the NHS, with less consideration on which types of patients and interventions are affected and why the decision are taken. NICE cannot be expected to reflect what is likely to be marked variation between local commissioners and providers in how they react to an effective reduction in their budget as a result of positive guidance. Given NICE's remit, it is the expected health effects (in terms of length and quality of life) of the average displacement within the current NHS (given existing budgets, productivity and the quality of local decisions) that is relevant to the estimate of the threshold.

- The methods used should not be a 'once and for all' effort but should facilitate regular updates to reflect changes in the broader NHS context such as changes in the overall real budget and productivity. This requires the use of data sources that are currently routinely available, are
expected to become so in the future or could be made available at reasonable cost. It may be possible to glean some idea of how the threshold may change in the future by studying how it has changed in the past, which would require routine data sources to extend back over a period of time. Periodic updating using explicit scientific methods would encourage accountability through scrutiny of estimates by relevant stakeholders. It would also provide more predictability in likely changes to the threshold for the investment decisions of technology manufacturers.
- The nature of the displacement of existing services (and hence the magnitude of the health forgone) will depend on the scale of the budget impact coming through NICE guidance. Therefore, the methods used to estimate the threshold should ideally be able to reflect this budget impact.
- The methods should recognise the inevitable uncertainty relating to the evidence currently available for threshold estimation and translate this into an expression of the uncertainty in the estimate of the threshold. As well as providing information with which NICE can-determine the appropriate implications for its choice of a threshold value, this consideration of uncertainty can help to prioritise further research or the collection of routine data.


### 2.4. An introduction to study methods

The current study has sought to develop methods consistent with these desired characteristics. This section provides a summary of the methods used. Further details are provided in each of the later chapters relating to the various components of work, and in the associated appendices. The general approach taken is to use routinely available data to look at the relationship between overall NHS expenditure and patients' health outcomes. By exploiting differences between PCTs in expenditure and outcomes, it is possible to infer the costs of generating health improvement from NHS services at the margin. In principle, this is what is needed as the basis of the NICE cost effectiveness threshold as it provides an indication of the health forgone through the services displaced by the additional budget effect of the Institute's guidance.

### 2.4.1 Past work

The study was able to build on some key existing research relating to the relationship between NHS expenditure and mortality.[57-59] Since 2003 data on expenditure on health care across 23 programme budgeting categories (PBCs) of care have been available for each PCT in the NHS in England. These programme budgeting data seek to allocate, to broad areas of illness according to the primary diagnosis (using ICD10 codes) all items of NHS expenditure, including expenditure on inpatient care, outpatient care, community care, primary care and pharmaceuticals and devices.

For the purposes of this study, the merit of these data is that they open up the possibility of examining the relationship between differences in local spending and associated disease-specific mortality outcomes routinely avaitable from the National Centre for Health Outcomes Development. In each programme, the elasticity ofoutcome with respect to changes in expenditure was estimated controlling for differences between PCTs in need. Changes in mortality were then transformed into life-years gained using assumptions regarding life expectancy without the change in expenditure. This provides estimates of the marginal cost per life-year gained on average across the NHS by PBC.

This work focused largely on spending and outcomes in two of the largest programmes: circulatory disease and cancer, [60] but has also informed the link across other programme categories. [58, 61] Estimates of the cost per life year gained for 2006/07 were $£ 15,387$ for cancer, $£ 9,974$ for circulation problems, $£ 5,425$ for respiratory problems, $£ 21,538$ for gastro-intestinal problems and $£ 26,428$ for diabetes. These estimates were based on a straightforward, though carefully constructed, theoretical model of health production which informs the specification and estimation of a set of equations. These dealt with the challenge of there being alternative plausible directions of causation - for example, between expenditure and health outcomes within a programme. This problem of endogeneity was addressed by identifying and testing suitable instrumental variables. In doing so, they accounted for variation in the
clinical needs of the local population relevant to each programme together with broader local environmental factors relevant to the costs of care and outcomes.

This earlier work provides a strong foundation for the current study through its consideration of the average marginal elasticity of outcome with respect to programme expenditure. However, to estimate the threshold suitable for NICE decision making, a number of further elements of research are necessary, and these are described below.

### 2.4.2 Further econometric analysis

This further econometric research is covered in Chapter 3, with full details in Appendix B. The earlier work estimated the cost per life-year gained for the major programme areas. The NICE threshold needs to relate to the whole NHS and will, therefore, depend on all the programmes of care where disinvestment takes place. Given that each programme of care has been estimated separately, it is not clear how expenditure on particular programmes changes with the overall budget. For example, does disinvestment tend to fall on respiratory care or diabetes following a budget impact from NICE guidance? Therefore, the current study has further developed the econometric analysis to reflect the need for PCTs to operate within a fixed overall budget. This provides an estimate of the 'budget elasticity of expenditure' in each PBC, and facilities estimates of the impact of marginal increases (or decreases) in overall PCT budgets on spending in each PBC.

As well as indicating budgetary influences on programme spending these have then been linked to changes in mortality outcomes by programme. These changes are used to estimate years of life lost taking account of the fact that some of the observed deaths would have occurred anyway (had the same population not been at risk in the particular PBC); that is, taking account of unobserved counterfactual deaths. This takes into account how such budgetary changes (such as those imposed by NICE guidance) translate through local decisions into changes in expenditure on programmes of care and then to health outcomes.

Changes in budgets are in practice incremental rather than marginal, and it may be the case that the outcome elasticities of programme expenditure in times of budgetary increase (when new initiatives are introduced) are not the same as in times ofbudgetary decrease (when the focus is on disinvestment). The possible effects of non-marginal changes have, therefore, been explored. The project has also sought to explore how both expenditure and outcome elasticities, and hence the threshold, vary over time, and this has been assessed by generating relevant estimates for three sets of data.

A development from earlier work has been to relate expenditure in period $t$ to mortality in periods $t, t+1$ and $t+2$. Whilst the data used are largely cross-sectional, mortality data are linked so as to follow expenditures. Given the inevitable uncertainty relating to assumptions in the analysis, extensive sensitivity analysis is undertaken to consider the implications for the estimates.

### 2.4.3 Moving from life-years to quality-adjusted life-years gained

A key element of the research has been to take the results of the econometric work linking NHS spending and mortality, and to translate this into effects on life years and quality adjusted life years (QALYs). This is achieyed using three sequential steps:
i. Translate the estimated effects on mortality from the econometrics work into life years by exploring the limitations of the mortality data available at PCT level and the published years of life lost (YLL) figures used in the econometric analysis, and by considering how to improve the estimates using additional data and analysis.
ii. Consider how estimates of life year effects can be adjusted for the quality of life in which they are lived, taking account of the gender and the age at which life years are gained or lost as well as the quality of life implications of particular diseases.
iii. Explore ways to take account of those effects on health not directly associated with mortality and life year affects (i.e., the 'pure' quality of life effects) to estimate an overall cost per QALY threshold.

This aspect of the analysis is described in Chapter 4 with further details provided in Appendix C.
The central or 'best' estimate is based on two assumptions relating to the health effects associated with expenditure, one conservative and the other more optimistic. The first assumption is that the health effects of changes in one year of expenditure are restricted to one year. This is implicit in the estimates of outcome elasticities estimated in the econometric analysis. This is likely to underestimate effects on mortality since expenditure that reduces mortality risk for an individual in one year may well also reduce their risk over subsequent years, and expenditure may also prevent disease in future patient populations. Therefore, total health effects will be underestimated and the cost per life year or QALY threshold will be overestimated. Although undoubtedly conservative, it may be offset to some extent by the more optimistic assumption. It is assumed that any death averted by expenditure in one year will return the individual to the mortality risk of the general population, i.e., the years of life gained associated with each death averted are based on what would have been their life expectancy taking account of their of age and gender (using life tables for the general population).

The extreme upper and lower bounds for cost per life year and cost per QALY thresholds are based on making both of these assumptions either optimistic (providing the lower bound for the threshold) or conservative (an upper bound for the threshold). The lower bound is based on assuming that health effects are not restricted to one year but apply to the remaining disease duration for the population at risk during the expenditure year. The upper bound is based on the combination of assuming that health effects are restricted to one year and that any death averted is only averted for the minimum duration consistent with the mortality data used to estimate the outcome. It is very important to note that the lower and upper bounds are very much extreme values with limited plausibility.

### 2.5. Conclusions

A cost effectiveness threshold is needed to inform decisions by NICE, the NHS more generally or the Department of Health which reflects the fact that opportunity costs fall on services and population health at a local level. Given that it is (and will continue to be) unfeasible to know precisely which services are displaced across all localities within the NHS, the threshold should reflect the average implications on health of actual local decisions about marginal changes in local service caused by changes in expenditure. The absence of an empirical estimate of the threshold which reflects these principles lies behind the project. Using data routinely collected in the NHS or available data that could be routinely updated, the study is organised into two major parts. The first updates earlier analysis to estimate the relationship between NHS expenditure and mortality, and the second seeks to translate these mortality effects into the more general measure of health - the QALY.

# Chapter 3: The link between NHS spending, mortality and the cost of a life year 

### 3.1 Introduction

This section presents an overview of the econometric work undertaken to estimate the link between NHS spending and mortality and how this is used to calculate the cost of a life year. As well as providing the analytical foundations for estimates of cost per QALY threshold presented in Chapter 4 and 5 this work also contributes to the on-going debate about the extent to which additional health care expenditure yields improved patient health outcomes.

The work presented in this report takes advantage of the availability of two new datasets to examine the relationship between National Health Service (NHS) expenditure and mortality rates for various disease categories. One dataset contains mortality rates for various disease categories at the level of geographically defined local health authorities, Primary Care Trusts (PCTs). The other dataset presents NHS expenditure by PCT on 23 broad programmes of care. This dataset embraces nhost items of publicly funded expenditure, including inpatient, outpatient and community care, and pharmaceutical prescriptions. NHS revenue derives almost entirely from national taxation, and access to the system is generally free to the patient. The system is organized geographically, with responsibility for the local administration of the NHS devolved to PCTs. ${ }^{2}$ PCTs are allocated fixed annual budgets by the Department of Health, within which they are expected to manage the health care in the locality.

We employ a model that assumes that each PCT receives an annual financial lump sum budget and allocates its resources across the 23 programmes of care to maximize the health benefits associated with that expenditure. Estimation of this model using the expenditure and mortality data facilitates two related studies: first, a study of how changes in the NHS budget impact on expenditure in each care programme; and second, a study of the link between expenditure in a programme and the health outcomes achieved, notably in the form of disease specific mortality rates. The latter also permits the calculation of the cost of an additional life year for individual programmes of expenditure.

The work presented here draws heavily upon on previous studies using these data $[57,59,60,62,63]$ and innovates in four major ways: (1) we relate expenditure in time period $t$ to outcomes in periods $t, t+1$, and $t+2^{3}$; (2) we present plausible outcome models for a large number of budgeting categories - previous studies have tended to focus on the four largest care programmes; (3) we present estimates of the cost of a life year for the enlarged number of programmes and, importantly, with the aid of assumptions about the productivity of programmes without a meaningful mortality-based outcome indicator, we extend our individual programme estimates to incorporate expenditure across all programmes of care; and (4) while the models we present appear well specified according to appropriate statistical tests, we subject our results to a substantial sensitivity analysis.

The next section presents a brief review of the relevant literature upon which the study builds. This is followed by a summary overview of our approach to estimating the cost per life year across the various prograrmmes of care and the results obtained using Programme Budgeting data provided by the Department of Health. Further details of all aspects of the modelling approach, description of the data, the results we derive and calculation of costs per life year are set out in Appendix B. This section is intended to be supported by the information contained within Appendix B.

[^2]
### 3.2. Previous studies

One of the most fundamental yet unresolved issues in health policy is the extent to which additional health care expenditure yields patient benefits, in the form of improved health outcomes. The work of health technology agencies such as the English National Institute for Health and Clinical Excellence (NICE) has greatly improved our understanding at the micro-level of the costs and benefits of individual technologies. However, there remains a dearth of reliable evidence at the macro-level on the benefits of increased health system expenditure.

The empirical problems of estimating the link between spending and health outcomes are manifest. If one relies on a time series of health outcome data for an individual health system it is difficult to disentangle the impact of expenditure from a wide range of other temporal influences on health, such as technological advances, epidemiological changes, and variations in broader economic circumstances. Similar methodological difficulties arise if one attempts a cross-sectional comparison of different health systems. In particular, when seeking to draw inferences from international comparisons, researchers might have failed to adjust for all the potential external influences on health outcomes and this might account in part for their findings. For example, in an early cross-sectional study of 18 developed countries, Cochrane et al.[64] use regression analysis to examine the statistical relationship between mortality rates on the one hand and per capita GNP and per capita consumption ofinputs such as health care provision on the other. They found that the indicators of health care provision were generally not associated with outcomes in the form of mortality rates. Thereafter, the failure to identify strong and consistent relationships between health care expenditure and healthoutcomes (after controlling for other factors) has become a consistent theme in the literature, whilst, in contrast, socioeconomic factors are often found to be good determinants of health outcomes.[65-67]

There is furthermore the possibility that indicators of heatth system inputs, such as expenditure, are endogenous, in the sense that they have to some extent been influenced by the levels of health outcome achieved. And the difficulty of satisfactorily estimating the impact of health system inputs on outcomes is compounded by the great heterogeneity of health care, the multiple influences on outcomes, and the rather general nature of the outcome mortality neasure traditionally used. Consequently, the failure to detect a significant positive relationship between expenditure and health outcome might reflect the difficulties associated with any such study rather than the absence of such a relationship. For example, Gravelle and Backhouse[68] examine some of the methodological difficulties associated with empirical investigation of the determinants of mortality rates. These include simultaneous equation bias and the associated endogeneity problem (that the level of health care input might reflect the level of health outcome achieved in the past), and that a lag may occur between expenditure and outcomes (studies typically assume that expenditure has an immediate effect on mortality).

To avoid the difficulties imposed by data heterogeneity inherent in international analyses, the study by Cremieux et al.[69] examines the relationship between expenditure and outcomes across ten Canadian provinces over the fifteen-year period 1978-1992. They find that lower healthcare spending is associated with a significant increase in infant mortality and a decrease in life expectancy. Although challenging the received empirical wisdom, one difficulty with the Cremieux et al.[69] study is that the estimated regression equation consists of a mixture of potentially endogenous variables (such as the number of physicians, health spending, alcohol and tobacco consumption, expenditure on meat and fat) and exogenous variables (such as income and population density). The authors' chosen estimation technique (GIS) does not allow for this endogeneity and consequently the coefficients on the endogenous variables may be biased.[68] Similarly, Nixon and Ulmann's[70] study, which uses three health outcome measures and various explanatory variables (such as per capita health expenditure) for 15 EU countries over the period 1980-1995, does not allow for the possibility that some of the explanatory variables may be endogenous.

More recently, studies have started to address the endogeneity issue.[71, 72] Bokhari, Gai and Gottret[71] estimate a cross-section model for 127 countries using data for 2000. They employ two health outcome indicators (the under-five mortality rate and the maternal mortality rate). Bokhari et al.
allow for the endogeneity of health expenditure via the use of instrumental variable techniques, and they estimate the elasticity of these indicators with respect to total government health expenditure conditional on the level of education and basic infrastructure (such as road transport and sanitation). They find that health expenditure has a statistically significant negative impact on both under-five mortality and maternal mortality. The authors do note, however, that their focus on child and maternal mortality implicitly assumes that these outcome indicators are in some way representative of outcomes across all activities financed by government health care expenditure. Data permitting, it would be preferable to relate health care expenditure on those aged under five years to under-five mortality, and expenditure on maternal care to maternal mortality.

In this study we relate expenditure in a specific disease area to mortality associated with those diseases. We also address the endogeneity issue through the use of instrumental variables and, unlike previous studies; we examine the sensitivity of our results to questions of instrument validity. Moreover, although previous empirical work has been loosely based on the notion of a health production function, it has rarely been informed by an explicit theoretical model. This is understandable, as the processes giving rise to the observed health outcome are likely to be very complex, and any theoretical model might become rather unwieldy. However, this absence of a theoretical model has sometimes led to an atheoretical search for measures of health inputs demonstrating a statistically 'significant' association with health outcomes. In contrast, in this study we inform our empirical modelling with a theoretical framework. We believe that this may lead to a more convincing and better specified model of health outcomes than that used in many previous studies, and this model is outlined in the next section.

### 3.3. Modelling framework

In the literature on the relationship between health expenditure and health outcomes, the statistical model estimated often contains a mixture of exogenous variables (such as income and population density) and endogenous variables (such as health spending, the number of doctors, and spending on cigarettes and alcohol). In such circumstances, the application of ordinary least squares will lead to biased coefficients on the endogenous variables. To avoid this problem, Gravelle and Backhouse [68] recommend that analysts model, even if only informally, the decision making process which generates the observed data set.

To avoid the problem of simultaneous equation bias we have constructed a very basic model of the budgeting and outcomes data generation processes. In places, the model makes some heroic assumptions (which we hope to relax in future work) but the framework reveals some of the more salient features of the data generation processes.

We assume - quite realistically - that each PCT, $i$, receives an annual financial lump sum allocation, $y_{i}$, from the Department of Health and that total within year expenditure for each PCT cannot exceed this amount. We also assume - less realistically - that this lump sum is allocated across the $J$ programmes of care $(J=23)$ by a single decision maker (although we know that in practice the programme budget data will in part reflect the myriad of individual clinical decisions that health care professionals take every day and that theseare decisions over which PCTs exercise little control).

We assume that each PCT adheres to a social welfare function, $W($.$) , that incorporates the health$ outcome ( $h$ ) across all 23 programmes of care so that for each PCT

$$
\begin{equation*}
\mathrm{W}=\mathrm{W}\left(h_{1}, h_{2}, \ldots, h_{J}\right) \tag{Equation 3.1}
\end{equation*}
$$

Health outcomes might be measured in a variety of ways, but the most obvious is to consider some measure of improvement in life expectancy, possibly adjusted for quality of life, in the form of a quality adjusted life year.

We assume that, for each PCT and for each programme of care, there is a 'health production function'
that indicates the link between local spending on programme $j\left(x_{j}\right)$ and health outcomes in the same programme $\left(h_{j}\right)$. Two such production functions are illustrated in Figure 3.1. We assume that increased expenditure yields improvements in health outcomes, as expressed, for example, in local mortality rates, but at a diminishing rate. Clearly the shape of the curve might depend on the health needs of the local population (such as epidemiological conditions) and other local circumstances, such as socio-economic conditions and local service input prices. Note that in Figure 3.1 the cost of securing a given level of health outcome is - for whatever reason - higher in $\mathrm{PCT}_{\mathrm{a}}$ than $\mathrm{PCT}_{\mathrm{b}}$.


Figure 3.1: The health production function for programme $j$ in two PCTs
In algebraic form, each PCT seeks to maximise total welfare across all $J$ programmes of care $(J=23)$ subject to the health production function for each programme of care of the form

$$
h_{j}=f_{j}\left(x_{j}, n_{j}, z_{j}\right)
$$

Equation 3.2
where $n_{j}$ is the need for health care in programme $\mathrm{j}, x_{j}$ is PCT expenditure on programme j , and $z_{j}$ represents environmental variables affecting the production of health outcomes in programme $j$ (which might include private (non-PCT) health care expenditure in the disease area). Each PCT's problem is to select an expenditure level for each programme $\left(x_{j}^{*}\right)$, so as to maximise the utility function in equation (3.1) subject to the health production functions in Equation 3.2 and the budget constraint that total expenditure on all programmes should not exceed PCT income (y).

Algebraically, the budget constraint is:

$$
x_{1}+x_{2}+\ldots+x_{23} \leq y
$$

## Equation 3.3

Solving this maximisation problem yields the result that the optimal level of PCT expenditure in each category, $x_{j}^{*}$, is a function of the need for health care in each category $\left(n_{1}, n_{2}, \ldots, n_{23}\right)$, environmental variables affecting the production of health outcomes in each category $\left(z_{1}, z_{2}, \ldots, z_{23}\right)$, and PCT income (y). Thus:

$$
\begin{aligned}
& x_{1}^{*}=x_{1}\left(n_{1}, n_{2} \ldots, n_{23}, z_{1}, z_{2} \ldots, z_{23}, y\right) \\
& x_{2}^{*}=x_{2}\left(n_{1}, n_{2} \ldots, n_{23}, z_{1}, z_{2} \ldots, z_{23}, y\right) \\
& \ldots \\
& x_{23}^{*}=x_{23}\left(n_{1}, n_{2} \ldots, n_{23}, z_{1}, z_{2} \ldots, z_{23}, y\right)
\end{aligned}
$$

$$
\text { Equation } 3.4
$$

These results imply that each PCT will allocate expenditure across the 23 programmes of care so that the marginal utility of the last pound spent in each programme of care is the same. Of course, this does not mean that each programme receives the same amount of cash; financial allocations will depend on both the relationship between utility and outcomes, and on the relationship between outcomes and expenditure for each programme of care. If we assume that one extra unit of health outcome improves managerial utility by the same amount irrespective of the programme of care, then the decision maker simply allocates expenditure across all programmes to maximise total health outcomes. This is achieved by ensuring that the marginal health outcome benefit (measured perhaps in QALYs) is the same for the last pound spent across all programmes of care.

Thus, for each programme of care, there exists an expenditure equation (equation 3.4) explaining the expenditure choice of PCTs and a health outcome equation (equation 3.2) which models the associated health outcomes achieved. As presented, our basic model is static in the sense that the health production function (equation 3.2) assumes that all health benefits occur contemporaneously with expenditure. We acknowledge that for some programmes of care benefits might occur one or more years after expenditure has occurred. This is particularly likely to be the case for those programmes aimed at encouraging healthy lifestyles, where some benefits may occur decades after the actual programme expendifure. For other programmes, such as maternity/reproductive conditions and neonate conditions, benefits may be largely contemporaneous with expenditure. However, while our data are largely cross-seetional in nature, we are able to link mortality data in such a way that this follows expenditures. Accordingly, for our empirical modelling we estimate models using expenditure for period $t$ with mortalify data for periods $t, t+1$, and $t+2$. Appendix B presents a number of sensitivity checks on these assumptions including models where mortality data precedes expenditure data ${ }^{4}$ and shows that these results are fairly consistent with the results presented here.

### 3.4 Data

### 3.4.1 Programme budgeting in England

Prior to October 2006, there were 303 PCTs in England with an average population of about 160,000 people. In October 2006 the 303 PCTs became 152 PCTs. Some PCT boundaries remained unchanged while other PCTs were merged with one or more neighbours to form a new, larger, PCT. In a few cases the geographic area covered by an existing PCT was split between two or more new PCTs. These 152 PCTs have an average population of about 330,000 people. PCTs are allocated fixed annual budgets within which they are expected to meet expenditure on most aspects of health care, including inpatient, outpatient and community care, primary care and pharmaceutical prescriptions.

Programme budgeting data collection was initiated by the Department of Health in April 2003 when each PCT was required to prepare expenditure data disaggregated according to 23 programmes of health care. These programmes are defined by reference to the International Classification of Diseases (ICD) Version 10 codes at the four digit level, and most programme budget categories reflect ICD 10 chapter headings (e.g., cancer and tumours, circulation problems, renal problems, neonates, problems associated with the skin, problems associated with vision, problems associated with hearing, etc). In some cases, the 23 categories ate broken down into further sub-areas to achieve a closer match with the various National Service Frameworks (NSFs): for example, the large mental health category is broken down into 'substance abuse', dementia', and 'other'.

Programme budgeting seeks to allocate all types of PCT expenditure to the various programme budget categories, including secondary care, community care and prescribing. However, the system acknowledges that a medical model of care may not always be appropriate, and two specific non-clinical groups --

[^3]'Healthy Individuals' and 'Social Care Needs' -- have been created. These are intended to capture the costs of disease prevention programmes and the costs of services that support individuals with social rather than health care needs. In addition, in some cases it is not possible to assign activity by medical condition, preventative activity, or social care need and, in these cases, expenditure is assigned to a residual category (PBC 23) entitled 'Other'. The most important element of this residual programme is expenditure on general practitioner services ( PBC 23 a ). In principle, it should be possible to allocate each GP consultation to a particular care programme. However, at the moment the available data information systems do not permit such an allocation and so all primary care expenditure is allocated to this residual programme. The use of this residual category ensures that all expenditure is assigned to a programme of care.[73]

The aim of the programme budget classifications is to identify the entire volume of health care resources assigned to broad areas of illness according to the primary diagnosis associated with an intervention. It serves a number of purposes, most notably to assist in the local planning of health care. But for this study its crucial merit is that it opens up the possibility of examining the statistical relationship between local programme spending and the associated disease-specific outcome. Various forms of data collection and analysis are required to map PCT expenditure on acute, community and other services to the 23 programme budget categories. From the PCT perspective, however, the construction of each PCT's return largely involves collating information provided by other bodies and drawing on other information already in the PCT's own annual accounts. Details of how expenditure is assigned to programmes of care can be found in Section B4.2 of Appendix B.

Table 3.1 shows the expenditure per head and the growth in this expenditure for each programme budget category for 2003/04 to 2008/09.5 Year on year comparisons of expenditure in each group are complicated by the fact that the algorithms used to allocate activity to PBCs are regularly revised. 6 However, by 2008/9 total PCT expenditure per person had increased to $£ 1,531$ (up $28 \%$ from 2004/5). The residual 'other' category (programme budget category 23) still accounted for the largest share of expenditure ( $14.9 \%$ ) with per capita expenditure of almost $£ 228$, of which $£ 145$ was accounted for by primary care expenditure. Mental health (budget category 5) accounted for just over $12 \%$ of expenditure, but the expenditure share recorded by circulation problems (budget category 10) had fallen from 10.2\% to $8.5 \%$. Other categories recording a fall in budget share of more than one half of one percentage point included: the gastro-intestinal system (down from $6.1 \%$ to $5.1 \%$ ), the musculo-skeletal system (down from $6.0 \%$ to $5.2 \%$ ), trauma and injuries (down from $6.0 \%$ to $4.2 \%$ ), and maternity (down from $4.6 \%$ to $3.9 \%$ ). Categories recording an increase in budget share of more than one half of one percentage point included neurological problems (up from $2.9 \%$ to $4.4 \%$ ) and dental problems (up from $1.1 \%$ to $4.1 \%$ ).

Some of these changes will partly reflect revisions to the algorithms used to allocate expenditure to particular PBCs. For example in 2006/7 expenditure per person on musculo-skeletal problems fell by $11 \%$ and expenditure on trauma and injuries fell by $25 \%$. In the same year, expenditure on neurological problems increased by $35 \%$. This suggests that some types of activity, which were previously allocated to musculo-skeletalproblems and/or trauma and injuries, were re-allocated to neurological problems.

Similarly, up to and including 2006/7 expenditure that was not directly attributable to a particular programme category was apportioned using admitted patient care percentages. ${ }^{7}$ In other words, if $x^{\%} \%$ of totaladmitted patient care expenditure was allocated to PBC 1, then $x \%$ of all expenditure that was not directly attributable to a particular programme category was also allocated to PBC 1. With effect from 2007 /8, however, NHS organisations were asked to select an appropriate basis for the apportionment of this non-programme specific expenditure and that, where no reasonable basis existed, such expenditure was to be allocated to the 'Other - Miscellaneous' (PBC 23X) category. These two changes to the

[^4]algorithm used to allocate expenditure to particular PBCs illustrate that year-on-year comparisons of expenditure need to be interpreted with care.

Expenditure per head on any given programme varies from one PCT to another and Table 3.2 presents some statistics that indicate the degree of variation in expenditure levels across PCTs by programme budget category. The first four columns of Table 3.2 present descriptive statistics for PCT expenditure per person. These reveal that, for example, PCT per capita expenditure in the cancer programme averaged $£_{9} 96.30$ across all PCTs, with the minimum spend being $£ 62.90$ and the maximum being £155.70.

Some PCTs will be spending more than other PCTs simply because they face higher input costs. The second set of four columns in Table 3.2 present descriptive statistics for PCT per capita expenditure that has been adjusted for the unavoidable geographical variation in costs (input prices) faced by PCTs. ${ }^{8}$ However, if anything this adjustment appears to increase the variation in expenditure across PGTs, for example, the range of per capita expenditure on cancer increases from between $£ 62.90$ and $f(155), 70$ (unadjusted) to between $£ 59.10$ and $£ 163.10$ (adjusted for local health care input prices).

Another cause of the variation in expenditure levels is the fact that the need for health care varies from one PCT to another. For example, areas with a relatively large proportion of eldenfy residents, or PCTs operating in relatively deprived locations, can be expected to experience relatively high levels of spending. The Department of Health has a well-developed methodology for estimating the relative health care needs, which it uses as the basis for allocating health care funds to.[74]

The final set of four columns in Table 3.2 present descriptive statistics for PCT per capita expenditure that has been adjusted for both the unavoidable geographicat variation in costs and the local need for health care faced by PCTs. ${ }^{\text {P }}$ For virtually every PBC, this adjustment reduces the variation in expenditure across PCTs; for example, the standard deviation of PCT per capita expenditure falls from £. 19.70 to $£ 15.30$ for the cancer programme. Although this adjustment reduces the variation in expenditure levels across PCTs, this decline is quite modest and there are still substantial differences in expenditure even after allowing for differences in local cost and need. For example, expenditure per head in the circulation problems category varies between $£ 78$ and $\times 328$ using cost adjusted expenditure data, but falls between $£ 76$ and $£ 327$ using cost and need adjusted population data.

The variation in expenditure across $\mathrm{PCT}_{s}$ has led some commentators to question the reliability of the programme budgeting data. The (National Audit Office[75] undertook a survey of Trusts, PCTs and SHAs to assess the quality of the data. They concluded that while the processes for collecting the budgeting data were well defined in most areas, there remained scope for improvements to the robustness of some of the data (e.g.non-admitted patient care). Appleby et al.[76] also considered the issue of data reliability in variations in spending on cancer services and noted some large year-on-year changes. However, the authors point out that it is difficult to define what might be either an implausible level of expenditure or an implausibly large change in expenditure. This is complicated by the fact that the Department of fealth makes regular improvements to the way in which activity is matched to programme categories.

As with most datasets, there are likely to be recording and other errors associated with the programme budgeting data. However, while we note that the allocation of programme budgeting data might not be perfect there is no systematic evidence of this. Accordingly, for each disease category, we observe that PCT expenditure per person varies considerably and this variation - holding constant input prices and the

[^5]need for health care - offers the opportunity to examine whether PCTs that spend more on health care achieve a better outcome and, if so, at what cost. Empirical estimates of the strength of this relationship for several programmes of care are presented in this report.

### 3.4.2 Health outcome data

Most studies of the relationship between expenditure and outcome have used some measure of mortality as an indicator of the latter. We also employ mortality as an outcome measure. First, it is a relevant (albeit not comprehensive) measure of the outcome of health care expenditure; and second, it is available for more disease areas than any other outcome measure at PCT level.

Although mortality is available (by PCT) for several disease areas, it is not available for just over one-half of all programmes not least because it is simply not relevant for these programmes (e.g., for learning disabilities, vision problems, hearing problems, dental problems, and skin problems). Moreover, even where a mortality measure is available, the ICD10 coverage of the mortality data often falls short of the coverage of the expenditure data. For some programmes, therefore, we have combined the published mortality rates for two or more disease areas in an attempt to match the ICD10 coverage of the mortality data with that of the expenditure data.

Table B5.1 (Appendix B) shows how we have attempted to marry the mortality data (column c) and the expenditure data (column a). ICD10 coverage of the component mortality rates for some PBCs falls short of the expenditure data and the extent of this shortfall is illustrated by the ratio reported in the final column of Table 3.3. For example, the cancers and tumours programme covers all expenditure associated with ICD10 codes C00-C97 and D00-D49 but the PCT-based mortality data only relates to ICD10 codes C00-C97. At the national (all England) level, figures are avaitable which show that, in 2008, there were 62,072 deaths of those aged under 75 years from codes COO-C97) and that there were 63,076 deaths from codes C00-C97 and D00-D49 combined. In other words, the PCT level mortality data reflects $98.4 \%$ of all deaths associated with the expenditure codes. We adjust our cost of life (year) estimates for this mismatch.

We acknowledge that mortality is a more relevant outcome indicator for some programmes (e.g., for circulatory problems) than for others (e.g., for epilepsy) and, for this reason, we would expect better results in some programmes than others. We also acknowledge that this focus on mortality ignores the impact of expenditure aimed at chronic care and at palliative care. Nevertheless, our focus on mortality is purely practical: it is both a widely available measure and it is clearly a relevant outcome indicator. ${ }^{10}$

The mortality data provide us with a number of possible outcome indicators including the under 75 years of age standardised mortality rate (SMR) and the (under 75 years) standardised years of life lost rate (SYLLR). The SMR gives equal weight to all deaths irrespective of the age at which they occur but the SYLLR gives greater zueight to deaths that occur at earlier ages. For our purposes we focus on a measure of the avoidable years of life lost (YLL). ${ }^{11}$ This is calculated by summing over ages 1 to 74 years the number of deaths at each age multiplied by the number of years of life remaining up to age 75 years. The crude YLDrate is simply the number of years of life lost divided by the resident population aged under 75 years. Eike conventional mortality rates, the crude YLL rate can be age standardised to eliminate the effects of differences in population age structures between areas, and this (age) standardised YLL rate is the health outcome variable generally employed in this study.[77]
${ }^{10}$ The approach adopted here is extendable in principle to other non-mortality based outcome indicators. We illustrate such an application in Section B8.8 of Appendix B where we use EQ-5D utility scores pre- and post- an operative procedure from the PROMs programme to generate a non-mortality-based outcome indicator, and we use this indicator to estimate our outcome model.
${ }^{11}$ One exception to this is the mortality rate for the trauma and injuries programme where initially only SMRs were available.

Table 3.1: National (all PCT) expenditure per head ( $£$ ) and growth in expenditure (\%) by PBC group, 2003/4-2008/9

| PBC \# | PBC description | Spend <br> (£) per head <br> 2003/4 | Spend <br> (£) per <br> head <br> 2004/5 | Spend <br> (£) per <br> head <br> 2005/6 | Spend <br> (£) per <br> head <br> 2006/7 | Spend <br> (£) f per <br> head <br> 2007/8 | Spend <br> (£) per <br> head <br> 2008/9 | Growth <br> (\%) <br> 2004/5 | Growth <br> (\%) <br> 2005/6 | Growth <br> (\%) <br> 2006/7 | Growth $(\%)$ $2007 / 8$ | Growth (\%) $2008 / 9$ | Share of total spend (\%) 2004/5 | Share of total spend (\%) 2008/9 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | Infectious diseases | 17.95 | 20.22 | 23.61 | 20.88 | 22.08 | 23.46 | 13 | 17 | 12 | 6 | 6 | 1.7\% | 1.5\% |
| 2 | Cancers and tumours | 64.95 | 75.54 | 83.24 | 81.67 | 90.21 | 94.55 | 16 |  |  | 10 | 5 | 6.3\% | 6.2\% |
| 3 | Blood disorders | 14.08 | 17.00 | 17.48 | 16.58 | 19.44 | 19.50 | 21 | (3) | -5 | 17 | 0 | 1.4\% | 1.3\% |
| 4 | Endocrine, nutritional | 28.96 | 31.86 | 37.26 | 36.70 | 39.39 | 43.38 | 10 | 1 | -1 | 7 | 10 | 2.7\% | 2.8\% |
| 5 | Mental health | 133.31 | 146.83 | 158.95 | 166.53 | 180.90 | 191.21 | 10 | 8 | 5 | 9 | 6 | 12.2\% | 12.5\% |
| 6 | Learning disability | 37.93 | 43.37 | 46.54 | 48.36 | 54.20 | 56.11 |  | 7 | 4 | 12 | 4 | 3.6\% | 3.7\% |
| 7 | Neurological | 29.83 | 35.09 | 41.06 | 55.27 | 62.43 | 67.64 | 18 | 17 | 35 | 13 | 8 | 2.9\% | 4.4\% |
| 8 | Vision problems | 24.61 | 27.65 | 28.24 | 26.97 | 30.69 | 32.95 | 12 | 2 | -4 | 14 | 7 | 2.3\% | 2.2\% |
| 9 | Hearing problems | 5.73 | 6.32 | 6.27 | 6.21 | 8.07 | 8.16 | 10 | -1 | -1 | 30 | 1 | 0.5\% | 0.5\% |
| 10 | Circulatory disease | 110.12 | 122.37 | 124.28 | 122.06 | 124.77 | (29.94 | 11 | 2 | -2 | 2 | 4 | 10.2\% | 8.5\% |
| 11 | Respiratory system | 54.60 | 62.71 | 69.56 | 65.07 | 67.68 | 77.97 | 15 | 11 | -6 | 4 | 15 | 5.2\% | 5.1\% |
| 12 | Dental problems | 10.78 | 13.55 | 24.91 | 51.93 | 59.45 | 62.44 | 26 | 84 | 108 | 14 | 5 | 1.1\% | 4.1\% |
| 13 | Gastro intestinal system | 63.56 | 73.22 | 81.30 | 73.30 | 75.05 | 77.89 | 15 | 11 | -10 | 2 | 4 | 6.1\% | 5.1\% |
| 14 | Skin problems | 20.98 | 24.90 | 26.84 | 28.31 | 30.41 | 32.34 | 19 | 8 | 5 | 7 | 6 | 2.1\% | 2.1\% |
| 15 | Musculo Skeletal system | 61.36 | 71.72 | 74.74 | 66.75 | 75.91 | 79.68 | 17 | 4 | -11 | 14 | 5 | 6.0\% | 5.2\% |
| 16 | Trauma and Injuries | 62.31 | 72.13 | 76.41 | 57.29 | 57.56 | 63.54 | 16 | 6 | -25 | 0 | 10 | 6.0\% | 4.2\% |
| 17 | Genito Urinary system | 55.32 | 62.38 | 67.38 | 68.98 | 67.83 | 73.78 | 13 | 8 | 2 | -2 | 9 | 5.2\% | 4.8\% |
| 18 | Maternity | 52.28 | 55.04 | 60.42 | 57.6 | 57.09 | 60.44 | 5 | 10 | -5 | -1 | 6 | 4.6\% | 3.9\% |
| 19 | Neonate conditions | 11.72 | 13.93 | 13.42 | 13.17 | 15.15 | 17.23 | 19 | -4 | -2 | 15 | 14 | 1.2\% | 1.1\% |
| 20 | Poisoning | 9.68 | 12.32 | 14.25 | 14.59 | 15.84 | 18.31 | 27 | 16 | 2 | 9 | 16 | 1.0\% | 1.2\% |
| 21 | Healthy individuals | 20.29 | 22.77 | 26.18 | 26.85 | 31.44 | 35.74 | 12 | 15 | 3 | 17 | 14 | 1.9\% | 2.3\% |
| 22 | Social care needs | 24.81 | 30.93 | 33.59 | 30.29 | 35.29 | 36.58 | 25 | 9 | -10 | 17 | 4 | 2.6\% | 2.4\% |
| 23 | Other (includes GMS/PMS) | 136.94 | 157.75 | 171.82 | 209.70 | 232.02 | 227.71 | 15 | 9 | 22 | 11 | -2 | 13.2\% | 14.9\% |
| 1 to 23 | All PBCs | 1052.12 | 1199.60 | 1307.76 | 1345.10 | 1452.91 | 1530.59 | 14 | 9 | 3 | 8 | 5 |  |  |

Notes: (i) The population figures for $2003 / 4,2004 / 5$ and 2005 ( 6 are identical (the total for England is 49,175,998).
(ii) The corresponding figure for $2006 / 7$ is $50,476,231$, for $2007 / 8$ it is $50,695,989$, and for $2008 / 9$ it is $51,220,531$.
(iii) The spend per head figures are calculated by summing expenditure across all PCTs and dividing by the national population.
(iv) All figures are at current prices.

Table 3.2: PCT expenditure per head by PBC, 2008/9: (a) unadjusted; (b) adjusted for local costs; and (c) adjusted for local costs and local need.

|  | Programme budget category | Spend per head (unadjusted), $£$ |  |  |  | Spend per head (cost adjusted), $£$ |  |  |  | Spend per head (cost and need adjusted), $£$ |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Mean | StdDev | Min | Max | Mean | StdDev | Min | Max | Mean | StdDev | Min | Max |
| 1 | Infectious diseases | 26.5 | 24.6 | 8.6 | 151.8 | 25.7 | 21.7 | 8.6 | 136.7 | 25.0 | 21.4 | 9.5 | 139.5 |
| 2 | Cancers and tumours | 96.3 | 16.9 | 62.9 | 155.7 | 96.7 | 19.7 | 59.1 | 163.1 | 94.2 | 15.3 | 55.2 | 154.0 |
| 3 | Blood disorders | 20.3 | 7.0 | 7.7 | 49.4 | 20.2 | 6.5 | 8.0 |  | 19.7 | 6.0 | 8.2 | 44.2 |
| 4 | Endocrine, nutritional | 44.6 | 8.8 | 28.9 | 74.8 | 44.7 | 9.5 | 27.4 | 77. | 43.3 | 6.1 | 29.9 | 61.5 |
| 5 | Mental health | 201.4 | 60.0 | 118.9 | 474.1 | 200.3 | 54.0 | 122.8 | 422.8 | 194.0 | 41.9 | 132.3 | 362.0 |
| 6 | Learning disability | 56.8 | 18.8 | 7.7 | 125.9 | 57.0 | 19.4 | 6.8 | 123.6 | 55.7 | 18.8 | 6.7 | 136.6 |
| 7 | Neurological | 68.5 | 13.8 | 41.1 | 133.8 | 68.8 | 15.6 |  | 137.5 | 66.9 | 12.1 | 41.5 | 125.2 |
| 8 | Vision problems | 33.2 | 6.7 | 16.7 | 57.7 | 33.4 | 7.5 | 14.8 | 59.2 | 32.5 | 6.1 | 15.6 | 48.3 |
| 9 | Hearing problems | 8.6 | 3.7 | 0.9 | 24.0 | 8.7 |  | -0.9 | 25.5 | 8.3 | 3.3 | 0.8 | 22.0 |
| 10 | Circulatory disease | 131.6 | 26.7 | 88.0 | 317.3 | 132.2 | 30.5 | 78.2 | 327.6 | 128.5 | 24.4 | 75.7 | 326.9 |
| 11 | Respiratory system | 80.5 | 17.4 | 48.0 | 141.2 | 80.9 | 4 | 42.7 | 145.3 | 78.1 | 12.4 | 48.2 | 126.0 |
| 12 | Dental problems | 64.8 | 13.4 | 28.0 | 111.9 | 64.9 | 14.1 | 24.9 | 115.8 | 63.0 | 10.7 | 28.1 | 97.1 |
| 13 | Gastro intestinal system | 80.0 | 14.5 | 46.7 | 119.6 | 80.4 | 16.8 | 41.5 | 124.6 | 78.0 | 11.3 | 41.6 | 114.4 |
| 14 | Skin problems | 33.1 | 8.0 | 18.1 | 66.4 | 33 | 8.6 | 16.5 | 69.1 | 32.2 | 6.3 | 16.0 | 57.7 |
| 15 | Musculo Skeletal system | 79.9 | 17.6 | 43.3 | 127.3 | 80.4 | 19.9 | 39.6 | 132.5 | 78.2 | 16.6 | 41.0 | 116.4 |
| 16 | Trauma and Injuries | 63.2 | 16.7 | 12.5 | 139.3 | 63.4 | 17.4 | 11.5 | 125.0 | 61.8 | 15.6 | 10.4 | 103.6 |
| 17 | Genito Urinary system | 75.7 | 13.7 | 49.9 | 112.3 | 75.6 | 13.6 | 48.4 | 108.9 | 73.7 | 10.1 | 50.6 | 105.5 |
| 18 | Maternity | 63.3 | 16.7 | 24.6 | 124.4 | 63.1 | 15.8 | 21.9 | 117.9 | 61.4 | 12.8 | 24.4 | 96.5 |
| 19 | Neonate conditions | 18.4 | 7.3 |  | - 46.4 | 18.2 | 6.8 | 6.6 | 43.7 | 17.8 | 6.6 | 5.8 | 47.8 |
| 20 | Poisoning | 18.6 | 4.2 | 10.8 | 31.2 | 18.7 | 4.7 | 9.6 | 32.3 | 18.2 | 3.9 | 10.1 | 33.1 |
| 21 | Healthy individuals | 38.4 | 18.1 | 9.7 | 125.0 | 38.4 | 17.8 | 8.9 | 115.6 | 36.7 | 14.5 | 9.4 | 104.5 |
| 22 | Social care needs | 40.8 | 56.6 | Q. 1 | 415.2 | 41.2 | 59.2 | 0.1 | 432.9 | 39.7 | 55.0 | 0.0 | 411.5 |
| 23 | Other (includes GMS/PMS) | 230.8 |  | 138. | 396.1 | 230.2 | 42.4 | 140.7 | 356.5 | 226.8 | 45.8 | 134.1 | 346.0 |
| All | All PBCs | 1,575.6 | 190.7 | 1,225.7 | 2,079.9 | 1,576.3 | 217.3 | 1,183.0 | 2,173.1 | 1,534.0 | 86.2 | 1,390.1 | 1,987.0 | each PCT's population.

### 3.4.3 Other variables

We employ an instrumental variables (IV) estimation technique to our empirical models of the outcome and expenditure equations as described in the next section. This is due to (i) own programme expenditure is likely to be endogenous in the outcome equation and (ii) other programme need is likely to be endogenous in the own programme expenditure equation. Endogeneity of programme expenditure results from expenditure levels being responsive to levels of outcomes and/or unobserved need rendering expenditure correlated with the residuals in an OLS regression of outcomes on expenditure. Due to limitations of data other programme need in the expenditure equation is proxied by death rates (minus that due to the programme under investigation). This will be influenced by expenditure decisions, including expenditure in other programmes and is treated as endogenous in the expenditure model.

IV estimation basically involves replacing the endogenous variable in the equation of interest with its predicted value from an OLS regression which regresses the endogenous variable on a set of instrumental variables. These instruments should be good predictors of the endogenous variable (i.e., they should be relevant and strong predictors) but should be appropriately excluded from the equation of interest (i.e., they should be valid instruments).

We have a number of potential instruments available, mostly derived from 2001 Population Census. In our earlier studies we found that a small sub-set of these instruments proved sufficient to generate plausible results. These included: the proportion of the population providing unpaid care; the proportion of households that are one pensioner households; index of multiple deprivation; proportion of the population in the white ethnic group.

We also had available a further set of potential instruments and, where our more limited set of instruments failed to generate plausible results, we extended our instrument search to include this wider set of variables. This extended set of instruments is shown in Table 3.3. 12
Our instruments reflect factors, such as socio-economic deprivation and the availability of informal care in the community, which might indirectly impact upon mortality rates and/or health care expenditure levels. As we shall see, although our instruments 'pass' the appropriate statistical tests, some commentators claim that such tests may have 'low power' to detect the presence of invalid instruments. Consequently in section B9 of Appendix B we examine how sensitive our results are to the presence of invalid instruments.

Table 3.3 reports descriptive statistics for the socio-economic and needs variables used in the study (these statistics are for the variables in absolute form). For example, on average, lone pensioner households comprise $14 \%$ of all households, the 'white ethnic' group accounts for $89 \%$ of the population and $10 \%$ of the population provide unpaid care.

In addition to the instrumental variables, Table 3.3 also reports descriptive statistics for the Department of Health's 'need for health care' index, ${ }^{13}$ its need for HIV services index, and its need for maternity services index. The latter two indices are used to either supplement or replace the all service measure of need when estimating ourmodels. The 'need for health care' index averages about 1 but varies substantially, with some PCTs having a needs index more than $25 \%$ below the national average and others facing a need for health care more than $30 \%$ above the national average. The Table also reports descriptive statistics for some disease prevalence rates (e.g., for diabetes and for epilepsy) and, again, these are used to either supplement or replace the all service measure of need when estimating our models. Finally, the MFF index shows that input prices in the most expensive PCT are almost $20 \%$ above those in the least expensive PCT.

[^6]
### 3.5. Approach to model estimation

The theoretical framework suggests the specification and estimation of a system of equations, with an expenditure and health outcome equation for each of the 23 programmes of care. However, this approach makes infeasible data demands, requiring variables to identify expenditure, need, environmental factors and health outcomes in each of the 23 programmes of care. Moreover, mortality rates are available for less than half of the 23 programmes. Rather than estimate a system of equations, we proceed on a programme-byprogramme basis, estimating health outcome and expenditure equations for those programmes for which mortality data are available.

In line with the theoretical framework presented above, we specify the following expenditure (equation 3.5) and health outcome (equation 3.0) models for each of the 23 programmes of care. Accordingly, for the $j$-th programme of care we have:
$x_{i}=\alpha+\beta n_{i}+\gamma m_{i}+\theta y_{i}+\varepsilon_{i}, \quad i=1, \ldots, 152$
Equation 3.5
$h_{i}=\gamma+\delta n_{i}+\pi x_{i}+\epsilon_{i}, \quad i=1, \ldots, 152$
Equation 3.6
where $x_{i}$ is expenditure; $n_{i}$ is the own programme need for care; $m_{i}$ is the need for care in other programmes; $y_{i}$ is the total budget and $h_{i}$ is the health gain in $\mathrm{PCT}_{\mathrm{i}}$.

Ideally we should employ a programme specific indicator of the level of need for each care programme $\left(n_{i j}\right)$ but these are not readily available. When estimating both the outcome and expenditure models we therefore proxy the own programme health care need using the 'needs' component of the Department of Health's resource allocation formula. ${ }^{14}$ This needs element is specifically designed to adjust PCT allocations for local health care needs and accordingly, ceteris paribus, we would expect a positive relationship between expenditure and need for each programme of care. We would also expect a positive relationship between need and adverse health outcomes. 15 .

The expenditure model includes both the own programme health care need (which is proxied using the 'needs' component of the Department of Health's resource allocation formula) and the need for health care in all other programmes. In the absence of programme-specific measures of need, we use the 'all cause mortality rate excluding the mortality rate in the programme of interest', $m_{i}$, as the proxy for need in other programmes of care.

All variables have been log transformed so that parameter estimates can be interpreted as elasticities. In other words, a regression coefficient of 0.5 implies that a $1 \%$ increase in the regressor is associated with a $0.5 \%$ increase in the dependent variable.

### 3.5.1. IV estimation

Other programe need, $m_{i}$, in the expenditure equation 3.5 and expenditure, $x_{i}$, in the outcome equation 3.6 are both likely to be endogenous rendering OLS both biased and inconsistent. Endogeneity of programme expenditure results from expenditure levels being responsive to levels of outcomes and/or unobserved need. Other programme need in the expenditure equation is proxied by death rates which is influenced by expenditure decisions and hence is treated as endogenous. To deal with this endogeneity we -
${ }^{14}$ However, we do experiment replacing and supplementing this all service measure of need with more programme specific measures where these are available (e.g., the diabetes and epilepsy prevalence rates).
${ }^{15}$ Whilst need is a function of mortality/morbidity in the resource allocation formula, the relationship is not sufficiently strong enough for us to be concerned about the endogeneity of the need in any individual care programme.
employ instrumental variables (IV) estimation and implement two-stage least squares (2SLS). Unlike OLS, IV is a consistent estimator in the presence of an endogenous regressor and, although in finite samples the IV estimator will be biased, with the bias (providing certain assumptions are met) being less than that associated with OLS.

For the health outcome equation, IV estimation can be viewed as finding variables (instruments) that are good predictors of programme expenditure but which are appropriately excluded from the outcome equation of interest (that is, from equation 3.6) because they are not predictive of outcome. The assumption is that these instruments impact upon health outcome through their impact on expenditure only, and that they do not have a direct effect on the outcome. ${ }^{16}$

Similarly, for the expenditure equation, IV estimation can be viewed as finding variables (instruments) that are good predictors of the proxy for other programme need $\left(m_{i}\right)$ but which do not belong in the expenditure equation of interest (that is, equation 3.5). The assumption is that these predictors impact upon own programme expenditure only through their impact on other programme need and that they do not have a direct effect on own programme expenditure.

The outcome and expenditure equations for any given programme may contain different instrumental variables because these instruments are trying to predict different variables (own programme expenditure and other programme mortality respectively). In addition, the instrument set for, say, the expenditure equation may vary across programmes because the other programme need variable will reflect need in a different basket of programmes for each expenditure equation.

We have a number of potential instruments available, mostly derived from 2001 Population Census. In previous studies, we have often found that a small sub-set (œur) of these instruments often proved sufficient to generate plausible results. However, if plausible results were not obtainable with some combination of these four instruments, we employed an extended instrument set. Further details of the identification of suitable instruments for each model can be found in Section B7.3 of Appendix B.

Table 3.3: Descriptive statistics for the instrumental and other variables

| Description | Obs | Mean | Std. <br> Dev. | Min | Max |
| :--- | :--- | :--- | :--- | :--- | :--- |
| Proportion of residents born outside the European Union | 151 | 0.0794 | 0.0876 | 0.0088 | 0.3817 |
| Proportion of population in white ethnic group | 151 | 0.8927 | 0.1299 | 0.3942 | 0.9926 |
| Proportion of population of working age (16-74) with LLT illness | 151 | 0.1182 | 0.0250 | 0.0709 | 0.1798 |
| Proportion of population providing unpaid care | 151 | 0.0990 | 0.0118 | 0.0662 | 0.1221 |
| Proportion of population providing unpaid care (<20 hrs week) | 151 | 0.0667 | 0.0079 | 0.0461 | 0.0817 |
| Proportion of population providing unpaid care (20-49 hrs week) | 151 | 0.0113 | 0.0025 | 0.0065 | 0.0195 |
| Proportion of population providing unpaid care (>50 hrs week) | 151 | 0.0210 | 0.0051 | 0.0093 | 0.0353 |
| Proportion of population aged 16-74 with no qualifications | 151 | 0.2960 | 0.0642 | 0.1301 | 0.4555 |
| Proportion of population aged 16-74 that are full-time students | 151 | 0.0720 | 0.0270 | 0.0425 | 0.1626 |
| Proportion of households without a car | 151 | 0.2932 | 0.1046 | 0.1325 | 0.5761 |
| Proportion of owner occupied households | 151 | 0.6692 | 0.1128 | 0.2891 | 0.8205 |

${ }^{16}$ IV estimation of say, equation 3.6, involves a first-stage regression of the endogenous expenditure variable, x , on the instrument, $\mathbf{z}$, and the set of exogenous regressors in equation 3.6, n . Predictions, $\hat{\mathrm{x}}$, from this model can then be included in a second-stage regression of equation 3.6 as a replacement for the endogenous regressor, x .

| Proportion of households in rented social (LA/HA) housing | 151 | 0.2071 | 0.0918 | 0.0817 | 0.5356 |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Proportion of households in rented private housing | 151 | 0.0924 | 0.0449 | 0.0349 | 0.2961 |
| Proportion of lone pensioner households | 151 | 0.1434 | 0.0184 | 0.0979 | 0.1942 |
| Proportion of one parent households | 151 | 0.0684 | 0.0180 | 0.0401 | 0.1207 |
| Proportion of population aged 16-74 that are permanently sick | 151 | 0.0574 | 0.0213 | 0.0242 | 0.1215 |
| Proportion of population aged 16-74 are long-term unemployed | 151 | 0.0113 | 0.0052 | 0.0036 | 0.0287 |
| Proportion of 16-74 in employment that are in agriculture | 151 | 0.0117 | 0.0119 | 0.0016 | 0.0668 |
| Proportion of those aged 16-74 that are in professional occupations | 151 | 0.2672 | 0.0688 | 0.1470 | 0.495 |
| Index of Multiple Deprivation 2007 | 151 | 23.8098 | 9.1168 | 8.0857 | 48.2627 |
| Need index (incorporates CARAN formula) | 151 | 1.0253 | 0.1334 | 0.7311 | 3479 |
| MFF index for HCHS and prescribing | 151 | 1.0021 | 0.0559 | 0.9410 | 1.1243 |
| Diabetes prevalence rate 2007/8 (\%, over 17 years) | 151 | 5.4872 | 0.7982 | 3.22 | 8.51 |
| Epilepsy prevalence rate 2007/8 (\%, over 18 years) | 151 | 0.7884 | 0.1489 | 0.41 | 1.09 |
| HIV need index | 151 | 1.1848 | 1.4984 | 0.1648 | 8.3332 |
| Chronic kidney disease 2007/8 (\%, over 18 years) | 151 | 4.1687 | 1.2711 | 1.35 | 8.41 |
| Maternity need index | 151 | 1.0345 | $0.2106$ | 0.6845 | 1.8129 |
| Raw (unadjusted) population 2007/8 | 151 | 335,735 | 196,501 | 90,142 | 1,264,298 |

[^7]The available instruments reflect factors, such as socio-economic deprivation and the availability of informal care in the community, which might indirectly impact upon mortality rates and/or health care expenditure levels. The set of instruments associated with each estimated equation was selected on both technical and pragmatic grounds. From a pragmatic point of view, we require a parsimonious set of instruments that satisfy the necessary technicalcriteria. These are, firstly, that they have face validity, that is, that they are plausible determinants of the endogenous variable being instrumented, and secondly, that the instruments are both relevant and valid. The relevance of an instrument set refers to its ability to predict the endogenous variable of concern, whereas validity refers to the requirement that instruments should be uncorrelated with the error term in the equation of interest.

Should the instrument set be strong, relevant and valid, 2SLS will produce consistent estimates of the parameters of the reduced form models. We subject the instrument sets to tests for validity using the Sargan-Hansen test of over identifying restrictions.[78] The joint null hypothesis is that the instruments are valid instruments, i.e., they are uncorrelated with the error term, and that the instruments are correctly excluded from the outcome equation of interest. A rejection of the null hypothesis casts doubt on the validity of the instruments. We test for instrument relevance using Shea's [79] partial R-squared measure; this reflects the correlation between the excluded instruments and the endogenous regressor. However, even where valid and relevant, a non-zero but small correlation between the set of instruments and the endogenous regressors can lead to the problem of weak instruments, again rendering IV estimation biased. We test for the presence of weak instruments using the procedures set out in Stock and Yogo[80] and the Kleibergen-Paap LM statistic. A general test of model specification is provided through the use of Ramsey's[81] reset test for OLS and an adapted version of the test for instrumental variables[82].

Finally, we check that the presumed endogenous variable is in fact endogenous using the test proposed by Durbin.[83] If the null hypothesis of exogeneity cannot be rejected, then we revert to using OLS. While, in general, our instruments 'pass' the appropriate statistical tests, some commentators claim that such tests may have 'low power' to and hence may fail to reject the validity of the instruments when this is false in
small samples. Consequently in Section B9 of Appendix B we examine how sensitive our results are to the relaxation of the assumption that the instruments are valid.

Further details of our approach to IV estimation are set out in Appendix B.

### 3.6. Results

The work presented here builds on previous studies of the link between expenditure and health outcomes. Martin, Rice and Smith[60] reported outcome elasticities for two programmes (cancer and circulatory disease) using expenditure data for 2004/5 and pooled mortality data for 2002, 2003, and 2004. ${ }^{17}$ This work was extended in a subsequent study[63] to include several other programme and updated expenditure data (2005/6). However, the authors struggled to obtain sensible outcome models for some programmes of care. Attempts to improve model estimates by considering alternating measures of the population need for health care ${ }^{18}$ and an extended set of potential instrumental variables are presented in Section B7 of Appendix B. This work forms the basis for the set of key results from the empirical modelling of health care expenditures and outcomes using more contemporaneous data presented in the following sections. Details of all results presented are set out in Appendix B.

### 3.6.1. $2006 / 7$ expenditure data and mortality data for 2006/2008

This section presents results that relate expenditure in 2006 to mortality in the same year and in the two following years (i.e., in 2006, 2007 and 2008). Throughout our measure of the need for health care is derived from the Department of Health's resource allocation model based on the CARAN needs formula.[84] This represents a more up-to-date needs adjustment than the AREA based model[85] that has been applied in previous studies $[60,63]$ and is directly applicable to the 152 PCTs in existence in the 2006/7 expenditure year. Expenditure data has been adjusted for differences in input prices using the market forces factors (MFFs for HCHS and prescribing). ${ }^{19}$ The outcome and expenditure results for the big four programmes are shown in Table 3.4 with the releyant outcome and expenditure elasticities highlighted.

In all four outcome models expenditure has a significant negative effect on mortality and the all service measure of need has a significant positive effect. The all service measure of need squared is also positive and significant in the cancer outcome equation. In the respiratory outcome model, there is an additional indicator of need - the proportion of the population that are permanently sick - and this is both positive and statistically significant. The diagnostic statistics suggest that, in all four cases, own programme expenditure is endogenous and that the instruments are valid. They also suggest that the instruments are relevant. There is no evidence that the instruments are weak in three of the four outcome results. The Pesaran-Taylor test suggests that there is no evidence of model mis-specification.

However, the Kleibergen-Paap F statistic for the respiratory disease outcome model is 7.022 and this is less than the 'critical' target of 10.0. This indicates that the instruments may be weak and not good predictors of the programme expenditure. However, if we re-estimate this model having dropped the least significant instrument, the coefficient on own programme expenditure becomes - 2.622 and is significant at the $1 \%$ level. Moreover, there is now no evidence of weak instruments (the Kleibergen-Paap F statistic is 11.025)

[^8]and it is this coefficient that we use for the respiratory outcome model in the cost of a life year calculations below.

In three of the four expenditure models both the need and budget variables have a positive and significant effect on own programme expenditure. In addition, the proxy for need in other programmes is negative and significant in all four cases. The diagnostic statistics suggest that, for all four expenditure models, expenditure is endogenous and the instruments are valid. They also suggest that the instruments are relevant and there is no evidence that the instruments are weak. The Pesaran-Taylor test suggests that there is no evidence of model misspecification.

### 3.6.1.1. Cost of a life year

The outcome and expenditure elasticities presented in Table 3.4 can be used to calculate the costrof a life year in each programme. These calculations -- for both the big four programmes as well as for the other six programmes with mortality based outcome indicator -- are shown in Table 3.5. The cost of a life (year) estimates presented in Table 3.5 assume a $1 \%$ increase in each PCT's budget calculated as:
the cost of an additional life in a particular programme
$=$ the change in expenditure in that programme / the change in mortalityinthat programme
$=$ (annual spend $*$ expenditure elasticity) $/$ (annual mortality $*$ outcome elasticity)
and
the cost of an additional life year in a particular programme
$=$ the change in expenditure in that programme / the change in life years lost in that programme
$=$ (annual spend $*$ expenditure elasticity) / (annual mortality * outcome elasticity)
Thus an integral part of the calculation of the costof life year is the annual mortality (life years lost) figure associated with a particular programme. Ideally, the ICD10 coverage of the expenditure data should coincide with that of the mortality data. Hoyvever, as shown in Table B5.1 of Appendix B, the ICD10 coverage of the mortality data typically falls short of that for the expenditure data. Unless we adjust the annual mortality figure so that its ICD10 coverage approximates that of the expenditure data, our cost of life (year) estimates will be too large because they will underestimate the mortality gain.

Table 3.5 incorporates this ICD 10 coverage. ${ }^{20}$ The results show that the cost of a life year for the big four PBCs is estimated as $£ 10,604$, and for all ten programmes with a mortality outcome measure, the estimate is $£ 19,965$. For all programmes, assuming a zero gain for the 13 PBCs without an outcome indicator, the corresponding estimate is $£, 73,457$.

If we assume that PBC23 generates a zero health gain and that the gain attributable to the remaining 12 programmes is, on average, the same as that attributable to those with a mortality outcome measure, then the cost of a life year across all programmes is $£ 22,565 .{ }^{21}$

### 3.6.1.2. Non-PCT Department of Health funded expenditure

PCT expenditure accounts for a large proportion of Department of Health expenditure but PCTs do not account for the Department's entire budget. In 2006/7 the Department of Health's gross expenditure totalled $£ 83.5 \mathrm{bn}$. Charges raised $£ 3.4 \mathrm{bn}$ so net expenditure totalled $£ 80.1 \mathrm{bn}$. Of this net expenditure, PCTs accounted for $£ 67.3 \mathrm{bn}$ (that is, $84 \%$ ) and various other bodies accounted for the remaining $£ 12.8 \mathrm{bn}$.

[^9]A breakdown of this gross and net expenditure by major body is shown in Table B8.24 of Appendix B. The Department of Health has allocated net non-PCT expenditure across the 23 PBCs. Of the additional $£ 12 \mathrm{bn}$ of net expenditure, $£ 11.2 \mathrm{bn}(93 \%)$ has been allocated to PBC23. This largely reflects: (a) the allocation of almost all Strategic Health Authority expenditure to either PBC23B ('other: SHAs including workforce development committees') or PBC23X ('other: miscellaneous'), and (b) the allocation of almost two-thirds of Department of Health expenditure to PBC23X ('other: miscellaneous'). The remaining $£ 0.8 \mathrm{bn}$ of additional net expenditure is spread across all PBCs according to various allocation rules and although this approach avoids allocating expenditure to the 'Other: Miscellaneous' category, this allocation of expenditure does not necessarily reflect actual expenditure.

The cost of a life (year) estimates presented above are based on the impact of a $1 \%$ exogenous change in total net PCT spend. All of our outcome and expenditure models have been estimated using net PCT expenditure, and all of our elasticities relate to this expenditure. Implicitly we assume that any budgetary shock only affects PCT funding and that it leaves non-PCT funding unchanged. Suppose instead we assume a $1 \%$ exogenous change in the Departmental budget. We have no information on how this Departmental budgetary shock is likely to be split between PCT and non-PCTs budgets. One might assume that the non-PCT budget is as responsive to a Departmental budgetary shock as is the PCT budget. If this was the case then it would add $17.7 \%$ to our cost of a life year estimate for 2006/7. However, in the absence of any information about the responsiveness of the non-DCT budget, it is difficult to come to any firm conclusion about the impact of non-PCT expenditure on our cost of a life year estimates.

### 3.6.2. $2007 / 8$ expenditure data and mortality data for $2007 / 2009$

Outcome and expenditure models were estimated using updated data for expenditure (from 2006/7 to $2007 / 8$ ) and updated mortality data (from 2006/2007/2008 to 2007/2008/2009). Appendix B, Section B10 presents detailed discussion of the findings including tables of results.

### 3.6.2.1. Outcome models

As before we model outcome as a function of own programme expenditure and a measure of health care need, where the latter is proxied by the measure of need as employed by the Department of Health for resource allocation purposes. ${ }^{22}$ There are, however, a few exceptions. For the respiratory programme we further included the square of the measure of need to improve model fit. In some other PBCs we found that the all service measure of need performed poorly and we replaced or supplemented this measure with either a more programme specific measure (e.g., the epilepsy prevalence rate for neurological mortality) or with a better performing proxy for need (e.g., the percentage of residents born outside the EU for maternity/neonate mortality). These amendments improved model specification ${ }^{23}$. Full results for all programmes are presented in Table B10.1 Appendix B; below is a summary of the findings.
${ }^{22}$ Using the CARAN model (Department of Health (2009).
${ }^{23}$ In addition to respiratory and neurological programmes the other programmes where the all service measure of need was replaced are: endocrine: IMD07 and diabetes prevalence rate; genitor-urinary: lone parent households; infectious diseases: IMD07 and HIV need per head and its square; maternity and neonates: proportion born outside EU and proportion of population with no qualification aged 16 to 74 . For trauma and injuries, the all service measure of need was supplemented with the proportion of households without a car and proportion of full time students.

Table 3.4: Outcome and expenditure models for the big four programmes using spend data for 2006/7 (two MFFs) and mortality data for 2006/7/8

|  | (1) <br> PBC 2 <br> cancer <br> outcome model | (2) <br> PBC 2 <br> cancer <br> spend model | (3) <br> PBC 10 <br> circulation outcome model | (4) <br> PBC 10 <br> circulation spend model | (5) <br> PBC 11 <br> respiratory outcome model | (6) <br> PBC 11 respiratory spend model | (7) <br> PBC 13 <br> gastro outcome model | (8) <br> PBC 13 <br> gastro spend model |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| own programme spend per head | $\begin{gathered} -0.342 * * * \\ {[0.099]} \end{gathered}$ |  | $\begin{gathered} -1.434 * * * \\ {[0.218]} \end{gathered}$ |  | $\begin{gathered} -2.029 * * * \\ {[0.636]} \end{gathered}$ |  | $\begin{gathered} -1.536^{* * *} \\ {[0.468]} \end{gathered}$ |  |
| need CARAN per head | $\begin{gathered} 0.995 * * * \\ {[0.106]} \end{gathered}$ | $\begin{gathered} 1.626 * * * \\ {[0.343]} \end{gathered}$ | $\begin{gathered} 2.860 * * * \\ {[0.252]} \end{gathered}$ | $\begin{gathered} 2.306^{* * *} \\ {[0.372]} \end{gathered}$ | $\begin{gathered} 2.696 * * * \\ {[1.044]} \end{gathered}$ | $\begin{gathered} 1.449 * * * \\ {[0.331]} \end{gathered}$ | $\begin{gathered} 4.160 * * * \\ {[0.577]} \end{gathered}$ | $\begin{gathered} 2.040^{* * *} \\ {[0.378]} \end{gathered}$ |
| need CARAN per head squared | $\begin{gathered} 1.163^{* * *} \\ {[0.348]} \end{gathered}$ |  |  |  | $\begin{gathered} 2.451 \\ {[1.561]} \end{gathered}$ |  |  |  |
| SYLLR all deaths exclude cancer |  | $\begin{gathered} -0.855^{* * *} \\ {[0.191]} \end{gathered}$ |  |  | $N$ |  |  |  |
| PCT budget per head |  | $\begin{gathered} 0.465 \\ {[0.300]} \end{gathered}$ |  | $\begin{aligned} & 0.540^{*} \\ & {[0.299]} \end{aligned}$ |  | $\begin{gathered} 0.679 * * * \\ {[0.251]} \end{gathered}$ |  | $\begin{aligned} & 0.446^{*} \\ & {[0.263]} \end{aligned}$ |
| SYLLR all deaths exc circulatory |  |  |  | $-1.666 * * *$ |  |  |  |  |
| permanently sick |  |  |  |  | $\begin{gathered} 0.759 * * \\ {[0.367]} \end{gathered}$ |  |  |  |
| SYLLR all deaths exc respiratory |  |  |  |  |  | $\begin{gathered} -0.672^{* *} \\ {[0.305]} \end{gathered}$ |  |  |
| SYLLR all deaths exclude gastro |  |  |  |  |  |  |  | $\begin{gathered} -1.206^{* * *} \\ {[0.314]} \end{gathered}$ |
| lone pensioner households |  |  |  |  |  |  |  |  |
| Constant | $\begin{gathered} 6.501^{* * *} \\ {[0.436]} \end{gathered}$ | $\begin{gathered} 5.913 * * * \\ {[2.815]} \end{gathered}$ | 11.413*** $[1.046]$ | $\begin{gathered} 10.696^{* * *} \\ {[2.379]} \end{gathered}$ | $\begin{gathered} 13.756 * * * \\ {[3.279]} \end{gathered}$ | $\begin{gathered} 3.346 \\ {[2.075]} \end{gathered}$ | $\begin{gathered} 9.719^{* * *} \\ {[2.009]} \end{gathered}$ | $\begin{gathered} 8.370 * * * \\ {[2.299]} \end{gathered}$ |
| Endogeneity test statistic | 13.695 | 19.421 | 42.548 | 24.461 | 17.687 | 8.439 | 16.373 | 15.211 |
| Endogeneity p-value | 0.000215 | $1.05 \mathrm{e}-05$ | $6.90 \mathrm{e}-11$ | $7.58 \mathrm{e}-07$ | $2.60 \mathrm{e}-05$ | 0.00367 | $5.20 \mathrm{e}-05$ | $9.61 \mathrm{e}-05$ |
| Hansen-Sargan test statistic | 0.685 | $0.021 \sim$ | 0.949 | 1.262 | 1.462 | 0.302 | 2.761 | 0.0164 |
| Hansen-Sargan p-value | 0.408 | 0.084 | 0.814 | 0.261 | 0.227 | 0.583 | 0.0966 | 0.0898 |
| Shea's partial R-squared | 0.164 | 0.445 | 0.300 | 0.296 | 0.0785 | 0.327 | 0.140 | 0.356 |
| Kleibergen-Paap LM test statistic | 17.85 | 41.88 | 32.37 | 32.02 | 10.02 | 34.98 | 14.86 | 35.72 |
| Kleibergen-Paap p-value | 0.000133 | $8.04 \mathrm{e}-10$ | $1.61 \mathrm{e}-06$ | 1.11e-07 | 0.00666 | $2.54 \mathrm{e}-08$ | 0.000592 | $1.75 \mathrm{e}-08$ |
| Kleibergen-Paap F statistic | 13.28 | 56.69 | 17.14 | 31.84 | 7.022 | 20.94 | 11.63 | 22.40 |
| Pesaran-Taylor reset statistic | 0.00537 | 0.18 | 0.136 | 0.00349 | 0.0120 | 1.497 | 1.669 | 0.007 |
| Pesaran-Taylor p-value | , 0.242 | 0.668 | 0.712 | 0.953 | 0.913 | 0.221 | 0.196 | 0.935 |

Note: robust standard errors in brackets. *** $\mathrm{p}<0.01,{ }^{* *} \mathrm{p}<0.05,{ }^{*} \mathrm{p}<0.1$.

Two sets of models were estimated for three of the big four programmes (i.e., for cancer, circulatory disease, respiratory problems and gastro-intestinal problems). One of the two models used two instruments and so we report the instrument validity test statistic. In all three cases we failed to reject the null hypothesis of instrument validity. However, there is some evidence of weak instruments (at least in the respiratory and gastro-intestinal programmes) and if we dropped one instrument and re-estimated the model, evidence of instrument weakness disappeared. The removal of one instrument has little impact on the coefficient on expenditure and it is this coefficient that we use below in our cost of a life year calculations reported in Table 3.5.

For the big four programmes the need variable has a positive and significant effect on mortality, and expenditure has the anticipated negative effect. The diagnostic statistics reveal that, in all four PBCs, own programme expenditure is endogenous and that the instruments are valid. They also suggest that the instruments are relevant and there is no evidence that they are weak in the models with one excluded instrument. The Pesaran-Taylor test reveals no evidence of model miss-specification.

The outcome results for the other programmes are similar to but more diverse than those for the big four programmes. This is to be anticipated because mortality is a much rarer outcome in these programmes than it is in the big four programmes. Own programme expenditure is not endogenous in four of these programmes but we retain the IV estimator for three of these four because this vields more plausible results than the OLS estimator (the results are more plausible in the sense that the signs on the coefficients are more in line with our prior expectations) ${ }^{24}$.

Expenditure has the anticipated negative effect on mortality in the endocrine problems programme but this is not statistically significant. The all service measure of need is not relevant for this PBC; instead, we find that the diabetes prevalence rate is positively associated with mortality, as is a measure of deprivation (the IMD2007).
Mortality from epilepsy is negatively and significantly associated with expenditure in the neurological programme. Both the all service need for health care and the epilepsy prevalence rate are positively and significantly associated with mortality in this programme.
Expenditure has a negative and statistically significant effect on mortality (from renal problems) in the genitor-urinary problems programme. The prevalence of lone parent households is positively associated with mortality.

Expenditure has the anticipated negative effect on mortality in the infectious disease programme and this is statistically significant. The all service measure of need is not relevant for this PBC; instead, we find that a measure of need associated with HIV is positively associated with mortality, as is a measure of deprivation (the IMD2007).

Expenditure has the anticipated negative effect on mortality in the maternity \& neonates programme but the estimated coefficient is not statistically significant. In this PBC the generic all service measure of need has been replaced with two other indicators of deprivation - the proportion of residents born outside the EU and the proportion of those aged 16-74 without any qualifications - both of these are positively associated with mortality.

Finally, expenditure and need have the anticipated effects on mortality in the trauma and injuries programme. In addition, the proportion of households without access to a car is negatively associated with mortality from fractures (perhaps access to a car facilitates involvement in serious road traffic accidents), and the proportion of residents that are students is positively associated with mortality from fractures.

The relevant statistical test suggests that expenditure is endogenous in six of the ten programmes but we have retained the IV estimates for three of the other four programmes because they provide plausible results. The Hansen-Sargen test suggests that the selected instruments are valid, and the Kleibergen-Paap
${ }^{24}$ The four programmes are: endocrine, infectious diseases, maternity/neonates and trauma/injuries.

LM statistic suggests that they are relevant (i.e., correlated with the endogenous regressor). With the possible exception of the trauma and injuries programme, the Kleibergen-Paap F statistic suggests that we do not have a problem with weak instruments. ${ }^{25}$ Finally, the Pesaran-Taylor/Ramsey reset test statistics reveal no evidence of misspecification.

### 3.6.2.2. Expenditure models

The majority of the expenditure models contain the three variables: the PCT budget, a proxy for the own programme need for health care, and a proxy for the need for health care in other programmes. The budget term is positive in all eleven models and it is statistically significant in eight of these eleven models.

The usual proxy for the own programme need for health care (i.e., the all service measure of need) is present in six of the models and it is significant in five of them. Its presence is supplemented with the addition of its squared value to improve model fit in the respiratory problems programme. In some programmes (e.g., the endocrine, metabolic \& nutritional programme and the neurological programme) ${ }^{26}$, we have replaced and/or supplemented the all service measure of need with a more programme specific measure (e.g., the diabetes prevalence rate and the epilepsy prevalence rate) and these meastures of need have the anticipated positive impact on expenditure.

In addition, in a couple of other programmes we have used alternative proxies for the own programme need (e.g., with the use of the Department of Health's measure of maternity need in the maternity/neonates expenditure equation). Full results for all programmes are presented in Table B10.2 Appendix B; below is a summary of the findings.

For eight of the eleven programmes we have used the all cause mortality rate less own programme mortality rate as the proxy for the need for health care in other programmes, and the coefficient on this term is negative in seven programmes and statistically significant in six of the seven. In three programmes -- maternity/neonates, GMS/PMS and trauma \& injuries programmes -- we have used the all cause mortality rate as the proxy for the need for health care in other programmes due to difficulties associated with the measurement of the own programme mortality rate. The coefficient on this term is not significant in any of the three models.

The relevant statistical test suggests that expenditure is endogenous in six of the eleven programmes but we have retained the IV estimates for two other programmes (GMS/PMS and trauma \& injuries) because the IV estimator provides more plausible results. In the other three programmes we report OLS results.

The Hansen-Sargen test suggests that the selected instruments are valid, and the Kleibergen-Paap LM statistic suggests that they are relevant (i.e., correlated with the endogenous regressor). The KleibergenPaap F statistic suggests that we do not have a problem with weak instruments. Finally, the PesaranTaylor reset test statistics and the Ramsey reset F statistics reveal no evidence of model misspecification.

### 3.6.2.3. Calculation of the cost of a life and life year

Expenditure and outcome elasticities for preferred models are used to calculate the cost of a life year, both for individual programmes and for all programmes collectively. The relevant figures are summarised in Table 3.5.27 The cost per life year gained is $£, 13,830$ for the big four programmes and $£ 28,983$ for all ten programmes with a mortality-based outcome indicator. These represent $30 \%$ and $45 \%$ increases on

[^10]the respective costs for the previous year (i.e., using expenditure data for 2006/7 and mortality data for 2006/2007/2008).

If we assume that the other 13 programmes (all without a mortality based outcome indicator) offer no health gain, then the cost per life year across all PCT expenditure is $£ 82,765$. This is up from $£ 73,457$ using data for the previous year (an increase of $13 \%$ ).

In addition, if we assume that PBC23 generates a zero health gain and that the gain attributable to the remaining 12 programmes is, on average, the same as that attributable to those with a mortality outcome measure, then the cost of a life year across all programmes is $£ 31,846$ (it was $£ 22,565$ using data for the previous year).

The next section presents outcome and expenditure models using PB data for 2008/9 and mortality data for 2008/9/10, and it explores the reasons for the increase in the cost of an additional life yearidentified in this section.

### 3.6.3. $2008 / 9$ expenditure data and mortality data for 2008/2010

Outcome and expenditure models were estimated using updated data for expenditure (from 2007/8 to $2008 / 9$ ) and updated mortality data (from 2007/2008/2009 to 2008/2009/2010). Detailed results for the outcome model and expenditure model are shown in Tables B11.1 and B11.2, Appendix B respectively. First stage regressions for these IV models can be found in Tables BA. 9 and BA. 10 in the annex to Appendix B.

### 3.6.3.1. Outcome models

The majority of the outcome models contain the two variables: own programme expenditure and a measure of the need for health care (the measure of need as employed by the Department of Health for resource allocation purposes ${ }^{28}$ ). For the respiratory disease programme we have added the square of the need measure to improve the model fit. In other PBCs (e.g., for the endocrine, metabolic and nutritional programme), we found that the all service measure of need performed poorly and we have replaced it with a more programme specific measure (e.g., the diabetes prevalence rate) or with a better performing proxy for need (e.g., the percentage of residents born outside the EU for maternity/neonate mortality). ${ }^{29}$

The relevant statistical test suggests that expenditure is endogenous in six of the ten programmes but we have retained the IV estimates for the other four because they provide plausible results. The HansenSargen test suggests that the selected instruments are valid, and the Kleibergen-Paap LM statistic suggests that they are relevant (i.e., correlated with the endogenous regressor). The Kleibergen-Paap F statistic suggests that we do not have a problem with weak instruments. Finally, the Pesaran-Taylor reset test statistics reveal no evidence of misspecification.

In all of the big four programmes the need for health care variable has a positive and significant effect on mortality, and expenditure has the anticipated negative effect. As we have noted before, the outcome results for the other programmes are similar to but more diverse than those for the big four programmes. This is to be anticipated because mortality is a much rarer outcome in these programmes than it is in the big four programmes.

[^11]Table 3.5: Cost of life and life year estimates using expenditure data for 2006 and outcome data for 2006/7/8 (assumes zero health gain for 13 programmes) adjusted for the ICD10 coverage of the expenditure and outcome data


Expenditure has the anticipated negative effect on mortality in the endocrine problems programme and this is statistically significant. The all service measure of need is not relevant for this PBC; instead, we find that the diabetes prevalence rate is positively associated with mortality, as is a measure of deprivation (the IMD2007).

Expenditure has a negative but statistically insignificant impact on mortality from epilepsy in the neurological programme, and the all service indicator of the need for health care is positively and significantly associated with mortality in this programme.
Expenditure also has a negative but not statistically significant effect on mortality (from renal problems) in the genitor-urinary problems programme. The prevalence of lone parent households is positively associated with mortality.

Expenditure has the anticipated negative effect on mortality in the infectious disease programmerand this is statistically significant. The all service measure of need is not relevant for this PBC; instead, we find that a measure of need associated with HIV is positively associated with mortality, as is a measure of deprivation (the IMD2007).

Expenditure has the anticipated negative effect on mortality in the maternity \& neonates programme. In this PBC the coefficient on the generic all service measure of need is positive but not significant. It has been supplemented with two other indicators of deprivation - the proportion of residents born outside the EU and the proportion of those aged 16-74 without any qualifications - and both of these are positively associated with mortality.

Finally, we were unable to develop a plausible outcome model for the trauma and injuries programme.

### 3.6.3.2. Expenditure models

The majority of expenditure models contain the three yariables: the PCT budget, a proxy for the own programme need for health care, and a proxy for the need for health care in other programmes.

The budget term is positive and statistically significant in ten of the eleven models.
The usual proxy for the own programe need for health care (i.e., the all service measure of need) is positive and significant in five of the eleven results. In a couple of programmes (respiratory disease and endocrine problems) we have added the squared value of need to improve the model fit and in both cases this term is positive and significant. In some programmes (e.g., the endocrine PBC and the neurological PBC), we have replaced and or supplemented the all service measure of need with a more programme specific measure (e.g., the diabetes and the epilepsy prevalence rates) and these usually have a positive and significant impact on expenditure. In addition, in a couple of programmes we have used alternative proxies for ownprogramme need (e.g., with the use of the Department of Health's measure of maternity need in the maternity/neonates expenditure equation and the use of HIV need in the infectious diseases programme) 30

For eight of the eleven programmes we have used the all cause mortality rate less the own programme mortality rate as the proxy for the need for health care in other programmes, and the coefficient on this term is hegative in seven programmes and statistically significant in six of the seven. In three programmes -- maternity/neonates, GMS/PMS and trauma \& injuries programmes -- we have used the all cause mortality rate as the proxy for the need for health care in other programmes due to difficulties

[^12]associated with the measurement of the own programme mortality rate. The coefficient on this term is negative but not significant in these three models.

The relevant statistical test suggests that expenditure is endogenous in five of the eleven programmes but we have retained the IV estimates for two further programmes (endocrine problems and maternity/neonates) because the IV estimator provides more plausible results than the OLS estimator. In the other four programmes we report OLS results.

The Hansen-Sargen test suggests that the selected instruments are valid, and the Kleibergen-Paap LM statistic suggests that they are relevant (i.e., correlated with the endogenous regressor). The Kleibergen Paap F statistic suggests that we do not have a problem with weak instruments. Finally, the Pesaran= Taylor reset test statistics and the Ramsey reset F statistics reveal no evidence of model misspecification,

### 3.6.3.3. Calculation of the cost of a life and life year

Expenditure and outcome elasticities for our preferred models are used to calculate the cost of a life year, both for individual programmes and for all programmes collectively. This results in the cost per life year gained having increased slightly compared with that using the previous expenditure and mortality data set (i.e., for 2007 and 2007/8/9 respectively): increasing from $£ 13,830$ to $£ 14,650$ forthe big four programmes and from $£ 28,983$ to $£ 30,883$ for all ten programmes with a mortality-based outcome indicator. If we assume that the other 13 programmes offer no health gain, then the cost per life year across all PCT expenditure has increased from $£ 82,765$ in $2007 / 8$ to $£ 84,974$ in 2008/9.

In addition, if we assume that PBC23 generates a zero health gain and that the gain attributable to the remaining 12 programmes is, on average, the same as that attributable to those with a mortality outcome measure, then the cost of a life year across all programmes in $2008 / 9$ is $£ 33,333$. This is a $5 \%$ increase on the figure $(£, 31,846)$ for the previous year.

### 3.6.4. Comparing the cost of life year estimates associated with different data sets

Table 3.6 presents expenditure and outcome elasticities for the five combinations of expenditure and outcome data that have been used to estimate our model. It also reports the corresponding unadjusted cost of life year estimates (i.e., estimates that are unadjusted for the mismatch in the ICD10 coverage of the expenditure and mortality data). It is clear from this Table (see row 13) that the (unadjusted) cost of a life year for the ten programmes with a mortality based outcome indicator fluctuated around $£ 22,000$ for the first three sets of estimations (see columns M-O). However, using the two most recent sets of expenditure data (i.e., for $2007 / 8$ and then for 2008/9), the figures in the table suggest that this cost has increased to about $£ 38,000$.

What are the proximate causes of this increase? Recall that the cost of a life year is calculated as
the change in expenditure associated with a $1 \%$ budget increase
the change in the number of life years lost associated with this increase
For 2006/7 (using mortality data for 2006/7/8) and for the ten programmes with a mortality based outcome indicator, the cost of a life year is calculated as

$$
(£ 32,911 \mathrm{~m} * 0.01 * 0.561) /(1,667,250 * 0.01 * 0.465)=£ 184.53 \mathrm{~m} / 7,760=£ 23,780
$$

For 2007/8 (using mortality data for 2007/8/9) and for the ten programmes with a mortality based outcome indicator, the cost of a life year is calculated as

$$
(£, 34,434 \mathrm{~m} * 0.01 * 0.749) /(1,635,716 * 0.01 * 0.414)=£ 257.94 \mathrm{~m} / 6,768=£, 38,110
$$

It is clear that the $60 \%$ increase in the cost of a life year between 2006/7 and 2007/8 is largely attributable (a) to the $40 \%$ increase in the additional expenditure directed towards these 10 programmes corresponding to a $1 \%$ budget increase and (b) to the $12 \%$ decline in the number of life years saved by this increase in expenditure.

The rise in the share of the budget increase directed towards these programmes can be attributed to the increase in the expenditure elasticity associated with these ten programmes (up from 0.561 to 0.749 ). The decrease in the number of years of life saved can be largely attributed to the $12 \%$ decline in the outcome elasticity associated with these programmes, down from -0.877 to -0.778 (see row 13, columns J and K of Table 3.6). ${ }^{31}$ However, it is not clear why such rather dramatic changes should have taken place.

If we correct the cost of life year estimates adjusting for the mismatch in the ICD10 coverage of the expenditure and mortality data, these reveal similar increases in the cost of a life year between 2006/7 on the one hand and 2007/8 and 2008/9 on the other. The cost of a life year increased from $£ 19,965$ in $2006 / 7$ to $£ 28,983$ in 2007/8 for the ten programmes with mortality rate, an increase of $45 \%$, and it increased from $£ 22,565$ to $£ 31,846$ for all programmes if we assume a zero health gain in PBC23 and the same gain in the other 12 programmes as in the ten with a mortality rate (an increase of $41 \%$ ).

A potential reason for this apparent step change in the cost of a life year is the adjustment that was made to the methodology for the collection of the 2007/8 programme budgeting data. In previous years expenditure that was not directly attributable to a particular programme category was apportioned using admitted patient care percentages. ${ }^{32}$ In other words, if $x \%$ of total admitted patient care expenditure was allocated to PBC 1, then $\mathrm{x} \%$ of all expenditure that was not directly attributable to a particular programme category was also allocated to PBC 1. With effect from 2007/8, however, NHS organisations were asked to select an appropriate basis for the apportionment of this non-programme specific expenditure and that, where no reasonable basis existed, such expenditure was to be allocated to the 'Other - Miscellaneous' (PBC 23X) category.

The Department of Health estimates that this allocation rule change increased the amount of expenditure attributed to PBC 23 X by $£ 700$ million. It will also, of course, have reduced expenditure across other programmes by the same amount in total. Howeyer, not all programmes will have been equally affected; PBCs that are more heavily inpatient based would have 'lost' expenditure while others, such as learning disabilities, social care, and mental health, will have 'lost' considerably less. In addition, not all PCTs will have been equally affected because each will have employed different apportionment rules for the nonprogramme specific expenditure. [86]

Although this allocation rule change has considerably increased the estimated cost of a life year, we believe that this rule change has led to a more accurate allocation of expenditure across PBCs, and that the more recent estimates of the cost of a life year (for 2007/8 and 2008/9) are more accurate than those for the earlier years (for 2005/6 and 2006/7).

### 3.6.5. Adjusting the cost of a life year estimates to constant prices

The estimates of the cost of a life year presented above are all at current prices. To put them on a constant price basis, we need an index of pay and price inflation for the labour and goods/services purchased by the NHS. Curtis[87] reports a pay and prices index for Hospital and Community Health

[^13]Table 3.6: Expenditure and outcome elasticities for five combinations of expenditure and outcome data, and corresponding (unadjusted) cost of life year estimates


Notes:
(i) that the spend and outcome elasticities reported for groups of programmes are the implied elasticites calculated from the totals for the relevant individual programmes (i.e., group spend elasticity $=\sum$ (PBC spend $* P B C$ spend elasticity) $/ \sum$ PBC spend, and group outcome elasticity $=\sum\left(\mathrm{PBC}\right.$ mortality $* \mathrm{PBC}$ outcome elasticity) $\sum \sum \mathrm{PBC}$ mortality). For the purpose of the calculation of the group outcome elasticvity, we have used the years of life lost as the mortality indicator.
(ii) for each individual programme: the cost of an additionalife year $=\%$ change in spend ${ }^{*}$ annual spend/(outcome elasticity $*$ annual life years lost)
(iii) for a group of programmes: the overall cost of andditional life year $=\sum$ (annual spend* ${ }^{*}$ spend elasticity) $/ \sum$ (spend elasticity*outcome elasticity*annual life years lost)

Services and this implies an inflation rate of $3.7 \%$ in 2006/7, $2.9 \%$ in $2007 / 8$, and $3.9 \%$ in 2008/9. ${ }^{33}$ If we assume that similar inflation rates also apply to the purchase of pharmaceuticals and the provision of primary care (items that are excluded from the HCHS index), then we can use these figures to put the estimates of the cost of a life year on a constant price basis.

For example, if we assume that PBC23 generates a zero health gain and that the gain attributable to the 12 programmes without a mortality indicator is, on average, the same as that attributable to those with a mortality outcome measure, then the cost of a life year across all programmes in 2008/9 is $£ 33,333$ at current (2008/9) prices. The cost for $2007 / 8$ is $£ 31,846$ at current $(2007 / 8)$ prices or $£ 33,088$ at constant (2008/9) prices, and the figure for $2006 / 7$ is $£ 22,565$ at current $(2006 / 7)$ prices or $£ 24,125$ at constant $(2008 / 9)$ prices. The conversion of the costs from a current to constant price basis has relatively little impact because the inflation rate over the relevant period is quite small.

### 3.7. Summary and concluding remarks

The findings presented in this report build on four previous studies. These studies and the results presented here draw on the availability of two new data sets to obtain empirical estimates of the relationship between mortality and expenditure across all English local health authorities.

In this research we have extended the previous studies in several ways. First, we have derived plausible outcome and expenditure models for a larger number of programmes (ten) than previous studies.

Second, we relate expenditure in time period $t$ to mortality in that period $(t)$ and in the next two periods ( $t+1$ and $t+2$ ). In other words, we assume that the health benefits associated with expenditure occur either in the same period as the expenditure or in the next two periods. This is an improvement on past practice where data constraints forced researchers to relate expenditure to the current and two previous periods. ${ }^{34}$ When we re-estimated our models using expenditure data for 2006/7 and mortality data for $2006 / 7 / 8$, we found that the cost of a life year across the ten programmes with a mortality based outcome indicator is $£ 23,780$ (up from $£ 20,893$ when expenditure data for 2006/7 is combined with mortality data for 2004/5/6; an increase of $14 \%$ ),

Third, we have noted the mismatch in the ICD10 coverage of the expenditure and mortality data. If we adjust the calculation of the cost of a life year for $2006 / 7$ for this mismatch then the cost of a life year across the ten programmes with a mortality based outcome indicator declines from $£ 23,780$ to $£ 19,965$ (a decrease of $16 \%$ ).

Fourth, previous estimates of the cost of a life year have been for individual programmes of care. In this report we have presented estimates of the cost of a life year for an enlarged number of programmes and, with the aid of assumptions about the productivity (health gain) of programmes without a meaningful mortality-based outcome indicator, we have extended our individual programme estimates to incorporate expenditure across all programmes of care. Thus for 2006/7, the cost of a life year for those PBCs with a mortality based outcome indicator is $£ 19,965$. If we assume that (a) that the health gain associated with PBC23, which includes primary care and workforce training expenditure, are reflected in the mortality rates for disease specific programmes and (b) that the average health gain across the other programmes without a mortality based outcome indicator is the same as that for those PBCs with a mortality based outcome indicator, then the cost of life year across all programmes is $£ 22,565$.

Fifth, we have extended our cost of life year estimates beyond 2006/7. Re-estimation of our model using budgeting expenditure for 2007/8 generates an all programme cost of a life year estimate of $£ 31,846$, and

[^14] (Curtis, 2011, p209).

[^15]re-estimation of our model using budgeting expenditure for 2008/9 generates a similar cost of a life year estimate ( $£ 33,333$ ). Together, the last two estimates suggest that there has been step change in the cost of a life year, and that this appears to have occurred between 2006/7 and 2007/8. The cost of a life year estimates are very similar up to and including 2006/7, and they are very similar for 2007/8 and 2008/9. However, there is a substantial difference between the figures for 2004/5, 2005/6 and 2006/7 on the one hand (at about $£ 22 \mathrm{k}$ ), and for $2007 / 8$ and $2008 / 9$ on the other (at about $£ 33 \mathrm{k}$ ). The reason for this step change is not obvious but it might be due to changes in the algorithm used by the Department of Health to allocate non-admitted patient care activity to budget categories. Although this allocation rule change has considerably increased the estimated cost of a life year, we believe that this rule change has led to a more accurate allocation of expenditure across PBCs, and that the more recent estimates of the cost of a life year (for 2007/8 and 2008/9) are more accurate than those for the earlier years (for 2005/6 and $2006 / 7$ ). A summary of the estimates of the cost of a life year adjusted for the mismatch between ICD10 chapters for expenditure and mortality are provided in Table 3.7.

Virtually all of the cost of a life year estimates presented in this report are calculated at current prices. However, it is possible to put them on a constant price basis using the Hospital and Community Health Services pay and prices index.[87] For 2006/7, 2007/8 and 2008/9 this index recorded an annual rate of inflation of about $3.5 \%$ and so the impact of this constant price adjustment is fairly minimal. For example, if we assume that PBC23 generates a zero health gain and that the gain attributable to the 12 programmes without a mortality indicator is, on average, the same as that attributable to those with a mortality outcome measure, then the cost of a life year across all programmes at constant 2008/9 prices is $£ 33,333$ for 2008/9, $£ 33,088$ for $2007 / 8$, and $£ 24,125$ for 2006/7.

Finally, although previous results and our current models 'pass' the appropriate statistical tests and, in particular, the Hansen-Sargen test for valid instruments, we are aware that this test might be unable to detect the presence of invalid instruments in some circumstances and that the validity of instrumental variables is often open to question. Responding to this, several studies [88, 89] have suggested that researchers using IV techniques should subject the estimated coefficient on the endogenous variable to a sensitivity analysis. We undertake a comprehensive sensitivity analysis for the outcome equation for each of the big four models. This sensitivity analysis reveals that uncertainty associated with instrument validity has little effect on our estimate of the cost of a life year but it does increase the degree of uncertainty associated with this estimate.

We recognize that this study has a number of limitations. The estimates of the cost of an additional life year for programmes with a mortality-based outcome indicator are unadjusted for the quality of life during the additional year. Accordingly, the quoted costs will be an under-estimate of the QALY-adjusted cost of a life year to the extent that additional life years are not in perfect health. In previous studies we have noted that a rudimentary adjustment for this issue using HODaR data increased the cost of a life year by about $50 \%$ to $60 \%$. $[60,63]$

At the same time, however, the estimated costs will exaggerate the cost of an additional QALY-adjusted year for those programmes with a mortality-based outcome indicator because they ignore any health benefits that are not associated with a reduction in mortality. In other words, expenditure that improves the qualityof life (e.g., cancer palliative care) but which does not extend the length of life is implicitly givenazero health gain value.

In-addition, the expenditure data relates to expenditure on all patients whereas the mortality data is based on a life expectancy of 75 years. Thus implicitly our calculations attribute a zero health gain to all expenditure on those aged over 75 . To illustrate the magnitude of the potential health gain ignored by this restriction, note that in a recent study of costs associated with all inpatient and outpatient activity (excluding mental health), those aged over 75 years accounted for $25 \%$ of all costs in 2007/8[90] for details of this study).

The results presented in this study are all from the estimation of the relationship between expenditure and mortality using data for a single time period. With the availability of several years of data for both expenditure and mortality, we wanted to estimate a panel data model because a panel can offer advantages
over a one period model (e.g., it is better able to handle any unobserved heterogeneity across PCTs). However, most of the instruments employed here are based on the 2001 Census and thus time invariant rendering them of little use in panel data modelling.

Table 3.7: Adjusted cost of life year estimates for various combinations of programmes
A
C


Cost per life year

Programme budgeting category
(adjusted for ICD10 coverage of spend and mortality data) 2006/7
£. 16,121
2 Circulatory disease
3 Respiratory problems
4 Gastro-intestinal problems
5 All big four programmes

6 Other six programme with a mortality rate
7 All ten PBCs with a mortality rate
(a) If we assume a zero health gain in those PBCs without a mortality rate. 8 All 23 programmes
...or (b) if we assume a zero gain in PBC23 and that the average gain from the
the 10 PBCs with a mortality rate is applied to the remaining programmes
9 All 23 programmes
C
$£ 22,565$
£31,846
£33,333

Note that the figures for 2006/7 relate to the use of mortality for 2006/2007/2008 combined.

# Chapter 4: Translating mortality effects into life years and quality adjusted life years 

### 4.1 Introduction

This chapter presents an overview of how the results of the econometric work undertaken to estimate the link between NHS spending and mortality, which was summarised in the previous chapter and detailed in Appendix B, can be translated in to effects on life years and quality adjusted life years (QALYs).

In this chapter we present three sequential steps of analysis which lead to estimates of the overall cost per QALY threshold for the NHS:
i. In section 4.2 we reconsider how the estimated effects on mortality from the econometrics work conducted in Chapter 3 might better translate in to life years by exploring the limitations of mortality data available at PCT level and the published years of life lost (YLL) figures presented. We explore how these estimates might be improved using additional data and analysis.
ii. In section 4.3 we consider how these estimates of life year effects might be adjusted for the quality of life in which they are lived, taking account of the genderand the age at which life years are gained or lost as well as the disutility associated with particular diseases.
iii. In section 4.4 we explore ways to also take account of those effects on health not directly associated with mortality and life year affects (i.e., the 'pure' quality of life effects) to estimate an overall cost per QALY threshold.

This sequence of analysis is set out and explained based on the analysis of 2006 expenditure and mortality data from 2006 to 2008. In section 4.5 we present estimates for 2008 expenditure and 2008 to 2010 mortality data using the same methods and discuss the uncertainties associated with these estimates. As in the previous chapter much of the detail of data and analysis that supports this overview is presented in an appendix (see Appendix C). At the end of each section we present a summary which includes a central 'best' estimate as well as extreme lower and upper bounds for the cost per life year and cost per QALY threshold.

The core assumptions which underpin these three values are common across sections 4.2 to 4.5 . The central or 'best' estimate is based on two assumptions one conservative and the other more optimistic with respect to the health effects associated with expenditure. The first is that the health effects of changes in one year of expenditure are restricted to one year. This is implicit in the estimates of outcome elasticities presented in the previous chapter. ${ }^{35}$ This is likely to underestimate effects on mortality since expenditure that redaces mortality risk for an individual in one year may well also reduce their risk over subsequent years; possibly over the whole of their remaining disease duration. Expenditure may also prevent disease in future patient populations. Therefore, total health effects will be underestimated and the cost per life year or QALY threshold will be overestimated. Although undoubtedly conservative, it may be offset to some extent by the more optimistic assumption used to translate mortality effects into life years. Any death averted by expenditure in one year is assumed to return the individual to the mortality risk of the general population, i.e., the years of life gained associated with each death averted are based on what would have been their life expectancy taking account of their of age and gender (using life tables for the general population).

The extreme upper and lower bounds for cost per life year and cost per QALY thresholds are based on making both assumptions either optimistic (providing the lower bound for the threshold) or both

[^16]conservative (an upper bound for the threshold). The lower bound is based on assuming that health effects are not restricted to one year but apply to the remaining disease duration for the population at risk during the expenditure year (although this still does not account for the effects of expenditure on preventing disease). The upper bound is based on the combination of assuming that health effects are restricted to one year and that any death averted is only averted for the minimum duration consistent with the mortality data used to estimate the outcome elasticities in Chapter 3 (see Section 4.2 .5 for a more detailed discussion). It is very important to note that the lower and upper bounds represent extreme values rather than alternative but plausible views that could reasonably be taken. We discuss this in more detail in Section 4.5 and explain why establishing narrower bounds, which might retain some plausibility, has not been possible given the data available and therefore the analysis that has been feasible.

### 4.2 From mortality to life years

In this section we summarise our examination of a number of issues associated with available PCT-based mortality data and the associated published estimates of YLL. We then examine how, given the limited information available about the population at risk in each PBC we might take proper account of the fact that some of the observed deaths would have occurred anyway (had the same population not been at risk in the particular PBC ) when estimating YLL, i.e., taking account of unobserved counterfactual deaths. This allows us to estimate the YLL that better reflects the effect of expenditure on the mortality observed in each PBC, and infer the excess deaths associated with each PBC. Finally we present cost per death averted and cost per life year which accounts for the issues raised in this section.

### 4.2.1 Mortality and YLL coverage

The mortality data that is available at PCT level does not offer full coverage of all deaths across all the ICDs that make up each PBC (see Table B5.1 in Appendix) B for how three-digit ICD-10 are mapped to PBCs). However, national (English) data is available that covers all deaths associated with all the ICDs that make up each PBC. Therefore, it is possible to adjust the incomplete reporting of mortality at PCT level (see section 3.2 in Chapter 3) before applying the estimated outcome elasticities to calculate the deaths averted due to expenditure. ${ }^{36}$ Applying published estimates of YLL per death to all the deaths averted provides the estimate of the cost per life year reported in Chapter 3.

The published estimates of YLL (NHS IC) used in Chapter 3 only include deaths below 75 years (but exclude deaths below 1 year) and are based on the difference between age 75 and the age of each death below 75. These estimates have the same limited coverage as PCT level mortality data so are not available for all the ICDs that make up each PBC. Therefore, applying the available estimates of YLL per death to the estimated number of deaths averted requires an assumption that the YLL per death is similar for those groups of ICDs covered and not covered by the published YLL figures.

This can be examined by using national ONS data to calculate YLL in the same way as NHS IC, but with full coverage of all the ICDs that make up each PBC. ${ }^{37}$ Although ONS data provides complete coverage and reports gender; age at death is only reported in 5 year ranges (these data are not available at PCT level so could not be used when estimating outcome elasticities in Chapter 3). Therefore, using ONS data to estimate YLL requires taking the midpoint of each range as the age of death, i.e., assuming reported deaths are equally likely over the range in which they are reported. For this reason it is not possible to precisely recover the published YLL figures using ONS data for those ICD groupings that can be
${ }^{36}$ This does assume that the proportionate effects on mortality due to changes in expenditure are similar for mortality that is and is not recorded at PCT level. This seems more reasonable than assuming no effect of expenditure on mortality that happens not to be recorded at PCT level.
${ }^{37}$ Although published estimates of YLL are available from NHS IC for PBC16 (Trauma and injuries), ONS does not provide the information required to calculate YLL for this PBC. However, the estimated outcome elasticity was zero for 2006 expenditure and 2006 to 2008 mortality. Therefore, this PBC does not contribute any changes in health outcomes due to changes in expenditure in subsequent estimates of cost per life year and QALY thresholds anyway.
precisely matched to the NHS IC coverage. However, the differences are small (see Table C2, Appendix C), suggesting that taking the midpoint of each range as the age of death may be a reasonable approximation.

The differences between estimates of YLL based on ONS and NHS IC data are, however, much more significant and are reported in Table 4.1. These reflect differences in the distribution of ages at death between those groups of ICDs covered and not covered in the NHS IC figures. For example, NHS IC figures available at PCT level for PBC7 (neurological problems) have low coverage of all deaths in this PBC ( 0.136 in column 1). The deaths that are reported in NHS IC are associated with epilepsy and the YLL (22,046 in column 2) reflects the generally younger age at death in this group. When adjusted for full coverage $(22,046 / 0.136=162,100$ in column 3) the estimated YLL is much greater than the YLL based directly on all deaths by age group reported in ONS. This difference in YLL reflects the fact that the deaths in PBC7 which are not covered by NHS IC figures tend to be in older age groups so generate fewer YLL.

Table 4.1. Estimates of YLL for NHS IC and ONS

| PBC |  | Coverage of mortality data relative to spend data [1] | $\begin{gathered} \mathrm{YLL}_{<75} \\ (\underset{\text { NHS IC }}{ } \text { (2] } \end{gathered}$ | YLL<75 <br> adjusted <br> (NHS IC) <br> [3] | YLL no adjustment needed (ONS) [4] | Difference from adjusted NHS IC to ONS [5] |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | Infectious diseases | 1.00 | 35,517 | 35,517 | - 40,928 | 15\% |
| 2 | Cancer | 0.98 | 735,674 | 747,636 | 758,804 | 1\% |
| 4 | Endocrine problems | 0.63 | 19,224 | 30,322 | 41,548 | 37\% |
| 7 | Neurological problems | 0.14 | 22,046 | 162,100 | 93,755 | -42\% |
| 10 | Circulatory | 0.99 | 453,878 | 4,457,538 | 481,246 | 5\% |
| 11 | Respiratory | 0.77 | 108,074 | 139,812 | 147,465 | 6\% |
| 13 | Gastro-intestinal | 0.57 | 115,303 | 201,931 | 177,532 | -12\% |
| 17 | Genito-urinary | 0.17 | 3,343 | 19,438 | 17,380 | -11\% |
| 18+19 | Maternity \& neonates | 0.68 | 164,200 | 241,826 | 15,409 | -94\% |

Using ONS data also allows deaths under the age of one year to be appropriately assigned to PBCs via the ICD in which they occurred (NHS IC YLL figures exclude deaths under one year), rather than assigning them all to PBC18 \& 19 as in the previous Chapter. ${ }^{38}$ This explains the large reduction in YLL for PBC18 \& 19 (Maternity and neonates) as much of the mortality is re-assigned to ICDs which contribute to other PBCs. Since most of the deaths that are re-assigned are allocated to PBC1 (infectious diseases) the YLL for this PBC increases despite complete reporting of deaths at PCT level and full coverage by NHS IC figures (see also Table C4 in Appendix C).

### 4.2.2 Life expectancy and $Y L L$

As noted above the NHS IC estimates of YLL only include deaths below 75 years and are based on the difference between age 75 and the age of each death below 75 . Implicitly this treats 75 as the appropriate normal life expectancy for males and females for the population at risk in each PBC. However, with the exception of maternity and neonates most deaths in PBCs occur above the age of 75 and life expectancies are signifieantly greater than 75 . For example, based on 2006 to 2008 data, life expectancy for the general population is 80.7 for males and 84.4 for females (considering age distribution) and even life expectancy at birth is greater than 75 ( 77.74 for males and 81.88 for females). ${ }^{39}$

[^17]Based on ONS data YLLs can be re-recalculated using gender specific life expectancy for the general population. ${ }^{40}$ When increasing life expectancy (LE) two effects occur, both of which tend to increase estimates of YLL. Firstly, more deaths are included in the YLL calculation (those that occur between age 75 and LE) and secondly, each death previously counted below 75 will generate 5.7 or 9.4 more YLL for males and females respectively. The effect on the number of deaths and the YLL for each PBC of using the life expectancy of the general population is reported in Table 4.2 (see columns 1, 2 and 3).

Table 4.2. The difference in YLL by life expectancy

*LE male=80.7, female=84.4
The number of deaths counted below LE increases for every PBC except for maternity \& neonates because, as expected, all deaths are below age 75 in PBC18 \& 19. However, YLL increases for all PBCs reflecting the additional years otherwise expected to be lived to an older LE. Of course including more of the deaths observed in each PBC and the greater YLL associated with them will generate more deaths averted and more life years gained when applying the same proportionate effects from the outcome elasticities estimated in Chapter 3. Therefore, the costper death averted and cost per life year threshold are lower using these figures than those reported in Chapter 3 (see Table 4.6 below and Table C7 in Appendix C for a summary of the effects on the thresholds). However, there are good reasons why YLL figures calculated as the sum across all deaths below LE of the difference between age of death and LE are overestimated. This is dealt with in the next section (Section 4.2.3). In Section 4.2.4 we take account of the fact that some of the deaths observed in a PBC would have occurred anyway in a similar 'normal' population (i.e., the counterfactual population not at risk through membership of the PBC) so not all observed deaths are 'excess' and generate YLL.

### 4.2.3 YLL and accounting for counterfactual deaths

The estimates of YLL based on ONS data overcome many of the limitations of the published NHS IC figures. However, the YLLs reported in Tables 4.1 and 4.2, are calculated in the same way as the NHS IC figures, by taking the difference between a fixed LE and the age at death of deaths observed below that LE. This will tend to-overestimate the YLL for two reasons: i) it does not account for the fact that not all deaths observed below LE are 'excess' deaths in the sense that some deaths would have occurred (at the same age) in a similar population not at risk in the PBC and ii) some of the deaths observed above LE may be 'excess' deaths that would not otherwise have occurred at that age. The YLL associated with deaths that are in fact not 'excess' below LE will be greater than excess death not counted above LE. However, the overall effect on YLL, and the cost per life year, will depend on the number of deaths aboye and below LE that are excess. Therefore, estimates of YLL are required which take account of the 'counterfactual' deaths that would have occurred even if the population in the PBC was not at risk through membership of the ICD codes that make it up, but faced the same mortality risks as the general population, accounting for the age and gender distribution of the PBC population.

[^18]Ideally, with reliable information about the size of the population at risk in each PBC and its age and gender distribution it would be possible to estimate the number of deaths that would be expected to occur had this population not been at risk, based on mortality data for the general population. The difference between deaths observed across all ages and the deaths expected to have occurred in this matched 'normal' population would provide the number of 'excess' deaths by age and gender. ${ }^{41}$ The YLL associated with each of these excess deaths is the life expectancy conditional on gender and on surviving to the age at which the excess death occurred. The total YLL for the at risk population is simply the sum of these YLLs over all excess deaths, which could occur at any age. This YLL is equivalent to the area between the survival curve for the population at risk in a PBC and the counterfactual survival curve for the same population but not at risk from membership of the PBC. The difficultly is that routinely available data do not provide any information about the size of the population at risk or its age and gender distribution. All that is routinely available are observed deaths (by age and gender). Therefore, it is not possible to directly estimate excess deaths or compare survival curves.

Even if the size of the at risk population is unknown we can still use information that might be ayailable about its age and gender distribution (or make reasonable assumptions) to estimate a matched 'normal' LE using life tables for the general population - such a LE summarises the area under the counterfactual survival curve. Unfortunately, it is not possible to also calculate the LE for the population at risk in the PBC (or represent the survival curve) without information about the size of the at risk population - if it was possible the difference between these life expectancies would approximate the YLL per patient at risk in a PBC.

Fortunately, we can still recover a consistent estimate of YLL using observed deaths and a LE that represents the normal LE of a matched population that is not at risk. This requires all observed deaths both those that occur below and those that occur above this DE to be taken into account. Those deaths occurring below LE generate YLL - compared to the ayerage of a matched population not at risk. However, we must also account for those deaths that occur at ages above LE. These deaths generate life years 'gained' (YLG) compared to the average of a matched population not at risk. Therefore, the appropriate estimate is a net YLL (i.e., YLL - YLG). In effect, by subtracting YLG from YLL we take account of the fact that not all deaths below LE/are excess deaths but some deaths above LE are (see Appendix C for more formal explanation of the equivalence of these ways of calculating YLL). ${ }^{42}$

## Using the life expectancy of the general population

Routinely available data provides the age and gender of observed deaths but no information about the age and gender distribution of the at risk population itself. Using observed age and gender at death as an indication of the distribution of the at risk population will significantly overestimate the LE of a normal matched population insofar as a disease may be chronic (not all PBC mortality occurs on entry into the at risk population), and that PBC related mortality risk may increase with age (see Table C14 Appendix C). ${ }^{43}$
${ }^{41}$ These 'counterfactual' deaths will occur in the other PBCs insofar as all deaths are recorded in an ICD codes. Therefore, we take account of the unavoidable fact that everyone must die of something at some time. For example, even if all observed cancer mortality was avoidable and could in principle be eliminated with sufficient expenditure, lives would not be 'saved' but deaths delayed and reallocated to other causes. Note that the outcome elasticities are based on PBC mortality that is sensitive to changes in expenditure (i.e., is avoidable) at the margin so no assumptions about how much of the PBC mortality is avoidable is required.
$\int_{5}^{42}$ Simply taking the difference between a fixed LE and the age at death of deaths that occur below LE and ignoring those death that occur above LE, would only provide the correct figure if it is reasonable to assume that no deaths would have otherwise occurred prior to LE (so all 'normal' deaths must occur at LE) and that there are no deaths (survivors) beyond LE in the at risk population, i.e. all deaths below LE are excess deaths and there are no excess deaths above LE.
${ }^{43}$ If risk increases over the disease duration more deaths would be observed in groups that have been prevalent for some time (i.e., are older) than those that are incident. Also if PBC related mortality is higher for older age groups they will be overrepresented in observed deaths compared to a matched normal population. For both reasons LE, YLL and cost per life year would be overestimated using age at death as a proxy for the age distribution of the at risk population.

In the absence of additional external information the net YLL could be based on the life expectancy of the general population, reflecting its current age and gender distribution. These are reported in Table 4.3 and illustrate the impact of accounting for counterfactual deaths in the way described above. The YLL reported in column 5 of Table 4.3 are calculated the same way and are the same as the figures previously reported (column 5 of Table 4.2). That is, they do not account for deaths that would have otherwise occurred below LE or the very many deaths that occur above LE. With the exception of PBC18\&19 many death occur above the LE of the general population (see column 4 in Table 4.3) in all PBCs. As a consequence there are LYG associated with all other PBCs (see column 6) so the net YLL in column 7 are lower than YLL based on the same life expectancy. Therefore, failure to account for counterfactual deaths would lead to an overestimate of the YLL associated with a PBC and the effects of expenditure on YLL. Consequently the cost per life year threshold would be underestimated (see Table 4.6).

Table 4.3. Net YLL using life expectancy of the general population

| PBC |  |  |  | Average 2006-2008 |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | LE of Males [1] | LE of Females [2] | Deaths $<$ LE <br> [3] | Deaths $>$ LE <br> [4] | YLL <br> [5] | $\begin{aligned} & \text { YLG } \\ & \text { I6 } \end{aligned}$ | Net YLL |
| 1 | Infectious diseases | 80.7 | 84.4 | 3,710 | 3,248 | 62,052 | 18,796 | 43,256 |
| 2 | Cancer | 80.7 | 84.4 | 95,213 | 35,597 | 1,345,038 | 175,350 | 1,169,689 |
| 4 | Endocrine | 80.7 | 84.4 | 4,000 | 2,764 | 65,016 | 15,864 | 49,152 |
| 7 | Neurological | 80.7 | 84.4 | 8,975 | 6,378 | 145,529 | 34,621 | 110,908 |
| 10 | Circulatory | 80.7 | 84.4 | 82,099 | 77,752 | 916,192 | 444,694 | 471,498 |
| 11 | Respiratory | 80.7 | 84.4 | 30,500 | 34,945 | 310,334 | 215,829 | 94,505 |
| 13 | Gastro-intestinal | 80.7 | 84.4 | 15,827 | 8,320 | 273,308 | 45,295 | 228,012 |
| 17 | Genito-urinary | 80.7 | 84.4 | 4,198 | 6,427 | 39,099 | 40,530 | -1,431 |
| $18+19$ | Maternity \& neonates | 80.7 | 84.4 | 226 | 0 | 17,167 | 0 | 17,167 |

However, these figures are only correct insofar as the distribution of age and gender in each PBC is similar to the general population. For example, if the at risk population tends to be younger the correct LE for the PBC will be lower and the net YLL will atso tend to be lower. Similarly if the at risk population tends to be older than the general population the correct LE will be higher and net YLL will also tend to be higher. ${ }^{44}$ This explains the apparent net gain in YLL (negative net YLL) for PBC17 (Genito-urinary) where most deaths occurat ages greater than the LE of the general population so that LYG exceeds YLL. As we are able to show later (see Table 4.5) this is because the age distribution in this PBC tends to be older than the general population, i.e., the LE for a matched normal population should be higher with fewer deaths above and more below this LE.

## Using additional information about age and gender distribution

It is evident that estimates of KL L require some account to be taken of counterfactual deaths. In the absence of routinely a a ailable information this requires examination of alternative sources of information which might provide a basis for more credible assumptions about the age and gender distribution of the PBC population than either, the distribution of observed deaths or the general population. ${ }^{55}$ The WHO Global Burden of Disease (GBD) study, updated in 2008 using 2004 data (see Addendum 1 in Appendix C for more details $)^{46}$ provides a range of summary health indicators for the UK, which are, in part, based
${ }^{44}$ A higher (lower) LE will mean that there are more (less) deaths below LE, each generating more (fewer) YLL and fewer (more) deaths above LE each generating fewer (more) LYG.
${ }^{45}$ Although this research was not funded to purchase access to GPRD data we were able to examine a sample of it which comprised of $22,313,086$ rows/patient-ICD10 events (3 digit) representing 4,229,910 patients with data on new diagnosis of diseases observed between 1 Jan 2006 and 24 June 2011 (see Addendum 1 in Appendix C).
Although GPRD data could, in principle, provide this type of information the difficulties of reliability, face validity and interpretation of the sample data in the form available to us meant that it was not directly useful. We discuss the potential value of other sources of information, including GPRD in Chapter 5.
${ }^{46}$ We are aware that the 2000-2002 WHO GBD study and the update which was published in 2008 using 2004 data has itself recently been updated. However, the report and tools where not publically available at the time this research was conducted. We discuss the potential of future sources of information in Chapter 5.
on estimates of the incidence of sequelae associated with different types of disease by age and gender ${ }^{47}$. Therefore, the type of information used by WHO in the GBD Study to generate summary estimates for the UK can also be used to improve the assumptions required about the age and gender distribution of the PBC populations. Importantly, at this stage, we do not need to rely on estimates of the absolute size of the at risk population, but only the relative 'share' by age and gender.

GBD classifies diseases by U-codes, which are groups of three digit ICD-10 codes (see Addendum 1 in Appendix C for details of how U-codes map to ICD-10 codes). ${ }^{48}$ Since we know which ICD codes contribute to each PBC we can map information from U-codes to PBCs via the ICD codes that contribute to each. The resulting average age and life expectancy for each PBC is reported in columns 3 and 4 of Table 4.4 using the information available from GBD in combination with life tables for the general population.

Table 4.4. Average age and life expectancy for PBCs based on GBD

| PBC |  | Sex | Average age of general population [1] | LE of general population [2] | Average age in PBC (GBD) [3] | LE of at risk population (GBD) [4] |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | Infectious diseases | m | 38.5 | 80.7 | 28.6 | $\checkmark 79.6$ |
| 1 | Infectious diseases | f | 40.8 | 84.4 | 30.2 | 83.6 |
| 2 | Cancer | m | 38.5 | 80.7 | 61.3 | 83.0 |
| 2 | Cancer | f | 40.8 | 84.4 | 52.3 | 84.7 |
| 4 | Endocrine | m | 38.5 | 80.7 | 44.2 | 81.0 |
| 4 | Endocrine | f | 40.8 | 84.4 | 50.8 | 84.7 |
| 7 | Neurological | m | 38.5 | 80.7 | 24.8 | 79.6 |
| 7 | Neurological | f | 40.8 | 84.4 | 23.5 | 83.3 |
| 10 |  | m | 38.5 | 80.7 | 55.4 | 83.0 |
| 10 | Circulatory | f | 40.8 | 84.4 | 57.9 | 86.5 |
| 11 |  | m | 38.5 | 80.7 | 32.1 | 80.3 |
| 11 | Respiratory | f | 40.8 | 84.4 | 33.7 | 84.0 |
| 13 | Gastro-intestinal | m | 38.5 | 80.7 | 35.8 | 80.6 |
| 13 | Gastro-intestinal | f | 40.8 | 84.4 | 41.9 | 84.5 |
| 17 | Genito-urinary | m | ${ }^{38.5}$ | 80.7 | 63.2 | 83.5 |
| 17 | Genito-urinary | f | - 40.8 | 84.4 | 47.3 | 85.6 |
| $18+19$ | Maternity \& neonat |  | 38.5 | 80.7 | 3.0 | 78.7 |
|  |  |  | 40.8 | 84.4 | 24.1 | 83.1 |

These summary estimates suggest that some of the PBC populations may be on average be older than the general population (e.g., Cancer, Circulatory and Genito-urinary PBCs) or younger (e.g., Maternity \& neonates, Infectious diseases and Neurological). However, when trying to interpret these summaries it should be noted that the average age is the average over the ages at which sequelae occur across the ICDs contributing to the PBC , weighted by incidence of the sequelae at each age. Therefore, a similar average age can reflect very different age distributions. Some reflect a markedly bimodal distribution, e.g., Respiratory, where there is high incidence at very young and older ages, or very different age distributions across the type of diseases that contribute to the PBC. For example PBC7 (Neurological) includes dementia which accounts for the vast majority of the PBC population older than 70 . However, a greater proportion of the population is in much younger age groups with other conditions, especially migraine (see Addendum 1 Appendix C). When interpreting these summary estimates it should also be noted that the reported life expectancies are not the life expectancies at the average ages reported in column 3, but
${ }^{47} \mathrm{WHO}$, through the National Burden of Disease toolkit reports UK specific information about the incidence and duration of sequelae associated with different types of disease by age and gender. Since it is possible that a patient may experience more than one of the types of sequelae reported in GBD we use the gender and age distribution of the sequelae with the highest prevalence (evaluated as incidence x duration) to evaluate the age and gender distribution within each disease, i.e., the minimum estimate of prevalence consistent with these figures (see Section C2.1.3 and Addendum 1 in Appendix C).
${ }^{48}$ Throughout the analysis in Chapter 4 mortality, life years and QALY were not assigned to procedural ICD codes (Section C2.1.3 Appendix C) as these are likely to be evident in other ICD codes related to the procedure.
the average over the life expectancies for each age group within the contributing ICDs weighted by the age distribution of sequelae from GBD U-codes.

The implications for net YLL of using these PBC specific estimates of 'normal' life expectancy are reported in Table 4.5. As expected, the net YLL for those PBC with a LE greater than the general population are higher than those reported in column 5 in Table 4.3 (e.g., PBC10 Circulatory and PBC17 Genito-urinary, which now has positive net YLL). Similarly those PBCs with a LE less than the general population have lower net YLL than reported in column 5 in Table 4.3 (e.g., PBC1 Infectious diseases and PBC18 \& 19 Maternity \& neonates, where the effect of a lower LE is more modest as there are no deaths above either of the estimates of LE).

Table 4.5. Net YLL using life expectancy for each PBC

| PBC |  | LE of Males [1] | LE of Female <br> s <br> [2] | Average 2006-2008 |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | Deaths |  | YLL <br> [5] | $\begin{gathered} \text { YLG } \\ {[6]} \\ \hline \end{gathered}$ |  |
|  |  |  |  | $\begin{gathered} <\mathrm{LE} \\ {[3]} \\ \hline \end{gathered}$ | $\begin{gathered} >\text { LE } \\ {[4]} \\ \hline \end{gathered}$ |  |  |  |
| 1 | Infectious diseases | 79.6 | 83.6 | 3,498 | 3,460 | 58,686 | 21,724 | 36,962 |
| 2 | Cancer | 83.0 | 84.7 | 101,203 | 29,607 | 1,473,733 | 126,549 | 1,347,184 |
| 4 | Endocrine | 81.0 | 84.7 | 4,068 | 2,696 | 66,283 | 15,058 | 51,225 |
| 7 | Neurological | 79.6 | 83.3 | 8,370 | 6,983 | 135,686 | 41,770 | 93,917 |
| 10 | Circulatory | 83.0 | 86.5 | 96,694 | 63,157 | 1,102,020 | 278,251 | 823,768 |
| 11 | Respiratory | 80.3 | 84.0 | 29,549 | 35,897 | 298,343 | 230,313 | 68,030 |
| 13 | Gastro-intestinal | 80.6 | 84.5 | 15,824 | 8,323 | 273,117 | 45,414 | 227,703 |
| 17 | Genito-urinary | 83.5 | 85.6 | 4,969 | 5,655 | 47,229 | 29,101 | 18,127 |
| 18+19 | Maternity \& neonates | 78.7 | 83.1 | 226 | 0 | 16,801 | 0 | 16,801 |

The impact on the cost per life year threshold of the issues discussed in Sections 2.1, 2.2 and 2.3 are summarised in Table 4.6 (see Table C16 in Appendix C for detailed breakdown of changes in spend and YLLs across PBCs).

Table 4.6. Summary of cost per life year threshold

|  | Using cut-off in estimating YLL (ONS) cut-off of 75 cut-off of LE of the GP <br> [1] <br> [2] |  | Using net YLL estimates |  |
| :---: | :---: | :---: | :---: | :---: |
|  |  |  | Using LE of the GP <br> [3] | Using LE of the PBC population (GBD) <br> [4] |
| big 4 PBCs | $) ¢ 10,398$ | £,5,487 | £10,421 | £,8,080 |
| 11 PBCs (with mortality) | $\underbrace{2} 20,031$ | £10,660 | £19,928 | £15,628 |
| All 23 PBCs (zero health effects for remaining 12 PBCs ) | £73,697 | £39,218 | £73,317 | $\AA 57,497$ |
| All 23 PBCs (non-zero health effects for remaining 12 PBCs, except GMS)* | £22,639 | £12,048 | $£ 22,523$ | $£ 17,663$ |

* in PBCs without a mortality signal, health effects were estimated by valuing changes in expenditure at the same rate as observed in PBCs for which there was a mortality signal.

Using ONS data to calculate YLL in the same way as the published NHS IC figures, but overcoming some of the issues associated with the reporting of mortality at PCT level and the coverage of published estimates 0 f 1 LL (see Section 4.2.1), generates similar estimates of a cost per life year threshold (see column 1 Table 4.6) to those reported in Chapter 3. Calculating YLL in the same way, but based on the life expectancy of the general population significantly overestimates YLL for the reasons set out in Section 4.2.2 so underestimates the cost per life year threshold (see column 2). Taking account of counterfactual deaths by calculating net YLL based on the life expectancy of the general population (see column 3) provides similar estimates to those reported in Chapter 3. Assuming that PBC populations have the same age and gender distribution as the general population when the, albeit limited, information that is available suggests otherwise, seems inappropriate. Therefore, our preferred central estimate of the cost per life year threshold is reported in column 4. These are lower than those based on the general population, reflecting the impact on net YLL of evidence that the population at risk in some key PBCs (especially PBCs 2 and 10) tend to be older than the general population. In Section 4.2 .5 we consider extreme upper and lower bounds that might be placed on this central estimate.

### 4.2.4 Inferring excess deaths

We have been able to establish a measure of net YLL, which takes account of deaths that would have occurred anyway below a normal LE for the PBC population (i.e., not all deaths observed in a PBC are excess) and that some deaths observed above this LE would not otherwise have occurred at that age (i.e., some of these deaths are excess). As explained in Section 4.2.3, net YLL calculated in this way is equivalent to first establishing the number of excess deaths at each age, then calculating YLL for each excess death (based on the LE conditional on the age at which each excess death occurred) and then summing these YLL across all excess deaths (i.e., across all ages). In other words, the estimates of net YLL imply a number of excess deaths required to generate them in each PBC. Therefore, it is possibleto solve for the total number of excess deaths based on the net YLL and the average YLL per observed death. ${ }^{49}$ The net YLL divided by the average YLL per death provides the number of excess deaths required, which on average will generate the estimated net YLL. ${ }^{50}$

The implied excess deaths associated with net YLL based on the LE of the PBCs (see column 1 Table 4.5) are reported in Table 4.7. With the exception of PBC18\&19, excess deaths are some proportion of total observed deaths in each PBC. The proportion of excess deaths differs by PBC reflecting the distribution of deaths relative to the LE of the PBC. ${ }^{51}$ For example, in those PBCs where a large proportion of deaths occur below LE (see column 3 and 4) excess deaths tend to be greater proportion of total deaths (e.g., PBC2, 13 and 10). Where most deaths occur above LE excess deaths as a proportion of total deaths tend to be lower (e.g., PBC1, 11 and 17).

Table 4.7: Excess deaths implied by net YLL.

| PBC |  | Net YLL [1] | YLL per observed death [2] | Excess deaths [3] | Total deaths [4] | \% excess deaths [5] |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | Infectious diseases | 36,962 | 13.4 | 2,797 | 6958 | 40\% |
| 2 | Cancer | 1,347,184 | 14.1 | 95,715 | 130810 | 73\% |
| 4 | Endocrine | 51,225 | 13.7 | 3,769 | 6764 | 56\% |
| 7 | Neurological | 93,917 | 13.7 | 6,909 | 15353 | 45\% |
| 10 | Circulatory | 823,768 | 10.5 | 79,218 | 159851 | 50\% |
| 11 | Respiratory | 68,030 | 9.2 | 7,386 | 65445 | 11\% |
| 13 | Gastro-intestinal | 227,703 | 15.2 | 15,199 | 24147 | 63\% |
| 17 | Genito-urinary | 18,127 | 8.3 | 2,172 | 10625 | 20\% |
| 18+19 | Maternity \& neonates | 16,801 | 73.9 | 226 | 226 | 100\% |

Estimates of net YLL and changes in life years due to expenditure (see Table 4.5 and 4.6) have already accounted for the fact that notall deaths are excess and don't generate YLL. Nevertheless, solving for the number of implied excess deaths associated with these net YLL estimates allows a comparison of the cost per excess and observed PBC death avoided and an examination of the interpretation that can be placed of the life years expected to be gained from an excess or observed death averted. Since only deaths obseryed in the PBC can used to estimate the effects of expenditure (excess deaths are not directly observed since they rely on an unobserved counterfactual population and would occur outside the PBC), the outcome elasticities can be interpreted as the proportionate change in observed PBC mortality due to a proportionate change in PBC expenditure. Equally, however, they can also be interpreted as the

[^19]proportionate effect on excess death due to a proportionate change in expenditure so can be applied to either total observed or total excess deaths. ${ }^{52}$

The cost per excess death and the cost per PBC death averted are reported in Table 4.8 (see Table C19 in Appendix C for a detailed breakdown of changes in spend and excess or PBC deaths across PBCs). The cost per PBC death averted is, of course; significantly lower than the cost per excess death as excess deaths are only a proportion of total deaths (see Table 4.7). Also the cost per PBC death averted are substantially lower than those reported in Chapter 3 (see, Tables B8.22 and B8.23 in Appendix B), since these estimates do not restrict the effects of expenditure to PBC deaths under $75 .{ }^{53}$ The cost per PBC or excess death averted (or life saved) should not be over interpreted because they are of little direct policy interest since lives are never saved (death is only delayed) and the significance of a death averted depends critically on how long it is averted for (the life years gained - see Table 4.6) and the quality of life in which additional years are lived (see Section 4.3).

Table 4.8. Summary of the cost per death averted threshold

|  | Cost per excess death averted, $£$ [1] | Cost per PBC death averted, $£$ [2] |
| :---: | :---: | :---: |
| big 4 PBCs | £ 91,129 | £32,864 |
| 11 PBCs (with mortality) | $£ 177,692$ | £ 64,774 |
| All 23 PBCs (zero health effects for remaining 12 PBCs ) | £653,748 | £238,310 |
| All 23 PBCs (non-zero health effects for remaining 12 PBCs , except GMS)* | ¢,200,829 | £,73,208 |

* in PBCs without a mortality signal, health effects were estimated by valuing changes in expenditure at the same rate as observed in PBCs for which there was a mortality signal.

However, establishing the number of excess and PBC deaths averted which are associated with net YLL is useful because it enables an assessment of the number of life years gained associated with each death averted. On average across all 11 PBCs each excess death averted is associated with 11.4 life years gained. These are reported for each PBC in Table C21 in Appendix C and range from 74.3 years per excess death for PBC 18 \& 19 Maternity \& neonates to 8.3 for PBC17Genito-urinary. However, clinicians or the evaluative literature cannot distinguish whethey an observed death is excess or not. What can be observed is whether groups of similar patients twith and without access to a treatment survive and for how long. Therefore, it is the life years associated with each observed death that provides a context that can be interpreted based on experience and evidence of how effective those interventions that could be invested or disinvested tend to be. The ayerage life years expected to be gained associated with each observed PBC deaths averted takes account of that fact that some deaths that are avoided in the PBC are
${ }^{52}$ Observed PBC mortality that is sensitive to changes in expenditure can be regarded as 'avoidable' and it is only this mortality that contributes to the estimates of outcome elasticities (not all observed mortality is necessarily avoidable and sensitive to expenditure - such mortality will not contribute to the estimates). Not all observed mortality is excess when compared to the counterfactual population but this is unrelated to the question of how sensitive it is to expenditure, i.e., observed mortality will be just as sensitive to expenditure whether or not it is regarded as excess. Therefore, the estimated outcome elasticities can be applied to either observed PBC deaths or excess PBC deaths
${ }^{53}$ Recall from Chapter 3 and appendix B that the measure of mortality that is available at PCT level and used to estimate the outcome elasticities is restricted to deaths under 75, as are the published estimates of YLL associated with) them (see Section 4.2.2). However, to restrict effects only to those under 75 would imply that there is no excess mortality above 75 or equivalently that there are no health effects of PBC expenditure above 75 . Rather than assume no affects of NHS activity in older populations we apply the effects that can be observed to the whole PBC but account for deaths that would otherwise occurred in our estimate of net YLL in Section 4.2.3. In many respects whether or not PBC deaths at older ages are as sensitive to changes in expenditure is not critical since any observed deaths that might be averted at older ages are less likely to generate life years gained because they are more likely to have occurred anyway in that year (i.e., are excess so generate zero life years gained anyway). Therefore, they will have very limited impact on cost per life year or subsequently on cost per QALY estimates in Sections 4.3 and 4.4). For this, and the reasons given in the text, it is the cost per life year rather than cost per death averted, whether excess or observed, that is of primary interest.
not delayed for very long but quickly occur ${ }^{54}$ elsewhere and do not generate LY gained (i.e., they were not excess deaths). These are also reported for each PBC in Table C21 in Appendix C and range from 74.3 years per observed death for PBC $18 \& 19$ Maternity \& neonates ${ }^{55}$ to 1.0 for PBC11 Respiratory problems, i.e., the YLL per PBC death are much lower for those PBCs where a small proportion of observed deaths are excess. On average across all 11 PBCs each PBC death averted is associated with 4.1 life years gained.

### 4.2.5 Summary of cost per life year estimates

The sequence of analysis set out above has enabled an examination of the impact of the limitations associated with the incomplete reporting mortality data at PCT level and incomplete coverage of published YLL estimates. We have also been able to consider effects above 75 while taking account of that fact that many deaths would have occurred anyway, despite the limited information available about the population at risk within a PBC. The GBD Study does provide some information about the age and gender distribution of the population at risk in a PBC so offers some improvement over the other assumptions that would otherwise be required (i.e., that the distribution of age and gender is the same as the general population or follows the distribution of observed deaths). For this reason the cost per life year threshold in column 4 of Table 4.6 and repeated in lines 1 to 4 in Table 4.9 are regarded as the central or best estimates given the evidence available and the credibility of altennative assumption that could be made. As explained in Section 4.1, these are based on the conservative assumption that any health effects of changes in expenditure are restricted to one year, which, to some extent, may be offset by the more optimistic assumption any death averted returns the individual to the mortality risk face by the general population, matched for age and gender.

Table 4.9: Summary of the cost per life year threshold with upper and lower bounds

| Effect of expenditure on mortality: YLL per PBC death averted: | $\begin{gathered} \text { Best estimate } \\ 1 \text { year } \\ \sim 4.1 \mathrm{YLL} \end{gathered}$ |  |
| :---: | :---: | :---: |
| big 4 PBCs | £8,080 | [1] |
| 11 PBCs (with mortality) | £15,628 | [2] |
| All 23 PBCs (zero health effects for remaining 12 PBCs) | £57,497 | [3] |
| All 23 PBCs (non-zero health effects for remaining 12 PBCs, except GMS)* | £,17,663 | [4] |
| Effect of expenditure on mortality: YLL per PBC death averted: | Lower bound Remainder of disease $\sim 4.1$ YLL ${ }^{* *}$ |  |
| big 4 PBCs | £3,846 | [5] |
| 11 PBCs (with mortality) | £6,106 | [6] |
| All 23 PBCs (zero health effects for remaining 12 PBCs ) | $£ 22,463$ | [7] |
| All 23 PBCs (non-zero health effects for remaining 12 PBCs, except GMS)* | £6,901 | [8] |
| $\cdots$ | Upper bound |  |
| Effect of expenditure on mortality: | 1 year <br> 2 YL |  |
| big 4 PBCs | £16,432 | [9] |
| 11 PBCs (with mortality) | £32,387 | [10] |
| All 23 PBCs (zerohealth effects for remaining 12 PBCs ) | £,119,155 | [11] |
| All 23 PBCs (non-zero health effects for remaining 12 PBCs, except GMS)* | $£ 36,604$ | [12] |

* in PBCs without a mortality signal, health effects were estimated by valuing changes in expenditure at the same rate as observed in PBCs for which there was a mortality signal. ** see Tables C14, C15 and C18 in Appendix C

It does not seem credible to imagine that NHS expenditure has no health effects in the 12 PBC which do not have sufficient mortality reported at PCT level to estimate outcome elasticities - what is implied by
$\qquad$

[^20]the estimate reported in line 3. Therefore, it is the estimates reported in lines 2 and 4 that are of policy interest. The estimate of $£ 15,628$ per life year (line 2 ) is restricted to the effects of changes in expenditure in the 11 PBCs where outcome elasticities can be estimated. The threshold of $£ 17,663$ per life year uses the estimated health effects of expenditure in these PBC as a surrogate for health effects in the others, i.e., assuming that the effects that can be observed will be similar to those that cannot. However, no health effects are assigned to PBC23 (General Medical Services) on the basis that any health effects of this expenditure would be recorded in the other PBCs. ${ }^{56}$

The extreme upper and lower bounds for the cost per life year thresholds in Table 4.9 are based on making the necessary assumptions about duration of health effects and how long a death might be averted optimistic (providing the lower bound for the threshold) or conservative (an upper bound for the threshold). The lower bound (lines 5 to 8 ) is based on assuming that health effects are not restricted to one year but apply to the whole of the remaining disease duration of the population at risk in PBCs during the expenditure year. ${ }^{57}$ Although this combines optimistic assumptions, it is possible, indeed likely, that at least some expenditure may have effects on the health outcomes of future patients that are not currently part of the population at risk in a PBC, e.g., investments or disinvestment in prevention will have an impact on populations that are incident to PBCs in the future. Such effects are not captured in any of the estimates presented in this chapter so all are conservative with respect to this type of health effects from changes in expenditure.

The upper bound (lines 9 to 12) is based on the combination of assuming that health effects are restricted to one year for the population currently at risk and that any death averted is only averted for the minimum duration consistent with the mortality data. The econometrics work used the average of 3 years of mortality (2006 to 2008), so the estimated outcome elasticities are based on differences in mortality that remain after averaging over three years. Therefore the estimated effects are based on differences in observed PBC deaths that must have been sustained, on average, for more than a minimum of 2 years. ${ }^{58}$

### 4.3 Adjusting life years for quality of life

The central or best estimates of the cost perlife year threshold, which were presented in Table 4.9 (lines 2 and 4) take no account of the health related quality of life in which years of life, expected to be gained or lost through changes in expenditure, are likely to be lived. Even if attention is restricted to the direct health consequences of changes in mortality, estimates of the cost per life year will tend to overestimate the effects of changes in expenditure (underestimate the threshold) compared to a more complete measure of health that accounts for the quality in which of the years of life are expected to be lived. In this Section we examine the ways in which the life years reported in Section 4.2 can be adjusted for quality, taking account of information that is available about: i) how quality of life differs by age and gender (see Section 4.3.1), and ii) how the quality of life years associated with mortality changes might be effected by the types of diseases that make up each PBC (see Section 4.3.2). Throughout we continue to

${ }^{56}$ It would be inappropriate to assign all the change in GMS expenditure to the estimate of cost per life year based only on the 17) PBCs with outcome elasticities because it would imply that GMS only contributes to these PBCs. Restricting attention to the 11 PBCs with outcome elasticities but allocating part of the change in GMS expenditure to thembased on their proportional share of changes in overall expenditure would yield the same cost per life year as reported in line 4.
${ }^{57}$ Estimates of the duration of disease for each U-code are available from the GBD Study (see Table C22 and Addendum 1 in Appendix C). This information is also used in Sections 4.4.
${ }^{58}$ Variation in mortality the first year of data will only contribute to these estimates if differences are sustained for a minimum of 3 years. Similarly variation in mortality in the second (third) year will only contribute if it is sustained for a minimum of 2 (1) years. If differences in mortality are similar each year (contribute equally to the estimates) then estimated effects must have been sustained on average for a minimum of 2 years. Indeed, since some of the variation in mortality in $1^{\text {st }}$ year that is not sustained to the $3^{\text {rd }}$ year will nevertheless be sustained for 1 or 2 years, 2 life years per death averted represents somewhat less than the minimum consistent with restricting live years gain to the observed mortality data. Of course, this is minimum difference in observed rather than unobserved counterfactual excess deaths. Nonetheless it can be interpreted as an upper bound given the data available and therefore the analysis that has been feasible.
take account for counterfactual deaths in the way described in Section 4.2 .3 by making the adjustment for quality to the life years associated with every observed death before calculating a quality adjusted net YLL. The implications for a cost per quality adjusted life year (QALY) threshold that only accounts for the health effects of mortality changes are presented in Section 4.3.3. In Section 4.4 we explore the ways in which the likely direct effects of expenditure on quality of life (other than through mortality) might also be taken into account.

### 4.3.1 Quality of life based on the general population

The most commonly used metric of health related quality of life in the UK is EQ5D,[91] which is specified in the NICE reference case for methods of technology appraisal.[1] This metric has 5 dimensions of quality each with three possible levels. Each of these 243 possible health states is valued relative to a score of one, which represents full or best imaginable health (the best score across all 5 dimensions), and a score of zero, which represents death, based on a representative sample of the UK population.[92] Therefore, insofar as the years of life expected gained or lost through changes in) expenditure would be lived in this state of full health the cost per life year thresholds reported in Table 4.9 would also be the cost per QALY thresholds, albeit ones that only account for the health effects of mortality changes. However, unsurprisingly, there is good evidence that, on average, the general population is not in this state of full health. Therefore, the quality of life scoreassociated with the health states experienced by the general population are less than 1, decline with age and differ by gender. These quality of life 'norms' for the general population by age and gender are illustrated in Figure 4.1 based on an analysis of data from the Health Survey for England (HSE). ${ }^{59}$


Figure 4.1: Quality of life for the general population by age and gender
These quality of lifenorms can be applied to the YLL associated with all observed deaths in each PBC, taking account of gender and age at death. The results are reported in column 4 to 6 of Table 4.10. Recall from Section 4.2.3 that taking account of counterfactual deaths requires calculation of the YLL associated with deaths below LE (of a normal population matched to the age and gender distribution in the PB6) and the implied YLG of deaths that occur above this LE. There are two effects of adjusting life years for quality: i) since quality of life norms are always less than 1 the adjusted YLL and YLG are always lower than the unadjusted values in columns 1 and 2 (previously reported in Table 4.5); and ii) deaths aboye LE are necessarily at older ages with poorer quality of life norms than those below, so the difference between adjusted and unadjusted values is greater for YLG than YLL. The overall effect of quality adjustment on net YLL is the balance of these two effects. The overall effect of quality adjustment is to reduce the net YLL (compare Colum 6 and 3). ${ }^{60}$

[^21]Table 4.10: Net YLL adjust for the quality of life 'norms'

| PBC |  | Unadjusted life years |  |  | Quality adjusted life years |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | $\begin{array}{r} \text { YLL } \\ {[1]} \\ \hline \end{array}$ | $\begin{gathered} \text { YLG } \\ {[2]} \\ \hline \end{gathered}$ | $\begin{gathered} \text { Net YLL } \\ {[3]} \\ \hline \end{gathered}$ | $\begin{gathered} \text { YLL } \\ {[4]} \\ \hline \end{gathered}$ | $\begin{array}{r} \text { YLG } \\ {[5]} \\ \hline \end{array}$ | $\begin{aligned} & \text { Net YLL } \\ & {[6]} \\ & \hline \end{aligned}$ |
| 1 | Infectious diseases | 58,686 | 21,724 | 36,962 | 47,481 | 14,618 | 32,864 |
| 2 | Cancer | 1,473,733 | 126,549 | 1,347,184 | 1,143,445 | 84,036 | 1,059,409 |
| 4 | Endocrine | 66,283 | 15,058 | 51,225 | 52,856 | 9,973 | 42,883 |
| 7 | Neurological | 135,686 | 41,770 | 93,917 | 109,349 | 28,262 | 81,087 |
| 10 | Circulatory | 1,102,020 | 278,251 | 823,768 | 848,046 | 183,330 | 664,717 |
| 11 | Respiratory | 298,343 | 230,313 | 68,030 | 231,578 | 154,743 | 76,835 |
| 13 | Gastro-intestinal | 273,117 | 45,414 | 227,703 | 216,256 | 30,277 | 185,979 |
| 17 | Genito-urinary | 47,229 | 29,101 | 18,127 | 35,929 | 18,947 | 16,982 |
| 18+19 | Maternity \& neonates | 16,801 | 0 | 16,801 | 14,568 | 0 | 14,568 |

The quality adjusted net YLL figures in column 6 suggests that the health effects of mortality are lower than when relying only on unadjusted life years in Section 4.2. Therefore, the health effects of changes in expenditure on this more complete measure of health are lower. The implications of these adjustments on a cost per QALY threshold that only accounts for the direct health effects of mortality are reported in Table 4.11. As expected the cost per QALY threshold based on adjusting the life years gained or lost (column 2) is higher than a threshold based on unadjusted life years (column 1 and breviously reported in Table 4.9).

Table 4.11: Summary of cost per QALY threshold based on population norms and mortality effects

|  | Cost per life year threshold <br> [1] | Cost per QALY threshold Population norms [2] |
| :---: | :---: | :---: |
| big 4 PBCs | £8,080 | $£^{\text {¢ }}$, 631 |
| 11 PBCs (with mortality) | £15,628 | £18,622 |
| All 23 PBCs* | 1,17,663 | £21,047 |

* in PBCs without a mortality signal, health effects were estimated by valuing changes in expenditure at the same rate as observed in PBCs for which there was a mortality signal except GyIS.


### 4.3.2 Adjusting age related quality of life for disease decrements

Adjusting life years for age and gender related quality of life norms assumes that any life year gained through a change in expenditure would be lived in a similar quality of life to the general population. It is possible however, that patients benefiting from reduced mortality may, nevertheless, continue to be effected by the type of diseases that make up each PBC and experience the quality of life associated with the original disease.

The Health Outcome Data Repository (HODaR)[93] provides over 30,000 observations of EQ-5D measures of quality of life by ICD code and the age and gender of the patients in the sample (see Addendum 1 Appendix C). Although this is a rich UK data set, there were a limited number of observations for some of the less common ICD codes. For this reason HODaR was supplemented with informationfrom the Medical Expenditure Panel Survey (MEPS)[94] which also provides EQ-5D by ICD and reports the average age of respondents (see Addendum 1 Appendix C). These data provided a means of estimating the quality of life associated with each ICD code at the average age of respondents in the pooled sample. ${ }^{61}$ The quality of life associated with each PBC can be expressed as an average of the quality of life associated with its component ICDs. ${ }^{62}$ The quality of life effects of being in each PBC can

[^22]then be expressed as a disease related decrement compared to the population norms at the same age (see Table C29 in Appendix C). This is illustrated for PBC1 (Infectious disease) in Figure 4.2, where the weighted average of quality of life scores across the component ICD codes was 0.667 , at an average age average age of 54 for male respondents. Since the quality of life norms for males age 54 is 0.859 this suggests a decrement associated with membership of PBC1 of 0.192 , which can then be applied to quality of life norms by age. ${ }^{63}$


Figure 4.2: Quality of life for males in PBC1 (Infectious disease) and the general population by age

Quality of life norms adjusted for disease related decrements can be applied to the YLL associated with observed deaths in each PBC, taking account of gender and age at death in the same way as Section 4.3.1. ${ }^{64}$ The results are reported in column 4 to 6 of Table 4.12. The overall effect of quality adjustment that also applies a disease related decrement is to reduce the net YLL to a greater extent than adjustment with population norms alone (compare column 6 in Table 4.12 and 4.10).

Table 4.12: Net YLL adjusted for disease and age related quality of life

| PBC |  | Unadjusted life years |  |  | Quality adjusted life years |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | YLL <br> [1] | $\begin{array}{r} \text { YLG } \\ {[2]} \end{array}$ | $\begin{gathered} \text { Net YLL } \\ {[3]} \\ \hline \end{gathered}$ | $\begin{gathered} \text { YLL } \\ {[4]} \\ \hline \end{gathered}$ | $\begin{array}{r} \text { YLG } \\ \text { [5] } \\ \hline \end{array}$ | $\begin{aligned} & \text { Net YLL } \\ & {[6]} \end{aligned}$ |
| 1 | Infectious diseases | 58,686 | 21,724 | 36,962 | 37,055 | 10,793 | 26,262 |
| 2 | Cancer | 1,473,733 | 126,549 | 1,347,184 | 955,690 | 67,930 | 887,760 |
| 4 | Endocrine | -66,283 | 15,058 | 51,225 | 43,394 | 7,844 | 35,550 |
| 7 | Neurological | 135,686 | 41,770 | 93,917 | 68,893 | 15,842 | 53,050 |
| 10 | Circulatory | 1,102,020 | 278,251 | 823,768 | 656,145 | 135,241 | 520,905 |
| 11 | Respiratory | 298,343 | 230,313 | 68,030 | 169,269 | 106,505 | 62,764 |
| 13 | Gastro-intestinal | 273,117 | 45,414 | 227,703 | 163,593 | 21,677 | 141,916 |
| 17 | Genito-urinary | 47,229 | 29,101 | 18,127 | 29,749 | 15,152 | 14,598 |
| 18+19 | Maternity \& neonates | 16,801 | 0 | 16,801 | 13,662 | 0 | 13,662 |


codesp within a PBC and the information about which ICDs are more likely to contribute to the effects of changes in PBC expenditure are explored.
${ }^{63}$ In principle it would be possible to estimate disease related disutility by age rather than assume a fixed decrement. HODaR does provide age for each reported quality of life score but MEPs only provides average age of respondents in published summaries. However, even with access to 'raw' scores and the age and gender of each, it is very unlikely that there would be sufficient data to estimate age related decrements in each of the component ICDs. It would, however, be possible to assume a proportionate rather than fixed decrement by age. Since the average age of respondents in the pooled HODaR and MEPs sample tends to be older than the age distribution of the PBC populations (see Table C29 and C13 in Appendix C) this would tend to increase the quality adjusted net YLL and reduce the cost per QALY threshold compared to the fixed decrement applied here.
${ }^{64}$ The quality of life score was applied to each observed death considering the age at which each life year was gained or lost (from ONS) the ‘PBC decrements’ from HODaR and MEPS.

It should be noted that combining quality of life adjustments for both population norms and disease related decrements assumes that any life years gained due to a reduction in mortality will be lived in the diseased state until life expectancy, i.e. that all diseases are not just chronic but disease duration is lifelong. Inevitably this assumption means that the health effects of changes in mortality will be reduced. Consequently the cost per QALY threshold reported in Table 4.13 (column 2) will be higher than adjusting life years gained for population norms in Table 4.11.

Table 4.13: Summary of cost per QALY threshold based on disease related decrements

|  | Cost per life year threshold <br> $[1]$ | Cost per QALY gained <br> Disease related decrements <br> $[2]$ |
| :--- | :---: | :---: |
| big 4 PBCs | $£ 8,080$ | $£_{12,109}^{23,395}$ |
| 11 PBCs (with mortality) | $£ 15,628$ | $£_{2}^{26,441}$ |

* in PBCs without a mortality signal, health effects were estimated by valuing changes in expenditure at the same rate as observed in PBCs for which there was a mortality signal except GMS.


### 4.3.3 Summary of the cost per QALY threshold based only on mortality effects

The analysis to this point is summarised in Table 4.14. The three estimates of a cost per QALY threshold are based on assuming that each life year gained is either: lived in full health (see column 1, equal to the cost per life year estimates in Table 4.9), lived in a quality of life that reflects age and gender norms of the general population (column 2); or lived in a quality of life that reflects the original disease state (column $3)$.

Assuming that life years gained are lived in full health is not credible and should be regarded as an underestimate of the threshold, given what is known about quatity of life norms for the general population (see Figure 4.1). Equally, assuming that all hife years gained are lived in the quality of life of the original disease state does not seem credible either and is likely to overestimate the threshold since it assumes that all disease is not only chronic but lifelong and all life years would be lived in the diseased state until death. ${ }^{65}$ Therefore, adjusting life years gained for the quality of life of the general population taking account of age and gender (in column 2) is regarded as the best estimate of a cost per QALY threshold, which only reflects the health effects of changes in mortality. ${ }^{66}$ The lower and upper bounds are based on combining optimistic and pessimistic assumptions about the duration of health effects and how long a death might be averted as described in Section 4.2.5.

However, it should be noted that these cost per QALY thresholds only account for the direct health effects of changes in mortality due to changes in expenditure. Insofar as much, or at least some, of NHS activity and expenditure is intended to improve quality of life, not just mortality, then these estimates will underestimate total health effects and overestimate a cost per QALY threshold based on a more complete measure of possible health effects. In Section 4.4 we explore the ways in which the likely effects of expenditure on quality of life (other than through mortality) might also be taken into account.
${ }^{65}$ The information that is available about disease duration suggests that many types of disease that comprise the PBCs are not chronic and certainty not lifelong (see Table C22 in Appendix C). In Section 4.4 we take account of quality of life experienced while alive in the diseased state.
${ }^{66}$ In section 4.4.2 measures of QALY burden are used as the basis of estimating the health effects of changes in expenditure. This analysis assigns disease decrements to any life years gained during the duration of disease, i.e., insofar as more patients survive as a consequence of increased expenditure they do so in the diseased state for the remaining duration of the disease. Quality of life norms are assigned for any life years lived beyond this duration. Although this is conservative with respect to health effects, we regard it as the best estimate of quality of life effect of changes in mortality.

Table 4.14: Summary of QALY threshold estimates based only on mortality effects

|  | $\begin{gathered} {[1]} \\ (Q o L \text { score =1) } \end{gathered}$ | [2] <br> (QoL norm) | [3] (QoL diseased) |  |
| :---: | :---: | :---: | :---: | :---: |
| Effect of expenditure on mortality: <br> YLL per death averted: QALYs per death averted <br> big 4 PBCs <br> 11 PBCs (with mortality) <br> All 23 PBCs* | Best estimate |  |  |  |
|  | 1 year | 1 year | 1 year |  |
|  | $\sim 4.1$ YLL ** | $\sim 4.1$ YLL ** | $\sim 4.1$ YLL ** |  |
|  | $\sim 4.12 \mathrm{ALYs}$ | $\sim 3.5 \mathrm{QALYs}$ | $\sim 2.8 Q A L Y s$ |  |
|  | £,8,080 | £, 0,631 | £12,109 | [1] |
|  | £15,628 | £18,622 | £23,395 | [2] |
|  | £ 17,663 | $\underbrace{〔} 21,047$ | $£ 26,441$ | [3] |
| Effect of expenditure on mortality: YLL per PBC death averted: QALYsper death averted |  Lower bound  <br> Remainder of disease Remainder of disease Remainder of disease <br> $\sim 4.1 \mathrm{YLL} * *$ $\sim 4.1 \mathrm{YLL} * *$ $\sim 4.1 \mathrm{YLL}(* *$ <br> $\sim 4.12 A L Y s$ $\sim 3.5 Q A L Y s$ $\sim 2.8 Q A L Y s$ |  |  |  |
|  |  |  |  |  |
|  |  |  |  |  |
|  |  |  |  |  |
| big 4 PBCs <br> 11 PBCs (with mortality) <br> All 23 PBCs* | £3,846 | £4,252 | £,5,319 | [4]$[5]$$[6]$ |
|  | £6,106 | £6,852 | ¢8,568 |  |
|  | £6,901 | £7,744 | +2,683 |  |
| Effect of expenditure on mortality: YLL per PBC death averted: QALYs per death averted |  Upper bound <br> 1 year 1 year <br> 2 YLL 2 YLL <br> $2 Q A L Y s$ $\sim 1.9 Q A L Y s$ |  |  |  |
|  |  |  |  |  |  |
|  |  |  |  |  |  |
|  |  |  |  |  |  |
| big 4 PBCs | £16,432 | £,17,456 | £21,747 | [7] |
| 11 PBCs (with mortality) | £ 32,387 | 634,492 | £ 42,967 | [8] |
| All 23 PBCs* | £36,604 | 638,983 | £.48,561 | [9] |

* in PBCs without a mortality signal, health effects were estimated by valuing changes in expenditure at the same rate as observed in PBCs for which there was a mortality signal except GMS.
** see Tables C14, C15 and C18 in Appendix C


### 4.4. Including quality of life effects during disease

The cost per QALY thresholds presented in Section 4.3 only account for the health (QALY) effects of changes in mortality due to changes in expenditure. It does not seem credible to suppose that all NHS activity and expenditure only influences mortality with no effect on the quality of life while alive and experiencing a disease. Insofar as changes in NHS expenditure will also affect quality of life as well as mortality then total health effects will be underestimated and the thresholds presented in Table 4.14 will overestimate the cost per QALY threshold compared to a more complete picture of the likely effects of changes in NHS expenditure. In this section we explore ways to also take account of those effects on health not directly associated with mortality and life year affects (i.e., the 'pure' quality of life effects) to estimate an overall cost per QALY threshold.

The routine reporting of quality of life outcomes are increasingly available at PCT level (see Addendum 1 in Appendix C for a description of these data). In principle, the variation in such measures of outcome across PCTs could be used to estimate outcome elasticities for quality of life rather than mortality effects using similar econometric methods to those described in Chapter 3 (see Section B8.8 in Appendix B for the results of exploratory econometric analysis of these data). However, the currently limited coverage of routine reporting of these outcomes means that it is not feasible to estimate quality of life effects across all the PBCs using these data. In Chapter 5 we discuss how these data might be used to improve estimates of the threshold as the coverage and routine reporting of quality of life outcomes improves and how the analysis presented in Section 4.5 might help prioritise reporting in particular areas (i.e., those PBCs and ICD codes that have the greatest influence on estimates of the threshold).

Therefore, in this section we explore how estimates of effects of expenditure that can be observed (i.e., on mortality) can be used to infer the likely effects on what cannot be directly observed (quality of life), rather than making extreme assumptions that are not credible (e.g., assuming that changes in expenditure will have no effects on quality of life outcomes).

In Section 4.4.1 we use three alternative estimates of the ratio of QALYs to life years lost due to different types of disease as a means of inferring the change in QALYs that is likely to be associated with the estimated change in YLL, i.e., essentially applying the estimated proportionate effect on life years to total

QALYs. This is consistent with regarding the estimates of the mortality and life year effects as a surrogate for a more complete measure of the health effects of a change in expenditure.

However, the ratios of QALYs lost to life years lost due to disease in those PBC where outcome elasticities could not be estimated cannot inform estimates of the threshold (there are no estimated life year effects with which to apply the ratios). Nonetheless, the sources of information on which ratios are based also provides much of the information required to calculate the QALY burden of disease in these areas, which can be used to inform estimates of the threshold. Therefore, in section 4.4 .2 we use estimates of the QALY burden of disease, infer a proportionate effect on burden from the estimated effects on life years, and then apply this proportionate effect to the measures of QALY burden for all the other PBCs. In this way we can use all the information available about the mortality and quality of life effects of the different types of disease that make up each PBC, including those where mortality based outcome elasticities are not available.

### 4.4.1 Using ratios of QALYs to YLL

The ratio of the total QALYs to years of life lost (YLL) due to a disease indicates the number of QALYs associated with each YLL. Therefore, any change in YLL is likely to generate a number of QALYs indicated by the ratio - if it is reasonable to interpret the estimated effects on mortality and life years as a surrogate for a more complete measure of total health effects. For example, a disease with a ratio greater than 1 suggests that each YLL across the at risk population is associated with more than one QALY, i.e., where there are significant quality of life effects while experiencing the disease. ${ }^{67}$ Therefore, a change in expenditure that leads to 1 life year gained in this type of disease maybe expected to generate more than one QALY and a greater QALY effect than the same life year effects in a disease where this ratio is less than 1, i.e., where most of the effect of disease is on mortality rather than quality of life. Therefore, information which allows these ratios to be estimated for the diseases that make up each PBC provides a means of accounting for the likely effect on quality of life other than through effects on mortality.

To understand the differences between the three ratios presented below it is useful to regard the total QALY lost to YLL ratio ( R ) for a particular disease as the sum of two ratios: i) the QALYs lost due to premature death to YLL ratio $\left(\mathrm{R}_{\text {death }}\right)^{68}$ and ii) the $\widehat{Q} A L Y s$ lost during disease (while alive) to YLL ratio ( $\mathrm{R}_{\text {alive }}$ ) (see Section C2.3.1 in Appendix © for more detailed explanation).

## DALY to YLL ratios

The WHO GBD study provides UK specific estimates of the years of life lived with disability and the years of life lost due to different types of disease (classified by U-codes that can be mapped to ICD-10, see Section 4.2 and Addendum 1 in Appendix C). GBD uses Disability Adjusted Life Years (DALYs) as a measure of the burden of disease. This DALY measure has two components: i) the years of life lived with disability (YLD), which incorporates weights (between zero and one) to reflect the scale of disability experienced each year and the number of years lived with disability over the durations of disease; and ii) the years of (ife lost (YLL). The total DALY associated with a disease is simply YLL+YLD. Therefore, the DAL§ to YLL ratio is (YLL+YLD)/YLL or equivalently YLL/YLL + YLD/YLL. Since the first term (YLL YLL $\left.=\mathrm{R}_{\text {death }}\right)$ must equal one and the second $\left(\mathrm{R}_{\text {alive }}=\mathrm{YLD} / \mathrm{YLL}\right)$ must be $\geq 0$, a ratio based on DALYs must necessarily be bounded by below by one. This is illustrated in Table 4.15a for four different types of diseases (U-codes) which reflect diseases where mortality is the major component (e.g., U016) and where the impact of disease on the quality of life while alive is the major component (e.g., U141).

[^23]Table 4.15a: Examples of DALY to YLL ratios

| Ucode | DALY <br> ratios | $\left(\mathbf{R}_{\text {death }}+\mathbf{R}_{\text {alive }}\right)$ |
| :--- | :---: | :--- |
| U037 (Other infectious diseases) | 1.23 | $(1+0.23)$ |
| U016 (Tetanus) | 1.00 | $(1+0)^{*}$ |
| U061 (Mouth and oropharynx cancers) | 1.05 | $(1+0.05)$ |
| U141 (Spina bifida) | 2.34 | $(1+1.34)^{* *}$ |

* Given the short disease duration, it is only mortality effects that contribute to the ratio
* Quality of life effects during disease contribute significantly to estimates of the ratio


## Adjusting DALYs for quality of life norms

The use of DALY ratios bounded below by one essentially assumes that YLL would have otherwise been lived in a state of full health. As was discussed in section 4.3 .1 this is not credible given information available about the quality of life in the general population (see Figure 4.1). It would lead to over estimating the QALYs associated with mortality and life year effects and underestimating the cost per QALY threshold. Therefore, it is important to adjust these DALY ratios for the quality of life norms by age and gender in the same way as described in Section 4.3.1. The effect of this adjustment ${ }^{60}$ is illustrated in Table 4.15b. Now those types of disease where mortality rather than quality of life with the disease is the major component can have ratios less than one. Indeed the first term of these ratios $\left(\mathrm{R}_{\text {death }}\right)$ is consistent with, and is implied by, the analysis in Section 4.3.1 where the ratio of quality adjusted net YLLs to unadjusted net YLLs represents this ratio on average for each PBC.

Table 4.15b: Examples of modified DALY to YLL ratios

| Ucode | Modified <br> DALY ratios | $\left(\mathbf{R}_{\text {death }}+\mathbf{R}_{\text {alive }}\right)$ |
| :--- | :---: | :--- |
| U037 (Other infectious diseases) | 1.01 | $(0.7+0.23)$ |
| U016 (Tetanus) | 0.78 | $(0.78+0$ |
| U061 (Mouth and oropharynx cancers) | 0.83 | $(0.78+0.05)$ |
| U141 (Spina bifida) | 2.18 | $(0.85+1.34)$ |

Using quality of life estimates (based on HODAR (and MEPS)
The disability weights used in the DALY measure (and in $\mathrm{R}_{\text {alive }}$ ) are not based on the same description of health states as the EQ5D measure, nor are the weights based on a representative sample of the UK population responding to choice based elicitation questions. EQ5D based quality of life decrements (adjustments to age related quality of life norms) associated with different types of disease can be estimated from HODaR and MEPS data (previously described in Section 4.3.2). ${ }^{70}$ These disease related quality of life decrements be can be expressed for each U-code so can be used to replace the DALY disability weights in $\mathrm{R}_{\text {alive }}$ reported in Tables 4.14 a and $4.14 \mathrm{~b} .{ }^{71}$ This final adjustment is illustrated in Table 4.1.4c and turns, what were originally, DALY ratios into EQ5D QALY ratios. ${ }^{72}$ For these reasons we

[^24]regard the QALY to YLL ratios rather than DALY or modified DALY ratios as the preferred basis of estimating a cost per QALY threshold that provides a more complete picture of the likely health effects of changes in expenditure.

Table 4.15c: Examples of QALY to YLL ratios (HODaR and MEPS)

| Ucode | QALY ratios <br> (HoDAR and <br> MEPs $)$ | $\left(\mathbf{R}_{\text {death }}+\mathbf{R}_{\text {alive }}\right)$ |
| :--- | :---: | :--- |
| U037 (Other infectious diseases) | 1.37 | $(0.78+0.60)$ |
| U016 (Tetanus) | 0.78 | $(0.78+0$ |
| U061 (Mouth and oropharynx cancers) | 0.80 | $(0.78+0.02)$ |
| U141 (Spina bifida) | 1.88 | $(0.85+1.03)$ |

## Allocating effects at PBC level to ICD codes

Table 4.15 illustrate how QALY ratios can be calculated for and differ by U-code and therefore the ICD codes that make them up. ${ }^{73}$ Unsurprisingly, these ratios differ across the type of diseases that make up each PBC (see Table C45 in Appendix C). Therefore, when using this information to estimate a cost per QALY threshold the mortality and life year effects observed at PBC level must be allocated in some way to the component ICD codes before ratios are applied to LY effects and the resulting QALY effects are summed across all the contributing ICD codes. ${ }^{74}$

For this reason it is important to consider how other information might inform the different ways in which the effects observed at PBC level might be generated by the distribution of impacts at ICD level, i.e., where investment or disinvestment is likely to occur within the PBC and therefore which ICDs are likely to contribute most to overall health effects.

An important and complementary element to the econometric analysis of routinely reported information at PBC level was to investigate whether other information, commonly available at a local level within the NHS, might provide a useful indication of where, within a PBC, investment or disinvestment is more likely across the NHS. The details of this investigation and the rather disappointing results for the purposes of this analysis are reported in Addendurn 2 in Appendix C. In the absence of useful information at a local level it is possible to assume that a change in PBC expenditure will be allocated equally (on a per patient basis) acros $\$$ the component ICD codes, i.e., any investment or disinvestment is equally likely (occurs at random) across the population at risk within the PBC. However, information is available that gives some indication of which areas are more likely to have been subject to investment or disinvestment across PCTs. HES provides information about the costs associated with each ICD by PCT so it is possible to establish which ICDs contribute most to the variability in PBC costs across PCTs.


[^25]Those that contribute most to this variance may be expected to be more likely to have been subject to differential investment or disinvestment across PCTs. ${ }^{75}$

There are very marked differences in relative weight assigned to ICD based on the size of the population or its contribution to variance in costs (see Addendum 1 in Appendix C). One would expect investment or disinvestment within a PBC to focus on areas of marginal value rather than be allocated at random, therefore, the health effects of a change in PBC expenditure are likely to be overestimated and a cost per QALY threshold underestimated when allocating effects equally across the population at risk within each PBC. This is confirmed by the results of this analysis reported in Table C41 in Appendix C. For these reasons our preferred analysis uses contribution to variance to 'weight' the different ICD codes within a PBC (allocate the life year effects), before applying the QALY ratios associated with each ICD. This is also conservative, with respect to the health effects of changes in expenditure, compared to alternative assumptions that could be made about how PBC level effects might be allocated to ICD codes. The implications for a cost per QALY threshold that uses the estimated mortality and life year effects as a surrogate for a more compete measure of the likely heath effects (i.e., that includes quality of life as well as quality adjusted life year effects) is summarised in Table 4.16.

Table 4.16: Summary of the QALY threshold using QALY to YLL ratios

|  | DALY ratios [1] | Modified DALY ratios <br> [2] | $\qquad$ |
| :---: | :---: | :---: | :---: |
| big 4 PBCs | £,5,402 | £6,419 | £,5,990 |
| 11 PBCs (with mortality) | £ 9 , 958 | $£ 11,718$ | £10,297 |
| All 23 PBCs | £,11,254 | £,13,244 | £,11,638* |

* Preferred analysis

The QALY to YLL ratio implied by this analysis for all 11 PBC with outcome elasticities is 1.52 , which suggests that every (unadjusted) life year is associated with 1.52 QALYs on average across these PBCs. However, this implied QALY ratio differs across these PBCs, ranging from 0.79 in PBC2 to 15.05 in PBC18+19 (see Table C43 in Appendix C). Since all the analysis in this Section seeks to use the estimated mortality and life year effects as a surrogate for a more complete measure of likely health effects, it is the cost per QALY threshold for all 23 PBCs that is most relevant. As expected this threshold ( $£ 11,638$ ), is lower than a cost per QALY threshold based only the quality adjusted life year effects ( $£ 21,047$ in Table 4.14 that assumes no effects of NHS expenditure on quality of life itself). This difference gives some indication of the relative importance of QALY effects due to avoidance of premature death and the QALY effects of avoiding disability during disease.

Table 4.17 reports how the estimated QALY effects for each PBC can be decomposed into that part associated with quality adjusted life year effects and that part associated with 'pure' quality of life effects. These results appear credible for the first 11PBCs, where those for which mortality is the major concern have a much greater share of total QALY effects associated with avoidance of premature death (e.g., PBC2 and PBC(10) compared to those where quality of life is the major concern (e.g., PBC 7). ${ }^{76}$
Table 4.17: Decomposing estimated QALY effects by PBC

|  | QALY <br> change <br> (total) | QALY <br> change <br> (death) | due to <br> avoidance of <br> premature death | due to avoidance <br> of disability <br> while alive |
| :--- | :---: | :---: | :---: | :---: |

${ }^{75}$ Although the costs from HES data are only a component of total PBC costs they are an important one.
Unfortunately total PBC costs are not available at ICD level across PCTs so could not be used for this purpose. However, the assumption is not that HES cost are representative of total PBC costs but that those ICDs that contribute most to variability in HES costs are also likely to contribute most to variability in total PBC costs as well. ${ }^{76}$ It should be noted that the implied QALY ratio of 1.52 for the 11 PBC with outcome elasticities is a ratio of QALYs to unadjusted YLL. The proportion of total QALY effects due to premature deaths for the same PBCs ( $50 \%$ in Table 4.17) also implies a ratio - equal to two. However, this is a ratio of total QALY effects to quality adjusted YLL. The difference between these two ratios is the denominator, i.e., quality adjusted YLL are lower than unadjusted YLL.


Recall that the ratios of QALYs to YLL due to disease in those PBC where outcome elasticities could not be estimated cannot be used to inform estimates of the threshold because there are no estimated life year effects with which to apply the ratios. Therefore, as in previous sections, the estimated effect of expenditure on health for the 11 PBCs with outcome elasticities is applied to the estimated changes in PBC expenditure for the other 12 PBCs (excluding GMS for the reasons given in Section 4.2), i.e., assuming that the health effects that can be observed of a change in expenditure will be similar to those that cannot. However, the use of QALY ratios also implies that the share of total health effects between quality adjusted life year effects and that part associated with 'pure' quality of life effects are also similar to those PBC with estimated outcome elasticities. Summing the different types of health effects across these 11 PBCs suggests that $50 \%$ is due to avoidance of premature death and $50 \%$ due to avoidance of disability. This is clearly not credible when applied to the other PBCs, e.g., mental health, vision and hearing are likely have a much greater share of total health effects associated with quality of life effects and very little associated with premature mortality.

The problem is that using QALY to YLL ratios means that much of the information that is available about the other 12 PBCs cannot be used to inform the estimates of the cost per QALY threshold. Fortunately, the sources of information on which ratios are based also provide much of the information required to calculate the QALY burden of disease in these areas. Section 4.4.2 explores how measures of burden can be used to estimate a cost per QALY threshold that captures the likely effects of a change in expenditure on all aspects of health while using all the information that is available about all the PBCs.

### 4.4.2 Using estimates of the QALY burden of disease

In this Section we use estimates of the QALY burden of disease, infer a proportionate effect on burden from the estimated effects on life years, and then apply this proportionate effect to the measures of QALYburden for all PBCs. In this way we can use all the information available about the mortality and quality of life effects of the different types of disease that make up each PBC, particularly those where mortality based outcome elasticities are not available. ${ }^{77}$

[^26]The total QALY burden of disease for the population with disease in a particular year includes: i) the quality adjusted years of life lost due to all the disease related mortality that could occur in this population over their remaining duration of disease and ii) the reduction in quality of life while alive also for their remaining disease duration. However, applying the estimated proportionate effects on mortality and life years to such a measure of total burden would provide an estimate of the effects of a change in expenditure, not just in one year, but in all the remaining years of disease for the population at risk in that year. Recall from Section 4.2 that we have adopted the conservative assumption that changes in expenditure will only have health effects in one year for the population with disease in that year. Therefore, it is not a measure of total burden that is required, but a measure of the QALY burden of disease during one year for the population with disease (prevalent and incident) in that year. The estimated outcome elasticities can then be appropriately applied to this measure of burden. ${ }^{78}$

The information from GBD used to derive QALY ratios in Section 4.4.1 includes information about the YLL and duration of disease for those incident to a U-code, i.e., the measure of QALY burden from the information included in the ratios is a measure of the total burden of the disease but only for the population that is incident (rather total population with disease) in one year. Assuming that incidence is stable over the disease duration this is also equivalent to the QALY burden of disease during one year for the population with disease (i.e., those that are incident and prevalent) in that year. ${ }^{79}$

However, in moving from ratios to absolute measures of burden it becomes more important to examine and then adjust for any inconsistency between information about YLL and size of the incident population from GBD (which is available by U-codes and can be mapped to ICDs), and the information about net YLL and observed deaths for each PBC based on ONS data as described in Section 4.2.3 (see Table C44 in Appendix C). ${ }^{80}$
The implications for the cost per QALY threshold of using information about the QALY burden of disease for all PBCs rather than QALY ratios for those where an outcome elasticity can be estimated are reported in Table 4.18.
years lived beyond this duration. Although this is conservative with respect to health effects, we regard it as the best estimate of quality of life effect of changes in mortality.
The QALY effects of a change in PBC expenditure are a weighted average of the QALY effects within each of the ICDs that contribute to the PBC. The figures reported in this section are based on weighing the effects at ICD level by the proportion of the total PBC population within each contributing ICD code, rather than the contribution to variance in PBC expenditure, Estimates based on weighting ICD level effects by the contribution to variance are very similar and are reported in Section 2.3 of Appendix C.
${ }^{78}$ Of course it woutd be possible to solve for a lower outcome elasticity that could be applied to total burden which would return the required estimate of total QALY effects restricted to one year - see Section 2.1 in Appendix C
${ }^{79}$ So long as estimates of the quality of life decrement of disease from HODaR and MEPS are representative of average effects across those earlier (incident) and later (prevalent) in their disease duration an assumption of constant quality of life decrement with respect to disease duration is not required.
${ }^{80}$ There are a number of reasons for potential inconsistencies: i) GBD is based on earlier years of mortality data; ii) the imprecision of mapping from U-codes to PBC via ICD codes; and iii) the YLL reported in GBD are calculated in the same way as published NHS IC estimates (see Section 4.2.2 and 4.2.3) and will tend to overestimate the net YLE (see Table 52 in Appendix C). Therefore, the YLL by U-code, reported in GBD, that are mapped to ICDs are adjusted by these proportionate differences to ensure that the YLLs associated with all contributing ICD codes are consistent with (do not over estimate) the net YLL for the PBC as a whole. However, due to the earlier years of data and imprecision in mapping from U-codes to ICDs there might also be some inconsistency in estimates of the total incidence of disease for a PBC. Insofar as disease related mortality risk is stable, the same number of deaths should be observed in GBD and ONS data for the same at risk population. The PBC deaths recorded in GBD and those observed in ONS data (see Table 52 in Appendix C) are similar but nonetheless the proportionate difference is used to adjust the scale of quality of life burden while alive based on GBD information (equivalent to adjusting estimates of incidence). Notable exceptions are PBC1 and PBC18+19 where the discrepancies are due to imperfect mapping from U-code to PBC via ICD codes.

Table 4.18: Summary of the cost per QALY threshold
$\left.\begin{array}{|l|cc|}\hline & \begin{array}{c}\text { Cost per QALY gained* } \\ \text { QALY Burden } \\ \text { QALY ratios, }\end{array} \\ \text { (HoDAR and MEPs) } \\ \text { [1] }\end{array} \quad \begin{array}{c}\text { (HoDAR and MEPs) } \\ \text { [2] }\end{array}\right]$

* Preferred analysis

The cost per QALY threshold for the 11PBCs with outcome elasticities is lower using a measure of QALY burden $(£, 5,128)$ rather than the QALY ratios $(£ 10,297)$ described in Section 4.2.1. This is because GBD calculates YLL in the same way as published NHS IC figures so will tend to overestimate a net YLL which accounts for counter factual deaths (see Section 4.2.3). This will make little difference to the first term in the QALY ratio ( $\mathrm{R}_{\text {death }}$ ) used in Section 4.2.1 since an overestimate of YLL affects both denominator and numerator of the ratio. However, the second term $\left(\mathrm{R}_{\text {alive }}\right)$ is likely to be underestimated. Therefore the ratios used in section 4.4.1 will tend to underestimate the QALY effects of expenditure and overestimate the cost per QALY threshold (see Table 4.18). We are able to adjust the GBD based measure of QALY burden for this overestimation of net YLL in calculating the QAICY threshold reported in column 2). ${ }^{81}$

Since the purpose of this Section is to use the estimated mortality and life year effects as a surrogate for a more complete measure of likely health effects, it is the cost per QALY threshold for all 23 PBCs that is of most relevance. The cost per QALY threshold for all 23 PBCs is based on applying the proportionate effects on the QALY burden of disease, based on the observed effects of changes in expenditure on mortality in the 11 PBC with outcome elasticities, ${ }^{82}$ to the QALY burden of disease in the other PBCs. This generates a much higher cost per QALY threshold ( $\delta 15,701$ ) than one based on applying the estimated QALY effects of changes in expenditure, using QALY ratios for the 11 PBC with outcome elasticities, to changes in expenditure in the others ( 611,638 ). The reason is that the QALY burden of disease in the other PBC is, in general, lower than the QALY burden of disease across those PBCs where outcome elasticities can be estimated (see Table C45 in Appendix C). Therefore, applying the same proportionate effects to a lower QALY burden generates a smaller health effect of a change in expenditure. ${ }^{83}$ In essence the difference between these estimates is that in column 1 the absolute effect on health associated with an absolute change in expenditure is extrapolated to the other PBCs, where as in column 2 it is the relative effect on health of an absolute change in expenditure that is extrapolated. Since we know that QALY burden differs between (and within) PBCs and especially between the groups of PBCs with and without estimated outcome elasticities (see Table C45 in Appendix C), ${ }^{84}$ it is the values based on QALY burden in cotymn 2 that are regarded as most credible and represent our central or best estimate.

A detailed breakdown of changes in expenditure and changes in QALYs across all PBCs is provided in Table C48 in Appendix C when the analysis is based on QALY ratios and when based on QALY burden of disease. A comparison of these values confirms that QALY effects for the other PBC are lower and therefore the cost per QALY for each of these PBCs are in general much higher when based on a
$\qquad$
${ }^{81}$ See previous footnote and Table 52 in Appendix C.
${ }^{82}$ Note that this is the ratio of total change in health to total change in expenditure across these PBC (rather than an average ratio) and the contribution that each of these PBCs make to these total effects on health and expenditure depends on the estimated expenditure as well as outcome elasticities.
${ }^{83}$ Indeed, applying the absolute health effect of expenditure from the 11 PBCs with outcome elasticities implies different (higher) proportionate effects in the other PBCs
${ }^{84}$ The QALY burdens per incident patient are reported in this Table for each PBC, including the median and range across the contributing ICD codes. However, these values should not be over interpreted as the 'average' QALY burden for the PBC depends on how PBC effects are allocated to ICDs (i.e., those which have the higher contribution to variance in PBC costs) and the 'average' burden for groups of PBCs depends on how a change in overall expenditure is shared between them, i.e., the expenditure elasticities estimated for each PBC in Chapter 3 and Appendix B.
proportionate effect on QALY burden. Of course, we have not directly observed quality of life effects in these PBC but inferred them from the proportionate effects that we can observe. Insofar as investment and disinvestment opportunities in these PBCs might have been more valuable (offered greater improvement in quality of life ${ }^{85}$ than suggested by the implied PBC thresholds, then overall QALY effects will tend to be underestimated and the cost per QALY threshold overestimated. For the reasons discussed in previous sections, we regard all the cost per QALY threshold reported in column 2 of Table 4.18 as on balance conservative with respect to overall health effects of a change in expenditure. However, the estimate of $£ 15,701$ maybe especially conservative with respect to health effects (i.e., overestimated) based, as it is on an extrapolation of the proportionate effects to measures of burden on these PBC, rather than observations of the direct impact of changes in expenditure on quality of life in these types of disease. This is especially so in PBC 5 Mental Health Disorders, which accounts for a large proportion of the change in overall expenditure ( $30 \%$ ) and where a review of the evidence suggests that the investment and disinvestment opportunities in this PBC are likely to have been more valuable than the implied PBC cost per QALY of $£ 60,111$ (see Addendum 3 Appendix C) ${ }^{86}$. The lower costper QALY threshold for the 11PBCs with outcome elasticities $(£ 5,128)$ might be regarded as more secure ini this respect but they only account for a proportion ( $27 \%$ ) of any change in overall expenditure (see Table C53 in Appendix C).

Table 4.19 reports how the estimated QALY effects based on measures of QALYBurden for each PBC can be decomposed into that part associated with life year effects adjusted for quality and that part associated with 'pure' quality of life effects. These results are very similar to those reported in Table 4.17 which were based on QALY ratios for the 11 PBCs with an estimated outcome elasticity. Those PBCs for which mortality is the major concern have a much greater share of total QALY effects associated with avoidance of premature death (e.g., PBC2 and PBC10) compared to those where quality of life is the major concern (e.g., PBC 7). The differences tend to favourQALY's gained though avoidance of disability, which reflects the underestimation of the effects on 'pure' quality of life when using QALY ratios based on estimates of YLL from GBD (see the discussion above). ${ }^{87}$ The QALY to YLL ratios that are implied by this analysis are reported in Table C50 Appendix C. As expected the implied QALY ratio across all 11PBCs with outcome elasticities is higher ( $3.05^{88}$ ) then reported in Section 4.4.1 because the previous bias against quality of life effects by using QALY ratios based on unadjusted GBD information has been removed.

Recall that in Section 4.4.1 the ratios of QALYs to YLL due to disease in those PBC where outcome elasticities could not be estimated could not be used to inform estimates of the threshold or indicate how any total health effects in these other PBCs are likely to be 'shared' between life year effects adjusted for quality and that part associated with 'pure' quality of life effects (see Table 4.17). By applying the observed proportionate effects of changes in expenditure to measures of QALY burden of disease in these other PBCs the likely share of any effects on QALYs between avoidance of premature mortality and avoidance of disability more closely reflect the nature of these types of diseases (see Table 4.19). As expected, a much greater proportion of QALY effects are associated with quality of life during the disease compared to the 11 PBCs where mortality based outcome elasticities could be estimated. The share of effects in particular PBCS are also much more credible. For example, in PBC5 Mental Health Disorders the overwhelming share of QALY effects are associated with quality of life itself and for others, such as PBC12 Dental problems, PBC9 Problems of Hearing and PBC8 Problems of Vision; almost all effects are
${ }^{85}$ See Addendum 3 in Appendix C for an examination of the value of investment and disinvestments that may have been available in PBC5 (Mental Health Disorders), which accounts for much of the change in overall expenditure. This qualitative analysis suggests that these may well be more valuable than the implied PBC cost per QALY of $£ 60,111$ reported in Table C56 in Appendix C.
${ }^{86}$ See footnote above.
${ }^{87}$ The exception is PBC $18 \& 19$. The reason is that there are significant adjustments made based on differences in observed and recorded mortality (to adjust for differences in classification when mapping from U codes to PBCs via ICDs) as well as differences in YLL due to the GBD method of calculation (see Table 52 in Appendix C).
${ }^{88}$ The implied QALY ratios across these 11 PBCs range from 0.70 in PBC2 Cancer to 14.86 in PBC7 Neurological.

Table 4.19: Decomposing estimated QALY effects by PBC

associated with quality of life rather than mortality and life years. For this, and the other reasons discussed above, the analysis based on measures of QAL Mburden are regarded as the best estimate of a cost per QALY ratio that reflects a more complete picture of the likely health effects of changes in overall expenditure.

### 4.4.3 Summary of the cost per QALY threshold

The results of the three sequential steps of analysis described in this Chapter are summarised in Table 4.20. In Section 4.2 we explored ways in which the estimated effects on mortality from the econometrics work in Chapter 3 might be better translated in to life year effects by overcoming some of the limitations of mortality data available at PCT level and taking account of counterfactual deaths. The results of this analysis were reported in Table 4.9 and are repeated in column 1 of Table 4.20.89 In Section 4.3 we considered how the estimated life year effects might be adjusted for the quality of life in which they are likely to be lived, taking account of the gender and the age at which life years are gained or lost (see Table 4.14). The results of this analysis are repeated in column 2 below. Finally in Section 4.4 we explored ways to also take account of the likely effects of changes in expenditure on quality of life during disease as well as the effects associated with mortality and life years (see column 3). These estimates provide our central estimate of cost per QALY threshold, because they make best use of available information while the assumptions required, which on balance are likely conservative with respect to health effects, appear more reasonable than the other alternatives available. ${ }^{90}$

[^27]Table 4:20: Summary of cost per QALY threshold estimates

|  | [1] | [2] | [3] |  |
| :---: | :---: | :---: | :---: | :---: |
| QoL associated with life extension: QoL during disease: | $\begin{aligned} & 1 \\ & 0 \end{aligned}$ | $\begin{gathered} \text { Norm } \\ 0 \end{gathered}$ | Norm Based on burden |  |
| Effect of expenditure on mortality: <br> YLL per death averted: QALYs per death averted: | $\begin{gathered} 1 \text { year } \\ \sim 4.1 \mathrm{YLL} \\ \sim 4.1 \text { QALY } \end{gathered}$ | $\begin{gathered} 1 \text { year } \\ \sim \\ \sim \\ \sim \\ \sim \end{gathered} .5 \text { YLL } \text { QALY }^{1} .$ | $\begin{gathered} \text { Best estimate } \\ 1 \text { year } \\ \sim 4.1 \mathrm{YLL} \\ \sim 12.6 \text { QALY } \end{gathered}$ |  |
| big 4 PBC's <br> 11 PBCs (with mortality) <br> All 23 PBCs | $\begin{gathered} £ 8,080 \\ £, 15,628 \\ £, 17,663 \\ \hline \end{gathered}$ | $\begin{gathered} £, 6,631 \\ £ 18,622 \\ £ 21,047 \end{gathered}$ | $\begin{aligned} & £, 0,036 \\ & £, 5,128 \\ & £, 15,701 \end{aligned}$ | $\begin{aligned} & {[1]} \\ & {[2]} \\ & {[3]} \end{aligned}$ |
| Effect of expenditure on mortality: <br> YLL per death averted: QALYs per death averted: | Remainder of disease duration $\begin{gathered} \sim 4.1 \mathrm{YLL} \\ \sim \\ \sim \end{gathered}$ | Remainder of disease duration ~ 4.1 YLL <br> ~ 3.5 QALY | Lower bound <br> Remainder of disease duration ~ 4.1 YLL <br> ~ 12.6 QALY |  |
| big 4 PBC's <br> 11 PBCs (with mortality) <br> All 23 PBCs | $\begin{aligned} & £ 3,846 \\ & £ 6,106 \\ & £ 6,901 \end{aligned}$ | $\begin{aligned} & £ 4,252 \\ & £ 6,852 \\ & £ 7,744 \end{aligned}$ | $\begin{gathered} £ 674 \\ £ 860 \\ £ 2,785 \end{gathered}$ | [ 14$]$ |
| Effect of expenditure on mortality: <br> YLL per death averted: QALYs per death averted: | $\begin{gathered} 1 \text { year } \\ 2 Y L L \\ \sim 2 \text { QALY } \end{gathered}$ | $\begin{gathered} 1 \text { year } \\ 2 \text { YLL } \\ \sim 1.9 \text { QALY } \end{gathered}$ | Upperbound 1 rear 21 LL 6.1 QALY |  |
| big 4 PBC's <br> 11 PBCs (with mortality) <br> All 23 PBCs | $\begin{aligned} & £ 16,432 \\ & £ 32,387 \\ & £, 36,604 \end{aligned}$ | $\begin{aligned} & £ 17,456 \\ & £ 34,492 \\ & £, 38,983 \end{aligned}$ | $\begin{array}{r} £ 6,292 \\ £ 10,626 \\ £, 32,537 \end{array}$ | $[7]$ <br> $[8]$ <br> $[9]$ |

The estimate of $£ 5,128$ per QALY (line 2) is restricted to the effects of changes in expenditure in the 11 PBCs where outcome elasticities can be estimated. Although this might be regarded as more secure these PBCs only account for a proportion of a change in overall expenditure (approximately $28 \%$, see Table 61 in Appendix C). The threshold of $£ 15,701$ uses the estimated proportionate effects of expenditure on the QALY burden of disease in these PBC as a surrogate for proportionate effects in the others, i.e., assuming that the effects that can be observed will be similar to those that cannot. As discussed in Section 4.4.2 there are reasons to suspect that this may underestimate health effects in these PBCs which have most influence on the overall threshold. As in previous sections, no health effects are assigned to PBC23 (General Medical Services) on the basis that any health effects of this expenditure would be recorded in the other PBC5.9 Therefore, the best or central estimate of cost per QALY threshold is $£ 15,701$ (column 3, line 3). However, this estimate reflects changes in undiscounted QALYs associated with changes in expenditure. Although all the health effects of a change in expenditure are restricted to one year (so no discounting is necessary) some of the quality adjusted life year effects of a change in mortality in that year vill occur in future years, so in principle should be discounted. However, discounting these life year effects, even at the higher rate of $3.5 \%$ recommended by NICE, only increases the cost per QALY threshold to $£ 15,940$ (see Table C52 in Appendix C for discounted values).

As in previous Sections of this Chapter, the upper and lower bounds for the cost per QALY thresholds in column 3 are based on making the necessary assumptions about duration of health effects and how long a death might be averted optimistic (providing the lower bound for the threshold) or conservative (an upper bound for the threshold). The lower bound (lines 4 to 6 ) is based on assuming that health effects are not restricted to one year but apply to the whole of the remaining disease duration of the population at risk in PBCs during one year. Although this combines optimistic assumptions, it is possible that at least some part of a change in expenditure may prevent disease so will have an impact on populations that are incident to PBCs in the future. Such effects are not captured in any of the estimates presented in this Chapter so all are conservative with respect to this type of health effects of expenditure. The upper

[^28]bound (lines 7 to 9 ) is based on the combination of assuming that health effects are restricted to one year for the population currently at risk and that any death averted is only averted for 2 years (see Section 4.2.5).

## Chapter 5: Implications for a policy threshold

### 5.1 Introduction

The three sequential steps of analysis, which provide a cost per life year threshold (see Section 4.2 of Chapter 4) through a cost per life year adjusted for quality (see Section 4.3) to a cost per QALY threshold (see Section 4.4), have been explained in this Chapter using the analysis of 2006 expenditure and mortality data from 2006 to 2008 (see Section 3.5.1 in Chapter 3 and Section B8.5 in Appendix B) to illustrate the implications for the threshold estimates. At each step we explored the different ways that routinely available data could be used and how additional information could improve our estimates. In doing so we identified a preferred analysis at each stage based on which made the best use of available information, whether the necessary assumptions appeared more reasonable than the alternatives available, and which provided a more complete picture of the likely health effects of a change in expenditure.

## 5. 2 Re-estimating the cost per QALY threshold using more recent data

The same methods of analysis can be applied to the econometric analysis of the 2008 expenditure and 2008 to 2010 mortality data (see Section 3.5.3 in Chapter 3 and Section B11 in Appendix B). The differences between the 2006 analysis reported in Chapter 4 and the analysis of expenditure in 2008 reported below are the: i) total PBC expenditure; ii) estimated expenditure elasticities; iii) estimated outcome elasticities; iv) observed PBC deaths by age and gender; and v) life expectancy by age and gender. The other information about quality of life norms (see Section 4.3.1), disease related decrements in quality of life (see Section 4.3.2) and the information from GBD about incidence and duration of disease remain unchanged between 2006 and 2008 (we discuss in this chapter how these estimates might be improved through access to more recent and better data). (-)

It should be noted that important improvements were made to the classification and collection of PBC expenditure data that took place after the 2006 data were collected. Therefore, the differences in threshold estimates between 2006 and 2008 partly reflect this (see Section 3.5.4 and B11.4 in Appendix B) so should not be over interpreted. The results of the analysis of 2007 and 2008 expenditure are comparable in this respect, providing insights into how the threshold might change over time and with changes in the overall budget. The implications of this analysis on the need for periodic reassessment are discussed in Section 4.5.5. For the purposes of this methodological research the 2008 expenditure and 2008 to 2010 mortality data were the latest to be analysed. Since it is the analysis of the most recent data that is of most policy relevance, our discussion throughout this Section is based on analysis of 2008 expenditure, although the same sensitivity analysis (see Section 4.5.2) and analysis of uncertainty (see Section 4.5.3) is available for 2006 and 2007 expenditure (see Section C.2.5 in Appendix C).

For these reasons it is unnecessary to repeat all the analysis presented in Sections 4.2 to 4.4 (the details of each stage of the analysis of 2008 data can be found in Appendix C). Instead the results of the three sequential steps of analysis are summarised in Table 5.1. They include: i) the cost per life year (column 1) ${ }^{92}$ based on the methods of analysis outlined in Section 4.2; ii) the cost per life year adjusted for quality of life (column 2) ${ }^{93}$ based on the methods of analysis outlined in Section 4.3; and iii) the cost per QALY (column 3) based on the methods of analysis outlined in Section 4.4. These estimates, in column 3, take account of the likely effects of changes in expenditure on quality of life during disease as well as the effects associated with mortality and life years; making best use of available information, while the

[^29]assumptions required appear more reasonable than the other alternatives available. For this reason these estimates remain our central or best estimates for all the waves of expenditure and mortality data.

Table 5.1: Summary of cost per QALY threshold estimates (2008)

|  | [1] | [2] | [3] |  |
| :---: | :---: | :---: | :---: | :---: |
| QoL associated with life extension: QoL during disease: | $\begin{aligned} & 1 \\ & 0 \end{aligned}$ | $\begin{gathered} \text { Norm } \\ 0 \end{gathered}$ | norm <br> Based on burden |  |
| Effect of expenditure on mortality: <br> YLL per death averted: QALYs per death averted: | $\begin{aligned} & 1 \text { year } \\ \sim & 4.5 \mathrm{YLL} \\ \sim & 4.5 \text { QALY } \end{aligned}$ | $\begin{gathered} 1 \text { year } \\ \sim 4.5 \mathrm{YLL} \\ \sim \\ \sim \end{gathered}$ | $\begin{gathered} \text { Best estimate } \\ 1 \text { year } \\ \sim 4.6 \mathrm{YLL} \\ \sim 12.7 \text { QALY } \end{gathered}$ |  |
| big 4 PBC's <br> 11 PBCs (with mortality) <br> All 23 PBCs | $\begin{aligned} & £ 10,220 \\ & £ 23,360 \\ & £ 25,214 \end{aligned}$ | $\begin{aligned} & £ 12,338 \\ & £ 28,045 \\ & £, 30,270 \end{aligned}$ | $\begin{array}{r} £^{\infty}, 8,872 \\ £, 8,308 \\ £, 18,317 \end{array}$ | $\begin{array}{\|l} {[1]} \\ {[2]} \\ {[3]} \\ \hline \end{array}$ |
| Effect of expenditure on mortality: <br> YLL per death averted: QALYs per death averted: | Remainder of disease duration $\begin{gathered} \sim 4.5 \mathrm{YLL} \\ \sim 4.5 \text { QALY } \end{gathered}$ | Remainder of disease duration $\begin{gathered} \sim 4.5 \mathrm{YLL} \\ \sim 3.8 \text { QALY } \end{gathered}$ | Lower bound <br> Remainder of disease duration ~ 4.6 YTL <br> ~ 12.7 QAL | 2 |
| big 4 PBC's <br> 11 PBCs (with mortality) <br> All 23 PBCs | $\begin{aligned} & \AA, 5,083 \\ & £, 8,579 \\ & £ 9,260 \end{aligned}$ | $\begin{gathered} \AA^{\infty}, 811 \\ £, 9,861 \\ £ 10,644 \end{gathered}$ | $\begin{array}{r} f 1,194 \\ 6,1,175 \\ \mathbf{f}, 2,832 \end{array}$ | [4] <br> [5] <br> [6] |
| Effect of expenditure on mortality: <br> YLL per death averted: QALYs per death averted: | $\begin{aligned} & 1 \text { year } \\ & 2 \text { YLL } \\ & \sim 2 \text { QALY } \end{aligned}$ | $\begin{gathered} 1 \text { year } \\ 2 Y L L \\ \sim \\ 1.4 Q A L Y \end{gathered}$ | $\begin{gathered} \text { Upper bound } \\ 1 \text { year } \\ 2 \mathrm{YLL} \\ \sim 5.6 \text { QALY } \end{gathered}$ |  |
| big 4 PBC's <br> 11 PBCs (with mortality) <br> All 23 PBCs | $\begin{aligned} & £ 23,346 \\ & £ 52,936 \\ & £, 57,136 \end{aligned}$ | $\begin{array}{r} 26,138 \\ \boxed{59,151} \\ 6,63,844 \\ \hline \end{array}$ | $\begin{aligned} & £ 11,040 \\ & £ 18,827 \\ & £ 41,507 \end{aligned}$ | [7] <br> [8] <br> [9] |

Recall that the estimate of $£ 8,308$ per QALY (column 3, line 2) is restricted to the effects of changes in expenditure in the 11 PBCs where outcome elasticities can be estimated. However, these PBCs only account for a proportion of a change in overall expenditure (approximately $35 \%$, see Table 5.2 below). As was explained in Section 4.4.2 and 4.4.3 the QALY threshold of $£ 18,317$ (column 3, line 3); the estimated proportionate effects of expenditure on the QALY burden of disease in the 11 PBCs were used as a surrogate for proportionate effects in the others, (i.e., assuming that the effects that can be observed will be similar to those that cannot) and tepresents our central or best estimate. As in previous sections, no health effects are assigned to PBC 23 (General Medical Services) on the basis that any health effects of this expenditure would be recorded in the other PBCs. ${ }^{94}$ Although this estimate of $£ 18,317$ reflects changes in undiscounted QALYS associated with changes in expenditure, discounting the quality adjusted life year effects only increases the cost per QALY threshold to $£ 18,613.95$

As in previous sections of this chapter, the upper and lower bounds for the cost per QALY thresholds in column 3 in Table 5.1 are based on making the necessary assumptions about duration of health effects of expenditure and how long a death might be averted optimistic (providing the lower bound for the thresholdeor conservative (an upper bound for the threshold). The lower bound (lines 4 to 6 ) is based
${ }^{94}$ It would be inappropriate to assign all the change in GMS expenditure to the estimate of cost per QALY based only on the 11 PBCs with outcome elasticities because it would imply that GMS only contributes to these PBCs.
Restricting attention to the 11 PBCs with outcome elasticities but allocating part of the change in GMS expenditure to them based on their proportional share of changes in overall expenditure would yield a slightly higher cost per QALY than reported in line 2.
${ }^{95}$ The effects of discounting are modest because: i) the health effects of a change in expenditure are restricted to one year (where no discounting is necessary); ii) most of the total QALY effect occurs in that year; iii) it is only some of the life year effects (adjusted for quality) of a change in mortality in that year that occur in future years that need to be discounted; and iv) these need to be discounted only over 4.6 years on average (see Tables C89 and C90 in Appendix C for discounted values).
on assuming that the health effects of expenditure are not restricted to one year but apply to the whole of the remaining disease duration of the population at risk in PBCs during one year. Although this combines optimistic assumptions, it is possible that at least some part of a change in expenditure may prevent disease so will have an impact on populations that are incident to PBCs in the future. Such effects are not captured in any of the estimates presented in this report so all estimates are conservative in this respect (the possibility of a longer and more complex lag structure for the effects of expenditure are discussed in this chapter). The upper bound (lines 7 to 9 ) is based on the combination of assuming that health effects are restricted to one year for the population currently at risk and that any death averted is only averted for 2 years (see Section 4.2.5).

As previously shown, the estimated QALY effects associated with each PBC can be decomposed into that part due to life year effects adjusted for quality and that part associated with effects on quality of life during disease. The proportionate share of these different aspects of the total health effect are the same as reported in Table 4.19; where those PBCs for which mortality is the major concern have a much greater share of total QALY effects associated with avoidance of premature death (e.g., PBC2 and PBC10) compared to those where quality of life is the major concern (e.g., PBC 7).

### 5.3 Which PBCs matter most?

Which PBCs have the greatest influence on the overall threshold depends, to a large extent, on how a change in overall expenditure is allocated to the different PBCs (see column 1 in Table 5.2), ${ }^{96}$ i.e., those that account for a greater share of the change in expenditure will tend to have the greater influence. However, it also depends on the proportionate effect of a change in PBC expenditure on the QALY burden associated with the $\mathrm{PBC}^{97}$ and the scale of the QALY burden (for the population at risk) associated with the type of diseases that make up each $\mathrm{PBC}^{88}$. These determine the cost per QALY associated with each PBC (see column 4 below and Table C80 in Appendix C). The share, attributable to each PBC, of the total health effects of a change in overall expenditure (see column 2 of Table 5.2) is the combined effect of all of these. The proportionate impact on the overall cost per QALY threshold of a $10 \%$ change in PBC health effects in column 3 gives an indication of how sensitive the overall threshold is to the estimate of health effects associated with each PBC. It starts to suggest where further efforts to improve estimates of the overall threshold might be most usefully directed.

Although the 11 PBCs where outcome elasticities could be estimated only account for $36 \%$ of the change in overall expenditure they account for $80 \%$ of the overall health effects. Within this group some PBCs contribute more than others. For example, PBC11 (Respiratory) accounts for a greater share of total health effects and has a higher elasticity ( $3.05 \%$ ) than PBC10 (Circulatory) even though the latter accounts for a greater part of a change in overall expenditure. The reason is that the cost per QALY associated with changes in expenditure in PBC11 is lower than PBC10 and much lower than the overall threshold (so generates more health effects for the same, or even smaller, change in expenditure). ${ }^{99}$ The elasticities in column 3 are instructive, e.g., the elasticity for PBC11 suggests that even if the health effects of a change in expenditure in this PBC were over estimated by $30 \%$ the overall threshold would increase by $9.15 \%$ to 19,993 . All other PBCs have much less influence in this respect. Nonetheless PBC10 is importantcompared to others as it does contribute a large share of total health effects and has one of the
${ }^{96}$ Which is determined by the estimated expenditure elasticities (the proportionate change in PBC expenditure due to a change in overall expenditure) and total PBC expenditure (see Chapter 3 and section B11 in Appendix B)
97 Which are determined by the outcome elasticities (the proportionate effects on mortality and YLL of a proportionate change in PBC expenditure (see Section 4.4.2 for details of how these estimates can be applied to measures of QALY burden in all PBCs).
${ }^{98}$ See Section 4.4 for how PBC level effects can be allocated to the contributing ICD codes and how measures of QALY burden for each ICD code can be established
${ }^{99}$ Within PBC11: Chronic lower respiratory diseases (J40-J47) accounts for $85 \%$ of the QALY effects of a change in PBC expenditure; Lung diseases due to external agents (J60-J70), 4\%; Other diseases of upper respiratory tract (J30J 39 ), $4 \%$; Other respiratory diseases principally affecting the interstitium (J80-J84), $1 \%$; and Other diseases of pleura (J90-J94), 1\%. The other ICD codes each contribute less but together account for $4 \%$ of the health effects of a change in PBC11 expenditure.
highest elasticities (1.43\%). ${ }^{100}$ Also PBC7 (Neurological), although accounting for a smaller share of a change in overall expenditure, does contribute a large share of total health effects with an elasticity of $1.45 \%$ and a relatively low cost per QALY associated with changes in PBC expenditure. ${ }^{101}$

Table 5.2: Impact of each PBC on the overall cost per QALY threshold (2008)


* The proportionate change in the overall cost per QALY threshold due to a $10 \%$ increase or decrease in the health effects associated with the PBC. These elasticities are correct up to a $50 \%$ change in health effects.

The other 12 PBCs , where outcome elasticities could not be estimated account for the greater part of a change in overall expenditure ( $64 \%$ ) but only $20 \%$ of the overall health effects, i.e., the cost per QALYs associated with a change in expenditure in these PBCs is, in general, much higher. Of course, we have not directly observed quality of life effects in these PBCs but inferred them from the proportionate effects that we can observe. Insofar as investment and disinvestment opportunities in these PBCs might have been more valuable (offered greater improvement in quality of life) than suggested by the implied PBC thresholds in column 4, the overall QALY effects will tend to be underestimated and the overall cost per QALY threshold will be overestimated.

The overall threshold of $£ 18,317$ maybe especially conservative (i.e., likely to be overestimated) with respect to healtheffects in PBC5 (Mental Health Disorders), which accounts for a large proportion of the change in oyerall expenditure ( $25 \%$ ) and contributes most to the overall health effects ( $9 \%$ ) compared to these other PBCs. The cost per QALY associated with this PBC $(£ 49,835)$ is based on an extrapolation of estimated proportionate effects to a population based measures of QALY burden in this PBC, rather
${ }^{100}$ Within PBC10: Ischemic heart diseases (I20-I25) accounts for $55 \%$ of the QALY effects of a change in PBC expenditure; Cerebrovascular diseases (I60-I69), $21 \%$; Other forms of heart disease (I30-I52), $7 \%$; Congenital malformations and deformations circulatory system (Q20-Q28), 3\%; and Diseases of veins, lymphatic vessels and lymph nodes, not elsewhere classified (I80-I89), $3 \%$. The other ICD codes each contribute less but together account for $8 \%$ of the health effects of a change in PBC 10 expenditure.
${ }^{101}$ Within PBC7 Episodic and paroxysmal disorders (G40-G47) accounts for $73 \%$ of the QALY effects of a change in PBC expenditure; Extrapyramidal and movement disorders (G20-G26), 8\%; Other degenerative diseases of the nervous system (G30-G32), $5 \%$; Other disorders of the nervous system (G90-G99), 3\%; and Nerve, nerve root and plexus disorders (G50-G59), $2 \%$. The other ICD codes each contribute less but together account for $9 \%$ of the health effects of a change in PBC 7 expenditure.
than observations of the direct impact of changes in expenditure on quality of life in the types of diseases that make up the PBC. Evidence that is available suggests that the investment and disinvestment opportunities in this PBC are likely to have been much more valuable than this implied cost per QALY.
A review of the evidence of the cost effectiveness of the investment and disinvestment opportunities that have been available in mental health during this period is reported in Addendum 3 Appendix C. A search for evidence about interventions in those ICD codes that contribute most to the PBC (based on prevalence or the contribution to the variance in PBC costs), suggests that pharmacological, psychological and social interventions for depression are all more cost effective (in general much less than $£ 10,000$ per QALY) than the overall threshold and significantly more valuable than the implied QALY threshold for this PBC. Based on the contribution that each ICD makes to variance in PBC costs across PCTs, it is schizophrenia that contributes most. Although interventions that may have been invested or disinyested in schizophrenia are, in general, less cost effective (in general less than $£ 24,000$ per QALY) than those available for depression, they are still much more valuable than the implied cost per QALY of this PBC in Table 5.2. ${ }^{102}$

It is very important not to misinterpret the cost per QALY associated with each PBC in column 4 of Table 5.2. These are not cost effectiveness thresholds. That is, they do not represent the QALYs likely to be forgone due to costs imposed (e.g., by the approval of a new and more costly technology by NICE) in a particular PBC because NHS expenditure is not devolved and constrained to PBC specific budgets. Rather the overall expenditure is constrained through government decisions about public expenditure, but within the NHS resources (at the margin at least) are fungible in anything other than the very short run across different activities and disease areas. For example, the additional net NHS costs of approving a new but more costly technology in PBC10 (Circulatory) will not be restricted to the circulatory PBC ( $5.5 \%$ will, see column 1 in Table 5.2) but are likely to be reallocated in the same way as an equivalent reduction in overall expenditure (i.e., the shares of a change in overall expenditure in column 1). ${ }^{103}$ Therefore, the relevant cost per QALY threshold for a teehnology in the Circulatory PBC is not $£, 7,038$ but the overall threshold of $£, 18,317$.

### 5.4 How uncertain are the estimates and what are the implications?

There are a number of sources of uncertainty which contribute to an assessment of how uncertain a central or best estimate of the cost per QALYthreshold might be. There are three reasons why uncertainty in the estimate of the threshold might be of policy interest: i) the uncertainty in the parameters that determine the threshold might influence the mean or expected value of the threshold if they have a non linear relationship to the threshold or when they have a multi linear relationship but are correlated with each other; ii) the consequences of over or underestimating the threshold differ so the uncertainty may have an influence on the extent to which a policy threshold (one that can be compared to the incremental cost effectiveness ratio of a new technology) should differ from the mean or expected


102 Although the published evidence suggests that investment and disinvestment opportunities in this PBC tend to be much more valuable than the imputed cost per QALY, in common with other PBCs, there will inevitably be inefficient, ineffective or even iatrogenic practice (e.g., due to poor diagnosis and inappropriate prescribing). Insofar as these types of activities are sensitive to changes in PBC expenditure this will tend to increase the cost per QALY associated with changes in expenditure in this PBC. Whether both the extent of these inefficiencies and their sensitivity to changes in expenditure are sufficient to increase the cost per QALY above $£ 49,835$ is unclear, although it seems unlikely. Note that the effects of the scale and sensitivity to expenditure of inefficient or even harmful practice in the other PBCs where outcome equations could be specified are already captured in the estimated outcome elasticities.
${ }^{103}$ In principle at least, with sufficient panel data which would allow a more complex lag structure and simultaneous estimation of expenditure and outcome elasticities across all PBCs ; it might be possible to isolate the short run effects of a change in expenditure in one PBC across all the others. In the absence of such data and so long as adjustments are expected take place quickly relative to the time horizon of the effects of the new technology on NHS cost and outcomes (i.e., marginal NHS resource is fungible in the medium term) using the overall cost per QALY threshold for technologies relevant to any PBC is reasonable and more so than other alternative assumptions that might be made.
value of the central or best estimate; and iii) in conjunction with other methods of analysis ${ }^{104}[95]$ it can indicate the potential value of gathering more information to improve these estimates in the future. Of course, hypothesis testing and the traditional rules of inference associated with it, such as statistical significance, p -values and confidence intervals, have no relevance when making unavoidable decisions about policy relevant quantities based on information currently available and the best use thereof.[96]

## An assessment of parameter uncertainty

Two sets of parameters are critical to the threshold, the expenditure elasticities estimated for each of the 23 PBCs , and the outcome elasticities estimated for 11 of these. These parameters are estimated with uncertainty, indicated by the standard errors on the relevant coefficients in the econometric analysis outlined in Chapter 3 and detailed in Appendix B. Since these statistical models estimate coefficients using normality on the relevant scale, normal distributions can be assigned to each of these estimated coefficients, each with a mean and standard deviation based on the results of the econometric analysis. ${ }^{105}$ These distributions, represent the uncertainty in the mean estimate of each of the parameters and can be propagated through the various calculations required to estimate and overall cost per QALYthreshold (i.e., through the sequence of analysis detailed in Section 4.2 to 4.4 ) using Monte Carlo simulation which randomly samples from the assigned distributions. The results of each random sample represent one possible realisation of the overall threshold, given the uncertainty in estimates of the mean parameter values that determine it. By repeatedly sampling, a distribution of potential values that the overall threshold might take can be revealed. The results of this simulation are illustrated in Figure 4.3 which illustrates the cumulative probability density function for a cost per QALY threshold based only on the 11 PBC with estimated outcome elasticities and for all 23 PBCs . It represents the probability (on the y axis) that the threshold lies below a particular value.

It has already been noted that restricting attention only tochanges in expenditure in those 11 PBCs where an outcome elasticity can be estimated is much lower than considering all changes in expenditure across all PBCs - the threshold value on the $x$-axis that corresponds to a probability of 0.5 is much lower in Figure 4.3 for these 11 PBCs (the mean is very similar but slightly greater than median values - see Section C.2.3.1 in Appendix C). This lower estimate of $£ 8,308$ per QALY is much less uncertain but these PBCs only account for $36 \%$ of a change in overall expenditure, so it is the higher estimate, for all 23PBCs, that is of most relevance for policy (see Sections 4.4.3 and 5.2). The fact that this estimate is more uncertain simply reflects the quality and quantity of data currently available. Since useful analysis should endeavour to faithfully characterise uncertainty in policy relevant quantities, rather than select those quantities or questions for which precise estimates are possible, it is the more uncertain estimate for all 23 PBCs that should be of primarily interest. The values that are used to generate Figure 5.1 are available in Table C81 in Appendix C. They indicate that the probability that the overall threshold is less than $£ 20,000$ per QAL is 0.64 and the probability that is less than $£ 30,000$ is 0.92 .
${ }^{104}$ A form of value of information analysis could be applied to these estimates in subsequent research, ideally capturing some of the other sources of uncertainty. Such analysis has firm foundations in statistical decision theory and has been applied to health care decisions. More recently it has been applied to the decisions faced by NICE when considering whether there is sufficient evidence to support the approval of a new technology ${ }^{105}$ The Monte Carlo simulation is in essence Bayesian, where the standard errors from the frequentist econometric analysis are used to assign normal prior distributions with means equal to the point estimates and a standard deviation equal to the estimated standard errors. This is equivalent to a fully Bayesian analysis with initially uninformative priors which are updated through the analysis of expenditure and mortality data.


Figure 5.1: Cumulative probability density function for the cost per QALY threshold

## The implications of uncertainty

Integrating this parameter uncertainty into the estimates of the overall threshold does not change the mean or expected value of the cost per QALY threshold. ${ }^{\text {tog }}$ This is to be expected as the expenditure and outcome elasticities have a multi linear relationship to the overall threshold and the analysis sampled independently from the distributions assigned to estimated coefficient. We did investigate the potential correlation between the expenditure and outcome elasticities by repeatedly re-estimating both based on randomly sampling with replacement the 152 PCTs - creating bootstrapped data sets where the original PCTs could appear more than once or not at allin the re-sampled data. This analysis indicated a small positive correlation between outcome and expenditure elasticities in 4 PBCs using 2006 expenditure data (see Section B. 10 in Appendix B). Such levels of correlation will tend to have a modest but positive influence on the mean value of the cost per QALY threshold. ${ }^{107}$ Uncertainty in the estimate of the overall threshold means that a policy threshold set at its mean or expected value may be inappropriate. Insofar as the consequences (to the NHS) of under or over estimation are symmetrical, then the expected or mean value would be the appropriate policy threshold irrespective of the scale of uncertainty. However, the consequences of overestimating the threshold are more serious than underestimating it. This is illustrated in Figure 5.2 which is similar to Figure 2.1 presented in Chapter 2.
${ }^{106}$ Note that the mean of the simulated values is not the mean of the sampled ratios but the ratio of the mean sampled values for the numerator and denominator. Deterministic and simulated values are the same for 2006, 2007 and 2008 expenditure data (other than negligible Monte Carlo error from 1000 samples). Also note that in constructing the cumulative probability density function in Figure 5.1 and the histograms of values in Appendix C it is important to identify whether sampled negative values favour a low value for the threshold or and unbounded one (there were no negative values sampled in the simulation of values for all 23 PBCs ).
${ }^{107}$ Positive correlation suggests that a high spend elasticity will be associated with a high outcome elasticity (i.e., less negative, implying a smaller heath effect of a change in expenditure) resulting in a higher estimate of the threshold. It also suggests that when spend elasticity is low outcome elasticity will also tend to be lower (i.e., more negative, implying a larger health effect of a change in expenditure) resulting in a lower estimate of the threshold. Although realisations of spend elasticities higher and lower than the mean estimate are equally likely, higher spend elasticities provide a greater 'weight' associated with higher estimates of the threshold (where outcome elasticity is also high) when calculating the mean threshold. For these reasons a positive correlation will tend to increase the mean estimate of the threshold.


Figure 5.2: Consequences of over and underestimating the overall threshold
It shows the impact on net health benefit if the central estimate of $£ 2 \theta, 000$ is in fact an overestimate and the threshold should be $£ 10,000$ per QALY. In these circumstances the technology should not have been approved at price $\mathrm{P}^{2}$. This overestimation leads to a loss of net health benefit of 2 QALYs as a consequence. Alternatively, the central estimate of $£ 20,000$ may be an underestimate and the threshold should be $£ 30,000$ per QALY. In these circumstances the technology could just as easily have been rejected or approved based on the central estimate and price $\mathrm{P}^{2}$. However, it should in fact have been approved at the threshold of $£ 30,000$. This underestimation leads to a loss of net health benefit of $2 / 3$ of a QALY as a consequence, i.e., less than the loss associated with the same scale of overestimation. If the scale of under or overestimation of the central estimate is equally likely (the distribution of possible values of the threshold is symmetrical) then using the mean or expected value as a policy threshold (one that can be compared to the incremental costeffectiveness ratio of a new technology) will lead to a loss of net benefit. A policy threshold that represents the maximum the NHS can afford to pay for QALY gains offered by a technology will be lqwer than the mean of the cost per QALY threshold (i.e., lower than $£ 18,317$ ) to compensate for the more serious consequences of overestimating the 'true' value. ${ }^{108}$ Importantly this remains the case even if effects are expressed in terms of their equivalent consumption value (net money benefit based on 'willingness to pay') rather than a measure of net health benefit. ${ }^{109}$ How much lower a policy threshold should be set below the mean or expected value depends on three considerations: i) the scale of uncertainty in the estimate of the threshold (greater the uncertainty implies a lower policy threshold); ii) the scale of the incremental costs relative to incremental health benefits

[^30]offered by the technology (policy threshold should only be equal to mean estimate if there are no additional NHS costs associated with the technology); and iii) the skewness of the distribution of cost per QALY threshold (a positive skew tends to offset these effects - see Figure C8 in Appendix C). The overall scale of the impact on a policy threshold will be specific to the additional NHS costs associated with a technology as well as the other sources of uncertainty discussed below and possible correlations between expenditure and outcome elasticities discussed above. We have not quantitatively integrated all these considerations in to an analysis of an appropriate policy threshold, although this maybe possible in future research.

## Other sources of uncertainty

The uncertainty associated with the parameters estimated in the econometric models is only one, and not necessarily the most important, source of uncertainty associated with the cost per QALY threshold. The parameter uncertainty presented above is conditional on the econometric model being 'correct'. In particular, that the instruments used to identify the causal effect on health of changes in expenditure are valid. Although all the models passed the relevant tests of validity, there remains some uncertainty about the validity of the instruments used, i.e., there remains structural or model uncertainty (see Chapter 3 for an overview).[97] For this reason we undertook an analysis of how sensitive estimates of outcome elasticities might be to instrumental validity (see Section B9.4 in Appendix B). We were also able to specify a distribution for the measure of instrumental validity used in this sensitivity analysis, i.e., how 'likely' each value might be (see Section B9.5 in Appendix B). Therefore, there are two 'levels' of uncertainty: i) the parameter uncertainty (uncertainty in estimated coefficients given a particular 'level' of instrumental validity) and the structural uncertainty in the level of instrumental validity. Both sources of uncertainty were integrated by randomly sampling the distribution of measures of instrumental validity and then, conditional on this sampled value, re-estimating outcome equations and sampling the estimated coefficients. This analysis in Section B9.5 of Appendix B shows that model or structural uncertainty constitutes a greater part of the overall uncertainty associated with the outcome elasticities, so fully integrating this source of uncertainty is likely to have a significant impact on the extent to which a policy threshold should be lower than the mean or expected value of the cost per QALY threshold. Importantly, this additional structural uncertainty has little effect on the point estimates of the outcome elasticities, i.e., the central estimate of the cost per QALY threshold is robust to uncertainty in instrumental validity in the econometric models.

Of course the parameter and structural uncertainty associated with the econometrics work outlined in Chapter 3 is itself only one source of uncertainty associated with the estimated cost per QALY threshold. Each of the steps of analysis insection 4.2 to 4.4 explored the different ways routinely available data could be used and how additional information could improve the estimates. We identified a preferred analysis (or scenario) at each stage based on which made the best use of available information, whether the assumptions required appeared more reasonable than the other alternatives available, and which provided a more complete picture of the likely health effects of a change in expenditure. Insofar as the preferred analysis is the only plausible scenario, there would be no other sources of uncertainty. However, other assumptions and judgments are possible, which although they may be judged less credible might nontheless have some probability of being the most credible (given evidence currently available). Therefore, there will be uncertainty between these alternative 'scenarios' as well as within each (the parameter and model uncertainty described above).[95] Although in principle this can be integrated into the analysis even in the absence of data to test alternative views[98]- we do not do so here since assigning prebabilities to alternative scenarios would be somewhat speculative and inevitably disputed. Instead we offer a summary of the qualitative considerations. Of course any increase in the uncertainty associated with the central estimate of the cost per QALY will impact on the extent to which a policy threshold should be lower than the mean. However, a critical issue is whether consideration of other 'scenarios' might change this central estimate, e.g., if scenarios that lead to a lower estimate are judged more credible than those that lead to higher ones. In other words the question is whether on balance the central or best estimate of $£ 18,317$ in Table 5.1 is likely to be an under or overestimate of the cost per QALY threshold.

Most of the considerations have been discussed in detail throughout Chapter 4 so are only briefly summarised here. On the one hand, there are some reasons why the health effects might be
overestimated and the central estimate of the QALY threshold underestimated. Recall from Section 4.2.3 and 4.2.4 that in calculating the life years lost that account for deaths that would have otherwise occurred is equivalent to assuming that those deaths averted by a change in expenditure returns the individuals to the mortality risk of the general population (matched for age and gender) and that the life years gained as a consequence will be lived in the same quality of life as the general population (again adjusted for age and gender) (see Table 4.14 in Section 4.3.3). Although these appear more credible than the other alternative assumptions that could be made, they are, however, optimistic with respect to the health effects of a change in expenditure, tending to underestimate the cost per QALY threshold. In addition integrating the small positive correlation between expenditure and outcome elasticities for all 11 PBCs is likely to have a modest but positive impact on the expected value of the threshold.

On the other hand there are a number of reasons why the central estimate might be overestimated. The health effects of a change in expenditure are restricted to the population at risk during one year. This is undoubtedly pessimistic in three respects: i) it means that effect on quality of life during disease only occur for one year (the effect of investment that might have long term effects on quality of life, e.g., hip replacement are excluded); ii) mortality effects are also restricted to one year, so the full effect investments that reduce mortality for patients throughout their disease duration, notjust in the first year, will not be captured; and iii) changes in expenditure that reduce incidence into the at risk population in the future (i.e., prevention of disease) will not be captured either. A more formal and longer lag structure in the estimation of outcome elasticities would be likely to capture more health effects of a change in expenditure.

The observed effects of a change in expenditure on mortality and life years in the 11 PBCs where outcome elasticities could be estimated was used as a surrogate for health effects in the other 12 PBCs (excluding GMS), i.e., the estimated effects of a change in expenditure that could be observed were used to inform those effects that currently, at least, cannot. This approach is not necessarily optimistic with respect to overall health effects. In fact there are good reasons to believe it may underestimate then (overestimate the threshold). As discussed previously in Sections 4.4.3 and 5.2; if this means of extrapolating from observed to unobserved effects is rejected then threshold estimate could be based only on the health effects of changes in expenditure in those PBCs where outcome elasticities can be estimated. This generates a much lower cost per $\widehat{Q} A L Y$ threshold $(£ 8,308)$ even if that portion of GMS expenditure was allocated to these 11 PBCS (see Section 4.2.5). Alternatively, taking account of the greater proportion of the change in expenditure allocated to the other 12 PBCs but assuming that there are no health effects of expenditure in all these other PBCs is not plausible. In fact the evidence that is available about the value of investment and disinvestment opportunities in the most important of these other PBCs (PBC 7 Mental Health Disorders), suggests that the health effects of changes in expenditure in this PBC may be underestimated and the central estimate of the threshold overestimated. (see Section 5.3 and Addendum 3 in Appendix C).

In addition, we have also shown that the uncertainty associated with our central estimate (from all sources) means that an appropriate policy threshold is likely to be below its mean or expected value. Finally, in Section 5 we explore how the threshold is likely to differ when considering opportunities to make investments (i.e., an increase in overall expenditure, or cost saving accruing to the NHS) and when disinvestment is required (a reduction in overall expenditure or costs imposed on the NHS). This analysis shows that a cost per QALY threshold relevant to technologies which impose costs on the NHS is likely tobe less than our central estimate of $£ 18,317$. Therefore, although other assumptions and judgments are possible that retain some level of plausibility, they do not all favour a higher threshold. Indeed, when considered together, they suggest that on balance the central or best estimate of $£, 18,317$ presented in Table 5.1 is, if anything, likely to be an overestimate. In Section 5.8 we discuss how some of these remaining uncertainties might be resolved through access to additional and better data and the type of analysis that would then be possible.

### 5.5 Impact of investment, disinvestment and non marginal effects

The central estimate of the cost per QALY threshold in Table 5.1 is based estimates of the health effects of changes in expenditure across all 152 PCTs, some of which will be making investments (where
expenditure is increasing) and others making disinvestments (where expenditure is reduced or growing more slowly). The cost per QALY threshold, however, is likely to differ across these different types of PCTs. This is illustrated in Figure 5.3 where the total observed variation in expenditure includes the impact of disinvestment ( $-\Delta \mathrm{E}$ ), e.g., where costs are imposed on the NHS by the approval of a more costly technology; and investment ( $\Delta \mathrm{E}$ ), e.g., where cost savings are accruing to the NHS. The central estimate of the cost per QALY threshold is the health effect of a change in expenditure across this variation in expenditure $\left(\mathrm{k}_{1}\right)^{110}$. One would expect that, other things equal, more expenditure (expanding the budget from B1) would increase health but at a diminishing rate. Therefore, the amount of health displaced by disinvestment, or a reduction in expenditure, would be expected to be greater, i.e., the threshold associated with $-\Delta \mathrm{E}\left(\mathrm{k}_{1}-\right)$ will be lower than the central estimate, $\mathrm{k}_{1}$. Equally, the health gained from investments, or an increase in expenditure, would be expected to be lower, i.e., the threshold associated with $\Delta \mathrm{E}\left(\mathrm{k}_{1}+\right)$ will be higher than $\mathrm{k}_{1}$.


Figure 5.3: Investment, disinvestment and budget impact
We have been able to examine this by re-estimating the outcome and expenditure elasticities separately for those PCTs where their actual budget is under the target allocation from the Department of Health resource allocation formula (i.e. , those under greater financial pressure and more likely to be disinvesting than investing), and those that are over target (under less financial pressure and more likely to be investing than disinvesting). The detail of this analysis (based on 2006 expenditure and restricted to the 'big 4' PBCs) are reported in Section B8.9 in Appendix B. The results confirm what would be expected given Figure 5.3 and the discussion above - the outcome elasticities are smaller (in absolute terms) for all 4 PBCs in the group of PCTs above their target allocation and larger for all 4 PBCs in those below. Therefore, the health effects of changes in expenditure are greater in all these PBCs when PCTs are under more financial pressure and are more likely to be disinvesting then investing. The cost per life year estimates for these PBCs are reported in Appendix B: $£ 10,604$ for all PCTs combined ( $k_{1}$ ); $£ 8,441$ for those PCTs under their target allocation (i.e., $\mathrm{k}_{1}$ - associated with $-\Delta \mathrm{E}$ ); and $£ 14,083$ for PCTs over their target allocation (i.e., $\mathrm{k}_{1}+$ associated with $+\Delta \mathrm{E}$ ). Although these cost per life year estimates are not based on the same calculations as Section 4.2, they do start to indicate the scale of the effect on a threshold that is most relevant for new technologies that impose net costs on the NHS.

[^31]Expenditure elasticities for these PBCs also differ between these groups of PCTs - they are higher for those under their target allocation. These PBCs together consistently offer the greatest value in terms of cost per death averted, life year or QALY (see Table 5.1 and 5.2). This suggests that budget impact not only displaces more valuable activities within each PBC (outcome elasticities are larger) but that overall expenditure tends to be reallocated to more valuable PBCs. The effect of this reallocation on the overall threshold is not captured in the cost per life year estimate reported above, which are restricted to these 4 PBCs. Therefore, extending this type of analysis to all PBCs in future research is likely to show that the effect on the cost per QALY threshold of both the sign and scale of changes in overall expenditure will be greater. Subsequent work might enable a quantitative assessment of how the relevant threshold should be adjusted for the scale of the budget impact of technologies appraised by NICE.

Although further work is needed to fully specify the quantitative effect of the scale of non marginal impact of new technologies on an appropriate threshold, the qualitative impact seems clear. Firstly, the central estimate of the threshold is likely to be an overestimate for all technologies which impose net costs on the NHS (almost all technologies appraised by NICE have positive incremental NHS costs and all effective technologies that will be subject to value based pricing will impose net costs on the NHS).[4, 5,13 ] Secondly the appropriate threshold to apply should be lower for technologies which have a greater impact on NHS costs.

### 5.6 How does the threshold change with overall expenditure?

The same methods of analysis can be applied to the econometric analysis of the 2007 expenditure and 2007 to 2009 mortality data (see Section B10 in Appendix B). This provides an opportunity to consider how the cost per QALY threshold is likely to have changed from 2007 to 2008 as overall expenditure has increased. This can provide some insights into how the threshold might be expected to change over time, as, for example, overall expenditure changes and productivity in the NHS might be expected to rise with innovation in health technologies, clinical practice and service delivery. This has implications for a judgement about the appropriate the frequency of periodic reassessment of the cost per QALY threshold.

It is not necessary the case that the threshold will rise with overall expenditure or even with NHS prices. This is illustrated in Figure 5.4 where the threshold at budget B1 is represented by $\mathrm{k}_{1}$. If overall expenditure increases to B 2 then, over things equal, the threshold would also be expected to increase (i.e., $\mathrm{k}_{1}$ now overestimates the health effects of a change in expenditure at B2). ${ }^{111}$ Increasing overall expenditure from B 1 to B 2 is equivalent to eliminating the same amount of waste in Figure 5.4, i.e., by re-allocating resources devoted to activities unproductive of health. Again, other things equal, the threshold would be expected to increase ( $\mathrm{k}_{1}$ now overestimates the health effects of a change in expenditure at B 1 ) once the waste has been eliminated. However, insofar as the productivity of those activities that are valuable to the NHS also improve through innovation in health technologies, clinical practice and service delivery, the threshold will tend to fall. Figure 5.4 illustrates a situation where the effects of eliminating waste (NHS stopping doing things it should not be doing) and, at the same time, improving productivity (NHS getting better at doing things it should do) means that the overall threshold is unchanged.
In makingan assessment of whether the threshold is likely to increase with the NHS budget it is also necessary to consider whether there is discretion over how additional resources can be spent. For example, if any growth in the overall budget is spent on national initiatives or other activities that cannot or cannot easily be disinvested, then the additional costs of technologies approved by NICE must be accommodated by displacing other activities elsewhere. Therefore, it is growth in expenditure on more 'discretionary' parts of NHS expenditure and changes in the productivity and input prices of those health care activities which more likely to be displaced which are most relevant.

Over recent years much of the real budget growth in the UK NHS has been devoted to national initiatives that are not easily displaced, e.g. new contracts for General Practitioners and consultants, national waiting time targets, information technology initiatives, etc.[99] It also includes Technology

[^32]Appraisal guidance issued by NICE itself, which has a funding mandate. Therefore, any real growth in what remains may have been more modest, so it is more likely to have been offset by any growth in the productivity of displaceable activities, e.g. drugs, devices, procedures and other services. Similarly, although there has been a general rise in input prices for the UK NHS, much of this inflation has been driven by staff as well as capital and overhead costs, some of which cannot be easily displaced. What are more relevant are the prices of inputs which could be displaced, an important element of which is drug prices. Although branded drug prices have tended to rise, at the same time there has been generic entry on patent expiry with dramatic reductions in prices for important classes of drugs.[100] Therefore, it is not self evident that the threshold has grown over recent years, despite real increases in the NHS budget.


Figure 5.4: Impact of changes in budget and productivity
The central estimates of the cost per QALY threshold for 2007 and 2008 expenditure years are reported in Table 5.3. In comparing these estimates of the QALY threshold it should be noted that important improvements were made to the classification and collection of PBC expenditure data that took place after the 2006 data were collected. Therefore, the differences in threshold estimates for 2006 and 2007 partly reflect this (see 3.5.4 in Chapter 3 and B11.4 in Appendix B) so should not be over interpreted. The results of the andysis of 2007 and 2008 expenditure are comparable in this respect.

Although oyerall expenditure increased by $6 \%$ between 2007 and 2008 which represented real growth of $2 \%$ in 2007 prices, ${ }^{112}$ the overall threshold for all 23 PBCs fell by $2 \%$ in nominal terms and by $5 \%$ in real terms.

Table 5.3 Growth in the cost per QALY threshold (2007 to 2008)


[^33]The reasons are complex but reflect changes in productivity, which differs across PBCs (changes in outcome elasticities), but also a general reallocation of a change in overall expenditure (changes in expenditure elasticities) towards those PBCs that appear more valuable in 2008. ${ }^{113}$ Given the sources of uncertainty described above, subtle differences between 2007 and 2008 should not be over interpreted. However, this analysis does suggest that the overall threshold will not necessary increase with growth in the real or even nominal NHS budget. In conjunction with the results of the analysis described in Section 5.4 it does suggest that the threshold is more likely to fall at a time when real budget growth is flat or falling and PCTs find themselves under increasing financial pressure.

Within the NICE Technology Appraisal process, the future incremental costs of a technology are expressed in real terms (at current prices) prior to discounting. Therefore, the estimates that are relevant to NICE decisions are: i) the nominal threshold in the current year ${ }^{114}$ and ii) some assessment of the real growth in the threshold over the time horizon where incremental NHS costs are incurred. If there is an expectation of real growth (or fall) in the threshold over time then one way to incorporate this is through a higher (lower) discount rate applied to future cost.[101] Indeed, an expectation of changes in the real threshold over time also suggests something about the social rate of time preference heath reyealed by budget allocations decisions.[102] However, incorporating an expected growth or decline in the threshold over time by adjusting discount rates is likely to be problematic once it is recognised that the expected incremental costs imposed by a technology are rarely uniform over time

This discussion and the results reported in Table 5.3 suggest that thereis little empirical support for an assumption that there will have been growth in the nominal threshold between 2008 and 2012.15 Growth in the nominal or real threshold seems much less likely in the future with the prospect of reduced budget growth, increased pressures to improve productivity and downward pressure on input prices. Since how the nominal or real threshold is likely to change over time cannot be assumed to follow prices or overall expenditure nor empirical estimates or theoretical predictions of a growth in the private consumption value of health (willingness to pay), it becomes especially important to be able to regularly update estimates of the cost per QALY threshold based of routinely available data (See section 5.8).

### 5.7 What type of health forgone by approval of a new technology?

The methods of analysis described in Chapters 3 and 4 and discussed in this chapter can identify, not only how many QALYs are likely to be forgone across the NHS as a consequence of approving a technology which imposes incremental costs on the NHS, it can also indicate where those QALYs are likely to be forgone and how they are made up, i.e., the additional deaths, life years lost (unadjusted and adjusted for quality of life) and the quality of life impacts on those with disease.

For example, in 2011, MICE considered whether ranibizumab for the treatment of diabetic macular oedema should be approved for widespread use in the NHS (TA237).[103] Initially this technology was rejected by NICE onthe grounds that, at its current price, it would be unlikely to be cost effective. In 2012, however, rapid review of TA237 approved ranibizumab if use was restricted to the most cost effective sub ©roup (those with central retinal thickness $\geq 400$ micrometres) and after a Patient Access Scheme (PAS) for this subgroup of patients was offered (details of the PAS which provides a discounts to the ${ }^{\text {NHS }}$ is commercial in confidence).[104]

[^34]The appraisal and guidance documents[103-105] ${ }^{116}$ provide the information required to estimate the additional NHS costs of treating this sub group of patients each year (see Addendum 4 to Appendix C for details of this example). Up to 44,000 NHS patients would be eligible for treatment with ranibizumab each year based on its licensed indication.[105] However, the subgroup of patients where ranibizumab was ultimately approved is likely to be 23,000 each year. This suggests that the approval of ranibizumab in this subgroup at the original appraisal price set in 2011 (i.e., without a PAS) would impose just over $£ 80 \mathrm{~m}$ of additional NHS costs for treating the eligible population each year.

Based on the 2008 central estimate of the cost per QALY threshold ( $£ 18,317$ in Table 5.1) the approval of ranibizumab without a PAS would have been likely to displace 4,367 QALYs elsewhere in the NHS. However, the analysis which underpins the threshold estimate can also be used to identify where the additional NHS cost of $£ 80 \mathrm{~m}$ are likely to impact and where and what type of health effects are likely to be forgone. These are illustrated in Table 5.4.

The results reported in Table 5.4 suggests that approval is likely to result in 295 additional deaths (most of which are likely to occur in Circulatory, Respiratory and Cancer PBCs - see column 2), and 1, 337 life years forgone (most of which are likely to occur in Circulatory, Cancer and Gastro-intestinal PBCs - see column 3). ${ }^{117}$ However, the impact of approval of this technology on QALYs forgone due to premature death (column 5) only accounts for a proportion of the total QALY effects (column4). Most $(3,509)$ are associated with quality of life forgone during disease (column 6). These quality of life impacts are most likely to occur in Respiratory, Neurological and Mental Health PBCs. The PBC level effects in Table 5.4 can also be examined at ICD level, whilst recognising the caveats discussed in Section 4.3 and 4.4. ${ }^{118}$ For example within in the respiratory PBC it appears to be Chronic lower respiratory diseases (J40-J47) where most additional deaths, life years and quality of life are forgone. In the Mental Health PBC the additional deaths appear to be associated with disorders due to psychoactive substance use (F10-F19) and Mood (affective) disorders (F30-F39) (see Addendum 4 in Appendix O). However, it should be recognised that these effects which are based on the central estimate in Table 5.1 are likely to underestimate the health forgone given the discussion in Section 5.4 and especially in 5.5.

The impact of a reduction in the price of this technology, either through value based pricing or the PAS that was offered during the rapid review, [1047 can also be examined in the same way. The PAS was commercial in confidence but we will consider a scenario where a $30 \%$ reduction in NHS costs was applied for this subgroup of patients. Such a discount would be expected to save 1,310 QALYs including 89 deaths averted, 401 life years ( 258 when adjusted for quality) and quality of life effects during disease equivalent to 1,053 QALYs, when compared to approval of the technology at the original price (see Addendum 4 Appendix C for more details on this scenario analysis).

In many respects this starts to make 'real' the previously abstract notion that additional NHS costs are the health and opportunities of other unknown NHS patients. The methods of analysis presented in this report go some hay to proving a empirically based and explicit quantification of the scale of opportunity costs the NHS faces when considering whether the health benefits associated with new technologies are expected to Offset the health that is likely to be forgone elsewhere in the NHS. It also starts to make the other NHS patients, who ultimately bear the opportunity costs of such decisions, less abstract and more 'known' in social decisions. Since who happens to be known or unknown is only a matter of perspective,
${ }^{146}$ All relevant documentation is available at http:// guidance.nice.org.uk/TA237 and
http://guidance.nice.org.uk/TA/Wave23/41
${ }^{117}$ The differences in contribution to deaths compared to life years reflects the distribution of age at death and the age and gender distribution of the population at risk in the ICD codes that contribute to each PBC (see Section 4.2 and addendum 1 in Appendix C).
${ }^{118}$ Recall that information about the age, gender and the incidence of sequelae associated with different diseases within a PBC are only available for u-codes which can be mapped to groups of three digit ICD codes. Also allocating PBC level effects to ICD codes was based on the contribution they made to the variance in PBC costs across PCTs based on HES data since total PBC costs are not recorded at ICD level (also see Addendum 1 in Appendix A).
time and ignorance, [106] ethical and coherent social decisions require that both should be treated in the same way. The methods of analysis discussed in this chapter have contributed to removing some of the 'ignorance' and making the unknown more real.

Table 5.4 Heath forgone across PBCs due to the approval of ranibizumab ( $£ 80 \mathrm{~m}$ budget impact)

| $\begin{aligned} & \text { PB } \\ & \text { C } \end{aligned}$ | PBC description | change in spend (m) <br> [1] | Additional Deaths <br> [2] | $\begin{gathered} \begin{array}{c} \text { Life } \\ \text { years } \\ \text { forgone } \end{array} \\ \\ \\ \hline[3] \\ \hline \end{gathered}$ | Total <br> QALYs <br> forgone | QALYs forgo Due to premature death [5] | Quality of life effects <br> [6] |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 2 | Cancer | $£ 2.59$ | 22 | 217 | 153 | 141 | 11 |
| 10 | Circulatory problems | £.4.40 | 132 | 672 | 625 | 427 | 198 |
| 11 | Respiratory problems | $£ 2.66$ | 78 | 93 | 1,330 | 58 | 1,272 |
| 13 | Gastro-intestinal | $¢_{£} 1.86$ | 15 | 143 | 255 | 94 | 161 |
|  | Big 4 | ¢,12 | 246 | 1,126 | 2,362 | 721 | 1,641 |
| 1 | Infectious diseases | £1.89 | 4 | 31 | 91 | 21 | 70 |
| 4 | Endocrine problems | $£ 1.10$ | 4 | 29 | 351 | 19 | - 332 |
| 7 | Neurological problems | £. 3.47 | 7 | 38 | 632 | 25 | 608 |
| 17 | Genito-urinary problems | £2.69 | 13 | 19 | 61 | 12. | 49 |
| 16 | Trauma \& injuries | $£ .4 .46$ | 0 | 0 | 0 | $\checkmark$ | 0 |
| 18+19 | Maternity \& neonates | $\ldots 3.96$ | 0 | 2 | 1 | $\bigcirc 1$ | 0 |
|  | 11 PBCs | £29 | 275 | 1,245 | 3,500 | ( 798 | 2,701 |
| 3 | Disorders of Blood | £2.33 | 1 | 6 | 82 | 4 | 78 |
| 5 | Mental Health Disorders | $£^{2} 20.25$ | 12 | 55 | 400 | 35 | 371 |
| 6 | Learning Disability | $£ 1.18$ | 1 | 4 | 15 | 3 | 12 |
| 8 | Problems of Vision | £2.20 | 0 | 2 | 29 | 1 | 28 |
| 9 | Problems of Hearing | £0.99 | 0 | 1. | 52 | 0 | 52 |
| 12 | Dental problems | £3.27 | 0 | a) | 59 | 0 | 59 |
| 14 | Skin | $£ 2.23$ | 2 | 4 | 13 | 5 | 8 |
| 15 | Musculo skeletal system | £. 4.11 | $3<$ | 15 | 203 | 10 | 193 |
| 20 | Poisoning and AE | $£ 1.05$ | 0 | 2 | 6 | 1 | 5 |
| 21 | Healthy Individuals | $¢_{6} 4.01$ | 0 |  | 3 | 0 | 2 |
| 22 | Social Care Needs | $\ldots 3.41$ | 1 |  | 0 | 0 | 0 |
| 23 | Other | $\mathrm{f}_{6} 5.88$ | 0 | 0 | 0 | 0 |  |
|  | All (23 PBCs) | $£ 80$ | - 295 | 1,337 | 4,367 | 859 | 3,509 |

### 5.8 Future research and improving estimates of the threshold

There are a number of ways in which this research could be usefully extended based on existing data and the information currently available, most of which have been discussed in previous sections of this chapter. Here we consider the scale of the evaluation problem in this context, examining what, in principle, would be required to resolve some of the key uncertainties discussed in Section 5.4, before a more detailed examination on of how additional routine data, greater access to existing data or data that are likely to become available might improve estimates of the cost per QALY threshold in the future. Recall from Section 4.1 and 5.4 that two important questions remain when attempting to translate the estimated proportionate effects on mortality due to a change in expenditure into a more complete measure of the health effects. These are: i) whether the health effects of a change in expenditure in one year should be restricted to one year or extend over a longer period; and ii) the extent to which any death averted by expenditure in one year returns an individual to the mortality risk of the general population matched for age and gender. The central or 'best' estimates presented in Chapter 4 and 5 are based on combining the conservative assumption that the health effects of changes in one year of expenditure are restricted to one year ${ }^{119}$ with the more optimistic assumption that any death averted by expenditure in one

[^35]year returns the individual to the mortality risk of the general population. ${ }^{120}$ The combination of assumptions that underpin the central estimates appear to be on balance conservative (see discussion in Section 5.4) and are certainty more credible than the implausibly pessimistic or optimistic assumptions that underpin the upper and lower bounds for the threshold that are also reported in Chapter 4 and 5. Key questions remain, however: why cannot routine data resolve some of these remaining uncertainties and what would be required to find a central estimate of the cost per life year or cost per QALY threshold only on econometric estimates rather than, in part at least, resting on judgments about the credibility of these alternative assumptions? ${ }^{121}$

A brief reiteration of the scale of this evaluation problem and the approaches to estimation that might be taken illustrates the quite profound difficulties and, therefore, the unavoidable need for explicit and accountable judgement and assumptions. ${ }^{122}$

## A longer and more complex lag structure

Of course, a longer and more complex lag structure exploiting the PBC panel data set (i.e., both cross section and time series observations) that is becoming available over time, could in principle at least, identify the effect of a change in expenditure taking place in year $t$ on health in years $t, t+1, \ldots, t+n$. However, the health effects in subsequent years would need to be isolated from the effects of change in expenditure also occurring in subsequent years (which would also have both immediate and lagged health effects). Depending on the length of time series data available it may be possible to specify and estimate a richer empirical model to account for the lagged health effects of past expenditure and of lagged expenditure effects of past health outcomes. ${ }^{123}$

Although this is not a problem of principle it does pose difficatties as there are very real limits to the current time series because: i) there are a limited number of observations in the cross section (152 PCTs); ii) the definition (and boundaries) of PCTs has changed and is due to change again; iii) there are a limited number of years of observation in the time series (especially if lags are long); and iv) as noted in Chapter 3 and 5, the quality of PBC reporting has changed oven time (recall that estimates from 2006 and 2007 PBC expenditure were not as comparable as 2007 and 2008). Nonetheless, as the panel data evolves over time there will be more opportunities to explore whether judgments about the duration of effects on mortality can be informed using the type of analysis presented in the report. Insofar as there are later lagged health effects this will tend to reduce the estimate of the cost per death averted and cost per life year and cost per QALY threshold.

In many respects the problem of duration of mortality effects is a relatively straight-forward one compared to the second issue of how changes in mortality (whether immediate or lagged) translate into life years. In principle, estimating the effect of change in expenditure on life years is really estimating the effect of changes on expenditure on the survival curves of the population at risk through membership of
mortality in one year due to a proportionate change in expenditure. This is likely to underestimate effects on mortalitt since expenditure that reduces mortality risk (or reduces the QALY burden of disease) for an individual in one year may well also reduce their risk (reduce QALY burden) over subsequent years; possibly over the whole of their remaining disease duration. Expenditure may also prevent disease in future patient populations. Therefore, total health effects will be underestimated and the cost per life year or QALY threshold will be overestimated.
120 Therefore, the years of life gained associated with each death averted are based on what would have been their life expectancy taking account of their age and gender (using life tables for the general population).
${ }^{121}$ It should be recognised that the purpose is to inform an assessment of the threshold for decisions that have not yet been made (i.e., prediction for decisions not yet made rather than a description of the past). Therefore, irrespective of the availability of evidence or the sophistication of analytic methods, the need for assumptions or scientific value judgements can never be avoided but only informed.
${ }^{122}$ The nature of prediction to inform decisions and combined with the reality of a forever unobserved counterfactual makes judgement unavoidable - see footnote above.
${ }^{123}$ For example, a more structural approach of estimating an outcome equation jointly with an expenditure equation, both with appropriately specified lag structures and controlling for unobserved PCT effects might be possible.

ICD codes that contribute to each PBC. Even if the issue of lags is set aside, and attention is restricted to mortality effects in the expenditure year, translating these effects into life years would require observations on the entire survival curve of the at-risk population. This poses two profound difficulties i) we would need detailed information about the members of the at-risk population (patient identifiers) and ii) sufficient time to follow up the entire cohort from expenditure change to death (also accounting for other changes that are likely to take place during that time). Even if these data were available and such heroic estimation was possible, any estimate would be so historic that it would be of limited policy relevance. This is not a problem unique to this research but remains a problem for all estimates of the years of life lost due to disease. It may be possible to use external, non-routine, historic data sources where patient identifiers are available to inform a judgement about whether changes in mortality in critieal ICD codes (e.g., respiratory) tend to return patients to mortality risks similar to those of the general population or not. If historic evidence suggests that they remain at higher mortality risk it might indicate the likely scale of over-estimation if life year effects are based on the mortality risk of the general population. However, this would not be without major problems of distinguishing causality from selection effects.

The evolving panel data do have another advantage that could be exploited in the future. Currently it is only cross sectional variation (i.e., between PCTs) that contributes to the estimates of outcome elasticities. This means that changes in expenditure that all PCTs tend to make together, that might have very large health effects (they all tend to invest in obviously valuable activities at the same time) or limited health effects (they all disinvest in some activities that are not valuable at the same time) may not be fully reflected in the current estimates. ${ }^{124}$ However, using variation in expenditure and outcome in both cross section and time series could more confidently pick up the full effects of simultaneous investment and disinvestment. The likely net effect on the overall threshold is unclear and will depend on whether PCTs tend to be more coordinated when investing in valuable activities (tending to reduce the threshold) or when disinvesting in ineffective ones (tending to increase the threshold). ${ }^{125}$

## Simultaneous estimation across PBCs

Although expenditure equations are estimated for all 23 PBCs and outcome equations for the 11 PBCs where there are sufficient mortality data, these are estimated separately; each accounting for other PBC expenditure and other PBC need (see Section 3.3). The correlations between expenditure and outcome elasticities within each PBC were also estimated by repeatedly re-sampling the data set and re-estimating expenditure and outcome elasticities (see B8.11 in Appendix B). Although the estimate of the overall threshold accounts for changes in expenditure across all 23 PBCs with health effects estimated in 11 and inferred in the others, it is possible that changes in expenditure in one PBC may have health effects in others. Although totaldeaths across all 23 PBCs are accounted for, unless the possible 'external' mortality effects in other PBCs happen to be associated with variation in expenditure in those PBCs then these health effects will not be reflected in the estimated outcome elasticities. This seems likely to underestimate the total health effects of changes in expenditure unless positive health effects are thought to be offset by expenditure in one PBC damaging health outcome in others (e.g., adverse events associated with treatment or other iatrogenic effects). ${ }^{126}$ To account properly for these possible effects would require estimating the interaction of changes in expenditure in each PBC on all the others while still accounting for possible endogeneity. Unfortunately, with only 151 observations in the cross section (PCTS), this type of simultaneous estimation is currently not feasible.

Recall that throughout Chapters 3, 4 and 5 we have not imputed health effects for PBC 23 (General Medical Services) on the grounds that the health effects of this type of expenditure will appear in ICD

[^36]codes that contribute to other PBCs. However, the health effects of this type of expenditure will only be reflected in the estimated outcome elasticities insofar as the variation in outcomes reported in other PBCs, due to variation in GMS expenditure, happens to be associated with variation in expenditure in those other PBCs. Therefore our approach to GMS expenditure is likely to be conservative with respect to overall health effects; tending to overestimate the cost per life year and cost per QALY threshold.

## Exogenous shocks and quasi experiments

One response to these difficulties would be to look for exogenous budgetary shocks to the whole health care system and then estimate the health effects of the shock at a macro level. In principle this is very attractive since it would avoid all the difficulties of endogeneity and identifying valid instruments, exploring sensitivity and structural uncertainty. If a complete measure of health outcome were available> at a health system level it would also avoid much of the complexity of working at a PBC and ultimately at ICD level.

Unfortunately there are a number of difficulties. Although the NHS budget is set eachyear through an essentially political process (so each year's change in budget might be regarded as an exogenous shock), insofar as public expenditure decisions are to some extent influenced by public sector performance then these apparent 'shocks' are endogenous in a very similar way to PCT expenditure decisions about particular PBCs but just at a higher level of aggregation. However, even if some arbitrary exogenous change to overall expenditure could be identified there are other serious difficulties. There is no comprehensive measure of outcome relevant to all NHS activities currently reported. This has two implications: i) the mortality data that are available is only relevant to approximately $36 \%$ of a change in overall expenditure (see Section 5.3); and ii) how mortality translates into life years and QALY depends critically on where those effects occur (the ICD codes that contribute to each PBC). In addition there are very good reasons why one would expect covariates (especially measures on need) and instruments to differ between different programmes of care. For all these reasons this research has focused on using routinely available data at its lowest level of aggregation.

By doing so we not only provide an estimates of a threshold based on a more complete measure of health effects, we are also able to indicate what type of health is effected and where they are most likely to occur. This provides a means to update estimates of the threshold should other aspects of social value be applied to measures of health or other aspects of social value be included in the future (e.g., consumption and other public expenditure effects). For example any 'weights' that might be assigned to different types of QALY gains or consumption and other public expenditure effects associated with health effects and the patient characteristics associated with ICD codes (e.g., QALY burden, years of life lost or other patient characteristics, such as age and gender) can be included in the current framework and a threshold reestimated for 'weighted' QALYs or, give an estimate of the consumption value of a QALY, a threshold benefit cost ratio that includes consumption as well as health effects.

Evolving Programme Budget Category Data
Each year offers another wave of PBC expenditure data which means that a potentially useful panel data set is developing. This offers some useful opportunities that have been described above. However, with only 151 PCTs in the cross section, there is a limit to how much of the remaining uncertainty might be resolved. The utility of this evolving panel will also be limited by the move to Clinical Commissioning Groups rather than PCTs as an important locus of expenditure decisions.

Of course it would also be useful to be able to observe PBC expenditure at a lower level of aggregation (ideally at ICD code) as this would avoid the assumption necessary to allocate PBC level effects to ICD codes based on either estimates of the size of the at risk population or the crude (unadjusted for covariates) contribution to variance in PBC expenditure. Since the only expenditure data that are available by ICD (and therefore PBC) for each PCT are HES based estimates of cost, the relevance of measures of contribution to variance in PBC expenditure depends on what proportion of PBC costs are accounted for by HES; which quite naturally differs by PBC. Greater disaggregation within PBCs would
be particularly useful as the examination of information routinely collected by PCTs was not particularly helpful in identifying what investment and disinvestments within a PBC explain the variation in PBC expenditure across PCTs (see Addendum C2 in Appendix C).

## Extending measures of health outcome

Currently the only routinely collected health outcome data that can be matched to expenditure by PBC category at PCT level is mortality. For this reason outcome equations could only be estimates for 11 of the 23 PBCs. As discussed in Sections 4.3 and 4.4, this represents only one aspect of health outcome and is not particularly relevant to many disease categories and much of the care that the NHS offers when the primary purpose is to improve health experience and quality of life rather than to extend survival. Therefore, the estimated proportionate effects of expenditure on the QALY burden of disease in these 11 PBCs (which only account for $36 \%$ of the change in overall expenditure, although $80 \%$ of the health effects) were used as a surrogate for proportionate effects in the others, i.e., assuming that the proportionate effects that can be observed will be similar to those that cannot (see Section-5.2).127

Of course, with access to a more complete measure of health outcome, which is routinely reported at PCT level and that can also be associated with PBC expenditure, it would be possible to use the same econometric methods to estimate the health effects of a change in expenditure across all PBCs, rather than imputing them in those PBCs where mortality is not the most relevant measure of health outcome.

The English NHS Patient Reported Outcomes (PROMs) programme was introduced in 2009 and routinely collects self-reported health status of patients receiving surgery for four elective procedures: knee and hip replacement, groin hernia repair, and varicose vein surgery. The data that are collected include both condition specific questions (the Oxford Hip Seore, Oxford Knee Score and the Aberdeen Varicose Vein score; no condition specific instrument is ayailable for hernia) as well as the generic instrument, the EQ-5D (both the EQ-5D profile, and the patient's global assessment of their health, the EQ-VAS. Patient-level data from the PROMs programme are freely available and can be linked to the HES database which provides a potential link to PBCs. Standardised reports on the PROMs data, including the average (case-mix adjusted) performance of providers, are regularly published by the NHS Information Centre, currently on a quarterly basis. Although currently offering very limited coverage for our purposes, there are plans to extend the PROMs programme in the future, with work underway or being planned around the potential use of PROMs in a wide range of long term conditions, primary care, in cancer survivorship, cardiovascular services, muscular skeletal, and cosmetic surgery.

In Appendix B, Section B8.8 we demonstrate how the econometric methods set out in Chapter 3 can be extended to these other non-mortality based outcome measures. EQ-5D utility scores (pre and post an operative procedure) from the PROMs programme are used to generate a non-mortality-based outcome measure, which we use to estimate our outcome model. Although the Department of Health does not report the number of patients undergoing an eligible operation by commissioner (PCT) it was possible to use the HES dataset to obtain this information. Routine reporting of procedure or intervention by commissioner in the PROMs data-set would seem a simple but important and valuable extension, especially as data are extended to primary care where HES cannot be used to substitute for this omission.

With data for both the average health gain per operation and the number of operations, we were able to estimate 'the health gain per head of population' for hip and knee replacements as defined above. This estimated outcome elasticity can then be used as an outcome measure for changes in expenditure in the 'problems of the musculoskeletal system' programme (i.e., PBC15) ${ }^{128}$. However, translating the short term impact of an intervention on quality of life, which can be estimated from PROMs data, into an estimate of the longer term effects on quality of life remains problematic.

[^37]Table B8.26 in Appendix B reports the estimated outcome equation for PBC15 (musculoskeletal system) using the PROMs based outcome measure. The result is intuitively plausible; an increase in expenditure improves health outcomes but, for a given spend, more need reduces the gain. The diagnostic statistics suggest that expenditure is endogenous and that the instruments are valid. They also suggest that the instruments are relevant and there is no evidence that the instruments are weak. Therefore, it is feasible to extend our modelling approach beyond those programmes with mortality outcomes should PROMs be extended more widely. Insofar as PROMs can contribute to a more secure estimate of the overall cost per QALY threshold in the future, the sensitivity analysis discussed in Section 5.3 starts to indicate where this type of information might be most useful.

Musculoskeletal is an important PBC, accounting for over $5 \%$ of a change on overall expenditure and almost $5 \%$ of the change in health outcomes. However, of those PBCs without mortality outcomes, it is PBC5 (Mental Health) that is most critical (see Table 5.2 in Section 5.3).
Measures of anxiety and depression are already routinely collected before, during and at the end of interventions as part of Improving Access to Psychological Therapies (IAPT), which is an NHS programme rolling out services across England offering interventions approved by NICEfor the treatment of depression and anxiety disorders. By March 2011 IAPT services were offered in 142 of 151 PCTs. A requirement of the programme is to complete the Patient Health Questionnaire-9 (PHQ9, a measure of depression) [107] and Generalised Anxiety Disorder Assessment 7 (GAD7, a measure of anxiety)[108]. Both of these disease-specific measures can be linked to SF20 and further work could, in principle, link these scores to EQ5D. This is a rich, valuable and evolving data-set which potentially provides much of the information required to extend the econometric modelling to the mental health PBC. The experience with PROMs data suggests that this would be feasible, and the analysis in Section 5.3 indicates that this could make a significant contribution to strengthening the assessment of the overall threshold. It would also contribute to an assessment of the COst effectiveness of this programme both nationally and by PCT, which would be of value in its own right. Unfortunately, despite the collection of these data for every patient encounter for a number of years, unlike PROMs, these data have not yet been made publically available. ${ }^{129}$ Of course, the services offered by the IAPT programme do not account for all the variation in expenditure in the mental health PBC. Nevertheless, access to data that have been and continue to be collected by practitioners and NHS patients, could provide estimates of changes in mental health outcomes due to changes in some types ofmental health expenditure, which would be a significant advance. ${ }^{130}$

## Incidence and duration of disease



Section 4.2 sets out the series of steps required to translate mortality effects into life years while taking account of competing risks or counterfactual deaths. This analysis used ONS data on deaths by age and gender in the ICD codes that contribute to each PBC, as well as life expectancies by age and gender for the general population. Some information was also required about the age and gender distribution of the population at risk in the ICD codes that contribute to each PBC (see Table 4.4 and 4.5). In Section 4.2.3 this was based on age and gender distribution of estimates of incidence from the WHO Global Burden and Disease (GBD) study. The same information was also used in Section 4.3.1 to adjust life years for the quality offife norms of the general population by age and gender. In Section 4.4.2 the measures of

129 These data have only been collated centrally since April 2012 despite IAPT sites collecting these data at individual patient encounters for many years. In April 2012, the IAPT data standard was approved by the NHS Information Standards Board as a nationally mandated data standard. Data is now collected centrally on a monthly basis from over 200 service locations. The first report on the quality of IAPT data was published in November 2012 but the quarterly IAPT data reports, which were scheduled to be released at the same time, do not appear not to have been made available. There appears to be a commitment to make the dataset publicly available during 2012/13, although the timing and details of what will be available (summaries or patient level data and whether it will include the waves of data collected since 2006) and who might have access (commissioners, service providers or independent researchers) remains unclear (see www.iapt.nhs.uk and www.ic.nhs.uk/iapt).
${ }^{130}$ Similar difficulties will arise, however, when translating the observed impact of a therapy on quality of life, before and immediately after the intervention, into longer term effects.

QALY burden of disease also used information about the duration as well as incidence of disease from the same WHO study. These estimates, published in 2008, were based on 2004 UK data and proved to be the best available source of this type of information given the resources available for this research. However, the WHO study has recently been updated with the findings first publically presented in December 2012.[109] The methodology of the new study as well as sources of information used have been much improved and any subsequent research on the threshold could integrate these new and improved estimates.

However, the WHO GBD study is not the only potential source of information about estimates of incidence of disease by age and gender and disease duration across all the ICD codes that contribute to the 23 PBCs. For example, the General Practice Research Database (GPRD) (recently renamed the Clinical Practice Research Datalink (CPRD)) contains over 3 million active patient records drawn from approximately 400 primary care practices in the UK. GPRD is jointly funded by the NHS National Institute for Health Research (NIHR) and the Medicines and Healthcare products Regulatory Agency (MHRA).The database has clinical and prescription data and can provide information to support) pharmaco-vigilance (indication, utilization, and risk/benefit profiles of drugs) and formal pharmacoepidemiologic studies, including information on demographics, medical symptoms, therapy (medicines, vaccines, devices), and treatment outcomes.

Although this research was not funded to purchase access to GPRD data, we were able to examine a sample which comprised of $22,313,086$ rows/patient-ICD10 events (3 digit) representing 4,229,910 patients with data on diagnosis of diseases observed between 1 Jan 2006 and 24 June 2011 (see Addendum C1 in Appendix C). Although GPRD data could, in principle, provide the type of information required the difficulties faced and the interpretation of the sample of data in the form available to us meant that it was not directly useful. The particular problems faced included: i) read rather than ICD codes reported in the data set, although mapping is and was possible; ii) being able to identify when an episode of disease ended; iii) estimating duration of disease from the sample of data when observations were censored by the limited years of data available to us; and iv) confidently identifying incident patients in diseases of longer duration despite two years of wash out prior to extracting the sample. GPRD is quite clearly a rich and valuable data set. However, our experience suggests that, to make best use of these data, specialist knowledge and experience of these data is really needed as well as access to a much larger sample than we were able to acquire with the limited resources available. Therefore, although GPRD could well help to improve estimates of incidence by age and gender and duration of disease, it would require additional well-resourced research including excess to specialist expertise and experience with this particular data set.

## References

1. NICE, Guide to the methods of technological appraisal. ref: N1618, 2008.
2. Health, D.o., A New Value-Based Approach to the Pricing of Branded Medicines - a Consultation. . London Department of Health, 2010.
3. Health, D.o., A New Value-Based Approach to the Pricing of Branded Medicines: Government Response to Consultation. London: Department of Health, 2011.
4. Claxton, K., et al., V alue based pricing for NHS drugs: an opportunity not to be missed? British Medical Journal, 2008. 336(7638): p. 251-254.
5. Claxton, K., M.J. Sculpher, and S. Carroll, Value-based pricing for pharmaceuticals: Its role, specification and prospects in a newly devolved NHS. CHE Research Paper 60.
http://www.york.ac.uk/media/che/documents/papers/researchpapers/CHERP60.pdf. York: Centre for Health Economics, University of York, 2011.
6. Culyer, A., et al., Searching for a threshold, not setting one: the role of the National Institute for Health and Clinical Excellence. J Health Serv Res Policy, 2007. 12(1): p. 56-8.
7. McCabe, C., K. Claxton, and A.J. Culyer, The NICE cost-effectiveness threshold - What it is and what that means. Pharmacoeconomics, 2008. 26(9): p. 733-744.
8. Committee, H.o.C.H., NICE: First report of the Health Committee 2007-2008. HC27-I. London: Stationery Office, 2008.
9. NICE, Guide to the methods of technological appraisal. ref: N0514, 2004.
10. Devlin, N. and D. Parkin, Does NICE bave a cost-effectiveness threshold and what other factors influence its decisions? A binary choice analysis. Health Economics, 2004. 13(5):p. 437-452.
11. Gafni, A. and S. Birch, Guidelines for the adoption of new technologies - a prescription for uncontrolled growth in expenditures and how to avoid the problem. Canadian Medical Association Journal, 1993. 148(6): p. 913-917.
12. Williams, A., What could be nicer than NICE? OHE annual lecture (book available), 2004.
13. Health, D.o., A new value-based approach to the pricing of branded medicines: A consultation. 2010.
14. NICE, First report of the Health Committee 2007-2008. HC27-I. London: Stationery Office, 2008.
15. Weinstein, M. and R. Zeckhauser, Critical ratios and efficient allocation. Journal of Public Economics, 1973. 2: p. 147-157.
16. Stinnett, A.A. and A.D. Paltiel, Mathematical programming for the efficient allocation of health care resources. Journal of Health Economics, 1996. 15(5): p. 641-653.
17. Epstein, D., et al., Efficiency, equity and budgetary policies: informing decisions using mathematical programming. Medical Decision Making, 2007. 27: p. 128-37.
18. Abelson, P., The value of life and health for public policy. Economic Record, 2003. 79: p. S2-S13.
19. Bobinac, A., et al., WVillingness to pay for a Quality-Adjusted Life-Year: The individual perspective. Value in Health, 2010. 13(8): p. 1046-1055.
20. Byrne, M.M., K. O'Malley, and M.E. Suarez-Almazor, Willingness to pay per quality-adjusted life year in a study of knee osteoarthritis. Medical Decision Making, 2005. 25(6): p. 655-666.
21. Dolan, P., et al., QALY maximisation and people's preferences: a methodological review of the literature. Health Economics, 2004. 14(2): p. 197-208.
22. Green, C. and K. Gerard, Exploring the social value of health-care interventions: a stated preference discrete choire experiment. Health Economics, 2009. 18(8): p. 951-976.
23. Groot, W. and H.M. van den Brink, The value of health. Bmc Health Services Research, 2008. 8.
24. Gyrd-Hansen, D., Willingness to pay for a QALY. Health Economics, 2003. 12(12): p. 1049-1060.
25. Gyrd-Hansen, D., Willingness to pay for a QALY - theoretical and methodological issues. Pharmacoeconomics, 2005. 23(5): p. 423-432.
26. Johnson, F.R. and M. Backhouse, Eliciting stated preferences for health-technology adoption criteria using paired comparisons and recommendation judgments. Value in Health, 2006. 9(5): p. 303-311.
27. King, J.T., et al., Willingness to pay for a quality-adjusted life year: Implications for societal health care resource allocation. Medical Decision Making, 2005. 25(6): p. 667-677.
28. Lieu, T.A., et al., Willingness to pay for a QALY based on community member and patient preferences for temporary health states associated with Herpes Zoster. Pharmacoeconomics, 2009. 27(12): p. 1005-1016.
29. Mason, H., M. Jones-Lee, and C. Donaldson, Modelling the monetary value of a qaly: a new approach based on uk. data. Health Economics, 2009. 18(8): p. 933-950.
30. Luis Pinto-Prades, J., G. Loomes, and R. Brey, Trying to estimate a monetary value for the QALY. Journal of Health Economics, 2009. 28(3): p. 553-562.
31. Shiroiwa, T., et al., International survey on willingness-to-pay (wtp) for one additional qaly gained: what is the threshold of cost effectiveness? Health Economics, 2010. 19(4): p. 422-437.
32. Yaesoubi, R. and S.D. Roberts, A game-theoretic framework. for estimating a bealth purchaser's willingness-to-pay for bealth and for expansion. Health Care Management Science, 2010. 13(4): p. 358-377.
33. Polsky, D., Does willingness to pay per quality-adjusted life year bring us closer to a useful decision rule for costeffectiveness analysis? Medical Decision Making, 2005. 25(6): p. 605-606.
34. Smith, R.D. and J. Richardson, Can we estimate the 'social' value of a QALY? Four core issues to resolve. Health Policy, 2005. 74(1): p. 77-84.
35. Hirth, R.A., et al., Willingness to pay for a quality-adjusted life year: In search of a standard. Medical Decision Making, 2000. 20(3): p. 332-342.
36. Haninger, K. and J. Hammitt, Willingness to pay for Quality-Adjusted Life Years: empiricalinconsistency between cost-effectiveness analysis and economic welfare theory. OECD, 2006.
37. Baker, R., et al., Weighting and valuing quality-adjusted life-years using stated preference methods: preliminary results from the Social V alue of a QALY Project. Health Technology Assessment, 2010. 14(27): p. 1-+.
38. Rawlins, M.D. and A.J. Culyer, National Institute for Clinical Excellence and its value judgments. British Medical Journal, 2004. 329(7459): p. 224-227.
39. Rawlins, M.D., D. Barnett, and A. Stevens, Pharmacoeconomics: NICE's approach to decision-making. British Journal of Clinical Pharmacology, 2010. 70(3): p. 346-249.
40. Tappenden, P., et al., A stated preference binary choice experiment to explore NICE decision making. Pharmacoeconomics, 2007. 25(8): p. 685-693.
41. NICE, Appraising Life Extending End-of-Life Treatments. London: NICE, 2009.
42. NICE, Draft Guide to the Methods of Technology Appraisal. Iondon: NICE, 2012.
43. Committee, H.o.C.H., NICE response to the first report of session 2007-2008. HC550. London: Stationery Office, 2008.
44. Committee, H.o.C.H., The government's response to the Health Select Committee's first report of session 2007-08 on the National Institute for Health and Clinical Excellence. Cm7331. London: Stationery Office, 2008.
45. Braithwaite, R. and M. Roberts, $\$ 50,000$ per QALY: inertia, indifference, or irrationality? presented at: Annual Meeting of the Society for Medical Decision Making, 2004.
46. Birch, S. and A. Gafni, The biggest bang for the buck or bigger bucks for the bang: the fallacy of the costeffectiveness threshold. Journal of health services research \& policy, 2006. 11(1): p. 46-51.
47. Collier, J., Parliamentary revien askes NICE to do better still. British Medical Journal, 2008. 336(56-57).
48. Towse, A., Should NICE's threshold range for cost per QALY be raised? Yes. British Medical Journal, 2009. 338.
49. Appleby, J., et al., Searching for cost effectiveness thresholds in the NHS. Health Policy, 2009. 91(3): p. 239-245. (V)
50. Hughes, D.A. and R.E. Ferner, New drugs for old: disinvestment and NICE. British Medical Journal, 2010.340.
51. Buxton, M., How much are health-care systems prepared to pay to produce a QALY? European Journal of Health Economic, 2005. 6(4): p. 285-28.
52. Elshaug, A., et al., Identifying existing health care services that do not provide value for money. The Medical Journal of Australia, 2009. 190(5): p. 269-73.
Laupacis, A., et al., How attractive does a new technology bave to be to warrant adoption and utilization tentative guidelines for using clinical and economic evaluations. Canadian Medical Association Journal, 1992. 146(4): p. 473-481.
53. Birch, S. and A. Gafni, Cost-effectiveness ratios - in a league of their own. Health Policy, 1994. 28(2): p. 133-141.
54. Drummond, M., G. Torrance, and J. Mason, Cost-effectiveness league tables - more barm than good. Social Science \& Medicine, 1993. 37(1): p. 33-40.
55. Appleby, J., N. Devlin, and D. Parkin, NICE's cost effectiveness threshold - How high should it be? British Medical Journal, 2007. 335(7616): p. 358-359.
56. Martin, S., N. Rice, and P. Smith, The link between bealth care spending and bealth outcomes: evidence from English programme budgeting data. CHE Research Paper 24, 2007a.
57. Martin, S., N. Rice, and P. Smith, Further evidence on the link between bealth care spending and bealth outcomes in England. CHE Research Paper 32, 2007b.
58. Martin, S., N. Rice, and P. Smith, The link between bealth care spending and bealth outcomes for the new English Primary Care Trusts. CHE Research Paper 42, 2008b.
59. Martin, S., N. Rice, and P. Smith, Does health care spending improve health outcomes? Journal of Health Economics, 2008a: p. 826-842.
60. Martin, S. and P. Smith, How good at commissioning bealth are English primary care trusts? A preliminary statistical analysis. Report to the Health Foundation, 2009.
61. Martin, S., N. Rice, and P. Smith, Panel data estimates of the link between health care spending and health outcomes for English Primary Care Trusts. Mimeo, 2010.
62. Martin, S., N. Rice, and P. Smith, Comparing costs and outcomes across programmes of health cares Health Economics, 2012: p. 316-337.
63. Cochrane, A.L., A.S.S. Leger, and F. Moore, Health service "input" and mortality "output" in developed countries (Reprinted from Journal of Epidemiology and Community Health vol 32, pg 200-205, 1968). Journal of Epidemiology and Community Health, 1997. 51(4): p. 344-348.
64. Young, F.W., An explanation of the persistent doctor-mortality association. Journal of Epidemiology and Community Health, 2001. 55(2): p. 80-84.
65. St Leger, S., The anomaly that finally went away? Journal of Epidemiology and Community Health, 2001. 55(2): p. 79-79.
66. Nolte, E. and M. McKee, Does health care save lives? The Nuffield Trust, London, 2004.
67. Gravelle, H.S.E. and M.E. Backhouse, International cross-section analys sis of the determination of mortality. Social Science \& Medicine, 1987. 25(5): p. 427-441.
68. Cremieux, P.Y., P. Ouellette, and C. Pilon, Health care spending as deterninants of health outcomes. Health Economics, 1999. 8(7): p. 627-639.
69. Nixon, J. and P. Ulmann, The relationship between bealth care expenditure and health outcomes. Evidence and caveats for a causal link. The European journal of health economics : HEPAC : health economics in prevention and care, 2006. 7 (1):p. 7-18.
70. Bokhari, F.A., Y. Gai, and P. Gottret, Government health expenditures and health outcomes. Health Econ, 2007. 16(3): p. 257-73.
71. Moreno-Serra, R. and P.C. Smith, The Effects of Health Coverage on Population Outcomes: A CountryLevel Panel Data analysis. Results fornDevelopment Institute, Washington D.C., Working Paper, 2011.
72. Health, D.o., NHS finance manual. December 2005 edition. See bttp:/ / www.dh.gov.ule/assetRoot/04/13/18/26/04131826.pdf. 2005a.
73. Smith, P.C., N, Rice, and R. Carr-Hill, Capitation funding in the public sector. Journal of the Royal Statistical Society Series a-Statistics in Society, 2001. 164: p. 217-241.
74. Office, N.A. Good governance report: review of programme budgeting. London: NAO, 2008.
75. Appleby, Harrison, T., Foot, C., Smith, A. and Gilmour, S. , Explaining variations in primary care trusts' spending on cancer services. The King's Fund, London, 2011.
76. Lakhani, A., H. Olearnik, and D.e. Eayres, Compendium of clinical and health indicators: data definitions and user guide for computer files. London, NCHOD, 2006.
77. Batm, C.F., M.E. Schaffer, and S. Stillman, ivreg2: Stata module for extended instrumental variables/2SLS, GMM and AC/HAC, LIML and k-class regression. See http://ideas.repec.org/c/ boc/ bocode/s425401.html. 2010.
78. Shea, J., Instrumental relevance in multivariate linear models: a simple measure. Review of Economics and Statistics, 1997. 79: p. 348-352.
79. Stock, J.H. and M. Yogo, Testing for weak instruments in linear IV regression. NBER Technical Working Paper 284, 2002.
80. Ramsey, J.B., Tests for specification errors in classical linear least-squares regression analysis. Journal of the Royal Statistical Society Series B-Statistical Methodology, 1969. 31(2): p. 350-\&.
81. Pesaran, M.H. and L.W. Taylor, Diagnostics for IV regressions. Oxford Bulletin of Economics and Statistics, 1999. 61(2): p. 255-+.
82. Durbin, J., Errors in variables. Review of the International Statistical Institute, 1954. 22: p. 23-32.
83. Health, D.o., PCT recurrent revenue allocations exposition book: 2009/10 and 2010/11. Department of Health, London, 2009.
84. Health, D.o., Recurrent resource allocations: 2006/07, 2007/08. Department of Health, London, 2005c.
85. Health, D.o., Personal communications. 2012.
86. Curtis, L., Unit costs of health and social care 2011. PSSRU, University of Kent., 2011.
87. Conley, T.G., C.B. Hansen, and P.E. Rossi, Plausibly exogenous. Review of Economics and Statistics, 2012. 94(1): p. 260-272.
88. Small, D.S., Sensitivity analysis for instrumental variables regression with overidentifying restrictions. Journal of the American Statistical Association, 2007. 102(479): p. 1049-1058.
89. Dixon, J., et al., A person based formula for allocating commissioning funds to general practices in England: development of a statistical model. British Medical Journal, 2011. 343.
90. Wailoo, A.D., S. Tosh, J, The incorporation of health benefits in cost utility analysis using the eq-5d: report by the decision support unit. School of Health and Related Research, University of Sheffield, 2010.
91. Dolan, P., et al., A social tariff for EuroQol: results from a UK general population survey, CHE discussion paper 138, University of York, 1995.
92. Currie, C.J., et al., The routine collation of health outcomes data from hospital treated subjectsin the Health Outcomes Data Repository (HODaR): descriptive analysis from the first 20,000 subjects. Value in health : the journal of the International Society for Pharmacoeconomics and Qutcomes Research, 2005. 8(5): p. 581-590.
93. Cohen, J.W., et al., The Medical Expenditure Panel Survey: a national health information resource. Inquiry, 1996. 33(4): p. 373-89.
94. Claxton, K., et al., Informing a decision framework for when NICE should recommend the use of health technologies only in the context of an appropriately designed programme of evidence development. Health Technology Assessment, in press 2012.
95. Claxton, K., The irrelevance of inference: a decision-making approach to the stochastic evaluation of health care technologies. J Health Econ, 1999. 18(3): p. 341-64.
96. Jackson, C., et al., A Framework for Addressing Structural Uncertainty in Decision Models. Medical Decision Making, 2011. 31(4): p. 662-674.
97. Soares, M.O., et al., Methods to elicit experts' beliefs over uncertain quantities: application to a cost effectiveness transition model of negative pressure wound therapy for severe pressure ulceration. Stat Med, 2011. 30(19): p. 2363-80.
98. Maynard, A. and A. Street, Seven jears of feast, seven years of famine: boom to bust in the NHS? BMJ, 2006. 332(7546): p. 906-908.
99. Trading, O.o.F., The Pharmaceutical Price Regulation Scheme. An OFT market study. VBP Report 2007. London: OFT, 2007.
100. Claxton, K., et al., Discounting and decision making in the economic evaluation of health care technologies. Health Economics, 2011. 20:2-15. DOI: 10.1002/hec.1612.
101. Paulden, M. and L. Claxton, Budget allocation and the revealed social rate of time preference for health. Health Economics 2011. 24 MAR: DOI: 10.1002/hec. 1730 .
102. NICE, TA237: Ranibizumab for the treatment of diabetic macular oedema. 2011.
103. NICE, Macular oedema (diabetic) - ranibǐ̧umab (rapid review of TA237): appraisal consultation document. 2012.
104. Novartis, Single technology appraisal (STA) manufacturer submission: Lucentis ${ }^{\circledR}$ (ranibizumab) for the treatment of visual impairment due to diabetic macular oedema (DMO). 2010.
105. Broome, J., Trying to value a life. Journal of Public Economics, 1978. 9: p. 91-100.
106. Kroenke, K., R.L. Spitzer, and J.B. Williams, The PHQ-9: validity of a brief depression severity measure. J Gen Intern Med, 2001. 16(9): p. 606-13.
107. Spitzer, R.L., et al., A brief measure for assessing generalized anxiety disorder: the GAD-7. Arch Intern Med, 2006. 166(10): p. 1092-7.
108. WHO, Global Burden of Disease Study 2010. The Lancet, 2012.

## Appendix A

## SYSTEMATIC REVIEW OF THE LITERATURE ON THE COST-EFFECTIVENESS THRESHOLD

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## A. Systematic review approach

## A1. Introduction

In the initial stages of this systematic review it became clear that the "traditional" method of conducting systematic searches of existing literature on the topic of the cost-effectiveness threshold would be insufficient to deal with the requirements of this particular study. Here we refer to the "traditional" method as the practice of finding key terms and Medical Subject Headings (MeSH) that most accurately capture the range of literature relevant to the topic, while attempting to include as few irrelevant studies as is possible (making use of programs such as Medline).

The main weaknesses of using such an approach for a systematic review of this topic is that it requires a pre-existing knowledge of the terms used and topics covered in the current literature. This process has always required a degree of expertise (and luck) as to the strategy taken, including both knowledge of the literature to find likely search terms and skill in the construction of the strategies. The implications of excluding a single key term are potentially equivalent to ignoring vast areas of the literature. In addition, the traditional approach relies on key terms existing that suitably encapsulate the relevant literature. Finding common terms used in literature with potential relevance to the cost effectiveness threshold was found to be a significant problem as many relevant topics were not specifically aimed issues relating to the NICE cost-effective threshold (for example the Martin et al. Papers (1-3) which provide a precursor to this project). In addition, due to the wide range of coverage of topics such a "threshold" and "costeffective", any attempts at a systematic review would be either excessively large or result in a clearly limited snap-shot of the existing literature.

As a result a pragmatic approach was taken to the identification of relevant papers, one of "pearl growing" which can be defined here as the use of existing collections of studies to identify additional relevant parts of the literature. The approach uses a pool of "initial pearls" to grow the literature both through references and citations until all relevant papers have been discovered. This approach therefore relies on the expertise of the authors of the exiting literature to populate the pool of studies rather than the searcher's potentially limited knowledge.

While this approach of "pearl growing" was significantly limited by the existing software available and has a time consuming element, it represents an approach that corrects for many of the failings of traditional searches for topics that share the characteristics of the cost-effectiveness threshold.

## A2. Systematic Review Methods

The "pearl growing" method of systematic review can be characterised into five steps for the identification of relevant papers.

1. Identification and extraction of "initial pearls".

- "Initial pearls" were indentified through consultation with researchers with experience of the cost-effectiveness threshold literature. Fourteen initial pearls were indentified through this process. These publications were chosen for their wide ranging coverage of the topic as well as their anticipated significance

Extraction of Citations and References from "initial pearls".

- Citations: Web of Knowledge was selected to perform the citation searches. The reason for this selection was in part due to expert advice from an information specialist as well as brief and non-systematic investigations of citation results from a range of alternative software packages.
- References: Web of Knowledge was also used for the collection of papers' references.
- Both citations and references were exported into an EndNote library for the purpose of collection and further analysis (exclusion of repeats, title searching and review of the abstracts).

3. Identification of further "pearls" from cited and referenced papers.

- Once citations and references of the "initial pearls" had been collected, they were subjected to a set of investigations to identify further "pearls".
- Papers were excluded based on whether the titles or abstracts suggested the paper contained information on five topics of interest. These topics had been previously identified given the objectives of the project and from a review of the "initial pearls" and included papers were classified by whether they could inform,
i) introduction to the cost-effectiveness threshold topic and policy context,
ii) discussion and debate around the current value use of the threshold,
iii) potential methods suggested to find a suitable threshold value,
iv) specific values proposed,
v) the use of individual and societal valuations of health gains to inform the value of the threshold

4. Repetition of citation and reference searches.

- The process was then repeated for the "pearls" identified in step 3.
- This process was repeated until no new "pearls" were discovered by additional iterations.

5. Manual search of references

- To ensure as complete a search had been conducted as possible a retrospective manual search of all of the "pearls"" references was conducted. Any potentially relevant references not discovered previously (most likely due to a mix of user error and limitations with the software used) were added to the analysis at the relevant step and further pearl growing methods applied to them to ensure completeness of results.


## A3. Systematic review results

The "pearl growing" method of systematic review revealed 76 papers deemed relevant. The results from each stage of the process are reported in Figure A.1. The figure highlights that after four iterations no new relevant papers were identified by the systematic process.


See next page

Key:
Solid line-included
references
Dotted line- excluded references one in
stages


Figure A.1: graph showing process results from pearl growing systematic review

## B. Review of literature

## A4. Introduction and policy context

Due to the broad range of context which the relevant literature covers it is necessary to break down the literature review into several topics, these will be discussed independently. The 76 papers (see section E for all of these papers) identified by the systematic review were defined into five different categories:

1. literature covering the introduction to the cost-effectiveness threshold topic and policy context, 2. discussion and debate around the current value use of the threshold,
2. potential methods suggested to find a suitable threshold value,

These categories were chosen to reflect the broad range of relevant topics and areas of discussion covered by the cost-effectiveness threshold literature. It should be noted that the majority of the literature identified by the literature review fell into the first and last categories, with very few covering multiple categories sufficiently completely to be discussed in more than one section. The final category will only be discussed briefly as it can be seen as a separate, unrelated approach to the threshold required for purposes of decision making by the National Institute for Health and Clinical Excellence (NICE).

The majority of papers ( 34 of the 76 papers discovered) identified in the literature review could be characterised as introducing the idea of a cost-effectiveness threshold (these consist of the very early literature pre-dating NICE) or discussing the policy context through the years (4-37). This section will characterise the main areas of discussion in the literature and briefly describe the key parts of the literature development.

## A4.1. Definition of the cost effectiveness threshold

An important place to start is the consideration of how the literature has defined the cost-effectiveness threshold. This is important to analyse in the review as not only is it worth ensuring that a good definition has been presented; but it also allows us to assess uyhether the existing literature uses a definition that is both consistent and accurate.

One of the earliest definitions of something resembling the modern interpretation of the cost-effective threshold comes from Weinstein and Zeckhauser(36). Their paper identifies a "critical ratio" between monetary costs and a measure of health gains. This critical ratio was argued to represent 'a cut-off point for allocation' of an activity in a budget constrained public sector entity ((36), p.1.).

A similar, more recent approach to define the threshold is that taken by Devlin (35) where the author considered a hypothetical budget constrained health care sector, with a perfectly informed decision maker who only considers the cost per QALY of health technologies. Assuming perfect information, the decision maker is able to rank all of the potential health care activities based on their cost per QALY. A decision maker will implement as many of the relatively low cost per QALY activities as possible until the budget 11 used up. Eventually a point will be reached where society is not willing to pay for a further marginal increase in QALYs and would rather the funding used on other consumption. The cost per QALS at which this cut-off occurs can be described as the cost-effectiveness threshold as it represents the switching point between an activity being funded and not. As the budget is assumed to be fully responsive, any new technologies with a cost per QALY below this threshold will be funded in the future.

## A4.2. NICE and the cost effectiveness threshold

The use and valuation of a cost-effectiveness threshold by NICE has been controversial. Williams (37) highlighted three events that may be argued to have particularly muddied the water. Firstly, NICE did not set a threshold value by the government at the time of its inception in 1999. This meant that NICE was obliged to come up with a de novo estimate fairly rapidly. Through his set of discussions with NICE,

Williams stated that at the point of inception NICE came up with a value of 'roughly $£ 30 \mathrm{k}$ per QALY, plus or minus $£_{5 \mathrm{k} \text { depending on the specific circumstances’ ((37), p.7.) }}^{\text {( }}$

The second event which Williams refers to was NICE's initial resistance to acknowledging that any form of threshold value existed. Following analyses such as Towse and Pritchard (35) and Devlin(38) investigating previous NICE decisions and inferring an implicit threshold, NICE began to publish details of its approach to an ICER threshold. The major step was the 2004 Guide to the Methods of Technology Appraisal (30) that provided these details, although the definition of the $£ 20,000$ to $£ 30,000$ threshold range may be considered loose and open to interpretation. While the 2004 Guide was one of the first official references to the threshold, Sir Michael Rawlins did state at the 2001 NICE Annual General Meeting that the Institute would 'need to be very clear in its reasons for supporting technologies with cost-effectiveness ratios higher than $£ 30,000$ per QALY' (Littlejohns in (35)).

Williams' final event is the often quoted $£ 20,000$ to $£ 30,000$ threshold range having never been scientifically justified. Authors such as Rawlins and Culyer(39) have argued that there has never been an empirical basis for the values or any definitive meaning behind the range. They therefore argued that the threshold should not be the only tool for NICE to draw conclusions about new technologies.

## A4.3. The threshold as a range

The idea of such a threshold range has been part of the literature for some time. Kaplan and Bush (24) considered the idea of a less abrupt approach than that suggested by Weinstein and Zeckhauser(36). Kaplan and Bush (24) investigated a set of early Medicare adopfion decisions and presented broad criteria of acceptance based on a set of threshold ranges in terms ofcost per additional well year. These were defined as $<\$ 20 \mathrm{k} /$ well year (cost effective), $\$ 20 \mathrm{k}-\$ 100 \mathrm{k}$ (possibly controversial but justifiable), $>\$ 100 \mathrm{k}$ questionable when compared with other expenditure). However, the authors noted that a $\$ 100 \mathrm{k}$ cut-off was not relevant to the policy decisions at the time and that all results would need significant future investigation. Similarly Laupacis(26) presented five "grades of recommendation" for decisions about technological reimbursement in Canada.

The conclusions of both of these papers can be represented graphically by Figure A.2, which is also described or presented in much of the literature (see (39), Littlejohn in (35), (29), (38)). This graph represents the probability of rejection of new technology as a function of technology's ICER. The graph clearly shows two points of inflection (A and B in Figure A.2), these two points represent an interpretation of the lower and upper bounds of a cost-effectiveness threshold range.


Figure A.2: Probability of rejection with a 'soft' cost-effectiveness threshold

The literature often makes use of the terms "soft" and "hard" when referring to the threshold. The term "soft" is often used in a similar way to the threshold "range" (alternatively Akehurst's "smudge" (Akehurst in (35)). While the underlying idea is the same, a "soft" threshold has also been used to refer to a single threshold. For example, McCabe et al. (29) argued that it is both feasible and probably desirable to use a single threshold rather than a range, as the threshold should represent the point beyond which factors other than cost effectiveness are considered. This approach would suggest that all new technologies with an ICER below the threshold should receive funding (regardless of their impact on other factors such as equity of health). It is, however, unclear from this paper what the implications are for technologies with an ICER beyond the single threshold value.

In contrast, a "hard" threshold represents the situation where the ICER valuation is the sole relevant variable in an adoption decision, as demonstrated in Figure A.3(15). It is an important point that if a "hard" threshold is set no other factors can be considered in the decision maker's consideration of a new technology. The difference between a "hard" and a "soft" threshold is therefore largely based on whether the ICER reflects all considerations. As, assuming the decision maker is optimising health, it should represent the most effective allocation of a health care budget but cannot account for any equity concerns (such as the severity of the condition, unmet need and orphan diseases) that are not included in the calculation of the ICER. Authors such as Dolan et al. (40) have demonstrated that a "hard" threshold may not be able to suitably reflect the non-linearity of social or political values of QALYs to factors such as quality and length of life and for those with worse health prospects or dependents.


Figure A.3: graph showing a "hard cost" effectiveness threshold

## A4.4. What does the threshold represent?

Two broad lines of thought have developed on what the threshold represents, social willingness to pay (WTP) and shadow pricing. $(13,29,36,37,41,42)$ The key difference between the two is the budget that should be considered by those accepting or rejecting health technologies. The social WTP approach (usually implicitly) assumes that the budget of the health care sector is flexible to the value of health gains determined by society. So in this case it is the value society places on the health benefits (for example in QALYs) generated by new health care programmes and technologies is estimated first, and then the health care budget is the sum of society's wrillingness to pay for all treatments. In other words, the threshold is set exogenously with no reference to a budget constraint.

In contrast, the shadow pricing approach takes the budget as given (at least beyond the control of those who determine the cost-effectiveness threshold) (13, 29). The threshold is, therefore, endogenous based on the services currently provided within the system. When a new programme or technology is accepted into the system and imposes an additional cost onto the budget, the only way to meet those costs is to remove or down-scate existing services which will incur opportunity costs in terms of population health. Hence the threshold tepresents the ICER of the least cost effective existing service covered by the budget. In principle, it is this service which is removed to fund a new programme or technology. In practice, a range of criteriaj lisely to be used to identify appropriate services for displacement to make room in the budget for new interventions.

In the OK the main source of debate about which of these concepts of the threshold is the correct one lies in NICE's remit. Authors such as Culyer (13) have discussed NICE's position as a "searcher" or a "setter" of the threshold. The distinction between these two roles is that a threshold "searcher" does not set a threshold with the motivation of maximising social welfare under the assumption of a flexible NHS budget, but instead investigates the threshold value that is appropriate given current NHS activities and the fixed budget as set down by Parliament.

Much of the literature on this topic is founded in the discussion of the correct constitutional role of NICE, the potential negative implications of setting a threshold and the feasibility of identifying displaced activities. In 2007, Culyer et al. (13) argued that it is not appropriate for NICE to be characterised as a threshold setter. The authors argued that the setting of a threshold would effectively imply that NICE sets the NHS budget. The setting of the NHS budget, they highlight, is the constitutional responsibility of

Parliament, not NICE. Hence the paper argues that NICE should concern themselves with being threshold "searchers", seeking to identify 'an optimal threshold ICER, at the ruling rate of expenditure, that is consistent with the aim of the health service to maximise population health' ((13), p.4).

In a similar vein Appleby et al. (43) concluded that the threshold used by NICE should be consistent with the decisions made by local commissioners within the NHS. This is important given that NICE provides little guidance to the NHS regarding interventions suitable for disinvestment to release the funding necessary to cover the new technologies it recommends. If the threshold is set too high NICE may well accept new technologies which are less cost effective than the services which local commissioners displace to fund those technologies. Conversely, if the threshold is set too low, NICE is likely to reject services that are cost effective relative to existing services delivered from the NHS budget. The authors conclude that, in the short term, NICE have to act as a threshold "searcher" to ensure continuity in the NHS.

Alternative arguments have been put forward which reject the idea of NICE as a threshold "searcher". Firstly, some authors (such as Gafni and Birch $(17,18)$ ) have made the case that an implicit threshold has the potential to lead to spiralling inflation if new cost effective technologies are funded without sufficient disinvestment. However, McCabe et al.(29) argued that Culyer's characterisation of the NICE threshold could overcome this challenge if it were regularly reviewed so as to be flexible over time to changes in the NHS budget and the productivity of the sector, and if the threshold for newactivities with a nonmarginal budget impact was greater than those with a marginal impact. The issue of the inflationary pressure of a threshold is discussed further below.

Another concern raised about Culyer et al.'s characterisation of the NICE threshold is that of Towse et al. (44). They argue that a lack of knowledge of the true opportuity cost of new activities makes us unable to identify the value of those activities being displaced and, therefore, it is impossible for NICE to "search" for a threshold relating to activities displaced at the margin. The issue of the difficulty of identifying current activities at the margin in terms of cost-effectiveness will be dealt with later in this chapter.

## A4.5. Factors considered by NICE other than the comparison of the ICER and threshold

As was discussed in the section the threshold range, the suitable threshold approach is dependent on the policy context around it, specifically if the comparison of the ICER with the threshold represents the only relevant piece of information that informs an adoption decision (a "hard" threshold) or if it is simply one of many factors considered ("soft" threshold). In the case of the UK, NICE has openly stated the ICER of a technology is not the sole consideration of the committee in its adoption decisions (30).

Both NICE and a number of other authors have provided overviews of the other factors that are considered by NICE in the adoption decision, these are provided in Table A.1.

Table A.1: table showing factors other than ICER considered by NICE

| NICE (30, 31) | Rawlins et al. (45) | Tappenden et al. (34) | Devlin et al. (38) |
| :--- | :--- | :--- | :--- |
| Uncertainty of variables | Severity of illness | Uncertainty of the <br> ICER | Uncertainty of the <br> ICER |
| Availability of <br> comparators | End of life treatment | Availability of <br> comparators | Burden of disease |
| Clinical priorities (as set <br> by Secretary of State) | Stakeholder opinion | Severity of illness |  |
| Clinical need | Innovation | *age not significant |  |
| Availability of resources | Population <br> characteristics <br> (disadvantaged and <br> children) |  |  |
| Innovation |  |  |  |
| Disease characteristics <br> and population size |  |  |  |
| Wider social costs and <br> benefits |  |  |  |
| Length of benefit |  |  |  |

This table suggests that the threshold is only one consideration to decisions makers at NICE. However, , in principle, these other types of benefits could be added to health benefits and compared to potential treatments for displacement which also have wider social benefits. In other words, this wider set of considerations relating to the benefits of new technologies should arguably also be reflected in the threshold. ${ }^{1}$

## A4.6. Multiple thresholds

Similarly some have argued for using different thresholds for different situations (29, 47). The two main cases for using different thresholds are the size of the budgetary impact, or depending on if the decision represents an investment in additional activities or a disinvestment in current activities.

The topic of different thresholds for different budgetary impacts of a proposed technology has received very little analytical attention from the literature. McCabe et al. (29) argue that technologies with a large budgetary impact should be evaluated against a lower threshold than those with a relatively small impact. The reason for this is a large budgetary impact will require a greater displacement of current activities (assuming a fixed oyerall budget); this may result in displacement of non-marginal activities which may be associated with lower ICER than those at the margin.

Several authors have suggested the use of different threshold values depending on whether the decision represents an investment in additional activities or a disinvestment in current activities. O'Brien et al.'s 2002 (47) paper considers the difference in willingness to accept monetary compensation to forgo a health care program and willingness to pay for the same benefit and link it to the cost effectiveness threshold. This paper came from the perspective of the threshold representing social preferences rather than the shadow price of a fixed budget constraint and highlights that from a traditional 'welfarist' economics standpoint; a greater threshold value for disinvestment may be welfare maximising. Similarly both Devlin et al. (38) and Speight et al. (48) have suggested a threshold for disinvestment of currently

[^38]performed activities could be lower than for new activities, however, neither present any methodology for calculating the weight of a disinvested activity.

This is in contrast with the view that cost effectiveness analysis guides the decisions of health systems with the objective of maximising some measure of health benefit subject to a budget constraint. Hughes et al. (21) has argued that differential threshold with respect to investment and disinvestment would result in sub-optimal levels of population health. This is because a new technology that would improve health may be rejected under a policy of having different thresholds for investment and disinvestment but not if the threshold values were the same. The authors argue that this failure to maximise population health represents an avoidable inefficiency not related to the aim of the health care sector to maximise health and thus making the case for a single threshold value for disinvestment and investment. This point can be seen as a further case for the shadow price approach as opposed to the social WTP perspective as it highlights that, given a fixed NHS budget, the social WTP approach will not lead to a maximisation of health.

## A4.7. The need for an independent threshold panel

Related to the discussion over the correct role of NICE in determining a suitablecost effectiveness threshold for the NHS is the literature on the potential for an independent threshold panel. Such a panel has been characterised in a similar manner to the Monetary Policy Committee (the setters of Bank of England's interest rate who act independently of the Government of the United Kingdom), as an independent committee responsible for the setting and updating of the cost-effectiveness threshold used by NICE.

The papers covering this topic are consistent in their call for an independent threshold panel, with no papers identified arguing against it. The main case provided in the literature for an independent setter is the removal of political influence; Claxton et al. (10) argue that political influence may drive the threshold up as politicians seek to use the threshold as a means to encourage investment by pharmaceutical companies. Williams (37) suggests that NICE is biased in the setting of a threshold, as its political connections mean a higher threshold makes it is more popular with the "sellers" (the author defines sellers as not only the pharmaceutical industry but health care professionals and patient groups) by allowing more technologies to be aproved. Similarly papers by Appleby et al. (43) and Raftery et al. (49) call for the creation of an independent threshold setter. 2008 Health Select Committee (50) recommended that a body independent of NICE should be established to set and review the threshold. However, it is unclear if such a body would also be independent of political influence or just of the NICE structure.

## A4.8. Arguments against the use of a cost effectiveness threshold

A number of authors have argued against the use of a threshold. As mentioned earlier authors such as Gafni and Birch $(17,18)$ have suggested that the threshold approach risks leading to spiralling increases in inflationary pressures on health care spending, and present an alternative approach based on the use of league tables of cost-effectiveness. The reason, they argue, is that there is no guarantee that the activities displaced are less cost-effective than those new technologies imposing cost on the health system budget. This observation is coupled with the expectation of authors such as Cohen et al. (11) that pharmaceutical firms will inevitably price their drugs so as to ensure the ICER of their proposed new technology is sufficiently close to the threshold to ensure adoption and thereby gain maximum producer surplus. This observation implies that providers such as the NHS may be forced to pay above market costs of new technologies by revealing their maximum willingness to pay, in the form of the threshold. In addition the point raised in McCabe et al. (29) that the threshold should be adjusted regularly over time to ensure its efficiency seeks to address both of these arguments.

Other authors such as Eichler et al. (15) have raised and debated the issues around the theoretical base for the cost-effectiveness threshold, namely the assumption of perfect divisibility of healthcare programs, constant returns to scale and constant marginal opportunity costs $(15,17,19,51,52)$.

Bridges et al. (53) argues that a unique threshold value imposes impractical assumptions in the case of the US health care sector, and fails to account for supply and demand side variations in the market. As an alternative the authors propose a series of thresholds that reflect regional, dynamic, budgeting and general methodological differences. They conclude that the case for abandoning a fixed threshold outweighs those for keeping one in the US and that any threshold should vary across payer, over time, in the true budget impact of interventions and in the measurement of the effectiveness of interventions. This argument has clear links to the argument for shadow pricing of the threshold rather than the social WTP approach, as the shadow price approach is based on the view that the threshold is determined by budget and current efficiency which can be seen to differ over time and across payers. The unresolved issue here is the degree to which different sub-groups (e.g. by region or budget) require different threshold values

## A4.9. Identification of activities under the threshold

An important part of the literature is the discussion around the identification of activities with an ICER greater than the proposed threshold. The importance of this discussion stems from the requirement of new activities to displace current activities that are at the margin of what is cost-effectife. If it is not possible to identify these activities separately from others then threshold analysis is methodologically flawed, as the funding of a new activity may impact on an activity with an ICER aBoye the proposed threshold.

Most literature on this topic focuses on the importance of identifying activities to be displaced rather than the process and feasibility of doing so. For example, Hughes et al. (21) and McCabe et al. (29) highlight the implications of inconsistent displacement on geographic variations in health care provision and that the lack of consistency in the displacement process undercuts, the use of a single cost effectiveness threshold for the evaluation of new technologies. Similarly Buxton et al. (54) suggests that, in order to fully appreciate the opportunity cost of the implementation of a new technology, we must have a clear knowledge of those activities displaced at the cost effectiveness margin.

Few authors have sought to develop methods to identify the activities that should be displaced to free-up budget for new more cost-effective activities. Elshâg et al. (16) outlines a set of criteria for the identification of existing, potentially non-cost-effective practices which could then be further assessed to assess their cost-effectiveness using health technology assessment. The criteria suggested include factors such as: new evidence on safety; efficacy or cost-effectiveness, geographic variation that have become apparent since technology adoption, clinical heterogeneity in the clinical procedure, and technological development.

## A5. The current value of the threshold

Since it became evident that decision making bodies such as NICE are using (more or less explicit) costeffectiveness thresholds, there has been a significant level of debate over its appropriate value (35, 37-39, 43-45, 47-49,53-58). In this section we will present three areas of the debate:

The lack of empirical basis to the current value
Arguments over the value being generally too high or too low
If and how the threshold should change over time

## A5.1. Lack of empirical base to the current value

Since NICE made it clear that it uses an explicit threshold (30) there has been little hiding the lack of evidential justification behind the $£ 20,000$ to $£ 30,000$ range. Indeed the Health Select Committee (50) heard (during their enquiry into NICE in 2008) that the NICE threshold has no basis in hard science. Similarly Appleby (43) noted that "the uncomfortable truth is that NICE's threshold has no basis in either theory or evidence."

Similarly the US value of $\$ 50,000$ per QALY, which is often cited as the cost-effectiveness threshold relevant to resource allocation decisions in that country, is often attacked for its lack of empirical founding $(20,23,32,55)$. Some have suggested that the US figure is rooted in the cost-effectiveness of Hospital Renal Dialysis (20), although why this makes it suitable for use more generally is unclear.

## A5.2. The threshold changing over time

Another concern of current NICE practice is the apparent lack of change in the threshold value used since the body's inception. Many authors have argued that factors such as the NHS budget, price inflation, technological developments in the NHS and the discount rate applied to economic evaluations $(20,24,25,59)$ have all changed since the first use of the cost-effectiveness threshold. As such, the threshold should have changed to reflect this fact. Braithwaite et al. (55) sought to demonstrate the impact of budget and technological growth on the optimal threshold. By creating a computersimulation of the US Medicare system, the authors were able to demonstrate the impact of these factors. While there is no doubt in the literature that the NICE threshold should potentially change over time ${ }^{2}$ no papers have been identified which model the impact of any changes on the threshold.

Both Ubel et al. (57) and Raftery (49) discuss the principles behind the directional change the threshold should take over time. Ubel et al. (57) have argued that the optimal threshotd value needs to fall over time assuming medical innovation continues at roughly its current rate. Raftery (49) has noted that, in real terms, the threshold has been falling since 1999 as, in order to stay constant in real terms, it should have increased given inflation (up $40 \%$ in the time period) and increased NHS spending (up $90 \%$ ). The authors argue that this decline in the threshold should have been observed in the value used by NICE in decision making. They describe the suggestion of a rise in the threshold being linked to the observed growth of the NHS budget over the last decade as "audacious" (49). It is unclear to what extent the authors disagree with this interpretation of NHS efficiency as a releyant factor affecting the optimal threshold.

## A5.3. Value generally too high or low

The majority of the debate over the current use of the threshold in the UK (and elsewhere) has been centred on whether the current value is toolow or too high. The papers that will be discussed in this section focus on the general discussion of necessary directional change in the value rather than the presentation of a specific value; the latter is discussed in more detail in the following section on the proposed values of the threshold in the literature.

Vernon et al. (58) presented an analysis of the implications of the threshold being above or below its optimum value in terms of signals to the companies involved in research and development of new medical products. The authors concluded that if the threshold is set too low (below the economic value of the health benefit) it will result in research and development investment levels that are too low relative to their economic value (at the margin). The reason for this lies in a lack of returns to investments for the pharmaceutical companies. However, in the isolated case of the threshold relevant to the NHS (a small propertion of the world pharmaceutical market), the impact of changes to the threshold on the international pharmaceutical market equilibrium is unknown but likely to be small.

Similarly, thresholds set too high (above the economic value of the health benefit) will result in inefficiently high levels of research and development spending, such that the health care provider is funding projects that do not have a sufficient impact on social welfare.

The literature that argued the threshold is too high in the UK can be broadly characterised into three key papers. Alan Williams (37) made the case that, intuitively, the threshold should not be significantly greater

[^39]than the GDP per capita (roughly $£ 18,000$ in the UK in 2004). He made the case that, while it may be possible to provide a lot of the population with health care when the threshold is above the GDP per capita, it is not possible to provide health care for much of the population without imposing great hardship on those expected to foot the bill (the tax payer or government debt).

Secondly, Raftery (49) argued that, while the UK threshold has been historically too high, it does not need reducing as the real value has decreased since 1999 due to inflationary pressure and increases in the NHS budget. He also suggests that recent policies implemented by NICE, such as greater weight being given to the benefits of treatment accruing to patients at the end of their life, need to be offset by reductions in the threshold for all other treatments for expenditure to remain within the NHS budget. Finally, Raftery cites the opportunity cost analysis of trastuzumab (4) which showed that more cost-effective oncology services were being sacrificed to fund trastuzumab in breast cancer. This result suggests directly that, in some cases at least, the threshold value is too high.

Work by Martin et al. (1,3) investigated the cost per life year saved in a selection of the 23 programme budgeting categories used in the NHS; these results are presented in Table A.2. It is important to note that these results are presented as the cost per life year gained rather than the cost per QALY of the least cost-effective current activity. The authors and others have used these results to argue that the threshold used by NICE may be too high (28). Similarly, Collier's (12) report of the Health Select Committee suggests that the threshold used by NICE is higher than that used by PCTs.

Table A.2: table showing cost per life year gained results of Martin et al.(1-3) papers

| Programme budgeting category | cost per life year gained |  |
| :---: | :---: | :---: |
|  | 2005/6 data | 2004/5 data |
| Cancer | £13,137 | £13,931 |
| Circulation problems | £8,426 < | £7,979 |
| Respiratory problems | £7,397 < - | N/A |
| Gastro-intestinal problems | £18,999 | N/A |
| Diabetes | £26,453 | N/A |

In contrast, a range of authors have argued that the current NICE threshold is too low. Both Speight et al. (48) and Towse (44) argued that the inclusion of wider social costs/benefits and full consideration of social willingness to pay for additionat health gains show that the threshold should be significantly larger. Both cite recent NICE work by Mason et al. $(56,60)$ which suggested the threshold should be between $£^{6} 30,000$ and $£ 75,000$ per QALY based on attempts to model a willingness to pay based value of a QALY based on observations of the value of avoiding a statistical fatality. Similarly in the US Ubel et al. (57) have argued that, if inflation and willingness to pay valuations are taken into account, the relevant threshold in the US shoutd be closer to $\$ 200,000$ per QALY that the regularly cited $\$ 50,000$.

Those analyses which conclude the UK and US thresholds should be significantly higher have, at the core of their argument, the assumption that the respective health care budget is fully capable of responding to society's willingness to pay for additional health gains.

## 6. Potential methods for threshold estimation

There are broadly three approaches that can be taken to determine the threshold value (43, 54): social WTP, shadow pricing of the budget constraint and non analytical approaches such as expert elicitation. This project is concerned with the latter approach to estimating the cost effectiveness threshold. This is entirely consistent with the remit of the NHS in general and NICE in particular - they do not set the NHS budget but have to allocate those finite resources appropriately.

## A6.1. Papers seeking to elicit social WTP and non-analytical approaches

The majority of the literature that has presented a proposed value for the threshold (in the UK, US and elsewhere) has done so using valuation methods based on willingness to pay for an additional health benefit (40, 59-76). However other approaches have been suggested. For example the World Health Organisation's (WHO) 2002 report (77) suggested that interventions costing less than three times GDP per capita for each DALY averted represent good value for money.

Lee et al. (78) sought to update the US "dialysis standard" often claimed to be the base of the US Medicare threshold (20). The authors present a valuation of $\$ 129,090$ per QALY based on current dialysis practice in the US. Finally in an appendix to their edited book, Towse et al. (35) provide an interesting set of results drawn from a set of participants to the associated workshop (the majority of which were health economists). The participants were asked to anonymously record their view on what threshold NICE should apply. Eighteen responses were recorded with the average of all responses being $£ 29,000$ per QALY.

## A6.2 Papers considering the shadow price of the budget constraint

The systematic review only identified four different papers by three different authors that suitably fell into the category of shadow pricing of the budget constraint.

Williams (37) suggested investigating the cost effectiveness of NHS interventions that represent the majority of the budget (he speculated that some 300 interventions accounted for about $90 \%$ of the cost incurred by the NHS). The purpose of this would be to identify current NHS activities that might not be cost-effective. He acknowledged the implausibility of conducting full technological appraisals on such a large number of interventions (estimating this would take 10 years, at which point it would be necessary to re-evaluate the initial appraisals), and thus suggested relying on expert opinion and existing patient data to speed up the process.

While Williams recommendations related to identifying current interventions with a high cost per QALY as the basis for disinvestment, there is the potential to take this approach further and use it for a method to determine the cost effectiveness threshold even down to the level of a local decision maker. This was attempted by Appleby et al. (79) who conducted a feasibility experiment into the estimation of the appropriate NHS threshold by examining decision making in the NHS at a local level. The authors propose a structured model considering new technology's cost per weighted QALY gain in a table of all existing services. In an attempt to test the feasibility of this model they conducted interviews with senior NHS staff as well as investigating information on public health to construct a list of healthcare services introduced or discontinued in 2006/7. The authors found that it was feasible to identify decisions and to make the important step of estimating their cost-effectiveness; however, they noted that any attempts to fully evaluate sufficient decisions as to estimate a threshold would require a detailed understanding of the understanding of the decision structure at a local level as well as a significant number of observations.

The other key papers seeking to develop and implement methods for estimating the NHS threshold were those of Martin et al. (1-3). They aimed to establish a link between health care spending and health outcomes in the NHS after having adjusted for the need of the patient population. They made use data
around the observed mortality at PCT level in the NHS alongside data expenditure data on health care across 23 programmes of care based on ICD010 disease categories. As has been mentioned earlier in this chapter these papers present the cost per life year of a range of programme budgeting categories, however, the key result of these papers is that it is possible to make use of existing data to determine such valuations for current NHS interventions. The authors concluded that while their results are highly limited and do not present a single cost per QALY estimate for the optimal threshold they can "inform the decisions of NICE on whether their current threshold for accepting new technologies is set at an appropriate level" ( $p .37$ ). These studies are the precursor of analyses presented in this report, and further details can be found in Appendix B and in Chapter 3 of the main report.

In the area of the efficient allocation of healthcare it is also important to note the contribution of the earlier mathematical papers such as Stinnett and Paltiel (41) who outlined mathematical techniques to approach the problem through the use of a mixed integer programming approach. While there approach differs from the interpretation of the threshold as used in this study it represented an important step in the evaluation of the methodology of seeking to solve the optimisation problem apparent in healthcare.

## C. Conclusion

This systematic review of the literature surrounding the cost-effectiveness (threstold has highlighted the significant range and diversity of the literature. Despite the international and mature nature of the literature there are significant differences in the suggested methods torepresent a cost-effectiveness threshold. The main areas of debate relevant to this report have revolved around the role of NICE as a "searcher" or "setter" of the threshold ( 13,29 ). While some authors have implicitly argued for NICE to fulfil a role of a threshold "setter" by suggesting method of elicitation of social WTP valuations of a QALY, death or life year (40, 59-76) the literature of most relevance to this research has sought to consider estimation methods consistent with its role as a "searcher"( $1-3,43$ ).

## References

1. Martin S, Rice N, Smith P. The link between health care spending and health outcomes: evidence from English programme budgeting data. CHE Research Paper 24. 2007a.
2. Martin S, Rice N, Smith P. Further evidence on the link between health care spending and health outcomes in England. CHE Research Paper 32. 2007b.
3. Martin S, Rice N, Smith P. The link between health care spending and health outcomes for the new English Primary Care Trusts. CHE Research Paper 42. 2008b.
4. Barrett A, Roques T, Small M, Smith R. How much will Herceptin really cost? Br Med J. 2006;333:1118-20.
5. Birch S, Gafni A. Cost-effectiveness ratios - in a league of their own. Health Policy. 1994;28(2):133-41.
6. Birch S, Gafni A. The biggest bang for the buck or bigger bucks for the bang: the fallacy of the cost-effectiveness threshold. Journal of health services research \& policy. 2006;11(1):46-51.
7. Brock DW. How much is more life worth? Hastings Center Report. 2006;36(3):17-9.
8. Brouwer W, van Exel J, Baker R, Donaldson C. The new myth - The social value of the QALY. Pharmacoeconomics. 2008;26(1):1-4.
9. Chambers JD, Neumann PJ, Buxton MJ. Does medicare have an implicit cost-effectiveness threshold? Med Decis Mak. 2010;30(4):E14-E27.
10. Claxton K, Lindsay AB, Buxton MJ, Culyer AJ, McCabe C, WalkerS, et al. Value based pricing for NHS drugs: an opportunity not to be missed? Br Med J. 2008;336(7638):251-4.
11. Cohen J, Looney W. Re: how much is life worth: cetuximab, non-small cell lung cancer, and the $\$ 440$ billion question. Journal of the National Cancer Institute. 2010;102(15).
12. Collier J. Parliamentary review asks NICE to do better still.Br Med J. 2008;336(56-57).
13. Culyer A, McCabe C, Briggs A, Claxton K, Buxton M, Akehurst R, et al. Searching for a threshold, not setting one: the role of the National Institute for Health and Clinical Excellence. J Health Serv Res Policy. 2007;12(1):56-8.
14. Drummond M, Torrance G, Mason J. Cost-effectiveness league tables - more harm than good. Soc Sci Med. 1993;37(1):33-40.
15. Eichler HG, Kong SX, Gerth WG, Mavrồ P, Jonsson B. Use of cost-effectiveness analysis in health-care resource allocation decision-making: How are cost-effectiveness thresholds expected to emerge? Value in Health. 2004;7(5):518-28.
16. Elshaug A, Moss J, Littlejohns P. al. e. Identifying existing health care services that do not provide value for money. The Medical Journal of Australia. 2009;190(5):269-73.
17. Gafni A, Birch S. Guidelines for the adoption of new technologies - a prescription for uncontrolled growth in expenditures and how to avoid the problem. Can Med Assoc J. 1993;148(6):913-7.
18. Gafni A, BirchS. Incremental cost-effectiveness ratios (ICERs): The silence of the lambda. Soc Sci Med. 2006;62(9):2091-100.
19. GarberAA, Phelps C. Economic foundations of cost-effectiveness analysis. Journal of Health Economics. 1997;16:1-31.
20. Grosse SD. Assessing cost-effectiveness in healthcare: history of the $\$ 50,000$ per QALY threshold.Expert Rev Pharmacoecon Outcomes Res. 2008;8(2):165-78.
21. Hughes DA, Ferner RE. New drugs for old: disinvestment and NICE. Br Med J. 2010;340.
22. Johannesson M, Meltzer D. Some reflections on cost-effectiveness analysis. Health Econ. 1998;7(1):1-7.
23. Johnson FR. Einstein on willingness to pay per QALY: Is there a better way? Med Decis Mak. 2005;25(6):607-8.
24. Kaplan R, Bush J. Health-related quality of life measurement for evaluation research and policy analysis. Health Psychology. 1982;1(1):61-80.
25. Laufer F. Thresholds in cost-effectiveness analysis - More of the story. Value in Health.

2005;8(1):86-7.
26. Laupacis A, Feeny D, Detsky AS, Tugwell PX. How attractive does a new technology have to be to warrant adoption and utilization - tentative guidelines for using clinical and economic evaluations. Can Med Assoc J. 1992;146(4):473-81.
27. Mauskopf J, Rutten F, Schonfeld W. Cost-effectiveness league tables - Valuable guidance for decision makers? Pharmacoeconomics. 2003;21(14):991-1000.
28. Maynard A, Bloor K. The future role of NICE. Br Med J. 2010;341.
29. McCabe C, Claxton K, Culyer AJ. The NICE cost-effectiveness threshold - What it is and what that means. Pharmacoeconomics. 2008;26(9):733-44.
30. NICE. Guide to the methods of technological appraisal. ref: N0514. 2004.
31. NICE. Guide to the methods of technological appraisal. ref: N1618. 2008.
32. Polsky D. Does willingness to pay per quality-adjusted life year bring us closer to a useful decision rule for cost-effectiveness analysis? Med Decis Mak. 2005;25(6):605-6.
33. Rascati KL. The $\$ 64,000$ question- What is a quality-adjusted life-year worth? Clinical Therapeutics. 2006;28(7):1042-3.
34. Tappenden P, Brazier J, Ratcliffe J, Chilcott J. A stated preference binary choice experiment to explore NICE decision making. Pharmacoeconomics. 2007;25(8):685-93.
35. Towse A, Pritchard C, Devlin N. Cost-effectiveness thresholds: economic and ethicat issues. Office of Health Economics: Kings Fund. 2002.
36. Weinstein M, Zeckhauser R. Critical ratios and efficient allocation. Journal of Public)Economics. 1973;2:147-57.
37. Williams A. What could be nicer than NICE? OHE annual lecture (book ayailable). 2004.
38. Devlin N, Parkin D. Does NICE have a cost-effectiveness threshold and what other factors influence its decisions? A binary choice analysis. Health Econ. 2004;13(5):437-52.
39. Rawlins MD, Culyer AJ. National Institute for Clinical Excellenceand its value judgments. Br Med J. 2004;329(7459):224-7.
40. Dolan P, Shaw R, Tsuchiya A, Williams A. QALY maximisation and people's preferences: a methodological review of the literature. Health Econ. 2004;14(2):197-208.
41. Stinnett AA, Paltiel AD. Mathematical programmingfor the efficient allocation of health care resources. Journal of Health Economics. 1996;15(5):641-53.
42. Epstein D, Chalabi Z, Claxton K, Sculpher MI. Efficiency, equity and budgetary policies: informing decisions using mathematical programming. Med Decis Mak. 2007;27:128-37.
43. Appleby J, Devlin N, Parkin D. NICE's cost effectiveness threshold - How high should it be? Br Med J. 2007;335(7616):358-9.
44. Towse A. Should NICE's threshold range for cost per QALY be raised? Yes. Br Med J. 2009;338.
45. Rawlins MD, Barnett D, Stevens. A. Pharmacoeconomics: NICE's approach to decision-making. British Journal of Clinical Pharmacology. 2010;70(3):346-249.
46. Health Do. A new value-based approach to the pricing of branded medicines: A consultation.
2010.
47. O'Brien BJ, Gertsen K. Willan AR, Faulkner LA. Is there a kink in consumers' threshold value for cost-effectiveness in health care? Health Econ. 2002;11(2):175-80.
48. Speight J, Reaney M. Wouldn't it be NICE to consider patients' views when rationing health care? Br Med J. 2009;338.
49. Raftery/J. Should NICE's threshold range for cost per QALY be raised? No. Br Med J. 2009;338.
50. NICE. First report of the Health Committee 2007-2008. HC27-I. London: Stationery Office.
2008.
51. Sendi P, Briggs A. Affordability and costeffectiveness: decision-making on the costeffectiveness plane. Health Econ. 2001;10:675-80.
52. Sendi P, Al M. Revisiting the decision rule of costeffectiveness analysis under certainty and uncertainty. Soc Sci Med. 2003;57:969-74.
53. Bridges JFP, Onukwugha E, Mullins CD. Healthcare rationing by proxy cost-effectiveness analysis and the misuse of the $\$ 50000$ threshold in the US. Pharmacoeconomics. 2010;28(3):175-84.
54. Buxton M. How much are health-care systems prepared to pay to produce a QALY? European Journal of Health Economic. 2005;6(4):285-28.
55. Braithwaite R, Roberts M. $\$ 50,000$ per QALY: inertia, indifference, or irrationality? presented at: Annual Meeting of the Society for Medical Decision Making. 2004.
56. Mason AR, Drummond MF. Public funding of new cancer drugs: Is NICE getting nastier? European Journal of Cancer. 2009;45(7):1188-92.
57. Ubel PA, Hirth RA, Chernew ME, Fendrick AM. What is the price of life and why doesn't it increase at the rate of inflation? Arch Intern Med. 2003;163(14):1637-41.
58. Vernon JA, Goldberg R, Golec J. Economic evaluation and cost-effectiveness thresholds signals to firms and implications for R\&D investment and innovation. Pharmacoeconomics. 2009;27(10):797806.
59. Hirth RA, Chernew ME, Miller E, Fendrick AM, Weissert WG. Willingness to pay for a qualityadjusted life year: In search of a standard. Med Decis Mak. 2000;20(3):332-42.
60. Mason H, Jones-Lee M, Donaldson C. Modelling the monetary value of a qaly: a new approach based on uk data. Health Econ. 2009;18(8):933-50.
61. Abelson P. The value of life and health for public policy. Economic Record. 2003;79:S2-S13.
62. Baker R, Bateman I, Donaldson C, Jones-Lee M, Lancsar E, Loomes G, et al. Weighting and valuing quality-adjusted life-years using stated preference methods: preliminary results from the Social Value of a QALY Project. Health Technology Assessment. 2010;14(27):1-+.
63. Bobinac A, van Exel NJA, Rutten FFH, Brouwer WBF. Willingness to pay for a Quality-

Adjusted Life-Year: The individual perspective. Value in Health. 2010;13(8):1046-55.
64. Byrne MM, O'Malley K, Suarez-Almazor ME. Willingness to pay per quality-adjusted life year in a study of knee osteoarthritis. Med Decis Mak. 2005;25(6):655-66.
65. Green C, Gerard K. Exploring the social value of health-care interventions: a stated preference discrete choice experiment. Health Econ. 2009;18(8):951-76.
66. Groot W, van den Brink HM. The value of health. Bmc Health Services Research. 2008;8.
67. Gyrd-Hansen D. Willingness to pay for a QALY. Health Econ. 2003;12(12):1049-60.
68. Gyrd-Hansen D. Willingness to pay for a QALY - theoretical and methodological issues. Pharmacoeconomics. 2005;23(5):423-32.
69. Haninger K, Hammitt J. Willingness to pay for Quality-Adjusted Life Years: empirical inconsistency between cost-effectiveness analysis and economic welfare theory. OECD. 2006.
70. Johnson FR, Backhouse M. Eliciting stated preferences for health-technology adoption criteria using paired comparisons and recommendation judgments. Value in Health. 2006;9(5):303-11.
71. King JT, Tsevat J, Lave JR, Roberts MS. Willingness to pay for a quality-adjusted life year: Implications for societal health care resource allocation. Med Decis Mak. 2005;25(6):667-77.
72. Lieu TA, Ray GT, Ortega-Sanchez IR, Kleinman K, Rusinak D, Prosser LA. Willingness to pay for a QALY based on community member and patient preferences for temporary health states associated with Herpes Zoster. Pharmacoeconomics. 2009;27(12):1005-16.
73. Luis Pinto-Prades J, Loomes G, Brey R. Trying to estimate a monetary value for the QALY. Journal of Health Economics. 2009;28(3):553-62.
74. Shiroiwa T, Sung Y-K, Fukuda T, Lang H-C, Bae S-C, Tsutani K. International survey on willingness-to-pay (wtp) for one additional qaly gained: what is the threshold of cost effectiveness? Health Econ. 2010;19(4):422-37.
75. Smith RD, Richardson J. Can we estimate the 'social' value of a QALY? Four core issues to resolve. Health Policy. 2005;74(1):77-84.
76. Yaesoubi R, Roberts SD. A game-theoretic framework for estimating a health purchaser's willingness-to-pay for health and for expansion. Health Care Management Science. 2010;13(4):358-77.
77. WHO The World Health report: reducing risks, promoting healthy life. 2002.
78. Lee GP, Chertow GM, Zenios SA. An empiric estimate of the value of life: updating the renal dialysis cost-effectiveness standard. Value in Health. 2009;12(1):80-7.
79. (—) Appleby J, Devlin N, Parkin D, Buxton M, Chalkidou K. Searching for cost effectiveness thresholds in the NHS. Health Policy. 2009;91(3):239-45.

## Papers discovered by the literature review

Note: Not all of these papers are referenced in this Appendix and some references used were not discovered through the systematic review

## Initial Pearls

1. Appleby, J., N. Devlin, and D. Parkin, NICE's cost effectiveness threshold - How high should it be? BRITISH MEDICAL JOURNAL, 2007. 335(7616): p. 358-359.
2. Appleby, J., et al., Searching for cost effectiveness thresholds in the NHS. HEALTH POLICY, 2009. 91(3):P. 239-45.
3. Bridges, J.F.P., E. Onukwugha, and C.D. Mullins, Healthcare Rationing by Proxy Cost-Effectiveness Analysis and the Misuse of the $\$ 50000$ Threshold in the US. PHARMACOECONOMICS, 2010.28(3): p. 175-184.
4. Culyer, A., et al., Searching for a threshold, not setting one: the role of the National Institute for Healtb and Clinical Excellence. JOURNAL OF HEALTH SERVICES RESEARCH \& POLICX, 2007. 12(1): p. 56-8.
5. Devlin, N. and D. Parkin, Does NICE have a cost-effectiveness threshold and what other factors influence its decisions? A binary choice analysis. HEALTH ECONOMICS, 2004. 13(5): p. 437-452.
6. Gafni, A. and S. Birch, Incremental cost-effectiveness ratios (ICERs): The silence of the lambda. Social Science \& Medicine, 2006. 62(9): p. 2091-2100.
7. McCabe, C., K. Claxton, and A.J. Culyer, The NICE cost-effectiveness threshold - What it is and what that means. PHARMACOECONOMICS, 2008. 26(9): p. 733-744.
8. Raftery, J., Should NICE's threshold range for cost per QALY be raised? No. BRITISH MEDICAL JOURNAL, 2009. 338.
9. Towse, A., Should NICE's threshold range for cost per QAL Be raised? Yes. BRITISH MEDICAL JOURNAL, 2009. 338.
10. Braithwaite et al. What Does the Value of Modern Medicine Say About the $\$ 50,000$ per QualityAdjusted Life-Year Decision Rule? Medical Care, 2008, 46 (4) pp349
11. Grosse, S, Assessing cost-effectiveness in healthcare: history of the $\$ 50,000$ per QALY threshold. Expert Review of Pharmacoeconomics \& Outcomes Research, 2008, 8 (2) pp 165
12. Rawlins, M.D. and A.J. Culyer, National Institute for Clinical Excellence and its value judgments. British Medical Journal, 2004. 329(7459): p. 224-227.
13. Chambers, Neumann and Buxton Does Medicare Have an Implicit Cost-Effectiveness Threshold? Medical Decision Making, 2010, 30 (4) pp E14-27
14. Chambers, J.D., P.J. Neumann, and M.J. Buxton, Does Medicare Have an Implicit CostEffectiveness Threshold? Medical Decision Making, 2010. 30(4): p. E14-E27.

## Step 1 Results

1. Brouwer, W., et al., The new myth - The social value of the QALY. Pharmacoeconomics, 2008. 26(1): p. 14.
2. Claxton, K., et al., V alue based pricing for NHS drugs: an opportunity not to be missed? British Medical Journal, 2008. 336(7638): p. 251-254.
3. Cohen, J. and W. Looney, Re: How Much Is Life W orth: Cetuximab, Non-Small Cell Lung Cancer, and the $\$ 440$ Billion Question. Journal of the National Cancer Institute, 2010. 102(15).
4. Eichler, H.G., et al., Use of cost-effectiveness analysis in health-care resource allocation decision-making: How are
cost-effectiveness thresholds expected to emerge? Value in Health, 2004. 7(5): p. 518-528.
5. Green, C. and K. Gerard, EXPLORING THE SOCIAL V ALUE OF HEALTH-CARE INTERVENTIONS: A STATED PREFERENCE DISCRETE CHOICE EXPERIMENT. Health Economics, 2009. 18(8): p. 951-976.
6. Groot, W. and H.M. van den Brink, The value of health. Bmc Health Services Research, 2008. 8.
7. Hughes, D.A. and R.E. Ferner, New drugs for old: disinvestment and NICE. British Medical Journal, 2010. 340.
8. Lieu, T.A., et al., Willingness to Pay for a QALY Based on Community Member and Patient Preferences for Temporary Health States Associated with Herpes Zoster. Pharmacoeconomics, 2009. 27(12): p. 1005-1016.
9. Mason, A.R. and M.F. Drummond, Public funding of new cancer drugs: Is NICE getting nastier? European Journal of Cancer, 2009. 45(7): p. 1188-1192.
10. Mason, H., M. Jones-Lee, and C. Donaldson, MODELLING THE MONETARY V ALUE OF A QALY: A NEW APPROACH BASED ON UK DATA. Health Economics, 2009. 18(8): p. 933950.
11. Maynard, A and Bloor, K The future role of NICE. BMJ. $341:$ c6286, November 13, 2010.
12. Rascati, K.L., The $\$ 64,000$ question- What is a quality-adjusted life-year worth? Clinical Therapeutics, 2006. 28(7): p. 1042-1043.
13. Rawlins, M., D. Barnett, and A. Stevens, Pharmacoeconomics: NICE's approach to decision-making. British Journal of Clinical Pharmacology, 2010. 70(3): p. 346-349.
14. Shiroiwa, T., et al., INTERNATIONAL SURVEY ON WILLINGNESS-TO-PAY (WTP) FOR ONE ADDITIONAL QALY GAINED: WHAT IS THE THRESHOLD OF COST EFFECTIVENESS? Health Economics, 2010. 19(4): p. 422-437.
15. Speight, J. and M. Reaney, Wouldn't it be NICE to consider patients' views when rationing health care? British Medical Journal, 2009. 338.
16. Tappenden, P., et al., A stated preference binary choice experiment to explore NICE decision making. Pharmacoeconomics, 2007. 25(8): p. 685-693.
17. Weinstein, M.C., How much are Americans willing to pay for a quality-adjusted life year? Medical Care, 2008. 46(4): p. 343-345.
18. Yaesoubi, R. and S.D. Roberts, A game-theoretic framework for estimating a bealth purchaser's willingness-topay for health and for expansion. Health Care Management Science, 2010.13(4): p. 358-377.

## Step 2 Results

1. Appleby J, Devlin N, Parkin D, et al. Searching for locatNHS cost effectiveness thresholds: a feasibility study. NICE confer Manchester; 2007 Dees5-6[online] Available from URL: http://www.nice2007.co.uk/ApplebyDevlin.pdf
2. Birch S, Gafni A. The biggest bang for the buck or bigger bucks for the bang: the fallacy of the costeffectiveness threshold. J. Health Serv. Res. Policy11,46-51 (2006).
3. Braithwaite RS, Roberts MS. $\$ 50,000$ per QALY: inertia, indifference, or irrationality? Presented at: Annual Meeting of the Society for Mêdical Décision Making. Atlanta, GA, USA, 17-20 October, 2004.
4. Drummond M, Torrance G, MasonJ.Cost-effectiveness league tables: more harm than good? Soc. Sci. Med.37,33-40 (1993).
5. Gerard K, Mooney G. QALY league tables: handle with care. Health Econ 1993; 2 (1): 59-64
6. Gyrd-Hansen D. Willingness to pay for a QALY: theoretical and methodological issues. Pharmacoeconomics 2005; 23 (5): 423-32
7. Hammitt JK. The $\$ 64,000$ question: what are we willing to pay for a QALY. ISPOR Connections11(1), $7-9$ (2005).
8. Hirth RA, Cherney ME, Miller E, et al. Willingness to pay for a quality-adjusted life year: in search of a standard. Med Decis Making 2000; 20 (3): 332-42
9. King JT/JF, Tsevat J, Lave JR, Roberts MS. Willingness to pay for a quality-adjusted life year: implications for societal health care resource allocation. Med. Decis. Making25,667-677 (2005). [GrossRef] [Medline]
10. Lee C, Chertow G, Zenios S. An empiric estimate of the value of life: updating the renal dialysis cost-effectiveness standard. Value Health 2009; 12 (1): 80-7
11. Martin S, Rice N, Smith P. Further evidence on the link between health care spending and health outcomes in England [CHE 28. National Institute for Health and Clinical Excellence. NICE discussion paper 32]. York: University of York, 2007
12. Martin S, Rice N, Smith PC. The link between health care spending and health outcomes for the new English primary care trusts. CHE Research Paper 42, University of York, York; 2008.
13. Mauskopf J, Rutten F, Schonfeld W. Cost-effectiveness league tables: valuable guidance for decision makers? Pharmacoeconomics21,991-1000 (2003).
14. Smith RD, Richardson J. Can we estimate the 'social' value of a QALY? Four core issues to resolve. Health Policy74,77-84 (2005). [CrossRef] [Medline]
15. Stinnett AA, Paltiel AD. Mathematical programming for the efficient allocation of health care resources. J Health Econ 1996; 15 (5): 641-53
16. Towse A, Pritchard C, Devlin N, eds. Cost effectiveness thresholds: economic and ethical issues. London: Office of Health Economics, Kings Fund, 2002.
17. Ubel PA, Hirth RA, Chernew ME, et al. What is the price of life and why doesn't it increase at the rate of inflation? Arch Int Med 2002; 163: 1637-41
18. Williams A. What could be nicer than NICE? London: Office for Health Economics, 2004. OHE Annual Lecture 2004
19. Winkelymayer WC, Weinstein MC, Mittelman MA, Glynn RJ, Pliskin JS. Health economic evaluations: the special case of end-stage renal disease treatment. Med Decis Making 2002; 22:417 30.

## Step 3 Results

1. Baker, R., et al., Weighting and valuing quality-adjusted life-years using stated preference methods: preliminary results from the Social Value of a QALY Project. Health Technology Assessment, 2010. 14(27)
2. Bobinac, A., et al., Willingness to Pay for a Quality-Adjusted Life-Year: The Individual Perspective. Value in Health, 2010. 13(8): p. 1046-1055.
3. Brock, D.W., How much is more life worth? Hastings Center Report, 2006. 36(3): p. 17-19.
4. Byrne, M.M., K. O'Malley, and M.E. Suarez-Almazor, Willingness to pay per quality-adjusted life year in a study of knee osteoarthritis. Medical Decision Making, 2005, 25(6): p. 655-666.
5. Griffin, S., K. Claxton, and M. Sculpher, Decision analysis for resource allocation in health care. Journal of Health Services Research \& Policy, 2008. 13: p. 23-30.
6. Gyrd-Hansen, D., Willingness to pay for a QALY. Heatth Economics, 2003. 12(12): p. 1049-1060.
7. Harrison, S., A POLICY AGENDA FOR HEALTH-CARE RATIONING. British Medical Bulletin, 1995. 51(4): p. 885-899.
8. Laufer, F., Thresholds in cost-effectiveness analysis - More of the story. Value in Health, 2005. 8(1): p. 86-87.
9. Pinto-Prades, J.L., G. Loomes, and R. Brey, Trying to estimate a monetary value for the QALY. Journal of Health Economics, 2009. 28(3): p, 553-562.
10. Vernon, J.A., R. Goldberg, and J. Golec, Economic Evaluation and Cost-Effectiveness Thresholds Signals to Firms and Implications for R\&D Investment and Innovation. Pharmacoeconomics, 2009. 27(10): p. 797-806

## Step 4 Results

1. Abelson, P., The vatue of life and health for public policy. Economic Record, 2003. 79: p. S2-S13.
2. Fryback, D.G. and W.F. Lawrence, Dollars may not buy as many QALYs as we think:: A problem with defining qualizy-of -iffe adjustments. Medical Decision Making, 1997. 17(3): p. 276-284.
3. Gafni, A. and S, Birch, GUIDELINES FOR THE ADOPTION OF NEW TECHNOLOGIES - A PRESCRIPTION FOR UNCONTROLLED GROWTH IN EXPENDITURES AND HOW TO AVOD THE PROBLEM. Canadian Medical Association Journal, 1993. 148(6): p. 913-917.
4. Johnson, F.R., Einstein on willingness to pay per QALY: Is there a better way? Medical Decision Making, 2005. 25(6): p. 607-608.
5. Laupacis, A., et al., HOW ATTR ACTIVE DOES A NEW TECHNOLOGY HAVE TO BE TO W ARR ANT ADOPTION AND UTILIZ ATION - TENTATIVE GUIDELINES FOR USING CLINICAL AND ECONOMIC EV ALUATIONS. Canadian Medical Association Journal, 1992. 146(4): p. 473-481.
6. Polsky, D., Does willingness to pay per quality-adjusted life year bring us closer to a useful decision rule for costeffectiveness analysis? Medical Decision Making, 2005. 25(6): p. 605-606.
7. Martin, S., N. Rice, and P. Smith, The link between health care spending and health outcomes: evidence from English programme budgeting data. CHE Research Paper 24, 2007a.
8. Chambers, J.D., P.J. Neumann, and M.J. Buxton, Does Medicare Have an Implicit CostEffectiveness Threshold? Medical Decision Making, 2010. 30(4): p. E14-E27.

## Step 5 Results

1. Birch, S. and A. Gafni, COST-EFFECTIVENESS RATIOS - IN A LEAGUE OF THEIR OWN. Health Policy, 1994. 28(2): p. 133-141.
2. Johnson, F.R. and M. Backhouse, Eliciting stated preferences for health-technology adoption criteria using paired comparisons and recommendation judgments. Value in Health, 2006. 9(5): p. 303-311.

## Step 6 Results

1. Dolan, P., et al., QALY maximisation and people's preferences: a methodological review of the literature. Health Economics, 2004. 14(2): p. 197-208.
2. Baker, R., et al., Weighting and valuing quality-adjusted life-years using stated preference methods: preliminary results from the Social Value of a QALY Project. Health Technology Assessment, 2010. 14(27)
3. O'Brien, B.J., et al., Is there a kink in consumers' threshold value for cost-effectiveness in health care? Health Economics, 2002. 11(2): p. 175-180.
4. Buxton, M., How Much Are Health-Care Systems Prepared to Pay to Produce aQALY? European Journal of Health Economic, 2005. 6(4): p. 285-28.
5. Mason, A.R. and M.F. Drummond, Public funding of new cancer drugs: Is NICE getting nastier? European Journal of Cancer, 2009. 45(7): p. 1188-1192.

## Appendix B

## THE LINK BETWEEN NHS SPENDING AND MORTALITY: ESTIMATING THE COST OF A LIFE YEAR IN ENGLAND

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## THE LINK BETWEEN NHS SPENDING AND MORTALITY: ESTIMATING THE COST OF A LIFE YEAR IN ENGLAND ${ }^{1}$

## Prologue

This report presents, in a linear fashion, details of the econometric work undertaken to estimate the link between NHS spending and mortality. It also presents details of how the econometric work is used to calculate the cost of a life year. This report is designed to serve as a reference document in support of the main project report, which highlights the major findings from the project. As a supporting document this report provides far more detail than most interested parties will require. Nevertheless, those who seek more detail than that contained in the main project report may find the material here useful.

[^40]
## A BACKGROUND, MODEL, DATA, AND ESTIMATION APPROACH

## B1. Introduction

In a recent White Paper the new British Conservative government emphasized the importance of clinical outcomes. It notes that, in future, success will be measured, not through the achievement of process targets, such as short waiting times, but against outcomes such as cancer and stroke survival rates [1]. Although the NHS budget is ring-fenced against the on-going public sector deficit reduction programme, its budget is still likely to be under considerable pressure, and attention is likely to focus on the extent to which any additional health care expenditure yields genuine patient benefits in the form of improved health outcomes.

However, one of the most fundamental yet unresolved issues in health policy is the extent to which additional health care expenditure yields patient benefits, in the form of improved health outcomes. The work of health technology agencies, such as the English National Institute for Health and Clinical Excellence (NICE), has greatly improved our understanding at the micro-level of the costs and benefits of individual therapeutic technologies. However, there remains a dearth of evidence af the macro-level on the benefits of increased health system expenditure.

Recently a series of studies has taken advantage of the availability of two new datasets to examine the relationship between NHS expenditure and mortality rates for various disease categories[2-5]. One dataset contains mortality rates for various disease categories at the level of geographically defined local health authorities, known as Primary Care Trusts (PCTs). The other dataset presents NHS expenditure by PCT on 23 broad programmes of care. This dataset embraces mostitems of publicly funded expenditure, including inpatient, outpatient and community care, and pharmaceutical prescriptions.

Like previous studies, we employ a model that assumes that each PCT receives an annual financial lump sum budget from the national ministry and allocates its resources across the 23 programmes of care to maximize the health benefits associated with that expenditure. Estimation of this model using the expenditure and mortality data facilitates two related studies: first, a study of how changes in the NHS budget impact on expenditure in each care programme; and second, a study of the link between expenditure in a programme and the health outcomes achieved, notably in the form of disease specific mortality rates. The latter study also permits the calculation of the cost of an additional life year for individual programmes of expenditure.

The work presented here draws heavily upon previous studies. These were constrained in a number of ways and, in this analysis; we build on and improve these previous studies in four major ways:

- first, due to data limitations previous studies related expenditure in time period $t$ to mortality in periods $t, t-1$, and $t-2$. In doing this, such studies assumed that PCTs had reached some sort of equilibrium in the expenditure choices they make and the outcomes they secure. This is probably not an unreasonable assumption given the relatively slow pace at which both types of variable change but, with more recent mortality data now available, here we relate expenditure in time period $t$ to mortality in periods $t, t+1$, and $t+2$ (see section B8.5).
. second, previous studies have tended to focus on a very limited number of care programmes (e.g., for cancer, circulatory disease, gastro-intestinal problems and respiratory problems). Here we present plausible outcome models for a larger number of budgeting categories.
- third, previous estimates of the cost of a life year have been for individual programmes of care. Here we present estimates of the cost of a life year for an enlarged number of programmes and, importantly, with the aid of assumptions about the productivity of programmes without a meaningful mortality-based outcome indicator, we extend our individual programme estimates to incorporate expenditure across all programmes of care.
- finally, although previous results and our current models 'pass' the appropriate statistical tests, we subject our latest results to a substantial sensitivity analysis.

The structure of this report is as follows. Section B2 presents a brief review of previous empirical studies in this domain, which have often yielded conflicting results. A straightforward theoretical model of the budgetary problem faced by a PCT manager seeking to allocate limited funds between competing programmes of care is presented in section B3. The programme budgeting and health outcome (mortality) data are described in sections B4 and B5 respectively. Section B6 outlines our estimation methods and some of the issues surrounding them.

In section B7 we commence our empirical work by estimating well specified econometric models that outline (a) the budgetary expenditure choices and (b) the health outcomes achieved by PCTs using expenditure data for $2005 / 6$ and mortality data for $2002 / 3 / 4$. Section B8 presents results using expenditure data for $2006 / 7$ and mortality data for $2004 / 5 / 6$. It also presents results using the same expenditure data but updating the mortality data to $2006 / 7 / 8$. Several pieces of sensitivity analysis are also included in section B 8 , but the major piece of sensitivity analysis - examining the impact) of relaxing the instrument validity restriction - is reported in section B9.

In section B10 we re-estimate our model using updated expenditure and mortality data. In particular, we use the programme budgeting expenditure for 2007/8 and mortality data for 2007/2008/2009 to reestimate our outcome and expenditure equations. In section B11 we update the dataset again, and this time we employ programme budgeting expenditure data for 2008/9 and mortality data for $2008 / 2009 / 2010$. We also compare the elasticities and cost of a life year estimates that we have obtained using expenditure and mortality data for different years.

Finally, section B12 presents a summary of our findings and some concluding remarks.

## B2. Previous studies

There is a large literature on the determinants of international variations in health care spending in which income levels often play a central role [6]. However, whether more expenditure generates better outcomes - for example, in terms of reduced mortality - remains a matter of debate. For example, Fisher and Welch [7] note various ways in which more health care might harm patients and they cite various studies supporting their arguments. In a comprehensive review, Nolte and McKee [8] discuss many studies that examine the impact of health care and other explanatory variables on some measure of health care outcome. Nolte and McKee point out that researchers usually combine a production function approach with the application of regression analysis. For example, in an early cross-sectional study of 18 developed countries, Cochrane et al.[9] use regression analysis to examine the statistical relationship between mortality rates on the one hand and per capita GNP and per capita consumption of inputs such as health care provision on the other. They find that the indicators of health care provision were generally not associated with outcomes in the form of mortality rates. Thereafter, the failure to identify strong and consistent relationships between health care expenditure and health outcomes (after) controlling for other factors) has become a consistent theme in the literature, whilst, in contrast, socioeconomic factors are often found to be good determinants of health outcomes [8, (10, 11].

This failure to detect a significant positive relationship between expenditure and health outcome might reflect the difficulties associated with any such study rather than the absence of such a relationship. For example, Gravelle and Backhouse [12] examine some of the methodological difficulties associated with empirical investigation of the determinants of mortality rates. These include simultaneous equation bias and the associated endogeneity problem (that the level of health care input might reflect the level of health outcome achieved in the past), and that a lag may occur between expenditure and outcomes (studies typically assume that expenditure has an immediate effect on mortality). To avoid the difficulties imposed by data heterogeneity inherent in international analyses, the study by Cremieux et al[13] examines the relationship between expenditure and outcomes across ten Canadian provinces over the fifteen-year period 1978-1992. They find that lower healthcare spending is associated with a significant increase in infant mortality and a decrease in life expectancy.

Although challenging the received empirical wisdom, one difficulty with the Cremieux et al[13] study is that the estimated regression equation consists of a mixture of potentially endogenous variables (such as the number of physicians, health spending, alcohol and tobacco consumption, expenditure on meat and fat) and exogenous variables (such as income and population density). The authors' chosen estimation technique (GLS) does not allow for this endogeneity and consequently the coefficients on the endogenous variables may be biased [12]. Or's [14] study of the determinants of variations in mortality rates across 21 OECD countries between 1970 and 1995 may suffer from the same weakness. She finds that the contribution of the number of doctors to reducing mortality in OECD countries is substantial but her estimation technique assumes that the number of doctors is exogenous to the health system.

Nixon and Ulmann[15] provide a detailed review of 16 studies that have examined the relationship between health care inputs and health outcomes, using macro-level data. They also undertake their own study using data for 15 EU countries over the period 1980-1995. They employ three health outcomes measures - life expectancy at birth for males and females, and the infant mortality rate - and a dozen or more explanatory variables including: per capita health expenditure, number of physicians (per 10,000 head of population), number of hospital beds (per 1,000 head of population), the average length of stay in hospital, the in-patient admission rate, alcohol and tobacco consumption, nutritional characteristics, and environmental pollution indicators. Nixon and Ulmann conclude that although health expenditure and the number of physicians have made a significant contribution to improvements in infant mortality, '...health care expenditure has made a relatively marginal contribution to the improvements in life expectancy in the EU countries over the period of the analysis'. Again, however, the study does not allow for the possibility that some of the explanatory variables may be endogenous.

Although loosely based on the notion of a health production function, the traditional empirical study described above has rarely been informed by an explicit theoretical model. This is understandable, as the processes giving rise to the observed health outcome are likely to be very complex, and any theoretical
model might become rather unwieldy. However, this absence of a model has usually led to an atheoretical search for measures of health inputs demonstrating a statistically 'significant' association with health outcomes. In contrast, in this study we inform our empirical modelling with a theoretical framework. We believe that this may lead to a more convincing and better specified model of health outcomes than that used in many previous studies, and this model is outlined in the next section.

## B3. Theoretical model

Our modelling framework assumes that each PCT i receives an annual financial lump sum budget $y_{i}$ from the national ministry, and that annual total expenditure cannot exceed this amount. The PCT must then decide how to allocate its budget across the $J$ programmes of care ( $J=23$ in this case). For each programme of care there is a 'health production function' $f_{i}($.$) that indicates the link between local$ spending $x_{i j}$ on programme $j$ and health outcomes in that programme $h_{i j}$. Health outcomes might be measured in a variety of ways, but the most obvious is to consider some measure of improvement in life expectancy, possibly adjusted for quality of life, in the form of a quality adjusted life year.

The nature of the specific health production function confronted by a PCT will depend on two types of local factors: the clinical needs of the local population relevant to the programme of care (which we denote $n_{i j}$ ) and broader local environmental factors $z_{i j}$ relevant to delivering the programme of care (such as input prices, geographical factors, or other uncontrollable influences on outcomes). Both clinical and environmental factors may be multidimensional in nature. Increased expenditure then yields improvements in health outcomes, as expressed for example in improved local mortality rates, but at a diminishing rate. That is:

$$
\begin{equation*}
h_{i j}=f_{j}\left(x_{i j}, n_{i j}, z_{i j}\right) ; \partial f_{j} / \partial x>0 ; \partial^{2} f_{j} / \partial x^{2}<0 \tag{3.1}
\end{equation*}
$$

We assume there is a PCT social welfare function $W$ (.) that embodies health outcomes across the $J$ programmes of care. Assuming no interaction between programmes of cate, each PCT allocates its budget so as to maximise total welfare subject to the local budget constraint and the health production function for each programme of care:
$\max \quad W\left(h_{i 1}, h_{i 2}, \ldots, h_{i J}\right)$
subject to $\sum_{j} x_{i j} \leq y_{i}$

$$
\begin{equation*}
h_{i j}=f_{j}\left(x_{i j}, n_{i j}, z_{i j} ; \quad \quad j=1, \ldots \mathrm{~J}\right. \tag{3.2}
\end{equation*}
$$

It can of course quite plausibly be argued that decision-makers do not discriminate between health outcomes in different programmes of care, and that W (.) is merely the sum of such outcomes. However, there is no need for that assumption in our formulation.

Each PCT allocates expenditure across the 23 programmes of care so that the marginal benefit of the last pound spent in each programme of care is the same. This is represented diagrammatically in Figure B3.1, which illustrates the trade-off between just two programmes of care. The top left hand quadrant indicates the health production function for programme 1, whilst the bottom right hand quadrant indicates the health production function for programme 2, albeit in transposed form. The bottom left hand quadrant indicates the budget constraint: the expenditure choice must lie on the budget line. This means that for each feasible pair of expenditure choices (points on the budget constraint line), a pair of health outcomes in the two programmes emerges, which is traced out as the health production possibility frontier in the top right quadrant. The PCT will choose the point on this frontier that maximizes welfare. In this example, we have indicated a simple health maximizing approach (the maximum health summing across the two programmes), leading to optimal health outcomes $\left(\mathrm{H}_{1}{ }^{*}, \mathrm{H}_{2}{ }^{*}\right)$ and expenditure $\left(\mathrm{X}_{1}{ }^{*}, \mathrm{X}_{2}{ }^{*}\right)$.


Figure B3.1: graph showing optimal trade-off between two programmes of care
Solving the constrained maximisation problem yields the result that the optimal level of expenditure in each category, $x_{i j}^{*}$, is a function of the need for health care in each category ( $n_{i}, n_{i}, \ldots, n_{i j}$ ), environmental variables affecting the production of health outcomes in each category ( $\left.z_{i}, z_{i}, \ldots, z_{i}\right)$, and PCT income $\left(y_{i}\right)$. Thus

$$
\begin{equation*}
x_{i j}^{*}=g_{j}\left(n_{i 1}, \ldots n_{i J}, z_{i 1}, \ldots z_{i J}, y_{i}\right) \cdot \quad \mathrm{j}=1, \ldots, \mathrm{~J} \tag{3.3}
\end{equation*}
$$

Thus, for each programme of care there exists an expenditure equation (3.3) explaining expenditure choice of PCTs and a health outcome equation (3.1) that models the associated health outcomes achieved.

Our model is static in the sense that the health production function (3.1) assumes that all health benefits occur contemporaneously with expenditure. We acknowledge that for some programmes of care benefits might occur one or more years after expenditure has occurred. This is particularly likely to be the case for those programmes amed at encouraging healthy lifestyles, where some benefits may occur decades after the actual programme expenditure. For other programmes, such as maternity/reproductive conditions and neonate conditions, benefits may be largely contemporaneous with expenditure. Furthermore, we do not model the decision maker's time preferences.

For our empirical modelling, however, we are constrained by the data we have available, which are largely cross-sectional in nature. Due to data limitations, previous studies have had to relate expenditure in period $t$ to mortality data in periods $t, t-1$, and $t-2$ so that the mortality data precedes the expenditure data. This is not ideal. Implicitly previous studies have had to assume that the data represent a quasi long-run equilibrium position, and that relative expenditure levels and health outcomes within each РСТ have been reasonably stable over a period of time. As we shall see, this appears to be a reasonable assumption because we obtain similar results when we estimate our models using expenditure for period $t$ with either mortality data for periods $t, t-1$, and $t-2$ (section B8.4) or with mortality data for periods $t, t+1$, and $t+2$ (section B8.5).

Having outlined our model, the next section discusses the datasets used to estimate this model.

## B4. NHS programme budgeting in England

The English National Health Service (NHS) is the archetypal centrally planned and publicly funded health care system. Its revenue derives almost entirely from national taxation, and access to the system is generally free to the patient. Primary care is an important element of the system, and general practitioners act as gatekeepers to secondary care and pharmaceuticals. The system is organized geographically, with responsibility for the local administration of the NHS devolved to local health authorities known as Primary Care Trusts (PCTs). ${ }^{2}$ For the purposes of this study, there were 303 PCTs with an average population of about 160,000 people until October 2006. In October 2006 the 303 PCTs became 152 PCTs. Some PCT boundaries remained unchanged while other PCTs were merged with one or more neighbours to form a new, larger, PCT. In a few cases the geographic area covered by an existing PCT was split between two or more new PCTs. These 152 PCTs have an average population of about 330,000 people. ${ }^{3}$ PCTs are allocated fixed annual budgets by the national ministry, within which they are expected to meet expenditure on most aspects of health care, including inpatient, outpatient and community care, primary care and pharmaceutical prescriptions.

## B4.1 The rationale behind the construction of programme budget data

Traditionally, PCTs and their predecessors have reported expenditure on the basispf inputs (for example, total expenditure on pay and non-pay items). However, NHS policy makers have for some time realized that this approach does not create clinically meaningful financial data or help in the design and evaluation of programmes of patient care. The Department of Health therefore initiated a 'Programme Budgeting' project. This has sought to create an accounting system that is more atigned with the distinct outputs and health outcomes of the health care system. Since April 2003, in addition to its conventional accounting data, each PCT has prepared expenditure data disaggregated according to 23 programmes of health care. These programmes are defined by reference to the International Classification of Diseases (ICD) Version 10 codes at the four digit level, and most programme budget categories reflect ICD 10 chapter headings (e.g., cancer and tumours, circulation problems, renat problems, neonates, problems associated with the skin, problems associated with vision, problems associated with hearing, etc). In some cases, the 23 categories are broken down into further sub-areas to achieve a closer match with the various National Service Frameworks (NSFs): for example, the large mental health category is broken down into 'substance abuse', 'dementia', and 'other'.

Programme budgeting seeks to allocate all types of PCT expenditure to the various programme budget categories, including secondary care, community care and prescribing. However, the system acknowledges that a medical model of care may not always be appropriate, and two specific non-clinical groups -'Healthy Individuals' and Social Care Needs' -- have been created. These are intended to capture the costs of disease prevention programmes and the costs of services that support individuals with social rather than health care needs. In addition, in some cases it is not possible to assign activity by medical condition, preventative activity, or social care need and, in these cases, expenditure is assigned to a residual category (PBC 23) entitled 'Other'. The most important element of this residual programme is expenditure on general practitioner services ( PBC 23 a ). In principle, it should be possible to allocate each GP consultation to a particular care programme. However, at the moment the available data information systems do not permit such an allocation and so all primary care expenditure is allocated to this residual programme. The use of this residual category ensures that all expenditure is assigned to a programme of care (16].

The aim of the programme budget classifications is to identify the entire volume of health care resources assigned to broad areas of illness according to the primary diagnosis associated with an intervention. It serves a number of purposes, most notably to assist in the local planning of health care. But for this

[^41]study its crucial merit is that it opens up the possibility of examining the statistical relationship between local programme spending and the associated disease-specific outcome.

## B4.2 The collection of programme budgeting data

Programme budgeting information is collected centrally by the Department of Health as part of the annual accounts process. Each PCT is required to submit an annual programme budgeting return to the Department which shows how their total expenditure is allocated across the 23 programme budgeting categories.

Various forms of data collection and analysis are required to map PCT expenditure onto acute, community and other services to the 23 programme budget categories. From the PCT perspective, however, the construction of each PCT's return largely involves collating information provided by other bodies and drawing on other information already in the PCT's own annual accounts. Thus General/Personal Medical Service expenditure, which is already reported in PCT accounts, relates to direct primary care and is mapped in its entirety to programme budget category 23a (Other: GMS/PMS); General Ophthalmic Service expenditure (again from PCT accounts) maps directly to programme budget category 8 (eye/vision problems); and General Dental Service expenditure maps directly to programme budget category 12 (dental problems). Prescribing and pharmaceutical services expenditure is allocated to programme budget categories on the basis of an annual apportionment report provided by the Prescription Pricing Authority for each PCT as part of the annual accounts process. This apportionment report allocates each PCT's annual FHS prescribing expenditure across the 23 programme budget categories. The balance of any primary healthcare purchased by the PCT is allocated / apportioned across the 23 programme budget categories on the basis of local records, with any remaining expenditure allocated/apportioned in line with the distributions already made across the budget categories.

It is the responsibility of all NHS providers - which includes PCTs, NHS Trusts, and Foundation Hospitals - to allocate admitted patient care expenditure across the programme budgeting categories, specific to each PCT that utilises its services. These allocations are constructed using 'finished consultant episodes' (from the mandatory administrative Hospital Episode Statistics data set returned by each provider) each of which is assigned to a Healthcare Resource Group (HRG), an English version of DRGs. National grouping software automatically assigns each HRG to one of the 23 programme budgeting categories and attaches the provider's average reference cost for the relevant HRG to each record. For each PCT this information generates a split of inpatient care expenditure by programme budget category for each of its secondary healthcare providers.

There are numerous difficulties faced when attempting to allocate non-admitted patient care activity (that is, outpatients, community services, direct access, A\&E etc) to programme budget categories. The difficulties are primarily due to the absence of clear diagnostic codes. The 'primary reason for care' (equivalent to a diagnosis code) is not information that is routinely collected for community patients. Because of this, the approach prescribed is for service providers to produce a generic allocation analysis/report, for all PCTs making use of their services, for all non-admitted patient care costs across the 23 programme budget categories. Once derived, this generic allocation analysis/report is made available to PCTs at the same time as the unique (PCT specific) inpatient care information described above. Unlike the first apportionment report relating to admitted patient care, the non-admitted patient care apportionment report will not be unique to the PCT, but will represent the provider's overall experience. PCTs are expected to use this data to inform the apportionment of their own spend on nonadmitted patient care across the 23 programme budget categories.

The Department of Health recognises that this approach - the provision of a PCT specific breakdown of admitted patient care costs and a generic allocation of all PCTs non-admitted patient care spend by providers - is likely to generate a crude method for apportioning non-admitted patient care costs. PCTs and their providers are therefore encouraged to put in place other arrangements that allow a more sophisticated analysis of non-admitted patient care expenditure. Such arrangements may well rely on an activity sampling approach [16].

Mental Health providers may not need to complete and forward detailed admitted and non-admitted patient care apportionment reports to PCTs. The nature of the services they provide may be such that the entire spend with them relates exclusively to the Mental Health programme (budget category 5). Ambulance Trusts are required to provide non-admitted patient care information to those PCTs for whom they provide services. Where it is not possible to split the activity by PCT, a generic non-admitted patient care report is produced for all purchasers [16].

The Department of Health has been criticised for the rather simplistic way in which it has apportioned certain costs among categories, and there are obvious issues with the allocation of costs associated with patients who have multiple disorders. However, the programme budgeting project is very much work-inprogress and the Department is investigating ways to improve the accuracy with which costs are allocated across programmes (for example, the Department is investigating the possibility of allocating training expenditures to specific programmes rather than to the generic medical training programme PB\&23b).4

## B4.3 Programme budgeting expenditure, 2003/4-2008/9

National (all PCT) expenditure per head and the growth in this expenditure are shown for each programme budget category for 2003/04 to 2008/09 in Table B4.1. Comparable data for each programme budget sub-category is shown in Table BA. 1 in the annex. Year on yearcomparisons of expenditure in each group are complicated by the fact that the algorithms used to allocate activity to PBCs are regularly revised. For example, for $2006 / 7$ two major changes were made to the methods employed to construct the programme budgeting data. First, expert medical opinion was employed to re-evaluate the existing mapping from inpatient diagnosis codes to programme budget category. This led to the re-assignment of just over $10 \%$ of all diagnosis (ICD10) codes from one programme budgeting category to another. ${ }^{5} 6$ Second, activity to be costed used the newly introduced version 4 of the Healthcare Resource Group (HRG) software which, among other things, changed the methodology for calculating non-admitted patient care costs. HRG4 reflected advances in clinical practice and was designed to generate a much more accurate costing of complex cases. Other developments, such as the transfer of responsibility for dental funding from local dental boards to PCTs, also complicate the interpretation of comparisons through time (for example, per capita dental expenditure by PCTs increased from $£ 13.55$ in 2004/5 to $£ 51.93$ in 2006/7).

The expenditure figures for the first year (2003/4) are calculated on a slightly different basis to those for the other years (2004/5-2008/9). In particular, the figures for 2003/4 are on a 'net expenditure' basis while the figures for 2004/5-2008/9 are on an 'own population' basis. The 'own population' figure starts with net expenditure; it adds anvexpenditure funded from sources outside of the NHS; and then deducts any expenditure on other PCTs' populations incurred through lead/host commissioning arrangements. In $2006 / 7$ and across all PBCs, expenditure per head on an own population basis was $2.3 \%$ greater than expenditure on a net population basis.

In 2004/5 totalPGT expenditure per person was $£ 1,200$. The category attracting the most expenditure was the 'other' category (programme budget category 23) with per capita expenditure of almost $£ 158$ $(13.2 \%)$. This category included primary care expenditure, workforce training expenditure, and a range of

[^42]other miscellaneous expenditure items. Of these components, primary care expenditure was by far the largest element at $£ 127$ per head.

In 2004/5 there were two other categories with a budget share of over 10\%: mental health (budget category 5 ) attracted $12.2 \%$ of expenditure ( $£ 147$ per person), and circulation problems (budget category 10) recorded $10.2 \%$ of expenditure ( $£ 122$ per person). Seven programme budget categories - cancers and tumours ( $£ 76$ ), gastro-intestinal problems ( $(, 73)$, trauma and injuries ( $\AA_{7}$ ), musculo-skeletal problems ( $£ 72$ ), respiratory problems ( $£ 63$ ), genito-urinary problems ( $£ 62$ ), and maternity and reproductive conditions ( $£ 55$ ) - had expenditure shares of between $4.6 \%$ and $6.3 \%$. Finally, the 13 remaining categories - from hearing problems ( $£ 6$ ) to learning disability ( $£ 43$ ) - each account for between $0.5 \%$ and $3.6 \%$ of total expenditure.

By 2008/9 total PCT expenditure per person had increased to $£ 1,531$ (up $28 \%$ from 2004/5). The residual 'other' category (programme budget category 23) still accounted for the largest share of expenditure $(14.9 \%)$ with per capita expenditure of almost $£ 228$, of which $£ 145$ was accounted for by primary care expenditure. Mental health (budget category 5) still accounted for just over $12 \%$ of expenditure, but the expenditure share recorded by circulation problems (budget category 10) had fallen from $10.2 \%$ to $8.5 \%$. Other categories recording a fall in budget share of more than one half of one percentage point included: the gastro-intestinal system (down from $6.1 \%$ to $5.1 \%$ (), the musculo-skeletal system (down from $6 \%$ to $5.2 \%$ ), trauma and injuries (down from $6 \%$ to $4.2 \%$ ), and maternity (down from $4.6 \%$ to $3.9 \%$ ).

Categories recording an increase in budget share of more than one half of one percentage point included neurological problems (up from $2.9 \%$ to $4.4 \%$ ) and dental problems (up from $1.1 \%$ to $4.1 \%$ ).

Some of these changes will partly reflect revisions to the lgotithms used to allocate expenditure to particular PBCs. For example in 2006/7 expenditure per person on musculo-skeletal problems fell by $11 \%$ and expenditure on trauma and injuries fell by $25 \%$. In the same year, expenditure on neurological problems increased by $35 \%$. This suggests that some types of activity, which were previously allocated to musculo-skeletal problems and/or trauma and injuries, were re-allocated to neurological problems. Similarly, up to and including 2006/7, expendirure that was not directly attributable to a particular programme category was apportioned using admitted patient care percentages. ${ }^{7}$ In other words, if $x \%$ of total admitted patient care expenditure was allocated to PBC 1 , then $x \%$ of all expenditure that was not directly attributable to a particular programme category was also allocated to PBC 1 . With effect from 2007/8, however, NHS organisations were asked to select an appropriate basis for the apportionment of this non-programme specific expenditure and that, where no reasonable basis existed, such expenditure was to be allocated to the 'Other -Miscellaneous' (PBC 23X) category.

These two changes to the algorithm used to allocate expenditure to particular PBCs illustrate that year-on-year comparisons of expenditure need to be interpreted with care.

Obviously, expenditure per head on any given programme varies from one PCT to another and Table B4.2 presents some statistics that indicate the degree of variation in expenditure levels across PCTs by programme budget category. The first four columns of Table B4.2 present descriptive statistics for PCT expenditure per person by PBC. These reveal that, for example, PCT per capita expenditure in the cancer programme averaged $£ 96.30$ across all PCTs, with the minimum spend being $£_{6} 62.90$ and the maximum being £155.70.

Some PCTs will be spending more than other PCTs simply because they face higher input costs. The second set of four columns in Table B4.2 present descriptive statistics for PCT per capita expenditure

[^43]that has been adjusted for the unavoidable geographical variation in costs (input prices) faced by PCTs. ${ }^{8}$ However, if anything this adjustment appears to increase the variation in expenditure across PCTs; for example, the range of per capita expenditure on cancer increases from between $£ 62.90$ and $£ 155.70$ (unadjusted) to between $£ 59.10$ and $£ 163.10$ (adjusted for local health care input prices).

Another cause of the variation in expenditure levels will be the fact that the need for health care will vary from one PCT to another. For example, areas with a relatively large proportion of elderly residents, or PCTs operating in relatively deprived locations, can be expected to experience relatively high levels of spending. The Department of Health has a well-developed methodology for estimating the relative health care needs of PCTs, which it uses as the basis for allocating health care funds to PCTs[17]. Recent 'needs' formulae have been derived from an adjustment for the demographic profile of the PCT and a series of econometric analyses of the link between health care expenditure and other socio-economic factors at a small area level within England[18].

The final set of four columns in Table B4.2 present descriptive statistics for PCT per capita expenditure that has been adjusted for both the unavoidable geographical variation in costs and the local need for health care faced by PCTs. ${ }^{9}$ For virtually every PBC, this adjustment reduces the variation in expenditure across PCTs; for example, the standard deviation of PCT per capita expenditure falls from $£ 19.70$ to $£ 15.30$ for the cancer programme. Although this adjustment reduces the variation in expenditure levels across PCTs, this decline is quite modest and there are still substantial differences in expenditure even after allowing for differences in local cost and need. For example, expenditure per head in the circulation problems category varies between $£ 78$ and $£ 328$ using cost adjusted expenditure data, but falls between $£ 76$ and $£ 327$ using cost and need adjusted population data.

This variation in expenditure across PCTs has led some commentators to question the reliability of the programme budgeting data. In a good governance report, the National Audit Office [19] sought to '...examine the quality, timeliness and suitability of Programme Budgeting data to support [their] audit of the Department of Health Resource Account and determine whether the systems and processes in place to provide the data are accurate.' The NAO undertook a survey of Trusts, PCTs and SHAs. The NAO noted that a number of PCTs expressed concern about the accuracy of data supplied to them by their service providers and noted that this was believed to be because most Trusts did not use or find the data they supply to PCTs of any use to themselyes. Overall, the NAO's main conclusion was that while the processes for collecting the budgeting data were well defined in most areas, there remained scope for improvement to the robustness of some of the data (such as the non-admitted patient care data).

Appleby, Harrison, Foot, Smith and Gilmour [20] also considered the issue of data reliability in their study of variations in PCT spending on cancer services. They noted a rather dramatic variation in spending across PCTs for any given year, and that a relatively large number of PCTs report relatively large year-on-year changes in cancer expenditure. However, and as the authors point out, it is difficult to define what might be either an implausible level of expenditure or an implausibly large change in expenditure. Moreover, the interpretation of a large change in expenditure is complicated by the fact that the Department of Health makes regular changes (improvements) to the algorithm used to allocated activity tøprogramme budget categories (as detailed above).

As acase study of the reliability of the programme budgeting data, Appleby et al [20] report the results of West Kent PCT's use of an alternative approach to producing the programme budgeting data for cancer and tumours. This alternative approach identified similar levels of expenditure to the traditional method at the aggregate level, but there were differences between the two approaches at the sub-programme level (that is, for expenditure on specific cancer sites and in the residual 'other cancers' category).

[^44]As with any dataset, there are likely to be recording and other errors associated with the programme budgeting data. However, there is no evidence on the magnitude of such errors and we have no reason to believe that such errors are likely to bias our estimates in one particular direction (for example, we have no reason to believe that measurement errors are systematically related to other relevant factors such as mortality rates). In this study, our focus is on whole programme expenditure and thus we avoid the data reliability issues inherent in any analysis of the sub-programme expenditure data. ${ }^{10}$ Moreover, although we present estimates of the cost of a life year for individual programmes, our primary focus is on the cost of a life year across all programmes combined. The advantage of this is that the impact of a PCT reporting, for example, too little expenditure in one category might be offset by reporting too much expenditure in another.

While we note that the allocation of expenditure might not be consistent across PCTs there is no systematic evidence that the magnitude of any inconsistency is sufficiently large to cause concern. Accordingly, for each disease category, the observed variation in expenditure per person - holding constant input prices and the need for health care - offers the opportunity to examine whether PCTs that spend more on health care achieve a better outcome and, if so, at what cost. Empirical estimates of the strength of this relationship for both individual and all programmes of care are presented later in this report.
${ }^{10}$ The ACCA/Audit Commission (2011) looked at the reliability of the programme budgeting data for the diabetes sub-group within the endocrine and metabolic problems category. The ACCA/Audit Commission noted that programme budgeting data includes inpatient and prescribing expenditure, which are thought to be relatively reliably allocated to PBCs and to be consistently costed across PCTs, and outpatient and community service expenditure, which are thought to be less reliably allocated to PBCs and to be less consistently costed across PCTs. The ACCA/Audit Commission compared the variation in expenditure for inpatient and prescribing expenditure with that for total programme budget expenditure and found that the latter was far greater than the former. However, the interpretation of this result is not straightforward: as the ACCA/Audit Commission noted, it is difficult to know whether differences in programme budget spend are attributable to variation in service provision and efficiency, or simply to different approaches to cost allocation.

Table B4.1: table showing national (all PCT) expenditure per head ( $£$ ) and growth in expenditure (\%) by PBC group, 2003/4-2008/9

| PBC \# | PBC description | Spend <br> (f.) per <br> head 2003/4 | Spend <br> (£) per <br> head <br> 2004/5 | Spend <br> (£) per <br> head <br> 2005/6 | Spend <br> (£) per <br> head <br> 2006/7 | Spend <br> (£) per <br> head 2007/8 | Spend <br> (f) per <br> head <br> 2008/9 | $\begin{aligned} & \text { Growth } \\ & (\%) \\ & 2004 / 5 \end{aligned}$ | $\begin{aligned} & \text { Growth } \\ & (\%) \\ & 2005 / 6 \end{aligned}$ | Growth (\%) <br> 2006/7 |  | Growth <br> (\%) $2008 / 9$ | Share of <br> total <br> spend <br> (\%) <br> 2004/5 | Share of <br> total <br> spend <br> (\%) <br> 2008/9 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | Infectious diseases | 17.95 | 20.22 | 23.61 | 20.88 | 22.08 | 23.46 | 13 | 17 | 12 | 6 | 6 | 1.7\% | 1.5\% |
| 2 | Cancers and tumours | 64.95 | 75.54 | 83.24 | 81.67 | 90.21 | 94.55 | 16 | 10 |  | 10 | 5 | 6.3\% | 6.2\% |
| 3 | Blood disorders | 14.08 | 17.00 | 17.48 | 16.58 | 19.44 | 19.50 | 21 | 3 |  | 17 | 0 | 1.4\% | 1.3\% |
| 4 | Endocrine, nutritional | 28.96 | 31.86 | 37.26 | 36.70 | 39.39 | 43.38 | 10 | 17 |  | 7 | 10 | 2.7\% | 2.8\% |
| 5 | Mental health | 133.31 | 146.83 | 158.95 | 166.53 | 180.90 | 191.21 | 10 | $8 \times$ | 5 | 9 | 6 | 12.2\% | 12.5\% |
| 6 | Learning disability | 37.93 | 43.37 | 46.54 | 48.36 | 54.20 | 56.11 | 14 | 7 | 4 | 12 | 4 | 3.6\% | 3.7\% |
| 7 | Neurological | 29.83 | 35.09 | 41.06 | 55.27 | 62.43 | 67.64 | 18 |  | 35 | 13 | 8 | 2.9\% | 4.4\% |
| 8 | Vision problems | 24.61 | 27.65 | 28.24 | 26.97 | 30.69 | 32.95 | 12 | 2 | -4 | 14 | 7 | 2.3\% | 2.2\% |
| 9 | Hearing problems | 5.73 | 6.32 | 6.27 | 6.21 | 8.07 | 8.16 | 10 | -1 | -1 | 30 | 1 | 0.5\% | 0.5\% |
| 10 | Circulatory disease | 110.12 | 122.37 | 124.28 | 122.06 | 124.77 | 129.94 | 14 | 2 | -2 | 2 | 4 | 10.2\% | 8.5\% |
| 11 | Respiratory system | 54.60 | 62.71 | 69.56 | 65.07 | 67.68 | 78.97 | 15 | 11 | -6 | 4 | 15 | 5.2\% | 5.1\% |
| 12 | Dental problems | 10.78 | 13.55 | 24.91 | 51.93 | 59.45 | 62.44 | 26 | 84 | 108 | 14 | 5 | 1.1\% | 4.1\% |
| 13 | Gastro intestinal system | 63.56 | 73.22 | 81.30 | 73.30 | 75.05 | 78.89 | 15 | 11 | -10 | 2 | 4 | 6.1\% | 5.1\% |
| 14 | Skin problems | 20.98 | 24.90 | 26.84 | 28.31 | 30.41 | 32.34 | 19 | 8 | 5 | 7 | 6 | 2.1\% | 2.1\% |
| 15 | Musculo Skeletal system | 61.36 | 71.72 | 74.74 | 66.75 | 75.91 | 79.68 | 17 | 4 | -11 | 14 | 5 | 6.0\% | 5.2\% |
| 16 | Trauma and Injuries | 62.31 | 72.13 | 76.41 | 57.29 | 57.56 | 63.54 | 16 | 6 | -25 | 0 | 10 | 6.0\% | 4.2\% |
| 17 | Genito Urinary system | 55.32 | 62.38 | 67.38 | 68.98 | 67.83 | 73.78 | 13 | 8 | 2 | -2 | 9 | 5.2\% | 4.8\% |
| 18 | Maternity | 52.28 | 55.04 | 60.42 | 57.64 | 57.09 | 60.44 | 5 | 10 | -5 | -1 | 6 | 4.6\% | 3.9\% |
| 19 | Neonate conditions | 11.72 | 13.93 | 13.42 | 13.17 | 15.15 | 17.23 | 19 | -4 | -2 | 15 | 14 | 1.2\% | 1.1\% |
| 20 | Poisoning | 9.68 | 12.32 | 14.25 | 14.59 | 15.84 | 18.31 | 27 | 16 | 2 | 9 | 16 | 1.0\% | 1.2\% |
| 21 | Healthy individuals | 20.29 | 22.77 | 26.18 | 26.85 | 31.44 | 35.74 | 12 | 15 | 3 | 17 | 14 | 1.9\% | 2.3\% |
| 22 | Social care needs | 24.81 | 30.93 | 33.50 | 30.29 | 35.29 | 36.58 | 25 | 9 | -10 | 17 | 4 | 2.6\% | 2.4\% |
| 23 | Other (includes GMS/PMS) | 136.94 | 157.75 | 171.82 | 209.70 | 232.02 | 227.71 | 15 | 9 | 22 | 11 | -2 | 13.2\% | 14.9\% |
| 1 to 23 | All PBCs | 1052.12 | 1199.60 | 1307.76 | 1345.10 | 1452.91 | 1530.59 | 14 | 9 | 3 | 8 | 5 |  |  |

Notes: (i) The population figures for $2003 / 4,2004 / 5$ and $2005 /(6$ are identical (the total for England is $49,175,998$ ).
(ii) The corresponding figure for $2006 / 7$ is $50,476,231$, for $2007 / 8$ it is $50,695,989$, and for $2008 / 9$ it is $51,220,531$.
(iii) The spend per head figures are calculated by summing expenditure across all PCTs and dividing by the national population.
(iv) All figures are at current prices.

Table B4.2: table showing PCT expenditure per head by PBC, 2008/9: (a) unadjusted; (b) adjusted for local costs; and (c) adjusted for local costs and local need


Note: the above statistics relate to 152 PCTs and the mean expenditure figures will differ slightly from the national ones in Table B4.1 because the statistics across PCTs are not weighted for the size of each PCT's population.

## B5. Health outcome and other data

## B5.1 Health outcome data

Most studies of the relationship between expenditure and outcome have used some measure of mortality as an indicator of the latter. We too employ mortality as our outcome measure for two reasons: first, it is a relevant (but admittedly not comprehensive) measure of the outcome of health care expenditure; and second, it is available for more disease areas than any other outcome measure at PCT level.

Although mortality is available (by PCT) for several disease areas, it is not available for just over one-half of all programmes not least because it is simply not relevant for these programmes (e.g., for learning disabilities, vision problems, hearing problems, dental problems, and skin problems). Moreover, even where a mortality measure is available, the ICD10 coverage of the mortality data often falls short of the coverage of the expenditure data. For some programmes, therefore, we have combined the published mortality rates for two or more disease areas in an attempt to match the ICD10 coverage of the mortality data with that of the expenditure data.

Table B5.1 shows how we have attempted to marry the mortality data (column c) and the expenditure data (column a). However, and as Table B5.1 shows, the ICD10 coverage of the component mortality rates for some PBCs still falls short that of the expenditure data and the extent of this shortfall is illustrated by the ratio reported in the final column of Table B5.1. For example, the cancers and tumours programme covers all expenditure associated with ICD10 codes C00-C97 and D00-D49 but the PCTbased mortality data only relates to ICD10 codes C00-C97. At the national (all England) level, figures are available which show that, in 2008 , there were 62,072 deaths offthose aged under 75 years from codes C00-C97 and that there were 63,076 deaths from codes C00-697 and D00-D49 combined. In other words, the PCT level mortality data reflects $98.4 \%$ of all deaths associated with the expenditure codes. Initially, we did not adjust our cost of life (year) estimates for this mismatch but, as we will see in section B8.6, an adjustment has been made for this mismatch in the final calculation of the cost of a life (year) associated with expenditure for 2006/7. The same adjustment has also been applied to the cost of a life (year) estimates associated with expenditure for $2007 / 8$ and for 2008/9.

Of course, we acknowledge that mortality is a more relevant outcome indicator for some programmes (e.g., for circulatory problems) than for others (e.g., for epilepsy) and, for this reason, we would expect better results in some programmes than others. We also acknowledge that this focus on mortality ignores the impact of expenditure aimed at chronic care and at palliative care. Nevertheless, our focus on mortality is purely practical: it is both a widely available measure and it is clearly a relevant outcome indicator. Moreover, the approach adopted here is extendable in principle to other non-mortality based outcome indicators. We illustrate such an application in section B 8.8 where we use EQ-5D utility scores pre- and post- an operative procedure from the PROMs programme to generate a non-mortality-based outcome indicator, and we use this indicator to estimate our outcome model.

Table B5.1: table showing ICD10 coverage of the expenditure and outcome measures

|  | ICD 10 coverage of programme budgeting category | Number of deaths, <75years, 2008, England (ONS, VS3) corresponding to column aICD10 codes | ICD 10 coverage of best match PCT based mortality rate(s) | Number of deaths, <br> <75years, 2008, <br> ) England (ONS, VS3) <br> corresponding to column c ICD10 codes | Ratio <br> (d/b) |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  | column a | column b | column c | column d | column e |
| PBC 1 | Infectious diseases <br> (large parts of A00-B99)* | 1,968 | Infectious diseases (A00-B99) | 1,968 | 1.000 |
| PBC 2 | Cancers and tumours (C00-C97, D00-D49) | 63,076 | All cancers (C00-C97) | 62,072 | 0.984 |
| PBC 3 | Blood disorders (D500-D899) | 393 | No relevant mortality rate by PCT available | n/a | $\mathrm{n} / \mathrm{a}$ |
| PBC 4 | Endocrine, nutritional and metabolic problems (E000-E899) | $2,368$ | Diabetes <br> (E10-E14) | 1,501 | 0.634 |
| PBC 5 | Mental health <br> (F00-F69, Z55, Z56) | n/a | No relevant mortality rate available | n/a | $\mathrm{n} / \mathrm{a}$ |
| PBC 6 | Learning disability (F700-F739, F780-F849, F88-F90, Q90, Q91) | n/a <br> $\bigcirc 1$ | No relevant mortality rate available | n/a | $\mathrm{n} / \mathrm{a}$ |
| PBC 7 | Neurological system (G000-G999, Q000-Q079, R200-R999) | 5,238 | Epilepsy* <br> (G40-G41) | 713 | 0.136 |
| PBC 8 | Eye and vision problems (H000-H599, Q100-Q159) | $\mathrm{n} / \mathrm{a}$ | No relevant mortality rate available | n/a | $\mathrm{n} / \mathrm{a}$ |




Notes: (i) the listed ICD10 coverage of the programme budgeting expenditure data includes the major ICD10 codes covered.
(ii) the ICD10 coverage of PBC 1 includes large elements of codes A00-B99 but a substantial minority of these codes map to the respiratory (PBC11) and gastro-intestinal (PBC 13) programmes. We do not have the detailed deaths data to remove them from the total for A00-B99 and then to add them to the respiratory and gastro-intestinal programmes. Instead, we acknowledge that the number of deaths attributed to PBC 1 will be overstated (and that the adjustment ratio in column e will be too low), and that the number of deaths attributed to PBCs 11 and 13 will be understated (and that their adjustment ratios in column e will be too high) but this is the best that can be achieved given the available data.
(iii) the ICD10 coverage of the all England mortality data does not always match precisely that of the expenditure data or the PCT level mortality data; again, we have done the best that can be achieved given the available data. In particular: the national epilepsy mortality data relates to ICD10 G40 ( 687 deaths) but the PCT level data relates to G40 and G41 (annual average over 2007/8/9 is 713 deaths) ; the national renal failure mortality data relates to ICD10 N17-N19 ( 415 deaths) but the PCT level data relates to N18 (annual average over 2007/8/9 is 269 deaths); the national liver disease mortality data relates to ICD10 K70-K77 ( 6,020 deaths) buit the PCT level data relates to K70, K73-K74 (annual average over 2007/8/9 is 5,195 deaths); and there is no good ICD10 match for femur and skull fracture deaths using national VS3 data (the PCT level data relates to S72, S02, S06, T90: annual average over 2007/8/9 is 1,014 deaths). For these four cases we use the annual average number of deaths over 2007/8/9 from the PCT-level data as the numerator when calculating the coverage adjustment factor (column e).
(iv) the number of deaths in England for those aged under 75 years for the trauma, burns and injuries programme (column b) relates to 2004 and is for the secondary cause of death (Martin, Rice and Smith, 2012).
(v) the mortality rate for neonate conditions relates to deaths aged under 28 days for all ICD 10 codes but the expenditure data relates only to ' P ' ICD 10 codes. Hence the large adjustment factor of 8.213 because the coverage of the expenditure data is much smaller than that of the mortality data. However, at the very end of the project it became clear that although the number of deaths data for those aged under 75 years includes those dying at all ages under 75 years (including those at under 1 year), the disease specific years of life lost totals for those aged under 75 years excludes those dying at under 1 year of age and actually refers to those dying at ages 1 to 74 (the argument is that infant deaths are mostly a result of causes specific to the age and have different causes to disease specific deathslater in life). We therefore have two adjustment factors for the maternity and neonates programme: first, an adjustment factor for the number of deaths derived on the same basis as the adjustment factors for other programmes; and second an adjustment factor for the YLL that reflects both the YLL in the maternity and neonates programme, as well as the YLL associated with deaths that would have been attributed to other programmes had the individual died over 1 year of age. (NB The total number of deaths in England in 2008 of those aged under 1 year is 3,184 and if we divide 2,193 by ( $3,184+41$ ) we obtain the YLL coverage adjustment factor $(=0.679)$ for maternity and neonates.
(vi) the PCT level mortality rates are available from the NHS Information Centre website.

Previous studies using the programme budgeting data have employed two alternative mortality based outcome indicators: the under 75 years of age standardised mortality rate (SMR) and the under 75 years standardised years of life lost rate (SYLLR). The SMR gives equal weight to all deaths irrespective of the age at which they occur but the SYLLR gives greater weight to deaths that occur at earlier ages.

We employed both the SMR and the SYLLR when undertaking some preliminary sensitivity analysis (i.e., in section B8.2 when considering, for example, which measure of need to use), but elsewhere we have focussed solely on a measure of the avoidable years of life lost (YLL). ${ }^{11}$ This is calculated by summing over ages 1 to 74 years the number of deaths at each age multiplied by the number of years of life remaining up to age 75 years. The crude YLL rate is simply the number of years of life lost divided by the resident population aged under 75 years. Like conventional mortality rates, the crude YLL rate can be age standardised to eliminate the effects of differences in population age structures between areas, and this (age) standardised YLL rate is the health outcome variable generally employed in this study (Lakhani et al., 2006, p379).

Descriptive statistics for the SYLLRs employed in this study are shown in Table B5.2. For example, for all deaths over the three year period from 2006 to 2008, the annual SYLLR across allPCTs for those aged under 75 years averaged 467 years of life lost per 10,000 population, but this rate varied considerably across PCTs, ranging between 288 and 749 years of life lost per 10,000 population. Similarly large variations in the mortality rate across PCTs are evident for other disease groups. 12

[^45]Table B5.2: table showing descriptive statistics for the mortality variables



| trauma, SMR, $2008 / 9 / 10$ (sum of femur and skull fracture rates) | 151 | 1.9 | 0.8 | 0.1 | 4.4 |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| infant mortality rate, $<28$ days per 1,000 live births, $2002 / 3 / 4$ | 303 | 3.4 | 1.3 | 0.9 | 7.8 |
| infant mortality rate, $<28$ days per 1,000 live births, $2004 / 5 / 6$ | 130 | 3.4 | 0.9 | 1.2 | 6.2 |
| infant mortality rate, $<28$ days per 1,000 live births, $2006 / 7 / 8$ | 152 | 3.3 | 1.0 | 1.4 | 6.4 |
| infant mortality rate, $<28$ days per 1,000 live births, $2007 / 8 / 9$ | 151 | 3.2 | 1.0 | 1.2 | 6.9 |
| infant mortality rate, $<28$ days per 1,000 live births, $2008 / 9 / 10$ | 151 | 3.2 | 1.0 | 1.2 | 6.9 |

Note: the SYLLRs are directly age-standardised rates and are expressed as rates per 10,000 European Standard population. Source: NHS Information Centre website.

## B5.2 Other variables

We employ an instrumental variable (IV) estimation technique to estimate our outcome and expenditure equations because (i) own programme expenditure is likely to be endogenous in the outcome equation and (ii) other programme need is likely to be endogenous in the own programme expenditure equation. IV estimation is described in section B6.2 but basically it involves replacing the endogenous yariable in the equation of interest with its predicted value from an OLS regression which regresses the endogenous variable on a set of instrumental variables. These instruments should be good predictors of the endogenous variable (i.e., they should be relevant and strong predictors) but should be appropriately excluded from the equation of interest (i.e., they should be valid instruments).

We have a number of potential instruments available, mostly derived from 2001 Population Census. In our earlier studies we found that a small sub-set of these instruments proved sufficient to generate plausible results and these included:

- the proportion of the population providing unpaid care
- the proportion of households that are one pensioner households
- the index of multiple deprivation
- the proportion of the population in the white ethnic group.

We also had available a further set of potential instruments and, where our more limited set of instruments failed to generate plausible results, we extended our instrument search to include this wider set of variables. This extended set of instruments included:

- the proportion of residents borm outside the European Union
- the proportion of the population of working age (16-74) with a limiting long term illness
- the proportion of the population aged $16-74$ with no qualifications
- the proportion of the Population aged 16-74 that are full-time students
- the proportion of households without a car
- the proportion of households that are owner occupied
- the proportion of households that are rented from a LA or HA
- the propottion of households that are rented from private landlords
- the proportion of households that are lone parent households with dependent children
- the proportion of the population aged 16-74 that are permanently sick
- the proportion of those aged 16-74 that are long-term unemployed
. the proportion of those aged 16-74 in employment that are working in agriculture
- the proportion of those aged 16-74 in managerial and professional occupations.

Details of the construction of all instruments are shown in Table BA. 2 in the Annex.
Our instruments reflect factors, such as socio-economic deprivation and the availability of informal care in the community, which might indirectly impact upon mortality rates and/or health care expenditure levels. As we shall see, although our instruments 'pass' the appropriate statistical tests, some commentators claim that such tests may have 'low power' to detect the presence of invalid instruments. Consequently in section B9 we examine how sensitive our results are to the presence of invalid instruments.

Table B5.3 reports descriptive statistics for the socio-economic and needs variables as available for the regression analysis of programme budgeting (PB) expenditure data for 2007/8 and for 2008/9 (these statistics are for the variables in absolute form). For example, on average, lone pensioner households comprise $14 \%$ of all households, the 'white ethnic' group accounts for $89 \%$ of the population, and $10 \%$ of the population provide unpaid care.

In addition to the instrumental variables, Table B5.3 also report descriptive statistics for various other variables available for the regression analysis including the of Department of Health's 'need for health care' index (this incorporates the CARAN formula for HCHS and reflects need across all health care services), its need for HIV services index, and its need for maternity services index. The latter two indices are used to either supplement or replace the all service measure of need when estimating oun models. The 'need for health care' index averages about 1 but varies substantially, with some PCTs having a needs index more than $25 \%$ below the national average and others facing a need for health care more than $30 \%$ above the national average.

Table B5.3 also reports descriptive statistics for some disease prevalence rates (e.g., for diabetes and for epilepsy) and, again, these are used to either supplement or replace the all service measure of need when estimating our models

Finally, the MFF index shows that input prices in the most expensive PCT are almost $20 \%$ above those in the least expensive PCT.

Table B5.3: table showing descriptive statistics for the instrumental and other variables

| Description | Obs | Mean | Std. Dev. | Min | Max |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Proportion of residents born outside the European Union | 151 | 0.0794 | 0.0876 | 0.0088 | 0.3817 |
| Proportion of population in white ethnic group | 151 | 0.8927 | 0.1299 | 0.3942 | 0.9926 |
| Proportion of population of working age (16-74) with LLT illness | 151 | 0.1182 | 0.0250 | 0.0709 | 0.1798 |
| Proportion of population providing unpaid care | 151 | 0.0990 | 0.0118 | 0.0662 | 0.1221 |
| Proportion of population providing unpaid care ( $<20 \mathrm{hrs}$ week) | 151 | 0.0667 | 0.0079 | 0.0461 | 0.0817 |
| Proportion of population providing unpaid care (20-49 hrs week) | 151 | 0.0113 | 0.0025 | 0.0065 | 0.0195 |
| Proportion of population providing unpaid care ( $>50 \mathrm{hrs}$ week) | 151 | 0.0210 | 0.0051 | 0.0093 | 0.0353 |
| Proportion of population aged 16-74 with no qualifications | 151 | 0.2960 | 0.0642 | 0.1301 | 0.4555 |
| Proportion of population aged 16-74 that are full-time students | 151 | 0.0720 | 0.0270 | 0.0425 | 1626 |
| Proportion of households without a car | 151 | 0.2932 | 0.1046 | 0.1325 | 0,576 |
| Proportion of owner occupied households | 151 | 0.6692 | 0.1128 | 0.2891 | 0.8205 |
| Proportion of households in rented social (LA/HA) housing | 151 | 0.2071 | 0.0918 | 0.081 | 0.5356 |
| Proportion of households in rented private housing | 151 | 0.0924 | 0.0449 | 0.0349 | 0.2961 |
| Proportion of lone pensioner households | 151 | 0.1434 | 0.0184 | 0.0979 | 0.1942 |
| Proportion of one parent households | 151 | 0.0684 | 0.0180 | 0.0401 | 0.1207 |
| Proportion of population aged 16-74 that are permanently sick | 151 | 0.0574 | 0.0213 | 0.0242 | 0.1215 |
| Proportion of population aged 16-74 are long-term unemployed | 151 | 0.0113 | 0.0052 | 0.0036 | 0.0287 |
| Proportion of 16-74 in employment that are in agriculture | 51 | 0.0117 | 0.0119 | 0.0016 | 0.0668 |
| Proportion of those aged 16-74 that are in professional occupation |  | 0.2672 | 0.0688 | 0.1470 | 0.4958 |
| Index of Multiple Deprivation 2007 |  | 23.8098 | 9.1168 | 8.0857 | 48.2627 |
| Need index (incorporates CARAN formula) |  | 1.0253 | 0.1334 | 0.7311 | 1.3479 |
| MFF index for HCHS and prescribing | 151 | 1.0021 | 0.0559 | 0.9410 | 1.1243 |
| Diabetes prevalence rate 2007/8 (\%, over 17 years) | 151 | 5.4872 | 0.7982 | 3.22 | 8.51 |
| Epilepsy prevalence rate 2007/8 (\%, over 18 years) | 151 | 0.7884 | 0.1489 | 0.41 | 1.09 |
| HIV need index | 151 | 1.1848 | 1.4984 | 0.1648 | 8.3332 |
| Chronic kidney disease 2007/8 (\%, over 18 years) | 151 | 4.1687 | 1.2711 | 1.35 | 8.41 |
| Maternity need index | 151 | 1.0345 | 0.2106 | 0.6845 | 1.8129 |
| Raw (unadjusted) population 2007/8) | 151 | 335,735 | 196,501 | 90,142 | 1,264,298 |

[^46]
## B6. Estimation issues and strategy

## B6.1 Introduction

The theoretical framework suggests the specification and estimation of a system of equations, with an expenditure and health outcome equation for each of the 23 programmes of care. However, this approach makes infeasible data demands, requiring variables to identify expenditure, need, environmental factors and health outcomes in each of the 23 programmes of care. Moreover, mortality rates are available for less than half of the 23 programmes. Rather than estimate a system of equations, we proceed on a programme-by-programme basis, estimating health outcome and expenditure equations for those programmes for which mortality data is available.

In line with the theoretical framework presented in section B3, we specify the following expenditure (0.1) and health outcome (6.2) models for each of the $J$ programmes of care ( $J=23$ )

$$
\begin{array}{ll}
\mathrm{x}_{\mathrm{i}}=\mathrm{a}_{1}+\sum \mathrm{b}_{1 \mathrm{j}} \cdot \mathrm{n}_{\mathrm{ij}}+\mathrm{d} \mathrm{y}_{\mathrm{i}}+\mathrm{e}_{1 \mathrm{i}} & \mathrm{j}=1, \ldots, 23 \\
\mathrm{~h}_{\mathrm{i}}=\mathrm{a}_{2}+\mathrm{b}_{2} \mathrm{n}_{\mathrm{i}}+\mathrm{fx}_{\mathrm{i}}+\mathrm{e}_{2 \mathrm{i}} & \tag{6.2}
\end{array}
$$

where $\mathrm{x}_{\mathrm{i}}$ is the expenditure in $\mathrm{PCT}_{i}$ in the selected programme $\mathrm{n}_{\mathrm{i} j}$ is the need for care in $\mathrm{PCT}_{\mathrm{i}}$ in programme ${ }_{j}$ $\mathrm{y}_{\mathrm{i}}$ is the total budget for $\mathrm{PCT}_{i}$
$\mathrm{h}_{\mathrm{i}}$ is the health gain in $\mathrm{PCT}_{i}$ in the selected programme $\mathrm{n}_{\mathrm{i}}$ is the need for care in PCT $_{i}$ in the selected programme.

Ideally we should employ a programme specific indicator of the level of need for each care programme but these are not readily available. When estimating both the outcome and expenditure models we therefore proxy the own programme health care need using the 'needs' component of the Department of Health's resource allocation formula. ${ }^{13}$ This needs element is specifically designed to adjust PCT allocations for local health care needs and accordingly, ceteris paribus, we would expect a positive relationship between expenditure $x_{i}$ and need niforeach programme of care. We would also expect a positive relationship between need $n_{i}$ and adverse health outcomes $h_{i} .{ }^{14}$

The expenditure model includes both the own programme health care need (which is proxied using the 'needs' component of the Department of Health's resource allocation formula) and the need for health care in all other programmes. When estimating the expenditure model previous studies have proxied the need for health care in other(competing) programmes using the mortality rate in those other programmes. The precise definition of the programmes included in the 'other programme' mortality rate has varied a little, but here all of our preferred results from 2006/7 onwards use the 'all cause mortality rate excluding the mortality rate in the programme of interest' as the proxy for need in other programmes. ${ }^{15}$

## B6.2 IV estimation

We do not use OLS to estimate equations (6.1) and (6.2) because both are likely to contain an endogenous regressor. Expenditure in the outcome equation (6.2) and other programme need in the expenditure equation (6.1) are both likely to be endogenous and, in the presence of an endogenous

[^47]regressor, OLS is both a biased and an inconsistent estimator. Instead, we use instrumental variable (IV) estimation and implement two-stage least squares (2SLS) using the -ivreg2- routine in Stata v11[21]. Unlike OLS, IV is a consistent estimator in the presence of an endogenous regressor and, although in finite samples the IV estimator will be biased, the belief is that (providing certain assumptions are met) this bias will be less than that associated with OLS.

For the health outcome equation, IV estimation can be viewed as finding variables (instruments) that are good predictors of programme expenditure but which are appropriately excluded from the equation of interest (that is, from equation 6.2). The assumption is that the instruments and exogenous variables from the equation of interest impact upon the health outcome through their impact on expenditure only, and that they do not have a direct effect on the outcome. ${ }^{16}$ If, on the other hand, an instrument reflects unobserved factors that affect both expenditure and mortality directly, then the IV estimator becomes both biased and inconsistent. Such an instrument is said to be 'invalid' because it belongs in the equation of interest in its own right.

We have a number of potential instruments available, mostly derived from 2001 Population Census, and these are described in section B5.2. In our earlier studies we found that a small sub-set (four) of these instruments often proved sufficient to generate plausible results and we commenced our empirical work with these. If plausible results were not obtainable with some combination of theseffour instruments, we employed an extended instrument set. Further details of the identification of suitable instruments for each model can be found in section B7.3.

The available instruments reflect factors, such as socio-economic deprivation and the availability of informal care in the community, which might indirectly impact upon mortality rates and/or health care expenditure levels. The set of instruments associated with each estimated equation was selected on both technical and pragmatic grounds. From a pragmatic pointof (viey, we require a parsimonious set of instruments that satisfy the necessary technical criteria. These are, firstly, that they have face validity, that is, that they are plausible determinants of the endogenous variable being instrumented, and secondly, that the instruments are both relevant and valid. The relevance of an instrument set refers to its ability to predict the endogenous variable of concern, whereas validity refers to the requirement that instruments should be uncorrelated with the error term in the equation of interest. The set of instruments was modified if, for example, the Hansen-Sargan test suggested that the set under test was not valid.

Should the instrument set be strong, releyant and valid, 2SLS will produce consistent estimates of the parameters of the reduced form models. We subject the instrument sets to tests for validity using the Sargan-Hansen test of overidentifying restrictions. The joint null hypothesis is that the instruments are valid instruments, i.e., they are uncorrelated with the error term, and that the excluded instruments are correctly excluded from the estimated equation. A rejection of the null hypothesis casts doubt on the validity of the instruments. We test for instrument relevance using Shea's[22] partial R-squared measure; this reflects the correlation between the excluded instruments and the endogenous regressor. However, even if valid and relevant, non-zero but small correlations between the instruments and the endogenous regressors canlead to the problem of weak instruments. This can be the case even where correlations are shown tobe significant at conventional levels of testing and sample sizes are large[23]. The IV estimator becomes abiased estimator if the instruments are weakly correlated with the endogenous regressors, and the extent of the bias can be specified relative to the bias of the OLS estimator.

For the case of a single regressor, Staiger and Stock[24] suggest applying the criterion that if the first-stage F-statistic, testing the null hypothesis that the instrument set does not significantly predict the endogenous regressor, is less than 10 then the instruments can be thought to be weak. Stock and Yogo[25] extend these ideas to the case where there can be multiple endogenous regressors and propose a test for the null that the instruments are weak and provide appropriate critical values. This is an extension of the Cragg and Donald [26] test for instrument relevance. For the case of a single endogenous

[^48]regressor, the Cragg-Donald statistic is simply the F-statistic of the test of the hypothesis that the instruments do not enter the first-stage regression. Stock and Yogo provide critical values of the Fstatistic (and the Cragg-Donald statistic for multiple endogenous regressors) that tabulates the ratio of 2SLS bias to the bias of OLS. The weakness or otherwise of the instruments can then be assessed by the relative bias exceeding a given threshold (for example, 2SLS bias exceeding $5 \%$ of OLS bias). ${ }^{17}$

To ensure the robustness of our estimates to arbitrary heteroskedasticity, we estimate our models with Stata's -robust- option. The Cragg-Donald statistics are not valid in the presence of heteroskedascity. We therefore report the Kleibergen-Paap LM statistic (testing instrument relevance) and the Kleibergen-Paap F statistic (testing for weak instruments) which are valid in the presence of heteroskedascity. For further details of these tests see Baum, Schaffer and Stillman[21].

A general test of model specification is provided through the use of Ramsey's [27] reset test for OLS and an adapted version of the test for instrumental variables [28]. ${ }^{18}$ The tests are more properly thought of as tests of a linearity assumption in the mean function or a test of functional form restrictions and omitted variables[29] and can be useful as a general check of model specification.

Finally, we check that the presumed endogenous variable is in fact endogenous using the test proposed by Durbin [30]. If the null hypothesis of exogeneity cannot be rejected, then we also use the OLS estimator. And, although our instruments 'pass' the appropriate statistical tests, some commentators claim that such tests may have 'low power' to detect the presence of invalid instruments. Consequently in section B9 we examine how sensitive our results are to the relaxation of the assumption that the instruments are valid.

## B6.3 Other estimation issues

In this research we build on previous studies that have used the PB data to estimate the outcome and expenditure models described in section B6.1. This previous research was undertaken over a period of years and a number of changes were made between these studies (these were sometimes forced on the researchers by, for example, data availability considerations). Here we persevere with the previous approach used to analyse the $2005 / 6 \mathrm{~PB}$ data $[5]$, but we make some changes to the way in which the 2006/7 (and subsequent) PB data are analysed.

In the next section we start by re-visiting the results obtained by Martin, Rice and Smith [5] who used the 2005/6 PB data. In 2005/6 there were 303 PCTs but a series of mergers reduced this total to 152 in 2006/7. These mergers exacerbated greatly the difference in size between the PCTs and so from 2006/7 it makes less sense to give each PCT equal weight in any regression. This is discussed further in section B8.2 when we come to estimate out model using $2006 / 7 \mathrm{~PB}$ data.

Different PCTs face different costs when buying health care inputs. For example, some health economy input prices are ap to $40 \%$ higher in London and the south east of England than elsewhere. In a previous study[3], we used the Market Forces Factor Index (MFF) that feeds into the Payment by Results tariffs for 2007/8 to adjust programme budgeting expenditure in 2006/7 for local input prices[31]. This index only reflects costs associated with the purchase of HCHS services but this was the only index available for the new (post October 2006) set of PCTs at the time of that study. Since then, a more comprehensive set of MFF indices for the 152 PCTs has been published [32]. In section B8.2 we investigate the use of alternative weighted averages of the HCHS, prescribing, and GMS/PMS MFF indices with weights reflecting the national share of expenditure across these three categories (these weights are $76.3 \%, 12.4 \%$, and $11.3 \%$ respectively). ${ }^{19}$ For 2005/6, however, we persevere with the MFF employed in the original Martin, Rice and Smith study [5], namely the HCHS MFF [33].

[^49]Estimation of the expenditure equation for any individual programme requires a proxy for the need for health care across all other programmes. Previous studies of PB expenditure in 2004/5, 2005/6 and 2006/7 have used the circulatory disease mortality rate as a proxy for the need for health care in other programmes in the cancer expenditure equation, and the cancer mortality rate as the proxy for need in other programmes in the circulatory disease expenditure equation $[2,3,5]$. As these are both programmes that attract considerable expenditure and record considerable mortality, it is not implausible that mortality and expenditure in one of the programmes will impact upon expenditure in the other. For other programmes (e.g., respiratory problems and gastro-intestinal problems) Martin, Rice and Smith[3, 5] used the all cause mortality rate as a proxy for the 'need in other programmes' variable when analysing expenditure in both 2005/6 and 2006/7. Here, however, we persevere with the previous approach when using 2005/6 PB data but, from 2006/7, in all programmes we proxy the need for health care in other (competing) programmes using the mortality rate in those other programmes (i.e., the all cause mortality rate minus the own programme mortality rate).

Finally, one data transformation that has been applied in all previous studies and is applied here too is to log transform all variables so that parameter estimates can be interpreted as elasticities. In other words, a regression coefficient of 0.5 implies that a $1 \%$ increase in the regressor is associated with a $0.5 \%$ increase in the dependent variable.

## B. EMPIRICAL RESULTS

## B7. Analysis of programme budgeting expenditure for 2005/6

This work builds on previous studies. Martin, Rice and Smith [2] reported outcome elasticities for two programmes (cancer and circulatory disease) using expenditure data for 2004/5 and pooled mortality data for 2002, 2003 and 2004. Martin, Rice and Smith [5] extended their preliminary analysis to include several other programmes and, in this extension, they used updated expenditure data (for 2005/6). However, the authors found it difficult to obtain sensible outcome models for some programmes of care. Here we commence our empirical work with an attempt to obtain plausible outcome models for those programmes that defeated Martin, Rice and Smith in their study. [5]

## B 7.1 Construction of an alternative measure of need

Our preferred measure of the need for health care is calculated from the Department Health s programme budgeting ( PB ) dataset. This dataset includes programme budgeting expenditare for each care programme as well as the raw population and the 'unified weighted' population foreach PCT. The unified weighted population incorporates adjustments to the raw population for both the need for health care as well unavoidable variations in local input costs. The latter are captured via an index which is known as the market forces factor (MFF). By removing the raw population and MEF adjustment from the unified weighted population we are left with the implied level of need, and this is the measure of need that was initially used in the estimation of the model.[5]

The Department of Health PB measure of need associated with expenditure for 2005/6 incorporates the AREA resource allocation formula. This has since been replaced with the CARAN formula and recent work by colleagues at York and the Nuffield Trust has investigated the possibility of constructing a person based resource allocation (PBRA) measure of need) [34]. We therefore decided to investigate the possibility of applying PBRA methods to the constraction of an alternative measure of need.

The construction of all of these measures of need involves two steps. The first step requires the estimation of the econometric relationship betryeen the previous utilisation of services and the characteristics of the local areas as existed at the time of the utilisation (e.g., their demographic profile and other indicators of service need such as socio-economic measures of deprivation). The second step involves the use of this relationship to predict future health care use given predictions about future demographic characteristics and socio-eeonomic measures of deprivation.

The major difference between the AREA and CARAN formulae and the PBRA formula is that the former largely use small area based indicators of socio-economic characteristics as indicators of the need for health care, whereas the latter largely obviates the requirement for these through the extensive use of individual based indicators of need. In particular, the PBRA formula employed here is based on an analysis of inpatient and outpatient cost data for $2007 / 8$ for $10 \%$ of the entire population of England
[34]. As regressors the PBRA utilisation model includes:
(a) 38 agetgender dummies;
(b) 150 ICD10 morbidity markers for each patient reflecting their use of inpatient services in the previous two years (that is, in 2005/6 and 2006/7 combined);
(c) 4 hospital encounter variables for each patient reflecting the intensity of their use of both outpatient and inpatient services in the previous two years (that is, in 2005/6 and 2006/7 combined);
(d) 10 small area based indicators of either local deprivation or health care supply characteristics; and
(e) 151 PCT dummies (reflecting variations in health care supply).

The coefficients from this modelling procedure are applied to patient registration data as at 1 April of the year for which the measure of need is required. Here we are studying expenditure in 2005/6 and so we applied the results of the modelling to patient registration data as at 1 April 2005. This requires the construction of a dataset containing the patient registration details of all 50 million patients registered with an English practice as at this date. To this we added the patient's age and gender as at April 2005.

We also added each patient's ICD10 morbidity markers and their encounter variables for 2003/4 and 2004/5 combined. Each patient's address (LSOA) is also added to the dataset and this is used to attach the small number of indicators reflecting the LSOA's socio-economic and health care supply characteristics.

Given this dataset, the calculation of PCT need (given supply) proceeds as follows. First, calculate the national average supply effect. This is the sum of the products of the national average values of the supply variables for the population as at 1 April 2005 and the relevant the regression coefficients.

Second, ignore supply and calculate PCT need. This involves calculating the PCT average values of the needs variables by age and gender group for the population as at 1 April 2005. Next, for each PCT calculate need by age and gender as the sum of the products of the mean values of the needs variables and their respective regression coefficients. Then total PCT need is the sum of (need in each age/ gender group multiplied by the number of patients in that age/gender group).

Finally, need given supply is calculated as total PCT need plus the number of patients multiplied by the national average supply effect. PCT need per person is simply total PCT need divided by the PCT population. Further details of how to use the results of the PBRA modelling to derive PCT weighted needs indices are presented in Dixon et al.[34]

## B7.2 Re-estimation of models using a new measure of need

We re-estimated the outcome and expenditure models for the big four programmes as reported by Martin, Rice and Smith [5] using the new (PBRA based) measure of need. ${ }^{20}$ In summary, the results for the cancer programme were acceptable but not quite as good as previously obtained, and the results for circulation problems, gastro-intestinal problems, and respiratory) problems were poor (e.g., the signs on the expenditure and need variables in the outcome equation were counter-intuitive). These were unanticipated results and we were curious to know why our alternative measure of need performed less well than the more established measure.

We undertook a brief comparison of the two measures of need. Figure B7.1 provides a scatter plot of the PB and PBRA measures of need. There is a clear positive correlation between the two measures (correlation coefficient=0.6146), and the summary statistics in Table B7.1 suggest that they have similar ranges.

[^50]

Figure B7.1: graph showing scatter plot of PB measure of need and PBRA measure of need

Table B7.1: table showing summary statistics for PB and PBRA based measures of need

|  | Number <br> of PCTs | Mean | Std. Dev. | Min | Max |
| :--- | :--- | :--- | :--- | :--- | :--- |
| Variable | 295 | 1.0062 | 0.1511 | 0.6883 | 1.4889 |
| PB need | 303 | 1.0146 | 0.1448 | 0.6884 | 1.4554 |

Note that there are only 295 PCTs with a PB based measure of need because
only 295 of the 303 PCTs were used to estimate our outcome and expenditure
models (due to a lack of data for some PCTs).

Table B7.2 reports values for the PB and PBRA based measures of need for selected types of PCTs. These figures suggest that:

- the PBRA measure attributes more need to the least needy areas as defined by the PB measure (see Table B7.2a);
- the-PBRA measure attributes more need to the coastal/retirement areas than does the PB measure (see Table B7.2b); and
the PBRA measure attributes far less need to inner city areas than does the PB measure (see Table B7.2c).

Table B7.2: table showing values for the PB and PBRA based measures of need for selected types of PCT

| (a) Examples of more affluent areas | PB need | PBRA need |
| :---: | :---: | :---: |
| Wokingham PCT | 0.6883 | 0.7703 |
| Blackwater Valley and Hart PCT | 0.7376 | 0.8395 |
| Bracknell Forest PCT | 0.7410 | 0.8262 |
| Royston, Buntingford and Bishop's Stortford PCT | 0.7426 | 0.8484 |
| Windsor, Ascot and Maidenhead PCT | 0.7460 | 0.8404 |
| Woking PCT | 0.7499 | 0.8097 |
| Chiltern and South Bucks PCT | 0.7515 | 0.8530 |
| Uttlesford PCT | 0.7583 | 0.8861 |
| North East Oxfordshire PCT | 0.7588 | 0.8372 |
| South Cambridgeshire PCT | 0.7619 | 0.9647 |
| (b) Examples of coastal/retirement areas | PB need | PBRA need |
| Suffolk Coastal PCT | 0.9159 | 1.0815 |
| Western Sussex PCT | 0.9613 | 1.3248 |
| North Somerset PCT | 0.9651 | 1.1746 |
| Poole PCT | 0.9860 | 1.2045 |
| South and East Dorset PCT | 1.0079 | 1.2439 |
| Fylde PCT | 1.0304 | 1.2157 |
| Southport and Formby PCT | . 0657 | 1.2141 |
| North Norfolk PCT | 1.0658 | 1.3684 |
| Adur, Arun and Worthing PCT | 1.0716 | 1.2641 |
| East Devon PCT | 1.0870 | 1.3325 |
| (c) Examples of inner city areas | PB need | PBRA need |
| Brent PCT | 0.9848 | 0.6991 |
| Lambeth PCT | 1.0454 | 0.7512 |
| Islington PCT | 1.1222 | 0.9014 |
| Southwark PCT | 1.1412 | 0.8163 |
| Newham PCT | 1.1746 | 0.7897 |
| City and Hackney PCT | 1.1849 | 0.8472 |
| Bradford City PCT | 1.2131 | 0.8757 |
| Tower Hamlets POT | 1.2192 | 0.9299 |
| Heart of Birmingham Teaching PCT | 1.2466 | 0.9052 |
| Central Manchester PCT | 1.2965 | 0.9262 |
| Gentral Liverpool PCT | 1.4065 | 1.0948 |

Although these differences are at first perplexing, they become more understandable when it is noted that the PB and PBRA measures record the level of need across different baskets of services. The PB measure of need refers to all health care activity, that is, Hospital and Community Services (HCHS), prescribing, and GMS/PMS (primary care), but the PBRA model only incorporates hospital activity (and it excludes mental health and maternity from this).

The need for hospital based services is less related to deprivation than are other health care services. Hence the PBRA measure of need - because it only relates to hospital services - re-distributes need away from the more deprived PCTs and towards the more affluent ones. Moreover, expenditure on cancer
services is largely hospital based and hence a measure of need based on HCHS spend alone will be reasonably satisfactory for cancer (as indeed we found). However, such a measure of need will perform less well for other programmes (e.g., circulatory disease), where more of the expenditure is on prescribing and/or primary care.

To test these hypotheses we need to compare our PBRA measure of need with a PB measure of need that only relates to acute services (i.e., that excludes maternity and mental health, and all prescribing and GMS/PMS). The Department of Health's measure of need used for the 2005/6 allocations employs the AREA formula for HCHS. This formula does not permit a separation of acute and maternity need and so we cannot compare the PBRA measure of need for 2005/6 with the PB measure for 2005/6 for the same group of specialties (i.e., for acute services excluding maternity and mental health).

However, the CARAN formula, first implemented for the 2009/10 allocations, does distinguish between acute and maternity. But this formula has only been applied to the new (post October 2006, $n=152$ ) PCTs whereas our PBRA-based measure is for the old (pre October 2006, $\mathrm{n}=303$ ) PCTs because we are modelling PB expenditure in 2005/6. However, not all of the old PCTs were involved in mergers in October 2006. Thus for about half of all PCTs, we can compare our PBRA based measure of need for 2005/6 with the CARAN-based measure of need for 2009/10 for the same set of HCHS services (i.e., for acute excluding maternity and mental health).

The correlation between PBRA need and CARAN acute need is much higher (correlation coefficient $=0.8722$ ) than that between the PBRA and PB need measures. And an inspection of the values taken by the various need indices (e.g., for acute, maternity, and mental health) for the inner city PCTs (where the PBRA and PB measures of need diverge the most) supports the hypothesis that it is the different service coverage of the PBRA and PB measures of need that explains why they are so poorly correlated (see Table B7.3).

For example, the PB index suggests that per capita need in Newham PCT is $17 \%$ above the national average but the PBRA index suggests that it is $21 \%$ below the national average. We believe that this difference is due to the fact that the PB inder relates to all services whereas the PBRA index only relates to acute services. The separate figures for acute, maternity and mental health need from the CARAN formula confirm this hypothesis: CARAN acute need, like PBRA acute need, is well below the national average, but maternity and mental health need are well above it.

Table B7.3: table showing comparing PB, PBRA and CARAN need indexes for selected inner city PCTs

| PCT | PB need <br> (all <br> services) | PBRA need <br> (acute) | CARAN need |  | acute |
| :--- | :--- | :--- | :--- | :--- | :--- |

## B7.3 Re-estimation of poorly performing models with an extended instrument set

Martin, Rice and Smith [5] found it difficult to obtain sensible outcome models for some programmes of care. As we were unable to find an improved measure of need, we sought to improve the outcome and expenditure models reported in Martin, Rice and Smith[5] through the use of an extended set of regressors/instruments. Martin, Rice and Smith[5] had focussed on the use of four instruments but here we extend the modelling to include an additional 13 regressors/instruments (born outside EU, limiting long-term illness, no qualifications, full-time students, no car households, owner occupiers, privately rented, socially rented, lone parents, permanently sick, long-term unemployed, work in agriculture, work in professional occupation). Further details about these variables can be found in section B5.2 and precise details about how they were constructed can be found in Table BA. 2 in the annex.

For each PBC, our modelling strategy with these additional regressors/instruments was the same:

(a) first, estimate an IV model using our preferred set of regressors (with need, budget, and other programme need for the own programme spend model, and with need and spend for the outcome model) and preferred set of instruments (proportion of households that are lone pensioner households, per cent of the population providing unpaid care, the IMD 2000, and the per cent of the population in the white ethnic group). Then adjust this set of instruments if necessary (e.g., remove from the instrument set or add an instrument to the regressor set if the Hansen-Sargan test indicates that this is appropriate). Estimate an OLS version of the IV model if the theoreticallyendogenous regressor is exogenous according to the relevant statistical test.
(b) second, if (a) fails to generate a reasonable model, add the same additional variables to both the regressor and instrument sets. Then eliminate insignificant regressors (least significant first, but always retaining e.g., the budget and other need variables in the expenditure model, and own programme spend in the outcome model). Then eliminate insignificant instruments until a reasonable model is obtained. Again, estimate an OLS version of the IV model ifthe theoretically endogenous regressor is exogenous according to the relevant statistical test.

## B7.4 IV estimates of outcome and expenditure models

The above approach generates preferred outcome and expenditure models for each of the programmes with a mortality based outcome indicator. Outcome models are shown in Table B7.4 with expenditure models in Table B7.5. The corresponding first-stage regression results can be found in Tables BA. 3 and BA. 4 respectively in the annex.

The first four results in Table B7. 4 show the outcome model for the big four programmes (i.e., for cancer, circulatory disease, respiratory problems and gastro-intestinal problems). In all four programmes the need variable has positive and significant effect on mortality, and expenditure has the anticipated negative effect. The diagnostic statistics reveal that, in all four cases, own programme expenditure is endogenous and that the instruments are valid. They also suggest that the instruments are relevant and there is no evidence that the instruments are weak. The Pesaran-Taylor test suggests that there is no evidence of model mis-specification.

The results for the other programmes are similar to but more diverse than those for the big four programmes. This is to be anticipated because mortality is a much rarer outcome in these programmes than it is in, say, the cancer programme. Own programme expenditure is not endogenous in the next two programmes (infectious diseases and neurological problems) and we revert to the use of the OLS estimator. Expenditure has the anticipated negative effect on mortality in the infectious disease programme but this is not statistically significant. The all service measure of need is not relevant for this PBC; instead, we find that a measure of need associated with HIV is positively associated with mortality, as is a measure of deprivation (households with no car). Mortality from epilepsy is negatively associated with expenditure in the neurological programme. The need for health care variables has a positive and significant effect on mortality.

Expenditure and need have the anticipated effects on mortality in the trauma and injuries programme. In addition, the provision of unpaid care appears to be associated with an increase in mortality from fractures. This might be because the availability of care allows the elderly to continue to live in their own home and that they are more likely to fall and die from a fall at home than they are in alternative accommodation (such as in a residential home or sheltered housing).

Expenditure has the anticipated negative effect on mortality in the neonates programme where the generic all service measure of need has been replaced with two more programme specific indicators of need (the proportion of births that are low birth weight births and the proportion of households that are lone parent households).

The final two results both employ the OLS estimator. Expenditure in the genitor-urinary programme has a small negative effect on mortality (from renal problems). The prevalence of one parent households and non-white residents both seem to be positively associated with mortality.

Finally, expenditure has the anticipated negative effect on mortality in the endocrine problems programme where the generic all service measure of need has again been replaced with a more programme specific indicator of need (the diabetes prevalence rate). Mortality in this programme is also positively associated with the Index of Multiple Deprivation 2000.

The first four results in Table B7.5 show the expenditure model for the bof four programmes (i.e., for cancer, circulatory disease, respiratory problems and gastro-intestinal problems). In all four programmes both the need and budget variables have a positive and significant effect on own programme expenditure. In addition, the proxy for need in other programmes is negativejand significant in all four cases. In the circulatory expenditure programme the provision of unpaid care is associated with more expenditure (patients may buy care in more affluent areas), as is the proportion of residents in the white ethnic group (there might be some unmet need associated with circulatory problems in the non-white ethnic groups).

The PCT budget variable is positive in all of the remaining seven programmes and this variable is significant in six of the seven. The proxy for other programme need (SYLLR all deaths) has the anticipated negative sign in five of the seven programmes and, where it is positive, it is never statistically significant.

The all service proxy for own programme need is positive and significant in three programmes. In the other four programmes, however, it has been replaced various other socio-economic indicators of need: in the trauma programme, for example, with the provision of unpaid care is associated with a reduction in NHS expenditure and, in the neonates programme, the proportion of residents in the white ethnic group is negatively associated with expenditure.

The diagnostic statistics reveal that, for all seven IV models, expenditure is endogenous and the instruments are valid. They also suggest that the instruments are relevant and there is no evidence that the instruments are weak. The Pesaran-Taylor test suggests that there is no evidence of model misspecification.

## B7.5 IV estimates of outcome and expenditure models: the first-stage equations

For the health outcome equation, IV estimation involves finding variables (instruments) that are good predictors of programme expenditure but which are appropriately excluded from the equation of interest (that is, from the outcome equation). The assumption is that the instruments impact upon the health outcome through their impact on expenditure only, and that they do not have a direct effect on the outcome. If, on the other hand, an instrument reflects unobserved factors that affect both expenditure and mortality directly, then the IV estimator becomes both biased and inconsistent. Such an instrument is said to be 'invalid' because it belongs in the equation of interest in its own right.

In our outcome model we typically employ two instruments (call these z1 and z2) for expenditure. IV estimation assumes that these instruments do not belong in the outcome equation. In other words, IV estimation assumes that the coefficients $\gamma 1$ and $\gamma 2$ in the outcome model

$$
\begin{equation*}
y=\alpha+\beta 1 x+\beta 2 n+\gamma 1 z 1+\gamma 2 z 2+\epsilon \tag{7.1}
\end{equation*}
$$

are identically zero (where y is mortality, x is expenditure, and n is a measure of the own programme need for health care and all variables relate to a particular programme of care). Such exclusion restrictions can be debatable and researchers who employ IV techniques often devote considerable effort towards convincing the reader that their assumed exclusion restrictions are a good approximation [35, 36]. These efforts usually take two forms: first, researchers often offer a strong theoretical economic argument why their instruments do not belong in the equation of interest; and, second, statistical tests for the validity of the exclusion restrictions (Sargan 2SLS, Hansen J-test GMM) are routinely reported as part of the results for any study that employs IV techniques.

It is difficult for us to identify clear theoretical reasons why our instruments (such as the proportion of lone pensioner households, the provision of unpaid care, and an index of multiple deprivation) do not belong in the equation of interest (that is, that they will not directly affect mortality). Of necessity, therefore, we must be guided by the available statistical tests for the validity of the exclusion restrictions. However, although our outcome models 'pass' the relevant statistical test, some commentators have argued that the Sargan/Hansen test may have weak power and may fail to reject the null hypothesis of instrument validity even when an exclusion restriction is not valid. As we shall see in section B9, this is likely to be the case when the induced biases in the estimates of $\beta 1$ (the coefficient on the endogenous variable) are the same across all instruments. The Hansen-Sargan J test statistic will be small when the null hypothesis of valid instruments is correct; but it will alsepe small if the biases induced in $\widehat{\beta 1}$ by invalid instruments all coincide (i.e., the instruments all identify the same wrong parameter)[37]. In other words, for the Hansen-Sargan J test to have low power the use of any subset of instruments should generate the same asymptotic bias in $\widehat{\beta 1}$.

Our approach, implemented below, is to identify theoretical reasons why our instruments might belong in the first-stage expenditure equation butnotin the second-stage outcome equation. Even if our arguments are thought unconvincing, a critic would also have to argue that any subset of our selected instruments will each induce the same bias in the coefficient on the endogenous variable. This is because it is only in these circumstances that the Hansen-Sargan test will be unable to reject the null hypothesis of instrument validity even when an exclusion restriction is not valid.

The first stage regressions associated with the IV outcome results in Table B7.4 can be found in Table BA. 3 in the annex. A briefsummary of the first-stage regressions is provided below.

## Cancer programme of care

The instrument set for the cancer programme of care (see column 1 in Table BA.3) includes the proportion of households that are lone pensioner households and the proportion of the population providing unpaid care. These instruments have intuitive appeal. The first stage regression of cancer expenditure on the instruments and the need for health care (as an exogenous regressor in the 2SLS model) reveals a positive and significant coefficient on lone pensioners and a negative but non-significant coefficient on the proportion of unpaid carers. The proportion of lone pensioners is likely to reflect an additional adjustment for health care need specific to an elderly and needy population. The omission of this variable from the second-stage regression is plausible as the dependent variable relates to mortality under 75 years of age and some of the lone pensioners will be aged over 75 years, and members of this group are, by definition, relatively healthy individuals. Unpaid care might act as a substitute for the provision of health care services and, in these circumstances, a negative relationship with expenditure is to be expected. There is no obvious relationship between the provision of unpaid care and mortality.

## Circulatory disease programme of care

The two instruments used for cancer were also employed to predict expenditure in the circulatory disease programme and they were augmented with the addition of the population weighted index of multiple
deprivation (IMD 2000). The relevance of the latter variable is theoretically plausible as circulatory disease is more related to disadvantage than is cancer. In addition, we also employed the proportion of residents in the white ethnic group as an additional instrument for expenditure but its coefficient is very small and it is not statistically significant.

Increased expenditure on circulatory disease in the first stage regression is associated with a greater proportion of pensioners living alone and a greater proportion of unpaid carers. The latter may reflect an increased awareness and compliance with medical intervention, particularly preventative measures, brought about by carers but this will not affect our outcome model if the impact of this additional support is largely on the mortality of those aged over 75 years. Expenditure on circulatory problems is also negatively associated with the IMD 2000. As the IMD incorporates an access to medical services domain, this negative association might reflect some unmet need which largely affects mortality in those aged over 75 years.

## Respiratory problems programme of care

The IMD 2000 is negatively associated with expenditure on respiratory problems. As the IMD incorporates an access to medical services domain, this negative association might reflect some unmet need which largely affects mortality in those aged over 75 years. The proportion of the population aged 16-74 that is permanently sick has a positive association with expenditure but might not affect mortality in the under 75 s if expenditure is largely directed towards managing chronic disease.

## Gastro-intestinal problems programme of care



Increased expenditure on gastro-intestinal problems in the first stage regression is positively associated with the proportion of residents providing unpaid care. This may reflect an increased awareness and compliance with medical intervention, particularly preventafive measures, brought about by carers but this will not affect our outcome model if the impact of this additional support is largely on the mortality of those aged over 75 years.

## Trauma, burns and injuries programme of care

Increased expenditure on trauma, burns and injuries in the first stage regression is positively associated with the proportion of pensioners living alone. This may reflect longer stays in hospital and an increased need for community care. However, the proportion of pensioners living alone will have little effect on our mortality measure if most of this expenditure is associated with patients aged over 75 years of age.

## Neonate programme of care

The percentage of those aged $16-74$ that are long-term unemployed and the proportion of households that are in social rented housing are both positively associated with expenditure on neonate care. These are both indicators of socio- economic deprivation and might be associated with the presence of larger families (i.e., more children per family). This would affect expenditure per head of population but not necessarily mortality per 1,000 live births. The negative coefficient on the proportion of those aged 16-74 with no qualifications might reflect the 'emigration' of young adults from those areas that are particularly deprived. This would reduce expenditure per head of population but would have no impact on the mortality measure.

The first stage regressions associated with the IV expenditure results in Table B7.5 can be found in Table BA. 4 in the annex.

## Cancer programme of care

The first-stage equation for the cancer expenditure model includes two instruments - lone pensioners, and unpaid carers -- that are excluded as regressors from the second stage of estimation. In this model the first stage regression of other programme need (as proxied here by the circulatory disease mortality rate) on the instrument set generates a negative coefficient on both instruments excluded from the second-stage regression. A greater proportion of unpaid carers might reflect an increased level of care (and perhaps increased compliance with care programmes and drug regimes) resulting in a decrease in other programme deaths. The availability of unpaid care in the community might not have a direct effect on cancer expenditure if such care supplements rather than substitutes for NHS funded care. Conditional
on need and the total PCT budget, the negative coefficient on the proportion of lone pensioners may be indicative of the presence of increased networks of social support. If this additional support reduces other programme mortality but does not substitute for NHS care, then the lone pensioner variable will not belong in the expenditure equation.

## Circulatory disease programme of care

In the circulatory disease expenditure model, the first stage regression of other programme need (as proxied here by cancer mortality rate) on the instrument set results in a negative coefficient on one instrument (lone pensioners) and a positive coefficient on the other (the IMD 2000). As noted above, the negative coefficient on the proportion of lone pensioners may be indicative of areas with increased networks of social support. If this additional support does not substitute for NHS care then the lone pensioner variable will not belong in the expenditure equation. It is plausible that the IMD 2000 should have a positive effect on other programme need but not belong in the expenditure equation if, for example, there is some unmet need in another (but not the circulatory disease) care programme.

## Respiratory problems programme of care

In the respiratory disease expenditure model, the first stage regression of other programme need (as proxied here by the all cause SYLL rate) on the instrument set results in a negative coefficient on one instrument (unpaid care) and a positive coefficient on another (i.e., on the IMD2000). A greater proportion of unpaid carers might reflect an increased level of care (and perhaps increased compliance with care programmes and drug regimes) resulting in a decrease in other programme deaths. The availability of unpaid care might not have a direct effect on own programme expenditure if such care does not substitute for NHS funded care. It is plausible that the IMD 2000 should have a positive effect on other programme need but not belong in the expenditure equation if, for example, there is some unmet need in another (but not the respiratory disease) care programme.

## Gastro-intestinal problems programme of care

In the gastro-intestinal problems expenditure moder, the first stage regression of other programme need (as proxied here by the all cause SYLL rate) on the instrument set (including need and total budget) results in a negative coefficient on one instrument (lone pensioners) and a positive coefficient on the other (IMD2000). As noted above, the negative coefficient on the proportion of lone pensioners may be indicative of areas with increased networks of social support. If this additional support does not substitute for NHS care then the lone pensioner variable will not belong in the expenditure equation. It is plausible that the IMD 2000 should have a positive effect on other programme need but not belong in the expenditure equation if, for example, there is some unmet need in another (but not the gastrointestinal) care programme.

## Neurological problems programme of care

The first-stage equation for the neurological expenditure model includes three instruments - lone pensioners, unpaiid carers and IMD2000 -- that are excluded as regressors from the second stage of estimation. Explanations for the signs on these variables have been outlined above when discussing the other first stage regressions.

Trauma and injuries programme of care
The first-stage equation for the trauma expenditure model includes two instruments - lone pensioners and the IMD2000 -- that are excluded as regressors from the second stage of estimation. Explanations for the signs on these variables have been outlined above when discussing the other first stage regressions.

GMS/PMS programme of care
The first-stage equation for the GMS/PMS expenditure model includes three instruments - households with no car, lone parents, and permanently sick -- that are excluded as regressors from the second stage of estimation. All three are plausibly positively associated with other programme need (as proxied here by the all cause SYLL rate) but do not occur as regressors in the second stage GMS/PMS expenditure model. The latter includes at least one measure of deprivation - the proportion of people aged 16-74
without any qualification - and the Hansen-Sargan test suggests that three excluded instruments offer no additional explanatory power for observed variations in GMS/PMS expenditure.

We appreciate that not everyone will be convinced by our arguments about the validity of our instruments and so in section B9 we undertake a sensitivity analysis that examines the impact of weakening the instrument exclusion restriction.

Table B7.4: table showing preferred outcome models using 2005/6 expenditure data and mortality for 2002/2003/2004

|  | (1) | (2) | (3) | (4) | (5) | (6) | (7) | (8) |  | (10) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | PBC 2 <br> cancer | PBC 10 <br> circulation | PBC 11 <br> respiratory | PBC 13 <br> gastro-intestinal | PBC 1 <br> infectious disease | PBC 7 <br> neurological | PBC 16 trauma | PBC 19 <br> neonates | PBC 17 <br> genito-urinary | PBC 4 endocrine |
|  | 2005/6 | 2005/6 | 2005/6 | 2005/6 | 2005/6 | 2005/6 | 2005/6 | 2005,6 | 2005/6 | 2005/6 |
|  | outcome model | outcome model | outcome model | outcome model | outcome model | outcome model | outcome model | Outcome model | outcome model | outcome model |
|  | instrument spend | instrument spend | instrument spend | instrument spend | spend exogenous | spend exogenous | instrument spend | instrument spend | spend exogenous | spend exogenous |
|  | unweighted | unweighted | unweighted | unweighted | unweighted | unweighted | unweighted | unweighted | unweighted | unweighted |
| VARIABLES | second stage | second stage | second stage | second stage | OLS | OLS | second stage | second stage | OLS | OLS |
|  |  |  |  |  |  |  | $\bigcirc$ |  |  |  |
| own programme spend $\mathrm{p} /$ head | -0.394*** | $-1.370 * * *$ | -1.574*** | -2.018*** | -0.152 | -0.182 | . 33254 | -0.237* | -0.034 | -0.244* |
|  | [0.100] | [0.156] | [0.483] | [0.364] | [0.117] | [0.143] | [0.469] | [0.127] | [0.220] | [0.129] |
| need per head | 0.905*** | $2.628 * * *$ | 4.076*** | 4.254*** |  | 1.157*** | 1.588*** |  |  |  |
|  | [0.083] | [0.163] | [0.562] | [0.412] |  | [0.252] | [0.445] |  |  |  |

lone pensioner households
[0.083] [0.163]
born outside EU
no car households
HIV need per head
unpaid carers
low birth weight births
one parents households
white ethic group

[^51]$-0.930^{*}$
opulation weighted IMD 2000
diabetes prevalence rate 2004/5
diabetes prevalence rate squared


Shea's partial R-squared
leibergen-Paap LM test statisi
Kleibergen-Paap p-value
Kleibergen-Paap F statistic
Pesaran-Taylor reset statistic
Pesaran-Taylor p-value
Ramsey reset F statistic
Probability > F
0.133
26.59
$1.68 \mathrm{e}-06$
16.94
0.0347
0.852
0.311

| 0.0376 | 0.173 |
| :--- | :--- |
| 20.56 | 34.83 |
| $3.44 \mathrm{e}-05$ | $1.32 \mathrm{e}-0$ |
| 10.29 | 23.32 |
| 0.0929 | 2.196 |
| 0.761 | 0.138 |

0.112
26.97
$1.39 \mathrm{e}-06$
17.76
17.76
0.756

Notes: (i) Robust standard errors in brackets, ${ }^{* * *} \mathrm{p}<0.01$, ${ }^{* *} \mathrm{p}<0.05$, ${ }^{*} \mathrm{p}<0.1$;
(ii) for the endogeneity test the null is that the specified endogenous regressors can actually be treated as exogenous;
(iii) the instrument validity test is based on the Hansen-Sargan test. The joint null hypothesis is that the instruments are valid instruments, i.e., uncorrelated with the error term, and that the excluded instruments are correctly excluded from the estimated equation.
(iv) Shea's partial R-squared is an indicator of the degree of instrument relevance (i.e., of the correlation between the instruments and the endogenous regressor). It is the value of R-squared from a regression of the endogenous variable on the excluded instruments.
(v) A statistical test of instrument relevance is provided by the Kleibergen-Paap LM test. The null hypothesis is that the instruments are not relevant
(vi) Weak identification arises when the excluded instruments are correlated with the endogenous regressor, but only weakly. Estimators can perform poorly when instruments are weak. The Kleibergen-Paap Wald F statistic provides a formal test of weak identification. The null hypothesis is that the instruments are weak.

Table B7.5: table showing preferred expenditure models using 2005/6 expenditure data and mortality for 2002/2003/2004



See notes to Table B7.4

## B7.6 Calculation of the cost of a life and life year

The preferred models identified in Tables B7.4 and B7.5 indicate the responsiveness of mortality to changes in expenditure, and of own programme expenditure to changes in budget, using expenditure data for 2005/6. Together with information about programme expenditure and mortality, the coefficients on the own programme expenditure and PCT budget variables listed in Tables B7.4 and B7.5 can be used to calculate the cost of an additional life year for the ten programmes for which outcome and expenditure models are available. ${ }^{21}$ For a relatively small budget change:
the cost of an additional life in a particular programme
$=$ the change in expenditure in that programme $/$ the change in mortality in that programme
$=($ annual spend $*$ expenditure elasticity) $/$ (annual mortality $*$ outcome elasticity)
and
the cost of an additional life year in a particular programme

$=$ the change in expenditure in that programme / the change in life years lost in that programme
$=$ (annual spend $*$ expenditure elasticity) / (annual life years lost * outcome elasticity).
Table B7.6 presents the necessary information to calculate the cost of an additional life (or life year) for each of these ten programmes. There is an assumed small ( $1 \%$ ) increase in the national budget and it is also assumed that this increase is applied to each PCT's budget. The total additional spend in each programme associated with this injection (column E) is determined bythe initial level of expenditure in the programme (column B) and the programme's expenditure elasticity (column D). And this additional spend, in conjunction with the outcome elasticity (column H ) and the number of deaths in the programme (column G), determine the number of lives saved that is associated with the additional expenditure. If we divide the change in programme expenditure (column E) by the change in the number of lives lost (column I) we obtain the cost per life gained (column K ).

Alternatively, we can apply the outcome elasticity (column $H$ ) to the annual number of life years lost in the programme (column G) to determine the number of life years saved that is associated with the additional expenditure. If we divide the change in programme expenditure (column E) by the change in the number of life years lost (column $\mathbf{N}$ ) we obtain the cost per life year gained (column O ). Note that none of these figures are QALY adjusted and that all costs are at current (2005/6) prices.

The cost per life year associated with the cancer programme is $£, 13,741$ and this is almost identical to that calculated using expenditure data for 2004/5 but with the same mortality data as that employed here [2]. Similarly, the cost per life year associated with the circulatory disease programme is $£ 8,328$ and this is also almost identical to that calculated using expenditure data for 2004/5 but with the same mortality data as that employed here [2].) The cost per life year for the respiratory programme ( $£ 20,601$ ) and for the gastro-intestinal programme $(£, 18,303)$ are a little larger than these figures but are still of the same order of magnitude. Taken together, the cost per life year for these 'big four' programmes is $£ 12,855 .{ }^{22}$

Table B7.6 also contains cost per life year estimates for the six other programmes for which a mortalitybased outcome indicator is available. These cost estimates are much larger than those for the big four programmes. This is to be expected as mortality is a less relevant outcome indicator for these PBCs than for the big four programmes. The cost per life year across all ten programmes for which a mortalitybased outcome indicator is available is $£ 21,256$.

[^52]Although we have an estimate of the cost per life year for ten programmes, it is unclear how we should adjust this estimate for the expenditure associated with the other 13 programmes. We attempted to estimate an outcome and expenditure model for expenditure and mortality in all 13 of these programmes combined. ${ }^{23}$ However, this was not successful with, for example, counter-intuitive signs on some variables. ${ }^{24}$

Instead, we decided to make some assumption about the cost per life year associated with the other 13 programmes. We examined two possibilities. First, we assumed that the other 13 programmes generate no mortality gain at all. This is clearly unrealistic but it does provide an upper bound for the cost per life year across all programmes of care. Table B7.7 is similar to Table B7.6 but it incorporates this zero gain assumption for the 13 other programmes. ${ }^{25}$ It shows that the cost per life year across all 23 programmes - assuming a zero mortality gain in the 13 programmes without a mortality based indicator - is $£, 56,799$.

Second, the zero mortality gain assumption is an extreme one but possibly relevant for the residual programme (PBC 23) -- where about two-thirds of the expenditure is attributable to primary care -- if we assume that any mortality gain associated with primary care expenditure is reflected in mortality rates associated with other, more disease specific, programmes (e.g., cancer, circulatory disease, etc). But if we assume a zero mortality gain in PBC 23, what assumption should we make about the mortality gain associated with the remaining 12 programmes?

One possibility is to assume that the cost per life (year) in the remaining 12 programmes is on average the same as that associated with the ten programmes for which a mortality-based outcome indicator is available. At first this may sound strange as we have already noted that mortality is not regularly associated with these programmes whereas it is a normal outcome for the ten programmes for which a mortality-based outcome indicator is available (and this is ofcourse why mortality data at PCT level is available for these ten PBCs). However, if we broaden our interpretation of health gain to include nonmortality effects (such as those on the quality of life), then this assumption - that the cost per life (year) in the remaining 12 programmes is on average the same as that associated with the ten programmes for which a mortality-based outcome indicator is ayailable - becomes far more plausible.

Thus Table B7.8 is similar to Table B7.7 but incorporates: (a) a zero gain assumption for the residual (including primary care) programme (PBC23); and (b) an average gain assumption for the remaining 12 programmes for which no mortality based outcome indicator is available. Table B7.8 shows that the cost per life year across all 23 programmes (see row 15) is $£ 24,200$. This is, of course, slightly greater than the cost of a life year for the ten programmes for which a mortality-based outcome indicator is available $(£ 21,256)$ because a small proportion of expenditure (that on primary care) is assumed to have no health benefit beyond that captured by the more disease specific programmes (e.g., in cancer, circulatory disease, etc).

The costs quoted inTables B7.6, B7.7, and B7.8 make no QALY adjustment but such an adjustment would add between $50 \%$ and $66 \%$ to the costs quoted [5].
${ }^{23}$ We are grateful to Steve Morris for this suggestion.
${ }^{24}$ Instead of estimating programme specific models we also tried estimating an outcome model using the all cause mortality rate and expenditure across all programmes combined but this was not successful (again, counter-intuitive signs were obtained on some variables). We also investigated the possibility of using an overall measure of health derived from the Health Survey for England. Apart from sample size issues at PCT level (4,645 adults in England were interviewed for the 2009 survey), such surveys by definition only provide information about the health status of the living population and reveal nothing about the level of mortality.
${ }^{25}$ The cost of a life year for those 13 programmes where there is no health gain is, of course, undefined.

Table B7.6: table showing cost of life and life year estimates for 2005/6 for the ten programmes for which we have outcome and expenditure elasticities


Table B7.7: table showing cost of life and life year estimates for 2005/6 for all programmes (assumes that 13 PBCs offer no health gain)


Table B7.8: table showing cost of life and life year estimates for 2005/6 for all programmes (assumes GMS/PMS provides no gain, other PBCs provide average gain)


## B7.7 Summary and conclusion

In this section we have extended the results reported by Martin, Rice and Smith[5] by obtaining plausible outcome and expenditure models for all ten programmes of care with a mortality based outcome indicator. In addition, we have, for the first time, calculated the cost of a life year across the big four programmes combined $(£, 12,855)$ and across all ten programmes $(£ 21,256)$. Moreover, with the aid of an assumption about the productivity (health gain) of programmes without a meaningful mortality based outcome indicator, we have extended our individual programme estimates to incorporate expenditure across all programmes of care.

If we assume that the other 13 programmes without a mortality-based outcome indicator generate no health gain then the cost of an additional life year across all expenditure for 2005/6 is $£ 56,799$.

Alternatively, if we assume that any health care gain associated with primary care expenditure is reflected in mortality rates associated with other, more disease specific, and that the health gain associated with the remaining 12 programmes is, on average, the same as that recorded by the PBCs with a mortality based indicator, then the cost per life year across all expenditure for $2005 / 6$ is $£ 24,200$.

This concludes our analysis of the 2005/6 programme budgeting data. In the nextsection we apply our model to the 2006/7 programme budgeting data.

## B8. Analysis of programme budgeting expenditure for 2006/7

## B8.1 Construction of an alternative measure of need

The analysis of the 2005/6 programme budgeting data employed a measure of the need for health care that incorporated the AREA resource allocation formula for acute services. As was described in section B6.2, we attempted to construct a better measure of need using a recently developed person-based approach [34]. However, we were unable to construct a viable alternative PBRA based measure of need for use with the PB data for 2005/ 6 because the PBRA formula only relates to acute services yet the PB data incorporates elements for acute, maternity, mental health, prescribing and primary care, and we were unable to separate these components parts.

The construction of an alternative measure of need is, however, possible for use with the 2006/7 PB data. Spend and mortality data are available for the new (152) PCTs, and the Department of Health's resource allocation exposition book for 2009/10 (which employs the CARAN model) provides separate measures of need for acute, maternity, mental health, prescribing and GMS/PMS services. We can therefore replace the (CARAN-based) measure of acute need for the 2009/10 allocation with our own PBRA based measure of acute need (albeit for 2006/7) to calculate an alternative to the AREA based measure of need across all health care services.

The PBRA model was applied to all patients on Practice lists as at 1 April 2006 to generate a PCT level measure of acute need (see section B7.1 for a description of this approach as applied to patients on Practice lists as at 1 April 2005). The resulting PBRA measure of acute need can be compared with the CARAN based measure of acute need as reported in the Department of Health's resource allocation exposition book for 2009/10. The correlation coefficient for these two measures is 0.8514 and descriptive statistics for the two measures are shown below in Table B8.1.

Table B8.1: table showing summary statistics for the CARAN and PBRA based measures of acute need

| Variable | Number <br> of PCTs | Mean | Std. Dev. | Min | Max |
| :--- | :--- | :--- | :--- | :--- | :--- |
| CARAN_acute need | 152 | 1.0033 | 0.1113 | 0.7659 |  |
| PBRA_acute need | 152 | 1.0037 | 0.1218 | 0.7606 | 1.3420 |

The all service measure of need (which is a weighted average of the acute, maternity, mental health, prescribing and GMS/PMS measures) as reported in the Department of Health's resource allocation exposition book for 2009/10 can be re-calculated by replacing the CARAN-based acute measure with the PBRA-based acute measure of need. The correlation coefficient for these two all service measures of need is 0.9714 and Figure B8.1 shows a scatter plot of these two measures. Descriptive statistics for these two all service measures of need along with the (AREA-based) PB measure of need are shown below in Table B8.2.


Figure B8.1: graph showing scatter plot of all service measures of need: incorporating CARAN or PBRA based measures of acute need

Table B8.2: table showing all service measures of need: incorporating CARAN, PBRA or AREA based measures of acute need

| Variable | Obs | Mean | Std. Dev. | Min | Max |
| :--- | :--- | :--- | :--- | :--- | :--- |
| PCTneed_CARAN | 152 | 1.0240 | 0.1339 | 0.7311 | 1.3479 |
| PCTneed_PBRA | 152 | 1.0242 | 0.1395 | 0.7287 | 1.3769 |
| PCTneed_AREA | 152 | 1.0293 | 0.1380 | 0.7165 | 1.4006 |

The correlation coefficients for the three measures are shown in Table B8.3.
Table B8.3: table showing correlation coefficients for alternative measures of all service need

| Variable | PCTneed_AREA | PCTneed_PBRA | PCTneed_CARAN |
| :--- | :--- | :--- | :--- |
| PCTneed_AREA | 1 |  |  |
| PCTneed_PBRA | 0.9583 | 1 |  |
| PCTneed_CARAN | 0.9839 | 0.9714 | 1 |

B8.2 Estimation issues associated with the use of 2006/7 expenditure data
As well as having to select a preferred measure of need from the three available, the estimation of our model using PB data for 2006/7 requires the resolution of several other issues.

Estimation issue 1: 'net spend' or 'own population' spend?
The Department of Health reports two sets of PB spend data: the first is on a 'net spend' basis and the second is on an 'own population' basis. The 'own population' data starts with the 'net spend' figure, adds any expenditure funded from non-NHS sources, and adjusts for expenditure made under PCT lead/host commissioning arrangements. These adjustments are usually very small. For 2005/6 we used the net spend data (because only net spend data was produced in the first year and we were hoping to build a
panel) but given the now regular production of own population data this would seem to be the more appropriate data set to use as, for example, it includes all expenditure irrespective of its funding source.

## Estimation issue 2: to weight or not to weight?

OLS and IV estimation implicitly gives the same weight to each PCT when estimating our expenditure and outcome models. With the re-organisation of PCTs in October 2006, the number of such organisations was reduced from 303 to 152. However, far from making them more similar in terms of size (as measured by their population), this re-organisation actually increased the disparity in size between the largest and the smallest PCTs, with the largest PCT now being 14 times the size of the smallest. Unless we explicitly weight each observation (PCT) by its size, we will be giving the same weight (influence) to PCTs that are much smaller than other PCTs.

## Estimation issue 3: which MFF?

This study builds on previous work using PB data. Martin, Rice and Smith[3] report the results of the estimation of our model using PB data for 2006/7. One essential step in this estimation is the removal of the impact of unavoidable variations in local costs from the reported measure of the 'unified weighted' population. At the time of the earlier study the authors only had access to an MFF based on HCHS for the new 152 PCTs. Now, however, a more broadly-based MFF is available, that is one based on a weighted average of MFFs for HCHS, prescribing, and GMS/PMS. Should we use an MFF for HCHS only, or one that incorporates HCHS and prescribing, or one that incorporates HCHS , prescribing and GMS/PMS?

Estimation issue 4: SMRs or SYLLRs, and which proxy for the other programme need variable?
Previous studies have reported results using both SMRs and SYILRs but the sheer number of models being estimated requires that we focus on one measure only $\bigodot$ Various proxies for other programme need have been employed in previous studies (see section B6.3 for further discussion). In this sub-section we persevere with this variety but consistency demands that we focus in on a preferred proxy for other programme need.

This study builds on previous work using PB data. Martin, Rice and Smith[3] report the results of the estimation of our model using PB data for $2006 / 7$. With several alternative measures of need and MFF available, we undertook a preliminary empirical analysis of the $2006 / 7 \mathrm{~PB}$ data using the outcome and expenditure models for the big four programmes as reported in Martin, Rice and Smith[3] as our starting point. These models incorporated the AREA-based measure of need and an MFF based on HCHS only.

We first re-estimated the qutcome and expenditure models by replacing the AREA based measure of need with one incorporating the PBRA formula. Then we re-estimated these models again with a measure of need incorporating the CARAN model. The results suggest that: (a) for the outcome models, the use of the PBRA measure of need generates a smaller coefficient on expenditure than does the AREA measure of need; and (b) that for the spend models, the use of the PBRA measure of need generates a larger coefficient on PCT budget than does the AREA measure of need. For both the outcome and expendituremodels, the use of the CARAN measure of need generates outcome and expenditure elasticities that lie between those generated by the AREA and PBRA measures.

Next, the results reported by Martin, Rice and Smith[3] employ an MFF based on HCHS only to remove unavoidable variations in local costs from the reported measure of the (unified weighted) need for health care services. This was the only MFF available for the new PCTs at the time of that study. Now, however, a more broadly-based MFF is available (that is, one based on a weighted average of MFFs for HCHS, prescribing, and GMS/PMS).

To examine the consequences of using the CARAN MFF (i.e., a weighted average of the HCHS, prescribing, and GMS/PMS MFFs), this MFF was used to calculate the implied level of need given the unified weighted populations for 2006/7 which are reported alongside the PB spend data by the

Department of Health. ${ }^{26}$ We found that the use of an extended set of MFFs can sometimes affect the coefficient on the variable of interest.

Models were also estimated using a weighted average of the CARAN MFFs for HCHS and prescribing only. The latter results were very similar to those using all three of the CARAN MFFs (i.e., a weighted average of the HCHS, prescribing, and GMS/PMS MFFs).

We also tried re-estimating the outcome and expenditure models from Martin, Rice and Smith [3] using the 'own population' expenditure data rather than the 'net spend' data but this adjustment had very little effect on the results. In addition, the impact of 'weighting' each observation by PCT size was usually rather modest.

Because of the sheer number of variations possible, we decided to estimate 13 particular variants of our model and details of these variants are summarised in Table B8.4. These variants were estimated for each of the big four programmes using both the outcome and expenditure equations. The results are presented in Tables B8.5 to B8.12.

Table B8.4: table showing variants of the outcome and expenditure models estimated using 2006/7 spend data


Note: UWP=unified weighted population.

[^53]Table B8.5: table showing cancer spend models with various indicators of MFF and need

|  | (1) | (2) | (3) | (4) | (5) | (6) | (7) | (8) | (9) | (10) | (11) | (12) | (13) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | PBC 2 cancer | PBC 2 <br> cancer | PBC 2 cancer | PBC 2 cancer | PBC 2 <br> cancer | PBC 2 cancer | PBC 2 cancer | PBC 2 <br> cancer | $\text { PBC } 2$ cancer | PBC 2 <br> cancer | $\mathrm{PBC} 2$ | PBC 2 | $\text { PBC } 2$ |
|  | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 2006/7 | $2006 / 7$ | 2006/7 | 2006/7 | 2006/7 |
|  | spend model | spend model | spend model | spend model | spend model | spend model | spend model | spend model | spend model | spend model | spend model | spend model | spend model |
|  | uses SMR | uses SMR | uses SMR | uses SYLLR | uses SYLLR | uses SYLLR | uses SMR | uses SMR | uses SMR | uses SMR | uses SMR | uses SMR | uses SMR |
|  | no weighting | no weighting | no weighting | no weighting | no weighting | no weighting | weighted | no weighting | weighted | no weighting | weighted | no weighting | weighted |
|  | second stage | second stage | second stage | second stage | second stage | second stage | second stage | second stage | second stage | second stage | second stage | second stage | second stage |
|  | iAREA need 1 HCHS MFF | PBRA need HCHS MFF | CARAN need HCHS MFF | iAREA need 1 HCHS MFF | PBRA need HCHS MFF | CARAN need HCHS MFF | iAREA need 1 HCHS MFF | iAREA need 2 CARAN 3 MFFs | iAREA need 2 CARANO $\mathrm{MEFP}_{s}$ ) | iAREA need 3 CARAN 2 MFFs | iAREA need 3 <br> CARAN 2 <br> MFFs | CARAN need CARAN 3 MFFs | CARAN need CARAN 3 MFFs |
|  | HCHS MFF | HCHS MFF | HCHS MFF | HCHS MFF | HCHS MFF | HCHS MFF | HCHS MFF |  | MEFs | MFFs | MFFs | MFFs | MFFs |
| PCT budget per head | 0.353 | 0.681*** | 0.572** | 0.388 | 0.752*** | 0.618** | 0.326 | 0.250 | 1.245 | 0.246 | 0.241 | 0.552** | 0.544* |
|  | [0.273] | [0.235] | [0.247] | [0.272] | [0.238] | [0.247] | [0.362] | [0.284] | [0.357] | [0.284] | [0.357] | [0.239] | [0.308] |
| needAREA1 | 1.513*** |  |  | 1.557*** |  |  | 1.351*** | (1) |  |  |  |  |  |
|  | [0.288] |  |  | [0.284] |  |  | [0.367] | (1) |  |  |  |  |  |
| other programme need 1 | -0.654*** | -0.771*** | $-0.733 * * *$ |  |  |  | -0.604*** | . 749 | -0.661*** | -0.749*** | $-0.661 * * *$ | -0.680*** | -0.616*** |
|  | [0.124] | [0.125] | [0.124] |  |  |  | [0.131] | [0.152] | [0.154] | [0.151] | [0.154] | [0.126] | [0.139] |
| needPBRA |  | 1.320*** |  |  | 1.352*** |  |  |  |  |  |  |  |  |
|  |  | [0.261] |  |  | [0.265] |  | $\cdots$ |  |  |  |  |  |  |
| needCARAN |  |  | 1.431*** |  |  | 1.477*** |  |  |  |  |  | 1.347*** | 1.160*** |
|  |  |  | [0.287] |  |  | [0.289] |  |  |  |  |  | [0.246] | [0.294] |
| other programme need 2 |  |  |  | -0.649*** | -0.773*** | -0.728*** |  |  |  |  |  |  |  |
|  |  |  |  |  | .124] |  |  |  |  |  |  |  |  |
| needAREA2 |  |  |  |  |  | - |  | $\begin{aligned} & 1.778^{* * *} \\ & {[0.329]} \end{aligned}$ | $\begin{aligned} & 1.554 * * * \\ & {[0.389]} \end{aligned}$ |  |  |  |  |
| needAREA3 |  |  |  |  |  |  |  |  |  | 1.765*** | 1.545*** |  |  |
|  |  |  |  |  |  |  |  |  |  | [0.328] | [0.388] |  |  |
| Constant | 0.271 | 0.707 | 0.569 | 0.356 | 0.835 | 0,665 | 0.067 | 5.901*** | 5.559** | 5.932*** | 5.582** | 3.430** | 3.220 |
|  | [0.544] | [0.551] | [0.554] | [0.542] | [0.567] | [0.554] | [0.572] | [2.062] | [2.550] | [2.063] | [2.550] | [1.645] | [2.087] |
| Observations | 152 | 152 | 152 | 152 | 52 | 152 | 152 | 152 | 152 | 152 | 152 | 152 | 152 |
| Endogeneity test statistic | 13.112 | 18.683 | 18.420 | 13.313 | 8.736 | 18.716 | 11.940 | 13.098 | 11.708 | 13.017 | 11.504 | 16.985 | 13.460 |
| Endogeneity p-value | 0.000293 | $1.54 \mathrm{e}-05$ | $1.77 \mathrm{e}-05$ | 0.000264 | $1.50 \mathrm{e}-05$ | 1.52e-05 | 0.000549 | 0.000296 | 0.000622 | 0.000309 | 0.000695 | 3.77e-05 | 0.000244 |
| Hansen-Sargan test statistic | 0.870 | 1.139 | 0.560 | 0.504 | 0.748 | 0.281 | 1.089 | 1.730 | 1.875 | 1.711 | 1.857 | 0.381 | 0.321 |
| Hansen-Sargan p-value | 0.351 | 0.286 | 0.454 | 0.478 | 0.387 | 0.596 | 0.297 | 0.188 | 0.171 | 0.191 | 0.173 | 0.537 | 0.571 |
| Shea's partial R-squared | 0.607 | 0.526 | 0.586 | 0.570 | 0.482 | 0.548 | 0.612 | 0.511 | 0.537 | 0.510 | 0.536 | 0.572 | 0.583 |
| Kleibergen-Paap LM test statistic | 40.55 | 38.73 | 40.45 | 40. 49 | 38.17 | 41.14 | 38.91 | 38.41 | 37.13 | 38.38 | 37.11 | 43.82 | 41.36 |
| Kleibergen-Paap p-value | 1.57e-09 | $3.88 \mathrm{e}-09$ | 1.64e-09 | 1.61e-09 | 5.14e-09 | 1.16e-09 | $3.55 \mathrm{e}-09$ | 4.56e-09 | 8.66e-09 | $4.63 \mathrm{e}-09$ | $8.75 \mathrm{e}-09$ | 3.06e-10 | 1.05e-09 |
| Kleibergen-Paap F statistic | 73.17 | 63.72 | 68.50 ) | 67.14 | 51.14 | 60.78 | 72.14 | 57.74 | 61.44 | 58.09 | 61.58 | 68.29 | 66.96 |
| Pesaran-Taylor reset statistic | 0.233 | 0.299 | 0.027 | 0.198 | 0.211 | 0.00518 | 0.000324 | 0.00529 | 0.0391 | 0.0345 | 0.00971 | $9.41 \mathrm{e}-07$ | 0.0158 |
| Pesaran-Taylor p-value | 0.629 | 0.585 | 0.869 | 0.656 | 0.646 | 0.943 | 0.986 | 0.942 | 0.843 | 0.853 | 0.922 | 0.999 | 0.900 |

[^54](b) iAREA need 2=AREA unified weighted population/HCHS, prescribing \& GMS MFFs
(c) iAREA need $3=$ AREA unified weighted population/HCHS \& prescribing MFFs
(d) other programme need $1=$ circulatory disease SMR
(e) other programme need $2=$ circulatory disease SYLLR
(f) robust standard errors in brackets, *** $\mathrm{p}<0.01, * * \mathrm{p}<0.05, * \mathrm{p}<0.1$

Table B8.6: table showing circulatory disease spend models with various indicators of MFF and need

|  | (1) | (2) | (3) | (4) | (5) | (6) | (7) | (8) | (9) | (10) | (1) | (12) | (13) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | PBC 10 <br> circulation | PBC 10 <br> circulation | PBC 10 circulation | PBC 10 <br> circulation | PBC 10 <br> circulation | PBC 10 <br> circulation | PBC 10 <br> circulation | PBC 10 <br> circulation | PBC 10 <br> circulation | PBC 19 circulation | PBC 10 circulation | PBC 10 <br> circulation | PBC 10 <br> circulation |
|  | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 200677 | 2006/7 | 2006/7 | 2006/7 |
|  | spend model | spend model | spend model | spend model | spend model | spend model | spend model | spend model | spend model | spend model | spend model | spend model | spend model |
|  | uses SMR | uses SMR | uses SMR | uses SYLLR | uses SYLLR | uses SYLLR | uses SMR | uses SMR | uses SMR | usess SMR | uses SMR | uses SMR | uses SMR |
|  | no weighting | no weighting | no weighting | no weighting | no weighting | no weighting | weighted | no weighting | weighted | no weighting | weighted | no weighting | weighted |
|  | second stage | second stage | second stage | second stage | second stage | second stage | second stage | second stage | second stage | second stage | second stage | second stage | second stage |
|  | iAREA need1 | PBRA need | CARAN need | iAREA need1 | PBRA need | CARAN need | iAREA need1 | iAREA need2 CARAN 3 | iAREA need2 <br> CARAN3 | iAREA need3 CARAN 2 | iAREA need 3 CARAN 2 | CARAN need CARAN 3 | CARAN need CARAN 3 |
|  | HCHS MFF | HCHS MFF | HCHS MFF | HCHS MFF | HCHS MFF | HCHS MFF | HCHS MFF | MFFs | MFFs | MFFs | MFFs | MFFs | MFFs |
| other programme need 1 | -0.766** | -0.736** | -0.811** |  |  |  | -0.939*** | -0.776** | -0.927*** | ${ }^{-0.781 * *}$ | $-0.935 * * *$ | -0.831** | -1.112*** |
|  | [0.298] | [0.299] | [0.370] |  |  |  | [0.305] | ${ }^{[0.315]}$ ] | [0.324] | [0.316] | [0.326] | [0.367] | [0.392] |
| PCT budget per head | 0.861*** | 1.162*** | 1.035*** | 0.836*** | 1.191*** | 0.998*** | 0.719*** | 0.832** | 0.661** | 0.829*** | 0.657** | 0.983*** | 0.914*** |
|  | [0.240] | [0.218] | [0.219] | [0.229] | [0.220] | [0.210] | [0.259] | (00.242]) | [0.264] | [0.242] | [0.264] | [0.213] | [0.231] |
| needAREA1 | 0.624* |  |  | 0.732* |  |  | 0.967** |  |  |  |  |  |  |
|  | [0.355] |  |  | [0.389] |  |  | [0.378] |  |  |  |  |  |  |
| white ethnic group | 0.215*** | 0.187** | 0.207** | 0.232*** | 0.199** | 0.225** | $0.278 * * *$ | 0.219** | 0.284*** | 0.219** | 0.284*** | 0.209** | 0.286*** |
|  | [0.079] | [0.080] | [0.086] | [0.083] | [0.083] | [0.095] | [0.084] | [0.085] | [0.091] | [0.085] | [0.091] | [0.086] | [0.098] |
| provision of unpaid care | 0.457** | 0.554*** | 0.488** | 0.437** | 0.549*** | 0.466* | 0.239 | 0.527*** | 0.336 | 0.528*** | 0.335 | 0.477** | 0.200 |
|  | [0.205] | [0.186] | [0.227] | [0.212] | [0.183] | [0.247] | [0.227] | [0.190] | [0.210] | [0.190] | [0.211] | [0.227] | [0.268] |
| needPBRA |  | 0.250 |  |  | 0.295 | () |  |  |  |  |  |  |  |
|  |  | [0.275] |  |  | [0.280] | 1 |  |  |  |  |  |  |  |
| needCARAN |  |  | 0.480 |  |  | .610) |  |  |  |  |  | 0.546 | 0.925** |
|  |  |  | [0.401] |  |  |  |  |  |  |  |  | [0.369] | [0.399] |
| other programme need 2 |  |  |  | $-0.904 * *$ | $-0.871^{* *}$ | -0.956* |  |  |  |  |  |  |  |
| needAREA2 |  |  |  |  | - |  |  | 0.655* | 1.020** |  |  |  |  |
|  |  |  |  |  | , |  |  | [0.383] | [0.424] |  |  |  |  |
| needAREA3 |  |  |  |  | - |  |  |  |  | 0.652* | 1.018** |  |  |
|  |  |  |  |  |  |  |  |  |  | [0.381] | [0.422] |  |  |
| Constant | 2.380** | 2.377* | 2.621* | 3.246** | 3.244** | 3.533* | 2.744** | 3.763* | 5.275** | 3.803* | 5.337** | 2.827 | 4.028** |
|  | [1.212] | [1.251] | [1.469] | [1.572] | [1.553] | [2.088] | [1.222] | [2.133] | [2.306] | [2.143] | [2.319] | [1.863] | [1.995] |
| Observations | 152 | 152 | 152 | 152 | 152 | 152 | 152 | 152 | 152 | 152 | 152 | 152 | 152 |
| Endogeneity test statistic | 8.506 | 10.727 | 8.136 | 6.475 | 8.793 | 5.743 | 9.019 | 8.939 | 8.654 | 9.036 | 8.745 | 8.315 | 9.729 |
| Endogeneity p-value | 0.00354 | 0.00106 | 0.00434 | -0.0109 | 0.00302 | 0.0166 | 0.00267 | 0.00279 | 0.00326 | 0.00265 | 0.00310 | 0.00393 | 0.00181 |
| Hansen-Sargan test statistic | 2.454 | 0.640 | 1.841 | 2.364 | 0.423 | 2.166 | 2.993 | 1.770 | 2.225 | 1.792 | 2.237 | 1.777 | 2.030 |
| Hansen-Sargan p-value | 0.117 | 0.424 | 0.175 | 0.124 | 0.515 | 0.141 | 0.0836 | 0.183 | 0.136 | 0.181 | 0.135 | 0.183 | 0.154 |
| Shea's partial R-squared | 0.235 | 0.205 | 0.184 | 0.207 | 0.184 | 0.148 | 0.238 | 0.225 | 0.230 | 0.224 | 0.228 | 0.183 | 0.183 |
| Kleibergen-Paap LM test statistic | 22.99 | 24.15 | 20.59 | 21.32 | 25.01 | 17.46 | 26.79 | 23.47 | 27.59 | 23.63 | 27.59 | 20.91 | 24.11 |
| Kleibergen-Paap p-value | $1.02 \mathrm{e}-05$ | 5.70e-06 | 3.39\%-05 | 2.35--05 | 3.70e-06 | 0.000161 | 1.52e-06 | 8.02e-06 | 1.02e-06 | 7.41e-06 | 1.02e-06 | 2.88e-05 | 5.80e-06 |
| Kleibergen-Paap F statistic | 23.14 | 22.53 |  | 17.28 | 20.44 | 12.62 | 22.37 | 21.47 | 22.07 | 21.27 | 21.63 | 19.10 | 18.93 |
| Pesaran-Taylor reset statistic | 0.00329 | 0.156 | 0.102 | 0.0384 | 0.333 | 0.288 | 0.0270 | 0.123 | 0.152 | 0.189 | 0.235 | 0.0165 | 0.190 |
| Pesaran-Taylor p-value | 0.954 | 0.693 ) | 0.750 | 0.845 | 0.564 | 0.592 | 0.869 | 0.726 | 0.696 | 0.664 | 0.628 | 0.898 | 0.663 |

Notes: (a) iAREA need 1=AREA unified weighted population/HCHS MFF
(b) iAREA need $2=$ AREA unified weighted population/HCHS, prescribing \& GMS MFFs
(c) iAREA need $3=$ AREA unified weighted population/HCHS \& prescribing MFFs
(d) other programme need $1=$ cancer SMR
(e) robust standard errors in brackets, *** $\mathrm{p}<0.01,{ }^{* *} \mathrm{p}<0.05,{ }^{*} \mathrm{p}<0.1$

Table B8.7: table showing respiratory problems spend models with various indicators of MFF and need

|  | (1) | (2) | (3) | (4) | (5) | (6) | (7) | (8) | (9) | (10) | (11) | (12) | (13) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | PBC 11 respiratory | PBC 11 respiratory | PBC 11 <br> respiratory | PBC 11 respiratory | PBC 11 respiratory | PBC 11 respiratory | PBC 11 respiratory | PBC 11 <br> respiratory | PBC 11 <br> respiratory | PBC 11 <br> respiratory | PBC 11 respiratory | PBC 11 respiratory | PBC 11 respiratory |
|  | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 2006/7 | 2006/7 |
|  | spend model | spend model | spend model | spend model | spend model | spend model | spend model | spend model | spend model | spend model | spend model | spend model | spend model |
|  | uses SMR | uses SMR | uses SMR | uses SYLLR | uses SYLLR | uses SYLLR | uses SMR | uses SMR | uses SMR | uses SMR | uses SMR | uses SMR | uses SMR |
|  | no weighting | no weighting | no weighting | no weighting | no weighting | no weighting | weighted | no weighting | weighted | now weighting | weighted | no weighting | weighted |
|  | second stage | second stage | second stage | second stage | second stage | second stage | second stage | second stage | second stage | second stage | second stage | second stage | second stage |
| PCT budget per head | 0.781** | 0.992*** | 0.957*** | 1.045*** | 1.315*** | 1.204*** | 0.808** | 0.592** | 0.591* | 0.588** | 0.585* | 0.865*** | 0.958*** |
|  | [0.318] | [0.330] | [0.363] | [0.370] | [0.409] | [0.432] | [0.334] | [0.282] | 10.3 | [0.283] | [0.310] | [0.287] | [0.329] |
| needAREA1 | 1.714*** |  |  | 1.741*** |  |  | 1.813*** |  |  |  |  |  |  |
|  | [0.597] |  |  | [0.497] |  |  | [0.563] | (1) |  |  |  |  |  |
| lone pensioner households | -0.497 | -0.243 | -0.483 | -0.419* | -0.240 | -0.380 | -0.595* | (078) | -0.304 | -0.089 | -0.320 | -0.447 | -0.556 |
|  | [0.346] | [0.243] | [0.356] | [0.252] | [0.210] | [0.271] | [0.346] | [0.285] | [0.378] | [0.286] | [0.380] | [0.337] | [0.344] |
| other programme need 1 | -0.803** | -0.602** | -0.890** |  |  |  | -0.866** | - 0.391 | -0.664 | -0.407 | -0.687 | -0.834* | -0.931** |
|  | [0.397] | [0.294] | [0.439] |  |  |  | [0.392] | [0.364] | [0.478] | [0.364] | [0.481] | [0.428] | [0.437] |
| needPBRA |  | 1.176*** |  |  | 1.226*** |  | $\square$ |  |  |  |  |  |  |
|  |  | [0.391] |  |  | [0.366] |  |  |  |  |  |  |  |  |
| needCARAN |  |  | 1.686*** |  |  | 1.720*** |  |  |  |  |  | 1.680*** | 1.782*** |
|  |  |  | [0.627] |  |  | [0.536] |  |  |  |  |  | [0.609] | [0.561] |
| other programme need 2 |  |  |  | $\begin{aligned} & -1.109 * * \\ & {[0.455]} \end{aligned}$ | $\begin{aligned} & -0.955^{* *} \\ & {[0.406]} \end{aligned}$ | -1.102** |  |  |  |  |  |  |  |
| needAREA2 |  |  |  |  |  |  |  | 1.312** | 1.770** |  |  |  |  |
|  |  |  |  |  |  |  |  | [0.667] | [0.803] |  |  |  |  |
| needAREA3 |  |  |  |  |  |  |  |  |  | 1.325** | 1.788** |  |  |
|  |  |  |  |  | ) |  |  |  |  | [0.661] | [0.800] |  |  |
| Constant | -0.143 | -0.649 | 0.258 | 2.954 | 2.293 |  | -0.046 | 1.605 | 2.457 | 1.686 | 2.578 | 1.022 | 0.595 |
|  | [1.250] | [0.961] | [1.415] | [2.335] | [2.094] | [2.856] | [1.220] | [2.372] | [2.547] | [2.389] | [2.574] | [1.980] | [2.082] |
| Observations | 152 | 152 | 152 | 152 |  | 152 | 152 | 152 | 152 | 152 | 152 | 152 | 152 |
| Endogeneity test statistic | 7.821 | 8.431 | 9.089 | 9.15 | 10.215 | 9.242 | 6.984 | 4.679 | 4.326 | 4.853 | 4.475 | 9.016 | 6.863 |
| Endogeneity p-value | 0.00516 | 0.00369 | 0.00257 | 0.00248 | 0.00139 | 0.00236 | 0.00822 | 0.0305 | 0.0375 | 0.0276 | 0.0344 | 0.00268 | 0.00880 |
| Hansen-Sargan test statistic | 1.655 | 3.108 | 0.983 | 0.214 | 1.502 | 0.0135 | 0.615 | 4.704 | 2.922 | 4.621 | 2.855 | 0.866 | 0.156 |
| Hansen-Sargan p-value | 0.198 | 0.0779 | 0.321 | 0.644 | 0.220 | 0.908 | 0.433 | 0.0301 | 0.0874 | 0.0316 | 0.0911 | 0.352 | 0.693 |
| Shea's partial R-squared | 0.164 | 0.211 | 0.149 | 0.183 | 0.203 | 0.172 | 0.167 | 0.161 | 0.131 | 0.161 | 0.131 | 0.142 | 0.146 |
| Kleibergen-Paap LM test statistic | 18.58 | 21.08 | 18.17 | 22.04 | 17.51 | 18.27 | 18.78 | 20.00 | 13.94 | 20.14 | 13.90 | 17.12 | 16.62 |
| Kleibergen-Paap p-value | 9.22e-05 | 2.64e-05 | 0.000113 | 1.64e-05 | 0.000158 | 0.000108 | $8.34 \mathrm{e}-05$ | $4.55 \mathrm{e}-05$ | 0.000940 | 4.23e-05 | 0.000960 | 0.000192 | 0.000246 |
| Kleibergen-Paap F statistic | 8.163 | 14.31 | 72 | 8.729 | 8.513 | 6.885 | 8.383 | 13.56 | 7.241 | 13.77 | 7.297 | 7.776 | 7.220 |
| Pesaran-Taylor reset statistic | 0.238 | 0.0135 | 0.141 | 0.0704 | 0.0139 | 0.0164 | 1.083 | 2.231 | 2.206 | 2.311 | 2.283 | 3.699 | 4.984 |
| Pesaran-Taylor $p$-value | 0.625 | $0.907 \sim$ | 0.707 | 0.791 | 0.906 | 0.898 | 0.298 | 0.135 | 0.138 | 0.128 | 0.131 | 0.0545 | 0.0256 |

[^55](b) iAREA need $2=$ AREA unified weighted population/HCHS, prescribing \& GMS MFFs
(c) iAREA need $3=$ AREA unified weighted population/HCHS \& prescribing MFFs
(d) other programme need $1=$ SMR for all causes of death amenableto health care (see Martin, Rice and Smith 2012)
(e) other programme need $2=$ SYLLR for all causes of death
(f) robust standard errors in brackets, *** $\mathrm{p}<0.01, * * \mathrm{p}<0.05, * \mathrm{p}<0.1$

Table B8.8: table showing gastro-intestinal problems spend models with various indicators of MFF and need


Notes: (a) iAREA need $1=$ AREA unified weighted population/HCHS MF
(b) iAREA need $2=$ AREA unified weighted population/HCHS, prescribing \& GMS MFFs
(c) iAREA need 3=AREA unified weighted population/HCHS \& prescribing MFFs
(d) other programme need $1=$ SMR for all causes of death ammenable to health care (see Martin, Rice and Smith 2012)
(e) other programme need $2=$ SYLLR for all causes of death
(f) robust standard errors in brackets, ${ }^{* * *} \mathrm{p}<0.01, * * \mathrm{p}<0.05, *^{*} \mathrm{p}<0.1$

# Table B8.9: table showing cancer outcome models with various indicators of MFF and need 



Notes: (a) iAREA need $1=$ AREA unified weighted population/HCHS MFF
(b) iAREA need $2=$ AREA unified weighted population $/$ HCHS, prescribin \& GMS MFFs
(b) iAREA need $2=$ AREA unified weighted population/HCHS, prescribin \& GMS MFFs
(c) iAREA need $3=$ AREA unified weighted population/HCHS \& prescribing MFFs
(d) robust standard errors in brackets, ${ }^{* * *} \mathrm{p}<0.01, * * \mathrm{p}<0.05, * \mathrm{p}<0.1$.

Table B8.10: table showing circulatory disease outcome models with various indicators of MFF and need


Notes: (a) iAREA need $1=$ AREA unified weighted population/HCHS MFE
(b) iAREA need $2=$ AREA unified weighted population/HCHS, prescribing \& GMS MFFs
(c) iAREA need 3=AREA unified weighted population/HCHS \& prescibing MFFs
(d) robust standard errors in brackets, ${ }^{* * *} \mathrm{p}<0.01, * * \mathrm{p}<0.05, * \mathrm{p}<0.1$

Table B8.11: table showing respiratory disease outcome models with various indicators of MFF and need


[^56](b) iAREA need $2=$ AREA unified weighted population/HCHS, prescribing \& GMS MFFs
(c) iAREA need 3=AREA unified weighted population/HCHS \& preseribing MFFs
(d) robust standard errors in brackets, ${ }^{* * *} \mathrm{p}<0.01, * * \mathrm{p}<0.05, *(\mathrm{p}<0.4)$

Table B8.12: table showing gastro-intestinal disease outcome models with various indicators of MFF and need

|  | (1) | (2) | (3) | (4) | (5) | (6) | (7) | (8) | (9) | (10) | (11) | (12) | (13) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | PBC 13 gastrointestinal | PBC 13 gastrointestinal | PBC 13 gastrointestinal | PBC 13 gastrointestinal | PBC 13 gastrointestinal | PBC 13 gastrointestinal | PBC 13 gastrointestinal | PBC 13 gastrointestinal | PBC 13 gastrointestinal | PBC 13 gastrointestinal | PBC 13 gastrointestinal | PBC 13 gastrointestinal | PBC 13 gastro |
|  | 2006/7 outcome model | 2006/7 outcome model | 2006/7 outcome model | 2006/7 outcome model | 2006/7 outcome model | 2006/7 outcome model | 2006/7 outcome model | 2006/7 outcome model | 2006/7 <br> outcome model | $2006 \times 7$ <br> outcome <br> model | 2006/7 outcome model | 2006/7 outcome model | 2006/7 outcome model |
|  | uses SMR <br> instrument <br> spend | uses SMR <br> instrument <br> spend | uses SMR <br> instrument <br> spend | uses SYLLR <br> instrument <br> spend | uses SYLLR <br> instrument <br> spend | uses SYLLR <br> instrument <br> spend | uses SMR <br> instrument <br> spend | uses SMR instrument spend | uses SMR instrument <br> spend (5) | uses SMR <br> instrument <br> spend | uses SMR <br> instrument <br> spend | uses SMR <br> instrument <br> spend | uses SMR <br> instrument <br> spend |
|  | no weighting | no weighting | no weighting | no weighting | no weighting | no weighting | weighted | no weightin | weighted | no weighting | weighted | no weighting | weighted |
|  | second stage | second stage | second stage | second stage | second stage | second stage | second stage | second sta | second stage | second stage | second stage | second stage | second stage |
|  | iAREA need1 HCHS MFF | PBRA need HCHS MFF | CARAN need HCHS MFF | iAREA need1 HCHS MFF | PBRA need HCHS MFF | CARAN need HCHS MFF | iAREA need1 HCHS MFF | iAREA nee CARAN 3 MEES | iAREA need2 CARAN 3 MFEs | iAREA need3 CARAN 2 MFEs | iAREA need3 CARAN 2 MFFs. | CARAN need CARAN 3 MFFs | CARAN need CARAN 3 MFFs |
|  |  |  |  |  |  |  |  | ( |  |  |  |  |  |
| needAREA1 | $3.853^{* * *}$ |  |  | $3.966^{* *}$ |  |  | 3.779*** |  |  |  |  |  |  |
|  | [0.551] |  |  | [0.558] |  |  | [0.499] |  |  |  |  |  |  |
| gastro spend per head | -1.755*** | -1.420*** | $-1.641 * * *$ | $-1.544^{* * *}$ | -1.180*** | -1.404*** | 750** | -1.275*** | -1.317*** | $-1.275^{* * *}$ | $-1.315 * * *$ | $-2.056 * * *$ | $-2.192^{* * *}$ |
|  | [0.397] | [0.353] | [0.427] | [0.399] | [0.358] | [0.429] | 855 | [0.335] | [0.326] |  | [0.325] | [0.589] | [0.574] |
| needPBRA |  | $\begin{aligned} & 3.342^{* * *} \\ & {[0.486]} \end{aligned}$ |  |  | $\begin{aligned} & 3.413^{* * *} \\ & {[0.498]} \end{aligned}$ |  |  |  |  |  |  |  |  |
| needCARAN |  |  | 3.794*** |  |  | 3.887** |  |  |  |  |  | 4.140*** | 4.250*** |
|  |  |  | [0.612] |  |  | ${ }^{[0.621]}$ |  |  |  |  |  | [0.768] | [0.710] |
| needAREA2 |  |  |  |  |  | D) |  | $\begin{aligned} & 3.426^{* * *} \\ & {[0.466]} \end{aligned}$ | $\begin{aligned} & 3.479 * * * \\ & {[0.419]} \end{aligned}$ |  |  |  |  |
| needAREA3 |  |  |  |  |  |  |  |  |  | 3.393*** | 3.443*** |  |  |
|  |  |  |  |  |  |  |  |  |  | [0.462] | [0.415] |  |  |
| Constant | -2.155** | -1.251 | -1.838 | -0.954 | 0.028 | -0.566 | -2.166** | 7.919*** | 8.073*** | 7.916*** | 8.064*** | 11.273*** | 11.835*** |
|  | [1.047] | [0.928] | [1.121] | [1.054] | 10.943 | [1.127] | [1.016] | [1.431] | [1.391] | [1.430] | [1.387] | [2.524] | [2.460] |
| Observations | 152 | 152 | 152 | 152 | 52 | 152 | 152 | 152 | 152 | 152 | 152 | 152 | 152 |
| Endogeneity test statistic | 23.347 | 18.985 | 25.405 | 17.048 | 11.389 | 17.857 | 16.834 | 16.638 | 11.980 | 16.689 | 11.942 | 25.632 | 22.341 |
| Endogeneity p-value | 1.35-06 | $1.32 \mathrm{e}-05$ | $4.65 \mathrm{e}-07$ | $3.65 \mathrm{e}-05$ | 0.000739 | 2.38e-05 | 4.08e-05 | 4.52e-05 | 0.000538 | $4.40 \mathrm{e}-05$ | 0.000549 | $4.13 \mathrm{e}-07$ | 2.28e-06 |
| Hansen-Sargan test statistic | 3.067 | 4.604 | 1.555 | 4.936 | 7.575 | 2.637 | 7.476 | 5.029 | 8.762 | 4.907 | 8.714 | 1.284 | 3.554 |
| Hansen-Sargan p-value | 0.216 | 0.100 | 0.459 | Q, 0847 | 0.0227 | 0.268 | 0.0238 | 0.0809 | 0.0125 | 0.0860 | 0.0128 | 0.526 | 0.169 |
| Shea's partial R-squared | 0.193 | 0.231 | 0.200 | 0.193 | 0.231 | 0.200 | 0.191 | 0.208 | 0.198 | 0.208 | 0.198 | 0.139 | 0.135 |
| Kleibergen-Paap LM test statistic | 16.47 | 17.51 | 16.32 | 16.47 | 17.51 | 16.32 | 17.68 | 17.09 | 18.26 | 17.14 | 18.34 | 13.39 | 13.98 |
| Kleibergen-Paap p-value | 0.000910 | 0.000556 | 0.000974 | 0.000910 | 0.000556 | 0.000974 | 0.000511 | 0.000679 | 0.000389 | 0.000661 | 0.000375 | 0.00386 | 0.00293 |
| Kleibergen-Paap F statistic | 12.12 | 13.24 | 10.70 | 12.12 | 13.24 | 10.79 | 11.96 | 13.23 | 12.98 | 13.24 | 13.00 | 7.550 | 7.248 |
| Pesaran-Taylor reset statistic | 0.233 | 0.0427 | 0.0897 | 1.246 | 1.121 | 0.258 | 0.170 | 0.0935 | 0.443 | 0.0893 | 0.411 | 0.00841 | 0.117 |
| Pesaran-Taylor p -value | 0.629 | 0.836 | 0.765 | 0.264 | 0.290 | 0.611 | 0.680 | 0.760 | 0.506 | 0.765 | 0.521 | 0.927 | 0.732 |

Notes: (a) iAREA need $1=$ AREA unified weighted population/HCHS MFF
(b) iAREA need $2=$ AREA unified weighted population/HCHS, prescribing \& GMS MFFs
(c) iAREA need 3=AREA unified weighted population/HCHS \& prescribing MFFs
(d) robust standard errors in brackets, *** $\mathrm{p}<0.01, * * \mathrm{p}<0.05, * \mathrm{p}<0.1$

The assimilation of the impact of alternative measures of need, weights, and MFFs proved overwhelming. Instead, we approached the selection of the appropriate need $\sim$ weighting $\sim M F F$ combination from an a priori perspective. The AREA-based need formula has been replaced by the CARAN formula for the purposes of resource allocation and therefore it must be believed to be a better indicator of relative health care need. The PBRA approach is relatively new and has not been implemented yet. We therefore decided to use the CARAN based measure as our indicator of the level of need.

With some PCTs several times larger than others, it is difficult to justify giving them all the same weighting. It was therefore decided to weight all of our models by PCT size (where size is measured by the PCT's population).

We also decided to use the 'own population' expenditure data on the grounds that all NHS expenditure, irrespective of its funding source, should be included in the analysis (although there is the issue about how this income is split between PBCs).

Finally, it was decided to focus on the use of the SYLLR as the outcome indicator, and to proxy 'other programme need' in the expenditure equation using the all cause SYLLR minus the own programme SYLLR.

## B8.3 Model estimation using 2006/7 expenditure data and mortality data for 2004/2006: CARAN need and three MFFs

Initially, acceptable models were obtained using the CARAN measure of need and adjusting expenditure for local input prices using a weighted average of the MFFs for all three services (HCHS, prescribing, and GMS/PMS). The outcome and expenditure results for the big fout programmes are shown in Table B8.13 with the relevant outcome and expenditure elasticities highlighted.

In all four outcome models expenditure has a significant negative effect on mortality and, in three of these; the all service measure of need has a significant positive effect. In the respiratory outcome model, where the all service need term is not significant, there is another indicator of need - the proportion of the population that are permanently sick -and this is both positive and statistically significant. The diagnostic statistics suggest that, in all four cases, own programme expenditure is endogenous and that the instruments are valid. They also suggest that the instruments are relevant. There is some evidence that the instruments are slightly weak in one of the four outcome results (the respiratory model). ${ }^{27}$ The Pesaran-Taylor test suggests that there is no evidence of model mis-specification.

In all four expenditure models both the need and budget variables have a positive and significant effect on own programme expenditure. In addition, the proxy for need in other programmes is negative and significant in all four cases. In the gastro-intestinal expenditure programme the prevalence of lone pensioners households is associated with less NHS expenditure; there might be some unmet need here or perhaps this is a self-selecting group.

The diagnostic statistics suggest that, for all four expenditure models, the proxy for other programme need is endogenous and that the instruments are valid. They also suggest that the instruments are relevant and, with the possible exception of the gastro-intestinal expenditure result, there is no evidence that the instruments are weak. The Pesaran-Taylor test suggests that there is no evidence of model misspecification.

The elasticities shown in Table B8.13 can be used to calculate the cost of a life year in each programme and these calculations -- for both these four programmes as well as for the other six programmes with a mortality based outcome indicator -- are shown in Tables B8.14 and B8.15 (the full outcome and expenditure models for the other six programmes with a mortality based outcome indicator are not shown here).

[^57]Table B8.14 reveals that the cost of a life year for the big four programmes combined is $£ 11,298$. This is remarkably close to the figure obtained using expenditure data for 2005/6, an AREA-based measure of need, and a HCHS MFF ( $£ 12,855$ ). The cost of a life year for all ten programmes with a mortality based measure of need the cost of a life year is $£ 21,743$, which is even closer to the figure obtained using 2005/6 expenditure data ( $£ 21,256$ ). If we assume a zero gain in the 13 programmes without a mortality based indicator then the cost per life year across all 23 programmes is $£ 66,318$ (it is $£_{,} 56,799$ for 2005/6 data).

Alternatively, if we assume that PBC23 generates a zero health gain and that the gain attributable to the remaining 12 programmes is, on average, the same as that attributable to those with a mortality outcome measure, then Table B8.15 shows that the cost of a life year across all programmes is $£ 25,038$ (it is $£ 24,200$ for 2005/6 data).

Table B8.13: table showing outcome and expenditure models for the big four programmes using spend data (incorporating three MFFs) for 2006/7

| Regressors | (1) <br> PBC 2 <br> cancer <br> 2006/7 <br> outcome model | (2) <br> PBC 2 <br> cancer <br> 2006/7 <br> spend model | (3) <br> PBC 10 <br> circulation 2006/7 <br> outcome model | (4) <br> PBC 10 <br> circulation <br> 2006/7 <br> spend model | (5) <br> PBC 11 <br> respiratory <br> 2006/7 <br> outcome model | (6) <br> PBC 11 <br> respiratory <br> $2006 \times 7$ spend model | (7) <br> PBC 13 <br> gastro <br> 2006/7 <br> outcome model | (8) <br> PBC 13 <br> gastro <br> 2006/7 <br> spend model |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| all cause SYLLR excluding cancer |  | $\begin{aligned} & -0.952^{* * *} \\ & {[0.179]} \end{aligned}$ |  |  |  | V |  |  |
| budget per head (HPG MFF) |  | 0.542** |  | 0.694** | $\sim$ | 0.712*** |  | 0.650** |
|  |  | [0.242] |  | [0.292] | $\bigcirc$ | [0.252] |  | [0.289] |
| need CARAN | $0.958 * * *$ | 1.765*** | 2.830*** | 2.185*** | 1.64 | 1.371*** | 4.609*** | 2.696*** |
|  | [0.129] | [0.286] | [0.252] | [0.355] | [1.192] | [0.297] | [0.700] | [0.679] |
| own programme spend per head | $-0.351^{* * *}$ |  | $-1.441^{* * *}$ |  | $52.830^{* * *}$ |  | -2.125*** |  |
|  | [0.117] |  | [0.219] |  | [0.767] |  | [0.563] |  |
| all cause SYLLR excluding circulatory problems |  |  |  | $-1.782 * * *$ |  |  |  |  |
| permanently sick |  |  |  | $\square$ | $\begin{aligned} & 1.371 * * * \\ & {[0.405]} \end{aligned}$ |  |  |  |
| all cause SYLLR excluding respiratory problems |  |  |  |  |  | $\begin{aligned} & -0.670 * * \\ & {[0.288]} \end{aligned}$ |  |  |
| all cause SYLLR excluding gastro-intestinal problems |  |  | $\bigcirc \bigcirc$ |  |  |  |  | $-1.856^{* * *}$ |
|  |  |  |  |  |  |  |  | [0.612] |
| lone pensioner households |  |  |  |  |  |  |  | $-0.593^{* *}$ |
|  |  |  |  |  |  |  |  | [0.297] |
| Constant | 6.588*** | 5.937*** | 11.538*** | 10.299*** | 18.965*** | 3.117 | 12.208*** | 9.752*** |
|  | [0.515] | [1.775] | [1.050] | [2.384] | [3.853] | [1.976] | [2.416] | [3.053] |
| Observations | 152 | 152 | 152 | 152 | 152 | 152 | 152 | 152 |
| Endogeneity test statistic | 14.496 | 20.274 | 43.352 | 25.784 | 27.923 | 7.922 | 21.862 | 13.531 |
| Endogeneity p-value | 0.000140 | (6.71e-06 | 0 | 3.82e-07 | $1.26 \mathrm{e}-07$ | 0.00488 | 2.93e-06 | 0.000235 |
| Hansen-Sargan test statistic | 0.208 | 0.293 | 1.507 | 0.542 | 1.879 | 0.356 | 1.006 | 0.0267 |
| Hansen-Sargan p-value | 0.649 | 0.588 | 0.681 | 0.462 | 0.170 | 0.550 | 0.316 | 0.870 |
| Shea's partial R-squared | $0.163 \sim$ | 0.445 | 0.303 | 0.296 | 0.0802 | 0.366 | 0.142 | 0.206 |
| Kleibergen-Paap LM test statistic | $16.97 \bigcirc$ | 42.38 | 32.53 | 32.70 | 10.51 | 36.33 | 15.00 | 19.07 |
| Kleibergen-Paap p-value | 0.000207 | $6.28 \mathrm{e}-10$ | $1.49 \mathrm{e}-06$ | 7.93e-08 | 0.00523 | $1.29 \mathrm{e}-08$ | 0.000553 | 7.22e-05 |
| Kleibergen-Paap F statistic | 12.47 ) | 48.32 | 17.31 | 25.71 | 7.482 | 24.32 | 11.80 | 8.660 |
| Pesaran-Taylor reset statistic | 5.471 | 0.00111 | 0.0912 | 0.0183 | 3.090 | 1.915 | 0.267 | 0.0880 |
| Pesaran-Taylor p-value | 0.0193 | 0.973 | 0.763 | 0.892 | 0.0788 | 0.166 | 0.605 | 0.767 |

Pesaran-Taylor p -value
Note: all spend figures are on a net population basis and are adjusted for local prices using three MFFs from the Department of Health's resource allocation exposition book for $2009 / 10$. All estimated models use 152 PCTs and are weighted by PCT population. The SYLLR is the mortality indicator
There are several differences between the models estimated here and those reported in Martin, Rice and Smith (2008b):
(i) here we use net population spend data (not net spend data);
(ii) here we use three MFFs (not solely the HCHS MFF); and
(iii) here we use a consistent definition of the 'other programme need' proxy across all programmes (i.e., all cause SYLLR minus the own programme SYLLR).

Table B8.14: table showing cost of life and life year estimates using spend data for 2006/7 (three MFFs) and outcome data for 2004/06 (assumes zero gain for 13 programmes)


Note that the annual mortality figures reported in cells G5 \& G6 and G13 \& G14 are identical because we do not have mortality data for 2002/04.
Note that, for 2006/7, the neonate category has been merged with maternity to obtain plausible outcome and expenditure models.

Table B8.15: table showing cost of life and life year estimates using spend data for 2006/7 (three MFFs) and outcome data for 2004/06 (assumes some gain in other 13 programmes)


Assume zero health gain in PBC23, and gain in ten PBCs applies to other 12 PBCs

| 17 | PBC23 | £10,585 | 0.844 | £89.34 |  | 0.00 |  | 0.00 |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 18 | PBC23 2005/6 | £8,449 | 0.926 | £.78.24 |  | 0.00 |  | 0.00 |  |
| 19 | Other 12 programmes | £24,400 |  | £.367.01 |  | 1,499.56 | $£ 244,747$ | 16,880 | $£ 21,743$ |
| 20 | Other 12 PBCs 2005/6 | £,25,493 |  | $\ldots 324.20$ |  | 1,247.69 | £,259,838 | 15,252 | £,21,256 |
| 21 | All 23 programmes | £67,896 |  | 6.678 .96 |  | 2,409.11 | £281,830 | 27,118 | £25,038 |
| 22 | All 23 programmes 2005/6 | £64,310 |  | $£ 643.10$ |  | 2,173.90 | £295,827 | 26,575 | £24,200 |
|  | Note: | 2006/7 | 2005/6 |  |  |  |  |  |  |
| 23 | All 23 programme spend | £67,896 | 604,310 |  |  |  |  |  |  |
| 24 | \% change in budget | 1.00 | 1.00 |  |  |  |  |  |  |
| 25 | proportionate change | 0.01 | 0.01 |  |  |  |  |  |  |
| 26 | Change in budget | ¢678.96 | £ 643.10 |  |  | 14 immedi |  |  |  |

## B8.4 Model estimation using 2006/7 expenditure data and mortality data for 2004/2006: CARAN need and two MFFs

Further discussion by the project team noted that the PB data incorporates all PCT expenditure and that, as there is a separate category for GMS/PMS expenditure (PBC23a), it seems appropriate that the GMS/PMS MFF should be applied to this category. However, other categories of expenditure exclude GMS/PMS expenditure but incorporate both HCHS and prescribing expenditure. It therefore seems appropriate that a weighted averaged of the HCHS and prescribing MFFs should be applied to these other (non-GMS/PMS) categories of expenditure.

We therefore re-estimated the outcome and expenditure models for those programmes with a mortality based outcome indicator using the CARAN measure of need and adjusting expenditure for local input prices using the MFFs for HCHS and prescribing services. The outcome and expenditure results for the big four programmes are shown in Table B8.16 with the relevant outcome and expenditure elasticities again highlighted (the first-stage regressions associated with these results can be found in Table BA. 5 in the annex).

In all four outcome models expenditure has a significant negative effect on mortality and, in three of these, the all service measure of need has a significant positive effect. In the respiratory outcome model, where the all service need term is not significant, there is another indicator of need - the proportion of the population that are permanently sick - and this is both positive and statistically significant. The all service measure of need squared is also positive and significant in the canceroutcome equation. The diagnostic statistics suggest that, in all four cases, own programme expenditure is endogenous and that the instruments are valid. They also suggest that the instruments are relevant. There is a little evidence that the instruments are weak in one of the four outcome results, namely the respiratory model. Reestimation of the latter model but without the least significant instrument generates a coefficient of - 3.507 on expenditure and the Kleibergen-Paap F statistic now exceeds ten (it is 11.799). The Pesaran-Taylor test suggests that there is no evidence of model mis specification in any of the outcome models.

In all four expenditure models both the need and budget variables have a positive and significant effect on own programme expenditure. In addition, the proxy for need in other programmes is negative and significant in all four cases. In the gastro-intestinal expenditure programme the prevalence of lone pensioners households is associated with less NHS expenditure; there might be some unmet need here or perhaps this is self-selecting group.

The diagnostic statistics suggest that, for all four expenditure models, expenditure is endogenous and the instruments are valid. They also suggest that the instruments are relevant and, with the possible exception of the gastro-intestinal expenditure result, there is no evidence that the instruments are weak. Re-estimation of the gastro-intestinal expenditure model without the least significant instrument generates a coefficient of 0.667 on the budget variable and the Kleibergen-Paap F statistic now exceeds ten (it is 16.871). The Pesaran-Taylor test suggests that there is no evidence of model mis-specification.

The outcome and expenditure elasticities are little changed from those presented in Table B8.13 and, like those, these new elasticities can be used to calculate the cost of a life year in each programme. These calcutations -- for both these four programmes as well as for the other six programmes with a mortality based outcome indicator -- are shown in Tables B8.17 and B8.18 (the full outcome and expenditure models for the other six programmes with a mortality based outcome indicator are not shown here).

The figures for 2006/7 in Table B8.17 (which incorporate two MFFs) can be compared with those for 2006/7 in Table B8.14 (which incorporate three MFFs). Table B8.17 reveals that the use of a different MFF has little impact on the cost of a life year for the big four PBCs (it was $£, 11,298$, it is now $£, 10,783$ ) as well as on the cost of a life year for all programmes with a mortality outcome measure (was $£ 21,743$, now $£ 20,893$ ).

In addition, Table B8.18 shows that if we assume that PBC 23 generates a zero health gain and that the gain attributable to the remaining 12 programmes is, on average, the same as that attributable to those
with a mortality outcome measure, then the cost of a life year across all programmes is now $£ 23,697$ (it was $£ 25,038$ for 2006/7 in Table B8.15).

The figures in Table B8.18 also reveal that the cost of a life year in $2006 / 7$ for all programmes $(£ 23,697)$ is little changed from the comparable figure for 2005/6 ( $£ 24,200)$.

Table B8.16: table showing outcome and expenditure models for the big four programmes using spend data for 2006/7 (incorporating two MFFs) and mortality data for 2004/5/6

|  | (1) | (2) | (3) | (4) | (5) | (6) | (7) | (8) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | $\text { PBC } 2$ | $\text { PBC } 2$ | PBC 10 | PBC 10 | $\text { PBC } 11$ | PBC 11 | PBC 13 | PBC 13 |
|  | cancer | cancer | circulation | circulation | respiratory | respiratory | gastro | gastro |
|  | $2006 / 7$ | $2006 / 7$ | $2006 / 7$ | $2006 / 7$ | $2006 / 7$ | 2006/7 | $2006 / 7$ | $2006 / 7$ |
|  | outcome model | spend model | outcome model | spend model | outcome model | spend model | outcome model | spend model |
| own programme spend per head | -0.337*** |  | -1.447*** |  | -2.839*** |  | -2.137*** |  |
|  | [0.104] |  | [0.220] |  | [0.772] |  | [0.569] |  |
| need CARAN per head | 0.974*** | 1.772*** | 2.860*** | 2.191*** | 1.782 | 375* | $4.657 * * *$ | $2.697 * * *$ |
|  | [0.110] | [0.287] | [0.257] | [0.355] | [1.198] | 0.297] | [0.716] | [0.676] |
| needCARAN per head squared | 1.314*** |  |  |  | - |  |  |  |
|  | [0.352] |  |  |  | (0) |  |  |  |
| all cause SYLLR excluding cancer |  | $-0.951 * * *$ |  |  |  |  |  |  |
|  |  | [0.180] |  |  | N |  |  |  |
| PCT budget per head |  | 0.548** |  | 0.701** |  | 0.718*** |  | 0.655** |
|  |  | [0.242] |  | [0.292] |  | [0.253] |  | [0.289] |
| all cause SYLLR excluding circulatory disease |  |  |  | -1.778*** |  |  |  |  |
| permanently sick aged 16-74 |  |  |  | - | $1.385^{* * *}$ |  |  |  |
|  |  |  |  | ) | [0.405] |  |  |  |
| all cause SYLLR excluding respiratory problems |  |  |  |  |  | -0.663** |  |  |
|  |  |  | $\bigcirc$ |  |  | [0.288] |  |  |
| all cause SYLLR excluding gastro-intestinal problems |  |  | $\leq$ |  |  |  |  | -1.847*** |
|  |  |  | , |  |  |  |  | [0.609] |
| lone pensioner households |  |  | D |  |  |  |  | -0.590** |
|  |  |  |  |  |  |  |  | [0.295] |
| Constant | 6.506*** | 5.881*** | $11.567 * * *$ | 10.227*** | 19.047*** | 3.032 | $12.260^{* * *}$ | 9.664*** |
|  | [0.455] | [1.778] | [1.058] | [2.387] | [3.877] | [1.977] | [2.441] | [3.046] |
| Endogeneity test statistic | 15.173 | 20.248 | 43.405 | 25.854 | 27.876 | 7.863 | 21.853 | 13.607 |
| Endogeneity p-value | $9.81 \mathrm{e}-05$ | $6.80 \mathrm{e}-06$ | 0 | $3.68 \mathrm{e}-07$ | $1.29 \mathrm{e}-07$ | 0.00505 | $2.94 \mathrm{e}-06$ | 0.000225 |
| Hansen-Sargan test statistic | 0.00201 | 0.300 | 1.440 | 0.530 | 1.912 | 0.344 | 1.011 | 0.0294 |
| Hansen-Sargan p-value | 0.964 | 0.580 | 0.696 | 0.467 | 0.167 | 0.557 | 0.315 | 0.864 |
| Shea's partial R-squared | 0.164 | 0.445 | 0.300 | 0.296 | 0.0793 | 0.366 | 0.140 | 0.206 |
| Kleibergen-Paap LM test statistic | 17.85 | 42.38 | 32.37 | 32.70 | 10.42 | 36.33 | 14.86 | 19.07 |
| Kleibergen-Paap p-value | 0.000133 | $6.28 \mathrm{e}-10$ | $1.61 \mathrm{e}-06$ | $7.93 \mathrm{e}-08$ | 0.00545 | $1.29 \mathrm{e}-08$ | 0.000592 | $7.22 \mathrm{e}-05$ |
| Kleibergen-Paap F statistic | 13.28 | 48.32 | 17.14 | 25.71 | 7.390 | 24.32 | 11.63 | 8.660 |
| Pesaran-Taylor reset statistic | 0.00226 | 0.00178 | 0.0945 | 0.0215 | 3.139 | 1.908 | 0.266 | 0.0605 |
| Pesaran-Taylor p-value | 0.962 | 0.966 | 0.759 | 0.883 | 0.0764 | 0.167 | 0.606 | 0.806 |

Note: robust standard errors in brackets, ${ }^{* * *} \mathrm{p}<0.01$, ** $\mathrm{p}<0.05,{ }^{*} \mathrm{p}<0.1$

Table B8.17: table showing cost of life and life year estimates using spend data for 2006/7 (two MFFs) and outcome data for 2004/06 (assumes zero gain for 13 programmes) $\qquad$ D


Note that the annual mortality figures reported in cells G5 \& G6 and G13 \& G14 are identical because we do not have mortality data for $2002 / 04$.
Note that, for $2006 / 7$, the neonate category has been merged with maternity to obtain plausible outcome and expenditure models.

Table B8.18: table showing cost of life and life year estimates using spend data for 2006/7 (two MFFs) and outcome data for 2004/06 (assumes some


Note that the annual mortality figures reported in cells G5 \& G6 and G13 \& G14 are identical because we do not have mortality data for $2002 / 04$.
Note that, for $2006 / 7$, the neonate category has been merged with maternity to obtain plausible outcome and expenditure models.

## B8.5 Model estimation using 2006/7 expenditure data and mortality data for 2006/2008: CARAN need and two MFFs

One shortcoming with the models presented above is that they relate expenditure in $2006 / 7$ to mortality in the same period and in the two previous periods (i.e., in 2004, 2005 and 2006). The difficulty with this is that one would expect expenditure in year $t$ to affect mortality in year $t$ and possibly subsequent years $(t+1, t+2$, etc) but not mortality in previous years ( $t-1, t-2$, etc). However, if we assume that PCTs have reached some sort of equilibrium in the expenditure choices they make and the outcomes they secure, so that expenditure levels change relatively little from one year to the next, then mortality over the three year period $t, t-1$ and $t-2$ might be a good proxy for mortality in $t, t+1$ and $t+2$. Indeed, this is probably not an unreasonable assumption given the relatively slow pace at which both types of variable change.

Although this assumption of equilibrium is not an unreasonable one, it is one that ideally we would like to be able to drop. Fortunately, with the recent availability of more up-to-date mortality data, we have the opportunity to relate expenditure in 2006 to mortality in the same year and in the two following years (i.e., in 2006, 2007 and 2008). ${ }^{28}$ Thus the models reported in Table B8.16 were re-estimated replacing the mortality rate for $2004 / 5 / 6$ with that for $2006 / 7 / 8$. The outcome and expenditure results for the big four programmes are shown in Table B8.19 with the relevant outcome and expenditure elasticities again highlighted (the first-stage regressions associated with these results can be found in Table BA. 6 in the annex). These elasticities are similar to those presented previously in Table B8.16 but there are some changes (e.g, the outcome elasticity in the respiratory outcome equation falls from -2.839 to -2.029 ).

In all four outcome models expenditure has a significant negative effect on mortality and the all service measure of need has a significant positive effect. The all service measure of need squared is also positive and significant in the cancer outcome equation. In the respiratory outcome model, there is an additional indicator of need - the proportion of the population that are (permanently sick - and this is both positive and statistically significant. The diagnostic statistics suggest that, in all four cases, own programme expenditure is endogenous and that the instruments are valid. They also suggest that the instruments are relevant. There is no evidence that the instruments are weak in three of the four outcome results. The Pesaran-Taylor test suggests that there is no evidence of model mis-specification

However, the Kleibergen-Paap F statistic for the respiratory disease outcome model is 7.022 and this is less than the 'critical' target of 10.0. This indicates that the instruments may be weak. However, if we reestimate this model having dropped the least significant instrument, the coefficient on own programme expenditure is now -2.622 and this is significant at the $1 \%$ level. Moreover, there is now no evidence of weak instruments (the Kleibergen-Paap F statistic is 11.025 ) and it is this coefficient that we use for the respiratory outcome model in the cost of a life year calculations below.

In three of the four expenditure models both the need and budget variables have a positive and significant effect on oun programme expenditure. In addition, the proxy for need in other programmes is negative and significant in all four cases. The diagnostic statistics suggest that, for all four expenditure models, expenditure is endogenous and the instruments are valid. They also suggest that the instruments are relevant and there is no evidence that the instruments are weak. The Pesaran-Taylor test suggests that there is no eyidence of model mis-specification.

The outcome and expenditure elasticities presented in Table B8.19 can be used to calculate the cost of a life year in each programme. These calculations -- for both the big four programmes as well as for the other six programmes with mortality based outcome indicator -- are shown in Table B8.20. They show that the use of a more appropriate measure of mortality (i.e., for 2006/2007/2008 rather than for $2004 / 2005 / 2006$ ) slightly increases the cost of a life year for the big four PBCs (from $£ 10,783$ to $£ 12,333$ ) as well as for all ten programmes with a mortality outcome measure (from $£ 20,893$ to $£ 23,780$ ).

[^58]In addition, Table B 8.21 shows that if we assume that PBC 23 generates a zero health gain and that the gain attributable to the remaining 12 programmes is, on average, the same as that attributable to those with a mortality outcome measure, then the cost of a life year across all programmes is now $£ 26,876$ (it was $£^{2} 23,697$ using mortality for $2004 / 5 / 6$ ).

Table B8.19: table showing outcome and expenditure models for the big four programmes using spend data for 2006/7(two MFFs) and mortality data for 2006/7/8


Table B8.20: table showing cost of life and life year estimates using spend data for 2006/7 and outcome data for 2006/7/8 (assumes zero health gain for 13 programmes)


Table B8.21: table showing cost of life and life year estimates using spend data for 2006/7 and outcome data for 2006/7/8 (assumes average health gain for 12 other programmes)


## B8.6 Adjusting the cost of life (year) estimates for the mismatch in the ICD10 coverage of the expenditure and the mortality data

The cost of a life (year) estimates presented in Tables B8.20 and B8.21 assume a $1 \%$ increase in each PCT's budget and are calculated as:
the cost of an additional life in a particular programme
$=$ the change in expenditure in that programme / the change in mortality in that programme $=$ (annual spend $*$ expenditure elasticity) $/$ (annual mortality * outcome elasticity)
and
the cost of an additional life year in a particular programme
$=$ the change in expenditure in that programme / the change in life years lost in that programme
$=$ (annual spend $*$ expenditure elasticity) $/$ (annual mortality $*$ outcome elasticity $)$
Thus an integral part of the calculation of the cost of a life (year) is the annual mortality (life years lost) figure associated with a particular programme. Ideally, the ICD10 coverage of the expenditure data should coincide with that of the mortality data but, as know from Table B5.1, the 1CD10 coverage of the mortality data typically falls short of that for the expenditure data. Unless we adjust the annual mortality figure so that its ICD10 coverage approximates that of the expenditure data, our cost of life (year) estimates will usually be too large because they will usually underestimate the mortality gain.

Table B8.22 reproduces Table B8.20 but incorporates this ICD 10 coverage adjustment (see columns L and R in Table B8.22). This adjustment reduces the cost of a life year:

- for the big four programmes from $£ 12,333$ to $£ 10,604$
- for the ten programmes with a mortalitybased outcome indicator from $£ 23,780$ to $£ 19,965$
- for all programmes assuming a zero gain for the 13 PBCs without an outcome indicator from $£ 87,494$ to $£ 73,457$.

Similarly, Table B8.23 reproduces Table B8.21 but incorporates this ICD 10 coverage adjustment (see columns L and R again). If we assume a zero health gain in PBC 23 and an average gain in the other 12 PBCs without a mortality based outcome indicator, then this adjustment reduces the cost of a life year for all programmes from $£ 26,87 \hat{6}$ to $£ 22,565$.

TableB8.22: table showing Cost of life and life year estimates using expenditure data for 2006 and outcome data for 2006/7/8 (assumes zero health gain for 13 programmes) adjusted for the ICD10 coverage of the expenditure and outcome data

| A | B | C <br> Spend <br> (£m) <br> 2006/7 | D <br> Spend elasticity | $\begin{aligned} & \mathrm{E} \\ & =0.01 * \mathrm{C}^{*} \\ & \mathrm{D} \\ & \\ & \text { Change in } \\ & \text { spend } \\ & (\mathrm{f}, \mathrm{~m}) \\ & \hline \end{aligned}$ | F | G <br> Annual mortality, $<75$ years, 2006/08 | H <br> Outcome <br> elasticity <br> (without <br> negative sign | $\begin{array}{ll} \text { I } \quad \text { J } \\ =0.01^{*} D^{*} G^{*} \\ H \end{array}$ <br> Change in annual mortality | K <br> =E/I <br> Cost per life gained $(\AA)$ | L <br> Coverage of mortality data relative to spend data | M <br> $=I / L$ <br> Change in annual mortality adj for coverage | N <br> =E/M <br> Cost per life gained $\left(f_{\text {. }}\right)$ adj for coverage | P <br> Total life years lost, $<75 y e a r s$, 2006/08 | Q <br> $=0.01 * \mathrm{D} * \mathrm{H}$ <br> P/3 <br> $\uparrow$ <br> Change in annual life years lost | Coverage of mortality data relative to spend data | $=\mathrm{Q} / \mathrm{R}$ <br> Change in annual life years lost adj for YLL | T <br> =E/Q <br> Cost per life year gained ( $\AA$ ) | $=\mathrm{E} / \mathrm{S}$ <br> Cost per life year gained adj for YLL coverage ( $\epsilon_{\text {, }}$ ) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | Cancer | $¢_{6} 4,122$ | 0.465 | $£_{6} 19.17$ |  | 61,961 | 0.342 | 98.54 | £.194,520 | 0.984 | 100.14 | $£ 191,407$ | 2,207,021 | 1,170 | 0.984 | 1,189 | $£_{616,383}$ | $£_{6} 16,121$ |
| 2 | Circulatory problems | $£_{6,161}$ | 0.540 | £.33.27 |  | 41,106 | 1.434 | 318.31 | $£ 104,519$ | 0.992 | 320.88 | £103,683 | -1,361, 314 | 3,515 | 0.992 | 3,543 | £9,466 | £, 930 |
| 3 | Respiratory problems | $£_{6,285}$ | 0.679 | $£^{22.31}$ |  | 11,574 | 2.622 | 206.06 | $£ 108,248$ | 0.773 | 262.57 | $¢_{¢} 83,676$ | 324,223 | 1,924 | 0.773 | 2,489 | $¢_{6} 11,593$ | $¢_{68,961}$ |
| 4 | Gastro-intestinal problems | $\mathrm{E}^{\text {. }}$, 700 | 0.446 | £ ¢ $16.50^{1}$ |  | 6,160 | 1.536 | 42.20 | £ 391,048 | 0.571 | 73.90 | $\ldots 223,288$ | 345,908 | 790 | 0.571 | 1,383 | $\mathrm{E}^{2} 20,892$ | £ 11,929 |
|  | Big four programmes summary: |  |  |  |  |  |  |  |  |  |  | ( |  |  |  |  |  |  |
| 5 | Spend 2006 \& mortality 2006/8 | $£_{\text {¢17,268 }}$ |  | £91.24 |  | 120,801 |  | 665.10 | $£ 137,188$ |  | 761.49 | 119823 | 4,238,786 | 7,399 |  | 8,604 | $£_{\text {¢12,333 }}$ | $£_{6}^{10,604}$ |
| 6 | Spend 2006 \& mortality 2004/6 | $£_{6}^{17,268}$ |  | $£^{114.04}$ |  | 125,290 |  | 953.13 | £.119,650 |  |  | ( | 4,355,559 | 10,576 |  |  | $¢_{6} 10,783$ |  |
| 7 | Spend 2005 \& mortality 2002/4 | $£_{1} 17,625$ |  | £.141.22 |  | 125,290 |  | 909.96 | $£ .155,196$ |  |  |  | 4,516,953 | 10,986 |  |  | $¢_{6} 12,855$ |  |
| 8 | Infectious diseases | $£_{6}^{1,053}$ | 0.792 | $¢_{6.34}$ |  | 2,050 | 0.047 | 0.76 | $£_{6}^{10,928,905}$ | 1.000 | 0.76 | 10,928,905 | 106,552 | 13 | 1.000 | 13 | $£ 630,798$ | £630,798 |
| 9 | Endocrine problems | $£_{61,852}$ | 0.953 | $£ .17 .65$ |  | 1,542 | 0.842 | 12.37 | $£_{6}^{1}, 426,410$ | 0.634 | . 52 | $\ldots 904,344$ | 57,672 | 154 | 0.634 | 243 | $£ 114,416$ | $£_{6} 72,539$ |
| 10 | Neurological problems | $£_{\text {¢ } 2,790}$ | 0.616 | £.17.19 |  | 727 | 0.112 | 0.50 | $¢_{3} 34,265,082$ | 0.136 | 3.69 | £4,660,051 | 66,137 | 15 | 0.136 | 112 | £.1,129,960 | £153,675 |
| 11 | Genito-urinary problems | $\mathrm{E}_{6}, 482$ | 0.912 | £.31.76 |  | 294 | 0.051 | 0.14 | $£_{2} 232,226,224$ | 0.172 | . 80 | $£_{6} 39,942,910$ | 10,030 | 2 | 0.172 | 9 | $£^{2} 20,421,090$ | ¢.3,512,427 |
| 12 | Trauma \& injuries* | $£^{\text {£ } 2,892}$ | 0.358 | $£_{6}^{10.35}$ |  | 1,037 | 0 | 0.00 | \#DIV/0! | 0,15 |  | \#DIV/0! | 30,000 | 0 | 0.175 | 0 | \#Div/0! | \#Div/0! |
| 13 | Maternity \& neonates* | $¢_{6,574}$ | 0.224 | $\ldots 8.01$ |  | 2,189 | 0.482 | 2.36 | $¢_{6}{ }^{3}, 387,363$ | 8.213 | 0.29 | $¢_{2} 27,820,413$ | 492,600 | 177 | 0.679 | 261 | $¢_{4} 45,158$ | $\AA^{\ldots} 30,662$ |
|  | Other six programmes summary: |  |  |  |  |  |  |  |  | J |  |  |  |  |  |  |  |  |
| 14 | Spend 2006 \& mortality 2006/8 | $£_{15,643}$ |  | £93.29 |  | 7,839 |  | 16.14 | £.5,780,723 |  | 25.05 | $£ 3,724,129$ | 762,991 | 362 |  | 639 | $£ 258,046$ | £.146,108 |
| 15 | Spend 2006 \& mortality 2004/6 | $£_{15,643}$ |  | £.112.13 |  | 7,923 |  | 18.17 | E6, 6172,491 |  |  |  | 757,531 | 249 |  |  | $£ 449,706$ |  |
| 16 | Spend 2005 \& mortality 2002/4 | $¢_{122,743}$ |  | £99.44 |  | 7,923 |  | 16.26 | 6,115,621 |  |  |  | 751,009 | 337 |  |  | $\ldots 295,074$ |  |
|  | All ten programmes summary: |  |  |  |  |  |  |  | , |  |  |  |  |  |  |  |  |  |
| 17 | Spend 2006 \& mortality 2006/8 | \&32,911 | 0.561 | £184.53 |  | 128,640 | 0.945 | 681.24 | 1270,881 |  | 786.54 | £234,617 | 5,001,777 | 7,760 |  | 9,243 | ¢ 23,780 | $¢_{6} 19,965$ |
| 18 | Spend 2006 \& mortality 2004/6 | $¢_{3} 32,911$ | 0.687 | £.226.18 |  | 133,213 | 1.061 | 971.30 | 232,861 |  |  |  | 5,093,090 | 10,826 |  |  | $¢_{\text {¢ } 20,893}$ |  |
| 19 | Spend 2005 \& mortality 2002/4 | $¢_{630,368}$ | 0.792 | £240.67 |  | 133,213 | 0.877 | 926.22 | 259,838 |  |  |  | 5,267,962 | 11,322 |  |  | $£_{\text {¢ } 21,256}$ |  |
|  | Assume zero health gain in the other 13 programmes |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
|  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| 20 | Spend 2006 \& mortality 2006/8 | ${ }_{6} 34,985$ | 1.413 | £494.43 |  |  |  |  |  |  |  |  |  | 0 |  | 0 |  |  |
| 21 | Spend 2006 \& mortality 2004/6 | $¢_{6} 34,985$ | 1.294 | ¢452.78 |  |  | - |  |  |  |  |  |  | 0 |  |  |  |  |
| 22 | Spend 2005 \& mortality 2002/4 | $\ldots 33,942$ | 1.186 | $¢_{4} 402.43$ |  |  |  | 0.00 |  |  |  |  |  | 0 |  |  |  |  |
|  | All 23 programmes |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| 23 | Spend 2006 \& mortality 2006/8 | $£_{6}^{67,896}$ |  | £678.96 |  |  |  | 681.24 | $£_{\text {¢ }} 996,655$ |  | 786.54 | £863,228 |  | 7,760 |  | 9,243 | £ 87,494 | $£_{6} 73,457$ |
| 24 | Spend 2006 \& mortality 2004/6 | $¢_{6}^{67,896}$ |  | £678.96 |  |  |  | 971.30 | $£ 699,024$ |  |  |  |  | 10,826 |  |  | $¢_{662,718}$ |  |
| 25 | Spend 2005 \& mortality 2002/4 | $£_{6} 64,310$ |  | $£_{6} 643.10$ |  |  |  | 926.22 | $£ 694,330$ |  |  |  |  | 11,322 |  |  | ¢56,799 |  |
|  | Note: | 2006/7 | 2005/6 |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| 26 | All 23 programme spend | £67,896 | $£_{64,310}$ |  |  | Note that the annual mortality figures reported in cells $\mathrm{G} 6 \& \mathrm{G} 7$ and G 15 \& G 16 are identical because we do not have mortality data for 2002/04. |  |  |  |  |  |  |  |  |  |  |  |  |
| 27 | \% change in budget | 1.00 |  | ) |  | Note that we have been unable to obtain a satisfactory outcome model for trauma \& injuries and have assumed a zero outcome elasticity. |  |  |  |  |  |  |  |  |  |  |  |  |
| 28 | proportionate change | 0.01 | $\begin{aligned} & 0.01 \\ & 6 \\ & \hline \end{aligned}$ |  |  | Note that, for expenditure in $2006 / 7$, the neonate category has been merged with maternity to obtain plausible outcome and expenditure models. |  |  |  |  |  |  |  |  |  |  |  |  |
| 29 | Change in budget | $£_{6} 678.96$ |  |  |  | Note that the adjustment for the coverage of the YLL data relative to the spend data uses deaths under age 75 in England in 2008. |  |  |  |  |  |  |  |  |  |  |  |  |

Table B8.23: table showing cost of life and life year estimates using expenditure data for 2006 and outcome data for 2006/7/8 (assumes average health gain for 12 other programmes) adjusted for the ICD10 coverage of the expenditure and outcome data


Note:
29 All 23 programme spend
$30 \%$ change in budget
proportionate chang
32 Change in budget

2006/7 2005/6
$£_{6}^{64,310}$
$\begin{array}{ll}1.00 & 1.00 \\ 0.01 & 0.01\end{array}$
$\begin{array}{ll}0.01 & 0.01 \\ £ 678.96 & £ 643.10\end{array}$

Note that the annual mortality figures reported in cells $\mathrm{G} 6 \& \mathrm{G} 7$ and $\mathrm{G} 15 \& \mathrm{G} 16$ are identical because we do not have mortality data for $2002 / 04$ yet. Note that we have been unable to obtain a satisfactory outcome model for trauma \& injuries and have assumed a zero outcome elasticity.
Note that, for expenditure in $2006 / 7$, the neonate category has been merged with maternity to obtain plausible outcome and expenditure models.
Note that the adjustment for the coverage of the YLL data relative to the spend data uses deaths under age 75 in England in 2008.

## B8.7 Adjusting the cost of life (year) estimates for Department of Health funded expenditure that is not undertaken by PCTs

PCT expenditure accounts for a large proportion of Department of Health expenditure but PCTs do not account for all of the Department's budget. In 2006/7 the Department of Health's gross expenditure totalled $£ 83.5 \mathrm{bn}$. Charges raised $£ 3.4$ bn so net expenditure totalled $£ 80.1 \mathrm{bn}$. Of this net expenditure, PCTs accounted for $£ 67.3$ bn (that is, $84 \%$ ) and various other bodies accounted for the remaining $£ 12.8 \mathrm{bn}$. A breakdown of this gross and net expenditure by major body is shown in Table B8.24.

Table B8.24: table showing department of Health funded expenditure by major bodies, 2006/7

| Body | Gross spend £ billion | Income £ billion | Net spend £ billion |
| :---: | :---: | :---: | :---: |
| PCTs | 69.8 | 2.5 | 67.3 |
| Strategic Health Authorities | 3.8 | 0.0 | 3.8 |
| Special Health Authorities* | 2.8 | 1.3 | 1.5 |
| Department of Health Own Costs (eg PSS grants, grants to LAs) | 7.1 | -0.4 | $7.5$ |
| Total Department of Health | 83.5 | 3.4 | $80.1$ |

*This includes, for example, NICE, the NHS Business Services Authority, the Information Centre, the NHS Litigation Authority, and the National Patient Safety Agency.

The Department of Health has allocated net non-PCT expenditure across the 23 PBCs and the impact of this allocation on total spend by PBC is shown in Table B8. 25 below. No geographic breakdown (e.g., by PCT) of this expenditure is available.

Of the additional $£ 12 \mathrm{bn}$ of net expenditure, $£ 11,2 \mathrm{bn}(93 \%)$ has been allocated to PBC 23 . This largely reflects: (a) the allocation of almost all Strategic Health Authority expenditure to either PBC23B ('other: SHAs including workforce development committees') or PBC23X ('other: miscellaneous'), and (b) the allocation of almost two-thirds of Department of Health expenditure to PBC23X ('other: miscellaneous').

The remaining $£ 0.8 \mathrm{bn}$ of additional net expenditure is spread across all PBCs according to various allocation rules. For example, the majority of expenditure on Special Health Authorities is apportioned across programme categories on the basis of the PCT and SHA expenditure breakdown. The exception is the NHS Business Services Authority expenditure which is apportioned on the basis of Primary Care prescribing expenditure splits. Although this approach avoids allocating expenditure to the 'Other: Miscellaneous' category, this allocation of expenditure does not necessarily reflect actual expenditure. For example, NHS Eitigation Authority expenditure may not be incurred in the same areas as overall PCT expenditure [38].

Table B8.25: table showing net PCT and other Department of Health funded expenditure by PBC, 2006/7


Note: the figures in Tables B8.24 and B8.25 draw on yarious sources (e.g., Department of Health resource accounts and programme budgeting returns) and may (a) disagree slightly and (b) create some unusual results (e.g., the aggregate PCT figure for dental problems exceeds the all England level[38]).

It is clear that most of the non-PCT expenditure is not specific to any disease area and that, to avoid putting all of it into a residual category, the Department has identified what are reasonable but largely arbitrary rules to spread what is a relatively small proportion of this non-PCT expenditure across all PBCs.

The cost of a life (year) estimates presented above are based on the impact of a $1 \%$ exogenous change in total net PCT spend. All of our outcome and expenditure models have been estimated using net PCT expenditure, and all of our elasticities relate to this expenditure. Implicitly we assume that any budgetary shock only affects PCT funding and that it leaves non-PCT funding unchanged.

Suppose instead we assume a $1 \%$ exogenous change in the Departmental budget. How might this budgetary shock be split between PCT and non-PCT expenditure? There are two obvious options to consider. We could assume either: (a) that all of this change is applied to PCT budgets and that there is no change in the non-PCT budget (as we do implicitly at the moment) or (b) that the budgetary shock affects both PCT and non-PCT budgets.

If the non-PCT budget is wholly unresponsive to the exogenous shock then our cost of a life year estimates will be unchanged because this expenditure category attracts none of the budgetary change (although this expenditure will clearly contribute to a measure of average productivity).
If the non-PCT budget is to some degree responsive to the exogenous shock then it will affect our cost of a life (year) estimates. To calculate the size of this impact we would need to know:
(a) how responsive the non-PCT budget is to a total Departmental budgetary shock;
(b) how the responsive part of the non-PCT budget is allocated across PBCs;
and
(c) the size of the health effects associated with changes in the non-PCT budget at PBC level.

We have no evidence on how responsive the non-PCT budget is likely to be to a total budgetary shock. However, from Table B8.25 and the discussion about the rather arbitrary (but understandable) rules employed by the Department to allocate non-PCT expenditure to PBCs, it would seem reasonable to assume that any change in the non-PCT budget should all be allocated to PBC23. This 'solves' the problem of identifying the health gains associated with this change in the non-PCT budget because, in our cost of a life year calculations, we assume that expenditure in this category attracts no health gains.

Thus although we have no evidence on how responsive the non-PCT budget is likely to be to a total budgetary shock, we can present two scenarios. In the first scenario, the non-PCT budget is wholly unresponsive to a budgetary shock and any budgetary change is fully implemented via PCT expenditure. In this case, there is no impact on the cost of a life year.

In the second scenario, one might assume that the non-PCT budget is as responsive to Departmental budgetary changes as is the PCT budget. In this case a $1 \%$ change in the Departmental budget is translated into a $1 \%$ change in both the total PCT and total non-PCT budgets, and this will increase the cost of a life year by $17.7 \%$ for $2006 / 7$, that is, from $£ 22,565$ to $£ 26,553$. This percentage increase is, of course, the same figure as total non-PCT expenditure expressed as a percentage of PCT expenditure. This is because all of the additional non-PCT expenditure is allocated to PBC23 and the assumption is that all expenditure in this category offers no health gain.

We have no information on how any Departmental budgetary shock is likely to be split between PCT and non-PCTs budgets. Our cost of a life year estimates implicitly assume that the non-PCT budget is wholly unresponsive to any budgetary shock. This is clearly a possibility. Alternatively, one might assume that the non-PCT budget is as responsive to a Departmental budgetary shock as is the PCT budget. If this was the case then it would add $17.7 \%$ to our cost of a life year estimate for 2006/7. However, in the absence of any information about the responsiveness of the non-PCT budget, it is difficult to come to any firm conclusion about the impact of non-PCT expenditure on our cost of a life year estimates. We therefore persevere with the assumption that the non-PCT budget is wholly unresponsive to Departmental budgetary shocks.

## B8.8 Application of method to other non-mortality based outcome indicators

Not all health care expenditure will be directed towards the reduction of mortality but it is relatively easy to envisage how our methods might be applied to other, non-mortality based, outcome indicators. To illustrate how our approach might be applied to other such indicators we note that PROMs (health gain) data for various operations is available from the HES online website. For each PCT this data set reports the average health gain for those survey respondents who have had a specific operation (e.g., for hip replacement, for knee replacement, for varicose veins, and for groin hernia) over the survey period.

As a starting point, and to illustrate the principles involved, we focus on hip and knee replacements. As our outcome indicator for these procedures, we calculate

$$
\text { [(average health gain per hip operation*number of hip operations) }+
$$

(average health gain per knee operation*number of knee operations)]
/total PCT population
for each PCT (this ignores age standardisation). This health gain measure is broadly comparable with our usual mortality measure, which is a 'years of life lost' rate per 10,000 of population (again, ignoring age standardisation).

Ideally the expenditure, number of operations, and PROMs data should all relate to the same time period but here the PROMs data covers operations undertaken between April 2009 and October 2010 yet the expenditure and number of operations data relate to $2006 / 7$. Implicitly we are assuming that the average
gain per operation in 2006/7 is the same as over the PROMs survey period (although this is not particularly important as we are only illustrating principles here).

Unfortunately, the Department of Health does not report the number of patients undergoing an eligible operation by commissioner (PCT) so we use the HES dataset for 2006/7 to obtain this information. Eligible hip and knee operations are defined in Annex 1 of the 'Guide to PROMs methodology' (on the HES website) and we use these definitions (of eligible operation codes) to obtain a count of eligible hip and knee finished consultant episodes (FCEs) by PCT for 2006/7.

With data for both the average health gain per operation and the number of operations, we are now in a position to calculate 'the health gain per head of population' for hip and knee replacements as defined above. We can then use this as an outcome indicator for expenditure in the 'problems of the musculoskeletal system' programme (i.e., PBC15) because the vast majority of hip and knee replacements fare for osteoarthritis and this diagnosis is included in PBC15.

Table B8.26 reports the estimated outcome equation for PBC15 using the PROMs based outcome indicator. The result is intuitively plausible. More expenditure boosts the health gain but, for a given spend, more need reduces the gain. Of course we should remember that the health gain data relates to operations undertaken between April 2009 and October 2010 yet the expenditare and number of operations data (FCEs) relate to 2006/7. However, one might assume that the gain associated with each operation in 2009/10 is the same as the gain associated with each operation in 2006/7.

The diagnostic statistics suggest that expenditure is endogenous and that the instruments are valid. They also suggest that the instruments are relevant and there is no evidence that the instruments are weak. The Pesaran-Taylor test suggests that there is no evidence of model mis-specification.

This brief example illustrates the principles involved in extending our modelling approach beyond those programmes with a mortality indicator.

Table B8.26: table showing outcome model for the trauma and injuries programme, 2006/7


Notes: (i) the dependent variable is the health gain per head of population associated with eligible hip and knee operations undertaken during 2006/7; (ii) that there are only 143 observations and not the usual 152 because, for the other nine PCTs, there are fewer than 30 completed PROMs questionnaires on which to compute the average health gain and, as a result of such a low number of respondents, these PCTs have been dropped from the sample; (ii) the first-stage regression includes the IMD2007 (coefficient $=-0.439$, standard error= 0.144 ) and the proportion of residents providing unpaid care (coefficient $=0.219$, standard error=0.367); and (iii) robust standard errors in brackets, ${ }^{* * *} \mathrm{p}<0.01, * * \mathrm{p}<0.05,{ }^{*} \mathrm{p}<0.1$

## B8.9 Comparing outcome models for 'high' spending and 'low' spending PCTs

As we have already noted, not all PCTs spend the same amount in each programme of care. Even after allowing for differences in local circumstances (such as input prices and need), some PCTs spend more than others, and it is this variation in expenditure that facilitates the estimation of our outcome and expenditure models.

Figure B8.2 illustrates the familiar health gain production function; as expenditure increases so too does health output but it increases at a diminishing rate. If all PCTs face the same production function (having controlled for input prices and need), and all PCTs are wholly efficient, then we would expect those PCTs that spend more (e.g., at point B) to experience a lower outcome elasticity than those that spend less (e.g., at point A) simply because they are further along the production function and are experiencing greater diminishing marginal returns.


Health care spend
in programme $\left(\ell_{\mathrm{c}}\right)$
Figure B8.2: graph showing Health gain production function
To test this hypothesis we used the expenditure model for each of the big four programmes to divide the 152 PCTs into two groups: those whose predicted spend is greater than the average predicted spend in that programme (ceteris paribus), and those whose predicted spend is smaller than the average predicted spend (ceteris paribus). We then re-estimated our outcome model for each of these two groups of PCTs and the results of this re-estimation are shown in Table B8.27. ${ }^{29}$

The first column in Table B8.27 presents the IV regression results for the outcome model for all PCTs; the second column reports the results for the 'high spend' PCTs; and the third column reports the results for the 'low spend' PCTs. For all four programmes, the coefficient on the expenditure variable is larger (in an absolute sense) for the 'high spend' PCTs than for the (low spend' PCTs. This result contradicts our hypothesis that 'high spenders' will have a lower elasticity than 'low spenders'.

However, if we drop the assumption that all PCTs are equally efficient - so that that some lie within the frontier defined by the production function - then it is clearly possible for 'high' spending PCTs to experience a larger outcome elasticity than a 'low' spending one. And, of course, it is rather difficult to defend the assumption that all PCTs are equally efficient.

We can use the outcome elasticities reported in Table B8.27 to calculate the cost of a life year for 'high' and 'low' spenders in each of the big four programmes. These calculations are shown in Table B8.28. ${ }^{30}$ As is to be anticipated, they reveal that the cost of a life year is much smaller in 'high' spend' PCTs than it is in 'low spend' PCTs. For example, the cost of a life year in the cancer programme is $£ 16,383$ across all PCTs but for 'high spenders' it is much less than this ( $£ 11,350$ ) and for 'low spenders' it is much greater than this $(£, 76,620)$. Presumably 'high spending' PCTs are high spenders because the cost of a life year is relatively low and additional health gains in a particular programme can be had relatively cheaply. Similarly, 'low spending' PCTs are low spenders because the cost of a life year is relatively high and additional health gains are relatively expensive.

[^59]Table B8.27: table showing re-estimating the 2006/7 outcome model for 'high' spending and 'low' spending PCTs


Table B8.28: table showing calculation of the cost of a life year for the big four programmes in 2006/7 by type of PCT: 'high spenders' and 'low spenders'


Split PCTs according to whether they are 'high spender' ( $\mathrm{n}=76$ ) or 'low spenders' ( $\mathrm{n}=76$ )

| 1 | Cancers | All | 4,122 | 41.22 | 0.342 | 2,207,021 | 735,674 | 2,516 | 16,383 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 2 | Cancers | High spend | 2,080 | 20.8 | 0.488 | $1,126,580$ | 375,527 | 1,833 | 11,350 |
| 3 | Cancers | Low spend | 2,042 | 20.42 | 0.074 | 1,080,442 | 360,147 | 267 | 76,620 |
| 4 | Circulatory problems | All | 6,161 | 61.61 | 43 | 1,361,634 | 453,878 | 6,509 | 9,466 |
| 5 | Circulatory problems | High spend | 3,148 | 31.48 |  | 695,890 | 231,963 | 3,254 | 9,673 |
| 6 | Circulatory problems | Low spend | 3,012 |  | 99 | 665,744 | 221,915 | 2,208 | 13,641 |
| 7 | Respiratory problems | All | 3,285 | 32.85 | 2.622 | 324,223 | 108,074 | 2,834 | 11,593 |
| 8 | Respiratory problems | High spend | 1,645 | 16.45 | 3.697 | 174,639 | 58,213 | 2,152 | 7,644 |
| 9 | Respiratory problems | Low spend | ,640 | 16 | 1.461 | 149,584 | 49,861 | 728 | 22,513 |
| 10 | Gastro- problems | All | 3,700 | 37 | 1.536 | 345,908 | 115,303 | 1,771 | 20,892 |
| 11 | Gastro- problems | High spend | $1,868$ | 18.68 | 1.471 | 190,231 | 63,410 | 933 | 20,026 |
| 12 | Gastro- problems | Low spend | 1,832 | 18.32 | 0.819 | 155,676 | 51,892 | 425 | 43,106 |

[^60]'low spending' PCTs are those whose predicted spend per person is less than the average predicted spend per person (ceteris paribus).

## B8.10 Comparing outcome models for over target and under target PCTs

The Department of Health has a well-developed resource allocation formula that determines the size of each PCTs 'target' budget given local conditions (such as population size and the need for health care). Every few years an improved resource allocation formula is developed and this generates a new 'target' budget for each PCT. The new target might be quite different from the old target and the immediate implementation of the new formula might lead to a large change in the budget for some PCTs. To avoid the difficulties that sudden large budgetary changes might bring, actual annual financial allocations are gradually moved towards the latest target budget. This means that in any year some PCTs receive an actual allocation which is greater than their target allocation, and that others receive an actual allocation which is less than their target allocation.

To examine whether being over or under the target allocation has any impact on the results, we split the 152 PCTs into two groups: those that received a budget over their target allocation in 2006/7 and those that received a budget under their target allocation in 2006/7. The outcome elasticities from the) estimation of these models are shown in column E of Table B8.29, and these elasticities are used to calculate the cost of a life year for each of these two groups of PCTs for each of the big four programmes (see column I).

The results are consistent for each programme: PCTs whose budget is beyond their target allocation record a smaller outcome elasticity and a larger cost of a life year than PCTs whose budget is less than their target allocation. For example, in the cancer programme and across allPCTs the outcome elasticity is -0.342 and the cost of a life year is $£ 16,383$ (unadjusted for the ICD10 coverage of the mortality data). For PCTs with a budget that exceeds their target allocation, the outcome elasticity is smaller ( -0.179 ) and the cost of a life year is larger $(£ 32,365)$ than for all PCTs combined. However, for PCTs with a budget that falls short of their target allocation, the outcome elasticity is larger $(-0.476)$ and the cost of a life year is smaller $(£, 11,502)$ than for all PCTs combined.

One explanation for this result is that PCTs whose budget is beyond their target allocation are under less financial pressure than other PCTs, and that one consequence of this is that there is less pressure on them to behave in the most efficient manner possible. There is some evidence in the literature to support the hypothesis that the degree of PCT inefficiency is positively related to the amount by which a PCTs is over its target allocation. [39]

If we also re-estimate the expenditure models for both groups of PCTs we can calculate the cost of a life year for the big four programmes combined. The relevant expenditure elasticities are shown in column D of Table B8.30. These expenditure elasticities are far larger for under target PCTs than they are for over target PCTs. One reason for this might be that the big four programmes are priority ('hard') programmes. Over target PCTs are able to devote sufficient resources to the big four so that any additional budget is directed towards other ('softer') programmes which are less well funded than the priority programmes. In contrast, under target PCTs are struggling to devote sufficient resources to the priority programmes so that, when further funding does become available, this is directed towards the priority programmes.

These expenditure and outcome elasticities in Table B8.30 can be used to calculate the cost of a life year for the big four programmes combined (adjusted for the ICD10 coverage of the mortality data). This cost is:

- $£ 10,604$ for all PCTs combined
- $£ 14,083$ for PCTs whose budget is beyond its target allocation
- $£ 8,441$ for PCTs whose budget falls short of its target allocation.

Again, the cost of a life year is much smaller for PCTs whose budget falls short of its target allocation.

Table B8.29: table showing calculation of the cost of a life year for the big four programmes by type of PCT: over target and under target allocations
Type of PCT
C
D
C

|  |  |
| :--- | :--- |
| Spend $(£ \mathrm{~m})$ | $1 \%$ of spend <br> FY2006/7 |
| $(£ \mathrm{~m})$ |  |
| FY2006/7 |  |

E
F
G
H
A
PBC description
B

| Outcome |  |
| :--- | :--- |
| elasticity | Total life years |
| (without | lost, $<75$ years, |
| negative sign) | $2006 / 08$ |

Annual
average life
years lost
$(=\mathrm{F} / 3$

Change in annual life years ost associated with $1 \%$ increase in spend
$\left(=E^{*} G\right) / 100$

Cost ( $£$ ) per life year gained gained
$(=\mathrm{D} / \mathrm{H})$


Split PCTs according to whether they are over target allocation ( $\mathrm{n}=67$ ) or under target allocation ( $\mathrm{n}=85$ )


[^61]Table B8.30: table showing cost of life and life year estimates using spend data for 2006 and outcome data for 2006/08 for the big four PBCs for:
(i) all PCTs; (ii) PCTs that are over target; and (iii) PCTs that are under target. A B C $\quad$ D E
F G
H
I J

| J | K | L |
| :---: | :---: | :---: |
| $\begin{aligned} & =0.01 * \mathrm{D}^{*} \\ & \mathrm{G}^{*} \mathrm{I} / \mathrm{H} \end{aligned}$ |  | $=\mathrm{E} / \mathrm{J}$ |
| Change in annual mortality, adj for coverage |  | Cost per li gained adj for coverag ( $\AA_{\text {. }}$ |


|  | PBC description | $\begin{aligned} & (£ \mathrm{~m}) \\ & 2006 / 7 \\ & \hline \end{aligned}$ | Spend elasticity | $\begin{aligned} & \text { spend } \\ & (£ \mathrm{~m}) \\ & \hline \end{aligned}$ | $\begin{aligned} & \text { years, } \\ & 2006 / 08 \\ & \hline \end{aligned}$ | spend data | negative <br> sign) | adj for coverage | for coverage $\left(\epsilon_{0}\right)$ | $\begin{aligned} & \text { <75years, } \\ & 2006 / 08 \\ & \hline \end{aligned}$ | $\begin{aligned} & \text { years } \\ & \text { lost } \\ & \hline \end{aligned}$ | spend data | for coverage | life year <br> gained ( $\mathrm{f}_{\mathrm{t}}$ ) | coverage $\left(\ell_{0}\right)$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | All PCTs together |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| 1 | Cancer | £4,122 | 0.465 | £19.17 | 61,961 | 0.984 | 0.342 | 100.14 | $£ 191,407$ | 2,207,021 | 1,170 | 0.984 | 1,189 | £16,383 | £16,121 |
| 2 | Circulatory problems | $£_{6} 6,161$ | 0.540 | £33.27 | 41,106 | 0.992 | 1.434 | 320.88 | f103,683 | 1,361,634 | 3,515 | 1.000 | 3,543 | $£^{¢} 9,466$ | $£^{9}, 390$ |
| 3 | Respiratory problems | £3,285 | 0.679 | $£ 22.31$ | 11,574 | 0.773 | 2.622 | 266.57 | ¢83,676 | 324,223 | 1,924 | 0.773 | 2,489 | £11,593 | £8,961 |
| 4 | Gastro-intestinal problems | ¢ 3,700 | 0.446 | £16.50 | 6,160 | 0.571 | 1.536 | 73.90 | ¢ 2223,228 | 345,908 | 790 | 0.650 | 1,383 | £20,892 | £11,929 |
| 5 | Big four programmes summary: Spend 2006 \& mortality 2006/8 | £17,268 |  | $£ 91.24$ |  |  |  |  | £119,823 |  | 7,399 |  | 8,604 | $£ 12,333$ | £10,604 |
|  | For over target PCTs only ( $n=67$ ) |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| 6 | Cancer | £1,733 | 0.193 | $£ 3.34$ | 24,918 | 0.984 | 0.179 | 8.75 | £382,320 | 897,403 | 103 | 0.984 | 105 | £32,365 | £31,847 |
| 7 | Circulatory problems | $£ 2,587$ | 0.150 | $£ 3.88$ | 16,346 | 0.992 | 1,115 | 27.56 | $£ 140,806$ | 544,326 | 303 | 1.000 | 303 | £12,787 | £12,685 |
| 8 | Respiratory problems | £1,357 | 0.326 | $£ 4.42$ | 4,588 | 0.773 | 2.637 | 51.02 | £86,701 | 127,810 | 366 | 0.773 | 474 | £12,079 | $£^{9,337}$ |
| 9 | Gastro-intestinal problems | £1,566 | 0.090 | £1.41 | 2,525 | 0.571 | 0.569 | 2.26 | £622,378 | 142,281 | 24 | 0.650 | 43 | £ 58,030 | £33,135 |
| 10 | Big four programmes summary: Spend 2006 \& mortality 2006/8 | $£^{7,243}$ |  | $£ 13.06$ |  |  |  | 89.60 | £145,748 |  | 797 |  | 927 | $£ 16,378$ | $£ 14,083$ |
|  | For under target PCTs only ( $n=85$ ) |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| 11 | Cancer | $£_{2,390}$ | 0.785 | £18.76 | 37,043 | . 84 | 0.476 | 140.67 | $£ 133,377$ | 1,309,618 | 1,631 | 0.984 | 1,658 | £11,502 | £11,318 |
| 12 | Circulatory problems | £3,574 | 0.748 | $£ 26.73$ | 24,760 | 0.992 | 1.947 | 363.50 |  | 817,308 | 3,968 | 1.000 | 4,000 | £6,738 | £6,684 |
| 13 | Respiratory problems | $£_{1,982}$ | 1.035 | $£ 20.51$ | 6,986 | 0.773 | 2.674 | 250.12 | £82,015 | 196,413 | 1,812 | 0.773 | 2,344 | £11,321 | £ 8,751 |
| 14 | Gastro-intestinal problems | £2,134 | 0.592 | £.12.63 | 3,602 | 0.571 | 1.869 | 69.80 | £181,000 | 203,626 | 751 | 0.650 | 1,315 | £16,822 | $£^{9} 9605$ |
| 15 | Big four programmes summary: Spend 2006 \& mortality 2006/8 | £10,080 |  | $£ .78 .6$ |  |  |  | 824.09 | £ 95,429 |  | 8,162 |  | 9,317 | $£_{0,635}$ | $£ 8,441$ |
|  | Note: | 2006/7 | 2005/6 |  |  |  |  |  |  |  |  |  |  |  |  |
|  | All 23 programme spend | £67,896 | £64,310 |  |  |  |  |  |  |  |  |  |  |  |  |
|  | \% change in budget | 1.00 | 1.00 (L) |  |  |  |  |  |  |  |  |  |  |  |  |
|  | proportionate change | 0.01 | 0.01 |  |  |  |  |  |  |  |  |  |  |  |  |
|  | Change in budget | £678.96 | ¢643.10 |  |  |  |  |  |  |  |  |  |  |  |  |

[^62]
## B8.11 The correlation between the outcome and expenditure elasticities

To investigate the correlation between the outcome and expenditure elasticities for any given programme, a random sample (with replacement) of 152 PCTs was drawn from the population of 152 PCTs. In this random drawing, some of the original observations will appear once, some more than once, and some not at all. Using this re-sampled dataset, outcome and expenditure models for the selected programme were estimated (as per Table B8.19) and the outcome and expenditure elasticities saved. This step was repeated 500 times and the correlation coefficient for the outcome and expenditure elasticities was calculated. Table B8.31 shows these correlation coefficients for each of the big four programmes.

Table B8.31: table showing correlation coefficient for the outcome and expenditure elasticities

| Programme of care | Correlation coefficient between the <br> outcome and expenditure elasticities |
| :--- | :--- |
| Cancers and tumours | 0.1542 |
| Circulatory disease | 0.1968 |
| Respiratory problems | 0.0368 |
| Gastro-intestinal problems | 0.0611 |

Note: the estimated elasticities are from unweighted IV regressions because there is no weight option with the bootstrap command in Stata. However, weighting makes little difference to our IV results. For example, in the cancer outcome model the coefficient on spend is -0.342 with weighting applied but it is
-0.299 without any weighting applied. For the cancer spend model the coefficient on budget is 0.465 with weighting but 0.520 without weighting.

## B8.12 Summary and conclusion

In this section we have undertaken several tasks. First, we have identified and resolved several estimation issues relating to the appropriate measure of need, the appropriate price index to be used to adjust PCT expenditure for local variations in input prices, and the fact that PCTs vary in size.

Second, we have derived plausible outcome and expenditure models for ten care programmes using expenditure data for 2006/7 and mortality data for 2004/5/6. The cost of a life year across these ten programmes is $£ 20,893$ (it was $£ 21,256$ using expenditure data for 2005/6).

Third, we have re-estimated the outcome and expenditure models using the same expenditure data but replacing the mortality data for $2004 / 5 / 6$ with data for $2006 / 7 / 8$. The advantage of this is that it assumes that the health benefits associated with expenditure occur either in the same period as the expenditure or in the next two periods. This is an improvement on past practice where data constraints forced researchers to relate expenditure to the current and two previous periods. This re-estimation increased the cost of a life year across all ten programmes by $14 \%$, from $£ 20,893$ to $£ 23,780$.

Fourth, we have adjusted the cost of a life year calculations for the mismatch in the ICD10 coverage of the expenditure and mortality data. This reduces the cost of a life year for 2006/(7) for those 10 PBCs with a mortality indicator by $14 \%$, from $£ 23,780$ to $£ 19,965$ (a decrease of $16 \% 0$ ).

Fifth, we have noted that our cost of a life year estimates are based on the assumption that any Departmental budgetary change falls entirely on PCTs. Although PCTs account for most of the Department of Health's budget, non-PCTs still accounted for $15 \%$ of the budget in 2006/7. Because we have no information on how any budgetary change would be split between PCTs and non-PCTs, our estimates implicitly assume that any Departmental budgetary change falls entirely on PCTs. If, on the other hand, the non-PCT budget is responsive to changes in the Department's budget then our cost of a life year estimates will be slightly too low (for exampre, if the non-PCT budget is as responsive as the PCT budget, then our cost of a life year estimate for $2006 / 7$ will be increased by $17.7 \%$ (that is, from $£ 22,565$ to $£ 26,553$ ).

We have also illustrated how our modelling framework can be applied to other non-mortality based outcome indicators, and the cost of fife year estimates that are obtained if PCTs are split into different groups (e.g., those that under and those that are over their target budget allocations). In the next section we examine the impact of relaxing the instrument validity restriction on our results.

## B9. The sensitivity of the outcome elasticity to the validity of the instrument exclusion restrictions

## B9.1 Introduction

One of the crucial elements in the calculation of the cost of a life year for any programme of care is the coefficient on the expenditure variable in the outcome equation. This coefficient indicates the amount by which mortality changes following a (small) change in expenditure in that care programme. It is to be expected that this coefficient will have a negative sign so that as expenditure increases, for example, mortality will decline. If this coefficient is small (in an absolute sense) then it implies that any change in expenditure will have little effect on mortality and so the cost of a life year will be relatively large (ceteris paribus). Alternatively, if this coefficient is large (in an absolute sense) then any change in expenditure will have a large effect on mortality and so the cost of a life year will be relatively small (ceteris paribus) For this reason it is important that we correctly identify the magnitude of this 'treatment parameter'.

Our basic outcome model for each programme of care is

$$
\begin{equation*}
y=\alpha+\beta 1 x+\beta 2 n+\epsilon \tag{9.1}
\end{equation*}
$$

where y is mortality, x is expenditure, and n is a measure of the need for health care (with all variables relating to a particular programme of care). We are particularly interestedin the size of the coefficient on expenditure ( $\beta 1$ ). We do not use OLS to estimate this outcome model because expenditure $(\mathrm{x})$ is endogenous and, in the presence of an endogenous regressor, OLS will provide both a biased and an inconsistent estimator of $\beta 1$. Instead, we use instrumental variable (IV) techniques. Unlike OLS, IV will provide a consistent estimator of $\beta 1$ and, although in finite samples the IV estimator will be biased, the belief is that (providing certain assumptions are met) this bias will be less than that associated with OLS.

IV estimation involves finding variables (instruments) that are good predictors of expenditure ( x ) but which are appropriately excluded from the equation of interest (that is, equation 1 ). The assumption is that the instruments impact upon mortality (y) through their impact on expenditure ( x ) only, and that they do not have a direct effect on mortality (y). If, on the other hand, an instrument reflects unobserved factors that affect both expenditure and mortality directly, the use of this instrument will lead to a biased and inconsistent estimate of the coefficient on expenditure. Such an instrument is said to be 'invalid' because it belongs in the equation of interest in its own right.

In our outcome models we typically employ two instruments (call these z1 and z2) for expenditure. IV assumes that these instruments do not belong in the outcome equation (9.1). In other words, IV assumes that the coefficients $\gamma 1$ and $\gamma 2$ in the outcome model

$$
\begin{equation*}
y=\alpha+\beta 1 x+\beta 2 n+\gamma 1 z 1+\gamma 2 z 2+\epsilon \tag{9.2}
\end{equation*}
$$

are identicallyzero. Such exclusion restrictions can be debatable and researchers who employ IV techniques often devote considerable effort towards convincing the reader that their assumed exclusion restrictions are a good approximation [35, 36]. These efforts usually take two forms: first, researchers oftenoffer a strong theoretical economic argument why their instruments do not belong in the equation of interest; and, second, statistical tests for the validity of the exclusion restrictions (Sargan 2SLS, Hansen I-test GMM) are routinely reported as part of the results for any study that employs IV techniques.

It is difficult for us to identify clear theoretical reasons why our instruments (such as the proportion of lone pensioner households, the provision of unpaid care, and an index of multiple deprivation) do not belong in the equation of interest (that is, that they will not directly affect mortality). Of necessity, therefore, we must be guided by the available statistical tests for the validity of the exclusion restrictions. However, although our outcome models 'pass' the relevant statistical test, some commentators have argued that the Sargan/Hansen test may have weak power and may fail to reject the null hypothesis of instrument validity even when an exclusion restriction is not valid. Given our reliance on this test, it is important that we examine the circumstances in which this test may have weak power.

## B9.2 The Sargan-Hansen test of overidentifying restrictions: when will it have low power?

As we have one endogenous variable (expenditure) in our outcome model and more than one instrument available for health care expenditure, our estimating equation is said to be 'overidentified'. With more instruments than endogenous regressors, there is more than one way of using the instruments to estimate the parameter $\beta 1$ on the endogenous variable. The Sargan-Hansen J test of overidentifying (OID) restrictions calculates whether different instruments or different combinations of instruments generate significantly different values for the coefficient $(\beta 1)$ on the endogenous variable in the equation of interest. If significant differences are detected then the test will reject the null hypothesis that all instruments are jointly valid. Of course, the test does not reveal which instrument(s) is(are) invalid; instead, the test uses the fact that different instruments (or combinations thereof) generate different estimates of $\beta 1$ to infer that something is wrong with the set of instruments. Even if all of the instruments are invalid in the sense that they are all correlated with the error term in the equation of interest (and thus belong in the outcome equation as regressors), the test can detect this failure if the induced biases in the estimates of $\beta 1$ differ across instruments. This 'vector-of-contrasts' interpretation of the Sargan-Hansen test makes it clear when the $J$ test will lack power to reject the null hypothesis when it is false. The J statistic will be small when the null hypothesis of valid instruments is correct; but it will also be small if the biases induced in $\widehat{\beta 1}$ by invalid instruments all coincide (i.e., the instruments all identify the same wrong parameter). [37]

Most of our estimated models involve the use of two instruments. Kovandic, Schaffer and Kleck[37] point that when there are only two instruments '...the J test statistic is numerically identical to a Hausman test statistic that contrasts the estimator using both instruments with an estimator using just one instrument. The intuition [behind this result] is...straightforward: a Hausman test will reject the null hypothesis that the two estimators being contrasted are both consistent so long as the estimators converge to different values. It is not a requirement for one of the two estimators to be consistent for the Hausman test (and therefore the J test) to have power to reject the null.' One implication of this observation is that misspecification, in the conditional mean of the model, need not necessarily cause the Hansen-Sargan test to fail.

Kovandic, Schaffer and Kleck point out that these arguments suggest '...that the more unrelated the instruments are to each other, the more credible is a failure to reject the null that the instruments are exogenous, since a failure to reject would require that two unrelated instruments generate the same asymptotic bias in $\widehat{\beta 1}$ ' (p19).

Schaffer [40] argues that '[d]ifferent] sets of instruments are likely to have more or less power depending on where they come from. If all the instruments are minor variations on the same variable -- e.g., they are the same variable but lagged a few different periods -- then they are all likely to identify the same psuedoparameter. The critique of low power is going to be fairly convincing here.'

On the other hand, if the instruments are very different and, even better, there are ex ante reasons for thinking that if they are invalid, they are invalid in 'different ways', the J test will have more power. For example, suppose two instruments are available and it is thought that, if one is invalid, it will bias the estimated parameter upwards but, if the other instrument is invalid, it will bias the estimated parameter downards. If the Hansen-Sargan J test fails to reject in this setting, it is a convincing result.[40]

In this study we typically use any two from three available instruments when estimating our outcome equations. These three instruments are:
(a) the proportion of households that are lone pensioner households (from the 2001 Census);
(b) the proportion of residents providing more than one hour of unpaid care per week (from the 2001 Census); and
(c) the index of multiple deprivation (IMD) 2007.

For the Hansen-Sargan J test to have low power the use of any two of these instruments should generate the same asymptotic bias in $\widehat{\beta 1}$. However, it is far from obvious that this will be the case, particularly given that our outcome equation already includes a measure of the need for health care.

Nevertheless, we must admit that it is possible that our instruments are correlated with both expenditure and some unobserved factor which is directly influencing the mortality rate, and that the induced bias in $\widehat{\beta 1}$ is the same for both instruments. In the next section we therefore examine the sensitivity of the estimated outcome elasticity to the validity of the exclusion restrictions.

In summary:

- the Sargan-Hansen J test of overidentifying restrictions calculates whether different instruments or different combinations of instruments generate significantly different values for the coefficient $(\beta 1)$ on the endogenous variable in the equation of interest. If significant differencesare detected then the test will reject the null hypothesis that all instruments are jointly valid.
- the J test uses the fact that different instruments (or combinations thereof) generate different estimates of $\beta 1$ to infer that something is wrong with the chosen set of instruments.
- even if all of the instruments are invalid in the sense that they are all correlated with the error term in the equation of interest, the test can detect this failure if the induced biases in the estimates of $\beta 1$ differ across instruments.
- this 'vector-of-contrasts' interpretation of the SarganHansen test also makes it clear when the J test will lack power to reject the null hypothesis when it is false. The J statistic will be small when the null hypothesis of valid instruments is correct; but it will also be small if the biases induced in $\widehat{\beta 1}$ by invalid instruments all coincide (i.e., the instruments all identify the same wrong parameter).
- most of our outcome models use two from the following three instruments: lone pensioners, multiple deprivation and unpaid carers. Thus our Hansen-Sargan test statistics are likely to have low power if our selected pair of instruments are both inducing the same bias in $\widehat{\beta 1}$. It is far from obvious that these instruments will induce the same bias in the coefficient on expenditure.
- however, in case our instruments are imparting the same bias to $\widehat{\beta 1}$, the next section examines the sensitivity of the estimated outcome elasticity to the validity of the exclusion restrictions.


## B9.3 The value selection problem

Given that the Hansen-Sargan J test might be unable to detect the presence of invalid instruments in some (rather restrictive) circumstances, several studies have suggested that researchers using IV techniques should subject the estimated coefficient on the endogenous variable to a sensitivity analysis (e.g., Conley, Hansen and Rossi, 2012; Small, 2007). Recall that IV estimation involves the assumption that the instruments do not belong in the equation of interest (i.e., in the outcome equation). In other words, the assumption is that the coefficients $\gamma 1$ and $\gamma 2$ on the instruments z 1 and z 2 in the outcome model

$$
\begin{equation*}
y=\alpha+\beta 1 x+\beta 2 n+\gamma 1 z 1+\gamma 2 z 2+\epsilon \tag{9.3}
\end{equation*}
$$

are identically zero (where y is mortality, x is expenditure, and n is a measure of the need for health care). One suggestion is that investigators should relax the assumption that $\gamma 1$ and $\gamma 2$ are identically zero and examine the impact of this relaxation on the estimated value for $\beta 1$. This proposal, however, raises the issue of which non-zero values should be imposed upon $\gamma 1$ and $\gamma 2$.

Proponents of this approach suggest that prior information about the extent of deviations from the exact exclusion restriction might be drawn from other research studies or from subject matter experts[35, 36]. In the present context, however, we have no prior beliefs about the likely values for, or even the signs on, $\gamma 1$ and $\gamma 2$.

As a starting point we re-estimated the outcome model for the 2006/7 cancer programme 420 times, assuming a uniform distribution between -1 and 1 for both $\gamma 1$ and $\gamma 2 .{ }^{31} 32$ Table B9.1 shows the estimated coefficients on expenditure $(\widehat{\beta 1})$ in our cancer outcome equation associated with the various pairs of values imposed upon $\gamma 1$ and $\gamma 2$. The coefficients in this table indicate that the outcome elasticity is rather sensitive to the precise values assigned to $\gamma 1$ and $\gamma 2$. However, in the absence of any guidance from other research studies or from subject matter experts, we require a method that will identify a plausible range of values for both $\gamma 1$ and $\gamma 2$, and which we can use as the basis for our sensitivity analysis.

[^63]Table B9.1: table showing the impact of weakening the exclusion restrictions on the instruments in the cancer outcome equation

| Coefficients on expenditure $(\widehat{\boldsymbol{\beta} 1})$ |  | Imposed coefficient on IMD variable ( $\boldsymbol{\gamma}_{\mathbf{2}}$ ) |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | -1 | -0.9 | -0.8 | -0.7 | -0.6 | -0.5 | -0.4 | -0.3 | -0.2 | -0.1 | 0.0 | 0.1 | 0.2 | 0.3 | 0.4 | (0.5) | 0.6 | 0.7 | 0.8 | 0.9 | 1.0 |
| 1.000.900.800.70 |  | -4.47 | -4.22 | -3.96 | -3.71 | -3.45 | -3.20 | -2.95 | $-2.70$ | -2.45 | $-2.20$ | -1.96 | -1.71 | -1.47 | -1.23 | -0.99 | -0.75 | -0.50 | -0.24 | 0.03 | 0.30 | 0.57 |
|  |  | -4.31 | -4.06 | -3.80 | -3.54 | -3.29 | -3.04 | -2.79 | $-2.53$ | -2.29 | -2.04 | -1.80 | -1.55 | -1.31 | -1.07 | -0.83 | -0.59 | -0.33 | -0.07 | 0.20 | 0.47 | 0.74 |
|  |  | -4.15 | -3.90 | -3.64 | -3.38 | -3.13 | $-2.88$ | -2.62 | $-2.37$ | -2.13 | -1.88 | -1.64 | -1.40 | -1.16 | -0.92 | -0.67 | -0.42 | -0.16 | 0.10 | 0.37 | 0.65 | 0.92 |
|  |  | -3.99 | -3.74 | -3.48 | -3.22 | -2.97 | -2.72 | -2.46 | -2.21 | -1.96 | -1.72 | -1.48 | -1.24 | -1.00 | -0.76 | -0.51 | -0.26 | 0.01 | 0.28 | 0.55 | 0.82 | 1.10 |
|  |  | -3.83 | -3.58 | -3.32 | -3.06 | -2.81 | $-2.55$ | $-2.30$ | -2.05 | -1.80 | -1.56 | -1.32 | -1.08 | -0.84 | -0.60 | -0.35 | -0.09 | 0.18 | 0.45 | 0.73 | 1.00 | 1.28 |
| Imposed | 0.50 | -3.67 | -3.41 | -3.16 | $-2.90$ | -2.65 | -2.39 | -2.14 | -1.89 | -1.64 | -1.40 | -1.16 | -0.92 | -0.68 | -0.44 | -0.18 | 0.08 | 0.35 | 0.63 | 0.91 | 1.18 | 1.45 |
| coefficient | 0.40 | -3.51 | -3.25 | -3.00 | -2.74 | -2.49 | $-2.23$ | -1.98 | -1.73 | -1.48 | -1.23 | -0.99 | -0.76 | -0.52 | -0.27 | -0.01 | 0.26 | 0.53 | 0.81 | 1.09 | 1.36 | 1.63 |
|  | 0.30 | -3.35 | -3.09 | -2.84 | -2.58 | -2.32 | -2.07 | -1.82 | -1.57 | -1.32 | -1.07 | -0.83 | -0.60 | -0.35 | -0.10 | 0.16 | 0.44 | 0.71 | 0.99 | 1.26 | 1.53 | 1.80 |
| on lone | 0.20 | -3.19 | -2.93 | -2.68 | -2.42 | -2.16 | -1.91 | -1.65 | -1.40 | -1.15 | -0.91 | -0.67 | -0.43 | -0.19 | 0.07 | 0.34 | 0.62 | 0.90 | 1.17 | 1.44 | 1.71 | 1.97 |
| pensioner | 0.10 | -3.03 | -2.77 | -2.51 | -2.26 | -2.00 | -1.75 | -1.49 | -1.24 | -0.99 | -0.75 | -0.51 | -0.27 | -0.02 | 0.25 | 0.52 | 0.80 | 1.07 | 1.34 | 1.61 | 1.88 | 2.14 |
| households | 0.00 | -2.87 | $-2.61$ | -2.35 | -2.10 | -1.84 | -1.58 | -1.33 | -1.07 | -0.83 | -0.58 | -0.34 | -0.10 | 0.15 | 0.43 | 0.70 | 0.98 | 1.25 | 1.52 | 1.78 | 2.04 | 2.31 |
| variable | -0.10 | -2.71 | -2.45 | -2.19 | -1.93 | -1.68 | -1.42 | -1.16 | -0.91 | -0.66 | -0.42 | -0.18 | 0.07 | 0.33 | 0.61 | 0.88 | 1.15 | 1.42 | 1.68 | 1.95 | 2.21 | 2.47 |
|  | -0.20 | $-2.55$ | -2.29 | -2.03 | -1.77 | -1.51 | -1.26 | -1.00 | -0.75 | -0.50 | -0.25 | -0.01 | 0.24 | 0.51 | 0.79 | 1.06 | 1.32 | 1.59 | 1.85 | 2.11 | 2.37 | 2.63 |
| $\left(\boldsymbol{\gamma}_{1}\right)$ | -0.30 | -2.39 | -2.13 | -1.87 | -1.61 | -1.35 | -1.10 | -0.84 | -0.58 | -0.33 | -0.09 | 0.15 | 0.41 | 0.68 | 0.95 | 1.22 | 1.48 | 1.75 | 2.01 | 2.27 | 2.53 | 2.79 |
|  | -0.40 | $-2.23$ | -1.97 | -1.71 | -1.45 | -1.19 | -0.93 | -0.68 | -0.42 | -0.17 | 0.06 | 0.30 | 0.57 | 0.84 | 1.11 | 1.38 | 1.64 | 1.91 | 2.17 | 2.43 | 2.69 | 2.95 |
|  | -0.50 | $-2.07$ | -1.81 | -1.55 | -1.29 | -1.03 | -0.78 | -0.52 | -0.27 | -0.03 | 0.20 | 0.44 | 0.72 | 1.00 | 1.27 | 1.54 | 1.80 | 2.06 | 2.32 | 2.58 | 2.84 | 3.10 |
|  | -0.60 | -1.91 | -1.65 | -1.39 | -1.13 | -0.87 | -0.62 | -0.37 | -0.13 | 0.10 | 0.32 | 0.58 | 0.87 | 1.15 | 1.42 | 1.69 | 1.95 | 2.22 | 2.48 | 2.74 | 3.00 | 3.26 |
|  | -0.70 | -1.75 | -1.49 | -1.23 | -0.97 | -0.72 | -0.47 | -0.22 | 0.01 | 0.22 | 0.46 | 0.73 | 1.02 | 1.30 | 1.57 | 1.84 | 2.11 | 2.37 | 2.63 | 2.89 | 3.16 | 3.42 |
|  | -0.80 | -1.59 | -1.33 | -1.07 | -0.82 | -0.57 | -0.32 | -0.09 | 0.14 | 0.35 | 0.60 | 0.88 | 1.17 | 1.45 | 1.72 | 1.99 | 2.26 | 2.52 | 2.79 | 3.05 | 3.31 | 3.57 |
|  | -0.90 | -1.43 | -1.17 | -0.92 | -0.67 | -0.42 | -0.18 | 0.05 | 0.27 | 0.50 | 0.76 | 1.04 | 1.32 | 1.60 | 1.88 | 2.15 | 2.41 | 2.68 | 2.94 | 3.20 | 3.47 | 3.73 |
|  | -1.00 | -1.27 | -1.02 | -0.76 | -0.52 | -0.27 | -0.04 | 0.18 | 0.41 | 0.65 | 0.92 | 1.20 | 1.48 | 1.76 | 2.03 | 2.30 | 2.57 | 2.83 | 3.09 | 3.36 | 3.62 | 3.88 |

Notes: This spreadsheet shows the value of the coefficient on expenditure $(\widehat{\beta 1})$ when estimating the cancer outcome equation $\left[y^{*}=\left(y-\gamma_{1} z_{1}-\gamma_{2} z_{2}\right)=\alpha+\beta 1 x+\beta 2 n+\epsilon\right]$ using IV having imposed different pairs of values for $\gamma 1$ and $\gamma 2$ between -1 and 1 . Cells in the top left-hand quadrant contain negative values for the outcome elasticity. The outcome elasticity associated with our standard IV model ( -0.34 ) is shown in the central square where $\gamma 1$ and $\gamma 2$ are, of course, zero.

## B9.4 The identification of values to be imposed on the coefficients on the excluded instruments

Our outcome equations typically involve two instruments and one endogenous regressor. With this structure we can re-estimate our outcome model twice, each time including one of the previously excluded instruments to the equation of interest. In particular, we can estimate

$$
\begin{equation*}
y=\alpha+\beta 1 x+\beta 2 n+\gamma 1 z 1+\quad \epsilon \tag{9.4}
\end{equation*}
$$

and then

$$
\begin{equation*}
y=\alpha+\beta 1 x+\beta 2 n \quad+\gamma 2 z 2+\epsilon \tag{9.5}
\end{equation*}
$$

with the same set of (included and excluded) instruments ( $n, z 1$, and $z 2$ ) being used to instrument $x 1$ in both cases. This provides us with coefficient and variance estimates for $\gamma 1$ and $\gamma 2$ and we can sample from these point estimates and their distributions to examine the impact of different (non-zero) values for $\gamma 1$ and $\gamma 2$ on the outcome elasticity $(\widehat{\beta 1})$.

The sampling procedure is straightforward. We sample from these estimates and their distributions by drawing two random numbers from a standard normal distribution and we form the product of these numbers and the standard errors associated with our estimates of $\gamma 1$ and $\gamma 2$. Our sampled pair of values of $\gamma 1$ and $\gamma 2$ (call these sampled values $\tilde{\gamma}_{1}$ and $\tilde{\gamma}_{2}$ ) are then the sum of these products and the respective coefficient estimates of $\gamma 1$ and $\gamma 2$. Table B9.2 shows the relevant coefficient and variance estimates for $\gamma 1$ and $\gamma 2$ that are used as part of this sampling procedure.

Table B9.2: table showing various estimates associated with the excluded instruments from the outcome equation for the big four programmes

| Programme | Instrument | coefficient | Std error | variance | covariance |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Cancer | $z 1$ : lone pensione | -0.2074942 | 0.1773647 | 0.0314582 | 0.00494454 |
|  | z2: IMD2007 | -0.0827677 | 0.1141054 | 0.0130200 |  |
| Circulatory disease | z1: lone pensione | -0.2606290 | 0.2441101 | 0.059590 | 0.01122591 |
|  | z2:IMD2007 | -0.2105334 | 0.2879230 | 0.082900 |  |
| Respiratory probler | z1: long term une | 0.2642582 | 0.13273061 | 0.0176174 | -0.02136305 |
|  | limiting long- | -1.739808 | 1.611403 | 2.5966196 |  |
| Gastro-intestinal problems | z1: unpaid carers | 1.812286 | 2.347459 | 5.510564 | 0.08016639 |
|  | z2: IMD2007 | 0.5567431 | 0.2066839 | 0.0427822 |  |

The estimation of equations (9.4) and (9.5) does not generate estimates of $\gamma 1$ and $\gamma 2$ as part of the same model and so the sampling procedure outlined above implicitly assumes a zero covariance between these estimates. If we want to incorporate a covariance term into the sampling procedure this must be obtained from elsewhere. In the absence of an obviously better approach, we obtain a covariance term from the OLS estimation of our outcome model with the previously excluded instruments both included in the regression equation. Thus we estimate

$$
\begin{equation*}
y=\alpha+\beta 1 x+\beta 2 n+\gamma 1 z 1+\gamma 2 z 2+\epsilon \tag{9.6}
\end{equation*}
$$

where $y, x, n, z 1$ and $z 2$ have their usual meaning, and where $\beta 1$ is constrained to be equal to its value from the IV estimation of equation (9.1). The value for the covariance term between the estimates of $\gamma 1$ and $\gamma 2$ for each of the big four programmes is shown in the final column of Table B9.2.

The sampling procedure from our estimates of $\gamma 1$ and $\gamma 2$ with a non-zero covariance is essentially the same as that outlined above but it incorporates the presence of a covariance term for the estimates of $\gamma 1$ and $\gamma 2$. We can illustrate this procedure using data from the cancer outcome model. First, we form the implied variance-covariance matrix for the estimates of $\gamma 1$ and $\gamma 2$

|  | lone_pensioners | IMD2007 |
| :--- | :--- | :--- |
| lone_pensioners | 0.031458 | 0.004944 |
| IMD2007 | 0.004944 | 0.013020. |

Second, we form the product of a pair of random numbers from the standard normal distribution ( $\mathrm{r} 1, \mathrm{r} 2$ ) with the Cholesky decomposition matrix from the variance-covariance matrix for the estimates of $\gamma 1$ and $\gamma 2$. The latter is given by:

|  | lone_pensioners | IMD2007 |
| :--- | :--- | :--- |
| lone_pensioners | 0.177364 | 0 |
| IMD2007 | 0.027877 | 0.110647. |

Finally, we add this pair of products to the respective coefficient estimates of $\gamma 1$ and $\gamma 2$ to obtain our sampled pair of values of $\gamma 1$ and $\gamma 2$ (call these $\tilde{\gamma}_{1}$ and $\tilde{\gamma}_{2}$ ). In other words, for each pair of random numbers ( $\mathrm{r} 1, \mathrm{r} 2$ ), we calculate the sampled values $\tilde{\gamma}_{1}$ and $\tilde{\gamma}_{2}$ where


This sampling procedure is undertaken 1,000 times, both with a zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$, and again with a non-zero covariance.

These procedures generate two sets of 1,000 pairs of values for $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ (one set assumes a zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ and the other does not). These sets of values for $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ can be used to examine the sensitivity of the estimated outcome elasticity to alternative non-zero values for the coefficients on the excluded instruments.

## B9.5 Obtaining the outcome elasticities associated with sampled coefficients on the excluded instruments

For each pair of sampled values of $\gamma 1$ and $\gamma 2\left(\tilde{\gamma}_{1}\right.$ and $\left.\tilde{\gamma}_{2}\right)$, we can use IV techniques to estimate the model

$$
\begin{equation*}
y_{\text {new }}=y-\tilde{\gamma} 1 z 1-\tilde{\gamma} 2 z 2=\alpha+\beta 1 x+\beta 2 n+\epsilon \tag{9.7}
\end{equation*}
$$

with the usual instrument set ( $\mathrm{x} 2, \mathrm{z} 1$, and z 2 ) used to instrument the endogenous variable x 1 (expenditure). For each pair of sampled the values $\tilde{\gamma}_{1}$ and $\tilde{\gamma}_{2}$, we obtain a different outcome elasticity $(\widehat{\beta 1})$ and these different values can be plotted in a histogram. Such a plot illustrates the uncertainty associated with our point estimate of the outcome elasticity due to doubts about the validity of our instruments; we call this type of uncertainty 'level 2' uncertainty. This 'level 2' uncertainty is in addition to what we label 'level 1' uncertainty, that is, the uncertainty about the value of the outcome elasticity assuming the validity of our exclusion restrictions (remember that our estimated outcome elasticity is only a point estimate and that it has a distribution attached to it). To illustrate this 'level 1' uncertainty, we can sample from the distribution of the point estimate for the outcome elasticity from our basic IV model (where $\tilde{\gamma}_{1}$ and $\tilde{\gamma}_{2}$ are zero in equation (9.7)) and plot the sampled values.

Plots illustrating the degree of level 1 uncertainty for each of the big four programmes are shown as Figures B9.1a (cancer), B9.2a (circulatory disease), B9.3a (respiratory problems), and B9.4a (gastrointestinal problems) below. These level 1 uncertainty plots can be compared with plots of $\widehat{\beta 1}$ from the estimation of equation (9.7). The latter plots illustrate the degree of level 2 uncertainty, that is, the uncertainty associated with our point estimate of the outcome elasticity due to doubts about the validity of the instruments. Figures B9.1b (cancer), B9.2b (circulatory disease), B9.3b (respiratory problems), and B9.4b (gastro-intestinal problems) show plots of the outcome elasticity ( $\widehat{\beta 1}$ from equation (9.7)) assuming a zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4) and (9.5). Figures B9.1c (cancer), B9.2c (circulatory disease), B9.3c (respiratory problems), and B9.4c (gastro-intestinal problems) show plots of the outcome elasticity ( $\widehat{\beta 1}$ from equation (9.7)) assuming a non-zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4) and (9.5).

Finally, the point estimates $\widehat{\beta 1}$ from the estimation of equation (9.7) also have a standard error and we can sample from these distributions. These sampled values illustrate what we term 'level 3' uncertainty, that is, the uncertainty associated with the value of the outcome elasticity due to both level 1 (sampling) and level 2 (instrument invalidity) effects.

Plots illustrating the degree of level 3 uncertainty, assuming a zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4) and (9.5), are shown as Figures B9.1d (cancer), B9.2d (circulatory disease), B9.3d (respiratory problems), and B9.4d (gastro-intestinal problems) below. Plots illustrating the degree of level 3 uncertainty, assuming a non-zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4) and (9.5), are shown as Figures B 9.1 e (cancer), B9.2e (circulatory disease), B9.3e (respiratory problems), and B9.4e (gastrointestinal(problems) below.

Uncertainty and the value of the cancer outcome elasticity (Figures B9.1a-B9.1e)
Figure B9.1a plots 1000 values from the distribution of the point estimate for the cancer outcome elasticity. The mean value of these sampled values is -0.338 (the outcome elasticity in the basic IV model is -0.342 and its standard error is 0.099 ) and virtually all of them lie between 0 and -0.6.


Figure B9.1a: graph showing histogram for the sampled cancer outcome elasticity
The histogram in Figure B9.1b provides a plot of 1000 point estimates for the cancer outcome elasticity if we drop the assumption that the coefficients on the excluded instruments are exactly zero (and we also assume a zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4) and (9.5)). The mean value of these 1000 outcome elasticities is -0.209 and this is about one-third lower than the elasticity in the basic IV model (= -0.342). In addition, the mean value of the standard errors associated with these 1000 elasticities (0.109) is slightly greater than the standard error in the basic IV model $(0.099)$ so that about one-quarter of the outcome elasticities in Figure B9.1b take a non-negative value.

The histogram in Figures B9.1c provides a similar plot to that in Figure B9.1b but this time we assume a non-zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4) and (9.5). There is very little difference between the zero (Figure B9.1b) and non-zero (9.1c) covariance plots, with both the mean elasticity and mean standard errortirtually identical in both plots.

The histograms in Figures B9.1b and B9.1c provide plots of the point estimate for the cancer outcome elasticity if ye drop the assumption that the coefficients on the excluded instruments are exactly zero. Each pointestimate also has a standard error and we can sample from these estimates and their distubutions to obtain the histograms shown in Figures B9.1d and B9.1e. With the exception of a slight lengthening in the tail on the left hand side, these plots are very similar to the plots in Figures B9.1b and B9.1 c .


Figure B9.1b: graph showing histogram for estimated outcome elasticity associated with cancer outcome model (zero covariance between the coefficients on the excluded instruments)


FigureB9.1c: graph showing histogram for estimated outcome elasticity associated with cancer outcome model (non-zero covariance between the coefficients on the excluded instruments)


Figure B9.1d: graph showing histogram for sampled values of the cancer outcome elasticity (zero covariance between the coefficients on the exchuded instruments)
sampling from the 1000 outcome elasticities for cancer expenditure (that were generated using 1000 alternative pairs of coefficients on the excluded instruments with a non-zero covariance between these coefficients)


NB The mean value of these 1000 sampled outcome elasticities is -0.218 .
The outcome elasticity for cancer expenditure in the basic IV model is -0.342 .

Figure B9.1e: graph showing histogram for sampled values of the cancer outcome elasticity (non-zero covariance between the coefficients on the excluded instruments)

Figure B9.1f reproduces the three kernel density plots shown in Figures B9.1a, B9.1b and B9.1d (remember that Figures B9.1b and B9.1d assume a zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4) and (9.5)). Together these plots illustrate the impact of all three levels of uncertainty on our estimate of the cancer outcome elasticity. It is clear that the uncertainty induced by the instrument validity issue considerably increases the uncertainty associated with our estimate of the outcome elasticity (compare, for example, the density plot for level 1 uncertainty with those for both level 2 and level 3 uncertainty).

 The outcome elasticity for cancer expenditure in the basic IV model is -0.342 .

Figure B9.1f: graph showing comparing the three kernel density plots shown in Figures B9.1a, 9.1b and 9.1d

Figure B9.1g reproduces the three kernel density plots shown in Figures B9.1a, B9.1c and B9.1e (remember that Figures B9.1c and B9.1e assume a non-zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4) and (9.5)). As is the case for Figure B9.1f, these plots illustrate the impact of all three levels of uncertainty on our estimate of the cancer outcome elasticity. And again, it is clear that it is the uncertainty induced by the instrument validity issue that considerably increases the uncertainty associated with our estimate of the outcome elasticity. For example, the standard deviation associated with the level 1 uncertainty density plot is 0.099 but the standard deviation for the level $2(0.338)$ and level $3(0.379)$ uncertainty density plots are both considerably larger than this.
kernel density plots from Figures B9.1a, B9.1c and B9.1e: illustrating the uncertainty associated with the point estimate for the cancer outcome elasticity

 The outcome elasticity for cancer expenditure in the basic IV model is -0.342 .

Figure B9.1g: graph showing comparing the three kernel density plots shown in Figures B9.1a, B9.1c and B9.1e

Uncertainty and the value of the circulatory disease outcome elasticity (Figures B9.2a-B9.2e)
Figure B9.2a plots 1000 values from the distribution of the point estimate for the circulatory disease outcome elasticity. The mean value of these sampled values is -1.418 and virtually all of these values lie between -2.0 and -0.75 . The outcome elasticity in the comparable IV model is $-1.427 .{ }^{33}$

[^64]

NB The mean value of these 1000 sampled outcome elasticities is -1.418 .
The outcome elasticity for circulatory disease expenditure in the comparable IV model is $\mathbf{- 1 . 4 2 7 .}$

Figure B9.2a: graph showing histogram for the sampled circulatory disease outcome elasticity
The histogram in Figure B9.2b provides a plot of 1000 point estimates for the circulatory disease outcome elasticity if we drop the assumption that the coefficients on the excluded instruments are exactly zero (and we also assume a zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4) and (9.5)). The mean value of these 1000 outcome elasticities is -1.697 and this is about one-fifth larger than the elasticity in the comparable IV model (-1.427). Similarly, the mean value of the standard errors associated with these 1000 elasticities ( 0.269 ) is also about One-fifth larger than the standard error in the comparable basic IV model ( 0.228 ). Virtually all of the point estimates values lie between -4.0 and 0.0 , and there are very few non-negative values.

The histogram in Figure B9.2c provides a similar plot to that in Figure B9.2b but this time we assume a non-zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4) and (9.5). There is very little difference between the zero (Figure B9.2b) and non-zero (9.2c) covariance plots, with both the mean elasticity and mean standard error virtually identical in both plots.

The histograms in Figures B9.2b and B9.2c provide plots of the point estimate for the circulatory disease outcome elasticity if we drop the assumption that the coefficients on the excluded instruments are exactly zero. Eachpoint estimate also has a standard error and we can sample (with replacement) from these estimates and their distributions to obtain the histograms shown in Figures B9.2d and B9.2e. With the exception of a slight lengthening in the tail on the left hand side (as was also observed for the cancer programme), these plots are very similar to the plots in Figures B9.2b and B9.2c.


NB The mean value of the 1000 outcome elasticities is -1.697 (mean $\mathrm{SE}=0.269$ ).
The outcome elasticity for circulatory expenditure in the comparable IV model is -1.427 ( $\mathrm{SE}=0.228$ ).

Figure B9.2b: graph showing histogram for estimated outcome elasticity associated with the circulatory disease outcome model (zero covariance between the coefficients on the excluded instruments)

Sensitivity of the outcome elasticity for circulatory disease expenditure to 1000 alternative pairs of coefficients on the excluded instruments with non-zero covariance between the coefficients


NB The mean value of the 1000 outcome elasticities is -1.700 (mean $\mathrm{SE}=0.269$ ).
The outcome elasticity for circulatory expenditure in the comparable IV model is $-1.427(\mathrm{SE}=0.228)$.

Figure B9.2c: graph showing histogram for estimated outcome elasticity associated with the circulatory disease outcome model (non-zero covariance between the coefficients on the excluded instruments)
sampling from the 1000 outcome elasticities for circulatory disease expenditure (that were generated using 1000 alternative pairs of coefficients on the
excluded instruments with a zero covariance between these coefficients)


NB The mean value of these 1000 sampled outcome elasticities is -1.717 .
The outcome elasticity for circulatory expenditure in the comparable IV model is $\mathbf{- 1 . 4 2 7 .}$

Figure B9.2d: graph showing histogram for sampled values of the circulatory disease outcome elasticity (zero covariance between the coefficients on the excluded instruments)
sampling from the 1000 outcome elasticities for circulatory disease expenditure
(that were generated using 1000 alternative pairs of coefficients on the excluded instruments with a non-zero covariance between these coefficients)


NB The mean value of these 1000 sampled outcome elasticities is -1.718 .
The outcome elasticity for circulatory expenditure in the comparable IV model is $\mathbf{- 1 . 4 2 7 .}$

Figure 9.2e: graph showing histogram for sampled values of the circulatory disease outcome elasticity (non-zero covariance between the coefficients on the excluded instruments)

Figure B9.2f reproduces the three kernel density plots shown in Figures B9.2a, B9.2b and B9.2d (remember that Figures B9.2b and B9.2d assume a zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4) and (9.5)). Together these plots illustrate the impact of all three levels of uncertainty on our estimate of the circulatory disease outcome elasticity. It is clear that the uncertainty induced by the instrument validity issue considerably increases the uncertainty associated with our estimate of the outcome elasticity (note that the range of values increases dramatically from the density plot illustrating level 1 uncertainty to that illustrating level 2 uncertainty).


The outcome elasticity for circulatory disease expenditure in the comparable IV model is -1.427 .

Figure B9.2f: graph showing comparing the three kernel density plots shown in Figures B9.2a, B9.2b and B9.2d

Figure B9.2g reproduces the three kernel density plots from Figures B9.2a, B9.2c and B9.2e (remember that Figures B9.2c and B9.2e assume a non-zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4) and (9.5)). As is the case for Figure B9.2f, these plots illustrate the impact of all three levels of uncertainty on our estimate on the circulatory disease outcome elasticity. And again, it is clear that it is the uncertainty induced by the instrument validity issue that considerably increases the uncertainty associated with our estimate of the outcome elasticity. For example, the standard deviation associated with the level 1 uncertainty density plot is 0.228 but the standard deviation for the level $2(0.735)$ and level $3(0.843)$ uncertainty density plots are both considerably larger than this.


NB The mean value of the level 1/level 2/level 3 elasticities is $-1.418 /-1.700 /-1.718$.
The outcome elasticity for circulatory disease expenditure in the comparable IV model is $\mathbf{- 1 . 4 2 7 .}$

Figure B9.2g: graph showing comparing the three kernel density plots shown in Figures B9.2a, B9.2c and B9.2e

Uncertainty and the value of the respiratory disease outcome elasticity)(Figures B9.3a-B9.3e)
Figure B9.3a plots 1000 values from the distribution of the point estimate for the respiratory disease outcome elasticity (see column 5 of Table B8.19). The mean value of these sampled values is -2.004 (the outcome elasticity in the comparable IV model is -2.029 ) and all of these values lie between -4.0 and 0 .


Figure B9.3a: graph showing histogram for the sampled respiratory disease outcome elasticity

The histogram in Figure B9.3b provides a plot of 1000 point estimates for the respiratory disease outcome elasticity if we drop the assumption that the coefficients on the excluded instruments are exactly zero (and we also assume a zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4) and (9.5)). The mean value of these 1000 outcome elasticities from the respiratory disease outcome model ( -1.145 ) is almost one-half of the size of the elasticity in the comparable basic IV model (-2.029). And the mean value of the standard errors associated with these 1000 elasticities ( 0.489 ) is about one-quarter less than the standard error in the comparable basic IV model (0.636).

The histogram in Figure B9.3c provides a similar plot to that in Figure B9.3b but this time we assume a non-zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4) and (9.5). However, there is very little difference between the zero (Figure B9.3b) and non-zero (B9.3c) covariance plots, with both the mean elasticity and mean standard error virtually identical in these two plots.

The histograms in Figures B9.3b and B9.3c provide plots of the point estimate for the respiratory disease outcome elasticity if we drop the assumption that the coefficients on the excluded instruments are exactly zero. Each point estimate also has a standard error and we can sample from these estimates and their distributions to obtain the histograms shown in Figures B9.3d and B9.3e. With the exception of a slight lengthening of the tail on the left hand side, these plots are similar to those in Figures B9.3b and B9.3c so the sampling procedure would appear to have little impact on the distribution of the point elasticities.

Sensitivity of the outcome elasticity for respiratory expenditure to 1000 alternative pairs of coefficients on the excluded instruments with a zero covariance between these coefficients


NB The mean value of the 1000 outcome elasticities is -1.145 (mean $\mathrm{SE}=0.489$ ). The outcome elasticity for respiratory expenditure in the comparable IV model is -2.029 ( $\mathrm{SE}=0.636$ ).

Figure B9.3b: graph showing histogram for estimated outcome elasticity associated with respiratory disease outcome model (zero covariance between the coefficients on the excluded instruments)


NB The mean value of the 1000 outcome elasticities is -1.149 (mean $\mathrm{SE}=0.485$ ).
The outcome elasticity for respiratory expenditure in the comparable IV model is -2.029 ( $\mathrm{SE}=0.636$ ).

Figure B9.3c: graph showing histogram for estimated outcome elasticity associated with respiratory disease outcome model (non-zero covariance between the coefficients on the excluded instruments)
sampling from the 1000 outcome elasticities for respiratory disease expenditure
(that were generated using 1000 alternative pairs of coefficients on the excluded instruments with a zero covariance between these coefficients)


NB The mean value of these 1000 sampled outcome elasticities is -1.146 .
The outcome elasticity for respiratory expenditure in the comparable IV model is -2.029 .

Figure B9.3d: graph showing histogram for sampled values of the respiratory disease outcome elasticity (zero covariance between the coefficients on the excluded instruments)
sampling from the 1000 outcome elasticities for respiratory disease expenditure (that were generated using 1000 alternative pairs of coefficients on the excluded instruments with a non-zero covariance between these coefficients)



NB The mean value of these 1000 sampled outcome elasticities is -1.151 .
The outcome elasticity for respiratory expenditure in the comparable IV model is -2.029 .

Figure B9.3e: graph showing histogram for sampled values of the respiratory disease outcome elasticity (non-zero covariance between the coefficients on the excluded instruments)

Figure B9.3f reproduces the three kernel density plots shown in Figures B9.3a, B9.3b and B9.3d (remember that Figures B9.3b and B9.3d assume a zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4) and (9.5)). Together these plots illustrate the impact of all three levels of uncertainty on our estimate of the respiratory disease outcome elasticity. It is clear that the uncertainty induced by the instrument validity issue both shifts the density plot to the right and increases the uncertainty associated with our estimate of the outcome elasticity (e.g., the range of values increases from -4.0 to 0.0 at level 1 to -5.0 to 2.5 at level 3).

## kernel density plots from Figures B9.3a, B9.3b and B9.3d: illustrating the uncertainty associated with the point estimate for the respiratory disease outcome elasticity



NB The mean value of the level $1 / l e v e l$ 2/level 3 elasticities is $-2.004 /-1.145 /-1.146$.
The outcome elasticity for respiratory disease expenditure in the comparable IV model is -2.029 .
Figure B9.3f: graph showing comparing the three kernel density plots shown in Figures B9.3a, B9.3b and B9.3d

Figure B9.3g reproduces the three kernel density plots shown in Figures B9.3a, B9.3c and B9.3e (remember that Figures B9.3c and B9.3e assume a non-zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4) and (9.5)). As is the case for Figure B9.3f, these plots illustrate the impact of all three levels of uncertainty on our estimate of the respiratory disease outcome elasticity. And again, it is clear that the uncertainty induced by the instrument validity issue both shifts the density plot to the right and considerably increases the uncertainty associated with our estimate of the outcome elasticity. More precisely, the standard deviation associated with the level 1 uncertainty density plot is 0.636 but the standard deviation for the level $2(0.919)$ and level 3 (1.098) uncertainty density plots are both considerably larger than this.


NB The mean value of the level $1 /$ level $2 /$ level 3 elasticities is $-2.004 /-1.149 /-1.151$. The outcome elasticity for respiratory disease expenditure in the comparable IV model is -2.029.

Figure B9.3g: graph showing comparing the three kernel density plots shown in Figures B9.3a, B9.3c and B9.3e

Uncertainty and the value of the gastro-intestinal disease outcome elasticity (Figures B9.4a-B9.4e)
Figure B9.4a plots 1000 values from the distribution of the point estimate for the gastro-intestinal disease outcome elasticity (see column 7 of Table B8.19). The mean of these sampled values is -1.518 (the outcome elasticity in the comparable IV model is -1.536 ) and all of these values lie between -3.0 and 0.0 .
sampling 1000 values from the distribution of the point estimate for the gastro-intestinal disease outcome elasticity


NB The mean value of these 1000 sampled outcome elasticities is -1.518 .
The outcome elasticity for gastro-intestinal expenditure in the comparable IV model is -1.536 .
Figure B9.4a: graph showing histogram for the sampled gastro-intestinal disease outcome elasticity

The histogram in Figure B9.4b provides a plot of 1000 point estimates for the respiratory disease outcome elasticity if we drop the assumption that the coefficients on the excluded instruments are exactly zero (and we also assume a zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4) and (9.5)). The mean value of these 1000 outcome elasticities $(-2.365)$ is $50 \%$ larger than the size of the elasticity in the comparable IV model ( -1.536 ). And the mean value of the standard errors associated with these 1000 elasticities ( 0.853 ) is about $80 \%$ larger than the standard error in the basic IV model ( 0.468 ).

The histogram in Figure B9.4c provides a similar plot to that in Figure B9.4b but this time we assume a non-zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4) and (9.5). However, there is very little difference between/the zero (Figure B9.4b) and non-zero (B9.4c) covariance plots, with both the mean elasticity and mean standard error virtually identical in these plots.

The histogranss in Figures B9.4b and B9.4c provide plots of point estimates for the gastro-intestinal problems outcome elasticity if we drop the assumption that the coefficients on the excluded instruments are exactly zero. Each point estimate also has a standard error and we can sample from these estimates and their distributions to obtain the histograms shown in Figures B9.4d and B9.4e. With the exception of a slight extension to both tails, these plots are similar to the plots in Figures B9.4b and B9.4c.


NB The mean value of the 1000 outcome elasticities is -2.365 (mean $\mathrm{SE}=0.853$ ).
The outcome elasticity for gastro-intestinal expenditure in the basic IV model is -1.536 ( $\mathrm{SE}=0.468$ ).

Figure B9.4b: graph showing histogram for estimated outcome elasticity associated with gastrointestinal outcome model (zero covariance between the coefficients on the excluded instruments)

Sensitivity of the outcome elasticity for gastro-intestinal expenditure to 1000 alternative pairs of coefficients on the excluded instruments with non-zero covariance between the coefficients


NB The mean value of the 1000 outcome elasticities is -2.360 (mean $\mathrm{SE}=0.839$ ).
The outcome elasticity for gastro-intestinal expenditure in the basic IV model is -1.536 ( $\mathrm{SE}=0.468$ ).

Figure B9.4c: graph showing histogram for estimated outcome elasticity associated with gastrointestinal outcome model (non-zero covariance between the coefficients on the excluded instruments)


Figure B9.4d: graph showing histogram for sampled values of the gastro-intestinal problems outcome elasticity (zero covariance between the coefficients on the excluded instruments)


Figure B9.4e: graph showing histogram for sampled values of the gastro-intestinal problems outcome elasticity (non-zero covariance between the coefficients on the excluded instruments)

Figure B9.4f reproduces the three kernel density plots shown in Figures B9.4a, B9.4b and B9.4d (remember that Figures B9.4b and B9.4d assume a zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4)
and (9.5)). These plots illustrate the impact of all three levels of uncertainty on our estimate of the gastrointestinal outcome elasticity. It is clear that the uncertainty induced by the instrument validity issue both shifts the density plot to the left slightly and dramatically increases the uncertainty associated with our estimate of the outcome elasticity (e.g., the range of values increases from -3 to 0 at level 1, from -13 to 9 at level 2, and then further from -16 to 11 at level 3 ).


Figure B9.4f: graph showing comparing the three kernel density plots shown in Figures B9.4a, B9.4b and B9.4d

Figure B9.4g reproduces the three kernel density plots shown in Figures B9.4a, B9.4c and B9.4e (remember that Figures B9.4c and B9.4e assume a non-zero covariance between $\hat{\gamma}_{1}$ and $\hat{\gamma}_{2}$ in equations (9.4) and (9.5)). As is the case for Figure B9.4f, these plots illustrate the impact of all three levels of uncertainty on our estimate of the gastro-intestinal outcome elasticity. And again, it is clear that the uncertainty induced by the instrument validity issue both shifts the density plot to the left slightly and considerably increases the uncertainty (range) associated with our estimate of the outcome elasticity. More precisely, the standard deviation associated with the level 1 uncertainty density plot is 0.468 but the standard deviation for the level 2 (3.658) and level 3 (3.834) uncertainty density plots are both eight times larger than this.
kernel density plots from Figures B9.4a, B9.4c and B9.4e: illustrating the uncertainty associated with the point estimate for the gastro-intestinal outcome elasticity



NB The mean value of the level $1 /$ level $2 /$ level 3 elasticities is $-1.518 /-2.360 /-2.434$.
The outcome elasticity for gastro-intestinal expenditure in the comparable IV model is -1.536 .
Figure B9.4g: graph showing comparing the three kernel density plots shown in Figures B9.4a, B9.4c and B9.4e

## B9.6 Implications of uncertainty for the estimate of the cost of a life year

In the previous subsection, we have evaluated the outcome equation elasticities when uncertainty over the validity of instrument variables is considered (level 3' uncertainty), in contrast to assuming the instruments are valid ('level 1' uncertainty). This analysis showed that including level 3 uncertainty affects the central value of the outcome elasticities; however, it is difficult to predict its effect on the expectation of the threshold given the impact of expenditure on mortality appears reduced in some programmes but increased in others. In Table B9 3, the mean estimates of the outcome elasticities under level 3 uncertainty were used to calculate the threshold for the big four programmes of health. The results show that relaxing the assumption of validity of the instruments has little impact on the expectation of the threshold for the big 4 PBCs [thecost Der life year gained threshold changed from $£ 10,604$ (Table B8.22) to $£ 11,009$ in Table B9.3].

Table B9.3: Cost of life and life year estimates for the big four programmes using expenditure data for 2006 and outcome data for 2006/7/8 adjusted for the ICD10 coverage of the expenditure and outcome data


The assumption of validity of instruments is expected to affect significantly the level of uncertainty over the cost effectiveness threshold estimate. Illustrations of this source of uncertainty were presented in the previous section (B9.5) using empirical distributions derived from the sampling procedure implemented; these illustrations represent the uncertainty in the mean estimate for each of the elasticities. To characterise the effect of levels 1 and 3 uncertainty on the overall threshold we used the sets of simulated elasticities (one for each of the 4 programmes of care) to compute a threshold value; in doing so for all simulated sets, a sample of threshold values was obtained. In this way, uncertainty was propagated from the outcome elasticities to the threshold estimates, and an empiric distribution describing uncertainty over threshold estimates obtained. The cumulative density function can be used to display such uncertainty; this plots the probability ( y -axis) of the threshold being below certain values ( x -axis) in the simulated sample (this corresponds to a Bayesian interpretation of uncertainty). Figure B9.5a plots the cumulative density curve for the cost per life gained threshold when level 1 and level 3 uncertainty are considered in turn, and B9.5b for the cost per life year gained threshold.


In drawing the cumulative density function, negative threshold values were dealt with by evaluating whether it was the health component or the cost component that was negative. For simulations where health change was negative ( $0 \%$ were observed for both levels 1 and 3 ), the threshold was left as a negative value. Simulations showing a negative change in spend were

Figure B9.5a Cumulative density plot for the cost per life gained threshold for the big 4 PBCs (considers covariance between the coefficients on the excluded instruments).

Cost per life year gained (4 PBC)


In drawing the cumulative density function, negative threshold values were dealt with by evaluating whether it was the health component or the cost component that was negative. For simulations where health change was negative ( $0 \%$ were observed for both levels 1 and 3 ), the threshold was left as a negative value. Simulations showing a negative change in spend were assigned a very high positive threshold value - in this was an asymptote is generated in the plot (respectively $0.04 \%$ and $7.7 \%$ were observed for levels 1 and 3 ).

Figure B9.5b Cumulative density plot for the cost per life year gained threshold for the big 4 PBCs (considers covariance between the coefficients on the excluded instruments)

The probability that the overall threshold is less than $£ 7,500$ per life year is around 0.2 when uncertainty over the validity of instruments in considered (level 3), whereas when the instruments are assumed valid (level 1) this probability is 0 . Under level 1 uncertainty, we would be confident that the threshold is less than $£ 30,000$ (probability of 1 ), but when considering level 3 uncertainty there is some chance that the threshold is higher than $£ 30,000$ (probability of 0.2 ). These plots show that uncertainty on the validity of the instruments generates significant uncertainty over the threshold value.

## B9.7 Summary and conclusion

One of the crucial elements in the calculation of the cost of a life year for any care programme is the coefficient on the expenditure variable in the outcome equation. The endogenous nature of expenditure in our model means that OLS estimation is inappropriate and that instead IV techniques must be used. The application of these techniques requires the identification of variables that are good predictors of the endogenous variable (expenditure) but which do not have a direct effect on the dependent variable (mortality).

It is difficult to provide theoretical arguments why our selected instruments will not affect mortality directly. Instead, we rely on the widely used Hansen-Sargen test of instrument validity. Although our models 'pass' this test, some commentators have argued that this test has weak power and may fail to reject the null hypothesis of instrument validity even when an exclusion restriction is not valid. Given our reliance on this test, we noted that this test will only lack power if the biases induced in the coefficient on the endogenous variable by invalid instruments all coincide (i.e., the instruments all identify the same wrong parameter). However, it is far from obvious that this will be so in this case, particularly given that our outcome equation already includes a measure of the need for health care.

Nevertheless, it is possible that our instruments are correlated with both expenditure and some unobserved factor which is directly influencing the mortality rate, and that the induced bias in $\widehat{\beta 1}$ is the same for both instruments.

We therefore undertook an extensive sensitivity analysis of the estimated outcome elasticity to the validity of the exclusion restrictions. In summary, we found that both the central value and distribution of the outcome elasticity may change if we drop the assumption that the coefficients on the excluded instruments are identically zero.

This change in the central value of the outcome elasticity reduces the impact of expenditure on mortality in some programmes (e.g., for cancer the 'average' outcome elasticity falls from -0.338 to -0.210 , and for respiratory disease it falls from -2.004 to -1.151 ). However, in other programmes this change in the central value increases the impact of expenditure on mortality (e.g., for circulatory disease the 'average' outcome elasticity increases from -1.418 to -1.718 , and for gastro-intestinal problems it increases from 1.518 to -2.434 ).

Howeverin all four programmes the standard deviation associated with the distribution of the value for the outcome elasticity increased: for cancer it increased from 0.099 to 0.379 ; for circulatory disease it increased from 0.228 to 0.843 ; for respiratory disease it increased from 0.636 to 1.098 ; and for gastrointestinal disease it increased from 0.468 to 3.834 .

## B10 <br> Analysis of programme budgeting expenditure for 2007/8 and mortality data for 2007/2009

Outcome and expenditure models were estimated using updated data for expenditure (from 2006/7 to $2007 / 8$ ) and updated mortality data (from 2006/2007/2008 to 2007/2008 / 2009). Results for the outcome model are shown in Table B10.1 and results for the expenditure model are in Table B10.2. First stage regressions for these IV models can be found in Tables BA. 7 and BA. 8 in the annex.

## B10.1 Outcome models

Some of the outcome models in Table B10.1 contain just two variables: own programme expenditure and a measure of the need for health care. The latter is usually the measure of need as employed by the $\Omega$ Department of Health for resource allocation purposes and this incorporates the CARAN formula for acute services. For the respiratory programme we have added the square of this need measure to improve the model fit. In other PBCs we found that the all service measure of need performed poorly and we have replaced or supplemented it with either a more programme specific measure (e.g., the epilepsy prevalence rate for neurological mortality) or with a better performing proxy for need (e.g., the percentage of residents born outside the EU for maternity/neonate mortality).

Two results are reported for three of the big four programmes. One of these two results uses two instruments and so we report the instrument validity test statistic. We cannot reject the null hypothesis of instrument validity in all three cases. However, there is some evidence of weak instruments (at least in the respiratory and gastro-intestinal programmes) but, if we drop one instrument and re-estimate the model, the evidence of instrument weakness disappears (but of course there is no instrument validity test statistic with this re-estimation). The removal of one instrument has little impact on the coefficient on expenditure and it is this coefficient from this one instrument model that we use below in our cost of a life year calculations.

The first seven results in Table B10.1 show the outcome model for the big four programmes (i.e., for cancer, circulatory disease, respiratory problens and gastro-intestinal problems). In all four programmes the need variable has a positive and significant effect on mortality, and expenditure has the anticipated negative effect. The diagnostic statistics reveal that, in all four PBCs, own programme expenditure is endogenous and that the instruments are valid. They also suggest that the instruments are relevant and there is no evidence that the instruments are weak in the models with one excluded instrument. The Pesaran-Taylor test suggests that there is no evidence of model mis-specification.

The outcome results for the other programmes (in columns 8-13) are similar to but more diverse than those for the big four programmes. This is to be anticipated because mortality is a much rarer outcome in these programmes than it is in the big four programmes. Own programme expenditure is not endogenous in fourof these programmes but we retain the IV estimator for three of these four because this yields more plausible results than the OLS estimator (the results are more plausible in the sense that the signs on the coefficients are more in line with our prior expectations).

Expenditure has the anticipated negative effect on mortality in the endocrine problems programme but this is not statistically significant. The all service measure of need is not relevant for this PBC; instead, we find that the diabetes prevalence rate is positively associated with mortality, as is a measure of deprivation (the IMD2007).

Mortality from epilepsy is negatively and significantly associated with expenditure in the neurological programme. Both the all service need for health care and the epilepsy prevalence rate are positively and significantly associated with mortality in this programme.

Expenditure has a negative and statistically significant effect on mortality (from renal problems) in the genitor-urinary problems programme. The prevalence of lone parent households is positively associated with mortality.

Expenditure has the anticipated negative effect on mortality in the infectious disease programme and this is statistically significant. The all service measure of need is not relevant for this PBC; instead, we find that a measure of need associated with HIV is positively associated with mortality, as is a measure of deprivation (the IMD2007).

Expenditure has the anticipated negative effect on mortality in the maternity \& neonates programme but the estimated coefficient is not statistically significant. In this PBC the generic all service measure of need has been replaced with two other indicators of deprivation - the proportion of residents born outside the EU and the proportion of those aged 16-74 without any qualifications - and both of these are positively associated with mortality.

Finally, expenditure and need have the anticipated effects on mortality in the trauma and injuries programme. In addition, the proportion of households without access to a car is negatively associated with mortality from fractures (perhaps access to a car facilitates involvement in serious road traffic accidents), and the proportion of residents that are students is positively associated with mortality from fractures.

The relevant statistical test suggests that expenditure is endogenous in six of the ten programmes but we have retained the IV estimates for three of the other four programmes because they provide plausible results. The Hansen-Sargen test suggests that the selected instruments are valid, and the Kleibergen-Paap LM statistic suggests that they are relevant (i.e., correlated with the endogenous regressor). With the possible exception of the trauma and injuries programme, the Kleibergen-Paap F statistic suggests that we do not have a problem with weak instruments. ${ }^{34}$ Finally, the Pesaran-Taylor/Ramsey reset test statistics reveal no evidence of mis-specification.

## B10.2 Expenditure models

Most of the expenditure models in Table B10.2 contain just three variables: the PCT budget, a proxy for the own programme need for health care, and a proxy for the need for health care in other programmes.

The budget term is positive in all eleven models and it is statistically significant in eight of these eleven models.

The usual proxy for the own programme need for health care (i.e., the all service measure of need) is present in six of the models and it is significant in five of them. Its presence is supplemented with the addition of its squared value to improve model fit in the respiratory problems programme.

In some programmes (e.g., the endocrine, metabolic \& nutritional programme and the neurological programme), we have replaced and/or supplemented the all service measure of need with a more programme specificmeasure (e.g., the diabetes prevalence rate and the epilepsy prevalence rate) and these measures of need have the anticipated positive impact on expenditure.

In addition, in a couple of other programmes we have used alternative proxies for the own programme need (e.g., with the use of the Department of Health's measure of maternity need in the maternity/ neonates expenditure equation).

For eight of the eleven programmes we have used the all cause mortality rate less the own programme mortality rate as the proxy for the need for health care in other programmes, and the coefficient on this term is negative in seven programmes and statistically significant in six of the seven. In three programmes -- maternity/neonates, GMS/PMS and trauma \& injuries programmes -- we have used the all cause mortality rate as the proxy for the need for health care in other programmes due to difficulties associated with the measurement of the own programme mortality rate. The coefficient on this term is not significant in any of the three models.

[^65]The relevant statistical test suggests that expenditure is endogenous in six of the eleven programmes but we have retained the IV estimates for two other programmes (GMS/PMS and trauma \& injuries) because the IV estimator provides more plausible results. In the other three programmes we report OLS results.

The Hansen-Sargen test suggests that the selected instruments are valid, and the Kleibergen-Paap LM statistic suggests that they are relevant (i.e., correlated with the endogenous regressor). The KleibergenPaap F statistic suggests that we do not have a problem with weak instruments. Finally, the PesaranTaylor reset test statistics and the Ramsey reset F statistics reveal no evidence of model mis-specification.

## B10.3 Calculation of the cost of a life and life year

Expenditure and outcome elasticities for our preferred models are shown in Table B10.3 (see columns D and H ) and these are used to calculate the cost of a life and the cost of a life year, both for individual programmes and for all programmes collectively.

Column N reports the cost per life gained and column U reports the cost per life year gained. From the latter we can see that the cost per life year gained is $£ 13,830$ for the big four programmes and $£ 28,983$ for all ten programmes with a mortality-based outcome indicator. These represent $30 \%$ and $45 \%$ increases on the respective costs for the previous year (i.e., using expenditure data for $2006 / 77$ and mortality data for 2006/2007/2008).

If we assume that the other 13 programmes (all without a mortality based outcome indicator) offer no health gain, then the cost per life year across all PCT expenditure is $£ 82,765$. This is up from $£ 73,457$ using data for the previous year (an increase of $13 \%$ ).

In addition, Table B10.4 shows that if we assume that PB623 generates a zero health gain and that the gain attributable to the remaining 12 programmes is, on average, the same as that attributable to those with a mortality outcome measure, then the cost of life year across all programmes is $£ 31,846$ (it was $£ 22,565$ using data for the previous year).

## B10.4 Summary and conclusion

In this section we have estimated outcome and expenditure models using PB data for 2007/8 and mortality data for 2007/8/9. The cost of an additional life year for all ten programmes with a mortality based outcome is $£ 28,983$. This is a $45 \%$ increase on the cost $(£ 19,965)$ for the previous year (i.e., using expenditure data for $2006 / 7$ and mortality data for $2006 / 2007 / 2008$ ). The next section presents outcome and expenditure models using PB data for 2008/9 and mortality data for 2008/9/10, and it explores the reasons for the increase in the cost of an additional life year identified in this section.

Table B10.1: table showing outcome models using spend data for 2007/8 (two MFFs) and mortality data for 2007/8/9


[^66]Table B10.2: table showing expenditure models using spend data for 2007/8 (two MFFs) and mortality data for 2007/8/9

|  | (1) | (2) | (3) | (4) | (5) | (6) | (7) | (8) | (9) | (10) | (11) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| VARIABLES | PBC 2 <br> cancer <br> 2007/8 <br> spend model second stage | PBC 10 <br> circulation <br> 2007/8 <br> spend model <br> second stage | PBC 11 <br> respiratory <br> 2007/8 <br> spend model <br> second stage | PBC 13 <br> gastro-intestinal <br> 2007/8 <br> spend model second stage | PBC 1 <br> infectious disease <br> 2007/8 <br> spend model <br> OLS | PBC 4 <br> endocrine 2007/8 spend model second stage | PBC 7 <br> neurological <br> 2007/8 <br> spend model <br> second stage | PBC 17 <br> genito-urinary <br> 2007/8 <br> spend model <br> OLS | PBC 1819 maternity/neonates 2007/8 spend model OLS | PBC 23 <br> GMS/PMS etc 2007/8 <br> spend model <br> second stage | PBC 16 <br> trauma/injuries <br> 2007/8 <br> spend model <br> second stage |
| all cause SYLLR excluding cancer | $\begin{aligned} & -1.227^{* * *} \\ & {[0.220]} \end{aligned}$ |  |  |  |  |  |  |  |  |  |  |
| PCT budget per head | $\begin{aligned} & 0.890 * * \\ & {[0.431]} \end{aligned}$ | $\begin{aligned} & 0.293 \\ & {[0.350]} \end{aligned}$ | $\begin{aligned} & 0.536^{*} \\ & {[0.298]} \end{aligned}$ | $\begin{aligned} & 0.622^{*} \\ & {[0.321]} \end{aligned}$ | $\begin{aligned} & 1.435^{* * *} \\ & {[0.258]} \end{aligned}$ | $\begin{aligned} & 0.264 \\ & {[0.206]} \end{aligned}$ | $\begin{aligned} & 1.036^{* * *} \\ & {[0.307]} \end{aligned}$ | $\begin{aligned} & 1.004 * * * \\ & {[0.356]} \end{aligned}$ | $\begin{aligned} & 0.5 \widehat{4} 4^{*} \\ & {[0.264]} \end{aligned}$ | $\begin{aligned} & 0.563 \\ & {[0.344]} \end{aligned}$ | $\begin{aligned} & 1.686 * * * \\ & {[0.384]} \end{aligned}$ |
| need CARAN per head | $\begin{aligned} & 1.659 * * * \\ & {[0.430]} \end{aligned}$ | $\begin{aligned} & 3.117 * * * \\ & {[0.535]} \end{aligned}$ | $\begin{aligned} & 1.786 * * * \\ & {[0.334]} \end{aligned}$ | $\begin{aligned} & 1.982 * * \\ & {[0.422]} \end{aligned}$ |  | $\begin{aligned} & 0.925 * * * \\ & {[0.305]} \end{aligned}$ |  | $\begin{aligned} & 0.029 \\ & {[0.371]} \\ & \hline \end{aligned}$ |  |  |  |
| all cause SYLLR exc circulatory |  | $\begin{aligned} & -2.115 * * * \\ & {[0.397]} \end{aligned}$ |  |  |  |  |  |  |  |  |  |
| all cause SYLLR exc respiratory |  |  | $\begin{aligned} & -0.781 * * * \\ & {[0.236]} \end{aligned}$ |  |  |  |  | , |  |  |  |
| need CARAN per head squared |  |  | $\begin{aligned} & 1.687 * * * \\ & {[0.446]} \end{aligned}$ |  |  |  | (0) |  |  |  |  |
| all cause SYLLR exc gastro |  |  |  | $\begin{aligned} & -1.279 * * * \\ & {[0.333]} \end{aligned}$ |  |  |  |  |  |  |  |
| HIV need per head |  |  |  |  | $\begin{aligned} & 0.440^{* * *} \\ & {[0.025]} \end{aligned}$ |  |  |  |  |  |  |
| all cause SYLLR exc infect diseases |  |  |  |  | $\begin{aligned} & -0.543 * * \\ & {[0.249]} \end{aligned}$ |  |  |  |  |  |  |
| HIV need per head squared |  |  |  |  | $\begin{aligned} & 0.183 * * * \\ & {[0.021]} \end{aligned}$ |  |  |  |  |  |  |
| all cause SYLLR exc diabetes |  |  |  |  |  | $\begin{aligned} & -0,384^{*} \\ & 0,2181 \end{aligned}$ |  |  |  |  |  |
| diabetes prevalence rate 2007/8 |  |  |  |  | $l$ | $\begin{aligned} & 0.332 * * * \\ & {[0.123]} \end{aligned}$ |  |  |  |  |  |
| all cause SYLLR exc epilepsy |  |  |  |  | $\bigcirc$ |  | $\begin{aligned} & -0.259 \\ & {[0.223]} \end{aligned}$ |  |  |  |  |
| epilepsy prevalence rate 2007/8 |  |  |  |  | $\rangle \quad>$ |  | $\begin{aligned} & 0.571 * * * \\ & {[0.072]} \end{aligned}$ |  |  |  |  |
| all cause SYLLR exc renal |  |  |  |  | , |  |  | $\begin{gathered} -0.072 \\ {[0.168]} \end{gathered}$ |  |  |  |
| maternity need per head |  |  |  |  |  |  |  |  | $\begin{aligned} & 0.582^{* * *} \\ & {[0.098]} \end{aligned}$ |  |  |
| all cause SYLLR |  |  |  |  |  |  |  |  |  | $\begin{aligned} & -0.169 \\ & {[0.290]} \end{aligned}$ | $\begin{aligned} & -0.277 \\ & {[0.363]} \end{aligned}$ |
| lone pensioner households |  |  |  |  |  |  |  |  |  | $\begin{aligned} & -0.480 * * * \\ & {[0.182]} \end{aligned}$ |  |
| population working in agriculture |  |  |  |  |  |  |  |  |  |  | $\begin{aligned} & 0.132 * * * \\ & {[0.022]} \end{aligned}$ |
| Constant | $\begin{aligned} & 4.973 \\ & {[3.047]} \end{aligned}$ | $\begin{aligned} & 15.081 * * * \\ & {[3.303]} \end{aligned}$ | $\begin{aligned} & 4.986 * * \\ & {[2.342]} \end{aligned}$ | $\begin{aligned} & 7.488 * * * \\ & {[2.786]} \end{aligned}$ | $\begin{aligned} & -4.212^{* * *} \\ & {[1.034]} \end{aligned}$ | $\begin{aligned} & 3.555^{*} \\ & {[1.87} \end{aligned}$ | $\begin{aligned} & -1.684 \\ & {[1.130]} \end{aligned}$ | $\begin{aligned} & -2.675 \\ & {[2.562]} \end{aligned}$ | $\begin{aligned} & -1.222 \\ & {[1.388]} \end{aligned}$ | $\begin{aligned} & 1.413 \\ & {[1.373]} \end{aligned}$ | $\begin{aligned} & -5.960^{* * *} \\ & {[1.104]} \end{aligned}$ |
| Observations | 151 | 151 | 151 | 151 | 151 | 151 | 151 | 151 | 151 | 151 | 151 |
| Endogeneity test statistic | 20.985 | 19.454 | 11.612 | 15.477 |  | 2.846 | 4.958 |  |  | 0.060 | 1.769 |
| Endogeneity p-value | $4.63 \mathrm{e}-06$ | $1.03 \mathrm{e}-05$ | 0.000655 | $8.35 \mathrm{e}-05$ |  | 0.0916 | 0.0260 |  |  | 0.807 | 0.183 |
| Hansen-Sargan test statistic | 0.411 | 0.003 | 1.369 | 0.0201 |  | 0.510 | 2.748 |  |  | 1.091 | 1.121 |
| Hansen-Sargan p-value | 0.522 | 0.959 | 0.504 | 0.887 |  | 0.775 | 0.0974 |  |  | 0.296 | 0.571 |
| Shea's partial R-squared | 0.384 | 0.253 | - 0.398 | 0.325 |  | 0.402 | 0.518 |  |  | 0.416 | 0.364 |
| Kleibergen-Paap LM test statistic | 40.04 | 28.14 | 39.41 | 33.23 |  | 40.29 | 31.53 |  |  | 16.51 | 27.19 |
| Kleibergen-Paap p-value | 2.02e-09 | 7.76000 | $1.42 \mathrm{e}-08$ | ${ }^{6.09 \mathrm{e}-08}$ |  | $9.26 \mathrm{e}-09$ | $1.42 \mathrm{e}-07$ |  |  | 0.000260 | 5.37e-06 |
| Kleibergen-Paap F statistic | 51.44 | $29.097 \sim$ | 40.69 | 20.04 |  | 37.14 | 73.21 |  |  | 26.60 | 32.54 |
| Pesaran-Taylor/Ramsey test statisti | 2.262 | 00.0002 | ${ }_{0}^{0.0236}$ | 0.0341 | 0.721 | 2.351 | 0.619 | 1.297 | 1.018 | 1.757 | 0.193 |
| Pesaran-Taylor/Ramsey p-value | 0.133 | (0,988) | 0.878 | 0.854 | 0.541 | 0.125 | 0.432 | 0.278 | 0.387 | 0.185 | 0.660 |

[^67]Table B10.3: table showing cost of life and life year estimates using spend data for 2007/8 and outcome data for 2007/2008/2009 (assumes zero health gain for 13 programmes)


Table B10.4: table showing cost of life and life year estimates using spend data for 2007/8 and outcome data for 2007/2008/2009 (assumes zero health gain for PBC23 and 'average' gain for other 12 programmes)


## B11. Analysis of programme budgeting expenditure for 2008/9 and mortality data for 2008/2010

Outcome and expenditure models were estimated using updated data for expenditure (from 2007/8 to $2008 / 9$ ) and updated mortality data (from 2007/2008/2009 to 2008/2009 / 2010). Results for the outcome model are shown in Table B11.1 and results for the expenditure model are in Table B11.2. First stage regressions for these IV models can be found in Tables BA. 9 and BA. 10 in the annex.

## B11.1 Outcome models

Most of the outcome models in Table B11.1 contain just two variables: own programme expenditure and a measure of the need for health care. The latter is usually the measure of need as employed by the $\Omega$ Department of Health for resource allocation purposes and this incorporates the CARAN formula for acute services. For the respiratory disease programme we have added the square of the need measure to improve the model fit. In other PBCs (e.g., for the endocrine, metabolic and nutritional programme), we found that the all service measure of need performed poorly and we have replaced it with a more programme specific measure (e.g., the diabetes prevalence rate) or with a better performing proxy for need (e.g., the percentage of residents born outside the EU for maternity/neonate mortality).

The relevant statistical test suggests that expenditure is endogenous in six of the ten programmes but we have retained the IV estimates for the other four because they provide plausible results. The HansenSargen test suggests that the selected instruments are valid, and the Kleibergen-Paap LM statistic suggests that they are relevant (i.e., correlated with the endogenous regressor). The Kleibergen-Paap F statistic suggests that we do not have a problem with weak instruments (although the F statistic is marginally less than the conventional target value of ten in the genitor-urinary and infectious disease programmes). Finally, the Pesaran-Taylor reset test statistics reveal no evidence of mis-specification.

Results for the big four programmes are shown in the first five columns of Table B11.1. Two results are reported for the gastro-intestinal programme. The first of these (column 4) uses two instruments and so we report the instrument validity test statistic. However, one of these instruments is insignificant in the first-stage regression and, if we drop this instrument and re-estimate the model, we obtain the result in column 5 (but of course there is no instrument validity test statistic with this re-estimation). The removal of one instrument has little impact on the coefficient on expenditure but the Kleibergen-Paap F statistic is now much greater than ten.

In all of the big four programmes the need variable has a positive and significant effect on mortality, and expenditure has the anticipated negative effect. As we have noted before, the outcome results for the other programmes (in columns $6-10$ ) are similar to but more diverse than those for the big four programmes. This is to beanticipated because mortality is a much rarer outcome in these programmes than it is in the big foum programmes.

Expenditure has the anticipated negative effect on mortality in the endocrine problems programme and this is statistically significant. The all service measure of need is not relevant for this PBC; instead, we find that the diabetes prevalence rate is positively associated with mortality, as is a measure of deprivation (the IMD2007).

Expenditure has a negative but statistically insignificant impact on mortality from epilepsy in the neurological programme, and the all service indicator of the need for health care is positively and significantly associated with mortality in this programme.

Expenditure also has a negative but not statistically significant effect on mortality (from renal problems) in the genitor-urinary problems programme. The prevalence of lone parent households is positively associated with mortality.

Expenditure has the anticipated negative effect on mortality in the infectious disease programme and this is statistically significant. The all service measure of need is not relevant for this PBC; instead, we find
that a measure of need associated with HIV is positively associated with mortality, as is a measure of deprivation (the IMD2007).

Expenditure has the anticipated negative effect on mortality in the maternity \& neonates programme. In this PBC the coefficient on the generic all service measure of need is positive but not significant. It has been supplemented with two other indicators of deprivation - the proportion of residents born outside the EU and the proportion of those aged 16-74 without any qualifications - and both of these are positively associated with mortality.

Finally, we were unable to develop a plausible outcome model for the trauma and injuries programme.

Table B11.1: table showing uutcome models using spend data for 2008/9 (two MFFs) and mortality data for 2008/9/10


Note: robust standard errors in brackets, $* * * \mathrm{p}<0.01, * * \mathrm{p}<0.05, * \mathrm{p}<0.1$

## B11.2 Expenditure models

Most of the expenditure models in Table B11.2 contain just three variables: the PCT budget, a proxy for the own programme need for health care, and a proxy for the need for health care in other programmes.

The budget term is positive and statistically significant in ten of the eleven models.
The usual proxy for the own programme need for health care (i.e., the all service measure of need) is positive and significant in five of the eleven results. In a couple of programmes (respiratory disease and endocrine problems) we have added the squared value of need to improve the model fit and in both cases this term is positive and significant.

In some programmes (e.g., the endocrine PBC and the neurological PBC), we have replaced and/or supplemented the all service measure of need with a more programme specific measure (e.g., the diabetes and the epilepsy prevalence rates) and these usually have a positive and significant impact on expenditure.

In addition, in a couple of programmes we have used alternative proxies for own programme need (e.g., with the use of the Department of Health's measure of maternity need in the maternity/neonates expenditure equation and the use of HIV need in the infectious diseases programme).

For eight of the eleven programmes we have used the all cause mortality rate less the own programme mortality rate as the proxy for the need for health care in other programmes, and the coefficient on this term is negative in seven programmes and statistically significant in six of the seven. In three programmes -- maternity/neonates, GMS/PMS and trauma \& injuries programmes -- we have used the all cause mortality rate as the proxy for the need for health care in other programmes due to difficulties associated with the measurement of the own programme mortality rate. The coefficient on this term is negative but not significant in these three models.

The relevant statistical test suggests that expenditure is endogenous in five of the eleven programmes but we have retained the IV estimates for two further programmes (endocrine problems and maternity/neonates) because the IV estimator proyides more plausible results than the OLS estimator. In the other four programmes we report OLS results.

The Hansen-Sargen test suggests that the selected instruments are valid, and the Kleibergen-Paap LM statistic suggests that they are relevant (i.e., correlated with the endogenous regressor). The KleibergenPaap F statistic suggests that we do not have a problem with weak instruments. Finally, the PesaranTaylor reset test statistics and the Ramsey reset $F$ statistics reveal no evidence of model mis-specification.

Table B11.2: table showing expenditure models using spend data for 2008/9 (two MFFs) and mortality data for 2008/9/10



[^68]
## B11.3 Calculation of the cost of a life and life year

Expenditure and outcome elasticities for our preferred models are shown in Table B11.3 (see columns D and H ) and these are used to calculate the cost of a life and the cost of a life year, both for individual programmes and for all programmes collectively.

Again, column N reports the cost per life gained and column U reports the cost per life year gained. From the latter we can see that the cost per life year gained has increased slightly compared with that using the previous expenditure and mortality data set (i.e., for 2007 and 2007/8/9 respectively): it has increased from $£ 13,830$ to $£ 14,650$ for the big four programmes and from $£ 28,983$ to $£ 30,883$ for all ten programmes with a mortality-based outcome indicator.

If we assume that the other 13 programmes offer no health gain, then the cost per life year across all PCT expenditure has increased from $£ 82,765$ in $2007 / 8$ to $£ 84,974$ in 2008/9.

In addition, Table B11.4 shows that if we assume that PBC23 generates a zero health gain and that the gain attributable to the remaining 12 programmes is, on average, the same as that attributable to those with a mortality outcome measure, then the cost of a life year across all programmes in 2008/9 is $£ 33,333$. This is a $5 \%$ increase on the figure $(£ 31,846)$ for the previous year.

Table B11.3: table showing cost of life and life year estimates using spend data for 2008/9 and outcome data for 2008/2009/2010 (assumes zero health gain for 13 programmes)


Table B11.4: table showing cost of life and life year estimates using spend data for 2008/9 and outcome data for 2008/2009/2010 (assumes zero health gain for PBC23 and average gain for other 12 programmes)


All 23 programme spend
$\%$ change in budget
37 proportionate chang
38 Change in budget

| $2008 / 9$ | $2007 / 8$ | $2006 / 7$ | $2005 / 6$ |
| :--- | :--- | :--- | :--- |
| $£ 78,398$ | $£ .73,657$ | $£ .67,896$ | $£ 64,310$ |
| 1.00 | 1.00 | 1.00 | 1 |
| 0.01 | 0.01 | 0.01 | 0.01 |
| $£ .783 .98$ | $£ 736.57$ | $£ .678 .96$ | $£ .643 .10$ |

Note that the annual mortality figures reported in cells $\mathrm{G} 7 \& \mathrm{G} 8$ and G 17 \& G 18 are identical because we do not have mortality data for 2002/04
Note that the coverage of the YLL data relative to the spend data for trauma \& injuries is assumed to take a value of 1.0 (that is, the ICD coverage is the same). Note that, for expenditure in 2007/8, the neonate category has been merged with maternity to obtain plausible outcome and expenditure models.
Note that, for expenditure in $2007 / 8$, the neonate category has been merged with maternity to obtain plausible outcome and expenciture
Note that the adjustment for the coverage of the mortality data relative to the spend data uses deaths under age 75 in England in 2008 .

## B11.4 Comparing the cost of life year estimates associated with different data sets

Table B11.5 presents expenditure and outcome elasticities for the five combinations of expenditure and outcome data that have been used to estimate our model. It also reports the corresponding unadjusted cost of life year estimates (i.e., estimates that are unadjusted for the mismatch in the ICD10 coverage of the expenditure and mortality data). It is clear from this Table (see row 13) that the (unadjusted) cost of a life year for the ten programmes with a mortality based outcome indicator fluctuated around $£ 22,000$ for the first three sets of estimations (see columns M-O). However, using the two most recent sets of expenditure data (i.e., for 2007/8 and then for 2008/9), the figures in the table suggest that this cost has increased to about $£ 38,000$.

What are the proximate causes of this increase? Recall that the cost of a life year is calculated as
the change in expenditure associated with a $1 \%$ budget increase
the change in the number of life years lost associated with this increase
For 2006/7 (using mortality data for $2006 / 7 / 8$ ) and for the ten programmes with a mortality based outcome indicator, the cost of a life year is calculated as

$$
\begin{equation*}
(£ .32,911 \mathrm{~m} * 0.01 * 0.561) /(1,667,259 * 0.01 * 0.465)=£ .184 .53 \mathrm{~m} / 7,7(60)=£_{2} 23,780 \tag{11.1}
\end{equation*}
$$

For 2007/8 (using mortality data for 2007/8/9) and for the ten programmes with a mortality based outcome indicator, the cost of a life year is calculated as

$$
\begin{equation*}
(£ 34,434 \mathrm{~m} * 0.01 * 0.749) /(1,635,716 * 0.01 * 0.414)=£(257.94 \mathrm{~m} / 6,768=£ 38,110 \tag{11.2}
\end{equation*}
$$

It is clear that the $60 \%$ increase in the cost of a life year between $2006 / 7$ and $2007 / 8$ is largely attributable (a) to the $40 \%$ increase in the additional expenditure directed towards these 10 programmes following the $1 \%$ budget increase and (b) to the $12 \%$ decline in the number of life years gained associated with this increase in expenditure.

The rise in the share of the budget increase directed towards these programmes can be attributed to the increase in the expenditure elasticity associated with these ten programmes (up from 0.561 to 0.749 ). The decrease in the number of years of life gained can be largely attributed to the $12 \%$ decline in the outcome elasticity associated with these programmes, down from -0.877 to -0.778 (see row 13 , columns J and K of Table B11.5). ${ }^{35}$ However, it is not clear why such rather dramatic changes should have taken place.

Table B11.6 presents cost of life year estimates (adjusted for the mismatch in the ICD10 coverage of the expenditure and mortality data) for various combinations of programmes. These reveal similar increases in the cost of a life year between 2006/7 on the one hand and 2007/8 and 2008/9 on the other. The cost of a life year moreased from $£ 19,965$ in 2006/7 to $£ 28,983$ in $2007 / 8$ for the ten programmes with mortality rate, an increase of $45 \%$; and it increased from $£ 22,565$ to $£ 31,846$ for all programmes if we assume a zero health gain in PBC23 and the same gain in the other 12 programmes as in the ten with a mortality rate (an increase of $41 \%$ ).

One reason for this apparent step change in the cost of a life year might be the adjustment that was made to the methodology for the collection of the 2007/8 programme budgeting data. In previous years expenditure that was not directly attributable to a particular programme category was apportioned using admitted patient care percentages. ${ }^{36}$ In other words, if $x \%$ of total admitted patient care expenditure was

[^69]allocated to PBC 1, then $x \%$ of all expenditure that was not directly attributable to a particular programme category was also allocated to PBC 1. With effect from 2007/8, however, NHS organisations were asked to select an appropriate basis for the apportionment of this non-programme specific expenditure and that, where no reasonable basis existed, such expenditure was to be allocated to the 'Other - Miscellaneous' (PBC 23X) category.

The Department of Health estimates that this allocation rule change increased the amount of expenditure attributed to PBC 23X by $£ 700$ million. It will also, of course, have reduced expenditure across other programmes by the same amount in total. However, not all programmes will have been equally affected; PBCs that are more heavily inpatient based would have 'lost' expenditure while others, such as learning disabilities, social care, and mental health, will have 'lost' considerably less. In addition, not all PCTs will have been equally affected because each will have employed different apportionment rules for the nonprogramme specific expenditure. [38]

Although this allocation rule change has considerably increased the estimated cost of a life year, we believe that this rule change has led to a more accurate allocation of expenditure across PBCs, and that the more recent estimates of the cost of a life year (for 2007/8 and 2008/9) are more accurate than those for the earlier years (for 2005/6 and 2006/7).

## B11.5 Adjusting the cost of a life year estimates to constant prices

The cost of a life year estimates presented above are all at current prices. To put them on a constant price basis, we need an index of pay and price inflation for the labour and goods/services purchased by the NHS. Curtis [41] reports a pay and prices index for Hospital and Community Health Services and this implies an inflation rate of $3.7 \%$ in $2006 / 7,2.9 \%$ in $2007 / 8$, and $3.9 \%$ in $2008 / 9 .{ }^{37}$ If we assume that similar inflation rates also apply to the purchase of pharmaceuticals and the provision of primary care (items that are excluded from the HCHS index), then we can use these figures to put the cost of a life year estimates on a constant price basis.

For example, if we assume that PBC23 generates a zero health gain and that the gain attributable to the 12 programmes without a mortality indicator is, on average, the same as that attributable to those with $\AA$ mortality outcome measure, then the cost of a life year across all programmes in 2008/9 is $£ 33,333$ at current (2008/9) prices. The cost for $2007 / 8$ is $£ 31,846$ at current $(2007 / 8)$ prices or $£ 33,088$ at constant (2008/9) prices, and the figure for $2006 / 7$ is $£ 22,565$ at current $(2006 / 7)$ prices or $£ 24,125$ at constant $(2008 / 9)$ prices. The conversion of the costs from a current to constant price basis has relatiyely little impact because the inflation rate over the relevant period is quite small.

## B11.6 Summary and conclusion

In this section we have estimated outcome and expenditure models using PB data for 2008/9 and mortality data for $2008 / 9 / 10$. The cost of an additional life year for all ten programmes with a mortality based outcome is $f 30,883$. This is similar to the comparable figure $(28,983)$ for the previous year (i.e., using expenditure data for $2007 / 8$ and mortality data for $2007 / 2008 / 2009$ ). If we assume that PBC23 generates a zero health gain and that the gain attributable to the 12 programmes without a mortality indicator is, on average, the same as that attributable to those with a mortality outcome measure, then the cost of a life year across all programmes in $2008 / 9$ is $£ 33,333$ and this too is similar to the figure for the previous year ( $£ 31,846$ ).

We have also identified a pay and prices index that can be used to put the estimated costs on a constant price basis. This index has recorded an annual inflation rate of about $3.5 \%$ since 2005/6.

There appears to have been a step change in the cost of an additional life year. The cost of a life year estimates are very similar up to and including 2006/7, and they are very similar for 2007/8 and 2008/9. However, there is a substantial difference between the figures for 2004/5, 2005/6 and 2006/7 on the one hand, and for $2007 / 8$ and $2008 / 9$ on the other. The reason for this step change is not obvious but it might be due to changes in the algorithm used by the Department of Health to allocate non-admitted patient care activity to budget categories. Although this allocation rule change has considerably increased the estimated cost of a life year, we believe that this rule change has led to a more accurate allocation of expenditure across (PBCS, and that the more recent estimates of the cost of a life year (for 2007/8 and 2008/9) are more accurate than those for the earlier years (for 2005/6 and 2006/7).

[^70]Table B11.5: table showing expenditure and outcome elasticities for five combinations of expenditure and outcome data, and corresponding (unadjusted) cost of life year estimates

| A | B | C | D | E | F | G |  | I | J | K | L |  |  | O | P | Q |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | spend elasticities |  |  |  |  | outcome elasticities |  |  |  |  | costrof an additional life year (unadjusted for YLL coverage) |  |  |  |  |
|  |  |  |  |  |  | (e) using |  |  |  |  | (e) using | sing |  |  |  |  |
|  |  | spend for | spend for | spend for | spend for | $\begin{aligned} & \text { ppend for } \\ & 2008 \text { and } \end{aligned}$ | spend for | spend for | spend for | spend for | $\begin{aligned} & \text { spend for } \\ & 2008 \text { and } \end{aligned}$ | spend for | spend for | (c) using spend for | spend for | (e) using spend for |
|  |  | 2005 and mortality for 2002/4 | 2006 and mortality for 2004/6 | 2006 and mortality for 2006/8 | 2007 and mortality for 2007/9 | mortality <br> for <br> 2008/10 | 2005 and mortality for 2002/4 | 2006 and mortality for 2004/6 | 2006 and mortality for 2006/8 | 2007 and mortality for 2007/ | mortality 2008/10 | 2005 and mortality for 2002/4 | 2006 and mortality for 2004/6 | 2006 and mortality for 2006/8 | 2007 and mortality for 2007/9 | 2008 and mortality for 2008/10 |
| 1 | Cancer | 0.968 | 0.548 | 0.465 | 0.890 | 0.525 | -0.394 | -0.337 | -0.342 | -0.365 - 307 |  | £ 13,741 | £16,518 | £16,383 | £17,165 | £21,802 |
| 2 | Circulatory problems | 0.682 | 0.701 | 0.540 | 0.293 | 0.648 | -1.370 | -1.447 | -1.434 | $-1.277$ | -1.319 | £ 8,328 | ¢ 8,725 | £ ¢, 466 | £11,315 | £.11,779 |
| 3 | Respiratory problems | 0.849 | 0.718 | 0.679 | 0.536 | 0.652 | -1.574 | -3.507 | -2.622 |  | -1.808 | $£^{20,601}$ | £ 8,747 | £11,593 | $£ 14,798$ | $£_{21,307}$ |
| 4 | Gastro-intestinal problems | 0.772 | 0.667 | 0.446 | 0.622 | 0.456 | -2.018 | -2.137 | $-1.536$ | 1.328 | -1.364 | £18,303 | £,15,795 | $£_{20,892}$ | £25,034 | £25,662 |
| 5 | All big four PBCs | 0.801 | 0.660 | 0.528 | 0.559 | 0.579 | -0.941 | -1.083 | $-0.965$ | -0.872 | -0.825 | £12,855 | $£ 10,783$ | £12,333 | $£ .16,345$ | £16,688 |
| 6 | Infectious diseases | 0.742 | 0.731 | 0.792 | 1.436 | 1.545 | -0.152 | -0.030 | -0.047 | -0.548 | -0.504 | £215,054 | $£ 1,036,377$ | $£ 630,798$ | £57,742 | £.71,432 |
| 7 | Endocrine problems | 0.425 | 0.966 | 0.953 | 0.264 | 0.484 | -0.244 | . 812 | -. 842 | -0.566 | -1.170 | £ 371,601 | £.112,882 | £114,416 | £.190,745 | £104,008 |
| 8 | Neurological problems | 1.111 | 0.648 | 0.616 | 1.035 | 0.98 | -0.182 | 098 | . 112 | -0.339 | -0.417 | f.503,201 | £1,241,253 | £1,129,960 | £.431,749 | £.388,267 |
| 9 | Genito-urinary problems | 1.041 | 0.837 | 0.912 | 1.004 | 0.697 | -0.034 | 0.07 | -0.051 | -1.855 | -1.615 | $£^{29,144,918}$ | £12,384,965 | £20,421,090 | ¢652,096 | £ 877,038 |
| 10 | Trauma \& injuries* | 0.627 | 0.617 | 0.358 | 1.686 | 1.344 | . 332 | . 527 | 0 | -0.369 | 0 | £282,132 | £548,767 | n/a | £ 1,115,197 | $\mathrm{n} / \mathrm{a}$ |
| 11 | Maternity \& neonates* | 0.388 | 0.601 | 0.224 | 0.514 | 0.975 | -0,237 | $-0.035$ | -0.482 | -0.110 | -0.125 | £,17,490 | £631,700 | $¢_{4} 45,158$ | £.204,168 | £198,939 |
| 12 | All small six PBCs | 0.780 | 0.717 | 0.596 | 0.961 | 0.962 | . 262 | -0.122 | -0.392 | -0.254 | -0.300 | £295,074 | £449,706 | $£_{2}^{258,046}$ | £274,309 | £254,794 |
| 13 | All 10 PBCs with mortality | 0.792 | 0.687 | 0.561 | 0.749 | 0.762 | -0.84 | -0.940 | -0.877 | -0.778 | -0.747 | £21,256 | £20,893 | £23,780 | £38,110 | £.38,328 |
| 14 | All 23 PBCs assuming zero gain in PBCs without mortality indicator |  |  |  |  |  |  |  |  |  |  | £56,799 | $\oint 62,718$ | $£ .87,494$ | $£ 108,829$ | $£ 105,460$ |
| 15 | GMS/PMS | 0.926 | 0.759 | 0.739 | 0.563 | 0.494 | $\mathrm{n} / \mathrm{a}$ | n/a | n/a | n/a | n/a | n/a | n/a | n/a | n/a | n/a |
| 16 | All 23 PBCs assuming zero | in in PBC 23 | t average gai | other PBC | without a mor | lity indicat |  |  |  |  |  | $¢_{2} 24,200$ | £23,697 | $£_{26,876}$ | £ 411,875 | £ 41,369 |

Notes: (i) that the spend and outcome elasticities reported for groups of progranmes are the implied elasticites calculated from the totals for the relevant individual programmes (i.e., group spend elasticity $=\sum$ (PBC spend $* \mathrm{PBC}$ spend elasticity) $/ \sum \mathrm{PBC}$ spend, and group outcome elasticity $=\sum\left(\mathrm{PBC}\right.$ mortality $* \mathrm{PBC}$ outcome elasticity) $\sum \sum \mathrm{PBC}$ mortality). For the purpose of the calculation of the group outcome elasticity, we have used the years of life lost as the mortality indicator. The group elasticities are directly comparable with the individual programme elasticities as both exclude the impact of the relevant budget elasticities.
(ii) for each individual programme: the cost of an additional life year $=\%$ change in spend $*$ annual spend /(outcome elasticity*annual life years lost)
(iii) for a group of programmes: the overall cost of antadditional life year $=\sum$ (annual spend ${ }^{*}$ spend elasticity) $/ \sum$ (spend elasticity ${ }^{*}$ outcome elasticity ${ }^{*}$ annual life years lost)
(iv) that the results using expenditure for 2006/7 and mortality for 2004/6 incorporate MFFs for HCHS and prescribing (see Tables B8.17 and B8.18).

Table B11.6: table showing adjusted cost of life year estimates for various combinations of programmes


Note that the figures for 2006/7 relate to the use of mortality for 2006 /2007/2008 combined.

## B12. Summary and concluding remarks

The findings presented in this report build on four previous studies[2-5]. These studies and the results presented here draw on the availability of two new data sets to obtain empirical estimates of the relationship between mortality and expenditure across all English local health authorities.

In this research we have extended the previous studies in several ways. First, we have derived plausible outcome and expenditure models for a larger number of programmes (ten) than previous studies. The cost of a life year across all ten programmes with a mortality based outcome indicator using expenditure data for $2006 / 7$ and mortality data for 2004/5/6 is $£ 20,893$.

Second, we relate expenditure in time period $t$ to mortality in that period $(t)$ and in the next two periods ( $t+1$ and $t+2$ ). In other words, we assume that the health benefits associated with expenditure occur either in the same period as the expenditure or in the next two periods. This is an improvement on past practice where data constraints forced researchers to relate expenditure to the current and two previous periods. When we re-estimated our models having replaced mortality data for 2004/5/6 with that the $2006 / 7 / 8$, we found that the cost of a life year across the ten programmes with a mortality based outcome indicator using expenditure data for 2006/7 is $£ 23,780$ (up from $£ 20,893$, an increase of $14 \%$ ).

Third, we have noted the mismatch in the ICD10 coverage of the expenditure and mortality data. If we adjust the calculation of the cost of a life year for $2006 / 7$ for this mismatch then the cost of a life year across the ten programmes with a mortality based outcome indicator declines from $£ 23,780$ to $£ 19,965$ (a decrease of $16 \%$ ).

Fourth, previous estimates of the cost of a life year have been for individual programmes of care. In this report we have presented estimates of the cost of a life year for an enlarged number of programmes and, with the aid of assumptions about the productivity (health gain) of programmes without a meaningful mortality-based outcome indicator, we have extended our individual programme estimates to incorporate expenditure across all programmes of care. Thus for 2006/7, the cost of a life year for those PBCs with a mortality based outcome indicator is $£ 19,965$. If we assume that (a) that the health gain associated with PBC23, which includes primary care and workforce training expenditure, are reflected in the mortality rates for disease specific programmes and (b) that the average health gain across the other programmes without a mortality based outcome indicator is the same as that for those PBCs with a mortality based outcome indicator, then the cost of life year across all programmes is $£ 22,565$.

Fifth, we have extended our cost of life year estimates beyond 2006/7. Re-estimation of our model using budgeting expenditure for $2007 / 8$ generates an all programme cost of a life year estimate of $£ 31,846$, and re-estimation of our model using budgeting expenditure for $2008 / 9$ generates a similar cost of a life year estimate ( $£ 33,333$ ). Together, the last two estimates suggest that there has been step change in the cost of a life year, and that this appears to have occurred between 2006/7 and 2007/8. The cost of a life year estimates are very similar up to and including 2006/7, and they are very similar for 2007/8 and 2008/9. However, there is a substantial difference between the figures for 2004/5, 2005/6 and 2006/7 on the one hand (at about $f_{2} 2 \mathrm{k}$ ), and for $2007 / 8$ and $2008 / 9$ on the other (at about $£ 33 \mathrm{k}$ ). The reason for this step change is not obvious but it might be due to changes in the algorithm used by the Department of Health to allocate non-admitted patient care activity to budget categories. Although this allocation rule change has considerably increased the estimated cost of a life year, we believe that this rule change has led to a more accurate allocation of expenditure across PBCs, and that the more recent estimates of the cost of a lifeyear (for 2007/8 and 2008/9) are more accurate than those for the earlier years (for 2005/6 and 2006/7).

Virtually all of the cost of a life year estimates presented in this report are at current prices. However, it is possible to put them on a constant price basis using the Hospital and Community Health Services pay and prices index [41]. For 2006/7, 2007/8 and 2008/9 this index recorded an annual rate of inflation of about $3.5 \%$ and so the impact of this constant price adjustment is fairly minimal. For example, if we assume that PBC23 generates a zero health gain and that the gain attributable to the 12 programmes without a mortality indicator is, on average, the same as that attributable to those with a mortality
outcome measure, then the cost of a life year across all programmes at constant 2008/9 prices is $£ 33,333$ for 2008/9, $£ 33,088$ for $2007 / 8$, and $£ 24,125$ for 2006/7.

Finally, although previous results and our current models 'pass' the appropriate statistical tests and, in particular, the Hansen-Sargen test for valid instruments, we are aware that this test might be unable to detect the presence of invalid instruments in some (albeit rather restrictive) circumstances. Responding to this, several studies have suggested that researchers using IV techniques should subject the estimated coefficient on the endogenous variable to a sensitivity analysis. We do precisely this for the outcome equation for each of the big four models. This sensitivity analysis reveals that uncertainty associated with instrument validity has little effect on our estimate of the cost of a life year but it does increase the degree of uncertainty associated with this estimate.

We recognize that this study has a number of limitations. The cost of an additional life year estimates for those programmes with a mortality-based outcome indicator are unadjusted for the quality of life during the additional year; the quoted costs will be an under-estimate of the QALY-adjusted cost of alife year to the extent that additional life years are not in perfect health. In previous studies we havenoted that a rudimentary adjustment for this issue using HODaR data increased the cost of a life year by about $50 \%$ to $60 \% .[2,5]$

At the same time, however, the estimated costs will exaggerate the cost of an additional QALY-adjusted year for those programmes with a mortality-based outcome indicator because they ignore any health benefits that are not associated with a reduction in mortality. In other words, expenditure that improves the quality of life (e.g., cancer palliative care) but which does not extend the length of life is implicitly given a zero health gain value.

In addition, the expenditure data relates to expenditure on all patients whereas the mortality data is based on a life expectancy of 75 years. Thus implicitly our calculations attribute a zero health gain to all expenditure on those aged over 75. To illustrate the magnitude of the potential health gain ignored by this restriction, note that in a recent study of costs associated with all inpatient and outpatient activity (excluding mental health), those aged over 75 years accounted for $25 \%$ of all costs in 2007/8[34] for details of this study).

Moreover, our cost of a life year estimates are based on the assumption that any Departmental budgetary change falls entirely on PCTs. Although PCTs account for most of the Department of Health's budget, non-PCTs still accounted for $15 \%$ of the budget in 2006/7. Because we have no information on how any budgetary change would be split between PCTs and non-PCTs, we have assumed that that any Departmental budgetary change falls entirely on PCTs. If, on the other hand, the non-PCT budget is responsive to changes in the Department's budget then our cost of a life year estimates will be too low. If the non-PCT budget is as responsive as the PCT budget, then our cost of a life year estimate for 2006/7 will be increased by $1757 \%$ (that is, from $£ 22,565$ to $£ 26,553$ ).

The results presented in this study are all from the estimation of the relationship between expenditure and mortality asing data for a single time period. With the availability of several years of data for both expenditure and mortality, we wanted to estimate a panel data model because a panel can offer advantages overa gne period model (e.g., it is better able to handle any unobserved heterogeneity across PCTs). However, most of the instruments employed here are based on the 2001 Census and thus estimation of a panel model will not be possible until these instruments become time variant; this should occur later this year with release of the 2011 Census data at PCT level. This is one piece of work that we intend to pursue in the near future.

## References

1. Health, D.o., A new value-based approach to the pricing of branded medicines: A consultation. 2010.
2. Martin, S., N. Rice, and P. Smith, Does health care spending improve health outcomes? Journal of Health Economics, 2008a: p. 826-842.
3. Martin, S., N. Rice, and P. Smith, The link. between health care spending and bealth outcomes for the new English Primary Care Trusts. CHE Research Paper 42, 2008b.
4. Martin, S., N. Rice, and P. Smith, Panel data estimates of the link. between bealth care spending and bealth outcomes for English Primary Care Trusts. Mimeo, 2010.
5. Martin, S., N. Rice, and P. Smith, Comparing costs and outcomes across programmes of health care. Health Economics, 2012: p. 316-337.
6. Gerdtham, U. and B. Jonsson, International comparisons of health expenditure' in A. Culyer and J. Newhouse (eds). Handbook of Health Economics, 2000. Elsevier Amsterdam.
7. Fisher, E.S. and H.G. Welch, Avoiding the unintended consequences of growth in medical care - How might more be worse? Jama-Journal of the American Medical Association, 1999. 281(5): p. 446-453.
8. Nolte, E. and M. McKee, Does health care save lives? The Nuffield Trust, London, 2004.
9. Cochrane, A.L., A.S.S. Leger, and F. Moore, Health service "input" and mortality "output" in developed countries (Reprinted from Journal of Epidemiology and Community Health vol 32, pg 200-205, 1968). Journal of Epidemiology and Community Health, 1997. 51(4): p. 344-348.
10. Young, F.W., An explanation of the persistent doctor-mortality association. Journal of Epidemiology and Community Health, 2001. 55(2): p. 80-84.
11. St Leger, S., The anomaly that finally went away? Journal of Epidemiology and Community Health, 2001. 55(2): p. 79-79.
12. Gravelle, H.S.E. and M.E. Backhouse, International crosssection analysis of the determination of mortality. Social Science \& Medicine, 1987. 25(5): p. 427-441.
13. Cremieux, P.Y., P. Ouellette, and C. Pilon, Health care spending as determinants of health outcomes. Health Economics, 1999. 8(7): p. 627-639.
14. Or, Z., Exploring the effects of health care on mortality across OECD countries. OECD Labour Market and Social Policy Occasional Paper 46, OECD, Paris., 2001.
15. Nixon, J. and P. Ulmann, The relationship between health care expenditure and health outcomes. Evidence and caveats for a causal link. The European journal of health economics : HEPAC : health economics in prevention and care, 2006. 7(1): p. 7-18.
16. Health, D.o., NHS finance manual. December 2005 edition. See bttp:// www.dh.gov.uk/ assetRoot/04/13/18/26/04131826.pdf. 2005a.
17. Smith, P.C., N. Rice, and R. Carr-Hill, Capitation funding in the public sector. Journal of the Royal Statistical Society Series a-Statistics in Society, 2001. 164: p. 217-241.
18. Health, D.o., Recurrent resource allocations: 2006/07, 2007/08. Department of Health, London, 2005c.
19. Office, N.A. Good governance report: review of programme budgeting. London: NAO, 2008.
20. Appleby, J., Harrison, T., Foot, C., Smith, A. and Gilmour, S. , Explaining variations in primary care trusts' spending on cancer services. The King's Fund, London, 2011.
21. Baam, C.F., M.E. Schaffer, and S. Stillman, ivreg2: Stata module for extended instrumental variables/2SLS, GMM and $A C / H A C, L I M L$ and k-class regression. See bttp:// ideas.repec.org/c/ boc/ bocode/s425401. btml. 2010.
22. Shea, J., Instrumental relevance in multivariate linear models: a simple measure. Review of Economics and Statistics, 1997. 79: p. 348-352.
23. Bound, J., D.A. Jaeger, and R.M. Baker, Problems with instrumental variables estimation when the correlation between the instruments and the endogenous explanatory variable is weak. Journal of the American Statistical Association, 1995. 90(430): p. 443-450.
24. Staiger, D. and J.H. Stock, Instrumental variables regression with weak instruments. Econometrica, 1997. 65(3): p. 557-586.
25. Stock, J.H. and M. Yogo, Testing for weak instruments in linear IV regression. NBER Technical Working Paper 284, 2002.
26. Cragg, J.G. and S.G. Donald, Testing identifiability and specification in instrumental variable models. Econometric Theory, 1993. 9(2): p. 222-240.
27. Ramsey, J.B., Tests for specification errors in classical linear least-squares regression analysis. Journal of the Royal Statistical Society Series B-Statistical Methodology, 1969. 31(2): p. 350-\&.
28. Pesaran, M.H. and L.W. Taylor, Diagnostics for IV regressions. Oxford Bulletin of Economics and Statistics, 1999. 61(2): p. 255-+.
29. Wooldridge, J., Econometric analysis of cross section and panel data. The MIT Press. Cambridge, 2002.
30. Durbin, J., Errors in variables. Review of the International Statistical Institute, 1954. 22: p. 23-32.
31. Health, D.o., Payment by results: tariff information. Department of Health, London, 2007.
32. Health, D.o., PCT recurrent revenue allocations exposition book: 2009/10 and 2010/11. Department of Health, London, 2009.
33. Health, D.o., Unified exposition book: 2003/04, 2004/05 and 2005/06 PCT revenue resource limits. Department of Health, London, 2005b.
34. Dixon, J., et al., A person based formula for allocating commissioning funds to general practices in England: development of a statistical model. British Medical Journal, 2011. 343.
35. Conley, T.G., C.B. Hansen, and P.E. Rossi, Plausibly exogenous. Review of Economics and Statistics, 2012. 94(1): p. 260-272.
36. Small, D.S., Sensitivity analysis for instrumental variables regression with overidentifying restrictions. Journal of the American Statistical Association, 2007. 102(479): p. 1049-1058.
37. Kovandic, T., M. Schaffer, and G. Kleck, Estimating the causal effect of gun prevalence on bomicide rates: a local average treatment effect approach. IZA Bonn. Discussion paper 3589, 2008.
38. Health, D.o., Personal communications. 2012.
39. Martin, S. and P. Smith, How good at commissioning bealth are English primary care trusts? A preliminary statistical analysis. Report to the Health Foundation, 2009.
40. Schaffer, M.E., Personal communication. 2011.
41. Curtis, L., Unit costs of health and social care 2011. PSSRU, Oniversity of Kent., 2011.

## Annex

Table BA.1: table showing national (all PCT) expenditure per head (£) and growth in expenditure (\%) by PBC group and sub-group, 2003/4-2008/9


| 7 a | Chronic Pain |  |  |  | 19.31 | 22.12 | 15 | 22.79 | 3 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 7 x | Neurological (Other) |  |  |  | 35.96 | 40.31 | 12 | 44.85 | 11 |
| 8 | Problems of Vision | 24.61 | 27.65 | 28.24 | 26.97 | 30.69 | 14 | 32.95 |  |
| 9 | Problems of Hearing | 5.73 | 6.32 | 6.27 | 6.21 | 8.07 | 30 | 8.16 |  |
| 10 | Problems of Circulation | 110.12 | 122.37 | 124.28 | 122.06 | 124.77 | 2 | 129.94 | 4 |
| 10 a | Coronary Heart Disease |  |  |  | 38.91 | 40.32 | 4 | 41.20 | 2 |
| 10b | Cerebrovascular disease |  |  |  | 16.05 | 17.30 | 8 | 19.35 | 12 |
| 10c | Problems of Rhythm |  |  |  | 7.22 | 8.21 | 14 | 8.43 | 3 |
| 10x | Problems of circulation (Other) |  |  |  | 59.88 | 58.95 |  | 60.96 | 3 |
| 11 | Problems of the Respiratory System | 54.60 | 62.71 | 69.56 | 65.07 | 67.68 |  | 77.97 | 15 |
| 11a | Obstructive Airways Disease |  |  |  | 10.64 | 10.64 |  | 12.70 | 19 |
| 11b | Asthma |  |  |  | 14.04 | 15.73 | 12 | 16.99 | 8 |
| 11x | Problems of the respiratory system, other |  |  |  | 40.40 | (41. 31 | 2 | 48.27 | 17 |
| 12 | Dental Problems | 10.78 | 13.55 | 24.91 | 51.93 | 59.45 | 14 | 62.44 | 5 |
| 13 | Problems of Gastro Intestinal System | 63.56 | 73.22 | 81.30 | 73.30 | 75.05 | 2 | 77.89 | 4 |
| 13a | Upper GI |  |  |  | 19.88 | 19.51 | -2 | 19.89 | 2 |
| 13b | Lower GI |  |  |  | 20.46 | 21.92 | 7 | 22.63 | 3 |
| 13 c | Hepatobiliary |  |  |  | 11.26 | 12.23 | 9 | 12.90 | 5 |
| 13 x | Problmes of the gastro intestinal system |  |  | 2 | 21.69 | 21.39 | -1 | 22.46 | 5 |
| 14 | Problems of the Skin 20.98Burns |  |  |  | 28.31 | 30.41 | 7 | 32.34 | 6 |
| 14 a |  |  |  |  | 1.08 | 1.56 | 44 | 1.02 | -34 |
| 14x | Problems of the Skin |  |  |  | 27.23 | 28.86 | 6 | 31.32 | 9 |
| 15 | Problems of Musculo Skeletal System Problems due to Trauma and Injuries | 61.36 | 71.7 | 74.74 | 66.75 | 75.91 | 14 | 79.68 | 5 |
| 16 |  | 62.31 | 72.13 | 76.41 | 57.29 | 57.56 | 0 | 63.54 | 10 |
| 17 | Problems of Genito Urinary System | 55.32 | 62.38 | 67.38 | 68.98 | 67.83 | -2 | 73.78 | 9 |
| 17 a | Genital tract problems | ) 2 |  |  | 19.33 | 18.80 | -3 | 19.36 | 3 |
| 17 b | Renal problems |  |  |  | 21.54 | 19.74 | -8 | 22.29 | 13 |
| 17c | STD |  |  |  | 4.26 | 4.71 | 10 | 5.43 | 15 |
| 17x | Problems of Genito Urinary system, other |  |  |  | 23.85 | 24.58 | 3 | 26.69 | 9 |
| 18 | Maternity and Reproductive Health | 52.28 | 55.04 | 60.42 | 57.64 | 57.09 | -1 | 60.44 | 6 |
| 19 | Conditions of Neonates | 11.72 | 13.93 | 13.42 | 13.17 | 15.15 | 15 | 17.23 | 14 |
| 20 | Adverse effects and poisoning | 9.68 | 12.32 | 14.25 | 14.59 | 15.84 | 9 | 18.31 | 16 |
| 20 a | Unintended consequences of treatment |  |  |  | 10.54 | 12.14 | 15 | 12.96 | 7 |
| 20b | Poisoning |  |  |  | 2.13 | 2.44 | 15 | 2.91 | 19 |



20x
21
21a
21b
Poisoning and adverse effects
Healthy Individuals
NSF Prevention programme
NSF Mental health prevention
Healthy Individuals (Other)
Social Care Needs
Other
GMS/PMS
Training (WDCs)
23x Miscellaneous
1 to 23 All PBCs
$052.12 \quad 1199.60 \quad 1307.76$
$1345.10 \quad 1452.918$
1530.595

Notes.
(i) The population figures for $2003 / 4,2004 / 5$ and $2005 / 6$ are identical (the total for England is 49,175,998)
(ii) The corresponding figure for $2006 / 7$ is $50,476,231$, for $2007 / 8$ it is $50,695,989$, and for $2008 / 9$ it is $51,220,531$.
(iii) The spend per head figures are calculated by summing expenditure across all PCTs and dividing byt the national population.

Table BA.2: table showing set of socio-economic indicators available as potential instruments in the IV estimation

| Indicator name | Short description | Long description |
| :---: | :---: | :---: |
|  | Residents born outside the European | Residents born outside the European Union divided by all residents (census cell definition: |
| BORNEXEU | Union | KS005008/KS005001) |
| WHITEEG | Population in white ethnic group | Population in white ethnic group divided by total population (KS006002+KS006003+KS006004)/KS006001 |
| PCWALLTI | $\mathrm{Pc}$ | Proportion of population of working age with limiting ong term illness aged 16-74 (KS008003/KS09A001) |
| POPPUCAR | Unpaid care providers in population | Proportion of population providing unpaid care (KS008007/KS008001) |
| POPPUCA1 | Unpaid care (<20 hrs week) in population |  |
| POPPUCA2 | Unpaid care (20-49 hrs) in pop | Proportion of population providing unpaid care for 20-49 hours per week (KS008009/KS008001) |
| POPPUCA3 | Unpaid care ( $>50 \mathrm{hrs}$ week) in |  |
| POPPUCA3 | Proportion aged 16-74 with no |  |
| NQUAL174 | qualifications | Proportion of population aged 16-74 with no qualifications (KS013002/KS013001 Proportion of population aged 16-74 that are full-time students |
| FTSTUDEN | Proportion aged 16-74 full-time students | ((KS013008+KS013009)/KS013001) |
| HHNOCAR | Households without a car | Proportion of households without a car (KS017002/KS017001) Proportion of households that are owner occupied |
| OWNOCC | Owner occupied households | (KS018002+KS018003+KS018004)/KS018001) |
| LAHARENT | Rented social housing | Proportion of households that are rented from LA or HA ((KS018005+KS018006)/KS018001) |
| PRIVRENT | Rented private housing | Proportion of households that are rented from private landlords (KS018007/KS018001) |
| LONEPENH | Lone pensioner households | Proportion of households that are one pensioner households (KS020002/KS020001) Proportion of households that are lone parent households with dependent children |
| LONEPARH | Lone parent households | (KS020011/KS020001) |
| PERMSICK | Permanently sick of those aged 16-74 Long-term unemployed of those aged | Proportion of population aged 16-74 that are permanently sick (KS09A010/KS09A001) |
| PC74LTUN | $16-74$ | Proportion of those aged 16-74 that are long-term unemployed (KS09A015/KS09A001) Proportion of those aged 16-74 in employment that are working agriculture |
| WORKAGRI | Employed in agriculture | (KS11A002/KS11A001) |
| PROFOCCU | People in professional occupations | Proportion of those aged 16-74 in managerial and professional occupations ((KS14A002+KS14A003+KS14A004)/KS14A001) |

Table BA.3: table showing first stage regressions for outcome models associated with 2005/6 expenditure and mortality data for 2002/3/4


Table BA.4: table showing first stage regressions for expenditure models associated with 2005/6 expenditure and mortality data for 2002/3/4


Table BA.5: table showing first stage regressions for outcome and expenditure models associated with 2006/7 expenditure and mortality data for 2004/5/6


Note: these are the first-stage regressions for the IV models reported in Table B8.16. Robust standard errors in brackets, ${ }^{* * *} \mathrm{p}<0.01, * * \mathrm{p}<0.05, * \mathrm{p}<0.1$

Table BA.6: table showing first stage regressions for outcome and expenditure models associated with 2006/7 expenditure and mortality data for 2006/7/8


Table BA.7: table showing first stage regressions for outcome models associated with 2007/8 expenditure

|  | (1) | (2) | (3) | (4) | (5) | (6) | (7) | (8) | (9) | (10) | (11) | (12) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | PBC 2 cancer | PBC 2 cancer | PBC 10 circulation | PBC 11 respiratory | PBC 11 respiratory | PBC 13 gastrointestinal | PBC 13 gastrointestinal | PBC 4 endocrine | PBC 17 genitourinary | PBC 1 infectious disease | PBC 1819 Maternity \&neonates | PBC 16 Trauma \&injuries |
| VARIABLES | 2007/8 instrument spend weighted first stage | 2007/8 <br> instrument <br> spend <br> weighted <br> first stage | 2007/8 <br> instrument <br> spend <br> weighted <br> first stage | 2007/8 <br> instrument <br> spend <br> weighted <br> first stage | 2007/8 <br> instrument <br> spend <br> weighted <br> first stage | 2007/8 <br> instrument <br> spend <br> weighted <br> first stage | 2007/8 <br> instrument <br> spend <br> weighted <br> first stage | 2007/8 <br> instrument <br> spend <br> weighted <br> first stage | 2007/8 instrument spend weighted first stage | $\begin{aligned} & \text { 2007/8 } \\ & \text { instrument } \\ & \text { spend } \\ & \text { weighted } \\ & \text { first stage } \end{aligned}$ | 2007/8 <br> instrument <br> spend <br> weighted <br> first stage | 2007/8 instrument spend weighted first stage |
| need CARAN per head | 0.582** | 0.545*** | 0.724*** | 1.251*** | 1.196*** | 0.999*** | 1.047*** |  | $\bigcirc$ |  | 1.111*** | 0.856*** |
| no car households | [0.284] | [0.105] | [0.168] | [0.094] | [0.113] | [0.105] | [0.099] |  |  | $\begin{aligned} & 0.512^{*} \\ & {[0.267]} \end{aligned}$ | [0.259] | $\begin{aligned} & {[0.282]} \\ & 0.288^{* *} \\ & {[0.137]} \end{aligned}$ |
| lone pensioner households | $\begin{aligned} & 0.632^{* * *} \\ & {[0.148]} \end{aligned}$ | $\begin{aligned} & 0.644^{* * *} \\ & {[0.119]} \end{aligned}$ | $\begin{aligned} & 0.468 * * * \\ & {[0.100]} \end{aligned}$ | $\begin{aligned} & 0.360 * * * \\ & {[0.103]} \end{aligned}$ | $\begin{aligned} & 0.269 * * * \\ & {[0.100]} \end{aligned}$ | $\begin{aligned} & 0.199 \\ & {[0.133]} \end{aligned}$ |  |  |  |  |  |  |
| IMD 2007 | $\begin{aligned} & -0.012 \\ & {[0.088]} \end{aligned}$ |  |  |  |  |  |  | $\begin{gathered} -0.067 \\ {[0.053]} \end{gathered}$ |  | $\begin{aligned} & 0.325 \\ & {[0.224]} \end{aligned}$ |  |  |
| need CARAN per head squ |  |  |  | $\begin{aligned} & 1.332 * * * \\ & {[0.428]} \end{aligned}$ | $\begin{aligned} & 1.338 * * * \\ & {[0.425]} \end{aligned}$ |  |  | $5$ |  |  |  |  |
| provision of unpaid care |  |  | $\begin{aligned} & 0.441^{* *} \\ & {[0.174]} \end{aligned}$ |  | $\begin{aligned} & 0.200 \\ & {[0.160]} \end{aligned}$ |  |  |  |  |  |  |  |
| born outside EU |  |  |  |  |  | $\begin{aligned} & -0.054^{* * *} \\ & {[0.018]} \end{aligned}$ | $\left[\begin{array}{l} -0.067 * * * \\ {[0.017]} \end{array}\right.$ |  |  |  | $\begin{aligned} & 0.004 \\ & {[0.041]} \end{aligned}$ | $\begin{aligned} & -0.079 * * \\ & {[0.039]} \end{aligned}$ |
| diabetes prevalence rate 2007/8 |  |  |  |  |  |  |  | $\begin{aligned} & 0.358 * * * \\ & {[0.123]} \end{aligned}$ |  |  |  |  |
| permanently sick |  |  |  |  |  | $5$ |  | $\begin{aligned} & 0.307 * * * \\ & {[0.061]} \end{aligned}$ |  |  |  |  |
| lone parent households |  |  |  |  |  | $2$ |  |  | $\begin{aligned} & 0.029 \\ & {[0.124]} \end{aligned}$ |  |  |  |
| CKD prevalence rate 2007/8 |  |  |  |  | 人 |  |  |  | $\begin{aligned} & 0.123^{* *} \\ & {[0.061]} \end{aligned}$ |  |  |  |
| long-term unemployed |  |  |  |  | N |  |  |  | $\begin{aligned} & 0.146 * * \\ & {[0.061]} \end{aligned}$ |  |  |  |
| limiting long-term illness |  |  |  |  |  |  |  |  | $\begin{aligned} & 0.207 \\ & {[0.134]} \end{aligned}$ |  |  |  |
| HIV need per head squared |  |  |  |  |  |  |  |  |  | $\begin{aligned} & 0.128 * * * \\ & {[0.031]} \end{aligned}$ |  |  |
| HIV need per head |  |  |  |  |  |  |  |  |  | $0.300 * * *$ |  |  |
| work in agriculture |  |  |  | $\underset{\sim}{1}$ |  |  |  |  |  | [0.044] <br> 0.152** <br> [0.064] |  | $\begin{aligned} & 0.126^{* * *} \\ & {[0.033]} \end{aligned}$ |
| work in professional occupation |  |  |  |  |  |  |  |  |  | $\begin{aligned} & 0.647 * * * \\ & {[0.172]} \end{aligned}$ |  |  |
| no qualifications |  |  | ®n |  |  |  |  |  |  |  | $\begin{gathered} -0.214 \\ {[0.160]} \end{gathered}$ |  |
| maternity need per head |  |  |  |  |  |  |  |  |  |  | $\begin{aligned} & 0.647 * * * \\ & {[0.159]} \end{aligned}$ |  |
| full-time students |  |  |  |  |  |  |  |  |  |  |  | $\begin{aligned} & 0.126 \\ & {[0.095]} \end{aligned}$ |
| LA/HA accommodation |  | C |  |  |  |  |  |  |  |  |  | $\begin{aligned} & -0.197^{*} \\ & {[0.104]} \end{aligned}$ |
| Constant | 5.759*** | 5.747*** | 6.754*** | 4.886*** | 5.172*** | 4.531*** | 4.104*** | 4.224*** | 5.269*** | 4.226*** | 4.004*** | 4.766*** |
|  | [0.252] | [0.234] | [0.322] | [0.201] | [0.350] | [0.281] | [0.057] | [0.390] | [0.209] | [1.007] | [0.297] | [0.339] |
| Observations | 151 |  | 151 | 151 | 151 | 151 | 151 | 151 | 147 | 151 | 151 | 151 |
| R-squared | 0.369 | 0.369 | 0.653 | 0.659 | 0.664 | 0.531 | 0.524 | 0.436 | 0.296 | 0.724 | 0.407 | 0.361 |

[^71]

[^72]Table BA.9: table showing first stage regressions for outcome models associated with 2008/9 expenditure

|  | (1) | (2) | (3) | (4) | (5) | (6) | (7) | (8) | 9) $\stackrel{ }{ }$ | (10) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | PBC 2 cancer | PBC 10 circulation | PBC 11 respiratory | PBC 13 <br> gastro-intestinal | PBC 13 <br> gastro-intestinal | PBC 4 endocrine | PBC 7 neurological | PBC 17 <br> genito-urinary | PBC 1 <br> infectious disease | PBC 1819 <br> maternity/neonates |
|  | 2008/9 | 2008/9 | 2008/9 | 2008/9 | 2008/9 | 2008/9 | 2008/9 | 2008/9 | 2008/9 | 2008/9 |
|  | outcome model | outcome model | outcome model | outcome model | outcome model | outcome model | outcome model | outcome model | outcome model | outcome model |
|  | instrument spend weighted | instrument spend weighted | instrument spend weighted | instrument spend weighted | instrument spend weighted | instrument spend weighted | instrument spend weighted | instrument spend weighted | instrument spend weighted | instrument spend weighted |
| Regressors | first stage | first stage | first stage | first stage | first stage | first stage | first stage |  | first stage | first stage |
| need CARAN per head | 1.122*** | 1.274*** | 1.228*** | 0.989*** | 1.056*** |  | 0.373** |  |  | 0.659*** |
|  | [0.198] | [0.146] | [0.085] | [0.088] | [0.087] |  | [0.175] |  |  | [0.236] |
| lone pensioner households | 0.490*** | 0.426*** | 0.252** | 0.272** |  |  | 0.287*** |  |  |  |
|  | [0.127] | [0.090] | [0.110] | [0.109] |  |  | [0.109] |  |  |  |
| IMD 2007 | -0.145** |  |  |  |  | -0.082 | (2) |  | 0.236 |  |
|  | [0.060] |  |  |  |  | [0.091] | (0) |  | [0.214] |  |
| no car households |  | -0.188*** |  |  |  |  |  |  | 0.502** |  |
|  |  | [0.053] |  |  |  |  |  |  | [0.227] |  |
| need CARAN per head sq |  |  | 1.071** |  |  | $\wedge$ |  |  |  |  |
|  |  |  | [0.426] |  |  |  |  |  |  |  |
| provision of unpaid care |  |  | 0.339*** |  |  | $0.530 \% *$ |  |  | ${ }^{1.393 * * *}$ |  |
|  |  |  | [0.117] |  |  | $0.232]$ |  |  | [0.340] |  |
| born outside EU |  |  |  | $\begin{aligned} & -0.042^{* * *} \\ & {[0.016]} \end{aligned}$ | $\begin{aligned} & -0.060^{* * *} \\ & {[0.015]} \end{aligned}$ | $\begin{aligned} & 0.080 * * * \\ & {[0.026]} \end{aligned}$ |  |  |  | $\begin{aligned} & -0.031 \\ & {[0.032]} \end{aligned}$ |
| diabetes prevalence rate 2007/8 |  |  |  |  |  | 0.167 |  |  |  |  |
|  |  |  |  |  | ค) | [0.132] |  |  |  |  |
| permanently sick |  |  |  |  | ) | $0.380 * * *$ |  |  |  |  |
| epilepsy prevalence rate 2007/8 |  |  |  |  | $)$ | [0.104] | 0.486*** |  |  |  |
|  |  |  |  |  |  |  | [0.121] |  |  |  |
| owner occupied households |  |  |  | 0 |  |  | -0.235** |  |  |  |
|  |  |  |  | - |  |  | [0.113] |  |  |  |
| lone parent households |  |  |  |  |  |  |  | 0.175** |  | 0.013 |
|  |  |  |  | $\cdots$ |  |  |  | [0.080] |  | [0.106] |
| CKD prevalence rate 2007/8 |  |  |  | $71$ |  |  |  | 0.089*** |  |  |
|  |  |  |  |  |  |  |  | [0.033] |  |  |
| long-term unemployed |  |  |  | $5$ |  |  |  | $\begin{aligned} & 0.148 * * * \\ & {[0.045]} \end{aligned}$ |  |  |
| HIV need per head |  |  |  |  |  |  |  |  | 0.471*** |  |
|  |  |  |  |  |  |  |  |  | [0.050] |  |
| HIV need per head squared |  |  | $\bigcirc$ |  |  |  |  |  | 0.146*** |  |
|  |  |  | , |  |  |  |  |  | [0.027] |  |
| no qualifications |  |  | F |  |  |  |  |  | -0.751*** | -0.092 |
|  |  |  |  |  |  |  |  |  | [0.189] | [0.113] |
| work in agriculture |  |  |  |  |  |  |  |  | 0.150*** |  |
|  |  |  |  |  |  |  |  |  | [0.051] |  |
| maternity need per head |  |  |  |  |  |  |  |  |  | $[0.162]$ |
| Constant | 5.937*** | 5.435*** | 5.610*** | 4.752*** | 4.167*** | 6.379*** | 4.808*** | 5.363*** | 6.010*** | 4.171*** |
|  | [0.221] | [0.2301) | [0.238] | [0.236] | [0.048] | [0.768] | [0.193] | [0.121] | [1.513] | [0.375] |
| Observations | 151 | 151 | 151 | 151 | 151 | 151 | 151 | 148 | 151 | 151 |
| R-squared | 0.521 | 0,612 | 0.746 | 0.665 | 0.648 | 0.559 | 0.477 | 0.378 | 0.791 | 0.614 |

[^73]Table BA.10: table showing first stage regressions for expenditure models associated with 2008/9 expenditure


## Appendix C

## Translating mortality effects into life years and quality adjusted life years

C. 1 Introduction
C. 2 Analysis of 2006/07 expenditure and 2006 to 2008 mortality data
C. 3 Re-estimating the cost per QALY threshold using 2008/09 expenditure data
C. 4 Re-estimating the cost per QALY threshold using 2007/08 expenditure data

Addendum C1: Data sources
Addendum C2: The role of data on local NHS decisions
Addendum C3: Characterisation of the investment and disinvestment decisions in mental health: depression and schizophrenia

Addendum C4: What type of health is forgone by the approval of a new technology?

## References

## C. 1 Introduction

This Appendix describes how the results of the econometric work undertaken to estimate the link between NHS spending and mortality, which was detailed in Appendix B, can be translated in to effects on life years and quality adjusted life years (QALYs). This Appendix presents much of the detail of data and analyses that support Chapter 4 of the main report.

We present three sequential steps of analysis which lead to estimates of the overall cost per QALY threshold for the NHS:
i. In section C.2.1 we reconsider how the estimated effects on mortality from the econometrics work might better translate into life years by exploring the limitations of mortality data ayailable at PCT level and the published years of life lost (YLL) figures presented in the previous chapter. We explore how these estimates might be improved using additional data and analysis.
ii. In section C.2.2 we consider how these estimates of life year effects might be adjusted for the quality of life in which they are lived, taking account of the gender and the age at which life years are gained or lost as well as the disutility associated with particular diseases.
iii. In section C.2.3 we explore ways to also take account of those effects on health not directly associated with mortality and life year effects (i.e., the 'pure' quality of life effects) to estimate an overall cost per QALY threshold.

This sequence of analysis is set out and explained based on the analysis of 2006/07 expenditure and mortality data from 2006 to 2008. At the end of each section, we present a summary which includes a central 'best' estimate as well as extreme lower and apper bounds for the cost per life year and cost per QALY threshold. The core assumptions which underpin these three values are common across sections C.2.1 to C.2.3. The central or 'best' estimate is based on two assumptions one conservative and the other more optimistic with respect to the healtheffects associated with expenditure. The first is that the health effects of changes in one year of expenditure are restricted to one year. Recall that the analyses in Appendix B use 3 years of mortality data, but these are averaged to an annual value prior to estimating outcome elasticities. Therefore, the estimated outcome elasticities represent the proportionate effect on mortality in one year due to a proportionate change in expenditure. This is likely to underestimate effects on mortality since expenditure that reduces mortality risk for an individual in one year may well also reduce their risk over subsequent years; possibly over the whole of their remaining disease duration. Expenditure may also preventdisease in future patient populations. Therefore, total health effects will be underestimated and the cost per life year or QALY threshold will be overestimated. Although undoubtedly conservative, it may be offset to some extent by the more optimistic assumption used to translate mortality effects into life years. In common with YLL figures published by NHS IC and the WHO Globat Burden of Disease study it is assumed that any death averted by expenditure in one year will return the individual to the mortality risk of the general population, i.e., the years of life gained associated with each death averted are based on what would have been their life expectancy taking account of their of age and gender (using life tables for the general population).

The extreme upper and lower bounds for cost per life year and cost per QALY thresholds are based on making both assumptions either optimistic (providing the lower bound for the threshold) or both conservative (an upper bound for the threshold). The lower bound is based on assuming that health effects are not restricted to one year but apply to the remaining disease duration for the population at risk during the expenditure year (although this still does not account for the effects of expenditure on preventing disease). The upper bound is based on the combination of assuming that health effects are restricted to one year and that any death averted is only averted for the minimum duration consistent with the mortality data used to estimate the outcome elasticities in Appendix B. It is very important to note that the lower and upper bounds represent extreme values rather than alternative but plausible views that could reasonably be taken.

The three sequential steps of analysis, which provide a cost per life year threshold, through a cost per life year adjusted for quality to a cost per QALY threshold, are explained and detailed in sections C.2.1 to C.2.3, using the analysis of 2006 expenditure and mortality data from 2006 to 2008. In Section C.2.4, further analysis using these data highlight which PBCs have the greatest influence on the overall threshold. An exploration of the impact of the uncertainty over the outcome and spend elasticities in estimates of the threshold is also presented in Section C.2.5. The sequence of analyses is then applied to 2008/09 expenditure and 2008 to 2010 mortality data; results of the cost per QALY threshold for the most recent years of analysis are presented in Section C.3. In Section C. 4 we present our best estimate of the threshold cost per QALY based on the analysis of 2007/8 expenditure and mortality data from/2007 to 2009 .

## C. 2 Analysis of 2006/07 expenditure and 2006 to 2008 mortality data

## C.2.1 From mortality to life years

In this section we summarise our examination of a number of issues associated with available PCT-based mortality data and the associated published estimates of YLL. We then examine how, given the limited information available about the population at risk in each PBC we might take proper account of the fact that some of the observed deaths would have occurred anyway (had the same population not been at risk in the particular PBC ) when estimating YLL, i.e., taking account of unobserved counterfactual deaths. This allows us to estimate the YLL that better reflects the effect of expenditure on the mortality observed in each PBC, and infer the excess deaths associated with each PBC. Finally we present cost perdeath averted and cost per life year which accounts for the issues raised in this section.

## C.2.1.1 Mortality and YLL coverage

The mortality data that is available at PCT level does not offer full coverage of all deaths across all the ICDs that make up each PBC. Table C. 1 illustrates, using a few PBCs as examples, the mapping of three digit ICD-10 to PBCs (column 1) and the incomplete coverage of these ICDs in mortality data (column 2). A more detailed account of the extent of coverage is presented in Table B.5.1 in Appendix B.

Table C.1. Illustrating coverage

| PBC |  | ICD codes covered by the spend data <br> [1] | ICD codes coyered by the mortality data (NHS IC) [2] | Coverage of mortality data relative to spend data (2008) <br> [3] |
| :---: | :---: | :---: | :---: | :---: |
| 1 | Infectious diseases | large parts of A00-B99 A00-B99 <br> C00-C97, D00-D49 C00-C97 <br> E000-E899 E10-E14 <br> I00-I99, Q20-Q28 $\mathrm{I} 00-\mathrm{I} 99$ <br> A150-A169, * A190-A199, $\mathrm{J} 12-\mathrm{J} 18$, J40-J44, J45- <br> J000-J989, Q300-Q349, J 46 |  | 1.000 |
| 2 | Cancer |  |  | 0.984 |
| 4 | Endocrine |  |  | 0.634 |
| 10 | Circulatory |  |  | 0.992 |
| 11 | Respiratory |  |  | 0.773 |

National (English) data are, however, available that cover all deaths associated with all the ICDs that make up each PBC. Therefore, it is possible to adjust the incomplete reporting of mortality at PCT level (see section B5.1 in Appendix B) before applying the estimated outcome elasticities to calculate the deaths averted due to expenditure. Applying published estimates of YLL per death to all the deaths averted using coverage adjustment factors (as illustrated in column 3 of Table C.1) provides the estimate of the cost per life year reported in Appendix B. Note that the proportionate effects on mortality (due to changes in expenditure) are therefore assumed to be similar for mortality that is and is not recorded at PCT level. This seems more reasonable than assuming no effect of expenditure on mortality that happens not to be recorded at PCT level.

The published estimates of YLL (NHS IC) used in Chapter 3 only include deaths below 75 years (but exclude deaths below 1 year) and are based on the difference between age 75 and the age of each death below 75. These estimates have the same limited coverage as PCT level mortality data, so are not available for all the ICDs that make up each PBC. Therefore, applying the available estimates of YLL per death to the estimated number of deaths averted requires an assumption that the YLL per death is similar for those groups of ICDs covered and not covered by the published YLL figures.

This can be examined by using national ONS data to calculate YLL in the same way as NHS IC, but with full coverage of all the ICDs that make up each PBC. Although ONS data provides complete coverage and reports gender, age at death is only reported in 5 year ranges (these data are not available at PCT level so could not be used when estimating outcome elasticities in Chapter 3). Therefore, using ONS data to
estimate YLL requires taking the midpoint ${ }^{1}$ of each range as the age of death, i.e., assuming reported deaths are equally likely over the range in which they are reported. For this reason it is not possible to precisely recover the published YLL figures using ONS data for those ICD groupings that can be precisely matched to the NHS IC coverage. However, the differences are small (ranging from $-1 \%$ to $2 \%$ as shown in Table C. 2 below), suggesting that taking the midpoint of each range as the age of death is a reasonable approximation.

Table C.2. Estimates of YLL for NHS IC and ONS for those ICD groupings that can be precisely matched to the NHS IC coverage


* does not take into account coverage adjustment
+ deaths age $<1$ included in PBC $18+19$
Published estimates of YLL are available from NHS IC for PBC16 (Trauma and injuries) but ONS does not provide the information required to calculate YLL for this PBC. However, the estimated outcome elasticity could not be estimated for 2006/07 expenditure and 2006 to 2008 mortality and was assumed to be zero. Therefore, this PBC does not contribute any changes in health outcomes due to changes in expenditure in subsequent estimates of cost per life year and QALY thresholds anyway.

The differences between estimates of YLL based on ONS and NHS IC data are, however, much more significant and are reported in Table C.3. These reflect differences in the distribution of ages at death between those groups of ICDs covered and not covered in the NHS IC figures. For example, NHS IC figures available at PCT level for PBC7 (neurological problems) have low coverage of all deaths in this PBC ( 0.136 in column 1). The deaths that are reported in NHS IC are associated with epilepsy and the YLL (22,046 in column 2) reflects the generally younger age at death in this group. When adjusted for full coverage $(22,046 / 0.136=162,100$ in column 3) the estimated YLL is much greater than the YLL based directly on all deaths by age group reported for the PBC in ONS. This difference in YLL reflects the fact that the deaths in PBCZ which are not covered by NHS IC figures tend to be in older age groups so generate fewer YLL.

Table C.3. Estimates of YLL for NHS IC and ONS

| PBC | Coverage of mortality data relative to spend data [1] | $\begin{gathered} \mathrm{YLL}_{<75} \\ (\mathrm{NHS} \mathrm{IC}) \\ {[2]} \end{gathered}$ | YLL<75 <br> adjusted <br> (NHS IC) <br> [3] | YLL<75 no adjustment needed (ONS) [4] | Difference from adjusted NHS IC to ONS [5] |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 1 Infectious diseases | 1.000 | 35,517 | 35,517 | 40,928 | 15\% |
| 2 Cancer | 0.984 | 735,674 | 747,636 | 758,804 | 1\% |
| 4 Endocrine problems | 0.634 | 19,224 | 30,322 | 41,548 | 37\% |
| 7 Neurological problems | 0.136 | 22,046 | 162,100 | 93,755 | -42\% |
| 10 Circulatory | 0.992 | 453,878 | 457,538 | 481,246 | 5\% |
| 11 Respiratory | 0.773 | 108,074 | 139,812 | 147,465 | 6\% |
| 13 Gastro-intestinal | 0.571 | 115,303 | 201,931 | 177,532 | -12\% |
| 17 Genito-urinary | 0.172 | 3,343 | 19,438 | 17,380 | -11\% |
| 18+19 Maternity \& neonates* | 0.679 | 164,200 | 241,826 | 15,409 | -94\% |

${ }^{1}$ The calculated midpoints are as follows,

| Age range | $<1$ | $1-4$ | $5-9$ | $10-14$ | $15-19$ | $20-24$ | $25-29$ | $30-34$ | $35-39$ | $40-44$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| midpoint | 0.5 | 3.0 | 7.5 | 12.5 | 17.5 | 22.5 | 27.5 | 32.5 | 37.5 | 42.5 |
| Age range | $45-49$ | $50-54$ | $55-59$ | $60-64$ | $65-69$ | $70-74$ | $75-79$ | $80-84$ | $85-89$ | $90+$ |
| midpoint | 47.5 | 52.5 | 57.5 | 62.5 | 67.5 | 72.5 | 77.5 | 82.5 | 87.5 | 92.5 |

Using ONS data also allows deaths under the age of 1 year to be appropriately assigned to PBCs via the ICD in which they occurred (NHS IC YLL figures exclude deaths under one year), rather than assigning them all to PBC18 \& 19 as in Appendix B. ${ }^{2}$ This explains the large reduction in YLL for PBC18 \& 19 (Maternity and neonates) as much of the mortality is re-assigned to ICDs which contribute to other PBCs. Since most of the deaths that are re-assigned are allocated to PBC1 (infectious diseases) the YLL for this PBC increases despite complete reporting of deaths at PCT level and full coverage by NHS IC figures (see Table C.4).

Table C.4. Estimates of YLL for NHS IC and ONS including deaths age $<1$


Using ONS data to calculate YLL in the same way as the published NHS IC figures, but overcoming some of the issues associated with the reporting of mortality at PCT levet and the coverage of published estimates of YLL, generates similar estimates of a cost per life year threshold (see column 2 Table C.6) to those reported in Appendix B.

## C.2.1.2 Life expectancy and YLL



As noted above, the NHS IC estimates of YLJ only include deaths below 75 years and are based on the difference between age 75 and the age of each death below 75 . Implicitly, this treats 75 as the appropriate normal life expectancy for males and females for the population at risk in each PBC. However, with the exception of maternity and neonates, most deaths in PBCs occur above the age of 75 and life expectancies are significantly greater than 75 . Based on 2006 to 2008 data, life expectancy at birth is greater than 75 ( 77.74 for males and 81.88 for females). ${ }^{3}$ Given the need to reflect the normal life expectancy for the at risk population, it is more appropriate to use the age distribution of the general population, and calculate life expectancy conditional on age averaged over the general population's age distribution. General population life expectancies are estimated to be 80.7 for males and 84.4 for females, These life expectancy estimates will always be higher than life expectancies at birth.

Based on ONS data, YLLs can be re-recalculated using the above estimates of gender specific life expectancy for the general population. When increasing life expectancy (LE) two effects occur, both of which tend to increase estimates of YLL. Firstly, more deaths are included in the YLL calculation (those that occurbetween age 75 and LE) and secondly, each death previously counted below 75 will generate 5.7 or 9.4 more YLL for males and females respectively. The effect on the number of deaths and the YLL for each PBC of using the life expectancy of the general population is reported in Table C.5.

[^74]Table C.5. The difference in YLL by life expectancy

| PBC |  | Deaths<75 (ONS) [1] | Deaths<LE (ONS) <br> [2] | Difference in deaths due to increased LE [3] | YLL<75 (ONS) <br> [4] | $\begin{gathered} \mathrm{YLL}_{<\mathrm{LE}} \\ (\mathrm{ONS}) \\ {[5]} \\ \hline \end{gathered}$ | Difference in YLL due to increased LE [6] |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | Infectious diseases | 2,050 | 3,710 | 81\% | 40,928 | 62,051 | 52\% |
| 2 | Cancer | 62,944 | 95,212 | 51\% | 758,804 | 1,345,013 | 77\% |
| 4 | Endocrine | 2,367 | 4,000 | 69\% | 41,548 | 65,015 | 56\% |
| 7 | Neurological | 5,095 | 8,975 | 76\% | 93,755 | 145,526 | 55\% |
| 10 | Circulatory | 41,487 | 82,098 | 98\% | 481,246 | 916,170 | 90\% |
| 11 | Respiratory | 14,000 | 30,500 | 118\% | 147,465 | 310,326 | $110 \%$ |
| 13 | Gastro-intestinal | 10,611 | 15,827 | 49\% | 177,532 | 273,303 | $54 \%$ |
| 17 | Genito-urinary | 1,588 | 4,197 | 164\% | 17,380 | 39,098 | 125\% |
| 18+19 | Maternity \& neonates | 226 | 226 | 0\% | 15,409 | 17,167 | 11\% |

Life expectancy (LE): male=80.7, female $=84.4$
The number of deaths counted below LE increases for every PBC except for maternity \& neonates because, as expected, all deaths are below age 75 in PBC18 \& 19. However, YLL increases for all PBCs reflecting the additional years otherwise expected to be lived to an older LE. Of course including more of the deaths observed in each PBC and the greater YLL associated with them will generate more deaths averted and more life years gained when applying the same proportionate effects from the outcome elasticities estimated in Appendix B. Therefore, the cost per death averted and cost per life year thresholds are expected to be lower using these figures than those reported in Appendix B.

The impact on the cost per life year and cost per death averted thresholds is summarised in Table C.6. A detailed breakdown of the changes in spend and YLL across PBCs is presented in Table C.7. A listing of the spend and outcome elasticities used in threshold calculations throughout this section is in Table C.8.

Table C.6: Summary of cost per death averted and cost per life year threshold.

|  | Using 75 as the cut-off (ONS) |  | Using LE as the cut-off (ONS) |  |
| :---: | :---: | :---: | :---: | :---: |
|  | Cost per death averted 11 | Cost per LY gained [2] | Cost per death averted <br> [3] | Cost per LY gained <br> [4] |
| big 4 PBC's | ¢,122,756 | £10,398 | £63,426 | £,5,487 |
| 11 PBCs (with mortality) | $) £ 240,433$ | $\AA 20,031$ | £124,655 | £10,660 |
| All 23 PBCs (zero health effects for remaining 12 PBCs) | $£ 884,579$ | £73,697 | £458,620 | £39,218 |
| All 23 PBCs (non-zero health effects for remaining 12 PBCs, except GMS)* | $£ 271,739$ | $£ 22,639$ | £140,886 | £12,048 |

* in PBCs without a mortality signal, health effects were estimated by valuing changes in expenditure at the same rate as observed in PBCs for which there was a mortality signal.

Table C.7: Breakdown of the cost per death averted and cost per life year thresholds

|  |  |  | Using 75 as the cut-off (ONS) |  |  |  |  |  | U |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| PBC | PBC description | Change in spend, Em | N death $(<75)$ | Change in N deaths | Cost per death averted, $£$ | YLL | Change in YLL | Cost per LY gained, $£$ | N deaths $(<\mathrm{LE})$ | Change in N deaths |  |
| 2 | Cancer | £19 | 62944 | 100.10 | £191,500 | 758804 | 1207 | £15,885 | 95212 | 151.42 |  |
| 10 | Circulatory | £33 | 41487 | 321.26 | £103,560 | 481246 | 3727 | £,8,928 | 82098 | 635.73 |  |
| 11 | Respiratory | $£ 22$ | 14000 | 249.25 | £89,482 | 147465 | 2625 | £8,495 | 30500 | 543. |  |
| 13 | Gastro-intestinal $\operatorname{Big} 4$ | £17 | 10611 | 72.69 | $\begin{aligned} & £ 227,013 \\ & £, 122,756 \end{aligned}$ | 177532 | 1216 | $\begin{array}{r} £ 13,568 \\ \quad \AA 10,398 \\ \hline \end{array}$ | $15827$ | $108.42$ |  |
| 1 | Infectious diseases | $\AA 8$ | 2050 | 0.76 | £10,936,680 | 40928 | 15 | £547,796 | 3710 | 1.38 |  |
| 4 | Endocrine | £18 | 2367 | 18.99 | £,929,559 | 41548 | 333 | £52,957 | 4000 | 32.1 |  |
| 7 | Neurological | £17 | 5095 | 3.52 | £,4,889,114 | 93755 | 65 | £265,693 | 8975 | 6.19 |  |
| 17 | Genito-urinary | £32 | 1588 | 0.74 | £42,993,075 | 17380 | 8 | £3,928,251 | 4197 | 1.95 | , |
| 16 | Trauma \& injuries* | £10 | NA | 0.00 | NA | NA | 0 | NA | NA | 0 |  |
| 18+19 | Maternity \& neonates* First 11 PBC's | $\AA 8$ | 226 | 0.24 | $\begin{array}{r} £ 32,813,038 \\ £ 240,433 \\ \hline \end{array}$ | 15409 | 17 | $\begin{array}{r} f 481,261 \\ \& 20,031 \end{array}$ | 226 | 0.24 | , |
| 3 | Disorders of Blood | $£ 11$ |  | 46.57 | £240,433 |  | 559 | £20,031 |  | 89.83 |  |
| 5 | Mental Health | $£ 204$ |  | 849.17 | £240,433 |  | 10193 | £20,031 |  | 1637.87 |  |
| 6 | Learning Disability | £31 |  | 128.05 | £240,433 |  | $1537)$ | £20,031 |  | 246.98 |  |
| 8 | Vision | $£_{24}$ |  | 100.54 | £240,433 |  | 1207 | $\AA^{\text {¢ } 20,031}$ |  | 193.92 |  |
| 9 | Hearing | $¢_{6} 6$ |  | 26.60 | £240,433 |  | 319 | £20,031 |  | 51.3 |  |
| 12 | Dental | $£ 23$ |  | 97.72 | £240,433 |  | 1173 | $\AA 20,031$ |  | 188.48 |  |
| 14 | Skin | £11 |  | 43.72 | £240,433 |  | 525 | £20,031 |  | 84.34 |  |
| 15 | Musculo skeletal | $£ 15$ |  | 62.93 | £240,433 |  | 755 | £20,031 |  | 121.38 |  |
| 20 | Poisoning and AE | $\AA 4$ |  | 18.27 | £240,433 |  | 219 | £20,031 |  | 35.23 |  |
| 21 | Healthy Individuals | £18 |  | 76.27 | £240,433 |  | 915 | $\AA 20,031$ |  | 147.1 |  |
| 22 | Social Care Needs | £68 |  | 281.19 | f240,433 |  | 3375 | $£ 20,031$ |  | 542.35 |  |
| 23 | Other | £78 |  | 0 | NA |  |  | NA |  |  |  |
|  | All (23 PBCs) |  |  |  | ¢,271,739 |  |  | £22,639 |  |  |  |

Note that we have been unable to obtain a satisfactory outcome model for trauma \& injuries and have assumed a zero outcome elasticity.
Note that, for expenditure in 2006/7, the neonate category has been merged with maternity to obtain plausible outcome and expenditure models.

Table C.8: Outcome and spend elasticities.

${ }^{(i)}$ The spend elasticities reflect how a $1 \%$ increase in budget is distributed across PBCs; however, in the econometrics, these were estimated separately for each PBC (unadjusted estimates in column 2) and because of this, its direct application to spend generates a change in budget bigger than the $1 \%$. An adjustment was thus applied to the remaining 12 PBC's (except PBC23 that was left unchanged), by multiplying each by a common factor - the magnitude of the unadjusted spend elasticities is changed but proportionality to the original elasticities is maintained (ii) without the negative sign

The cost per death averted (or life saved) threshold should not be over interpreted because this is of little direct policy interest since lives are never saved (death is only delayed) and the significance of a death averted depends critically on how long it is averted and the quality of life in which additional years are lived (see Section C.2.2). However, establishing the number of deaths averted which are associated with net YLL is useful because it enables an assessment of the number of life years gained associated with each death averted. Table C, 9 presents the YLL saved for each death averted implied by the assumptions underlying calculations of the cost per life year threshold in Table C.7. For the 11 PBCs with mortality signal, each death averted is assumed to be associated with a gain of 11.7 YLL (when LE is used, column 2). This value is smaller than when using 75 years old as a cut-off (column 1 ) because a higher proportion of deaths closer to the cut-off age are being considered (i.e., with lower YLL associated).

Table C.9: Implied YLL per death averted for each PBC


There are good reasons why YLL figures calculated in this way are biased. This is dealt with in the next section (Section C.2.1.3). In Section C.2.1.4 we take account of the fact that some of the deaths observed in a PBC would have occurred anyway in a similar 'normal' population (i.e., the counterfactual population not at risk through membership of the PBC) so not all observed deaths are 'excess' and generate YLL.

## C.2.1.3 YLL and accounting for counterfactual deaths

The estimates of YLL based on ONS data overcome many of the limitations of the published NHS IC figures. However, the YLLs reported in Table C. 5 are calculated in the same way as the NHS FE figures, by taking the difference between a fixed LE and the age at death for deaths observed below thatLE. Simply taking the difference between a fixed LE and the age at death of deaths that occur below LE and ignoring those deaths that occur above LE, is only an accurate representation of the XLL if it is reasonable to assume that no deaths would have otherwise occurred prior to LE (so all 'normal' deaths must occur at LE) and that there are no deaths (survivors) beyond LE in the at risk population, i.e. all deaths below LE are excess deaths and there are no excess deaths above LE. The estimate of YLL in the previous section may thus be biased for two reasons: i) it does not account for the fact that not all deaths observed below LE are 'excess' deaths in the sense that some deaths would have occurred (at the same age) in a similar population not at risk in the PBC and ii) some of the deaths observed above LE may be 'excess' deaths that would not otherwise have occurred at that age (see breakdown of deaths below and above LE in Table C.10).

The overall effect on YLL, and on the cost per life year, will depend on the number of deaths above and below LE that are excess. However, it is more likely that deaths below LE are 'excess'. Also, the YLL associated with deaths that are in fact not 'excess' below LE are expected to be greater than the YLL of excess deaths not counted above LE. Therefore, YLL estimates considering only deaths below LE are likely to overestimate YLL. Estimates of YHL are required which take account of the 'counterfactual' deaths that would have occurred even if the population in the PBC was not at risk through membership of the ICD codes that make it up, but faced the same mortality risks as the general population, accounting for the age and gender distribution of the PBC population.

Table C.10. Number of deaths below and above LE in 2006/07/08, by PBC

| PBC |  | $\begin{gathered} >\text { LE } \\ 2006 \\ {[2]} \\ \hline \end{gathered}$ | $\begin{gathered} <\mathrm{LE} \\ 2007 \\ {[3]} \\ \hline \end{gathered}$ | $\begin{gathered} >\mathrm{LE} \\ 2007 \\ {[4]} \\ \hline \end{gathered}$ | $\begin{gathered} <\mathrm{LE} \\ 2008 \\ {[5]} \\ \hline \end{gathered}$ | $\begin{gathered} >\text { LE } \\ 2008 \\ {[6]} \\ \hline \end{gathered}$ | Annual N deaths $<$ LE [7] | Annual N deaths > LE <br> [8] |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 Infectious diseases | 3,824 | 3,420 | 3,902 | 3,735 | 3,403 | 2,589 | 3,710 | 3,248 |
| 2 Cancer | 95,549 | 34,192 | 95,331 | 35,455 | 94,758 | 37,144 | 95,213 | 35,597 |
| 4 Endocrine | 4,006 | 2,661 | 3,967 | 2,750 | 4,028 | 2,882 | 4,000 | 2,764 |
| 7 Neurological | 8,454 | 5,762 | 8,845 | 6,501 | 9,626 | 6,871 | 8,975 | 6,378 |
| 10 Circulatory | 84,909 | 78,369 | 80,610 | 78,481 | 80,779 | 76,407 | 82,099 | 77,752 |
| 11 Respiratory | 29,925 | 34,549 | 29,540 | 35,060 | 32,036 | 35,227 | 30,500 | 34,945 |
| 13 Gastro-intestinal | 15,893 | 8,311 | 15,658 | 8,376 | 15,930 | 8,274 | 15,827 | 8,320 |
| 17 Genito-urinary | 4,056 | 6,049 | 4,072 | 6,558 | 4,465 | 6,673 | 4,198 | 6,427 |
| 18+19 Maternity \& neonates | 195 | 0 | 216 | 0 | 267 | 0 | 226 | 0 |

Dife expectancy (LE): male $=80.7$, female $=84.4$
Ideally, with reliable information about the size of the population at risk in each PBC and its age and gender distribution it would be possible to estimate the number of deaths that would be expected to occur had this population not been at risk, based on mortality data for the general population. The difference between deaths observed across all ages in the PBC and the deaths expected to have occurred in this matched 'normal' population would provide the number of 'excess' deaths by age and gender. These 'counterfactual' deaths will occur in the other PBCs insofar as all deaths are recorded in an ICD code, taking account of the unavoidable fact that everyone must die of something at some time. For
example, even if all observed cancer mortality was avoidable and could in principle be eliminated with sufficient expenditure, lives would not be 'saved' but deaths delayed and reallocated to other causes. ${ }^{4}$

The YLL associated with each of these excess deaths is the life expectancy conditional on gender and on surviving to the age at which the excess death occurred. The total YLL for the at risk population is simply the sum of these YLLs over all excess deaths, which could occur at any age. We do not (and will never) know the counterfactual expected age of death for each individual patient. However, two perfectly matched populations of individuals, one at risk and another not at risk in the PBC can be compared in terms of their survival curves (Figure C.1). The area below each survival curve reflects the life expectancy and the area between the two survival curves returns the YLL. This is equivalent to comparing the average age of death across patients in the population at risk in the PBC ( $N$ patients), with the average age of death in the matched, not at risk, population (for simplicity assumed to be equally sized). Equation (1) describes the YLL per patient as the difference in the average age of death, age $e_{\text {death }}$, observed for each individual, $i$ (out of $N$ individuals), in each population. The YLL for the population is simply the per patient YLL multiplied by the size of the population N.


Figure C.1: Survival curve of a population at risk in a PBC and of a matched 'normal' population
The difficultly is that routinely available data do not provide any information about the size of the population at risk or its age and gender distribution (matching criteria). Thus a matched population cannot be generated, and the area between the two curves cannot be evaluated. Therefore, it is not possible to directly estimate excess deaths or compare survival curves. Even if the size of the at risk population is unknown we can still use information that might be available about its age and gender distribution (or make reasonable assumptions) to estimate a matched 'normal' LE using life tables for the general population - such a LE summarises the area under the counterfactual survival curve ( $L E^{\text {norm }}=$ $\frac{1}{N} \sum_{i=1}^{\mathrm{N}} \mathrm{age}_{\text {death }, i}^{\text {norm }}$ in equation 1). Unfortunately, it is not possible to also calculate the LE for the population at risk in the PBC (or represent the survival curve) without information about the size of the at risk population - if it was possible, the difference between these life expectancies would approximate the YLL per patient at risk in a PBC.

Fortunately, we can still recover a consistent estimate of YLL using the normal LE of a matched population that is not at risk (a summary of the counterfactual average age of death), alongside the death

[^75]data available for the PBC population. Equation (2) shows that population YLL can be approximated by subtracting the age at which each observed death in a PBC has occurred to the normal LE.
\[

$$
\begin{equation*}
\text { YLL }=\mathrm{N} \cdot\left(\mathrm{LE}^{\text {norm }}-\frac{1}{N} \sum_{i=1}^{\mathrm{N}} \operatorname{age}_{\text {death }, \mathrm{i}}^{\mathrm{PBC}}\right)=\sum_{i=1}^{\mathrm{N}}\left(\mathrm{LE}^{\text {norm }}-\operatorname{age}_{\text {death }, \mathrm{i}}^{\mathrm{PBC}}\right) \tag{2}
\end{equation*}
$$

\]

The data on the PBC observed deaths available expresses the ages at which deaths occurred in age groups ( $k$, out of $K$ groups). Following from Equation (2), the population YLL can be evaluated considering the number of patients dying in each of the age groups, $\mathrm{N}_{\text {die, }, \text {, as depicted in Equation (3). This is equivalent }}$ to comparing survival curves where age is discretized into intervals and the midpoint of the intervals used as age of death - this is illustrated in Figure C.2.

$$
\mathrm{YLL}=\sum_{k=1}^{K}\left(\mathrm{LE}^{\text {norm }}-\mathrm{age}_{\text {death }, k}\right) \cdot \mathrm{N}_{\mathrm{die}, k}
$$



Figure C.2: Area between the survival curves, discretized.
The calculations (in Equations 2 and 3) require all observed deaths - both those that occur below and those that occur above this LE - to be taken into account. Those deaths occurring below LE generate YLL - compared to the average of a matched population not at risk. However, we must also account for those deaths that occur at ages above LE. These deaths generate life years 'gained' (YLG) compared to the average of a matched population not at risk. Therefore, the appropriate estimate is a net YLL (i.e., YLL - YLG). In effect, by subtracting YLG from YLL we take account of the fact that not all deaths below LE are excess deaths but some deaths above LE are. Insofar as deaths above LE have been observed in a specific PBC, the net YLL estimate will always be lower than the estimate of YLL. Consequently, the estimates in Section C.2.1.3 overestimate YLL and hence underestimate the cost per life year threshold.

Ksing the life expectancy of the general population
Routinely available data provides the age and gender of observed deaths but no information about the age and gender distribution of the at risk population itself. Using observed age and gender at death (see columns 5 and 6 of Table C.11) as an indication of the distribution of the at risk population will significantly overestimate the LE of a normal matched population insofar as a disease may be chronic (not all PBC mortality occurs on entry into the at risk population). If mortality risk increases over the disease duration more deaths would be observed in groups that have been prevalent for some time (i.e., are older) than those that are incident. Older age groups will thus be overrepresented in observed deaths compared to a matched normal population. For these reasons LE and YLL would be overestimated using age at death as a proxy for the age distribution of the at risk population, and the cost per life year would be underestimated.

Table C.11: Average age and life expectancy for PBCs based on age of the general population


In the absence of additional external information the net YLL could be based on the life expectancy of the general population, reflecting its current age and gender distribution. Such net YLL estimates are reported in Table C.12, and illustrate the impact of accounting for counterfactual deaths in the way described above. The YLL reported in column 5 of Table C. 12 are calculated the same way and are the same as the figures previously reported (column 5 of Table C.5). That is, they do not account for deaths that would have otherwise occurred below LE or the very many deaths that occur above LE. With the exception of PBC18 \& 19 many death occurabove the LE of the general population (see column 4 in Table C.12) in all PBCs. As a consequence, there are LYG associated with all other PBCs (see column 6) so the net YLL in column 7 are lower than YLL based on the same life expectancy. Therefore, failure to account for counterfactual deaths would lead to an overestimate of the YLL associated with a PBC and the effects of expenditure on YLL. The cost per life year threshold would be underestimated (see Table C.15).

Table C.12. Net YLL using life expectancy of the general population

| PBC |  |  | Average2006-2008 |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | LE of <br> Males <br> [1] | LE of Females [2] | Deaths $<$ LE <br> [3] | $\begin{gathered} \text { Deaths } \\ >\mathrm{LE} \\ {[4]} \end{gathered}$ | YLL <br> [5] | $\begin{gathered} \text { YLG } \\ {[6]} \end{gathered}$ | Net YLL |
| 1 Infectious diseases | 80.7 | 84.4 | 3,710 | 3,248 | 62,052 | 18,796 | 43,256 |
| 2 Cancer | 80.7 | 84.4 | 95,213 | 35,597 | 1,345,038 | 175,350 | 1,169,689 |
| 4 Endocrine | 80.7 | 84.4 | 4,000 | 2,764 | 65,016 | 15,864 | 49,152 |
| (7) Neurological | 80.7 | 84.4 | 8,975 | 6,378 | 145,529 | 34,621 | 110,908 |
| 10 Circulatory | 80.7 | 84.4 | 82,099 | 77,752 | 916,192 | 444,694 | 471,498 |
| 11 Respiratory | 80.7 | 84.4 | 30,500 | 34,945 | 310,334 | 215,829 | 94,505 |
| 13 Gastro-intestinal | 80.7 | 84.4 | 15,827 | 8,320 | 273,308 | 45,295 | 228,012 |
| 17 Genito-urinary | 80.7 | 84.4 | 4,198 | 6,427 | 39,099 | 40,530 | -1,431 |
| 18+19 Maternity \& neonates | 80.7 | 84.4 | 226 | 0 | 17,167 | 0 | 17,167 |

However, these figures are only correct insofar as the distribution of age and gender in each PBC is similar to the general population. For example, if the at risk population tends to be younger the correct LE for the PBC will be lower. A lower LE will mean that there are less deaths below LE each generating fewer YLL, and more deaths above LE each generating more LYG. The net YLL will thus tend to be
lower. Similarly, if the at risk population tends to be older than the general population, the correct LE will be higher and net YLL will also tend to be higher.

This explains the apparent net gain in YLL (negative net YLL) for PBC17 (Genito-urinary) where most deaths occur at ages greater than the LE of the general population so that LYG exceeds YLL. As we are able to show later (see Table C.14) this is because the age distribution in this PBC tends to be older than the general population, i.e., the LE for a matched normal population should be higher with fewer deaths above and more below this LE.

## Using additional information about age and gender distribution

It is evident that estimates of YLL require some account to be taken of counterfactual deaths. In the absence of routinely available information this requires examination of alternative sources of information which might provide a basis for more credible assumptions about the age and gender distribution of the at risk population in each PBC than either, the distribution of observed deaths or of the general population. ${ }^{5}$ The WHO Global Burden of Disease (GBD) study, updated in 2008 using 2004 data (see Addendum 1 for more details ${ }^{6}$ ) provides a range of summary health indicators for the UK, which are, in part, based on estimates of the incidence and duration of sequelae associated with different types of disease by age and gender. Therefore, the type of information used by WHO in the GBD Study to generate summary estimates for the UK can also be used to improve the assumptions required about the age and gender distribution of the PBC populations. Importantly, at this stage, we do not need to rely on estimates of the absolute size of the at risk population, but only the relative 'share' by age and gender.

Specifically, the information reported by GBD (estimates specific to the UK provided in the National Burden of Disease toolkit) reported the incidence and duration of sequelae associated with different types of disease by age and gender. Since it is possible that a patient may experience more than one of the types of sequelae reported in GBD we use the gender and age distribution of the sequelae with the highest prevalence, i.e., the minimum estimate of prevalence consistent with these figures (Addendum 1 to this Appendix), to evaluate the age and gender distribution within each disease.

GBD classifies diseases by U-codes, which are groups of three digit ICD-10 codes (see Addendum 1 to this Appendix for details of how U-codes map to ICD-10 codes). ${ }^{7}$ Since we know which ICD codes contribute to each PBC we can map information from U-codes to PBCs via the ICD codes that contribute to each. The resulting average age and life expectancy for each PBC is reported in columns 3 and 4 of Table C. 13 using the information available from GBD in combination with life tables for the general population.

[^76]Table C.13. Average age and life expectancy for PBCs based on GBD


These summary estimates suggest that some of the PBC populations may be on average older than the general population (e.g., Cancer, Circulatory and Genito-urinary PBCs) or younger (e.g., Maternity \& neonates, Infectious diseases and Neurological). However, when trying to interpret these summaries it should be noted that the average age reflects the age distribution of the sequelae with the highest prevalence. Therefore, a similar average age cân reflect very different age distributions. Some reflect a markedly bimodal distribution, e.g., PBC11, Respiratory, where there is high incidence at very young and older ages (see Figure C.3), or very different age distributions across the type of diseases that contribute to the PBC. For example PBC7 (Neurological) includes dementia which accounts for the vast majority of the PBC population older than 70 . However, a greater proportion of the population is in much younger age groups with other conditions, espeeially migraine (see Addendum 1 to this Appendix for a detailed description of age and gender distributions in all PBCs). When interpreting these summary estimates it should also be noted that the reported life expectancies are not the life expectancies at the average ages reported in column 3 (Table C,13), but the average over the life expectancies for each age group within the contributing ICDs wetghted by the age distribution of the sequelae with maximum prevalence from GBD U-codes.

Figure C.3: Distribution of PBC11 prevalence by age, gender and contributing ICDs, alongside proportion of prevalent patients in the PBC and contribution to variance of each ICD


The implications for net YLL of using these PBC specific estimates of 'normal' life expectancy are reported in Table C.14. As expected, the net YLL for those PBCs with a LE greater than the general population are higher than those reported in column 7 in Table C. 12 (e.g., PBC10 Circulatory and PBC17 Genito-urinary, which now has positive net YLL). Similarly those PBCs with a LE less than the general population have lower net YLL than reported in column 7 in Table C. 12 (e.g., PBC1 Infectious diseases and PBC18 \& 19 Maternity \& neonates, where the effect of a lower LE is more modest as there are no deaths above either of the estimates of LE).

Table C.14. Net YLL using life expectancy for each PBC


The impact on the cost per life year threshold of the issues discussed in this Section is summarised in columns 3 and 4 of Table C. 15 .

Table C.15. Summary of cost per life year threshold


* in PBCs without a mortality signal, health effects were estimated by valuing changes in expenditure at the same rate as observed in PBCs for which there was a mortality signal.

Taking account of counterfactual deaths by calculating net YLL based on the life expectancy of the general population (seecolumn 3) provides similar estimates to those reported in Appendix B. Assuming that PBC populations have the same age and gender distribution as the general population when the, albeit limited, information that is available suggests otherwise, seems inappropriate. Therefore, our preferred central estimate of the cost per life year threshold is reported in column 4 (Table C.15). These are lower than those based on the general population, reflecting the impact on net YLL of evidence that the population at risk in some key PBCs (especially PBC2 and 10) tend to be older than the general population. A detailed breakdown of the changes in spend and YLLs across PBCs that originate this central estimate are presented in columns 5 to 7 of Table C.16. In Section C.2.1.5 we consider extreme upper and lower bounds that might be placed on this central estimate.

Table C.16: LY threshold using net YLL estimates (non-zero health effects for remaining PBCs except GMS).

| PBC | PBC description | Change in spend, $£ \mathrm{~m}$ [1] | Using LE of the GP |  |  | Using LE of the PBC population |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | $\begin{gathered} \text { Net YLL } \\ {[2]} \\ \hline \end{gathered}$ | Change in Net YLL [3] | Cost per LY gained, $£$ [4] | $\begin{gathered} \text { Net YLL } \\ {[5]} \\ \hline \end{gathered}$ | Change in net YLL [6] | Cost per LY gained, £ [7] |
| 2 | Cancer | £19 | 1169689 | 1860 | £10,305 | 1347184 | 2142 | £8,947 |
| 10 | Circulatory problems | £33 | 471498 | 3651 | ¢, 9,112 | 823768 | 6379 | £,5,216 |
| 11 | Respiratory problems | £22 | 94505 | 1683 | £13,256 | 68030 | 1211 | £18,415 |
| 13 | Gastro-intestinal problems <br> Big 4 | £17 | 228012 | 1562 | $\begin{aligned} & £ 10,564 \\ & \quad, 10,421 \end{aligned}$ | 227703 | 1560 | $\begin{array}{r} £ 10,579 \\ £, 8,080 \end{array}$ |
| 1 | Infectious diseases | £8 | 43256 | 16 | $£^{£ 518,314}$ | 36962 | 14 | ¢ 606,574 |
| 4 | Endocrine problems | £18 | 49152 | 394 | $\AA 44,765$ | 51225 | 411 | $\ldots 42,953$ |
| 7 | Neurological problems | £17 | 110908 | 77 | £224,601 | 93917 | 65 | f265,235 |
| 17 | Genito-urinary problems | £32 | - 1431 | - 1 | -£,47,709,995 | 18127 | 8 | \&3,766,371 |
| 16 | Trauma \& injuries* | $£ 10$ | NA | 0 | NA | NA | 0 | NA |
| $18+19$ | Maternity \& neonates* <br> First 11 PBC's | £8 | 17167 | 19 | $\begin{gathered} £ 431,977 \\ £, 19,928 \end{gathered}$ | 16801 | $18$ | $\begin{array}{r} £^{4} 441,387 \\ £, 15,628 \end{array}$ |
| 3 | Disorders of Blood | $£ 11$ |  | 562 | £19,928 |  | 716 | £15,628 |
| 5 | Mental Health Disorders | $£ 204$ |  | 10245 | £19,928 |  | 13064 | £15,628 |
| 6 | Learning Disability | $\ldots 31$ |  | 1545 | £19,928 |  | 1970 | £15,628 |
| 8 | Problems of Vision | £24 |  | 1213 | £19,928 |  | 1547 | £15,628 |
| 9 | Problems of Hearing | $£_{6} 6$ |  | 321 | £19,928 | , | 409 | £15,628 |
| 12 | Dental problems | £23 |  | 1179 | £19,928 |  | 1503 | £15,628 |
| 14 | Skin | £11 |  | 528 | £19,928 |  | 673 | £15,628 |
| 15 | Musculo skeletal system | £15 |  | 759 | £19,928 |  | 968 | £15,628 |
| 20 | Poisoning and AE | £ 4 |  | 220 | £19,928 |  | 281 | £15,628 |
| 21 | Healthy Individuals | £18 |  | 920 | £19,928 |  | 1173 | £15,628 |
| 22 | Social Care Needs | £68 |  | 3393 | *19,928 |  | 4326 | £15,628 |
| 23 | Other | £.78 |  | 0 | NA |  | 0 | NA |
|  | All (23 PBCs) |  |  |  | £22,523 |  |  | £,17,663 |

Note that we have been unable to obtain a satisfactory outcome model for trauma \& injuries and have assumed a zero outcome elasticity.
Note that, for expenditure in 2006/7, the neonate category has been merged with maternity to obtain plausible outcome and expenditure models.

## C.2.1.4 Inferring excess deaths

We have been able to establish a measure of net YLL which takes account of deaths that would have occurred anyway below a normallE for the PBC population (i.e., not all deaths observed in a PBC are excess) and that some deaths observed above this LE would not otherwise have occurred at that age (i.e., some of these deaths are excess). As explained in Section C.2.1.3, net YLL calculated in this way is equivalent to first establishing the number of excess deaths at each age, then calculating YLL for each excess death (based onf the LE conditional on the age at which each excess death occurred) and then summing these (1LD across all excess deaths (i.e., across all ages). In other words, the estimates of net YLL imply a number of excess deaths required to generate them in each PBC. Therefore, it is possible to solve forthe total number of excess deaths based on the net YLL and the average YLL per observed death (the atyerage of the sum of the YLLs for every observed death where the YLL for each observed deathis the difference between age at death and LE conditional on age of death). The net YLL divided by the average YLL per death provides the number of excess deaths required, which on average will generate the estimated net YLL.

In the absence of information about the age distribution of excess deaths, calculations assume that the average YLL associated with observed and excess deaths are similar. Insofar as excess deaths are thought likely to generate more YLL than observed deaths the number of excess deaths will tend to be overestimated. This would tend to underestimate the cost per excess death averted. However, the cost per life year estimates remain unchanged and do not require such an assumption.

The implied excess deaths associated with net YLL based on the LE of the PBCs (see column 7 Table C.14) are reported in Table C.17. With the exception of PBC18\&19, excess deaths are some proportion of total observed deaths in each PBC. The proportion of excess deaths differs by PBC reflecting the distribution of deaths relative to the LE of the PBC. For example, in those PBCs where a large proportion of deaths occur below LE (see column 3 and 4) excess deaths tend to be greater proportion of total deaths (e.g., PBC2, 13 and 10). Where most deaths occur above LE excess deaths as a proportion of total deaths tend to be lower (e.g., PBC11, 17 and 1). Nevertheless, the impact of the age distribution of deaths and the age distribution of the at risk population (summarised as LE) on the calculation of excess deaths is not always obvious as both will affect the numerator (net YLL) as well the denominator (average YLL per death) in this calculation.

Table C.17: Excess deaths implied by net YLL.


Excess deaths are calculated for each gender by dividing net YLLs by the YLbper death (column [3] = column [1] / column [2] ) * The number of excess deaths estimated in PBC18\&19 was initially estimated to be 230, higher than the number of total deaths. This is due to the use of approximations (i.e. in the life expectancy, or in using the net YLL) thus, for consistency, we assumed this to be $100 \%$ of the total deaths.

Estimates of net YLL and changes in life years due to expenditure (see Table C. 14 and C.15) have already accounted for the fact that not all deaths are excess and do not generate YLL. Nevertheless, solving for the number of implied excess deaths associated with these net YLL estimates allows a comparison of the cost per excess and observed PBC death avoided and an examination of the interpretation that can be placed of the life years expected to be gained from an excess or observed death averted.

Since only deaths observed in the PBC can be used to estimate the effects of expenditure (excess deaths are not directly observed since they rely on an unobserved counterfactual population and would occur outside the PBC), the outcome elasticities can be interpreted as the proportionate change in observed PBC mortality due to a proportionate change in PBC expenditure. Equally, however, they can also be interpreted as the proportionate effect on excess death due to a proportionate change in expenditure so can be applied to either total observed or total excess deaths. Observed PBC mortality that is sensitive to changes in expenditure can be regarded as 'avoidable' and it is only this mortality that contributes to the estimates of outcome elasticities (not all observed mortality is necessarily avoidable and sensitive to expenditure such mortality will not contribute to the estimates). Not all observed mortality is excess when compared to the counterfactual population but this is unrelated to the question of how sensitive it is to expenditure, i.e., observed mortality will be just as sensitive to expenditure whether or not it is regarded as excess. Therefore, the estimated outcome elasticities can be applied to either observed PBC deaths or excess PBC deaths

The cost per excess death and the cost per PBC death averted are reported in Table C.18, and a detailed breakdown of changes in spend and excess or total deaths across PBCs is shown in Table C.19. The cost per PBC death averted is, of course; significantly lower than the cost per excess death as excess deaths are only a proportion of total deaths (see Table C.17). Also the cost per PBC death averted are substantially lower than those reported in Appendix B (see Tables B8.22 and B8.23), since these estimates do not restrict the effects of expenditure to PBC deaths under 75.

Table C.18. Summary of the cost per death averted threshold

|  | Cost per excess death averted, $£$ [1] | Cost per PBC death averted, $£$ [2] |
| :---: | :---: | :---: |
| big 4 PBC's | $£(1,129$ | £32,864 |
| 11 PBCs (with mortality) | £177,692 | ¢ 64,774 |
| All 23 PBCs (zero health effects for remaining 12 PBCs ) | £653,748 | £238,310 |
| All 23 PBCs (non-zero health effects for remaining 12 PBCs, except GMS)* | £200,829 | £ 73,208 |

* in PBCs without a mortality signal, health effects were estimated by valuing changes in expenditure at the same rate as observed in PBCs for which there was a mortality signal.

Table C.19: Breakdown of the cost per death averted threshold.

| PBC | PBC description | Change in spend, £m [1] | Total <br> PBC <br> deaths <br> [2] | PBC dea Change in PBC deaths [3] | Cost per PBC death averted, £ [4] | Excess deaths [5] | Excess de Change in excess deaths [6] | hs <br> Cost per excess death averted, £ 21 [7] |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 2 | Cancer | £.19 | 130809 | 208.03 |  | 95715 | 152.22 | 6,125,934 |
| 10 | Circulatory problems | £33 | 159851 | 1237.82 | £26,878 | 79218 | 613.43 | £54,235 |
| 11 | Respiratory problems | £22 | 65446 | 1165.14 | £19,142 | 7386 | 131.49 | £,169,616 |
| 13 | Gastro-intestinal problems Big 4 | £.17 | 24148 | 165.42 | $\begin{array}{r} £ 99,757 \\ £ 32,864 \end{array}$ | $\begin{gathered} 15199 \\ 0 \end{gathered}$ | $104,12$ | $\begin{array}{r} \AA 158,488 \\ \AA 91,129 \\ \hline \end{array}$ |
| 1 | Infectious diseases | £8 | 6958 | 2.59 | £3,222,218 | 2797 | 1.04 | £8,014,595 |
| 4 | Endocrine problems | £18 | 6765 | 54.28 | £325,291 | 3769 | 30.24 | £583,830 |
| 7 | Neurological problems | £17 | 15353 | 10.59 | £1,622,486 | 6)909 | 4.77 | £3, ${ }^{\text {, }}$, 05,579 |
| 17 | Genito-urinary problems | £.32 | 10625 | 4.94 | £,6,425,694 | 2172 | 1.01 | $£ 31,430,287$ |
| 16 | Trauma \& injuries* | £10 | NA | 0 | NA | NA | 0 | NA |
| $18+19$ | Maternity \& neonates* First 11 PBC's | $\AA 8$ | 226 | 0.24 | $\begin{array}{r} £ 32,813,038 \\ 64,774 \end{array}$ | 226 | 0.24 | $\begin{array}{r} £ 32,813,038 \\ £ 177,691 \end{array}$ |
| 3 | Disorders of Blood | £.11 |  | 172.87 | - 564,774 |  | 63.01 | £,177,692 |
| 5 | Mental Health Disorders | £204 |  | 3152.02 | C64,774 |  | 1149.00 | £177,692 |
| 6 | Learning Disability | £31 |  | 475.30 | £64,774 |  | 173.26 | £177,692 |
| 8 | Problems of Vision | $£ 24$ |  | 373.19 | $¢_{6}^{64,774}$ |  | 136.04 | £177,692 |
| 9 | Problems of Hearing | $\mathrm{f}_{6} 6$ |  | 98.72 | £64,774 |  | 35.99 | £177,692 |
| 12 | Dental problems | £23 |  | 362.72 | $¢_{6} 64,774$ |  | 132.22 | £177,692 |
| 14 | Skin | £11 |  | 162.30 | ¢ 64,774 |  | 59.16 | £177,692 |
| 15 | Musculo skeletal system | £15 |  | 233.59 | ¢ 64,774 |  | 85.15 | £,177,692 |
| 20 | Poisoning and AE | $£ 4$ |  | 67.80 | $¢_{6} 64,774$ |  | 24.71 | £177,692 |
| 21 | Healthy Individuals | £18 |  | 283.09 | $¢_{6}^{6} 64,774$ |  | 103.19 | £177,692 |
| 22 | Social Care Needs | £68 |  | 1043.74 | £64,774 |  | 380.47 | £177,692 |
| 23 | Other | £ 78 |  | 0 | ¢ 0 |  | 0 | NA |
|  | All (23 PBCs) | S |  |  | £,73,208 |  |  | £200,828 |

Recall from Appendix B that the measure of mortality that is available at PCT level and used to estimate the outcome elasticities is restricted to deaths under 75, as are the published estimates of YLL associated with them (see Section)C.2.1.1). However, to restrict effects only to those under 75 would imply that there is no excess mortality above 75 or equivalently that there are no health effects of PBC expenditure above 75. Rather than assume no effects of NHS activity in older populations we apply the effects that can be observed to the whole PBC but account for deaths that would have otherwise occurred in our estimate of net YLL in Section C.2.1.3. Table C. 20 illustrates the number deaths averted for a $1 \%$ change in bagget implicit in the alternative calculations of the cost per death averted threshold.

Table C.20: Illustration of the number of deaths averted for a $1 \%$ change in budget

|  | $\begin{gathered} \text { Using deaths }<75 \\ \text { (Appendix B, Table B8.21) } \end{gathered}$ |  | Using excess deaths (Table C.18) |  | Using PBC deaths (Table C.18) |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Cost per death averted (<75), £ <br> [1] | Number of deaths averted ( $<75$ ) for a $1 \%$ change in budget [2] | Cost per excess death averted, $£$ [3] | Number of excess deaths averted for a $1 \%$ change in budget [4] | Cost per PBC death averted, $£$ [5] | Number of PBC deaths averted for a $1 \%$ change in budget [6] |
| big 4 PBC's | £137,188 | 665 | ¢ 917129 | 1,001 | £32,864 | 2,776 |
| 11 PBCs (with mortality) | £270,881 | 681 | £177,692 | 1,039 | ¢64,774 | 2,849 |
| All 23 PBCs (zero health effects for remaining 12 PBCs) | $£ 996,655$ | 681 | £653,748 | 1,039 | £238,310 | $2,849$ |
| All 23 PBCs (non-zero <br> health effects for remaining 12 PBCs, except GMS)* | £306,153 | 2,218 | $£ 200,828$ | 3,381 | £73,208 | $5,191$ |

* in PBCs without a mortality signal, health effects were estimated by valuing changes in expenditure at the same rate as observed in PBCs for which there was a mortality signal.

In many respects, whether or not PBC deaths at older ages are as sensitive to changes in expenditure is not critical since any observed deaths that might be averted at older ages are less likely to generate life years gained because they are more likely to have occurred anyway in that year (i.e., are excess so generate zero life years gained anyway). Therefore, they will have very limited impact on cost per life year or subsequently on cost per QALY estimates (in Sections C.2.2 and C.2.3). For this reason, it is the cost per life year rather than cost per death averted, whether excess or observed, that is of primary interest. The cost per PBC or excess death averted (or life saved) should thus not be over interpreted since lives are never saved (death is only delayed). However, establishing the number of excess and PBC deaths averted which are associated with net YLL is useful because it enables an assessment of the number of life years gained associated with each death averted. These are reported for each PBC in Table C. 21 and range from 74.3 years per excess death for PBC $18 \& 19$ Maternity \& neonates to 8.3 per excess death for PBC17 Genito-urinary. On average, across all 11 PBCs each excess death averted is associated with 11.4 life years gained.

Table C.21: Implied YLL per death averted for each PBC

| PBC PBC description |  | Implied YLL per PBC death averted <br> [2] |
| :---: | :---: | :---: |
| 2 Cancer | 14.07 | 10.30 |
| 10 Circulatory problems | 10.40 | 5.15 |
| 11 Respiratory problems | 9.21 | 1.04 |
| 13 Gastro-intestina problems | 14.98 | 9.43 |
| Big 4 | 11.28 | 4.07 |
| 1 Infectious diseases | 13.21 | 5.31 |
| 4 Endocrine problems | 13.59 | 7.57 |
| 7 Neurological problems | 13.59 | 6.12 |
| 17 Genito-urinary problems | 8.34 | 1.71 |
| 16 Trauma \& injuries | NA | NA |
| 18+19 Maternity \& neonates | 74.34 | 74.34 |
| First 11 PBC's | 11.37 | 4.14 |

Hoyvever, clinicians or the evaluative literature cannot distinguish whether an observed death is excess or not. What can be observed is whether groups of similar patients with and without access to a treatment survive and for how long. Therefore, it is the life years associated with each observed death that provides a context that can be interpreted based on experience and evidence of how effective those interventions that could be invested or disinvested tend to be. The average life years expected to be gained associated with each observed PBC deaths averted takes account of that fact that some deaths that are avoided in the PBC are not delayed for very long but quickly occur elsewhere and do not generate LY gained (i.e., they were not excess deaths). The portion of observed deaths that are regarded as excess depend on how time is discretised. The data available reports deaths in annual intervals so in this context 'quickly' means within one year. If deaths were reported in narrower time intervals then a greater proportion of observed
deaths would be regarded as excess and, in the limit, with continuous time all observed deaths would be excess. Of course, the average YLL associated with them would be smaller and is approximated by the net YLLs per observed death reported (the effects of approximation is likely to be small but unavoidable as it is due to deaths being reported in annual intervals).

However, establishing the number of excess and PBC deaths averted which are associated with net YLL is useful because it enables an assessment of the number of life years gained associated with each death averted. On average across all 11 PBCs each excess death averted is associated with 11.4 life years gained. These are reported for each PBC in Table C21 in Appendix C and range from 74.3 years per excess death for PBC $18 \& 19$ Maternity \& neonates to 8.3 for PBC17 Genito-urinary. However, clinicians or the evaluative literature cannot distinguish whether an observed death is excess or not. What can be observed is whether groups of similar patients with and without access to a treatment survive and for how long. Therefore, it is the life years associated with each observed death that provides a context that can be interpreted based on experience and evidence of how effective those interventions that could be invested or disinvested tend to be. The average life years expected to be gained associated with each observed PBC deaths averted takes account of that fact that some deaths that are avoided in the PBC are not delayed for very long but quickly occur elsewhere and do not generate LY gained (i.e., they were not excess deaths). What portion of observed deaths are regarded as excess depend on how time is discretised.

The data available reports deaths in annual intervals so in this context quickly means within one year. If deaths were reported in narrower time intervals then a greater proportion of observed deaths would be regarded as excess and in the limit with continuous time all observed deaths would be excess. Of course, the average YLL associated with them would be smaller and is approximated by the net YLLs reported in Table 4.5 per observed death (the effects of approximation is likely to be small but unavoidable as it is due to deaths being reported in annual intervals). These are also reported for each PBC in Table C 21 in Appendix C and range from 74.3 years per observed death for PBC $18 \& 19$ Maternity \& neonates ${ }^{8}$ to 1.0 for PBC11 Respiratory problems, i.e., the YLL per PBC death are much lower for those PBCs where a small proportion of observed deaths are excess. On average across all 11 PBCs each PBC death averted is associated with 4.1 life years gained.

## C.2.1.5 Summary of cost per life year estimates

The sequence of analysis set out above has enabled an examination of the impact of the limitations associated with the incomplete reporting mortality data at PCT level and incomplete coverage of published YLL estimates. We have also been able to consider effects above 75 while taking account of that fact that many deaths would have occurred anyway, despite the limited information available about the population at risk within a PBC. The GBD Study does provide some information about the age and gender distribution of the population at risk in a PBC so offers some improvement over the other assumptions that would otherwise be required (i.e., that the distribution of age and gender is the same as the general population or follows the distribution of observed deaths). For this reason the cost per life year threshold in column 4 of Table C. 15 and repeated in lines 1 to 4 in Table C. 22 are regarded as the central or best estimates given the evidence available and the credibility of alternative assumption that could be made. As explained in Section C.1, these are based on the conservative assumption that any health effects of changes in expenditure are restricted to one year, which, to some extent, may be offset by the more optimistic assumption any death averted returns the individual to the mortality risk faced by the general population, matched for age and gender.
${ }^{8}$ This is the same as life years associated with excess deaths since all observed deaths in this PBC are excess.

Table C.22: Summary of the cost per life year threshold with upper and lower bounds

| Effect of expenditure on mortality: YLL per PBC death averted: | Best estimate 1 year ~ 4.1 YLL |  |
| :---: | :---: | :---: |
| big 4 PBC's <br> 11 PBCs (with mortality) <br> All 23 PBCs (zero health effects for remaining 12 PBCs) <br> All 23 PBCs (non-zero health effects for remaining 12 PBCs, except GMS)* | $\AA 8,080$ $£ 15,628$ $£ 57,497$ $£ 17,663$ | $\begin{aligned} & {[1]} \\ & {[2]} \\ & {[3]} \\ & {[4]} \end{aligned}$ |
| Effect of expenditure on mortality: YLL per PBC death averted: | Lower bound Remainder of disease ~ 4.1 YLL |  |
| big 4 PBC's <br> 11 PBCs (with mortality) <br> All 23 PBCs (zero health effects for remaining 12 PBCs) <br> All 23 PBCs (non-zero health effects for remaining 12 PBCs, except GMS)* | $\begin{aligned} & £_{0} 3,846 \\ & £_{0}, 106 \\ & £_{2}^{22,463} \\ & £ 6,901 \\ & \hline \end{aligned}$ | $\begin{aligned} & {[5]} \\ & {[6]} \\ & {[7]} \\ & {[8]} \\ & \hline \end{aligned}$ |
| Effect of expenditure on mortality: YLL per PBC death averted: | Upper bound <br> 1 year <br> 2 YLL |  |
| big 4 PBC's <br> 11 PBCs (with mortality) <br> All 23 PBCs (zero health effects for remaining 12 PBCs) <br> All 23 PBCs (non-zero health effects for remaining 12 PBCs, except GMS)* | $\begin{gathered} £, 16,432 \\ £, 32,387 \\ £, 119,155 \\ £ 36,604 \end{gathered}$ | $\begin{gathered} {[9]} \\ {[10]} \\ {[11]} \\ {[12]} \\ \hline \end{gathered}$ |

* in PBCs without a mortality signal, health effects were estimated by valuing changes in expenditure at the same rate as observed in PBCs for which there was a mortality signal.

It does not seem credible to imagine that NHS expenditure has no health effects in the 12 PBCs which do not have sufficient mortality reported at PCT level to estimate outcome elasticities - what is implied by the estimate reported in line 3. Therefore, it is the estimates reported in lines 2 and 4 that are of policy interest. The estimate of $£ 15,628$ per life year (line 2 ) is restricted to the effects of changes in expenditure in the 11 PBCs where outcome elasticities can be estimated. The threshold of $£ 17,663$ per life year uses the estimated health effects of expenditure in these PBC as a surrogate for health effects in the others, i.e., assuming that the effects that can be observed will be similar to those that cannot. However, no health effects are assigned to PBC23 (General Medical Services) on the basis that any health effects of this expenditure would be recorded in the other PBCs.

It would be inappropriate to assign all the change in GMS expenditure to the estimate of cost per life year based only on the 11 PBCs with outcome elasticities because it would imply that GMS only contributes to these PBCs. Restricting attention to the 11 PBCs with outcome elasticities but allocating part of the change in GMS expenditure to them based on their proportional share of changes in overall expenditure would yield the same cost per life year as reported in line 4.

The extreme upper and lower bounds for the cost per life year thresholds in Table C. 22 are based on making the necessary assumptions about duration of health effects and how long a death might be averted optimistic (providing the lower bound for the threshold) or conservative (an upper bound for the threshold). The (lower bound (lines 5 to 8 ) is based on assuming that health effects are not restricted to one year but apply to the whole of the remaining disease duration of the population at risk in PBCs during the expenditure year. Estimates of the average disease durations across the PBCs used in this calculationare depicted in Table C. 23 (column 2). ${ }^{9}$ These were obtained from the GBD Study (see Addendum 1 to this Appendix). Although this lower bound for the threshold combines optimistic assumptions, it is possible, indeed likely, that at least some expenditure may have effects on the health outcomes of future patients that are not currently part of the population at risk in a PBC, e.g., investments or disinvestment in prevention will have an impact on populations that are incident to PBCs in the future. Such effects are not captured in any of the estimates presented in this chapter so all are conservative with respect to this type of health effects from changes in expenditure.

[^77]Table C.23: Disease duration by PBC (GBD).

| PBC | Duration of disease for <br> an incident patient <br> (years), GBD <br> $[1]$ | Remaining duration of <br> disease for at risk <br> population (years), GBD <br> $[2]$ |
| :---: | :---: | :---: |
| 1 | 6.21 | 3.11 |
| 2 | 1.19 | 0.59 |
| 3 | 1.07 | 0.53 |
| 4 | 24.83 | 12.42 |
| 5 | 7.41 | 3.70 |
| 6 | 3.46 | 1.73 |
| 7 | 30.91 | 15.45 |
| 8 | 13.96 | 6.98 |
| 9 | 16.40 | 8.20 |
| 10 | 3.21 | 1.61 |
| 11 | 11.24 | 5.62 |
| 12 | 0.33 | 0.17 |
| 13 | 0.27 | 0.13 |
| 14 | 1.01 | 0.50 |
| 15 | 9.56 | 4.78 |
| 16 | 3.74 | 1.87 |
| 17 | 1.11 | 0.56 |
| 18 | 0.58 | 0.29 |
| 19 | 9.71 | 4.86 |
| 20 | 0.93 | 0.47 |
| 21 | 1.07 | 0.53 |
| 22 | 3.74 | 1.87 |
| 23 | 3.74 | 1.87 |

The upper bound (lines 9 to 12 in Table C.22) is based on the combination of assuming that health effects are restricted to one year for the population currently at risk and that any death averted is only averted for the minimum duration consistent with the mortality data. The econometrics work used the average of 3 years of mortality ( 2006 to 2008), so the estimated outcome elasticities are based on differences in mortality that remain after averaging over three years. Therefore, the estimated effects are based on differences in observed PBC deaths that must have been sustained, on average, for more than a minimum of 2 years. This is because whilst variation in mortality the first year of data will only contribute to estimates if differences are sustained for a minimum of 3 years, variation in mortality in the second year will only contribute if it is sustaned for a minimum of 2 years, and in the third year only if sustained for 1 year. If differences in mortality are similar each year (the three years contribute equally to the estimates) then estimated effects must have been sustained, on average, for a minimum of 2 years. ${ }^{10}$ These estimates can be interpreted as an upper bound given the data available and therefore the analysis that has been feasible.

## C.2.2 Adjusting life years for quality of life

The central orbest estimates of the cost per life year threshold, which were presented in Table C. 22 (lines 2 and 4) take no account of the health related quality of life in which years of life, expected to be gained or lost through changes in expenditure, are likely to be lived. Even if attention is restricted to the direct health consequences of changes in mortality, estimates of the cost per life year will tend to overestimate the effects of changes in expenditure (underestimate the threshold) compared to a more complete measure of health that accounts for the quality in which the years of life are expected to be lived. In this Section we examine the ways in which the life years reported in Section C.2.1 can be adjusted for quality, taking account of information that is available about: i) how quality of life differs by age and gender (see Section C.2.1), and ii) how the quality of life years associated with mortality changes might be affected by the types of diseases that make up each PBC (see Section C.2.2). Throughout we continue to take account for counterfactual deaths in the way described in Section C.2.1.3 by making the adjustment for

[^78]quality to the life years associated with every observed death before calculating a quality adjusted net YLL. The implications for a cost per quality adjusted life year (QALY) threshold that only accounts for the health effects of mortality changes are presented in Section C.2.3.2. In Section C.2.3 we explore the ways in which the likely direct effects of expenditure on quality of life (other than through mortality) might also be taken into account.

## C.2.2.1 Quality of life based on the general population

The most commonly used metric of health related quality of life in the UK is EQ5D,[1] which is specified in the NICE reference case for methods of technology appraisal.[2] This metric has 5 dimensions of quality each with three possible levels. Each of these 243 possible health states is valued relative to a score of one, which represents full or best imaginable health (the best score across all 5 dimensions), and a score of zero, which represents death, based on a representative sample of the UK population. [3] Therefore, insofar as the years of life expected gained or lost through changes in expenditure would be lived in this state of full health, the cost per life year thresholds reported in Table C. 22 would also be the cost per QALY thresholds, albeit ones that only account for the health effects of mortality changes.

However, unsurprisingly, there is good evidence that, on average, the general population is not in this state of full health. Therefore, the quality of life score associated with the health states experienced by the general population are less than 1 , and are expected to decline with age and to differ by gender. These quality of life 'norms' for the general population by age and gender are illustrated in Figure C. 4 based on an analysis of data from the Health Survey for England (HSE, see Addendum 1 to this Appendix for a description on HSE data and the analysis of quality of life norms illustrated in Figure C.4).


Figure C.4: Quality of life for the general population by age and gender
These quality of life norms can be applied to the YLL associated with all observed deaths in each PBC, taking account of gender and age at death. The results are reported in column 4 to 6 of Table C.24.

Table C.24: Net YLL adjusted for quality of life 'norms'

|  |  | Unadjusted life years |  |  | Quality adjusted life years |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | $\begin{gathered} \text { YLL } \\ {[1]} \\ \hline \end{gathered}$ | $\begin{gathered} \text { YLG } \\ {[2]} \\ \hline \end{gathered}$ | Net YLL [3] | $\begin{gathered} \text { YLL } \\ {[4]} \\ \hline \end{gathered}$ | $\begin{gathered} \text { YLG } \\ \hline[5] \\ \hline \end{gathered}$ | Net YLL <br> [6] |
| $\checkmark 1$ | Infectious diseases | 58,686 | 21,724 | 36,962 | 47,481 | 14,618 | 32,864 |
| 2 | Cancer | 1,473,733 | 126,549 | 1,347,184 | 1,143,445 | 84,036 | 1,059,409 |
| 4 | Endocrine | 66,283 | 15,058 | 51,225 | 52,856 | 9,973 | 42,883 |
| 7 | Neurological | 135,686 | 41,770 | 93,917 | 109,349 | 28,262 | 81,087 |
| 10 | Circulatory | 1,102,020 | 278,251 | 823,768 | 848,046 | 183,330 | 664,717 |
| 11 | Respiratory | 298,343 | 230,313 | 68,030 | 231,578 | 154,743 | 76,835 |
| 13 | Gastro-intestinal | 273,117 | 45,414 | 227,703 | 216,256 | 30,277 | 185,979 |
| 17 | Genito-urinary | 47,229 | 29,101 | 18,127 | 35,929 | 18,947 | 16,982 |
| 18+19 | Maternity \& neonates | 16,801 | 0 | 16,801 | 14,568 | 0 | 14,568 |

Recall from Section C.2.1.3 that taking account of counterfactual deaths requires calculation of the YLL associated with deaths below LE (of a normal population matched to the age and gender distribution in the PBC) and the implied YLG of deaths that occur above this LE. There are two effects of adjusting life years for quality: i) since quality of life norms are always less than 1 the adjusted YLL and YLG are always lower than the unadjusted values in columns 1 and 2 (previously reported in Table C.5); and ii) deaths above LE are necessarily at older ages with poorer quality of life norms than those below, so the difference between adjusted and unadjusted values is greater for YLG than YLL (Table C. 25 illustrates these effects by showing the implied QoL scores applied to YLL and YLG). The overall effect of quality adjustment on net YLL is the balance of these two effects, and tends to reduce the net YLL (compare column 6 and 3 in Table C.24). The only exception is PBC11 (Respiratory) which has a large proportion of deaths occurring above the life expectancy of the PBC population (see Table C.14).

Table C.25: Implied quality of life score in the net YLL adjustment for quality of life 'norms'

| PBC |  | QoL score for YLL | QoL score for YLG |
| ---: | :--- | :---: | :---: |
| 1 | Infectious diseases | 0.81 | 0.67 |
| 2 | Cancer | 0.78 | 0.66 |
| 4 | Endocrine | 0.80 | 0.66 |
| 7 | Neurological | 0.81 | 0.68 |
| 10 | Circulatory | 0.77 | 0.67 |
| 11 | Respiratory | 0.78 | 0.67 |
| 13 | Gastro-intestinal | 0.79 | 0.65 |
| 17 | Genito-urinary | 0.76 |  |
| $18+19$ | Maternity \& neonates | 0.87 |  |

The quality adjusted net YLL figure in Column 6 suggest that the health effects of mortality are lower than when relying only on unadjusted life years in Section \&.2.1.3. Therefore, the health effects of changes in expenditure on this more complete measure of health will also be lower. The implications of these adjustments to a cost per QALY threshold that only accounts for the direct health effects of mortality are summarised in Table C.26, and detailed in Table C.27. As expected, the cost per QALY threshold based on adjusting the life years gained or lost (column 2, Table C.26) is higher than a threshold based on unadjusted life years (column 1 in Tables C.26, these results were previously reported in Tables C. 15 and C.22).

Table C.26: Summary of cost per QALYthreshold based on population norms and mortality effects

|  | Cost per life year threshold <br> [1] | Cost per QALY threshold Population norms [2] |
| :---: | :---: | :---: |
| big 4 PBC's | £,8,080 | £,9,631 |
| 11 PBCs (with mortality) | £15,628 | £18,622 |
| All 23 PBCs (zero health effects for remaining 12 PBCs) | $\uparrow 57,497$ | $f .68,513$ |
| All 23 PBCs (non-zero health effects for remaining 12 PBCs , except GMS )* | $£, 17,663$ | $£ 21,047$ |

* in PBCs without a mortality signal, health effects were estimated by valuing changes in expenditure at the same rate as observed in PBCs for which there was a mortality signal except GMS.

Table C.27: A breakdown of the cost per QALY threshold based on population norms


Table C. 28 depicts the judgements over life years, quality of life weights and total QALYs implicit in calculations of the threshold cost per QALY in Table C.26. Specifically, columns 1 and 2 of Table C. 28 report the number of life years associated with each death averted for each PBC; as expected, the values are equal to those in Table C. 21 as estimates rely on the net YLLs evaluated in Section C.2.1.3. In columns 3 and 4, the number of QALYs gained associated with each death averted are presented. These ranged from 64.46 QALYs gained per PBC death âverted for PBCs18\&19 (Maternity and Neonates) to 1.17 QALYs per PBC death averted for PBC11 (Respiratory) - column 4. In general, these values are expected to be smaller than the unadjusted YLL per PBC death averted in column 2 . The exception is PBC 11 (respiratory) - in this PBC, the number of YLL and YLG are more similar than in other PBCs (respectively, columns 1 and 2 of Table C.24), and given that YLGs are weighted more heavily (with lower QoL scores) than YLL, the netting of adjusted estimates returns a higher number than the netting of unadjusted estimates. On average, across all 11 PBCs each PBC death averted is associated with 3.5 QALYs gained.

Table C.28: Implied YLL per excess death averted and implied QoL score per YLL gained, for each PBC

|  | Implied YLL per <br> excess death averted <br> [1] | Implied YLL per PBC death averted [2] | Implied QALYs gained per excess death averted $[3]$ | ```Implied QALYs gained per PBC death averted [4]``` |
| :---: | :---: | :---: | :---: | :---: |
| 2 Cancer | 14.07 | 10.30 | 11.07 | 8.10 |
| 10. Circulatory | 10.40 | 5.15 | 8.39 | 4.16 |
| 11 Respiratory | 9.21 | 1.04 | 10.40 | 1.17 |
| 13 Gastro-intestinal | 14.98 | 9.43 | 12.24 | 7.70 |
| Pig 4 | 11.28 | 4.07 | 9.46 | 3.41 |
| 1 Infectious diseases | 13.21 | 5.31 | 11.75 | 4.72 |
| 4 Endocrine | 13.59 | 7.57 | 11.38 | 6.34 |
| 7 Neurological | 13.59 | 6.12 | 11.74 | 5.28 |
| 17 Genito-urinary | 8.34 | 1.71 | 7.82 | 1.60 |
| 16 Trauma \& injuries* | NA | NA | NA | NA |
| 18+19 Maternity \& neonates* | 74.34 | 74.34 | 64.46 | 64.46 |
| First 11 PBC's | 11.37 | 4.14 | 9.54 | 3.48 |

## C.2.2.2 Adjusting age related quality of life for disease decrements

Adjusting life years for age and gender related quality of life norms assumes that any life year gained through a change in expenditure would be lived in a similar quality of life to the general population. It is possible however, that patients benefiting from reduced mortality may, nevertheless, continue to be affected by the type of diseases that make up each PBC and experience the quality of life associated with the original disease.

The Health Outcome Data Repository (HODaR)[4] provides over 30,000 observations of EQ-5D measures of quality of life by ICD code and the age and gender of the patients in the sample (see Addendum 1 to this Appendix). Although this is a rich UK data set, there were a limited number of observations for some of the less common ICD codes. For this reason HODaR was supplemented with information from the Medical Expenditure Panel Survey (MEPS)[5] which also provides EQ-5Dby ICD and reports the average age of respondents (see Addendum 1 to this Appendix). These data provided a means of estimating the quality of life associated with each ICD code at the average age of respondents in the pooled sample (ICD estimates of the quality of life score and age were pooled across datasets by considering the number of patients from each dataset contributing to estimates, i.e. a weighted average). The quality of life associated with each PBC was then expressed as the average of the quality of life associated with its component ICDs. The average quality of life scores across IDDs which contribute to each PBC and the average age and gender of respondents were used to calculate a PBC disease related decrement (disutility) based on quality of life norms from the general population - it is important to note that by expressing the quality of life effects of different diseases as age related decrements we do not require the HODaR and MEPS samples to necessarily be representative of the age distribution of the population at risk in the PBCs.

Table C. 29 summarises the data from HoDAR and MEPS and the quality of life decrements used further in calculations of the threshold, namely: the number of patients for which quality of life scores were available (column 1), the average age of these patients by gender (columns 2 and 3), the average quality of life scores across PBCs (column 4), the quality of life scores for the population norms by gender (columns 5 and 6), and the calculated disease related decrements (columns 7 and 8).

Table C.29: QoL scores per PBC from different sources

no gender details were available from MEPS so assumed 50:50 split of frequency
only primary diagnosis is used from HoDAR data
a lower bound of 0 is assumed for disutility for each PBC

Figure C. 5 illustrates the use of the decrement to quality of life norms for PBC1 (Infectious disease) across a range of ages. For PBC1, the quality of life score was evaluated across the component ICD codes was evaluated to be 0.667 in HoDAR and MEPS, at an average age of 54 for male respondents. Since the quality of life norms for males age 54 is 0.859 this suggests a decrement associated with membership of PBC1 of 0.192 , which can then be applied to quality of life norms by age as illustrated in Figure C.5.


Figure C.5: Quality of life for males in PBC1 (Infectious disease) and the general population by age

In principle, it would be possible to estimate disease related disutility byage rather than assume a fixed additive decrement. HODaR does provide age for each reported quality of life score but MEPs only provides average age of respondents in published summaries. However, even with access to 'raw' scores and the age and gender of each, it is very unlikely that there would be sufficient data to estimate age related decrements in each of the component ICDs. It would, however, be possible to assume a proportionate rather than fixed decrement by age. However, the average age of respondents in the pooled HODaR and MEPs sample (columns 2 and 3 of Table C.29) tends to be older than the age distribution of the PBC populations (columns 3 and 4 of Table C.13). Given that older individuals are expected to have a lower quality of life (norm), relative decrements can overestimate the decrements observed in younger patients. By applying overestimated decrements, the quality adjusted net YLL would be underestimated and the cost per QALY threshold increased compared to the fixed decrement applied here.

Quality of life norms adjusted for disease related decrements can be applied to the YLL associated with observed deaths in each PBC, taking account of gender and age at death in the same way as Section C.3.1. To do so, the 'PBC decrements' calculated from HoDAR and MEPS were applied to each observed death and the age at whicheach life year was gained or lost (from ONS). The results are reported in columns 4 to 6 of Table C.30. The overall effect of quality adjustment that also applies a disease related decrement is to reduce the net YLL to a greater extent than adjustment with population norms alone (compare columns 6 in Table C. 30 and C.24).

Table C.30: Net YLL adjusted for disease and age related quality of life

| PBC |  | Unadjusted life years |  |  | Quality adjusted life years |  |  |
| :---: | :--- | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | YLL | YLG | YLL | YLG | YLL | YLG |
|  |  | $[1]$ | $[2]$ | $[3]$ | $[4]$ | $[6]$ |  |
| 1 | Infectious diseases | 58,686 | 21,724 | 36,962 | 37,055 | 10,793 | 26,262 |
| 2 | Cancer | $1,473,733$ | 126,549 | $1,347,184$ | 955,690 | 67,930 | 887,760 |
| 4 | Endocrine | 66,283 | 15,058 | 51,225 | 43,394 | 7,844 | 35,550 |
| 7 | Neurological | 135,686 | 41,770 | 93,917 | 68,893 | 15,842 | 53,050 |
| 10 | Circulatory | $1,102,020$ | 278,251 | 823,768 | 656,145 | 135,241 | 520,905 |
| 11 | Respiratory | 298,343 | 230,313 | 68,030 | 169,269 | 106,505 | 62,764 |
| 13 | Gastro-intestinal | 273,117 | 45,414 | 227,703 | 163,593 | 21,677 | 141,916 |
| 17 | Genito-urinary | 47,229 | 29,101 | 18,127 | 29,749 | 15,152 | 14,598 |
| $18+19$ | Maternity \& neonates | 16,801 | 0 | 16,801 | 13,662 | 0 | 13,662 |

The implied quality of life weights (considering the disease related decrements) for YLL and YLG are shown in Table C.31. Note that, as expected, the weights assume a lower value than in Table C. 25.

Table C.31: Implied QoL weights in the net YLL adjusted for disease and age related quality of life

| PBC |  | QoL weights <br> for YLL <br> $[1]$ | QoL weights <br> for YLG <br> $[2]$ |
| :--- | :--- | :---: | :---: |
| 1 | Infectious diseases | 0.63 | 0.50 |
| 2 | Cancer | 0.65 | 0.54 |
| 4 | Endocrine | 0.65 | 0.52 |
| 7 | Neurological | 0.51 | 0.38 |
| 10 | Circulatory | 0.60 | 0.49 |
| 11 | Respiratory | 0.57 | 0.46 |
| 13 | Gastro-intestinal | 0.60 | 0.48 |
| 17 | Genito-urinary | 0.63 | 0.52 |
| $18+19$ | Maternity \& neonates | 0.81 | NA |

Combining quality of life adjustments for both population norms and disease related decrements assumes that any life years gained due to a reduction in mortality will be lived in the diseased state until life expectancy, i.e. that all diseases are not just chronic but disease duration is lifelong. Inevitably this assumption means that the health effects of changes in mortality will be reduced. Consequently the cost per QALY threshold reported in Table C. 32 (column 2) will be higher than adjusting life years gained for population norms in Table C. 26 (column 2). A detailed breakdown of the cost per QALY threshold based on disease related disability and mortality effects is shown is Table C.33.

Table C.32: Summary of cost per QALY threshold based on disease related disutility

|  | Cost per life year threshold | Cost per QALY threshold <br> Disease related disutility <br> $[2]$ |
| :--- | :---: | :---: |
| big 4 PBC's <br> 11 PBCs (with motality) <br> All 23 PBCs (zero health effects for remaining | $£ 8,080$ | $£, 12,109$ |
| 12 PBCs ) |  |  |
| All 23 PBCS (non-zero health effects for <br> remaining 12PBCs, except GMS)* | $£ 15,628$ | $£ 23,395$ |

* in PBCs without a mortality signal, health effects were estimated by valuing changes in expenditure at the same rate as observed in PBEs for which there was a mortality signal except GMS.

Table C.33: Breakdown of the cost per QALY threshold based on disease related disutility


The number of life years gained associated with each death averted (columns 1 and 2 in Table C.34) is, again, consistent with previous estimates (Tables C. 28 and C.21). The average number of QALYs gained across all 11 PBCs is 2.8 QALY per death averted column 4 in Table C.34). As expected this value is lower than in the previous section (column 4 in Table C.28).

Table C.34: Implied YLL per death averted and implied QoL score per YLL gained, for each PBC

| PBC | PBC description | Implied YLL per excess death averted [1] | Implied YLL per PBC death averted [2] | Implied QALYs gained per excess death averted [3] | Implied QALYs gained per PBC death averted [4] |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 2 Cancer <br> 10 Circulatory <br> 11 Respiratory <br> 13 Gastro-intestinal <br> Big 4 |  | $\checkmark 14.07$ | 10.30 | 9.28 | 6.79 |
|  |  | 10.40 | 5.15 | 6.58 | 3.26 |
|  |  | 9.21 | 1.04 | 8.50 | 0.96 |
|  |  | 14.98 | 9.43 | 9.34 | 5.88 |
|  |  | 11.28 | 4.07 | 7.53 | 2.71 |
| 1 | Infectious diseases | 13.21 | 5.31 | 9.39 | 3.77 |
| 4 | Endocrine | 13.59 | 7.57 | 9.43 | 5.26 |
| 7 | Neurological | 13.59 | 6.12 | 7.68 | 3.46 |
| 17 | Genito-ufinary | 8.34 | 1.71 | 6.72 | 1.37 |
| 16 | Trauma \& injuries* | NA | NA | NA | NA |
| 18+19 Maternity \& neonates* |  | 74.34 | 74.34 | 60.45 | 60.45 |
|  | - First 11 PBC's | 11.37 | 4.14 | 7.60 | 2.77 |

C.2.2.3 Summary of the cost per QALY threshold based only on mortality effects

The analysis to this point is summarised in Table C.35. The three estimates of a cost per QALY threshold are based on assuming that each life year gained is either: lived in full health (see column 1, equal to the cost per life year estimates in Table C.22), lived in a quality of life that reflects age and gender norms of the general population (column 2); or lived in a quality of life that reflects the original disease state (column 3).

Table C.35: Summary of QALY threshold estimates based only on mortality effects

|  | $\begin{gathered} {[1]} \\ (Q o L \text { score }=1) \end{gathered}$ | [2] <br> (population norms) | $[3]$ (Disease related disutility) |  |
| :---: | :---: | :---: | :---: | :---: |
| Effect of expenditure on mortality: <br> YLL per death averted**: QALYsper death averted**: | $\begin{gathered} 1 \text { year } \\ \sim \\ \sim 4.1 \mathrm{YLL} \\ \sim 4.1 \text { QALYs } \end{gathered}$ | $\begin{gathered} \hline \text { Best estimate } \\ 1 \text { year } \\ \sim 4.1 \mathrm{YLL} \\ \sim 3.5 \text { QALYs } \end{gathered}$ | $\begin{gathered} 1 \text { year } \\ \sim \\ \sim 4.1 \mathrm{YLL} \\ \sim 2.8 \text { QALYs } \end{gathered}$ | $\begin{aligned} & {[1]} \\ & {[2]} \\ & {[3]} \end{aligned}$ |
| big 4 PBC's <br> 11 PBCs (with mortality) <br> All 23 PBCs* | $\begin{gathered} £ 8,080 \\ £ 15,628 \\ £, 17,663 \end{gathered}$ | $\begin{gathered} £, 631 \\ £ 18,622 \\ £ 21,047 \end{gathered}$ | $\begin{aligned} & £, 12,109 \\ & £ 23,395 \\ & £ 26,441 \end{aligned}$ |  |
| Effect of expenditure on mortality: <br> YLL per death averted**: QALYsper death averted**: | Remainder of disease <br> ~ 4.1 YLL <br> ~4.1 QALYs | Lower bound <br> Remainder of disease $\begin{gathered} \sim 4.1 \mathrm{YLL} \\ \sim 3.5 \mathrm{QALYs} \end{gathered}$ | Remainder of diseas $\begin{gathered} \sim 4.1 \text { YLE } \\ \sim 2.8 \text { QALYs } \end{gathered}$ |  |
| big 4 PBC's <br> 11 PBCs (with mortality) <br> All 23 PBCs* | $\begin{aligned} & £_{3,846} \\ & £ 6,106 \\ & £_{6} 6,901 \end{aligned}$ | $\begin{aligned} & £ 4,252 \\ & £ 6,852 \\ & £ 7,744 \end{aligned}$ | $\begin{array}{r} 6,5,319 \\ f, 568 \\ 6,683 \end{array}$ | [4] $[5]$ $[6]$ |
| Effect of expenditure on mortality: <br> YLL per death averted**: QALYsper death averted**: | $\begin{gathered} 1 \text { year } \\ 2 \text { YLL } \\ 2 \text { QALY } \end{gathered}$ | $\begin{gathered} \hline \text { Upper bound } \\ 1 \text { year } \\ 2 \mathrm{YLL} \\ \sim 1.92 .41 \end{gathered}$ | $\begin{gathered} 1 \text { year } \\ 2 \text { YLL } \\ \sim 1.5 \text { QALY } \end{gathered}$ |  |
| big 4 PBC's <br> 11 PBCs (with mortality) <br> All 23 PBCs* | £16,432 <br> £ 32,387 <br> £36,604 | $\begin{array}{r} £, 17,456 \\ 634,492 \\ 638,983 \\ \hline \end{array}$ | $\begin{array}{r} £ 21,747 \\ £ 42,967 \\ £, 48,561 \\ \hline \end{array}$ | $[7]$ $[8]$ $[9]$ |

* in PBCs without a mortality signal, health effects were estimated by valuing changes in expenditure at the same rate as observed in PBCs for which there was a mortality signal except GMS.
** see Tables C.20, C. 27 and C. 33
The weights reflecting the quality in which each of the years of life saved is lived implied in each of these three estimates is shown in Table C.36.

Table C.36: Implied QoL weight per YLL gained

| PBC | PBC description | Full health <br> [1] | Population <br> norms <br> [2] | Disease related disutility [3] |
| :---: | :---: | :---: | :---: | :---: |
| 2 | Cancer | 1 | 0.79 | 0.66 |
| 10 | Circulatory | 1 | 0.81 | 0.63 |
| 11 | Respiratory | 1 | 1.13 | 0.92 |
| 13 | Gastro-intestinal |  | 0.82 | 0.62 |
| Big 4 |  | 1 | 0.84 | 0.67 |
| 1 | Infectious diseases | - 1 | 0.89 | 0.71 |
| 4 | Endocrine | ) 1 | 0.84 | 0.69 |
| 7 | Neurological | 1 | 0.86 | 0.56 |
| 17 | Genito-urinary | 1 | 0.94 | 0.81 |
| 16 | Trauma \& injuries | NA | NA | NA |
| $18+19$ | Maternity \& neonates | 1 | 0.87 | 0.81 |
|  | First 11 PBC's | 1 | 0.84 | 0.67 |

Assuming that life years gained are lived in full health is not credible and should be regarded as an underestimate of the threshold given what is known about quality of life norms for the general population (see(Figure C.4). Equally, assuming that all life years gained are lived in the quality of life of the original disease state does not seem credible either and is likely to overestimate the threshold since it assumes that all disease is not only chronic but lifelong and all life years would be lived in the diseased state until death. The information that is available about disease duration suggests that many types of disease that comprise the PBCs are not chronic and certainty not lifelong (see Table C.23). Therefore, adjusting life years gained for the quality of life of the general population taking account of age and gender (in column 2, Table C35) is regarded as the best estimate of a cost per QALY threshold, which only reflects the health effects of changes in mortality. The lower and upper bounds are based on combining optimistic and pessimistic assumptions about the duration of health effects and how long a death might be averted as described in Section C.2.1.5.

However, it should be noted that these cost per QALY thresholds only account for the direct health effects of changes in mortality due to changes in expenditure. Insofar as much, or at least some part, of NHS activity and expenditure is intended to improve quality of life, not just mortality, then these estimates will underestimate total health effects and overestimate a cost per QALY threshold based on a more complete measure of possible health effects. In Section C.2.3 we explore the ways in which the likely effects of expenditure on quality of life (other than through mortality) might also be taken into account.

## C.2.3. Including quality of life effects during disease

The cost per QALY thresholds presented in Section C.2.2 only account for the health (QALY) effects of changes in mortality due to changes in expenditure. It does not seem credible to suppose that all NHS activity and expenditure only influences mortality with no effect on the quality of life while alive and experiencing a disease. Insofar as changes in NHS expenditure will also affect quality of life as well as mortality then total health effects will be underestimated and the thresholds presented in Table C. 35 will overestimate the cost per QALY threshold compared to a more complete picture of the likely effects of changes in NHS expenditure. In this section we explore ways to also take account of those effects on health not directly associated with mortality and life year affects (i.e., the 'pure' quality of life effects) to estimate an overall cost per QALY threshold.

The routine reporting of quality of life outcomes are increasingly available at PCT level (see Addendum 1 for a description of these data). In principle, the variation in such measures of outcome across PCTs could be used to estimate outcome elasticities for quality of life rather than mortality effects using similar econometric methods to those described in Appendix B (see Section B.8.8 for the results of an exploratory econometric analysis of these data). Howeyer, the currently limited coverage of routine reporting of these outcomes means that it is not feasible to estimate quality of life effects across all the PBCs using these data. Therefore, in this section we explore how estimates of effects of expenditure that can be observed (i.e., on mortality) can be used to infer the likely effects on what cannot be directly observed (quality of life), rather than making extreme assumptions that are not credible (e.g., assuming that changes in expenditure will have no effects on quality of life outcomes).

In Section C.2.3.1 we use three alternative estimates of the ratio of QALYs to life years lost due to different types of disease as a means of inferring the change in QALYs that is likely to be associated with the estimated change in YLL, i.e., essentially applying the estimated proportionate effect on life years to total QALYs. This is consistent with regarding the estimates of the mortality and life year effects as a surrogate for a more complete measure of the health effects of a change in expenditure.

However, the ratios of QALYs lost to life years lost due to disease in those PBC where outcome elasticities could not be estimated cannot inform estimates of the threshold (there are no estimated life year effects with which to apply the ratios). Nonetheless, the sources of information on which ratios are based also proyide much of the information required to calculate the QALY burden of disease in these areas, which can be used to inform estimates of the threshold. Therefore, in section C.2.3.2 we use an estimate of the QALY burden of disease, infer a proportionate effect on burden from the observed effects on life years, and then apply this proportionate effect to the measures of QALY burden for all the other PBCs. In this way we can use all the information available about the mortality and quality of life effects of the different types of disease that make up each PBC, including those where mortality based outcome elasticities are not available.

## C.2.3.1 Using ratios of QALYs to YLL

The ratio of the total QALYs to years of life lost (YLL) due to a disease indicates the number of QALYs associated with each YLL. Therefore, any change in YLL is expected to generate a number of QALYs indicated by the ratio - in this way, the estimated effects on mortality and life years are interpreted as a surrogate for a more complete measure of total health effects, which is reasonable. For example, a
disease with a ratio greater than 1 suggests that each YLL across the at risk population is associated with more than one QALY, i.e., there are significantly greater quality of life effects while experiencing the disease. Therefore, a change in expenditure that leads to 1 life year gained in this type of disease may generate a greater QALY effect than the same life year effects in a disease where this ratio is less than 1 , i.e., where most of the effect of disease is on mortality rather than quality of life. Therefore, using these ratios provides a means of accounting for the likely effect on quality of life other than through effects on mortality.

To understand the differences between the three ratios presented below it is useful to regard the total QALY lost to YLL ratio (R) for a particular disease as the sum of two ratios: i) the QALYs lost due to premature death to YLL ratio $\left(\mathrm{R}_{\text {death }}\right)$ and ii) the QALYs lost during disease (while alive) to YLL ratio ( $\mathrm{R}_{\text {alive }}$ ), as depicted in Equation 4.

$$
\begin{equation*}
R=\frac{Q A L Y \text { lost }}{Y L L}=\underbrace{\frac{Q A L Y \text { lost }_{\text {premature death }}}{Y L L}}_{R_{\text {death }}}+\underbrace{\frac{Q A L Y \text { lost } w \text { wile alive }}{Y L L}}_{R_{\text {alive }}} \tag{4}
\end{equation*}
$$

Insofar as YLL would not have been lived in full health, the quality of life effects eaptured in $\mathrm{R}_{\text {death }}$ are estimated to be lower than 1. Note that the analyses in Section C.2.2 already imply $2 \mathrm{R}_{\text {death }}$ ratio at PBC level. The second component of the ratio, $\mathrm{R}_{\text {alive, }}$, represents QALYs lost during disease for the at risk population as a proportion of the YLL observed in the same population - in diseases for which quality of life during disease is compromised but life expectancy is not changed significantly $\mathrm{R}_{\text {alive }}$ may thus assume high values. The ratios do not represent the balance of QALY gains due, to mortality and morbidity in a single patient, but rather in the population. Where $\mathrm{R}_{\text {death }}$ if lower than 1, only when the pure QALY effects offset the less than full quality of life of the YLL is the ratio greater than one. Therefore, ratios less than one are possible even when disease has measurable quality of life effects for those experiencing it.

## DALY to YLL ratios

The WHO GBD study provides UK specific estimates of the years of life lived with disability and the years of life lost due to different types of disease. Diseases in GBD are classified using U-codes that can then be mapped to ICD-10, as illustrated in Table C. 37 using a few examples (Addendum 1 provides more details on the mapping procedure). GBD uses Disability Adjusted Life Years (DALYs) as a measure of the burden of disease. This DALY measure has two components: i) the years of life lived with disability (YLD), which evaluates the number of years lived with disability over the durations of disease, and incorporates weights (between zero and one) to reflect the scale of disability experienced in each year; and ii) the years of life lost (YLL).

Table C.37: Illustration of the mapping between U-code and ICD

| U-code | ICDs |
| :--- | :--- |
| U037 (Other infectious diseases) | A02,A05,A20-A28,A31,A32,A38,A40-A49,A65-A70,A74-A79, A81,A82, |
|  | A83.1-A83.9, A84-A89,A92-A99, B00-B04,B06-B15,B25-B49,B58-B60, B64, |
| U016 (Tetanus) | B66-B72, B74.3-B74.9,B75,B82-B89,B92-B99, G04 |
| U061 (Mouth and oropharynx cancers) | A33-A35 |
| (U057 (Iron-deficiency anaemia) | D50-C14 |

The total DALY associated with a disease is simply YLL+YLD. Therefore, the DALY to YLL ratio is $(\mathrm{YLL}+\mathrm{YLD}) / \mathrm{YLL}$ or equivalently YLL/YLL + YLD/YLL. Since the first term (YLL/YLL $=\mathrm{R}_{\text {death }}$ ) must equal one and the second ( $\mathrm{R}_{\text {alive }}=\mathrm{YLD} / \mathrm{YLL}$ ) must be $\geq 0$, a ratio based on DALYs must necessarily be bounded by one.

$$
\begin{equation*}
R_{D A L Y}=\frac{D A L Y}{Y L L}=\frac{(Y L L+Y L D)}{Y L L}=1+\underbrace{\frac{Y L D}{Y L L}}_{R_{\text {death }}+R_{\text {alive }}} \tag{5}
\end{equation*}
$$

This is illustrated in Table C.38a for the four different diseases (U-codes) introduced in Table C. 36 which reflect diseases where mortality is the major component (e.g., U016) and where the impact of disease on the quality of life while alive is the major component (e.g., U141).

Table C.38a: Examples of DALY to YLL ratios

|  |  |  |
| :--- | :---: | :--- |
| Ucode | DALY ratios | $\left(\mathrm{R}_{\text {death }}+\mathrm{R}_{\text {alive }}\right)$ |
| U037 (Other infectious diseases) | 1.23 | $(1+0.23)$ |
| U016 (Tetanus) | 1.00 | $(1+0)^{*}$ |
| U061 (Mouth and oropharynx cancers) | 1.05 | $(1+0.05)$ |
| U141 (Spina bifida) | 2.34 | $(1+1.34)^{* *}$ |

* Given the short disease duration, it is only mortality effects that contribute to the ratio
** Quality of life effects during disease contribute significantly to estimates of the ratio
Note that the estimates of GBD YLL used here are derived using UK data on mortality (relating to the year 2004) by age and gender groups - we assume these data to be from ONS and thus consistent with the data used in this work. However, the calculation of YLLs in GBD differs from both the approach adopted by NHS IC and the approach adopted here of using net YLL. For each death obsevved in the data, GBD evaluates YLL by considering the life expectancy at the age at which the death occurred (and gender).[6] This is expected to overestimate net YLL (which accounts for counterfactual deaths, as detailed in Section C.2.2.3). This will make no difference to the first term in the QALY ratio ( $\mathrm{R}_{\text {death }}$ ) since an overestimate of YLL affects both denominator and numerator of the ratio. However, the second term $\left(\mathrm{R}_{\text {alive }}\right)$ is likely to be underestimated. Therefore the ratios will tend to underestimate the QALY effects of expenditure and overestimate the cost per QALY threshold. This will be adjusted for in Section C.2.3.2, where our preferred analysis based on burden of disease is presênted.


## Adjusting DALYs for quality of life norms

The use of DALY ratios bounded below by one essentially assumes that YLL would have otherwise been lived in a state of full health. As was discussed in section C.2.3.1 this is not credible given information available about the quality of life in the general population (see Figure C.4). It would lead to overestimating the QALYs associated with mortality and life year effects and underestimating the cost per QALY threshold. Therefore, it is important to adjust these DALY ratios for the quality of life norms by age and gender in the same way as described in Section C.2.3.1. Equation 6 shows how the adjusted ratio is formulated when YLLs are adjusted by the quality of life in the general population, $u_{n}$. This is a simplified representation of the adjustment as despite gender and age having been considered in calculations these are not shown in the notation below.

$$
\begin{equation*}
R_{D A L Y \text { adj }}=\frac{u_{n} Y L L}{Y L L}+\frac{Y L D}{Y L L}+=u_{n}+\underbrace{\frac{Y L D}{Y L L}} \tag{6}
\end{equation*}
$$

The effect of this adjustment (within each U-code, see Addendum 1) is illustrated in Table C.38b. Now those types of disease where mortality rather than quality of life with the disease is the major component can hare ratios less than one. Indeed the first term of these ratios $\left(\mathrm{R}_{\text {death }}\right)$ is consistent with (but not equivalent to) the analysis in Section C.2.3.1, where the ratio of quality adjusted net YLLs to unadjusted net YLLs represents this ratio on average for each PBC.

Table C.38b: Examples of adjusted DALY to YLL ratios

| Ucode | Adjusted DALY <br> ratios | $\left(\mathrm{R}_{\text {death }}+\mathrm{R}_{\text {alive }}\right)$ |
| :--- | :---: | :--- |
| U037 (Other infectious diseases) | 1.01 | $(0.78+0.23)$ |
| U016 (Tetanus) | 0.78 | $(0.78+0)$ |
| U061 (Mouth and oropharynx cancers) | 0.83 | $(0.78+0.05)$ |
| U141 (Spina bifida) | 2.18 | $(0.85+1.34)$ |

The disability weights used in the DALY measure (in $\mathrm{R}_{\text {alive }}$ ) are not based on the same description of health states as the EQ5D measure, nor are the weights based on a representative sample of the UK population responding to choice based elicitation questions. EQ5D based quality of life decrements (in relation to age adjusted quality of life norms) associated with different types of disease can be estimated from HODaR and MEPS data for the groups of ICD codes that make up each U-code. The calculations of the quality of life decrements from HODaR were conducted as previously described in Section C.2.2.2. In summary, the average quality of life scores across the ICDs which contribute to each U-code (see Table C. 36 and Addendum 1 for how ICD codes map to U-codes) and the average age and gender of respondents from HODaR and MEPS were used to calculate a disease decrement for each U-code, based on quality of life norms from the general population. Note that, by expressing the quality of life effects of different diseases as age related decrements (see Figure C.5), we do not require the HODaR and MEPS samples to necessarily be representative of the age distribution of the population at risk.

The disease related quality of life decrements can then be used to replace the DALY disability weights in $\mathrm{R}_{\text {alive }}$ reported in Tables C.38a and C.38b. This final adjustment is illustrated in Table C.38c: for example, the evidence about quality of life from HODaR and MEPS suggests that the impact of U037 on quality of life is greater than indicated by DALY disability weights. The quality of life effects of U141, although still very significant, are lower than indicated by DALY disability weights.

Table C.38c: Examples of QALY to YLL ratios (HODaR and MEPS)

| Ucode | QALY ratios <br> $(H o D A R ~ a n d ~ M E P s) ~$ | $\left(\mathrm{R}_{\text {death }}+\mathrm{R}_{\text {alive }}\right)$ |
| :--- | :---: | :---: |
| U037 (Other infectious diseases) | 1.37 | $(0.78+0.60)$ |
| U016 (Tetanus) | 0.78 | $(0.78+0$ |
| U061 (Mouth and oropharynx cancers) | 0.80 | $(0.78+0.02)$ |
| U141 (Spina bifida) | 1.88 | $(0.85+1.03)$ |

By turning what were originally DALY ratios into EQ5D QALY ratios, we regard the QALY to YLL ratios rather than DALY or modified DALY ratios as the preferred basis of estimating a cost per QALY threshold. We consider these estimates to provide a more complete picture of the likely health effects of changes in expenditure.


U-code QALY ratios to ICD QADI ratios
Information about the size and age and gender distribution is only available at U-code level. Therefore U-code ratios are applied to all the ICD codes that contribute to a particular U-code. Note that, unlike ICD codes, U-codes do not map directly to PBCs so some ICDs in different PBCs may belong to the same U-code and therefore have the same U-code ratio. Some ICDs are not included in the U-code classification of disease. Some of these are procedural codes (84 out of 1562)where mortality and QALY effects were not assigned mortality of QALY effects anyway (any health effects would be evident in other ICD codes) so it was not necessary to impute ratios for them. Of the others, some were associated with PBC16 ( 186 out of 1562 ) with a zero outcome elasticity so did not require imputation either. Imputation based on the median ratio across the ICDs within the PBC was required for the remaining (482 out of 1562). Eighty eight of these are not mapped into U-codes - these include three big categories of ICDs: Symptoms and signs (R00-R69), Abnormal clinical and laboratory findings, not elsewhere classified (R70-R99) and Ill-defined and unknown causes of mortality (R95-R99). The remaining 394 were associated with U-codes where the ratio was undefined because the denominator (YLL) was zero. In both these cases, values were also imputed based on the median ratio across the ICDs within the PBC. Since the distribution of ratios within a PBC tend to be highly positively skewed, imputation based on the median is likely to be conservative with respect to health effects and especially in the latter case where mortality effects appear to be a much less important aspect of the disease.
Table C. 39 illustrates the variation observed in the ratios (imputed) across ICDs within the same PBC.

Table C.39: Percentiles of the ratio across ICDs, by PBC


## Allocating effects at PBC level to ICD codes

Tables C.38a,b and cillustrate how QALY ratios can be calculated for and differ by U-code and therefore the ICD codes that make them up. Unsurprisingly, these ratios differ across the type of diseases that make up each PBC (Table C.39). Therefore, when using this information to estimate a cost per QALY threshold the mortality and life year effects observed at PBC level must be allocated in some way to the component ICD codes before ratios are applied to LY effects and the resulting QALY effects are summed across all the contributing ICD codes.

Alternatively, one could calculate an average of the ratios within a PBC and then apply this 'average ratio' to life year effects at PBC level, rather than calculate QALY effects at ICD level by applying the relevant ratio. This would be inappropriate for two reasons. Firstly, ratios should not be averaged; instead, the total QALYs lost and KLL should be summed across ICDs and the ratio of these sums used to represent a PBC level estimate (i.e., a ratio of averages). Secondly, and even if the appropriate estimate of the QALY to YLL ratio is calculated at the PBC level, this estimate would assume ICDs to be equally representative of the PBC - i.e., that expenditure would be equally likely to affect any of the ICDs that compose aparticular PBC. This is unlikely to be true not only due to the inherent differences in the disease described by the ICD coding, but also as ICDs are likely to differ significantly in what concerns the size of the at risk population they represent.

It is important to consider explicitly how other information might inform the different ways in which the effects observed at PBC level might be generated by the distribution of impacts at ICD level, i.e., where investment or disinvestment is likely to occur within the PBC and therefore which ICDs are likely to contribute most to overall health effects. An important and complementary element to the econometric analysis of routinely reported information at PBC level, was to investigate this by looking at local level information available within the NHS. The details of this investigation are reported in Addendum 2, and show, rather disappointingly, that the information available is not useful for the purposes of this analysis. In the absence of information, it is possible to assume that a change in PBC expenditure will be allocated equally (on a per patient basis) across the component ICD codes, i.e., any investment or disinvestment is equally likely (occurs at random) across the population at risk within the PBC.

However, there is another source of information that is able to give some indication of which areas are more likely to have been subject to investment or disinvestment across PCTs: the Hospital Episode Statistics (HES) dataset. This data source provides information about the costs associated with each ICD by PCT. The variation in per patient costs between PCTs (where total costs allocated to individual ICDs were divided by the number of patients using services in the PCT) was analysed to establish which ICDs contribute most to the variability in PBC costs across PCTs. The ICDs that contribute most to this variance may be expected to be more likely to have been subject to differential investment or disinvestment across PCTs. The costs from HES data are, however, only a component of total PBC costs but they are an important one. Unfortunately total PBC costs are not available at ICD level across PCTs so could not be used for this purpose. However, the assumption is not that HES cost are representative of total PBC costs but that those ICDs that contribute most to variability in HES costs are also likely to contribute most to variability in total PBC costs as well.

There are very marked differences in relative weight assigned to ICDs based solely on the size of the population or its contribution to variance in costs (see Addendum 1). One would expect investment or disinvestment within a PBC to focus on areas of marginal value rather than be allocated at random, therefore, the health effects of a change in PBC expenditure are likely to be overestimated and a cost per QALY threshold underestimated when allocating effects equally across the population at risk within each PBC. This is confirmed by the results of this analysis reported in footnote ${ }^{11}$. For these reasons our preferred analysis uses contribution to variance to 'weight' the different ICD codes within a PBC (allocate the life year effects), before applying the QALY ratios associated with each ICD. This is also conservative, with respect to the health effects of changes in expenditure, compared to alternative assumptions that could be made about how PBC level effects might be allocated to ICD codes. The implications for a cost per QALY threshold that uses the estimated mortality and life year effects as a surrogate for a more complete measure of the likely heatheffects (i.e., that includes quality of life as well as quality adjusted life year effects) is summarised in Table C. 40 and detailed in Table C.41.
${ }^{11}$ The table below reports the cost per QALY threshold using a relative weight based on the size of the ICD population to allocate health effects.

|  | Cost per QALY threshold |  |  |
| :---: | :---: | :---: | :---: |
|  | DALY ratios <br> [1] | Adjusted DALY ratios [2] | QALY ratios <br> (HoDAR and MEPs) <br> [3] |
| big 4 PBC's | £4,400 | £,5,100 | £2,340 |
| 11 PBCs (with mortality) | £,8,066 | $¢_{9,267}$ | £ 4,212 |
| All 23 PBCs | $\oint 9,117$ | £ 10,474 | £.4,760 |

Table C.40: Summary of the QALY threshold using ratios

|  | Cost per QALY threshold |  |  |
| :---: | :---: | :---: | :---: |
|  | $\begin{gathered} \text { DALY ratios } \\ {[1]} \\ \hline \end{gathered}$ | Adjusted DALY ratios <br> [2] | QALY ratios (HoDAR and MEPs) [3] |
| big 4 PBC's | £5,402 | £6,419 | £,5,990 |
| 11 PBCs (with mortality) | £, 9,958 | £11,718 | £10,297 |
| All 23 PBCs | ¢,11,254 | £,13,244 | £11,638* |

* Preferred analysis

Table C.41: Breakdown of the QALY threshold using ratios by PBC


Since all the analysis in this Section seeks to use the estimated mortality and life year effects as a surrogate for a more complete measure of likely health effects, it is the cost per QALY threshold for all 23 PBCs that is most relevant. As expected, this threshold $(£, 11,638)$, is lower than a cost per QALY threshold based only the quality adjusted life year effects ( $£ 21,047$ and $£ 26,441$ in Table C. 34 that assumes no effects of NHS expenditure on quality of life itself). This difference gives some indication of the relative importance of QALY effects due to avoidance of premature death and the QALY effects of avoiding disability during disease. Table C. 42 reports how the estimated QALY effects for each PBC can be decomposed into that part associated with quality adjusted life year effects and that part associated with 'pure' quality of life effects. These results appear credible for the first 11PBCs, where those for which mortality is the major concern have a much greater share of total QALY effects associated with avoidance of premature death (e.g., PBC2 and PBC10) compared to those where quality of life is the major concern (e.e., PBC 7).

Table C.42: Decomposing estimated QALY effects by PBC
$\left.\begin{array}{|c|cc|cc|}\hline & & \begin{array}{c}\text { QALY } \\ \text { change } \\ \text { (total) }\end{array} & \begin{array}{c}\text { QALY } \\ \text { change } \\ \text { (death) }\end{array} & \begin{array}{c}\text { QALY gained } \\ \text { due to avoidance } \\ \text { of premature } \\ \text { due to avoidance } \\ \text { death }\end{array} \\ \text { PBC disability while } \\ \text { alive }\end{array}\right]$

Recall that the ratios of QALYs to YLL due to disease in those PBC where outcome elasticities could not be estimated cannot be used to inform estimates of the threshold because there are no estimated life year effects with which to apply the ratios. Therefore, as in previous sections, the estimated effect of expenditure on health for the 11 PBCs with outcome elasticities is applied to the estimated changes in PBC expenditure for the other 12 PBCs (excluding GMS for the reasons given in Section C.2.1.5), i.e., assuming that the health effects that can be obseryed of a change in expenditure will be similar to those that cannot. However, the use of QALYratios also implies that the share of total health effects between quality adjusted life year effects and that part associated with 'pure' quality of life effects are also similar to those PBC with estimated outcome elasticities. Summing the different types of health effects across these 11 PBCs suggests that $50 \%$ is due to avoidance of premature death and $50 \%$ due to avoidance of disability. This is clearly not eredible when applied to the other PBCs, e.g., mental health, vision and hearing are likely have a much greater share of total health effects associated with quality of life effects and very little associated with premature mortality.

By comparing the change in QALY in each PBC (that originates cost per QALY threshold estimates, column 2 in Table C.43), with the corresponding change in YLL (column 6, Table C.16), we can infer the implied QALX to YLL ratio in each of the PBCs with a mortality signal. These are shown in Table C.40. The QALY to YLL ratio implied by the analysis using QALY ratios for all 11 PBC with outcome elasticities is 1.52 , which suggests that every life year is associated with 1.52 QALYs on average across these PBCs. However, this implied QALY ratio differs across these PBCs, ranging from 0.79 in PBC2 to $\sqrt{5} .05 \mathrm{in} \mathrm{PBC18+19}$ (see column 4 of Table C.43). It should be noted that the implied QALY ratio of 1.35 for the 11 PBC with outcome elasticities is a ratio of QALYs to unadjusted YLL. The proportion of total QALY effects due to premature deaths for the same PBCs ( $50 \%$ in Table C.41) also implies a ratio equal to two. However, this is a ratio of total QALY effects to quality adjusted YLL. The difference between these two ratios is the denominator, i.e., quality adjusted YLL are lower than unadjusted YLL.

Table C.43: Implied QALY to YLL ratios.

| PBC | PBC description | Adjusted DALY ratios |  |  | QALY ratios(HoDAR and MEPs) |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Implied QALY per LY gained <br> [1] | Implied QALY per excess death averted [2] | Implied QALY per PBC death averted [3] | Implied QALY per LY gained <br> [4] | Implied QALY per excess death averted [5] | Implied QALY per PBC death averted [6] |
| 2 | Cancer | 0.82 | 11.58 | 8.48 | 0.79 | 11.16 | 8.17 |
| 10 | Circulatory problems | 1.20 | 12.51 | 6.20 | 1.05 | 10.94 | 5.42 |
| 11 | Respiratory problems | 1.96 | 18.09 | 2.04 | 2.65 | 24.45 | 2.76 |
| 13 | Gastro-intestinal problems | 1.54 | 23.02 | 14.49 | 2.31 | 34.63 | 21.80 |
|  | Big 4 | 1.26 | 14.20 | 5.12 | 1.35 | 15.21 | 5.49 |
| 1 | Infectious diseases | 1.56 | 20.64 | 8.30 | 1.98 | 26.22 | 10.54 |
| 4 | Endocrine problems | 2.62 | 35.61 | 19.84 | 4.95 | 67.31 | 37.51 |
| 7 | Neurological problems | 4.56 | 61.99 | 27.90 | 5.27 | 71.69 | 32.26 |
| 17 | Genito-urinary problems | 1.75 | 14.56 | 2.98 | 1.44 | 11.98 | 2.45 |
| 16 | Trauma \& injuries* | NA | NA | NA | NA | NA | NA |
| 18+19 | Maternity \& neonates* | 6.88 | 511.33 | 511.33 | 15.05 | 1118.85 | 1118.85 |
|  | First 11 PBC's | 1.33 | 15.16 | 5.53 | 1.52 | $\bigcirc 17.26$ | 6.29 |

The problem is that using QALY to YLL ratios means that much of the information that is available about the other 12 PBCs cannot be used to inform the estimates of the cost per QALY threshold. Fortunately, the sources of information on which ratios are based also provide much of the information required to calculate the QALY burden of disease in these areas. Section C.2.3.2 explores how measures of burden can be used to estimate a cost per QALY threshold that captures the likely effects of a change in expenditure on all aspects of health while using all the information that is available about all the PBCs.

## C.2.3.2 Using estimates of the QALY burden of disease

In this Section we use estimates of the QALY burden of disease, infer a proportionate effect on burden from the observed effects on life years, and then apply this proportionate effect to the measures of QALY burden for all PBCs. In this way we can use all the information available about the mortality and quality of life effects of the different types of disease that make up each PBC, particularly for those where mortality based outcome elasticities are not available.

The total QALY burden of disease for the population with disease in a particular year includes: i) the quality adjusted years of life lost due to all the disease related mortality that could occur in this population over their remaining duration of disease and ii) the reduction in quality of life while alive also for their remaining disease duration. These components of burden represent, respectively, the QALY lost due to premature death (QAI Pl $_{\text {death }}$ ) and the QALY lost while alive $\left(\mathrm{QALYl} \mathrm{l}_{\text {alive }}\right)$ as a consequence of disease.

$$
\begin{equation*}
\text { Burden }=Q A L Y l_{\text {death }}+Q A L Y l_{\text {alive }} \tag{7}
\end{equation*}
$$

However, applying the estimated proportionate effects on mortality and life years to such a measure of total burdenwould provide an estimate of the effects of a change in expenditure, not just in one year, but in all the remaining years of disease for the population at risk in that year. Recall from Section C.2.1 that we have adopted the conservative assumption that changes in expenditure will only have health effects in one year for the population with disease in that year. Therefore, it is not a measure of total burden that is required, but a measure of the QALY burden of disease during one year for the population with disease (prevalent and incident) in that year. The estimated outcome elasticities can then be appropriately (and directly) applied to this measure of burden. Of course, it would be possible to solve for a lower outcome elasticity that could be applied to total burden which would return the required estimate of total QALY effects restricted to one year.

The information from GBD used to derive QALY ratios in Section C.2.3.2 includes information about the YLL and duration of disease for those incident to a U-code, i.e., the measure of QALY burden from the information included in the ratios is a measure of the total burden of the disease but only for the
population that is incident (rather than total population with disease) in one year. Assuming that incidence is stable over the disease duration this is also equivalent to the QALY burden of disease during one year for the population with disease (i.e., those that are incident and prevalent) in that year. This is valid as long as estimates of the quality of life decrement of disease from HODaR and MEPS are assumed representative of average effects across those earlier (incident) and later (prevalent) in their disease duration.

However, in moving from ratios to absolute measures of burden it becomes more important to examine and then adjust for any inconsistency between information about YLL and size of the incident population from GBD (which is available by U-codes and can be mapped to ICDs), and the information about net YLL and observed deaths for each PBC based on ONS data as described in Section C.2.2.3 - see Table C.44.

Table C.44: Comparing deaths and YLL from ONS and GBD.

|  |  | Excess deaths ONS [1] |  deaths <br> All All deaths <br> deaths GBD* $^{*}$ <br> ONS $[3]$ <br> $[2]$ 1,408 |  | adjustment factor (deaths) [4] | (12) |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Net estimates ONS <br> 151 |  |  | Total YLL GBD* <br> [6] | adjustment factor (YLL) [7] |
| 1 | Infectious diseases |  | 2,797 | 6,958 |  | 1,408 | 4.94 | 36,962 | 25,142 | 1.47 |
| 2 | Cancer | 95,715 | 130,810 | 140,124 | 0.93 | 1,347,184 | 1,932,637 | 0.70 |
| 4 | Endocrine | 3,769 | 6,765 | 7,509 | 0.90 | 51,225 | 95,401 | 0.54 |
| 7 | Neurological | 6,909 | 15,353 | 12,854 | 1.19 | 93,917 | 164,796 | 0.57 |
| 10 | Circulatory | 79,218 | 159,852 | 178,454 | 0.90 | 823,768 | 1,750,608 | 0.47 |
| 11 | Respiratory | 7,386 | 65,446 | 67,441 | 0.97 | 68,030 | 594,529 | 0.11 |
| 13 | Gastro-intestinal | 15,199 | 24,147 | 28,329 | 0.85 | 227,703 | 396,829 | 0.57 |
| 17 | Genito-urinary | 2,172 | 10,625 | 8,606 | 1.23 | 18,127 | 77,338 | 0.23 |
| $18+19$ | Maternity \& neonates | 226 | 226 | 2,211 | 0.10 | 16,801 | 149,868 | 0.11 |
|  | Total | 213,391 | 420,182 | 446,936 | 0.94 | 2,683,717 | 5,187,148 | 0.52 |

There are a number of reasons for the potential inconsistencies: i) GBD is based on earlier years of mortality data; ii) the imprecision of mapping from U-codes to PBC via ICD codes; and iii) the YLL reported in GBD are based on life expectancy at the age of death (see Section C.2.2 and C.2.3) and will overestimate the net YLL. Therefore, the YLL by U-code, reported in GBD, that are mapped to ICDs are adjusted by these proportionate differences (column 7 of Table C.44) to ensure that the YLLs associated with all contributing ICD codes are consistent with (do not overestimate) the net YLL for the PBC as a whole. The variation across ICDs in the adjusted QALY burden associated with mortality gains (for the population with disease in a particular year) is depicted in column 2 of Table C.45.

Table C.45: Variation across ICDs of the QALY burden of disease during one year for a patient with disease in a particular year


It is QALY burden during one year per patient with disease in that particular year that is reported in this Table, including the median and range across the ICD codes contributing to each PBC. However, these values should not be over interpreted as the 'average' QADY burden for the PBC, as this depends on how PBC effects are allocated to ICDs (i.e., those which have the higher contribution to variance in PBC costs) and the 'average' burden for groups of PBCs depends on how a change in overall expenditure is shared between them, i.e., the expenditure elasticities estimated for each PBC in Appendix B.

Due to the earlier years of data and imprecision in mapping from U-codes to ICDs there might also be some inconsistency in estimates of the total incidence of disease for a PBC. Insofar as disease related mortality risk is stable, the same number of deaths should be observed in GBD and ONS data for the same at risk population. The PBC deaths recorded in GBD and those observed in ONS data (columns 2 and 3 in Table C.44) are similar but nonetheless the proportionate difference is used to adjust the scale of quality of life burden while alive based on GBD information (equivalent to adjusting estimates of incidence). Notable exceptions are PBC1 and PBC18+19 where the discrepancies are likely to be due to imperfect mapping from 0 -code to PBC via ICD codes. Summaries of the ICD specific values of the adjusted burden of disease while alive are depicted in column 1 of Table C.45. Total burden (for the population with disease in a particular year) is the sum of the two components of burden (Table C. 46 presents a few examples for illustration).

Table C.46: Examples of QALY burden of disease for the population with disease in a particular year

| Ocode | QALY burden | (QALY lost ${ }_{\text {death }}+$ QALY lost $_{\text {alive }}$ ) |
| :--- | :---: | :--- |
| U037 (Other infectious diseases)* | 0.20 | $(0.09+0.11)$ |
| U016 (Tetanus) | 0.78 | $(2.73+0.00)$ |
| U061 (Mouth and oropharynx cancers) | 2.97 | $(2.87+0.10)$ |
| U141 (Spina bifida) | 0.65 | $(0.18+0.46)$ |

*Note that differential adjustments have been made to YLL (affecting QALY lost ${ }_{\text {death }}$ ) and to the incidence (affecting QALY lostalive), thus implied ratios from these burden estimates may differ from ratios presented is Section C.2.3.1.

The implications for the cost per QALY threshold of using information about the QALY burden of disease for all PBCs, rather than QALY ratios for those where an outcome elasticity can be estimated, are reported summarily in Table C. 47 and in detail in Table C.48.

Table C.47: Summary of the cost per QALY threshold

|  | Cost per QALY gained |  |
| :---: | :---: | :---: |
|  | QALY ratios (HoDAR and MEPs) [1] | QALY burden (HoDAR and MEPs) [2] |
| big 4 PBC's | £5,990 | £3,036 |
| 11 PBCs (with mortality) | £10,297 | £,5,128 |
| All 23 PBCs | £,11,638* | £,15,701* |

* Preferred analysis

The cost per QALY threshold for the 11PBCs with outcome elasticities is a little lower using a measure of QALY burden $(£, 5128)$ rather than the QALY ratios $(£ 10,297)$ described in Section C.3.2.1. This is because the way GBD calculates YLL overestimates net YLL (which accounts for counterfactual deaths, as detailed in Section C.2.2.3). This will make no difference to the first term in the QALY ratio ( $\mathrm{R}_{\text {death }}$ ) used in Section C.2.3.1 since an overestimate of YLL affects both denominator and numerator of the ratio. However, the second term $\left(\mathrm{R}_{\text {alive }}\right)$ is likely to be underestimated. Therefore the ratios will tend to underestimate the QALY effects of expenditure and overestimate the cost per QALY threshold (see Table C.47). We are able to adjust the GBD based measure of QALY burden for this overestimation in calculating the QALY threshold reported in column 2.

Since the purpose of this Section is to use the estimated mortality and life year effects as a surrogate for a more complete measure of likely health effects, it is the cost per QALY threshold for all 23 PBCs that is of most relevance. The cost per QALY threshold for all 23 PBCs is based on applying the proportionate effects on the QALY burden of disease, based on the observed effects of changes in expenditure on mortality in the 11 PBC with outcome elasticities, ${ }^{12}$ to the QALY burden of disease in the other PBCs. This generates a much higher cost per QALY threshold ( $£ 15701$ ) than the one based on applying the estimated QALY effects of changes in expenditure, using QALY ratios for the 11 PBC with outcome elasticities, to changes in expenditure in the others $(£ 11,638)$. The reason is that the QALY burden of disease in the other PBCs is, in general, lower than the QALY burden of disease across those PBCs where outcome elasticities can be estimated (see Table C. 45 above).

Therefore, applying the same proportionate effects to a lower QALY burden generates a smaller health effect of a change in expenditure. ${ }^{13}$ In essence the difference between these estimates is that in column 1 of Table C. 47 the absolute effect on health associated with an absolute change in expenditure is extrapolated to the other PBCs, where as in column 2 it is the relative effect on health of an absolute change in expenditure that is extrapolated. Since we know that QALY burden differs between (and within) PBCs and especially between the groups of PBCs with and without estimated outcome elasticities (see Table C.45), it is the values based on QALY burden in column 2 of Table C. 47 that are regarded as most credible and represent out central or best estimate.

A detailed breakdown of changes in expenditure and changes in QALYs across all PBCs is shown in Table C. 48 when the analysis is based on QALY ratios and on QALY burden of disease. A comparison of these values confirms that QALY effects for the other PBC are lower and therefore the cost per QALY for each of these PBCs are in general much higher when based on a proportionate effect on QALY burden. Of course, we have not directly observed quality of life effects in these PBC but inferred them from the proportionate effects that we can observe. Insofar as investment and disinvestment opportunities in these PBCs might have been more valuable (offered greater improvement in quality of life) than suggested by the implied PBC thresholds, then overall QALY effects will tend to be underestimated and the cost per QALY threshold overestimated.

[^79]Table C.48: Breakdown of the cost per QALY threshold


For the reasons discussed in previous sections, we regard all the cost per QALY threshold reported in column 2 of Table C. 47 to be on balance conservative with respect to overall health effects of a change in expenditure. However, the estimate of $£ 15701$ may be especially conservative with respect to health effects (i.e., overestimated) based, as it is on an extrapolation of the proportionate effects to measures of burden on these PBC, rather than observations of the direct impact of changes in expenditure on quality of life in these types of disease. This is especially so in PBC 5, Mental Health Disorders, which accounts for a large proportion of the change in overall expenditure ( $30 \%$ ) and where a review of the evidence suggests that the investment and disinvestment opportunities in this PBC are likely to have been more valuable than the implied PBCcost per QALY of $£ 60,111$ (see Addendum 3 to this Appendix). The lower cost per QALY threshold for the 11PBCs with outcome elasticities ( $£, 5128$ ) might be regarded as more secure in this respect but they only account for a proportion ( $28 \%$ ) of any change in overall expenditure (see Table C.53).

Table C. 49 reports how the estimated QALY effects based on measures of QALY burden for each PBC can be decomposed into that part associated with life year effects adjusted for quality and that part associated with 'pure' quality of life effects. These results are very similar to those reported in Table C. 40 which were based on QALY ratios for the 11 PBCs with an estimated outcome elasticity. Those PBCs for which mortality is the major concern have a much greater share of total QALY effects associated with avoidance of premature death (e.g., PBC 2 and PBC 10 ) compared to those where quality of life is the major concern (e.g., PBC 7). The differences tend to favour QALYs gained though avoidance of disability, which reflects the underestimation of the effects on 'pure' quality of life when using QALY ratios based on estimates of YLL from GBD (see the discussion above). The exceptions are PBC 1 and PBC $18 \& 19$. The reason is that there are significant adjustments made based on differences in observed and recorded mortality (to adjust for differences in recording) as well as differences in YLL due to the GBD method of calculation (see Table C.42).

Table C.49: Decomposing estimated QALY effects by PBC


The implied QALY per life year gained and death averted are reported in Table C.50. As expected, the implied QALY per PBC death averted across all 11PBCs with outcome elasticities is higher (12.6 QALY) than reported in Section C.2.3.1 (6.3 QALY) because of the previous bias against quality of life effects.

Table C.50: Implied QALY per excess death ayerted: using burden, contribution to variance

| PBC | PBC description |  | Implied QALY per excess death averted <br> [2] | Implied QALY per PBC death averted [3] |
| :---: | :---: | :---: | :---: | :---: |
| 2 | Cancer | 0.70 | 9.86 | 7.21 |
| 10 | Circulatory problems | 0.93 | 9.63 | 4.77 |
| 11 | Respiratory problems | 16.40 | 151.10 | 17.05 |
| 13 | Gastro-intestinal problems | 1.78 | 26.66 | 16.78 |
|  | Big 4 | 2.66 | 30.02 | 10.82 |
| 1 | Infectious diseases ${ }^{\text {a }}$ | 3.83 | 50.62 | 20.35 |
| 4 | Endocrine problems | 11.89 | 161.59 | 90.04 |
| 7 | Neurological problems | 14.86 | 202.05 | 90.92 |
| 17 | Genito-urinary problems | 2.85 | 23.80 | 4.87 |
| 16 | Trauma \& injuries | NA | NA | NA |
| 18+19 | Maternity \& neonates | 0.54 | 40.33 | 40.33 |
|  | First 11 PBC's | 3.05 | 34.65 | 12.63 |

Recall that in Section C.2.3.1, the ratios of QALYs to YLL due to disease in those PBCs where outcome elasticities could not be estimated could not be used to inform estimates of the threshold or indicate how any total health effects in these other PBCs are likely to be 'shared' between life year effects adjusted for quality and that part associated with 'pure' quality of life effects (see Table C.42). By applying the observed proportionate effects of changes in expenditure to measures of QALY burden of disease in these other PBCs the likely share of any effects on QALYs between avoidance of premature mortality and avoidance of disability more closely reflect the nature of these types of diseases (see Table C.49). As expected, a much greater proportion of QALY effects are associated with quality of life during the disease compared to the 11PBCs where mortality based outcome elasticities could be estimated. The share of effects in particular PBCs are also much more credible. For example, in PBC5 Mental Health Disorders the overwhelming share of QALY effects are associated with quality of life itself and for others, such as

PBC12 Dental problems, PBC9 Problems of Hearing and PBC8 Problems of Vision; almost all effects are associated with quality of life rather than mortality and life years. For this, and the other reasons discussed above, the analysis based on measures of QALY burden are regarded as the best estimate of a cost per QALY ratio that reflects a more complete picture of the likely health effects of changes in overall expenditure.

## C.2.3.3 Summary of the cost per QALY threshold

The results of the three sequential steps of analysis described in this Chapter are summarised in Table C.51. In Section C.2.1 we explored ways in which the estimated effects on mortality from the econometrics work in Appendix B might be better translated in to life year effects by overcoming some of the limitations of mortality data available at PCT level and taking account of counterfactual deaths. The results of this analysis were reported in Table C. 21 and are repeated in column 1 of Table C. 51. These results can be interpreted as cost per QALY thresholds conditional on the assumption that all life years are lived in full health and the quality of life with disease is zero (equivalent to death).

Table C.51: Summary of cost per QALY threshold estimates

|  | $\begin{gathered} {[1]} \\ \text { (Table C.20) } \\ \hline \end{gathered}$ | $\begin{gathered} {[2]} \\ \text { (Table C.34) } \\ \hline \end{gathered}$ | $(1 /[3]$ |  |
| :---: | :---: | :---: | :---: | :---: |
| QoL associated with life extension: QoL during disease: | $\begin{aligned} & 1 \\ & 0 \end{aligned}$ | Norm <br> 0 | $\begin{aligned} & \text { nased on burden } \end{aligned}$ |  |
| Effect of expenditure on mortality: <br> YLL per death averted: QALYs per death averted: | $\begin{gathered} 1 \text { year } \\ \sim \\ \sim 4.1 \mathrm{YLL} \\ \sim \end{gathered}$ | $\begin{gathered} 1 \text { year } \\ \sqrt{4.1 \mathrm{YLL}} \\ 3.52 A L Y^{1} \end{gathered}$ | $\begin{gathered} \text { Best estimate } \\ 1 \text { year } \\ \sim 4.1 \mathrm{YLL} \\ \sim 12.6 \text { QALY } \end{gathered}$ |  |
| big 4 PBC's <br> 11 PBCs (with mortality) <br> All 23 PBCs | $\begin{aligned} & \AA 8,080 \\ & £ 15,628 \\ & £ 17,663 \end{aligned}$ | $\begin{array}{r} £, 9,631 \\ £, 18,622 \\ \\ £ 21,047 \end{array}$ |  | [1] [2] [3] |
| Effect of expenditure on mortality: <br> YLL per death averted: QALYs per death averted: | Remainder of disease duration 4.1 YLL 4.1 QALY | Remainder of disease duration $\begin{gathered} \sim 4.1 \mathrm{YLL} \\ \sim 3.5 \text { QALY } \end{gathered}$ | Lower bound <br> Remainder of disease duration ~ 4.1 YLL <br> ~ 12.6 QALY |  |
| big 4 PBC's <br> 11 PBCs (with mortality) <br> All 23 PBCs | $\begin{aligned} & £ 3,846 \\ & £ 6,106 \\ & £ 6,901 \end{aligned}$ | $\begin{aligned} & £ 4,252 \\ & £ 6,852 \\ & £ 7,744 \end{aligned}$ | $\begin{gathered} £ 674 \\ £ 860 \\ £ 2,785 \end{gathered}$ | $[4]$ $[5]$ $[6]$ |
| Effect of expenditure on mortality: <br> RUDer death averted: QALY per death averted: | $\begin{gathered} 1 \text { year } \\ 2 \text { YLL } \\ \sim \\ 2 \text { QALY } \end{gathered}$ | $\begin{gathered} 1 \text { year } \\ 2 \text { YLL } \\ \sim 1.9 \text { QALY } \end{gathered}$ | $\begin{gathered} \hline \text { Upper bound } \\ 1 \text { year } \\ 2 \mathrm{YLL} \\ \sim 6.1 \text { QALY } \end{gathered}$ |  |
| big 4 PBC's <br> 11 PBCs (with mortality) <br> All 23 PBCs | $\begin{aligned} & £ 16,432 \\ & £ 32,387 \\ & £, 36,604 \\ & \hline \end{aligned}$ | $\begin{aligned} & £, 17,456 \\ & £ 34,492 \\ & £, 38,983 \\ & \hline \end{aligned}$ | $\begin{array}{r} £ 6,292 \\ £ 10,626 \\ £, 32,537 \\ \hline \end{array}$ | [7] <br> $[8]$ <br> $[9]$ |

In Sectionc.2.2 we considered how the estimated life year effects might be adjusted for the quality of life in which they are likely to be lived, taking account of the gender and the age at which life years are gained or lost (see Table C.34). The results of this analysis are repeated in column 2 below. Finally, in the current Section, C.2.3, we explored ways to also take account of the likely effects of changes in expenditure on quality of life during disease as well as the effects associated with mortality and life years (see column 3). These estimates provide our central estimate of a cost per QALY threshold, because they make best use of available information while the assumptions required, which on balance are likely conservative with respect to health effects, appear more reasonable than the other alternatives available. ${ }^{14}$

[^80]The estimate of $£ 5,128$ per QALY (line 2 ) is restricted to the effects of changes in expenditure in the 11 PBCs where outcome elasticities can be estimated. Although this might be regarded as more secure these PBCs only account for a proportion of the change in overall expenditure (approximately $28 \%$, see Section C.2.4). The threshold of $£ 15,701$ uses the estimated proportionate effects of expenditure on the QALY burden of disease in these PBC as a surrogate for proportionate effects in the others, i.e., assuming that the effects that can be observed will be similar to those that cannot. As discussed in Section C.2.3.2, there are reasons to suspect that this may underestimate health effects in these PBCs which have most influence on the overall threshold. As in previous sections, no health effects are assigned to PBC23 (General Medical Services) on the basis that any health effects of this expenditure would be recorded in the other PBCs. Therefore, the best or central estimate of cost per QALY threshold is $£ 15,701$ (column 3 , line 3 ).

This estimate reflects changes in undiscounted QALYs associated with changes in expenditure. Although all the health effects of a change in expenditure are restricted to one year (so no discounting is necessary) some of the quality adjusted life year effects of a change in mortality in that year will occur in future years, so in principle should be discounted. However, discounting these life year effects, even at the higher rate of $3.5 \%$ recommended by NICE, only increases the cost per QALY threshold to $£ 15,940$ (Table C.52).

Table C.52: Summary of QALY threshold, discounted.

|  | ${ }^{[1]}$ undiscounted |  |  |
| :---: | :---: | :---: | :---: |
|  | Best estimate |  |  |
| big 4 PBC's | £3,036 | £3,097 | [1] |
| 11 PBCs (with mortality) | £,5,128 | £5,218 | [2] |
| All 23 PBCs ${ }^{1}$ | £,15,701 | £,15,940 | [3] |

${ }^{1}$ in PBCs without a mortality signal, health effects were estimated by valuing changes in expenditure at the same rate as observed in PBCs for which there was a mortality signal.
${ }^{2}$ Only quality adjusted net YLL were discounted, and thus QALYs associated with gains in QoL during disease were not. The discounting factor has been calculated by applying a $3.5 \%$ discount rate to each year of life lost in the PBCs - the estimate of years of life lost used was the implied YLL per death averted in each PBC (in Table C. 18 column 4 and reproduced in Tables 28 column 2 and Table 35 column 2). This discounting factor was applied to net YLLs, before applying the outcome elasticity to calculate YLL averted.

As in previous Sections of this Chapter, the upper and lower bounds for the cost per QALY thresholds in column 3 are based on making the necessary assumptions about duration of health effects and how long a death might be averted optimistic (providing the lower bound for the threshold) or conservative (an upper bound for the threshold). The lower bound (lines 4 to 6 ) is based on assuming that health effects are not restricted to one year but apply to the whole of the remaining disease duration of the population at risk in PBCs during one year. Although this combines optimistic assumptions, it is possible that at least some part of a change in expenditure may prevent disease so will have an impact on populations that are incident to PBCs in the future. Such effects are not captured in any of the estimates presented in this Chapter so all are conservative with respect to this type of health effects of expenditure. The upper bound (lines 7 to 9) is based on the combination of assuming that health effects are restricted to one year for the population currently at risk and that any death averted is only averted for 2 years (see Section C.2.1.5).

## C.2.4. Which PBCs matter most?

Which PBCs have the greatest influence on the overall threshold depends, to a large extent, on how a change in overall expenditure is allocated to the different PBCs (see column 1 in table C.53), i.e., those that account for a greater share of the change in expenditure will tend to have the greater influence. ${ }^{15}$ However, the overall threshold also depends on the proportionate effect of a change in PBC expenditure on the QALY burden associated with the $\mathrm{PBC}^{16}$ and the scale of the QALY burden (for the population at

[^81]risk) associated with the type of diseases that make up each $\mathrm{PBC}^{17}$. These determine the cost per QALY associated with each PBC (see column 4 below). The share, attributable to each PBC, of the total health effects of a change in overall expenditure (see column 2) is the combined effect of all of these. The proportionate impact on the overall cost per QALY threshold of a $10 \%$ change in PBC health effects in gives an indication of how sensitive the overall threshold is to the estimate of health effects associated with each PBC (see column 3).

Table C.53: Impact of each PBC on the overall cost per QALY threshold


* Calculated using the effect on the threshold of a $10 \%$ increase (or decrease) in QALY change of the PBC.

Although the 11PBCs where outcome elasticities could be estimated only account for $27 \%$ of the change in overall expenditure they account for $83 \%$ of the overall health effects. Within this group some PBCs contribute more than others. Fon example, PBC11 (Respiratory) accounts for a greater share of total health effects and has a higher elasticity ( $4.60 \%$ ) than PBC10 (Circulatory) even though it accounts for a greater part of a change in overall expenditure. The reason is that the cost per QALY associated with changes in expenditure in PBC11 is lower than PBC10 and much lower than the overall threshold (so generates more health effects for the same, or even smaller, change in expenditure). The elasticities in column 3 are instructive, e.g., the elasticity for PBC11 suggests that even if the health effects of a change in expenditure in this PBC were overestimated by $30 \%$ the overall threshold would only increase by $13.8 \%$ to $£ 17,867$. All other PBCs have much less influence in this respect. Nonetheless PBC10 is importantcompared to others as it does contribute a large share of total health effects and has one of the highest elasticities ( $1.37 \%$ ).

The other 12 PBCs, where outcome elasticities could not be estimated, account for the greater part of a change in overall expenditure ( $73 \%$ ) but only $17 \%$ of the overall health effects, i.e., the cost per QALYs associated with a change in expenditure in these PBCs are, in general, much higher. Of course, we have not directly observed quality of life effects in these PBCs but inferred them from the proportionate effects that we can observe. Insofar as investment and disinvestment opportunities in these PBCs might

[^82]have been more valuable (offered greater improvement in quality of life) than suggested by the implied PBC thresholds in column 4, the overall QALY effects will tend to be underestimated and the overall cost per QALY threshold will be overestimated.

The overall threshold of $£ 15,701$ may be especially conservative (i.e., likely to be overestimated) with respect to health effects in PBC5 (Mental Health Disorders), which accounts for a large proportion of the change in overall expenditure ( $30 \%$ ) and contributes most to the overall health effects ( $7.85 \%$ ) compared to these other PBCs. The cost per QALY associated with this PBC $(£, 60,111)$ is based on an extrapolation of estimated proportionate effects to a population based measures of QALY burden in this PBC, rather than observations of the direct impact of changes in expenditure on quality of life in the types of diseases that make up the PBC. Evidence that is available suggests that the investment and disinvestment opportunities in this PBC are likely to have been much more valuable than this implied cost per QALY (Addendum 3 to this Appendix). A search for evidence about interventions in those ICD codes that contribute most to the PBC (based on prevalence or the contribution to the variance in PBC costs), suggests that pharmacological, psychological and social interventions for depression are all more cost effective (in general much less than $£ 10,000$ per QALY) than the overall threshold and significantly more valuable than the implied QALY threshold for this PBC. Based on the contribution that each ICD makes to variance in PBC costs across PCTs, it is schizophrenia that contributes most. Although interventions that may have been invested or disinvested in schizophrenia are, in general, less cost effective (in general less than $£ 24,000$ per QALY) than those available for depression, they are still much more valuable than the implied cost per QALY of this PBC in Table C. $52 .{ }^{18}$

## C.2.5. How uncertain are the estimates?

There are a number of sources of uncertainty which may contribute to an assessment of how uncertain a central or best estimate of the cost per QALY threshold might be. There are three reasons why uncertainty in the estimate of the threshold might be of policy interest: i) the uncertainty in the parameters that determine the threshold mightsinfluence the mean or expected value of the threshold if they have a non linear relationship to the threshold or when they have a multi linear relationship but are correlated with each other; ii) the consequences of over or underestimating the threshold differ so the uncertainty may have an influence on the extent to which a policy threshold (one that can be compared to the incremental cost effectiveness ratio of a new technology) should differ from the mean or expected value of the central or best estimate; and iii) in conjunction with other methods of analysis it can indicate the potential value of gathering more information to improve these estimates in the future. Such analysis, known as value of information analysis, has firm foundations in statistical decision theory and has been applied to health care decisions. A form of these analyses could be applied in subsequent research, ideally capturing some of the other sources of uncertainty. More recently it has been applied to the decisions faced by NICE when considering whether there is sufficient evidence to support the approval of a new technology.[7] Of course, hypothesis testing and the traditional rules of inference associated with it , such as statistical significance, p -values and confidence intervals, have no relevance when making unavoidable decisions aboat policy relevant quantities based on information currently available and the best use thereof $[8]$
(An assessment of parameter uncertainty
Two sets of parameters are critical to the threshold, the expenditure elasticities estimated for each of the 23 PBCs, and the outcome elasticities estimated for 11 of these. These parameters are estimated with uncertainty, indicated by the standard errors on the relevant coefficients in the econometric analysis

[^83]detailed in Appendix B. Since these statistical models estimate coefficients using normality on the relevant scale, normal distributions can be assigned to each of these estimated coefficients, each with a mean and standard deviation based on the results of the econometric analysis. These distributions represent the uncertainty in the mean estimate of each of the parameters and can be propagated through the various calculations required to estimate and overall cost per QALY threshold (i.e., through the sequence of analysis detailed in Section C.2.2 to C.2.4) using Monte Carlo simulation which randomly samples from the assigned distributions. The use of Monte Carlo simulation in this context is in essence Bayesian, where the standard errors from the frequentist econometric analysis are used to assign normal prior distributions with means equal to the point estimates and a standard deviation equal to the estimated standard errors. This is equivalent to a fully Bayesian analysis with initially uninformative priors which are updated through the analysis of expenditure and mortality data.

The results of each random sample from the Monte Carlo simulation represent one possible realisation of the overall threshold, given the uncertainty in estimates of the mean parameter values that determine it. By repeatedly sampling, a distribution of potential values that the overall threshold might take can be revealed. The results of this simulation are illustrated in Figure C. 6 showing a histogram of threshold values, and in Figure C. 7 showing the cumulative probability density function for a cost per QALY threshold based only on the 11 PBC with estimated outcome elasticities and for all 23 PBCs. It represents the probability (on the y axis) that the threshold lies below a particular value.


Figure C. 6 Distribution of the cost per QALY threshold (all 23 PBCs)


Figure C. 7 Cumulative probability density function for the cost per QALY threshold

It has already been noted that restricting attention only to changes in expenditure in those 11PBC where an outcome elasticity can be estimated is much lower than considering all changes in expenditure across all PBCs - the threshold value of the x axis that corresponds to a probability of 0.5 is much lower in Figure C. 6 for these $11 \mathrm{PBCs}-£ 5,144$ vs. $£ 15,607$. This lower estimate of $£, 5,144$ per QALY is much less uncertain but these PBCs only account for $27 \%$ of a change in overall expenditure, so it is the higher estimate, for all 23 PBCs , that is of most relevance for policy (see Sections C.2.3.3 and C.2.4). The fact that this estimate is more uncertain simply reflects the quality and quantity of data currently available. Since useful analysis should endeavour to faithfully characterise uncertainty in policy relevant quantities, rather than select those quantities or questions for which precise estimates are possible, it is the more uncertain estimate for all 23 PBCs that should be of primarily interest. The values that are used to generate Figure C. 6 are available in column 2 of Table C.54. They indicate that the probability that the overall threshold is less than $£ 20,000$ per QAL is 0.84 and the probability that is less than $£ 30,000$ is 0.99 .

Table C.54: Uncertainty over the QALY threshold.

| S |  | $\begin{gathered} \hline \text { All } 23 \text { PBCs } \\ {[2]} \\ \hline \end{gathered}$ |
| :---: | :---: | :---: |
| Best estimate (deterministic) | ¢,5,128 | £15,701 |
| Mean estimate (from the simulations) | ¢,5,114 | £,15,634 |
| Threshold value at the probability of (from the simulations): |  |  |
| 2.5\% | £3,553 | £10,963 |
| 5.0\% | £3,776 | £11,591 |
| 50.0\% | £,5,144 | £15,607 |
| 95.0\% | £ 7,154 | £25,159 |
| 97.5\% | £, 7,812 | $£ 28,212$ |
| Probability (from the simulations)of the threshold being smaller than: |  |  |
| £3,000 perQALY | 0.00 | 0.00 |
| \&4,000 per QALY | 0.09 | 0.00 |
| 6,5000 per QALY | 0.44 | 0.00 |
| 66,000 per QALY | 0.79 | 0.00 |
| $\bigcirc$ ¢ $£, 000$ per QALY | 0.94 | 0.00 |
| $£ 8,000$ per QALY | 0.98 | 0.00 |
| $£^{2} 9,000$ per QALY | 0.99 | 0.00 |
| £,10,000 per QALY | 1.00 | 0.01 |
| $£ 15,000$ per QALY | 1.00 | 0.44 |
| $\AA 20,000$ per QALY | 1.00 | 0.84 |
| $£ 25,000$ per QALY | 1.00 | 0.95 |
| $£ 30,000$ per QALY | 1.00 | 0.99 |
| $\AA 35,000$ per QALY | 1.00 | 0.99 |
| $\AA .40,000$ per QALY | 1.00 | 1.00 |

## C. 3 Re-estimating the cost per QALY threshold using 2008 expenditure data

The same methods of analysis can be applied to the econometric analysis of the 2008/09 expenditure and 2008 to 2010 mortality data (see Section 3.5.3 in Chapter 3 and Section B11 in Appendix B). The differences between the 2006 analysis reported above and the analysis of expenditure in 2008 reported below are the: i) total PBC expenditure ii) estimated expenditure elasticities; iii) estimated outcome elasticities; iv) observed PBC deaths by age and gender; and v) life expectancy by age and gender. The other information about quality of life norms (see Section C.2.2.2), disease related decrements (see Section C.2.2.3) and the information from GBD about incidence (by age and gender) and duration of disease (C.2.3) remain unchanged between 2006 and 2008.

It should be noted that important improvements were made to the classification and collection of PBC expenditure data that took place after the 2006 data were collected. Therefore, the differences in threshold estimates for 2006 and 2008 partly reflect this (see Section 3.5.4 and B11.4 in Appendix B) so should not be over interpreted. The results of the analysis of 2007 and 2008 expenditure are comparable in this respect, providing insights into how the threshold might change over time and with changes in the overall budget. For the purposes of this methodological research the 2008 expenditure and 2008 to 2010 mortality data were the latest to be analysed.

Table C.55: Outcome and spend elasticities (2008)

| PBC | PBC description | Total spend 2008/09, ( € $^{\text {) }}$ <br> [1] | Spend unadjusted $[2]$ | $\qquad$ | Outcome elasticities* [3] |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 2 | Cancer | £4,843 | 0.525 | 0.525 | 0.307 |
| 10 | Circulatory problems | £6,655 | 0.648 | 0.648 | 1.319 |
| 11 | Respiratory problems | £3,994 | 0.652 | 0.652 | 1.808 |
| 13 | Gastro-intestinal problems <br> Big 4 | $£^{2}, 19,481$ | 0.456 | 0.456 | 1.364 |
| 1 | Infectious diseases | £1,201 | ค 1.545 | 1.545 | 0.504 |
| 4 | Endocrine problems | £2,222 | 00.484 | 0.484 | 1.170 |
| 7 | Neurological problems | £3,466 | 0.980 | 0.980 | 0.417 |
| 17 | Genito-urinary problems | $£ 3,779$ | 0.697 | 0.697 | 1.615 |
| 16 | Trauma \& injuries* | £3,255 | 1.344 | 1.344 | - |
| 18+19 | Maternity \& neonates* <br> First 11 PBC's | $\begin{array}{r} £ 3,978 \\ \qquad 37,382 \end{array}$ | 0.975 | 0.975 | 0.125 |
| 3 | Disorders of Blood | ¢ 998 | 1.171 | 2.291 | - |
| 5 | Mental Health Disorders | ¢, 9,794 | 1.036 | 2.027 | - |
| 6 | Learning Disability | £2,874 | 0.205 | 0.401 | - |
| 8 | Vision | £1,688 | 0.654 | 1.279 | - |
| 9 | Hearing | $£ 417$ | 1.191 | 2.330 | - |
| 12 | Dental problems | £ 2,198 | 0.513 | 1.003 | - |
| 14 | Problems of the Skin | £1,657 | 0.674 | 1.318 | - |
| 15 | Musculo-skeletal system | £4,081 | 0.505 | 0.988 | - |
| 20 | Poisoning and AE | $\AA 938$ | 0.562 | 1.099 | - |
| 21 | Healthy Individuals | £1,831 | 1.097 | 2.146 | - |
| 22 | Social Care Needs | £1,874 | 0.911 | 1.782 | - |
| 23 | Other All (23 PBCs) | $\begin{aligned} & £ 11,666 \\ & £, 78,398 \\ & \hline \end{aligned}$ | 0.494 | 0.494 | - |

## C.3.1 From mortality to life years

In this section we summarise report the calculation of net YLL, which take account of the fact that some of the observed deaths would have occurred anyway (had the same population not been at risk in the particular PBC) when estimating YLL (unobserved counterfactual deaths). In summary, to obtain net YLL, all observed deaths - both those that occur below and those that occur above LE (Table C.56) - are taken into account. Those deaths occurring below LE generate YLL and those that occur at ages above LE generate life years 'gained' (YLG). By subtracting YLG from YLL to generate net YLL we take account of the fact that not all deaths below LE are excess deaths but some deaths above LE are.

Table C.56. Number of deaths above LE in 2008/9/10, by PBC

| PBC |  | $\begin{gathered} <\mathrm{LE} \\ 2008 \\ {[1]} \end{gathered}$ | $\begin{aligned} & >L E \\ & 2008 \end{aligned}$ <br> [2] | $\begin{gathered} <\mathrm{LE} \\ 2009 \\ {[3]} \end{gathered}$ | $\begin{gathered} >\mathrm{LE} \\ 2009 \\ {[4]} \end{gathered}$ | $\begin{aligned} & <\mathrm{LE} \\ & 2010 \end{aligned}$ [5] | $\begin{aligned} & >\text { LE } \\ & 2010 \end{aligned}$ <br> [6] | Annual N deaths $<$ LE [7] | Annual N deaths $>$ LE [8] |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | Infectious diseases | 3,406 | 2,586 | 3,044 | 2,190 | 2,667 | 1,894 | 3,039 | 2,223 |
| 2 | Cancer | 94,873 | 37,029 | 94,276 | 37,151 | 94,309 | 38,198 | 94,486 | 37,459 |
| 4 | Endocrine | 4,033 | 2,877 | 3,834 | 2,826 | 3,816 | 2,902 | 3,894 | 2,868 |
| 7 | Neurological | 9,638 | 6,859 | 9,445 | 6,939 | 9,951 | 7,480 | 9,678 | 7,093 |
| 10 | Circulatory | 80,894 | 76,292 | 76,048 | 73,342 | 74,035 | 73,719 | 76,992 | 74,451 |
| 11 | Respiratory | 32,083 | 35,180 | 29,912 | 33,304 | 29,691 | 33,176 | 30,562 | 33,887 |
| 13 | Gastro-intestinal | 15,945 | 8,259 | 15,361 | 8,161 | 15,595 | 8,372 | 15,633 | 8,264 |
| 17 | Genito-urinary | 4,471 | 6,667 | 4,378 | 6,900 | 4,453 | 7,166 | 4,434 | 6,911 |
| 18+19 | Maternity \& neonates | 267 | 0 | 281 | 1 | 247 | 0 | 265 | 0 |

The estimates of net YLL calculated considering estimates of the life expectancy for each PBC are detailed in Table C.57.

Table C.57. Net YLL using life expectancy for each PBC (2008)

| PBC |  | LE of Males <br> [1] | LE of Females [2] | Average2006-2008 |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | Deaths |  |  |  |  |
|  |  |  |  | $\begin{gathered} <\mathrm{LE} \\ {[3]} \end{gathered}$ | $\begin{gathered} >\mathrm{LE} \\ {[4]} \\ \hline \end{gathered}$ | YLL [5] |  | Net YLL <br> [7] |
| 1 | Infectious diseases | 79.6 | 83.6 | 2,919 | 2,344 | 53,926 | 15,132 | 38,794 |
| 2 | Cancer | 83.0 | 84.7 | 100,487 | 31,459 | 1,456,255 | 134,089 | 1,322,166 |
| 4 | Endocrine | 81.0 | 84.7 | 3,945 | 2,818 | -65,800 | 15,983 | 49,817 |
| 7 | Neurological | 79.6 | 83.3 | 9,112 | 7,659 | 137,791 | 47,722 | 90,069 |
| 10 | Circulatory | 83.0 | 86.5 | 89,434 | 62,009 | 1,049,459 | 278,421 | 771,038 |
| 11 | Respiratory | 80.3 | 84.0 | 29,828 | 34,621 | 306,838 | 229,403 | 77,434 |
| 13 | Gastro-intestinal | 80.6 | 84.5 | 15,612 | 8,286 | 271,395 | 46,141 | 225,254 |
| 17 | Genito-urinary | 83.5 | 85.6 | 5,058 | 6,287 | 49,036 | 32,528 | 16,508 |
| 18+19 | Maternity \& neonates | 78.7 | 83.1 | 265 | 0 | 19,783 | 1 | 19,781 |

The impact on the cost per life year threshold is summarised in column 2 of Table C.58, and a detailed breakdown in Table C. 59 .

Table C.58. Summary of cost per life year threshold (2008)

| $(())$ | $\begin{gathered} 2006 \\ {[1]} \\ \hline \end{gathered}$ | $\begin{gathered} 2008 \\ {[2]} \\ \hline \end{gathered}$ |
| :---: | :---: | :---: |
| big 4 PBC's | £,8,080 | £,10,220 |
| 11 PBCs (with mortality) | £15,628 | £23,360 |
| All 23 PBCs (zero health effects forremaining 12 PBCs ) | £57,497 | £64,275 |
| All 23 PBCs (non-zero health effects for remaining 12 PBCs, except GMS)* | $£ 17,663$ | $£ 25,214$ |

* in PBCs without a mortality signal, health effects were estimated by valuing changes in expenditure at the same rate as observed in PBCs for which there yas a mortality signal.

Table C.59: Breakdown of the cost per life year threshold (2008).

| PBC | PBC description | Change in spend, $£ \mathrm{~m}$ [1] | Using LE of the PBC population |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Net YLL <br> [5] | Change in net YLL <br> [6] | Cost per LY gained, $£$ [7] |
| 2 | Cancer | £25 | 1322166 | 2131 | £11,931 |
| 10 | Circulatory problems | £43 | 771038 | 6590 | £6,544 |
| 11 | Respiratory problems | £26 | 77434 | 913 | £28,528 |
| 13 | Gastro-intestinal problems Big 4 | £18 | 225254 | 1401 | $\begin{aligned} & £ 12,983 \\ & £, 10,220 \\ & \hline \end{aligned}$ |
| 1 | Infectious diseases | £19 | 38794 | 302 | £61,425 |
| 4 | Endocrine problems | £11 | 49817 | 282 | £38,122 |
| 7 | Neurological problems | £34 | 90069 | 368 | $¢ 92,282$ |
| 17 | Genito-urinary problems | £26 | 16508 | 186 | £141,746 |
| 16 | Trauma \& injuries* | $£ 44$ | NA | 0 | NA |
| 18+19 | Maternity \& neonates* <br> First 11 PBC's | £39 | 19781 | 24 | $\begin{array}{r} £ 1,608,817 \\ £, 23,360 \\ \hline \end{array}$ |
| 3 | Disorders of Blood | £23 |  | 979 | £23,360 |
| 5 | Mental Health Disorders | £198 |  | 8496 | £23,360 |
| 6 | Learning Disability | £12 |  | 493 | £23,360 |
| 8 | Problems of Vision | £22 |  | 924 | £23,360 |
| 9 | Problems of Hearing | $£ 10$ |  | 416 | £23,360 |
| 12 | Dental problems | £32 |  | 1374 | £23,360 |
| 14 | Skin | $£_{22}$ |  | 935 | £23,360 |
| 15 | Musculo skeletal system | $£ 40$ |  | 1726 | \&23,360 |
| 20 | Poisoning and AE | £10 |  | 441 | £23,360 |
| 21 | Healthy Individuals | £39 |  | 1682 | $\underset{\sim}{23,360}$ |
| 22 | Social Care Needs | £33 |  | 1430 | +23,360 |
| 23 | Other | $\ldots 58$ |  | 0 | NA |
|  | All (23 PBCs) |  |  | $\bigcirc$ | £25,214 |

Note that we have been unable to obtain a satisfactory outcome model for trauma \& injuries and haye assumed a zero outcome elasticity.
Note that, for expenditure in 2006/7, the neonate category has been merged with maternity to obtain plausible outcome and expenditure models.
The estimates of net YLL imply a number of excess deaths required to generate them in each PBC. The implied excess deaths associated with net YLL are reported in Table C.60.

Table C.60: Excess deaths implied by net YLL (2008).

| PBC |  | YLL per observed Net YLL death [1] [2] |  | Excess deaths <br> [3] | Total deaths $[4]$ | \% excess <br> deaths [5] |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | Infectious diseases | 38,794 | 13.4 | 2,934 | 5,262 | 56\% |
| 2 | Cancer | 1,322,166 | 14.1 | 93,917 | 131,945 | 71\% |
| 4 | Endocrine | 49,817 | 13.7 | 3,663 | 6,762 | 54\% |
| 7 | Neurological | 90,069 | 13.6 | 6,642 | 16,771 | 40\% |
| 10 | Circulatory | 771,038 | 10.5 | 74,217 | 151,443 | 49\% |
| 11 | Respiratory | 77,434 | 9.2 | 8,432 | 64,449 | 13\% |
| 13 | Gastro-intestinal | 225,254 | 15.2 | 15,049 | 23,897 | 63\% |
| 17 | Genito-urinary | 16,508 | 8.3 | 1,978 | 11,345 | 17\% |
| 18+19 | Maternity \& neonates | 19,781 | 74.1 | 265* | 265 | 100\% |

Excess deaths are calculated for each gender by dividing net YLLs by the YLL per death (column [3] = column [1] / column [2] )

* The number of excess deaths estimated in PBC18\&19 was initially estimated to be 265, higher than the number of total deaths. This is due to the use of approximations (i.e. in the life expectancy, or in using the net YLL) thus, for consistency, we assumed this to be $100 \%$ of the total deaths.

The cost per excess death and the cost per PBC death averted are reported in Table C.61, and a detailed breakdown of changes in spend and excess or total deaths across PBCs is shown in Table C.62. The cost per PBC death averted is, of course; significantly lower than the cost per excess death as excess deaths are only a proportion of total deaths (see Table C.61).

Table C.61. Summary of the cost per death averted threshold (2008)

|  | 2006-2008 |  | 2008-2010 |  |
| :---: | :---: | :---: | :---: | :---: |
|  | Cost per excess death averted, $£$ [1] | Cost per PBC death averted, $£$ [2] | Cost per excess death averted, $£$ [3] | Cost per PBC death averted, $\ell$ [4] |
| big 4 PBC's | $£^{91,129}$ | £32,864 | £115,234 | £46,692 |
| 11 PBCs (with mortality) | £177,691 | £64,774 | £265,784 | £105,872 |
| All 23 PBCs (zero health effects for remaining 12 PBCs) | $£ 653,744$ | $£ 238,310$ | £731,301 | $£ 291,305$ |
| All 23 PBCs (non-zero health effects for remaining 12 PBCs, except GMS)* | $£ 200,828$ | £73,208 | £286,872 | £114,272 |

* in PBCs without a mortality signal, health effects were estimated by valuing changes in expenditure at the same rate as observed in PBCs for which there was a mortality signal.

Table C.62: Breakdown of the cost per death averted threshold (2008).

| PBC | PBC description | Change in spend, £m [1] | Total <br> PBC <br> deaths <br> [2] | PBC dea Change in PBC deaths [3] | Cost per PBC death averted, $£$ <br> [4] | Excess deaths [5] | Excess Change in excess deaths [6] | Cost per excess death averted, $£$ [7] |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 2 | Cancer | £25 | 131945 | 212.66 | £119,559 | 93917 | 151.37 | £167,969 |
| 10 | Circulatory problems | $\AA 43$ | 151443 | 1294.40 | £33,316 | 74217 | 634.34 | £67,983 |
| 11 | Respiratory problems | £26 | 64449 | 759.74 | $\AA 34,276$ | 8432 | 99.40 | £261,992 |
| 13 | problems $\text { Big } 4$ | $£ 18$ | 23897 | 148.64 | $\begin{array}{r} £ 122,379 \\ £ 46,692 \end{array}$ | 15049 0 | 93.60 | $\begin{array}{r} £ 194,332 \\ £, 115,234 \\ \hline \end{array}$ |
| 1 | Infectious diseases | £19 | 5262 | 40.97 | £452,858 | $\checkmark 2934$ | 22.84 | £ 812,249 |
| 4 | Endocrine problems | £11 | 6762 | 38.29 | £280,856 | 3663 | 20.74 | £,518,533 |
| 7 | Neurological problems Genito-urinary | £34 | 16771 | 68.54 | ¢,495,603 | 6642 | 27.14 | £1,251,391 |
| 17 | problems | £26 | 11345 | 127.71 | £206,253 | 1978 | 22.27 | £1,182,744 |
| 16 | Trauma \& injuries* | $\AA 44$ | NA | 0 | NA | NA | 0 | NA |
| 18+19 | Maternity \& neonates* First 11 PBC's | £39 | 265 | 0.32 | $\begin{array}{r} 120,090,566 \\ £ 105,872 \\ \hline \end{array}$ | 265 | 0.32 | $\begin{array}{r} £ 120,090,566 \\ £, 265,784 \\ \hline \end{array}$ |
| 3 | Disorders of Blood Mental Health | $£ 23$ |  | 215.92 | £105,872 |  | 86.01 | £265,784 |
| 5 | Disorders | $£ 198$ |  | 874.69 | £105,872 |  | 746.76 | £265,784 |
| 6 | Learning Disability | £12 |  | 108.86 | £105,872 |  | 43.36 | £265,784 |
| 8 | Problems of Vision | £22 |  | 203.97 | £105,872 |  | 81.25 | £265,784 |
| 9 | Problems of Hearing | £10 |  | 91.76 | £105,872 |  | 36.55 | £265,784 |
| 12 | Dental problems | £.32 |  | 303.11 | £105,872 |  | 120.74 | £265,784 |
| 14 | Skin | f22 |  | 206.34 | £105,872 |  | 82.19 | £265,784 |
| 15 | Musculo skeletal system | $\mathscr{L} 40$ |  | 380.77 | £105,872 |  | 151.68 | £265,784 |
| 20 | Poisoning and AE | $£ 10$ |  | 97.40 | £105,872 |  | 38.80 | £265,784 |
| 21 | Healthy Individuals | £39 |  | 371.11 | £,105,872 |  | 147.83 | £265,784 |
| 22 | Social Care Needs | £33 |  | 315.43 | £105,872 |  | 125.65 | £265,784 |
| 23 | Other | ¢, 58 |  | 0 | NA |  | 0 | NA |
|  | All (23 PBCs) |  |  |  | £,114,272 |  |  | £286,872 |

The number of life years gained associated with each excess death averted are reported for each PBC in TableC. 63 (column 1) and range from 74.6 years for $\mathrm{PBC} 18 \& 19$ to 8.3 years for PBC17. On average, across all 11 PBCs each excess death averted is associated with 11.4 life years gained. The life years associated with each observed death are reported for each PBC in (column 2) and range from 74.6 years in PBC $18 \& 19$ to 1.2 for PBC17. On average across all 11 PBCs each PBC death averted is associated with 4.5 life years gained.

Table C.63: Implied YLL per death averted for each PBC (2008)

| PBC | PBC description | Implied YLL per excess death averted <br> [1] | Implied YLL per <br> PBC death averted <br> [2] |
| :---: | :---: | :---: | :---: |
| 2 | Cancer | 14.1 | 10.0 |
| 10 | Circulatory problems | 10.4 | 5.1 |
| 11 | Respiratory problems | 9.2 | 1.2 |
| 13 | Gastro-intestinal problems | 15.0 | 9.4 |
|  | Big 4 | 11.3 | 4.6 |
| 1 | Infectious diseases | 13.2 | 7.4 |
| 4 | Endocrine problems | 13.6 | 7.4 |
| 7 | Neurological problems | 13.6 | 5.4 |
| 17 | Genito-urinary problems | 8.3 | 1.5 |
| 16 | Trauma \& injuries | NA | NA |
| 18+19 | Maternity \& neonates | 74.6 | 74.6 |
|  | First 11 PBC's | 11.4 | 4.5 |

## Summary of cost per life year estimates

The cost per life year threshold in lines 1 to 4 in Table C. 64 are regarded as the central or best estimates given the evidence available and the credibility of alternative assumption that could be made. As explained in Section C.1, these are based on the conservative assumption that any health effects of changes in expenditure are restricted to one year, which, to some extent, may be offset by the more optimistic assumption any death averted returns the individual to the mortality risk face by the general population, matched for age and gender. See Section C. 2.5 for guidance in the interpretation of the upper and lower bound estimates.

Table C.64: Summary of the cost per life year threshold with upper and lower bounds (2008)

|  | $\underset{11]}{2006-2008}$ | 2008-2010 |  |
| :---: | :---: | :---: | :---: |
| Effect of expenditure on mortality: YLL per PBC death averted: | $\begin{gathered} 1 \text { year } \\ \sim 4.1 \text { YLL } \\ \\ \sim * \end{gathered}$ | ate $\begin{gathered} 1 \text { year } \\ \sim 4.5 \text { YLL } * * \end{gathered}$ |  |
| big 4 PBC's | £,8,080 | £10,220 | [1] |
| 11 PBCs (with mortality) | £15,628 | $£ 23,360$ | [2] |
| All 23 PBCs (zero health effects for remaining 12 PBCs) | £57,497 | £64,275 | [3] |
| All 23 PBCs (non-zero health effects for remaining 12 PBCs, except GMS)* | £17,663 | $£ 25,214$ | [4] |
| $\bigcirc$ | Lower bound |  |  |
| Effect of expenditure on mortality: YL per PBC death averted: | Remainder of disease ~ 4.1 YLL ** | Remainder of disease <br> $\sim 4.5$ YLL ** |  |
| big 4 PBC's | £3,846 | £5,083 | [5] |
| 11 PBCs (with mortality) | £6,106 | £8,579 | [6] |
| All 23 PBCs (zero health effects for remaining 12 PBCs) | $£ 22,463$ | $£ 23,605$ | [7] |
| All 23 PBCs (non-zero health effects for remaining 12 PBCs, except GMS)* | $£_{6} 6,901$ | £ ${ }^{9}$,260 | [8] |
|  | Upper bound |  |  |
| Effect of expenditure on mortality: YLL per PBC death averted: | $\begin{aligned} & 1 \text { year } \\ & 2 \text { YLL } \end{aligned}$ | $\begin{aligned} & 1 \text { year } \\ & 2 \text { YLL } \end{aligned}$ |  |
| big 4 PBC's | £16,432 | £23,346 | [9] |
| 11 PBCs (with mortality) | ¢ 32,387 | ¢52,936 | [10] |
| All 23 PBCs (zero health effects for remaining 12 PBCs) | £119,155 | £145,653 | [11] |
| All 23 PBCs (non-zero health effects for remaining 12 PBCs, except GMS)* | £36,604 | £57,136 | [12] |

* in PBCs without a mortality signal, health effects were estimated by valuing changes in expenditure at the same rate as observed in PBCs for which there was a mortality signal.
** see Table C. 63


## C.3.2 Adjusting life years for quality of life

The central or best estimates of the cost per life year threshold, which were presented in Table C. 64 (lines 2 and 4) take no account of the health related quality of life in which years of life, expected to be gained or lost through changes in expenditure, are likely to be lived. In this Section we examine the ways in which the life years reported in Section C.3.2 can be adjusted for quality, taking account of information that is available about: i) how quality of life differs by age and gender, and ii) how the quality of life years associated with mortality changes might be affected by the types of diseases that make up each PBC.

## Quality of life based on the general population

Quality of life norms (in Figure C.4) can be applied to the YLL associated with all observed deaths in each PBC, taking account of gender and age at death. The results are reported in column 4 to 6 of Table C.65.

Table C.65: Net YLL adjusted for quality of life 'norms' (2008)

| PBC |  | Unadjusted life years |  |  | Quality adjusted vife years |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | YLL | YLG | YLL | YLG | [LL | YLG |
|  | [1] | $[2]$ | $[3]$ | $[4]$ | $[5]$ | $[6]$ |  |
| 1 | Infectious diseases | 53,926 | 15,132 | 38,794 | 43,703 | 10,187 | 33,516 |
| 2 | Cancer | $1,456,255$ | 134,089 | $1,322,166$ | $1,129,191$ | 89,231 | $1,039,960$ |
| 4 | Endocrine | 65,800 | 15,983 | 49,817 | 52,465 | 10,598 | 41,867 |
| 7 | Neurological | 137,791 | 47,722 | 90,069 | 110,532 | 32,262 | 78,270 |
| 10 | Circulatory | $1,049,459$ | 278,421 | 771,038 | 807,893 | 183,796 | 624,097 |
| 11 | Respiratory | 306,838 | 229,403 | 77,434 | 237,981 | 154,300 | 83,680 |
| 13 | Gastro-intestinal | 271,395 | 46,141 | 225,254 | 214,756 | 30,811 | 183,945 |
| 17 | Genito-urinary | 49,036 | 32,528 | 16,508 | 37,178 | 21,190 | 15,989 |
| $18+19$ | Maternity \& neonates | 19,783 | 1 | 19,781 | 17,176 | 1 | 17,175 |

The implications of the quality adjustment to a cost per QALY threshold that only accounts for the direct health effects of mortality are summarised in Table C.66, and detailed in Table C.67.

Table C.66: Summary of cost per QALY threshold based on population norms and mortality effects (2008)

|  | Cost per life year <br> threshold | Cost per QALY <br> threshold <br> Population norms | Cost per life year <br> threshold | Cost per QALY <br> threshold <br> Population norms |
| :--- | :---: | :---: | :---: | :---: | :---: |
|  |  | $[1 N$ |  |  |

Table C.67: A breakdown of the cost per QALY threshold based on population norms (2008)


Table C. 68 depicts the judgements over life years, quality of life weights and total QALYs implicit in calculations of the threshold cost per QALY in Tabec.64.

Table C.68: Implied YLL per excess death ayerted and implied QoL score per YLL gained, for each PBC (2008)

| PBC | PBC description | Implied YLL per excess death averted [1] | Implied YLL per PBC death averted [2] | Implied QALYs gained per excess death averted [3] | Implied QALYs gained per PBC death averted [4] |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 2 | Cancer | 14.08 | 10.02 | 11.07 | 7.88 |
| 10 | Circulatory | 10.39 | 5.09 | 8.41 | 4.12 |
| 11 | Respiratory | 9.18 | 1.20 | 9.92 | 1.30 |
| 13 | Gastro-intestinal | 14.97 | 9.43 | 12.22 | 7.70 |
|  | Big 4 | 11.28 | 4.57 | 9.34 | 3.78 |
| 1 | Infectious diseases | 13.22 | 7.37 | 11.42 | 6.37 |
| 4 | Endocrine | 13.60 | 7.37 | 11.43 | 6.19 |
| 7 | Neurological | 13.56 | 5.37 | 11.78 | 4.67 |
| 17 | Genito-urinary | 8.34 | 1.46 | 8.08 | 1.41 |
| 16 | Trauma \& injuries* | NA | NA | NA | NA |
| 18+19 | Maternity \& neonates* | 74.65 | 74.65 | 64.81 | 64.81 |
|  | First 11 PBC's | 11.38 | 4.53 | 9.48 | 3.78 |

Adjusting age related quality of life for disease decrements
By using age related quality of life disease decrements (exemplified in Figure C.5) YLL can be adjusted for quality of life of disease. The results are reported in column 4 to 6 of Table C.69.

Table C.69: Net YLL adjusted for disease and age related quality of life (2008)

| PBC |  | Unadjusted life years |  |  | Quality adjusted life years |  |  |
| :---: | :--- | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | YLL | YLG | YLL | YLG | YLL | YLG |
|  |  | $[1]$ | $[2]$ | $[3]$ | $[4]$ | $[6]$ | 7,524 |
| 1 | Infectious diseases | 53,926 | 15,132 | 38,794 | 34,108 | 26,584 |  |
| 2 | Cancer | $1,456,255$ | 134,089 | $1,322,166$ | 943,650 | 72,197 | 871,452 |
| 4 | Endocrine | 65,800 | 15,983 | 49,817 | 43,063 | 8,334 | 34,729 |
| 7 | Neurological | 137,791 | 47,722 | 90,069 | 69,520 | 18,084 | 51,436 |
| 10 | Circulatory | $1,049,459$ | 278,421 | 771,038 | 625,150 | 135,622 | 489,527 |
| 11 | Respiratory | 306,838 | 229,403 | 77,434 | 173,953 | 106,200 | 67,754 |
| 13 | Gastro-intestinal | 271,395 | 46,141 | 225,254 | 162,441 | 22,060 | 140,380 |
| 17 | Genito-urinary | 49,036 | 32,528 | 16,508 | 30,770 | 16,949 | 13,820 |
| $18+19$ | Maternity \& neonates | 19,783 | 1 | 19,781 | 16,100 | 1 | 16,099 |

The implications of the quality adjustment to a cost per QALY threshold that only accounts for the direct health effects of mortality are summarised in Table C.70, and detailed in Table C.71.

Table C.70: Summary of cost per QALY threshold based on disease and age related quality of life and mortality effects (2008)


* in PBCs without a mortality signal, health effects were estimated by valuing changes in expenditure at the same rate as observed in PBCs for which there was a mortality signal except GMS.

Table C.71: A breakdown of the cost per QALY threshold based on disease and age related quality of life and mortality effects (2008)

|  | N |  | g LE of PBC |
| :---: | :---: | :---: | :---: |
| PBC PBC description | Change in spend, fm 11 | Change in QALY <br> [2] | Cost per QALY gained, $£$ [3] |
| 2 Cancer | £25 | 1405 | £18,102 |
| 10 Circulatory problems | £43 | 4184 | £10,307 |
| 11 Respiratory problems | £26 | 799 | $£ 32,604$ |
| 13 Gastro-intestinal problem | £18 | 873 | $\AA 20,833$ |
| Big 4 |  |  | £,15,534 |
| 1 Infectious diseases | £19 | 207 | £89,638 |
| 4 Endocrine problems | £11 | 197 | £54,685 |
| 7 Neurological problems | £34 | 210 | £161,594 |
| 17 Genito-urinary problems | £26 | 156 | £169,315 |
| 16 Trauma \& injuries* | £44 | 0 | NA |
| 18+19 Maternity \& neonates* | £39 | 20 | £1,976,769 |
| First 11 PBC's |  |  | £ 35,397 |
| (3) Disorders of Blood | $£_{23}$ | 646 | £35,397 |
| 5 Mental Health Disorders | £198 | 5607 | £35,397 |
| ) 6 Learning Disability | £12 | 326 | £35,397 |
| 8 Problems of Vision | £22 | 610 | £35,397 |
| 9 Problems of Hearing | £10 | 274 | £35,397 |
| 12 Dental problems | £.32 | 907 | £35,397 |
| 14 Skin | $\AA 22$ | 617 | £35,397 |
| 15 Musculo skeletal system | $\AA 40$ | 1139 | $\AA 35,397$ |
| 20 Poisoning and adverse effects | $£ 10$ | 291 | $£ 35,397$ |
| 21 Healthy Individuals | $£ 39$ | 1110 | £35,397 |
| 22 Social Care Needs | £33 | 943 | £35,397 |
| 23 Other | £.58 | 0 | NA |
| All (23 PBCs) |  |  | £38,206 |

Table C. 72 depicts the judgements over life years, quality of life weights and total QALYs implicit in calculations of the threshold cost per QALY in Table C.70.

Table C.72: Implied YLL per excess death averted and implied QoL score per YLL gained, for each PBC (2008)


## Summary of the cost per QALY threshold based only on mortality effects

The analysis to this point is summarised in Table C.73. The three estimates of a cost per QALY threshold are based on assuming that each life year gained is either: lived in full health (see column 1), lived in a quality of life that reflects age and gender norms of the general population (column 2); or lived in a quality of life that reflects the original disease state (column 3).

Table C.73: Summary of QALY threshold estimates based only on mortality effects (2008)

|  | $\begin{gathered} {[1]} \\ (Q o L \text { score }=1) \end{gathered}$ | [2] (QoL norm) | [3] (QoL diseased) |  |
| :---: | :---: | :---: | :---: | :---: |
| Effect of expenditure on mortality: <br> YLL per death averted *: QALYs per death averted *: |  | $\begin{gathered} \text { Best estimate } \\ 1 \text { year } \\ \sim 4.5 \mathrm{YLL} \\ \sim 3.8 Q \mathrm{ALY} \\ \hline \end{gathered}$ | $\begin{aligned} & 1 \text { year } \\ \sim & 4.5 \mathrm{YLL} \\ \sim & 3.0 \text { QALY } \end{aligned}$ |  |
| big 4 PBC's <br> 11 PBCs <br> All 23 PBCs | 10,220 $£ 23,360$ $£ 25,214$ | $\begin{aligned} & £ 12,338 \\ & £ 28,045 \\ & £ 30,270 \end{aligned}$ | $\begin{aligned} & £ 15,534 \\ & £ 35,397 \\ & £ 38,206 \end{aligned}$ | [1] [2] [3] |
| Effect of expenditure on mortality: <br> YLL per death averted *: QALYs per death averted *: | Remainder of disease $\begin{gathered} \sim 4.5 \mathrm{YLL} \\ \sim 4.5 \text { QALY } \end{gathered}$ | Lower bound <br> Remainder of disease <br> ~ 4.5 YLL <br> ~ 3.8 Q ALY | Remainder of disease <br> ~ 4.5 YLL <br> ~ 3.0 Q ALY |  |
| big 4 PBC's <br> 11 PBCs <br> All 23 PBCs | $\begin{aligned} & £ 5,083 \\ & £ 8,579 \\ & £, 9,260 \end{aligned}$ | $\begin{aligned} & £ 5,811 \\ & £, 9,861 \\ & £ 10,644 \end{aligned}$ | $\begin{gathered} £ 7,305 \\ £ 12,720 \\ £ 13,729 \end{gathered}$ | $[4]$ $[5]$ $[6]$ |
| Effect of expenditure on mortality: YLL per death averted *: QALYs per death averted *: | $\begin{aligned} & 1 \text { year } \\ & 2 Y L L \\ \sim & 2 \text { QALY } \end{aligned}$ | $\begin{gathered} \text { Upper bound } \\ 1 \text { year } \\ 2 \mathrm{YLL} \\ \sim 1.8 Q A L Y \end{gathered}$ | $\begin{gathered} 1 \text { year } \\ 2 \mathrm{YLL} \\ \sim \\ 1.4 \text { QALY } \end{gathered}$ |  |
| big 4 PBC's | £23,346 | £26,138 | £32,797 | [7] |
| 11 PBCs | £52,936 | £59,151 | £74,183 | [8] |
| All 23 PBCs | ¢,57,136 | £ 63,844 | ¢,80,069 | [9] |

. see Table C. 72

## C.3.3. Including quality of life effects during disease

In this section we explore how estimates of effects of expenditure that can be observed (i.e., on mortality) can be used to infer the likely effects on what cannot be directly observed (quality of life), rather than making extreme assumptions that are not credible (e.g., assuming that changes in expenditure will have no effects on quality of life outcomes). In Section C.2.3.2, we described the use of ratios of QALYs lost
to life years lost due to disease and explored how the use of the QALY burden of disease is preferable to inform estimates of the threshold. We here present only the results for he QALY burden approach.

In Table C.74, deaths and YLL from ONS (2008 to 2010 mortality data) compare to those from GBD. The factors used to adjust GBD information are reported in columns 4 and 7.

Table C.74: Comparing deaths and YLL from ONS and GBD. (2008)

|  |  | Excess deaths ONS [1] | deaths |  |  | YLL |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | All deaths ONS [2] | All deaths GBD* <br> [3] | adjustment factor (deaths) [4] | Net estimates ONS [5] | Total YLL GBD* <br> [6] | adjustment factor ( YLL ) (71) |
| 1 | Infectious diseases |  | 2,934 | 5,262 | 1,408 | 3.737 | 38,794 | 25,142 | 1,543 |
| 2 | Cancer | 93,917 | 131,946 | 140,124 | 0.942 | 1,322,166 | 1,932,637 | 0.684 |
| 4 | Endocrine | 3,663 | 6,762 | 7,509 | 0.901 | 49,817 | 95,401 | 0.522 |
| 7 | Neurological | 6,642 | 16,771 | 12,854 | 1.305 | 90,069 | 164,796 | 0.547 |
| 10 | Circulatory | 74,217 | 151,443 | 178,454 | 0.849 | 771,038 | 1,750,608 | 0.440 |
| 11 | Respiratory | 8,432 | 64,449 | 67,441 | 0.956 | 77,434 | 594,529 | 0.130 |
| 13 | Gastro-intestinal | 15,049 | 23,897 | 28,329 | 0.844 | 225,254 | 396,829 | 0.568 |
| 17 | Genito-urinary | 1,978 | 11,345 | 8,606 | 1.318 | 16,508 | 77,338 | 0.213 |
| 18+19 | Maternity \& neonates | 265 | 265 | 2,211 | 0.120 | 19,781 | 149,868 | 0.132 |
|  | Total | 207,097 | 412,140 | 446,936 | 0.92 | 2,610,861 | 5,187,148 | 0.50 |

The threshold cost per QALY based on burden associated with one year of disease derived from GBD are summarised in Table C. 75 and detailed in Table C.76.

Table C.75: Summary of the cost per QALY threshold (2008)

|  | $\begin{gathered} 2006 \\ {[1]} \\ \hline \end{gathered}$ | $2008$ <br> [2] |
| :---: | :---: | :---: |
| big 4 PBC's | £3,036 | £4,872 |
| 11 PBCs (with mortality) | £,5,128 | £8,308 |
| All 23 PBCs | £,15,701 | 618,317 |
| * Preferred analysis |  |  |

Table C.76: Breakdown of the cost per QALY threshold (2008)

| PBC PBC description | Change in spend, $£ \mathrm{~m}$ [1] |  | Y burden and MEPs) Cost per QALY gained, £ [5] |
| :---: | :---: | :---: | :---: |
| 2 Cancer | £25 | 1496 | £16,997 |
| 10 Circulatory problems | £43 | 6127 | £7,038 |
| 11 Respiratory problems | $£ 26$ | 13032 | £1,998 |
| 13 Gastro-intestinal problems <br> Big 4 | £18 | 2494 | $\begin{aligned} & £ 7,293 \\ & \quad \AA 4,872 \\ & \hline \end{aligned}$ |
| 1 Infectious diseases | $£ 19$ | 891 | £20,829 |
| 4 Endocrine problems | $£ 11$ | 3442 | £3,124 |
| 7 Neurotogical problems | $£ 34$ | 6198 | £5,480 |
| 17 Genito-urinary problems | £26 | 601 | £43,813 |
| 16 Trauma \& injuries* | $£ 44$ | 0 | NA |
| 18+19 Maternity \& neonates* | £39 | 13 | £2,969,208 |
| (2irst 11 PBC's |  |  | £,8,308 |
| 3 Disorders of Blood | $£ 23$ | 808 | £28,305 |
| ) 5 Mental Health Disorders | $£ 198$ | 3983 | £49,835 |
| 6 Learning Disability | £12 | 146 | £78,854 |
| 8 Problems of Vision | £22 | 281 | £76,850 |
| 9 Problems of Hearing | $£ 10$ | 509 | £19,070 |
| 12 Dental problems | $£ 32$ | 574 | $\AA 55,916$ |
| 14 Skin | £22 | 125 | £,174,775 |
| 15 Musculo skeletal system | $£ 40$ | 1990 | $£ 20,254$ |
| 20 Poisoning and adverse effects | $£ 10$ | 63 | £163,766 |
| 21 Healthy Individuals | $£ 39$ | 26 | £1,483,012 |
| 22 Social Care Needs | £33 | 0 | NA |
| 23 Other | £.58 | 0 | NA |
| All (23 PBCs) |  |  | £18,317 |

The results of the three sequential steps of analysis are summarised in Table C.77, for this year of analysis. They include: i) the cost per life year (column 1) based on the methods of analysis outlined in Section C.2.1; ii) the cost per life year adjusted for quality of life (column 2) based on the methods of analysis outlined in Section C.2.2; and iii) the cost per QALY (column 3) based on the methods of analysis outlined in Section C.2.3. These estimates, in column 3, take account of the likely effects of changes in expenditure on quality of life during disease as well as the effects associated with mortality and life years; making best use of available information, while the assumptions required appear more reasonable than the other alternatives available. For this reason these estimates remain our central or best estimates forall the waves of expenditure and mortality data.

Table C.77: Summary of cost per QALY threshold estimates (2008)


Recall that the estimate of $f, 3,308$ per QALY (line 2) is restricted to the effects of changes in expenditure in the 11PBCs where outcome elasticities can be estimated. However, these PBCs only account for a proportion of a change in overall expenditure (approximately $35 \%$, see Table C. 80 below). As was explained in Section 6.2.3 the QALY threshold of $£ 18,317$ (column 3, line 3) uses the estimated proportionate effects of expenditure on the QALY burden of disease in the 11PBCs were used as a surrogate for proportionate effects in the others, (i.e., assuming that the effects that can be observed will be similar to those that cannot) and represents our central or best estimate. As in previous sections, no health effects are assigned to PBC23 (General Medical Services) on the basis that any health effects of this expenditure would be recorded in the other PBCs. Although this estimate of $£ 18,317$ reflects changes in undiscounted QALYs associated with changes in expenditure, discounting the quality adjusted life year effects only increases the cost per QALY threshold to $£ 18,613$ (Table C.78). The effects of discounting are modest because: i) the health effects of a change in expenditure are restricted to one year (where no discounting is necessary); ii) most of the total QALY effect occurs in that year; iii) it is only some of the life year effects (adjusted for quality) of a change in mortality in that year that occur in future years that need to be discounted; and iv) these need to be discounted only over 4.5 years on average.

Table C.78: Summary of QALY threshold, discounted (2008).

|  | 2008-2010 <br> discounted |  |
| :--- | :---: | :---: |
|  | Best estimate |  |
| big 4 PBC's | $£ 4,998$ | $[1]$ |
| 11 PBCs | $£ 8,467$ | $[2]$ |
| All 23 PBCs | $£, 18,613$ | $[3]$ |

${ }^{2}$ Only quality adjusted net YLL were discounted, and thus QALYs associated with gains in QoL during disease were not. The discounting factor has been calculated by applying a $3.5 \%$ discount rate to each year of life lost in the PBCs - the estimate of years of life lost used was the implied YLL per death averted in each PBC (in Table C. 18 column 4 and reproduced in Tables 28 column 2 and Table 35 column 2). This discounting factor was applied to net YLLs, before applying the outcome elasticity to calculate YLL averted.

As in previous Sections of this Appendix, the upper and lower bounds for the cost per QALY thresholds in column 3 are based on making the necessary assumptions about duration of health effects and how long a death might be averted optimistic (providing the lower bound for the threshold) erconservative (an upper bound for the threshold). The lower bound (lines 4 to 6 ) is based on assuming that health effects are not restricted to one year but apply to the whole of the remaining diseaseduration of the population at risk in PBCs during one year. Although this combines optimistic assumptions, it is possible that at least some part of a change in expenditure may prevent disease so will have an impact on populations that are incident to PBCs in the future. Such effects are not capfured in any of the estimates presented in this report so all are conservative in this respect. The upper bound (lines 7 to 9 ) is based on the combination of assuming that health effects are restricted to one year for the population currently at risk and that any death averted is only averted for 2 years (see Section C.2.1.5).

As previously, the estimated QALY effects associated with @ach PBC can be decomposed into that part due to life year effects adjusted for quality and that part associated with effects on quality during disease (Table C.79). Those PBCs for which mortality is the major concern have a much greater share of total QALY effects associated with avoidance of premature death (e.g., PBC2 and PBC10) compared to those where quality of life is the major concern (e.g., PBC 7).

Table C.79: Decomposing estimated QALK effects by PBC (2008)


## C.3.4. Which PBCs matter most?

Table C.80: Impact of each PBC on the overall cost per QALY threshold (2008)

${ }^{*}$ Calculated using the effect on the threshold of a $10 \%$ increase (or decrease) in QALY change of the PBC.

## C.3.5. How uncertain are the estimates?

In Section 2.2.5, the impact of uncertainty over the spend and outcome elasticities on estimates of the cost per QALY threshold has been illustrated and interpreted in detail. We here repeat this analysis using expenditure data from 2008/09 and mortality data from 2008 to 2010. Figure C. 8 shows the histogram of threshold values from the Monte Carlo simulation (where each random sample from the simulation represents one possible realisation of the overall threshold), and Figure C. 9 shows the cumulative probability density function for a cost per QALY threshold based on the 11 PBC with estimated outcome elasticities and for all 23 PBCs.


Figure C.8: Histogram of simulation of undiscounted threshold (all 23 PBCs) (2008)


Figure C.9: Cumulative probability density function for the cost per QAIY threshold (2008)

Table C.81: Uncertainty over the QALY threshold (2008).

|  | $\begin{gathered} \text { 11PBCs } \\ {[1]} \end{gathered}$ | $\begin{gathered} \mathrm{All}(23 \mathrm{PBCs} \\ [2]) \end{gathered}$ |
| :---: | :---: | :---: |
| Best estimate (deterministic) | £,8,308 | 2,18,317 |
| Mean estimate (from the simulations) | £,8,330 | ¢,18,310 |
| Threshold value at the probability of (from the simulations): |  |  |
| 2.5\% | £6,329 | £.12,232 |
| 5.0\% | £6,670 | £12,907 |
| 50.0\% | \& 8,266 | £18,192 |
| 95.0\% | 6012,272 | £32,845 |
| 97.5\% | 6,13,602 | £ 38,099 |
| Probability (from the simulations)of the threshold being smaller than: |  |  |
| £5,000 per QALY | 0\% | 0\% |
| £6,000 per QALY | 1\% | 0\% |
| £7,000 per QALY | 12\% | 0\% |
| $£ 8,000$ per QALY $\bigcirc$ | 42\% | $0 \%$ |
| $\AA 9,000$ per QALY | 67\% | 0\% |
| £10,000 per QALY | 82\% | 0\% |
| £15,000 per QALY | 99\% | 21\% |
| $\chi^{20,000}$ per QALX | 100\% | 64\% |
| £25,000 per QALY | 100\% | 85\% |
| £30,000 per QALY | 100\% | 92\% |
| £35,000 per QALY | 100\% | 96\% |
| $\AA 40,000$ per QALY | 100\% | 98\% |
| ¢ 45,000 per QALY | 100\% | 98\% |
| 6,50,000 per QALY | 100\% | 99\% |

## C. 4 Re-estimating the cost per QALY threshold using 2007 expenditure data

The same methods of analysis were applied to the econometric analysis of the 2007/08 expenditure and 2007 to 2009 mortality data (see Section B10 in Appendix B). Given the detailed reporting of the methods and interpretation of the analyses for other expenditure years (see Sections C. 2 and C.3), we will here only present the necessary Tables of results.

Table C.82: Outcome and spend elasticties (2007)


Table C.83. Number of deaths above LE in 2007/8/9, by PBC

| PBC | $\begin{aligned} & <L E \\ & 2007 \end{aligned}$ | $\begin{aligned} & >\text { LE } \\ & 2007 \end{aligned}$ | $\begin{aligned} & <\mathrm{LE} \\ & 2008 \end{aligned}$ | $\begin{aligned} & >\text { LE } \\ & 2008 \end{aligned}$ | $\begin{aligned} & <\mathrm{LE} \\ & 2009 \end{aligned}$ | $\begin{aligned} & >\text { LE } \\ & 2009 \end{aligned}$ | Annual N deaths $<$ LE | Annual N deaths > LE |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 Infectious diseases | 3,906 | 3,731 | 3,404 | 2,588 | 3,042 | 2,192 | 3,451 | 2,837 |
| $2 \quad$ Cancer | 95,385 | 35,401 | 94,814 | 37,088 | 94,218 | 37,209 | 94,806 | 36,566 |
| 4 Endocrine | 3,970 | 2,747 | 4,031 | 2,879 | 3,832 | 2,828 | 3,944 | 2,818 |
| 7 Neurological | 8,852 | 6,494 | 9,632 | 6,865 | 9,439 | 6,945 | 9,308 | 6,768 |
| 10 Girculatory | 80,687 | 78,404 | 80,834 | 76,352 | 75,993 | 73,397 | 79,172 | 76,051 |
| 11 Respiratory | 29,571 | 35,029 | 32,059 | 35,204 | 29,890 | 33,326 | 30,507 | 34,520 |
| 13 Gastro-intestinal | 15,667 | 8,367 | 15,937 | 8,267 | 15,354 | 8,168 | 15,653 | 8,267 |
| 17 Genito-urinary | 4,077 | 6,553 | 4,468 | 6,670 | 4,375 | 6,903 | 4,307 | 6,709 |
| 18+19) Maternity \& neonates | 216 | 0 | 267 | 0 | 281 | 1 | 255 | 0 |

Table C.84: Net YLL using LE of the PBC (2007)

|  |  | Average2007-2009 |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| PBC |  | LE of | LE of | Deaths |  |  |  |  |
| Males | Females | $<$ LE | $>$ LE | YLL | YLG | Net YLL |  |  |
| 1 | Infectious diseases | 79.6 | 83.6 | 3,280 | 3,008 | 57,715 | 19,085 | 38,629 |
| 2 | Cancer | 83.0 | 84.7 | 100,810 | 30,561 | $1,464,726$ | 129,810 | $1,334,916$ |
| 4 | Endocrine | 81.0 | 84.7 | 4,004 | 2,759 | 66,575 | 15,386 | 51,189 |
| 7 | Neurological | 79.6 | 83.3 | 8,719 | 7,357 | 135,760 | 44,925 | 90,835 |
| 10 | Circulatory | 83.0 | 86.5 | 92,729 | 62,494 | $1,069,632$ | 276,368 | 793,264 |
| 11 | Respiratory | 80.3 | 84.0 | 29,668 | 35,359 | 304,168 | 230,245 | 73,922 |
| 13 | Gastro-intestinal | 80.6 | 84.5 | 15,640 | 8,280 | 271,092 | 45,500 | 225,593 |
| 17 | Genito-urinary | 83.5 | 85.6 | 5,008 | 6,007 | 47,656 | 30,931 | 16,725 |
| $18+19$ | Maternity \& neonates | 78.7 | 83.1 | 255 | 0 | 18,844 | 1 | 18,843 |

Table C.85: Comparing deaths and YLL from ONS and GBD (2007).

|  |  | Excess <br> deaths ONS |  deaths <br> All All deaths <br> deaths GBD* <br> ONS  |  | adjustment factor (deaths) |  |  | adjustment factor (YLL) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  |  |  |  |
| 1 | Infectious diseases | 2,925 | 6,288 | 1,408 | 4.47 | 38,629 | 25,142 | 1.54 |
| 2 | Cancer | 94,827 | 131,372 | 140,124 | 0.94 | 1,334,916 | 1,932,637 | 0.69 |
| 4 | Endocrine | 3,765 | 6,762 | 7,509 | 0.90 | 51,189 | 95,401 | 0.54 |
| 7 | Neurological | 6,692 | 16,076 | 12,854 | 1.25 | 90,835 | 164,796 | 0.55 |
| 10 | Circulatory | 76,322 | 155,223 | 178,454 | 0.87 | 793,264 | 1,750,608 | 0.45 |
| 11 | Respiratory | 8,034 | 65,027 | 67,441 | 0.96 | 73,922 | 594,529 | 0.12 |
| 13 | Gastro-intestinal | 15,064 | 23,920 | 28,329 | 0.84 | 225,593 | 396,829 | 0.57 |
| 17 | Genito-urinary | 2,005 | 11,016 | 8,606 | 1.28 | 16,725 | 77,338 | 0.22 |
| 18+19 | Maternity \& neonates | 255 | 255 | 2,211) | 0.12 | 18,843 | 149,868 | 0.13 |
|  | Total | 209,890 | 415,939 | 446,936 | 0.93 | 2,643,916 | 5,187,148 | 0.51 |

Table C.86: Summary of the cost per QALY threshold (2007)

|  | $\begin{gathered} 2006 \\ {[1]} \\ \hline \end{gathered}$ | $\begin{gathered} 2007 \\ {[2]} \end{gathered}$ | $\begin{gathered} 2008 \\ {[3]} \end{gathered}$ |
| :---: | :---: | :---: | :---: |
| big 4 PBC's | £3,036 | ¢ 4,549 | £4,872 |
| 11 PBCs (with mortality) | £,5,128 | $£ 8,8,513$ | £,8,308 |
| All 23 PBCs | £.15,70 | £.18,624 | £.18,317 |

Table 87: Breakdown of the cost per QALY threshold (2007)

| PBC | PBC description | Change in spend, $£ \mathrm{~m}$ [1] | Q (HoD <br> Change in QALY <br> [4] | Y burden $R$ and MEPs) <br> Cost per QALY gained, $£$ <br> [5] |
| :---: | :---: | :---: | :---: | :---: |
| 2 | Cancer | £41 | 3041 | £13,384 |
| 10 | Circulatory problems | £19 | 2756 | £6,724 |
| 11 | Respiratory problems | £18 | 13152 | £1,398 |
| 13 | Gastro-intestinal problems | £24 | 3316 | £7,137 |
| $\operatorname{Big} 4$ |  |  |  | ¢,4,549 |
| 1 | Infectious diseases | £16 | 1035 | £15,530 |
| 4 | Endocrine problems | $\AA 5$ | 910 | £5,796 |
| 7 | Neurological problems | £33 | 5111 | £6,409 |
| 17 | Genito-urinary problems | £35 | 974 | £ 35,449 |
| 16 | Trauma \& injuries* | £49 | 0 | NA |
| 18+19 | Maternity \& neonates* <br> First 11 PBC's | £19 | 6 | $\begin{array}{r} £^{3,250,386} \\ £ 8,513 \end{array}$ |
| 3 | Disorders of Blood | £28 | 878 | £ 32,310 |
| 5 | Mental Health Disorders | £165 | 4331 | $\AA 38,145$ |
| 6 | Learning Disability | £19 | 159 | £,119,676 |
| 8 | Problems of Vision | $£ 29$ | 306 | £ ¢93,716 |
| 9 | Problems of Hearing | ${ }_{6} 7$ | 554 | £11,960 |
| 12 | Dental problems | £20 | 624 | $£ 32,214$ |
| 14 | Skin | $£ 10$ | 136 | £76,382 |
| 15 | Musculo skeletal system | £49 | 2164 | £22,545 |
| 20 | Poisoning and adverse effects | $\AA 8$ | 68 | ¢123,247 |
| 21 | Healthy Individuals | £25 | 29 | \& 858,150 |
| 22 | Social Care Needs | £.52 | 0 | NA |
| 23 | Other | £66 | 0 | $1 \bigcirc \mathrm{NA}$ |
| All (23 PBCs) |  |  | $\bigcirc 1.18,624$ |  |

Table C.88: Decomposing estimated QALY effects by PBC (2007)


Table C.89: Summary of cost per QALY threshold estimates (2007)

| QoL associated with life extension: QoL during disease: | norm <br> Based on burden |  |
| :---: | :---: | :---: |
| Effect of expenditure on mortality: <br> YLL per death averted: QALYs per death averted: | $\begin{gathered} \text { Best estimate } \\ 1 \text { year } \\ \sim 4.6 \mathrm{YLL} \\ \sim 12.7 \text { QALY } \end{gathered}$ |  |
| big 4 PBC's <br> 11 PBCs (with mortality) <br> All 23 PBCs | $\begin{gathered} £,{ }^{\infty}, 549 \\ £, 8,513 \\ £, 18,624 \end{gathered}$ | [1] $[2]$ $[3]$ |
| Effect of expenditure on mortality: <br> YLL per death averted: QALYs per death averted: | Lower bound <br> Remainder of disease duration $\begin{gathered} \sim 4.6 \text { YLL } \\ \sim 12.7 \text { QALY } \end{gathered}$ |  |
| big 4 PBC's <br> 11 PBCs (with mortality) <br> All 23 PBCs | $\begin{aligned} & £ 1,116 \\ & £ 1,361 \\ & £, 3,247 \end{aligned}$ | $[4]$ $[5]$ $[6]$ |
| Effect of expenditure on mortality: <br> YLL per death averted: QALYs per death averted: | $\begin{gathered} \text { Upper bound } \\ 1 \text { year } \\ 2 Y \mathrm{LL} \\ \sim 5.6 \text { QALY } \end{gathered}$ |  |
| big 4 PBC's <br> 11 PBCs (with mortality) <br> All 23 PBCs | $\begin{aligned} & £ 10,965 \\ & £ 20,517 \\ & £ 44,889 \end{aligned}$ | [7] $[8]$ $[9]$ |

Table C.90: Summary of QALY threshold, discounted (2007)

|  | $\begin{gathered} 2006-2008 \\ {[1]} \\ \hline \end{gathered}$ | $2007-2009$ <br> [2] | $\begin{gathered} 2008-2010 \\ {[3]} \\ \hline \end{gathered}$ |  |
| :---: | :---: | :---: | :---: | :---: |
| big 4 PBC's | £3,036 | $£ 4,690 \sim$ | £4,998 | [1] |
| 11 PBCs | £,5,218 | £,8,718 | £,8,467 | [2] |
| All 23 PBCs | £,15,701 | £18,987 | ¢,18,613 | [3] |

## Appendix C: Addendum 1

## DATA SOURCES

## Contents

A. General Practice Research Database (GPRD)
B. Global Burden of Disease (GBD)
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F. Hospital Episode Statistics (HES)
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H. Tables of prevalence distribution within PBCs

## References

## A. General Practice Research Database (GPRD)

GPRD contains over 3 million active patient records drawn from approximately 400 primary care practices in the UK. The Medicine Control Agency manages the dataset. The database has clinical and prescription data and can provide information to sûpport pharmaco-vigilance (indication, utilization, and risk/benefit profiles of drugs) and formal pharmaco-epidemiologic studies, including information on demographics, medical symptoms, therapy (medicines, vaccines, devices), and treatment outcomes.

As of 29th March 2012 GPRD has become the Clinical Practice Research Database (CPRD), an expanded dataset that represents 'The All England Data and Interventional Research Service'. GPRD was approached to provide information on the prevalence of disease by ICD-10 disease code. A sample set of data was analysed by researchers at Pharmatelligence ${ }^{19}$ who were tasked with extracting data on prevalence of each disease state by ICD-10.

We were provided with access to data comprising of 22,313,086 rows/patient-ICD10 events ( 3 digit $)^{20}$ representing $4,229,910$ patients with data on new diagnosis of diseases observed between 1 Jan 2006 and 24 June 2011. Multiple events per patient are thus possible, and all patients are active in the dataset, i.e. patients had at least one new diagnosis in the period of interest. Newly diagnosed (incident) events were defined using a wash-in period of 24 months (or from registration to index date if lower than 24 months). The sample contains 1873 unique ICD codes in the dataset. 70 ICD codes account for $50 \%$ of the total number of events, 166 for $75 \%$ and 306 for $90 \%$.

[^84]Diagnoses are collected in GPRD using Read codes. These were mapped into three-character ICD-10 codes. Cross-mappings from Read V2 and Read V3 to ICD-10 were used in order to maximize the number of GPRD Read and ICD-10 codes included ( $33.2 \%$ of Read codes; $99.7 \%$ of ICD codes $)^{21}$.

Unfortunately due to the short collection period of GPRD it was not possible to directly observe prevalence only incidence over a period. Attempts were made to elicit prevalence estimate through observed incidence data from GPRD coupled with clinical expertise on expected disease duration (provided by Dr. Charlotte Haylock, Hull and East Yorkshire Hospitals NHS Trust). Our approach classified expected duration for all ICD-10 diseases by 3-digit code into one of five duration 'buckets'22? However, the limitations of the data were deemed too extensive to provide sufficient accuracy of estimates to represent a stronger estimate of prevalence than provide by GBD.

## B. Global Burden of Disease (GBD)

The WHO GBD project draws on a wide range of data sources to quantify globat and regional effects of diseases, injuries and risk factors on population health. We were provided with access to the beta version of the WHO's National Burden of Disease (NBD) toolkit for the United Kingdom which represents a set of metrics on World Health Organisation (WHO) prior estimates of mortality and burden of disease for WHO Member states for 2004 (based on the Global Burden of Disease: 2004 update[9]) ${ }^{23}$.

The metrics of interest to our analysis included disease incidence, prevalence, duration and mortality. These metrics were provided by U-code disease code which were mapped to ICD-10 using direct WHO mapping algorithms[10]. In addition, in many cases each U-code was sub-divided by disease sequela which represent disease sub-categories of each U-cøde[10]. As an individual may be represented in multiple sequela in a single U-code to avoid double counting in the event of multiple sequela in a given Ucode our analysis uses prevalence estimates based on the sequela with the largest prevalent population.

Our analysis uses two forms of prevalence data, "point prevalence" and "annual prevalence". "Point Prevalence" represents the instantaneous prevalence of a disease whereas "annual prevalence" represents the extent of the prevalence population oyer a given year. To calculate "annual prevalence" incidence of a disease was multiplied by expected disease duration rounded up to the nearest year.

All data was provided by age, given for both genders in fixed age buckets (either eight or nineteen buckets depending on the data of interest), as a result it was necessary to assume the relevant population could be represented by the mid-point of that bucket for the relevant metric.

## C. Health Survey for England (HSE)

The Health Survey for England (HSE) comprises a series of annual surveys beginning in 1991. This survey is how commissioned and published by The NHS Information Centre. It is designed to provide

[^85]regular information on various aspects of the nation's health. All surveys have covered the adult population aged 16 and over living in private households in England. ${ }^{24}$

In order to define the quality of life norms for the population of the UK required for the analysis detailed in section 4.3.1 data from six Health Surveys for England (1996, 2003-2006 and 2008) were pooled. Selfreported health status and EQ-5D data were extracted and used to generate mean health state utility values for the 'normal' population.

Surveys are not completed for people age under 16; as a result we have assumed that all persons age 0 to 15 have the same quality of life norms as a person age 16 . In addition the number of surveys recorded for persons over 91 years of age is relatively small, as a result all persons over 91 are assumed to have the same quality of life norm as a person age 91 . The quality of life norms for each age and gender are shown in figure 1 in section 4.3.1.

## D. Health Outcome Data Repository (HODaR)

HODaR represents a supplement of routine clinically coded data from the Cardiff and Vale NHS Hospitals Trust, UK, with survey data covering socio-demographic characteristics, QoL, utility, and resource use information[4]. HODaR data was collected for subjects treated at Cardiff and Vale NHS hospital from 2002 to 2004. Inpatients were surveyed 6 weeks post-discharge whilst outpatients are handed a survey package when they attend. More than 30,000 observations (aged above 18) are available relating to approximately 2,000 diagnoses of disease by ICD-10

We used HODaR to estimate Health Related Quality of Life (HRQoL) by ICD-10 diagnoses codes and age using EQ-5D. If data on a patient was provided with multiple diagnoses the primary condition was used.

## E. Medical Expenditure Panel Survey (MEPS)

MEPS is a national representative survey of the US civilian non-institutionalised population, collecting information on health care utilisation which began in 1996.[5] EQ-5D was employed to measure HRQoL of the population in years 2000 to 2002. There are about 38,000 adults (aged above 18) completing EQ5D relating to 700 ICD-10 diagnoses. MEPS consists of a household component and an insurance component, both aimed atidentifying the medical usage of individuals as well as how they are funded, their cost, and the scope and breadth of health insurance held and available.

As with HODaR, MEPS allowed us to estimate the HRQoL by ICD-10 code and age. If data on a patient was provided with multiple conditions the primary condition was used.

## F. Hospital Episode Statistics (HES)

HES represents a collection of data with details on all admissions to the NHS hospital in England. It contains admitted patient care data from 1989 onwards, with more than 12 million new records added each year, and outpatient attendance data from 2003 onwards, with more than 40 million new records added each year.

Expenditure by ICD-10 codes and PCT was used to estimate the contribution to variance of each PBC. This was done by calculating the contribution of that ICD to the variance in expenditure between PCTs

[^86]within a PBC (total costs allocated to individual ICDs were divided by the number of patients using services in the PCT). For our analysis we make use of HES data on the year 2007/08.

## G. Patient Reported Outcome Measures (PROMs)

Introduced in 2009, the English NHS Patient Reported Outcomes (PROMs) programme routinely collects self-reported health status of patients receiving surgery for four elective procedures: knee and hip replacement, groin hernia repair, and varicose vein surgery. Patients are invited to complete a questionnaire prior to surgery, and again six (or three) months after surgery.[11] Differences in their selfreported health status are used to explore differences between provider performance in improving patient health.[12] The data that are collected include both condition specific questions (the Oxford Hip Score, Oxford Knee Score and the Aberdeen Varicose Vein score; no condition specific instrument is available for hernia) as well as the generic instrument, the EQ-5D (both the EQ-5D profile, and the patient's global assessment of their health, the EQ-VAS - see.[13] All NHS patients receiving these surgical procedures are invited to complete the PROMs questionnaires - in practice, for a variety of reasons, some patients do not participate, or complete only the pre-surgery, or the post-surgery, questionnaire - so the data do not cover $100 \%$ of patients. However, good coverage rates have been achieved for example, the response rate from hip surgery patients to April 2012 was $78 \%$ for the pre-surgery questionnaire, and $81 \%$ on the post-surgery questionnaire.[14]

Patient-level data from the PROMs programme are freely available to download in anonymised form. Those data can also be linked to further information in the HES database, via requests to the NHS Information Centre. Standardised reports on the PROMs data, including the average (case-mix adjusted) performance of providers, is regularly published by the Information Centre, currently on a quarterly basis.

There are plans to extend the PROMs programme in the future, in keeping with the Government's NHS Outcomes Framework, and a number of pilot studies have been commissioned by the Department of Health in order to inform the roll out to other XHS services. There is currently work underway or being planned around the potential use of PROMs in a vide range of long term conditions; primary care; in cancer survivorship; cardiovascular services; muscular skeletal; and cosmetic surgery.
G. Tables of prevalence distribution within PBCs

Table C1.1: Distribution of PBC1 prevalence by age, gender and contributing ICD alongside proportion of prevalence patients and contribution to variance of each ICD


Table C1.2: Distribution of PBC2 prevalence by age, gender and contributing ICD alongside proportion of prevalence patients and contribution to variance of each ICD


Table C1.3: Distribution of PBC3 prevalence by age, gender and contributing ICD alongside proportion of prevalence patients and contribution to variance of each ICD


Table C1.4: Distribution of PBC4 prevalence by age, gender and contributing ICD alongside proportion of prevalence patients and contribution to variance of each ICD


Table C1.5: Distribution of PBC5 prevalence by age, gender and contributing ICD alongside proportion of prevalence patients and contribution to variance of each ICD


Table C1.6: Distribution of PBC6 prevalence by age, gender and contributing ICD alongside proportion of prevalence patients and contribution to variance of each ICD


Table C1.7: Distribution of PBC7 prevalence by age, gender and contributing ICD alongside proportion of prevalence patients and contribution to

A- Episodic and paroxysmal disorders (G40-G47)
B- Extrapyramidal and movement disorders (G20-G26)

C- Other degenerative diseases of the nervous system (G30-G32)
D- Other disorders of the nervous system (G90-G99)
E- Nerve, nerve root and plexus disorders (G50-G59)
F- Demyelinating diseases of the central nervous system (G35-G37)
G- Polyneuropathies and other disorders of the peripheral nervous system (G60-G64)

G83)
variance of each ICD



X - Symptoms and signs nervous and musculoskeletal systems (R25-R29)
Y- Symptoms and signs urinary system (R30R39)
Z- Symptoms and signs cognition, perception, emotional state and behaviour (R40-R46)
AA- Symptoms and signs speech and voice (R47-R49)
AB- General symptoms and signs (R50-R69) AC- Abnormal clinical and laboratory findings, not elsewhere classified (R70-R99) AD- Persons with potential health hazards related to family and personal history and certain conditions influencing health status (Z80-Z99)

Table C1.8: Distribution of PBC8 prevalence by age, gender and contributing ICD alongside proportion of prevalence patients and contribution to variance of each ICD


Table C1.9: Distribution of PBC9 prevalence by age, gender and contributing ICD alongside proportion of prevalence patients and contribution to variance of each ICD


Table C1.10: Distribution of PBC10 prevalence by age, gender and contributing ICD alongside proportion of prevalence patients and contribution to variance of each ICD


Table C1.11: Distribution of PBC11 prevalence by age, gender and contributing ICD alongside proportion of prevalence patients and contribution to variance of each ICD


Table C1.12: Distribution of PBC12 prevalence by age, gender and contributing ICD alongside proportion of prevalence patients and contribution to variance of each ICD


Table C1.13: Distribution of PBC13 prevalence by age, gender and contributing ICD alongside proportion of prevalence patients and contribution to variance of each ICD


Table C1.14: Distribution of PBC14 prevalence by age, gender and contributing ICD alongside proportion of prevalence patients and contribution to variance of each ICD


Table C1.15: Distribution of PBC15 prevalence by age, gender and contributing ICD alongside proportion of prevalence patients and contribution to variance of each ICD


Table C1.16: Distribution of PBC17 prevalence by age, gender and contributing ICD alongside proportion of prevalence patients and contribution to variance of each ICD


Table C1.17: Distribution of PBC18\&19 prevalence by age, gender and contributing ICD alongside proportion of prevalence patients and contribution to variance of each ICD


Table C1.18: Distribution of PBC20 prevalence by age, gender and contributing ICD alongside proportion of prevalence patients and contribution to variance of each ICD


Table C1.19: Distribution of PBC21 prevalence by age, gender and contributing ICD alongside proportion of prevalence patients and contribution to variance of each ICD


## Appendix C: Addendum 2

## THE ROLE OF DATA ON LOCAL NHS DECISIONS

## Contents

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B.5. NHS Right Care
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B.7. Annual reporting and strategic commissioning plans
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## A. The role of local data in this study

The aim of this research project, noted in Chapter 1, is to develop and demonstrate methods for threshold estimation that make best use of routinely available NHS data. The principal focus of that methodological development, as reflected in the main body of this report, has been the use of econometric methods to exploit observed variations in spending and health outcomes between PCTs, at the programme budget level of aggregation.

However, we also aimed to investigate, as a complementary element of the project, the extent to which there may be other, more disaggregated sources of evidence on investment and disinvestment decisions made by local NHS organisations which might inform our analysis.

Specifically, we set out to (a) identify and evaluate what data might be routinely available from local N1S organisations with respect to their decisions to increase or decrease spending on specific services, and (b) consider whether and how such evidence might contribute directly toward the quantitative estimates of the threshold - for example, by providing more granular, contextual information on spending decisions that might assist in the interpretation of model estimates. For example, we wished to explore whether there were any routinely collected data from local NHS organisations that could tell us something about which ICDs within a given PBC might be the focus of investment and disinvestment.

The work which was undertaken was therefore focused on the potential use of local data alongside the econometric analysis - rather than their potential use as an alternative means of identifying the marginal cost of a QALY in the NHS[15].

## B. Sources of publicly available data on PCT investment and disinvestment

To help us identify possible sources of data on NHS spending decisions, we began by consulting a number of experts within the NHS, identified for us by our collaborator, Professor David Parkin (Chief Economist at NHS South East Coast). These included Directors of Finance, Commissioning and Public Health. Those discussions helped direct us to a number of initiatives which involved the development of tools or evidence to inform resource allocation decisions, and helped to identify types and sources of documents published by PCTs that potentially contained relevant information on spending decisions. We then undertook a search for publicly available documents, in each case identifying what was available, and assessing its potential relevance for the purposes of this work outlined above. In evaluating each data source, the key considerations were:
(a) Whether the data were routinely collected: Routinely-collected data are preferred, as our overarching aim is to develop a set of methods to estimate the threshold, which can be readily updated from data routinely generated by the NHS.
(b) Whether the data were in the public domain: Published data are preferred to data that can only be obtained on request, because this would increase the cost and effort required in obtaining data from all relevant organisations.
(c) Whether the data were collected and reported in a systematic and consistent manner that would facilitate comparisons between PCTs, and with sufficient detail to enable us to link spending decisions to specific programme budgets or ICDs. This aspect of the work was undertaken during 2010.

The following were identified as potential sources of data:

## B.1. Programme Budgeting tools - quadrant analysis - spend outcome tool (SPOT)

Data are available for three years, 06/07, 07/08 and 08/0925 under the Spend Outcome Tool (SPOT) which is available to download. Expenditure data are organised by Programme Budget Category only,

[^87]with no lower level of disaggregation. The data shows, for each PCT, the spend per head this year, the Zscore of that spend, and the PCTs national ranking based on their Z-score.

Outcomes data have also been captured, with different outcome measures within each PB category. Again, for each outcome there is a related Z-score and the PCTs national ranking based on that Z-score.

The tool enables users to see graphically how one PCT compares to others nationally, by SHA and by those PCTs similar to it by cluster (eg. other PCTs in manufacturing towns). The quadrant analysis tool has to origin as the mean PCT for that PB category, with Z-score for both expenditure and outcome equal to zero. The $y$-axis shows outcome, and the $y$-axis expenditure, both by Z -score.

While a useful tool, this source added little to the data already used in the econometric analysis, as it does not provide any additional information on the allocation of resources within PBs.

## B.2. Lists of interventions not normally funded

Most PCTs provide information about interventions not normally funded. However, these were of limited usefulness because most of the procedures listed are those that might be expected (cosmetic surgery; tattoo removal, etc), and are not particularly informative about the marginal cost per QALY in the NHS. We did not find any information regarding whether any previously funded treatments had been added to these lists.

## B.3. Special therapeutic and cancer committees

These are regionally based (not PCT or SHA) specialised committees that make decisions regarding spending on new cancer medicines and other special therapeutic areas. While such decisions would be potentially of direct relevance, we were unable to find any public documentation on their processes or decision outcomes.

## B.4. Quality Innovation Productivity and Prevention (QIPP) published data on efficiency savings in the NHS

Introduced in 2009, QIPP addresses the quality and productivity challenge faced by the NHS. Developed by NICE, the Cochrane Quality and Productivity (QP) topics identify areas where resources could be significantly reduced or stopped completely without reducing the quality of NHS care, releasing cash and/or resources to other areas in the NHS. Each Cochrane topic has been established from systematic reviews undertaken by reviewers at the Cochrane Collaboration.

Every month the Cochrane Collaboration informs NICE as to new or existing Cochrane reviews where they have found that the existing treatment options(s) are harmful or ineffective and should not be used, or where evidence is unavailable or insufficient to support widespread use of that treatment in the NHS. NICE then completes an assessment of a Cochrane topic, to evaluate the efficiency savings that are likely against the QPP criteria of likely ease of implementation, impact on productivity, and on the quality of care.

Savings per 100,000 patients are calculated, and then efficiency gains per PCT can be calculated. Once a topic has been accepted as best practise, users (PCTs) are encourage to submit their experience of implementing the changes, and the users achieving the best efficiency gains become QIPP examples of best practise.

The data shows which procedures are considered inefficient use of resources, although to the extent these are based on means of achieving the same or improved outcomes but with lower resources, will not be revealing of the marginal cost of producing a QALY in the NHS. Further, there is incomplete information about the extent to which PCTs actually implement these recommendations.

## B.5. NHS Right Care

This website ${ }^{26}$, has a section on the NHS Atlas of variation in health care, which seeks to reduce unwarranted variations in health care, defined as "...variation in the utilisation of health care services that cannot be explained by variation in patients or patients preferences", to increase value and improve quality.

It also provides a Third Annual Population Value Review which uses programme budgeting and marginal analysis to deliver QIPP. This provides, amongst other things, a 10 -step, structured approach for PCTs to follow to establish where investment and disinvestment decisions could be made.

Further, it provides a tool for NHS Foundation Trusts to improve efficiency via Service Line Management.

While these tools may be being used by PCTs and Foundation Trusts, it was not clear to what extent that was the case, and there is no routine data on their use by NHS organisations or the decisions that resulted from that.

## B.6. Health Investment Network - case studies of PBMA

The NHS network, Health Investment Network, was established to provide the access to the latest knowledge and tools to help commissioners optimise their investment and disinvestment decisions. It provides case studies of PCTs which have used PBMA to identify efficiency gains. This includes examples of 'spend to save' decisions e.g. where an initial investment (eg in vascular checks for men in deprived areas) could be more than outweighed by savings. Such initiatives, while important, are not useful in identifying the marginal cost per QALY in the NHS. Other case studies identify 'wish lists' (areas which PCTs prioritise for additional spending, should budgets expand) and 'hit lists' (services that might be reduced, to free up resources for more cost effective services). These case studies provide useful selected examples - but do not provide a routine or systematic reporting of such decisions across all PCTs.

## B.7. Annual operating plans and strategic commissioning plans

PCTs are required to publish, each year, operating plans and strategic commissioning plans detailing their planning for the coming year, including information on the way that PCTs have made decisions concerning resource allocation. Because these reports are published annually, we considered that they constituted the most promising source of data, as they are produced routinely, and cover all PCTs.

Contact details and websites were identified for all 142 PCTs. Strategic Commissioning Plans were obtained for an initial 70 of these. These were used to identify any information provided about programmes of care or specific services where spending was planned to be increased or decreased. Those data were extracted and recorded into a spreadsheet, along with any relevant contextual information eg relating to the process by which the decision had been made.

Our review of the data from the first 70 of these showed that there was considerable variation between the documents in terms of the level of detail and specificity about the services which were the subject of changes in spending. In many cases, the services were described in terms of broad initiatives which might have related to multiple programme budgets and ICDs. There was also variation in, and occasionally a lack of clarity about, the way in which spending changes were described: in some cases these were described in terms of absolute changes in spending; in others, as net changes, once estimates of offsetting savings elsewhere had been taken into account; and in others it was not stated.

Given those concerns, the data were considered unlikely to be useful to complement the econometric analysis, and the research team decided not to proceed with further data extraction for the remaining PCTs.

[^88]
## C. Conclusions

The context within which this element of the work took place may be relevant to note. While the NHS was not subject to the budget cuts imposed on other areas of government activity in response to the financial crisis, the NHS was required to make substantial productivity improvements within its existing budgets. This gave rise to a number of initiatives in response to the 'productivity challenge' and, generally, heightened interest in the identification of ways to improve efficiency; potential areas for disinvestment; and areas for investment which were motivated by 'spend to save'. This may have made it more likely that we would observe disinvestment decisions. The NHS was also, during the course of this project, undergoing a period of restructuring. The transition from PCTs to clinical commissioning groups, and the disestablishment of strategic health authorities, may have had an effect on the availability of data and information relating to decision making. It may also have broader implications for the availability of data in the future, given the change in administrative units.

Our review of local data sources suggested that there is very little routinely collected data on investment and disinvestment by local NHS organisations beyond the high-level aggregate data onspending by PB which are used in the econometric analysis. More disaggregated data on spending decisions about specific services could, of course, be obtained by other means - for example, by survexing PCTs, or by requesting such information from them using a Freedom of Information request. However, that would impose data collection costs and would need to be designed carefully to ensure that such efforts yielded complete and consistent information.

## Appendix C: Addendum 3

## Characterisation of the investment and disinvestment decisions in mental health: <br> depression and schizophrenia

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## Systematic review strategies

## A. Introduction

As has been highlighted in the main body of this project, it was not possible to produce an outcome equation for PBC 5: mental health problems, because no relevant mortality data was available from the NHS IC by PCT. Mental health represents a significant incidence and expenditure within the NHS. As a result we investigated the direction of bias from the exclusion of mental health problems on our estimate of the cost-effectiveness threshold. To understand this bias we examined current investment decisions in mental health. Recent investments in treatments with ICERs above the estimated threshold would suggest that not including PBC 5 more directly in our calculation may underestimated the threshold, conversely if recent investment has ICERs below the estimated threshold it would suggest that its exclusion results in an overestimated threshold. We focussed on depression and schizophrenia because of their high prevalence and contribution to variance.

## B. Method employed

To evaluate the direction of bias of the exclusion of PBC 5 we followed four steps to make the connection from the identification of the most significant ICDs of PBC 5 to considering the costeffectiveness of the investment and disinvestment decisions made in the NHS around these disease areas. The strategy was as follows:

Step 1:

- Identify the mental health ICD codes that are most influential and suitable on which to focus our analysis
- Done from number of patients and contribution to variance calculations using HES.

Step 2:


- Determine the medications or treatments used in the NHS to treat each of the significant ICDs
- There is likely to be a large cross-over in the use of treatments for mental health areas, for example antipsychotics and cognitive behavioural therapy (CBT) are both widely used.
- We made use of the NHS Choices website coupled with clinical expertise for this identification.

Step 3:

- Identify the cost-effectiveness of the current treatments and medications used in the NHS. - This identification will be done from a range of sources including: published HTAs, published guidance, TUFTs, NHS EED and Medline searches.
- This step relies heavily on the literature published, literature tends to cover historical activities many of which represent treatments of interest for this analysis. The case could (be made that historical treatments that have not been evaluated have escaped evaluation due to their apparent cost-effectiveness, and are as such unlikely to be marginal activities. Further difficulties arose in the identification of the relevant cost-effectiveness figure. Ideally it would represent the cost-per QALY relative to what would be performed if that activity was no longer available to the NHS.

Step 4:
Connecting the available literature on the cost-effectiveness to recent investment and disinvestment decisions made in the NHS.

## C. Results of analysis

## C.1. Step 1: identification of relevant ICDs

We first rank ICDs by prevalence and contribution to variance. Prevalence is estimated from HES data. The contribution to variance is calculated as the variance in expenditure across PCTs for each ICD compared to the total variance in expenditure across PCTs for all ICDs within PBC 5. The most prevalent ICD was for depressive episode (F32) at $25.07 \%$ of all ICDs within PBC 5 (Table C3.1). The ICD with the greatest contribution to variance was for schizophrenia (F20) with $45.16 \%$ (Table C3.2).

Depression (F32) and (F33) and schizophrenia (F20) have been chosen as the focus of our evaluation as they represent two of the largest mental health ICDs in terms of proportion of patients as well as proportion of variance in expenditure, as shown in Tables C3.1 and C3.2 below. In addition they represent ICDs that involve interventions by the NHS that can be more clearly defined (in contrast to, for example, unspecified dementia and mental and behavioural disorders due to the use of alcohol${ }^{27}$.

Table C3.1: table showing ranking of mental health ICDs by prevalence from HES

| ICD | Description | $\%$ of Mental <br> health <br> prevalence <br> to variance |  |
| :---: | :---: | :---: | :---: |
| F10 | Mental and behavioural disorders due to use of alcohol | 27, 84\% | 9.70\% |
| F20 | Schizophrenia | - $10.01 \%$ | 45.16\% |
| F32 | Depressive episode | $\pm 9.96 \%$ | 6.91\% |
| F31 | Bipolar affective disorder | 6.19\% | 6.38\% |
| F41 | Other anxiety disorders | 4.92\% | 0.26\% |
| F60 | Specific personality disorders | 4.33\% | 14.11\% |
| F03 | Unspecified dementia | 3.93\% | 3.29\% |
| F01 | Vascular dementia | 3.32\% | 1.58\% |
| G30 | Alzheimer disease | $3.30 \%$ | 0.84\% |
| F33 | Recurrent depressive disorder | 2.83\% | 3.68\% |

Table C3.2: table showing ranking of mental health ICDs by contribution to variance

|  |  | \% of Mental <br> health <br> prevalence | Contribution <br> to variance |
| :--- | :--- | ---: | ---: |
| ICD | Description | $\mathbf{1 0 . 0 1 \%}$ | $\mathbf{4 5 . 1 6 \%}$ |
| F20 | Schizophrenia | $4.33 \%$ | $14.11 \%$ |
| F60 | Specific personality disorders | $27.84 \%$ | $9.70 \%$ |
| F10 | Mental and behavioural disorders due to use of alcohol | $\mathbf{9 . 9 6 \%}$ | $\mathbf{6 . 9 1 \%}$ |
| F32 | Depressive episode | $6.19 \%$ | $6.38 \%$ |
| F31 | Bipolaraffective disorder | $\mathbf{2 . 8 3 \%}$ | $\mathbf{3 . 6 8 \%}$ |
| F33 | Recurrent depressive disorder | $3.93 \%$ | $3.29 \%$ |
| F03 | Unspecified dementia | $3.32 \%$ | $1.58 \%$ |
| F01 | Vascular dementia | $3.30 \%$ | $0.84 \%$ |
| G30 | Alzheimer disease | $4.92 \%$ | $0.26 \%$ |
| F41 | Other anxiety disorders |  |  |

## C.2. Step 2: determination of treatment employed

Table C3.3 provided an overview of the main treatments for depression and schizophrenia. This list of treatments was identified using the NHS Choices website ${ }^{28}$ as well as discussion with our clinical

[^89]representative for each of the respective illnesses. This list was used to inform a literature search of costeffectiveness publications.

Table C3.3: table showing treatments for schizophrenia and depression in the NHS

| ICD | Disease | Treatments |
| :---: | :---: | :---: |
| F20 | Schizophrenia | 1. Typical antipsychotics <br> 2. Atypical antipsychotics <br> 3. Cognitive behavioural therapy (CBT) <br> 4. Crisis resolution teams (CRT) |
| F32 \& F33 | Depressive episode \& recurrent depressive episode | 1. Cognitive behavioural therapy (CBT) <br> 2. Interpersonal therapy (IPT) <br> 3. Selective serotonin reuptake inhibitors (SSRIs) .Serotonin-norepinephrine reuptake inhibitors (SNRIs) <br> 5. Tricyclic antidepressants (TCAs) <br> 6. Monoamine oxidase inhibitors (MAOIs) <br> 7. Lithium <br> 8. Electro-convulsive therapy (ECT) |

## C.3. Step 3: evaluation of the relevant cost-effectiveness literature

Using the treatment categories identified in step 2 of this work a systematic search was conducted to attempt to identify the range of literature on the cost-effectiveness of current NHS treatment of schizophrenia and depression. For both illnesses five online databases were searched: the CostEffectiveness Analysis (CEA) Registry of the Tufts Medical Centre, the NHS Economics Evaluation Database run by the Centre for Reviews and Dissemination (CRD) at the University of York, Medline, the NICE online database of Technical Appraisals (TA) andClinical Guidelines (CG), as well as NIHR's Health Technology Assessments (HTA). All searches were conducted on the 19th October 2011.

The search strategies employed to search for relevant cost-effectiveness literature and details of the results can be found in the search strategy section at the end of this addendum. For both schizophrenia and depression five sources of information were searched sequentially: the Cost-Effectiveness Analysis (CEA) Registry, the NHS Economic Evaluation Database (EED), Medline, NICE's online library of Technical Appraisals (TA) and Clinical Guidelines (C), and finally NIHR's online library of Health Technology Assessments (HTAs). For schizophrenia this approach identified 61 unique publications, five of which were deemed to be of broad relevance to this analysis. For depression 65 publications were discovered, ten of which were relevant. A paper of relevance to our analysis of mental health was deemed to be so if it presented cost-effectiveness results (in the form of a cost per QALY ICER) of a comparison of at least two of the treatments for either schizophrenia or depression identified in section C.2. These results could be from a de-novo analysis or from a systematic review of the relevant literature.

Table C3.4 reports the cost-effectiveness results of antipsychotics for schizophrenia as first line treatments. The NICE clinical guidelines for schizophrenia (CG82)[16] demonstrate that the differences in costs and effects of the 1st and 2nd generation treatments described are very similar with ICERs comparing each to no treatment ranging from $£ 21,517$ to $£ 23,237$ per QALY. Comparisons to active treatments result in ICERs of $£ 5,156$ to $£ 33,240$ per QALY[17, 18].

Table C3.4: table showing cost-effectiveness studies of antipsychotics for schizophrenia

| Study | Treatment | Comparator | Cost (f) | QALYs | ICER $(£ / \text { QALY })$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{aligned} & \hline \text { NICE } \\ & \text { CG82[16] } \end{aligned}$ | Zotepine ( $2^{\text {nd }}$ ) | No treatment | 139,170 | 6.468 | 21,517 |
|  | Paliperidone ( $2^{\text {nd }}$ ) | No treatment | 142,173 | 6.427 | 22,121 |
|  | Olanzapine ( $2^{\text {nd }}$ ) | No treatment | 141,212 | 6.42 | 21,996 |
|  | Risperidone ( $2^{\text {nd }}$ ) | No treatment | 149,112 | 6.417 | 23,237 |
|  | Haloperidol ( $1^{\text {st }}$ ) | No treatment | 143,406 | 6.413 | 22,362 |
|  | Aripiprazole ( $2^{\text {nd }}$ ) | No treatment | 145,697 | 6.4 | 22,765 |
|  | Amisulpride (2 $2^{\text {nd }}$ ) | No treatment | 147,920 | 6.392 | 23,141 |
| Knapp et al 2008 [17] | Olanzapine ( $2^{\text {nd }}$ ) | other antipsychotics |  |  | s,156 |
| $\begin{aligned} & \text { Davies et al } \\ & 2008 \text { [18] } \end{aligned}$ | Clozapine ( $2^{\text {nd }}$ ) | Other 2 ${ }^{\text {nd }}$ gen antipsychotics |  |  | $33,240$ |
|  | Aripiprazole then risperidone | risperidone then olanzapine |  |  | 9,440 |

The CG82 results are similar to the first line treatment results from Bagnall et al. [19], shown below in Table C3.5. The cost-effectiveness of antipsychotics compared to no treatment as second, third or final therapy are less than $£ 20,000$ per QALY.

Table C3.5: table showing cost-effectiveness studies of antipsychotics for schizophrenia

| Bagnall et al., 2003 [19] |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: |
| ICER (£/QALY) | Line of treatm | ent | ) |  |
| Antipsychotic | $1{ }^{\text {st }}$ | $2^{\text {nd }}$ | $3^{\text {rd }}$ | Final |
| Chlorpromazine (1 ${ }^{\text {st }}$ ) | 21,989 | 15,185 | 15,419 | 15,303 |
| Haloperidol (1 ${ }^{\text {st }}$ ) | 24,069 | $17,177$ | 17,211 | 17,022 |
| Clozapine ( $2^{\text {nd }}$ ) | 24,500 | $17,595$ | 17,577 | 17,402 |
| Olanzapine ( $2^{\text {nd }}$ ) | 25,719 | 18,869 | 18,808 | 18,865 |
| Quetiapine (2nd) | 26,316 | 19,090 | 18,751 | 19,096 |
| Zotepine (2 ${ }^{\text {nd }}$ ) | 22,769 | 16,350 | 16,360 | 16,400 |
| Risperidone ( $2^{\text {nd }}$ ) | 22,255 | 15,596 | 15,599 | 15,700 |
| Ziprasidone (2 $2^{\text {nd }}$ (V) | 21,935 | 15,192 | 15,191 | 15,224 |
| Amisulpride (2n) ${ }^{\text {nd }}$ ) | 23,174 | 15,941 | 15,945 | 15,962 |
| Sertindole (atypical) | 23,181 | 16,297 | 16,308 | 16,354 |

Onl) one study reported the cost-effectiveness of a psychological or social intervention for schizophrenia.
Barton et al. [20] conducted a randomised trial to estimate the clinical and cost-effectiveness of social recovery orientated cognitive behavioural therapy (SRCBT) against case management alone for people recently diagnosed with psychosis. SRCBT consisted of three stages of social recovery combined with CBT techniques including vocational case management. SRCBT was found to have an ICER of $£ 18,844$ per QALY compared to case management. However, it is not clear that all forms of CBT are well represented by this one study or that these results relate well to schizophrenia since this study was for the use of social recovery CBT for psychosis disorders in general.

Table C3.6: table showing cost-effectiveness of psychological/social interventions for schizophrenia

| Study | Treatment | Comparator | ICER <br> $(£ /$ QALY $)$ |
| :--- | :--- | :--- | :--- |
| Barton et al. 2009 [20] | SRCBT | case management | 18,844 |

Table C3.7 reports the cost-effectiveness results of publications identified in the systematic search of drug treatments for depression in the NHS. As was highlighted in table C3.3 a range of drug treatments are available for depression, broadly falling into five categories: selective serotonin reuptake inhibitors (SSRIs), Serotonin-norepinephrine reuptake inhibitors (SNRIs), tricyclic antidepressants (TCAs), monoamine oxidase inhibitors (MAOIs) and lithium.

The NICE guideline CG90 tested the cost-effectiveness of numerous treatments for moderate and severe depression. It was found that across all the treatments tested the mean QALYs for moderate depression had a range of 0.053 and severe depression had a range of 0.065 . The costs had a range of 498 for moderate and $£ 396$ for severe depression. The results suggest that mirtazapine has the lowest ICER for both moderate and severe depression. If mirtazapine is not a suitable treatment option than escitalopram or sertraline is preferred because escitalopram dominates venlafaxine and sertraline dominates the remaining antidepressants. The ICERs of escitalopram versus sertraline are $£, 32,087$ per QALY for moderate depression and $£ 27,172$ per QALY for severe. The authors thus suggest that according to these results escitalopram should be considered when mirtazapine and sertraline are not suitable. Other ICERs reported in CG90 can be found in Table C3.7. CG90 states that the economic evidence had limitations and these comparisons were considered insufficient to make specific recommendations for treatments.

ICERs in other studies range from $£ 2,172$ - $£ 20,600$ per QALY, with TCA being dominated by Lofepramine (TCA) in two cases and fluoxetine being dominated by amitriptyline (TCA).

Table C3.7: table showing cost-effectiveness of drug treatments for depression

| Study | Treatment | comparator | incremental cost | incremental QALY | $\begin{aligned} & \hline \text { ICER } \\ & \text { (£/QALY) } \end{aligned}$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{aligned} & \text { NICE CG } 90 \\ & \text { [21] } \end{aligned}$ | Combined SSRI and CBT (severe depression) | SSRI alone |  |  | £,5,558 |
|  | Duloxetine (SNRI) | SSRI |  |  | £, 6,300 |
|  | Duloxetine (SNRI) | Mirtazapine (TCA) |  |  | $£ 2,400$ |
|  | Duloxetine (SNRI) | Venlafaxine (SNRI) |  |  |  |
|  | escitalopram (moderate depression) (SSRI) | $\begin{aligned} & \text { Sertraline } \\ & \text { (SSRI) } \end{aligned}$ |  |  | $\text { £ } 32,987$ |
|  | escitalopram (severe depression) (SSRI) | $\begin{aligned} & \text { Sertraline } \\ & \text { (SSRI) } \end{aligned}$ |  | $1$ | £27,172 |
| Lenox-Smith et al. 2009[22] | Venlafaxine (major depression) (SNRI) | SSRI |  | N | $£ 20600$ |
|  | Fluoxetine (SSRI) | Amitriptyline (TCA) |  |  | dominated |
| Kendrick et al.$2006[23]$ | SSRI | TCA | $\checkmark$ |  | £2,692 |
|  | TCA | Lofepramine (TCA) |  |  | dominated |
|  | SSRI | Lofepramine (TCA) | $5$ |  | £5,686 |
| Hatziandreu et al. 1994[24] | Sertraline (SSRI) | Dothiepin $(\mathrm{TC})$ |  |  | £2,172 |
| Peveler, 2005[25] | SSRI | Lofepramine (TCA) | 0.035 | £199 | £,5,686 |
|  | TCA | Lofepramine (TCA) | -0.004 | £.93 | dominated |
|  | SSRI | TCA | 0.039 | £105 | £2,692 |
| Kendrick, 2009[26] | SSRI + SC | SC |  |  | 14,854 |

Table C3.8 provides the results of the combination therapies for moderate and severe depression presented in CG90 [21]and Simon et al.[27]. These studies considered the impact of combined SSRI and CBT versus SSRI alone. Both of these studies find combined CBT and antidepressant to have ICERs of less than $£ 8,000$ per QALY.

In addition table C 3.8 provides results of analyses of computerised CBT (CCBT) compared to treatment as usual or relaxation. The results generally find CBT and CCBT to be highly cost-effective, with the exception of BT Steps[28] all ICERs are found to be under $£ 18,000$ per QALY.

Table C3.8: table showing cost-effectiveness of psychological and social intervention for depression

| Study | Treatment | comparator | incremental cost | incremental QALY | $\begin{aligned} & \text { ICER } \\ & (£ / \text { QALY }) \end{aligned}$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| NICE CG 90[21] | Combined SSRI and CBT (moderate depression) | SSRI alone |  |  | £7,052 |
|  | Combined SSRI and CBT (severe depression) | SSRI alone |  |  | £,5,558 |
| Simon et al.$2006[27]$ | CBT + antidepressants in severe dep | fluoxetine | 1 |  | £,5,777 |
|  | CBT + antidepressants in moderate dep | fluoxetine |  |  | £14,540 |
| Kaltenthaler et al., 2002 [29] | Beat the Blues CCBT (BtB) | Treatment as usual (TAU) | $5$ |  | $\begin{array}{r} £ 1,209 \text { to } \\ £ 7,692 \\ \hline \end{array}$ |
| Kaltenthaler et al., 2006 [28] | BtB | TAU | £,147 | 0.08 | £,1,801 |
|  | Cope CCBT | TAU | £193 | 0.03 | £,7,139 |
|  | Overcoming Depression CCBT | TAU | £,64 | 0.01 | ¢,5,391 |
|  | FearFighter CCBT | Relaxation CBT | £,138 | 0.058 | £,2,380 |
|  | Therapist lead CBT | Relaxation | £,194 | 0.011 | £,17,604 |
|  | BT Steps CCBT | Relaxation | £ 360 | -0.01 | Dominated |
| Hollinghurst et al., 2010 [30] | Online CCBT | TAU |  |  | £ 17,173 |

## C.4. Step 4: connection to investment and disinvestment decisions

In this section we discuss the investment and disinvestment decisions made considering the costeffectiveness information in the previous section. If we believe that decision makers will invest in treatments below their cost-effectiveness threshold and disinvest in treatments above this threshold then by considering the ICERs of treatments subject to investment and disinvestment we can create a range for their cost-effectiveness threshold. This approach and its role in the consideration of a costeffectiveness threshold has been previously discussed by Appleby et al. [31]. With a view of the costeffectiveness threshold within PBC 5 we consider how its exclusion from our calculation of the threshold might influence our results.

To identify the broad areas of investment in the disease areas we make use of recent NICE guidance documents. While NICE clinical guidance does not definitively represent observed shifts in practice and are often not well implemented in mental health trusts[32] it can help to inform our evaluation. NICE guidance does not identify areas where disinvestment should occur within a disease; as a result we have consulted experts in the respective fields to gain an understanding of any significant recent disinvestment decisions. For schizophrenia we were provided expert opinion by Professor Tim Kendall (Centre for Psychological Services Research, University of Sheffield) and for depression by Professor Simon Gilbody (Health Sciences, University of York).

For both schizophrenia and depression we will briefly discuss the areas of investment and disinvestment in two care categories: (i) drug treatments, and (ii) psychological and social interventions.

## C.4.1. Results of step 4 for schizophrenia

## C.4.1.1. Analysis of drug treatment

Antipsychotics used for the treatment of schizophrenia can be broadly identified as first or second generational (typical and atypical antipsychotics). To a certain extent there is still a debate over the relative strengths and weaknesses of each[33], and the significance of the adverse events associated with the second generation may still not be fully understood (such as the impact on new-onset type-2 diabetes [34]). However, our clinical experts indicated that clinicians were largely concerned with the adverse effects associated with the second generational drugs, and the increasing evidence questioning the relative efficacy, such as Rosenheck [33] who suggests that the first generational drugs in many cases are just as effective. Recent NICE guidance leaves the choice of first or second generational drugs to the clinicianto decide[16].

When considering the impact on our estimate of the threshold of the possible shift to first generation from second generation antipsychotics we must attempt to generalise about the relative cost-effectiveness of the two. Clearly this is difficult as each generation represents many different drugs. However, from CG82 the costs and benefits of the mainstream antipsychotics are broadly similar Table C3.4. This would suggest that a shift away from the second generation back towards the first would have little impact on the overall threshold as the costs and benefits associated with each are very similar.

Olanzapine came off patent in the third quarter of 201129. Olanzapine and similar second generation antipsychotics are associated with a cost of around $£ 30$ million a year ${ }^{30}$, clearly the introduction of generics to the market would significantly reduce this cost and thus increases the cost-effectiveness of these drugs. While this shift does not fall within the years of our analysis, it will have a significant impact on the future value of the cost-effectiveness threshold.

The other significant area of debate, as identified by our clinician, is the role of the antipsychotic clozapine, which has often been viewed as the most effective antipsychotic drug for schizophrenia however has been connected with some severe adverse events (such as myocarditis[35], agranulocytosis[36] and central nervous system depression). This has lead to the NICE guidelines advising clozapine only if an array of other antipsychotics has failed[16]. While clozapine is highly clinically effective it is associated with a higher overall cost (a significant proportion due to the associated adverse events). As is shown in table C3.4Davies et al. [18] show clozapine to have an ICER of $£ 33,240$ when compared to other second generation antipsychotics. Disinvestment of clozapine suggests that the threshold is lower than $£ 33,240$ per QALY. However, current investment in other 1 st line antipsychotics suggests that the threshold in mental health is over $£ 23,237$ per QALY.

## C.4.1.2. Analysis of psychological and social intervention

In this section we discuss all non-drug related interventions for schizophrenia. The NICE guidelines[16] outline the provision of CBT , arts therapy and family interventions to treat schizophrenia, however, efficacy of these interventions is disputed[37] and little is known about their cost-effectiveness. The systematic review only yielded one paper that was relevant to our analysis, as is shown in Table C3.6. The Barton et al. [20] study found that SRCBT had an ICER of less than $£ 20,000$ per QALY relative to case management. However, as mentioned previously, this study may not represent all forms of CBT or schizophrenia.

Our clinical advisors informed us that CBT provision varies significantly across PCTs and therefore represents an intervention likely to be subject to investment and disinvestment at the margin. The yariation in CBT provision (and indeed other psychosocial/social interventions) is largely a result of the poor support for its efficacy and significant initial cost.

Other interventions of relevance to this investigation include art therapies and family interventions. As with CBT there is a significant variation in the provision of family interventions. No information on its cost-effectiveness was found from our search. Art therapies include: music therapy, art therapy, and body

[^90]movement or dance therapy. Our clinical advisors have highlighted increasing investigations into arts therapy, including the "Matisse trial" [38], publications around which have shown that art therapy as adjunctive therapy had little benefit over a comparator activity or treatment as usual[37]. No information on its cost-effectiveness was found from our search.

Early interventions in schizophrenia, which aims to identify and treat early symptoms associated with schizophrenia, have been a significant area of investment over recent years in the NHS. While we were unable to identify any relevant cost-effectiveness literature around early interventions in schizophrenia it is generally expected that these represent cost-effective interventions over the long term ${ }^{31}$.

While the lack of cost-effectiveness literature clearly limits the potential to directly associate these interventions with the wider cost-effectiveness threshold it is widely accepted that many social interventions for schizophrenia (specifically around CBT and family interventions) are cost-saving for the $\mathrm{NHS}^{32}$, as they reduce hospitalisation by reducing emergency hospital access and relapse rates that are high in schizophrenia representing the majority of related hospitalisations[39].

Investment in CBT with an expected ICER of $£ 18,844$ per QALY suggests that the threshold for mental health treatments is above this value.

## C.4.2. Results of step $\mathbf{4}$ for depression

As table C3.3 shows depression is associated with a wider range of treatment than schizophrenia, specifically a wider range of drug treatments are available. As with the schizophrenia section of this addendum we will deal with the treatments under the two categories of drug treatments and psychological/ social interventions. Electroconvulsive therapy (ECT), which is included in the treatment options available in the NHS as shown in table C3.3, is excladed from this analysis based on expert opinion on the grounds of it being a very rarely used butextreme treatment that is not likely to be further subject to substantial investment or disinvestment, so is not relevant for our analysis.

Recent NICE clinical guidance[21] highlights a range of key priorities for implementation. As with schizophrenia there is no guarantee that these are the areas of investment in depression care but it represents a suitable outline of the areas of interest. Several areas are highlighted:

- Early identification and diagnosis
- Low intensity psychological interventions (CBT, CCBT and group physical activity) for persistent sub-threshold depressive symptoms or mild to moderate depression
- Reduced routine use of antidepressant for sub-threshold depressive symptoms or mild depression
- Combination therapies (antidepressant and psychological) for moderate or severe depression
- Extension of thefapy (antidepressant and psychological) beyond remission to reduce relapse
- SSRIs are presented as the preferred type of antidepressant due to their equivalent efficacy and favourable risk-benefit ratio.

These are the areas of investment that our analysis will focus on.

## C.4.2.1. Analysis of drug treatment for depression

Our clinical advisors reported that the current area of activity in antidepressants is the creation of drugs such as escitalopram (an SSRI) and venlafaxine (an SNRI) that are relatively similar to generic treatments currently in the market. As these new drugs are covered by patents they are relatively expensive. Table C3.7 reports the results on the cost effectiveness of these two drugs from NICE CG90[21] as well as Lenox-Smith et al.[22]. In both reports the drugs are compared to alternative SSRIs in moderate and severe depression. In both cases the newer SSRIs were approved by NICE with an ICER for escitalopram of $£ 32,928$ per QALY and for venlafaxine of $£ 20,600$ per QALY. If mirtazapine and

[^91]sertraline are not suitable then the ICER of escitalopram for moderate depression is $£, 5,357$ per QALY compared to citalopram. While evidence was not available on whether clinicians were making use of these newer SSRIs, an investment in them away from alternative SSRIs may represent an increase in the cost-effectiveness threshold due to the relatively high ICERs reported in the two studies. However, the cost-effectiveness of each depends on what they displace and ICERs may be lower if the more costeffective treatments have failed.

Investment decisions in the NHS for antidepressants are likely to represent changes in the type of antidepressant being prescribed rather than a shift from no treatment to treatment. The majority of trials discovered by systematic review given in table C3.7 show that while the ICER of SSRIs versus TCAs is very low [25] this is largely driven by very small gains in QALYs but for a similarly small increase in cost. As a result any observed investment in SSRIs away from TCAs is likely to lead to a small decrease in an observed threshold for the NHS.

## C.4.2.2. Analysis of psychological and social intervention for depression

The NICE guidelines reported in CG90 place a lot of focus on the provision of psychological interventions such as CBT (and CCBT) over antipsychotics wherever possible. Table C3.8 provides the results of the combination therapies for moderate and severe depression presented in CG90 [21] and Simon et al. [27]. These considered the impact of combined SSRI and CBT versus SSRI alone and concluded that combined therapies in both populations had ICERs of less than $, 15,000$ per QALY. According to our clinicians, this is an area that is likely to have had significant investment in recent years.

The two HTAs reported in table C3.8 [28, 29] provide a good analysis of the cost-effectiveness of computer cognitive behavioural therapy (CCBT) versus treatment as usual (TAU). They show that the CCBTs investigated have ICER of less than $£ 8,000$ per QALY relative to TAU. Further analyses investigated different kinds of CCBT and found that compared to relaxation CBT ICERs ranged from $£ 2,380$ per QALY to dominated. Hollinghurst et al. report that two CBT interventions compared to TAU had ICERs of $£ 17,173$.

As NICE guidelines encourage the use of CBT and our clinical experts believe this has been an area of increased investment, this review suggests that the threshold in mental health is over $£, 17,173$ per QALY.

## D. Conclusion

There is very little accessible data on the investment and disinvestment decisions in specific areas of mental health and so we relied on the opinions of clinical experts. The NHS Information Centre has some information on prescriptions of mental health treatments, however it was not clear for which diseases these treatments were being used or for which line of therapy. As a result this data was not included in our analysis as it was decided it may not represent the investment and disinvestment decisions that we were seeking to identify.

Most treatments reviewed had an ICER of less than $£ 24,000$ per QALY. Two treatments had higher ICERs. Clozapine for the first line treatment of schizophrenia was found to have an ICER of $£ 33,240$ per QALY compared to other 2nd generation antipsychotics. NICE's recommendation to use clozapine only as a last line treatment suggests that the threshold is less than $£ 33,240$ per QALY. Escitalopram for moderate depression has been recommended by NICE and was reported to have an ICER of 32,987 per QALY compared to seratraline. Conclusions on the threshold from this finding are unclear. The costeffectiveness of escitalopram in the NHS will depend on its use. If it is used rather than seratraline then the threshold may be over $£ 32,987$, but if it is used as third line therapy than according to CG 90 its use is less costly and more effective than the next best options.

How well the actual threshold reflects the ICERs reported above depends on how well clinical practice matches the clinical guidelines i.e. whether the more cost-effective treatments are being used first.

## Search Strategies

## Search Strategy for schizophrenia

CEA Registry search:
Six keywords associated with the entire Schizophrenia, schizotypal and delusional disorders ICD10 subchapter were search for in the CEA Registry, these were: schizophrenia, schizotypal, delusional, psychotic, schizoaffective and psychosis. A search for any of these keywords in the Registry yielded 18 different papers at the time of searching, with four of these being deemed suitable for our investigation (Barton 2009, Davies 2008, Jarbrink 2009, Knapp 2008 and Davies 2007).

NHS EED search:


A single relatively simple search strategy was defined to investigate NHS EED, this was as follows: ((Schizophrenia) AND (cost effectiveness):TI) and Economic evaluation:ZDT and Abstract:ZPS) This result strategy yielded 28 hits, only one of which was both relevant to our search and not discovered in the CEA Registry search (Rosenheck 2007).

Medline search:
Medline was searched using the strategy:
cost benefit analysis and (schizophrenia or schizotypal personality disorder or delusions) and Great britain(MeSH)

This strategy yielded 13 hits, none of which were both relevant and had not been previously identified through the CEA Registry of NHS EED searches.

NICE Technical Appraisals (TA) and Clinical Guidelines (CG):
NICE's online database of published mental health related TAs and CGs
(http://www.nice.org.uk/guidance/index.jsp?action=byTopic\&o=7281) was searched for schizophrenia related publications. Only one was found to fulfil our criteria for schizophrenia: CG82.

## NIHR's HTAs:

Finally NIHR's database of published HTAs was searched. This activity discovered one additional relevant publication: HTA 00/20/01 - Bagnall, 2003.

## Search Strategy for Depression

## CEA Registry:

Two keywords were searched on the CEA Registry, they were: depression and depressive. These keywords yielded 17 papers, 5 of which were deemed relevant for our purposes (Hollinghurst 2010, Lenox-Smith 2009, Kendrick 2006, Simon 2006, and Hatzinandreu 1994).

## NHS EED search:

A search similar in structure to the search for schizophrenia papers was conducted in NHS EED: ((depressive OR depression):TI AND (cost-effectiveness):TI) and Economic evaluation:ZDT and Abstract:ZPS) IN NHSEED

This yielded 43 hits, none of which were both relevant and previously undiscovered by the CEA registry search. Due to the complete nature of the CEA Registry and NHS EED searches as well as time constraints on the systematic review, a Medline search was not conducted as it was decided it would not provide sufficient added value.

NICE Technical Appraisals (TA) and Clinical Guidelines (CG):
Searching the NICE database of TAs and CGs yielded one publication deemed relevant to the analysis: CG90- depression in adults.

NIHR's HTAs:

A search of the NIHR's online database of published HTAs yielded four relevant publications:

HTA 01/23/01- Bennett, 2000
HTA04/01/01- Kaltenthaler, 2006
HTA- 96/61/11-Peveler, 2005
HTA- 01/70/05-Kendrick, 2009

## Appendix C: Addendum 4

What type of health is forgone by the approval of a new technology?

The methods of analysis described in this work can identify not only how many QALYs are likely to be forgone across the NHS as a consequence of approving a technology which imposes incremental costs on the NHS, it can also indicate where those QALYs are likely to be forgone and how they are made up, i.e., the additional deaths, life years lost (unadjusted and adjusted for quality of life) and the quality of life impacts on those with disease. Based on the 2008 central estimate of the cost per QALY threshold, we will exemplify within this Addendum the likely health displaced elsewhere in the NHS as a consequence/ of approving a new technology.

## The example of ranibizumab for diabetic macular oedema

In 2011, NICE considered whether ranibizumab for the treatment of diabetic macular oedema in patients with central retinal thickness $\geq 400$ micrometres should be approved for widespread use in the NHS (TA237[40]). Initially this technology was rejected by NCE on the grounds that, atits current price, it would be unlikely to be cost effective. In 2012, however, a rapid review of TA237 [41] approved Ranibizumab if use was restricted to the most cost effective sub group (those with central retinal thickness $\geq 400$ micrometres) and after a Patient Access Scheme (PAS) for this subgroup of patients was offered (details of the PAS which provides a discounts to the NHS is commercial in confidence). The Committee concluded that the most plausible ICER for the subgroup of people with thicker retinas was likely to be higher than the manufacturer's estimate (of $£ 13,322$ per QALY), but would be under $£ 25,000$ per QALY gained.[41]

The appraisal and guidance documents (http://guidance.nice.org.uk/TA/Wave23/41) provide the information required to estimate the additional NHS costs of treating this sub group of patients each year. The original manufacturer submission presented an estimate of the numbers of patients in the NHS eligible to receive ranibizumab, based on its licensed indication[42]. These estimates are presented in Table C4.1. In the first year of implementation, up to 44,000 NHS patients would be eligible for treatment with ranibizumab based on its licensed indication. No consideration is made as to the size of the sub-population approved for treatment, however the RESTORE trial (that informs the submission) found approximately half of the participants in the study to be in this sub-population [114 of $217(52.5 \%)$ ].[40] The subgroup of patients where ranibizumab was ultimately approved is thus likely to be approximately 23,000 in the first year after approval.

Table C4.1 Estimated size of the NHS population eligible for ranibizumab [42]

| Licensed indication | $\mathbf{2 0 1 1}$ | $\mathbf{2 0 1 2}$ | $\mathbf{2 0 1 3}$ | $\mathbf{2 0 1 4}$ | $\mathbf{2 0 1 5}$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Prevalent cases |  |  |  |  |  |
| Incidenteases | 43,847 | 0 | 0 | 0 | 0 |
| Totaleligible number of patients | 43,847 | 4,481 | 5,481 | 5,481 | 5,481 |
| Sub-population approved for treatment by NICE |  | 54,809 | 60,290 | 65,771 |  |
| Prevalent cases | 23,020 | 0 |  |  |  |
| Incident cases | 0 | 2,878 | 2,878 | 2,878 | 2,878 |
| Total eligible number of patients | 23,020 | 25,897 | 28,775 | 31,652 | 34,530 |

The incremental costs associated with the new treatment (compared to laser monotherapy) in the initial submission (TA237) were $£ 3,506$ per patient[42]. Given estimates reported in the rapid review are not available (commercial in confidence), we will use this estimate of incremental costs for the subpopulation of interest. These data suggests that the approval of ranibizumab in this subgroup at the original appraisal in 2011 (i.e., without a PAS) would impose just over $£ 80 \mathrm{~m}$ of additional NHS costs for treating the eligible population each year.

Table C4.2 Estimated total budget impact of ranibizumab

|  | 2011 | 2012 | 2013 | 2014 | 2015 |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Total eligible number of patients | 23,020 | 25,897 | 28,775 | 31,652 | 34,530 |
| Total cost, without PAS ( $¢$ ) | £80,708,120 | £, $90,794,882$ | £100,885,150 | £110,971,912 | £,121,062,180 |
| Total cost, 30\% lower incremental costs( $¢$ ) | £.56,495,684 | £63,556,417 | £,70,619,605 | £.77,680,338 | £ 84,743,526 |

With introduction of the PAS, it is likely that a simple discount on the acquisition price of the new technology has been approved by the DH.[41] Given the scale of the discount is not available (commercial in confidence) we assumed that this discount would reduce incremental costs by $30 \%$ (to $£ 2,454$ per patient). After such a PAS, the approval of ranibizumab in this subgroup would impose just $£^{5} 56 \mathrm{~m}$ (rather than $£ 80 \mathrm{~m}$ ) of additional NHS costs for treating the eligible population in the first year.

## What type of health is forgone by approval of a new technology?

Based on the 2008 central estimate of the cost per QALY threshold ( $£ 18,317$ in Table 5.1 the approval of ranibizumab without a PAS would have been likely to displace 4,367 QALYs elsewhere in the NHS. However, the analysis which underpins the threshold estimate can also be used to identify where the additional NHS cost of $£ 80 \mathrm{~m}$ are likely to impact and where and what type of health effects are likely to be forgone. These are illustrated in Table C4.3.

Table C4.3: Heath forgone across PBCs due to the approval of ranibizumab ( $£ 80 \mathrm{~m}$ budget impact)

| PBC PBC description | change in spend (m) [1] | Additional deaths | Life years foregone | Total QALYs forgone [4] | QALYs foregone <br> Due to premature death [5] | Quality of life effects $[6]$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 2 Cancer | £2.59 | 22 | 217 | 153 | 141 | 11 |
| 10 Circulatory problems | £.4.40 | 132 | 672 | 625 | 427 | 198 |
| 11 Respiratory problems | $£_{2} 2.66$ | 78 | 93 | 1,330 | 58 | 1,272 |
| 13 Gastro-intestinal | £.1.86 | 15 | 143 | 255 | 94 | 161 |
| Big 4 | \&12 | 246 | 1,126 | 2,362 | 721 | 1,641 |
| 1 Infectious diseases | £1.89 | 4 | 31 | 91 | 21 | 70 |
| 4 Endocrine problems | £1.10 | 4 | 29 | 351 | 19 | 332 |
| 7 Neurological problems | ¢ 3.47 | 7 | 38 | 632 | 25 | 608 |
| 17 Genito-urinary problems | 2. 2.69 | 13 | 19 | 61 | 12 | 49 |
| 16 Trauma \& injuries | ¢ 4.46 | 0 | 0 | 0 | 0 | 0 |
| 18+19 Maternity \& neonates | \&.3.96 | 0 | 2 | 1 | 1 | 0 |
| 11 PBCs | $£^{29}$ | 275 | 1,245 | 3,500 | 798 | 2,701 |
| 3 Disorders of Blood | $£ 2.33$ | 1 | 6 | 82 | 4 | 78 |
| 5 Mental Health Disorders | $£ 20.25$ | 12 | 55 | 406 | 35 | 371 |
| 6 Learning Disability | $\AA 1.18$ | 1 | 4 | 15 | 3 | 12 |
| 8 Problems of Vision | $\AA 2.20$ | 0 | 2 | 29 | 1 | 28 |
| 9 Problens of Hearing | $£ 0.99$ | 0 | 1 | 52 | 0 | 52 |
| 12 Dental-problems | $£ 3.27$ | 0 | 0 | 59 | 0 | 59 |
| 14 Skin | $£ 2.23$ | 2 | 7 | 13 | 5 | 8 |
| 15 Musculo skeletal system | $£ 4.11$ | 3 | 15 | 203 | 10 | 193 |
| 20 Poisoning and AE | $£ 1.05$ | 0 | 2 | 6 | 1 | 5 |
| 21) Healthy Individuals | $£ 4.01$ | 0 | 1 | 3 | 0 | 2 |
| 22 Social Care Needs | £3.41 | 0 | 0 | 0 | 0 | 0 |
| 23 Other | ¢, 5.88 | 0 | 0 | 0 | 0 | 0 |
| All (23 PBCs) | $£ 80$ | 295 | 1337 | 4367 | 859 | 3509 |

Table C4.4: Heath forgone across specific PBCs and groups of ICDs due to the approval of ranibizumab ( $£ 80 \mathrm{~m}$ budget impact)

| Total change in spend analysed $=£ 80 \mathrm{~m}$ | change in spend <br> (m) <br> [1] | Life years foregone [2] | Total QALYs forgone [3] | QALYs foregone <br> Due to premature death [4] | Quality of life effects [5] |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Overall |  | 1337 | 4367 | 859 | 3509 |
| PBC specific <br> PBC2 (Cancer) <br> Malignant neoplasms, digestive organs (C15-C26) <br> Malignant neoplasms, respiratory system and intrathoracic organs (C30-C39) <br> Malignant neoplasms, breast and female genital organs (C50-C58) <br> Malignant neoplasms, stated or presumed to be primary, of lymphoid, haematopoietic and related tissue (C81-C96) <br> Malignant neoplasms, human male genital organsmale genital organs (C60-C63) Other ICD codes in this PBC | 62.59 | $\begin{gathered} 217 \\ 59 \\ 50 \\ 40 \\ 19 \\ 13 \\ 37 \\ \hline \end{gathered}$ | $\begin{gathered} 153 \\ 41 \\ 33 \\ 31 \\ 13 \\ 2 \\ 25 \end{gathered}$ | 141 38 33 25 13 8 24 | $\begin{gathered} 11 \\ 2 \\ 0 \\ 6 \\ 0 \\ 1 \\ 1 \\ \hline \end{gathered}$ |
| PBC10 (Circulatory) <br> Ischemic heart diseases (I20-I25) <br> Cerebrovascular diseases (I60-I69) <br> Other forms of heart disease (I30-I52) <br> Congenital malformations and deformations circulatory system (Q20-Q28) Diseases of veins, lymphatic vessels and lymph nodes, not elsewhere classified (I80-I89) <br> Other ICD codes in this PBC | £4.40 |  | 625 345 132 46 31 20 50 | $\begin{gathered} \hline 427 \\ 242 \\ 85 \\ 35 \\ 10 \\ 16 \\ 39 \\ \hline \end{gathered}$ | $\begin{gathered} \hline 198 \\ 103 \\ 47 \\ 11 \\ 22 \\ 4 \\ 11 \\ \hline \end{gathered}$ |
| PBC11 (Respiratory) <br> Chronic lower respiratory diseases (J40-J47) <br> Lung diseases due to external agents (J60-J70) <br> Other diseases of upper respiratory tract (J30-J39) <br> Other respiratory diseases principally affecting the interstitium (J80-J84) <br> Other diseases of pleura (J90-J94) <br> Other ICD codes in this PBC | $\epsilon^{2.66}$  | 93 41 6 5 2 2 37 | $\begin{gathered} 1330 \\ 1137 \\ 52 \\ 47 \\ 19 \\ 19 \\ 57 \\ \hline \end{gathered}$ | $\mathbf{5 8}$ 26 4 3 1 1 23 | $\begin{gathered} \hline 1272 \\ 1111 \\ 48 \\ 44 \\ 17 \\ 17 \\ 34 \\ \hline \end{gathered}$ |
| PBC7 (Neurological) <br> Episodic and paroxysmal disorders (G40-G47) <br> Extrapyramidal and movement disorders (G20-G26) <br> Other degenerative diseases of the nervous system (G30-G32) <br> Other disorders of the nervous system (G90-G99) <br> Nerve, nerve root and plexus disorders (G50-G59) <br> Other ICD codes in this PBC | $\int 63.47$ | $\begin{gathered} \hline 38 \\ 7 \\ 7 \\ 7 \\ 6 \\ 1 \\ 1 \\ 14 \\ \hline \end{gathered}$ | $\begin{gathered} \hline 632 \\ 464 \\ 52 \\ 32 \\ 16 \\ 14 \\ 54 \\ \hline \end{gathered}$ | 25 5 4 4 1 1 10 | 608 459 48 28 15 13 45 |
| PBC5 (Mental Health) <br> Mental and behavioural disorders due to psychoactive substance use (F10-F19) Mood (affective) disorders (F30-F39) <br> Organic, including symptomatic, mental disorders ( $\mathrm{FOO}-\mathrm{F} 09$ ) <br> Neurotic, stress-related and somatoform disorders (F40-F48) <br> Behavioural syndromes associated with physiological disturbances and physical factors (F50-F59) <br> Other ICD codes in this PBC | 620 | $\begin{gathered} 55 \\ 21 \\ 1 \\ 18 \\ 1 \\ 1 \\ 13 \end{gathered}$ | $\begin{gathered} \hline 406 \\ 166 \\ 69 \\ 48 \\ 38 \\ 23 \\ 63 \end{gathered}$ | $\begin{gathered} 35 \\ 15 \\ 1 \\ 11 \\ 0 \\ 1 \\ 8 \end{gathered}$ | $\begin{gathered} \hline 371 \\ 151 \\ 68 \\ 37 \\ 38 \\ 22 \\ 55 \end{gathered}$ |

The results reported in Table C4.3 suggests that approval is likely to results in 295 additional deaths (column 2) and 1,337 life years (column 3) forgone, most of which are likely to occur in Circulatory, Respiratory and Cancer PBCs. However, impact of approval of this technology on QALYs forgone due to premature death (column 5) only accounts for a proportion of the total QALY effects (column 4). Most $(3,509)$ are associated with quality of life forgone during disease (column 6). These quality of life impacts are most likely to occur in Respiratory, Neurological and Mental health PBCs. The PBC level effects in Table C4.3 can also be examined at ICD level (Table C4.4) whilst recognising the caveats discussed in Chapter 4. For example, within in the respiratory PBC, it appears to be Influenza and Pneumonia (J09-J18) where most additional deaths, life years and quality of life would be forgone. In the Mental Health PBC the additional deaths appear to be associated with disorders due to psychoactive substance use (F10-F19) and Schizophrenia, schizotypal and delusional disorders (F20-F29). The impact of a reduction in the price of this technology, either through value based pricing or the PAS that was offered during the rapid review, can also be examined in the same way. The PAS was commercial in confidence, so here we will consider the hypothetical case that a $30 \%$ reduction in NHS costs (incremental costs) would make this technology cost-effective for this subgroup of patients. Such a discount would be expected to save 1,310 QALYs including 89 deaths averted, 401 life years ( 258 when adjusted for quality) and quality of life effects during disease equivalent to 1,053 QALYs, compared to approval of the technology at the original list price (Table C4.5).

Table C4.5: Heath forgone before and after a hypothetical PAS scheme on ranibizumab

| PBC description | change in spend (m) [1] | Additional deaths [2] | Life years foregone [3] |  | ALYs foreg <br> Due to premature death [5] | Quality of life effects [6] |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Before PAS <br> big 4 <br> 11PBCs <br> all 23 | $\begin{aligned} & £ 12 \\ & £ 29 \\ & £ 80 \end{aligned}$ | $\begin{aligned} & 246 \\ & 275 \\ & 295 \end{aligned}$ | $\begin{aligned} & 1,126 \\ & 1,245 \\ & 1,337 \end{aligned}$ | $\begin{aligned} & 2,362 \\ & 3,500 \\ & 4,367 \end{aligned}$ | $\begin{aligned} & 721 \\ & 798 \\ & 859 \end{aligned}$ | $\begin{aligned} & 1,641 \\ & 2,701 \\ & 3,509 \end{aligned}$ |
| After PAS big 4 11PBCs all 23 | $\begin{aligned} & £ 8 \\ & £ 20 \\ & £_{5} 56 \end{aligned}$ |  | $\begin{aligned} & 788 \\ & 871 \\ & 936 \end{aligned}$ | $\begin{aligned} & 1,654 \\ & 2,450 \\ & 3,057 \end{aligned}$ | $\begin{aligned} & 505 \\ & 559 \\ & 601 \end{aligned}$ | $\begin{aligned} & 1,149 \\ & 1,891 \\ & 2,456 \end{aligned}$ |
| Difference <br> big 4 <br> 11PBCs <br> all 23 | $\begin{gathered} -63 \\ -f^{2} \\ -f^{24} \\ \hline \end{gathered}$ | $\begin{aligned} & -74 \\ & -82 \\ & -89 \end{aligned}$ | $\begin{array}{r} -338 \\ -373 \\ -401 \\ \hline \end{array}$ | $\begin{gathered} -709 \\ -1,050 \\ -1,310 \\ \hline \end{gathered}$ | $\begin{aligned} & -216 \\ & -239 \\ & -258 \\ & \hline \end{aligned}$ | $\begin{gathered} -492 \\ -810 \\ -1,053 \\ \hline \end{gathered}$ |

## References

1. Wailoo, A.D., S. Tosh, J, The incorporation of health benefits in cost utility analysis using the eq-5d: report by the decision support unit. School of Health and Related Research, University of Sheffield, 2010.
2. NICE, Guide to the methods of technological appraisal. ref: N1618, 2008.
3. Dolan, P., et al., A social tariff for EuroQol: results from a UK general population survey. CHE discussion paper 138, University of York, 1995.
4. Currie, C.J., et al., The routine collation of health outcomes data from hospital treated subjects in the Health Outcomes Data Repository (HODaR): descriptive analysis from the first 20,000 subjects. Value in health the journal of the International Society for Pharmacoeconomics and Outcomes Research, 2005. 8(5): p. 581-590.
5. Cohen, J.W., et al., The Medical Expenditure Panel Survey: a national health information resource. Inquiry, 1996. 33(4): p. 373-89.
6. Murray, C. and A.e. Lopez, The global burden of disease. Geneva, World Health Organization, Harvard School of Public Health, World Bank., 1996.
7. Claxton, K., et al., Informing a decision framework for when NICE should recommend the use of health technologies only in the context of an appropriately designed programme of evidence development. Health Technology Assessment, in press 2012.
8. Claxton, K., The irrelevance of inference: a decision-making approach to the stochastic evaluation of health care technologies. J Health Econ, 1999. 18(3): p. 341-64.
9. WHO, The global burden of disease: 2004 update. 2008.
10. Mathers, C., et al., Global burden of disease in 2002: data sources, methods and results. GPE Discussion Paper No. 54. Geneva, World Health Organization.,2003.
11. Appleby, J. and N. Devlin, Getting the most out of PROMs. Putting bealth outcomes at the heart of NHS decision-making The King's Fund, London, 2010.
12. Gutacker, N., et al., Truly inefficient or providing better quality of care? Analysing the relationship between risk-adjusted hospital costs and patients' health outsomes. Health Economics, 2012. doi: 10.1002/hec. 2871.
13. Feng, Y., D. Parkin, and N. Devlin, Assessing the Performance of the EQ-V AS in the NHS PROMs Programme. Office of Health Economics: Kings Fund, 2012. Research Paper 12/01.
14. Centre, N.I., Provisional Monthly Patient Reported Outcome Measures (PROMs) in England - April 2012. 2012.
15. Appleby, J., Harrison, T., Foot, C., Smith, A. and Gilmour, S., Explaining variations in primary care trusts' spending on cancer services. The King's Fund, London, 2011.
16. NICE, CG82: Core interventions in the treatment and management of schizophrenia in primary and secondary care. 2009.
17. Knapp, M., et al., Cost-utility analysis of treatment with Olanzapine compared with other antipsychotic treatments in patients with schizophrenia in the Pan-European SOHO study. Pharmacoeconomics, 2008. 26(4): p. 341-358.
18. Davies, L.M., et al., A randomized controlled trial of the cost-utility of second-generation antipsychotics in people Wirth psychosis and eligible for clozapine. Value Health, 2008. 11(4): p. 549-62.
19. Baghall, A.M., et al., A systematic review of atypical antipsychotic drugs in schizophrenia. Health Technol Assess, 2003. 7(13): p. 1-193.
20. Barton, G.R., et al., Cognitive behaviour therapy for improving social recovery in psychosis: cost-effectiveness analysis. Schizophr Res, 2009. 112(1-3): p. 158-63.
21. NICE, CG90: Depression the treatment and managementof depression in adults. 2009.
22. Lenox-Smith, A., et al., Cost effectiveness of venlafaxine compared with generic fluoxetine or generic amitriptyline in major depressive disorder in the UK. Clinical Drug Investigation, 2009. 29(3): p. 173-184.
23. Kendrick, T., et al., Cost-effectiveness and cost-utility of tricyclic antidepressants, selective serotonin reuptake inbibitors and lofepramine: randomised controlled trial. Br J Psychiatry, 2006. 188: p. 337-45.
24. Hatziandreu, E.J., et al., Cost utility of maintenance treatment of recurrent depression with sertraline versus episodic treatment with dothiepin. Pharmacoeconomics, 1994. 5(3): p. 249-68.
25. Peveler, R., et al., A randomised controlled trial to compare the cost-effectiveness of tricyclic antidepressants, selective serotonin reuptake inhibitors and lofepramine. Health Technol Assess, 2005. 9(16): p. 1-134, iii.
26. Kendrick, T., et al., Randomised controlled trial to determine the clinical effectiveness and cost-effectiveness of selective serotonin reuptake inhibitors plus supportive care, versus supportive care alone, for mild to moderate depression with somatic symptoms in primary care: the THREAD (THREshold for AntiDepressant response) study. Health Technol Assess, 2009. 13(22): p. iii-iv, ix-xi, 1-159.
27. Simon, J., et al., Treatment options in moderate and severe depression: decision analysis supporting a clinical guideline. Br J Psychiatry, 2006. 189: p. 494-501.
28. Kaltenthaler, E., et al., Computerised cognitive behaviour therapy for depression and anxiety update: a systematic review and economic evaluation. Health Technol Assess, 2006. 10(33).
29. Kaltenthaler, E., et al., A systematic review and economic evaluation of computerised cognitive behaviour therapy for depression and anxiety. Health Technol Assess, 2002. 6(22): p. 1-89.
30. Hollinghurst, S., et al., Cost-effectiveness of therapist-delivered online cognitive-bebavioural therapy for depression: randomised controlled trial. Br J Psychiatry, 2010. 197(4): p. 297-304.
31. Appleby, J., et al., Searching for cost effectiveness thresholds in the NHS. Health Policy, 2009. 91 (3): p. 239-245.
32. Mears, A., et al., Progress on NICE guideline implementation in mental health trusts: meta-analyses. The Psychiatrist, 32:383-387., 2008.
33. Rosenheck, R.A., et al., Cost-effectiveness of second-generation antipsychotics and perphenarine in a randomized trial of treatment for chronic schizophrenia. Am J Psychiatry, 2006. 163(12): p. 2080-9.
34. Lambert, M.T., et al., New-onset type-2 diabetes associated with atypical antipsychotic medications. Progress in Neuro-Psychopharmacology and Biological Psychiatry, 2006. 30(5): p. 919-923.
35. Haas, S.J., et al., Clozapine-associated myocarditis: a review of 116 cases of suspected myocarditis associated with the use of clozapine in Australia during 1993-2003. Drug Saf, 2007. 30(1): p. 47-57.
36. Alvir, J.M., et al., Clozapine-induced agranulocytosis. Incidence and risk factors in the United States. N Engl J Med, 1993. 329(3): p. 162-7.
37. Crawford, M.J., et al., Group art therapy as an adjunctive treatment for people with schizophrenia: multicentre pragmatic randomised trial. BMJ, 2012. 344.
38. Kendall, T., Treating negative symptoms of schirophrenia. BMJ, 2012. 344.
39. Tarrier, N., et al., The Salford Family Intervention Project: relapse rates of schizophrenia at five and eight years. The British Journal of Psychiatry, 1994. 165(6): p. 829-32.
40. NICE, TA237: Ranibizumab for the treatment of diabetic macular oedema. 2011.
41. NICE, Macular oedema (diabetic) - ranibǐ̌umab (rapid review of TA237): appraisal consultation document. 2012.
42. Novartis, Single technology appraisal (STA) manufacturer submission: Lucentis $\circledR_{~(r a n i b i z u m a b) ~ f o r ~ t h e ~}^{\text {( }}$ treatment of visual impairment due to diabetic macular oedema (DMO). 2010.

[^0]:    OKarl Claxton, Steve Martin, Marta Soares, Nigel Rice, Eldon Spackman, Sebastian Hinde, Nancy Devlin, Peter C Smith, Mark Sculpher

[^1]:    ${ }^{1}$ Thisis the case so long as all incremental costs are health care system costs or, as currently, the perspective adopted by NICE is commonly restricted to the health care system. If a broader perspective was to be adopted and, insofar as there are some incremental costs (or benefits) of adopting a technology that fall on private consumption, then $v$ does become relevant to decision making because it represents the value of these consumption effects in terms of health. In these circumstances it would be inappropriate either to compare an ICER which included consumption effects to k (because consumption costs do not displace health in the NHS), or to compare it to v (because some of the costs do not displace private consumption but displace health at rate k . The ratio of $\mathrm{k} / \mathrm{v}$ represents the value of NHS resources relative to private consumption. Observing $\mathrm{k}<\mathrm{v}$ would suggest a positive shadow price on NHS resources and public expenditure more generally, i.e., it would indicate that a public sector $\mathcal{E}$ is scarce relative to a private $\mathcal{f}$. See Claxton K., Walker S., Sculpher MJ. and Palmer S. Appropriate perspectives for heath care decisions. Centre for Health Economics, University of York. CHE Research Paper 54; 2010 for a more extended treatment of perspective, the implications for decision rules and the centrality of an estimate of the threshold, k.

[^2]:    ${ }^{2}$ Strictly speaking, these local health authorities are Primary Care Organisations (PCOs) but the vast majority of these are 'Trusts' and we retain this terminology throughout.
    ${ }^{3}$ Due to data limitations the cited studies were only able to relate expenditure in period $t$ to mortality in periods $t, t-$ 1 , and $t-2$. Such studies assumed that PCTs had reached some sort of equilibrium in the expenditure choices they make and the outcomes they secure.

[^3]:    ${ }^{4}$ Due to data availability constraints previous studies had to relate expenditure in period $t$ to mortality data in periods $\mathrm{t}, \mathrm{t}-1$, and t -2. Implicitly this assumes that data represent a quasi long-run equilibrium position, and that relative expenditure levels and health outcomes within each PCT have been reasonably stable over a period of time.

[^4]:    ${ }^{5}$ Comparable data for each programme budget sub-category is shown in Table BA. 1 in Appendix B.
    ${ }^{6}$ These revisions are documented in Appendix B, Section B4.3.
    ${ }^{7}$ Expenditure on, for example, community care, A\&E, ambulance services, and outpatients can be difficult to attribute a particular PBC. Critical care, rehabilitation, and specialised commissioning across care settings will also be difficult to attribute to a particular programme.

[^5]:    ${ }^{8}$ This cost adjustment reflects the fact that health economy input prices vary considerably across the country and, for some inputs, are up to $40 \%$ higher in London and the south east of England than elsewhere. We have used a weighted average of the three Market Forces Factor Indices (MFFs) for HCHS, for prescribing, and for GMS/PMS to adjust the raw expenditure figures in Table 2 for local input prices (see Department of Health, 2009)
    ${ }^{9}$ This needs adjustment incorporates the AREA resource allocation formula for HCHS (see Department of Health, 2005).

[^6]:    ${ }^{12}$ Details of the construction of all instruments are shown in Table A. 2 of Appendix B.
    ${ }^{13}$ This incorporates the CARAN formula for HCHS and reflects need across all health care services.

[^7]:    Note: these statistics are unweighted across PCTs and reflect the values for these variables as ayailable for the regression analysis of PB expenditure data for 2007/8 and for 2008/9.
    Sources: Population Census 2001, Department of Health (2009), NHS Information Centre website.

[^8]:    ( )

    Note that the mortality data precedes expenditure in these models. This was due to data limitations at the time of the study.
    ${ }^{18}$ Initial modelling work employed the Department of Health's resource allocation model of the need for health care based on the AREA report (Department of Health, 2005c). Subsequent refinements and updates to this model employed the implementation of the CARAN model (Department of Health, 2009) and the initial findings of a Person Based Resource Allocation study (Dixon et al, 2011). The use of these alternative models for the need for health care were explored.
    ${ }^{19}$ An exception to this is expenditure on GMS/PMS (PBC23a) which is adjusted using the GMS/PMS market forces factor.

[^9]:    ${ }^{20}$ Unadjusted cost of a life year figures can be found in Appendix B, Table B21a.
    ${ }^{21}$ Refer to Appendix B, Table B8.23.

[^10]:    ${ }^{25}$ The Kleibergen-Paap F statistic is very close to the target value of ten for both the genitor-urinary and infectious diseases outcome models.
    ${ }^{26}$ These are endocrine: all service measure of need and diabetes prevalence rate; neurological: epilepsy prevalence; GMS/PMS: proportion of lone pensioner households; trauma/injuries: proportion of population working in agriculture.
    ${ }^{27}$ Full details of these calculations can be found in Tables B10.3 and B10.4 of Appendix B.

[^11]:    ${ }^{28}$ The CARAN measure of service need.
    ${ }^{29}$ The amendments are: respiratory diseases: all service need and all service need squared; endocrine: IMD07 and diabetes prevalence rate; genitor-urinary: lone parent households; infectious diseases: IMD07 and HIV need per head and its square; maternity and neonates: all service need and proportion born outside EU and proportion of population with no qualification aged 16 to 74 .

[^12]:    ${ }^{30}$ These are infectious diseases: HIV need and its square; endocrine: all service measure of need, its square and diabetes prevalence rate; genitor-urinary: all service measure of need and proportion of residence born outside EU; maternity/neonates: maternity measure of need; GMS/PMS: all service measure of need, proportion of residents reporting permanent sickness ( $16 \mathrm{yrs}-74 \mathrm{yrs}$ ), proportion of lone pensioner households and proportion in professional occupations; trauma/injuries: proportion of population working in agriculture.

[^13]:    ${ }^{31}$ Note that the apparent outcome elasticities -- 0.465 and 0.414 -- shown in the calculations in equations 3.7 and 3.8 are not pure outcome elasticities but incorporate both the expenditure and outcome elasticities. For pure outcome elasticities see row 13 of Table 6.
    ${ }^{32}$ Expenditure on, for example, community care, $\mathrm{A} \& E$, ambulance services, and outpatients can be difficult to attribute to a particular PBC. Critical care, rehabilitation, and specialised commissioning across care settings will also be difficult to attribute to a particular programme.

[^14]:    ${ }^{33}$ With the index for $1987 / 8$ set equal to 100 , then $2005 / 6=240.9,2006 / 7=249.8,2007 / 8=257.0$, and $2008 / 9=267.0$

[^15]:    ${ }^{34}$ Such studies assumed that PCTs had reached some sort of equilibrium in the expenditure choices they make and the outcomes they secure.

[^16]:    ${ }^{35}$ Recall that although 3 years of mortality data are used in the analysis of each year of expenditure, these are averaged to an annual value prior to estimating outcome elasticities. Therefore, the estimated outcome elasticities represent the proportionate effect on mortality in one year due to a proportionate change in expenditure.

[^17]:    ${ }^{38}$ The YLL available from NHS IC represented all deaths from maternity and all deaths under 28 days across PBCs. The coverage factor ( 0.68 in column 1 of Table 4.1) adjusts this YLL to represent maternity and all deaths $<1$ year across PBCs. The calculation is described in Appendix B, footnote (v) of Table B5.1.
    ${ }^{39}$ Figures for England, from http://www.ons.gov.uk/ons/rel/subnational-health4/life-expec-at-birth-age-65/2004-06-to-2008-10/statistical-bulletin.html\#tab-National-life-expectancy

[^18]:    ${ }^{40}$ This is the life expectancy that reflects the age distribution of the general population, i.e., the average of the sum of the life expectancies conditional on age, over the current age distribution. It will always be higher than life expectancy at birth.

[^19]:    The average of the sum of the YLLs for every observed death where the YLL for each observed death is the difference between age at death and LE conditional on age of death.
    ${ }^{50}$ In the absence of information about the age distribution of excess death this assumes that the average YLL associated with observed and excess deaths are similar. Insofar as excess deaths are thought likely to generate more YLL than observed deaths the number of excess deaths will tend to be overestimated. This would tend to underestimate the cost per excess death averted. However, the cost per life year estimates remain unchanged and do not require such an assumption.
    ${ }^{51}$ The impact of the age distribution of deaths and the age distribution of the at risk population (summarised as LE) on the calculation of excess deaths is not always obvious as both will affect the numerator (net YLL) as well the denominator (average YLL per death) in this calculation.

[^20]:    ${ }^{54}$ What portion of observed deaths are regarded as excess depend on how time is discretised. The data available reports deaths in annual intervals so in this context 'quickly' means within one year. If deaths were reported in narrower time intervals then a greater proportion of observed deaths would be regarded as excess and in the limit with continuous time all observed deaths would be excess. Of course, the average YLL associated with them would be smaller and is approximated by the net YLLs reported in Table 4.5 per observed death (the effects of approximation is likely to be small but unavoidable as it is due to deaths being reported in annual intervals). ${ }^{55}$ This is the same as life years associated with excess deaths since all observed deaths in this PBC are excess.

[^21]:    ${ }^{59}$ See Addendum 1 in Appendix C for a description on HSE data and section C2.2.1 of appendix C for the analysis of quality of life norms illustrated in Figure 4.1.
    ${ }^{60}$ The only exception is PBC11 (Respiratory) which has a large proportion of deaths occurring above the life expectancy of the PBC population (see Table 4.5).

[^22]:    ${ }^{61}$ ICD estimates of the quality of life score and age were pooled across datasets by considering the number of patients from each dataset contributing to estimates, i.e. a weighted average.
    ${ }^{62}$ The average quality of life scores across the ICDs which contribute to each PBC and the average age and gender of respondents were used to calculate a PBC disease related decrement based on quality of life norms from the general population. This 'PBC decrement' could then be applied to each observed death and the age at which each life year was gained or lost. In Section 4.4 information about the relative share of different types of disease (U-

[^23]:    ${ }^{67}$ Insofar as YLL would not have been lived in full health (see Section 4.3), the quality of life effects during disease must offset the less than full quality of life of the YLL to generate a ratio greater than one. Therefore, ratios less than one are possible even when disease has measurable quality of life effects for those experiencing it.
    ${ }^{68}$ The analysis in Section 4.3 already implies an $\mathrm{R}_{\text {death }}$ ratio at PBC level - see the following main text.

[^24]:    ${ }^{69}$ Reflecting the quality of life norms for the general population in Figure 4.1 and the distribution of ages and gender within each U-code (see Addendum 1 in Appendix C).
    ${ }^{70}$ Since quality of life effects of different disease states are expressed as age related decrements (see Figure 4.2) we fonot require the HODaR and MEPS samples to necessarily be representative of the age distribution of the population at risk in the groups of ICD codes that make up each U-code.
    $\int_{\frac{7}{1}}{ }^{1}$ The average quality of life scores across the ICDs which contribute to each U-code (see Addendum 1 for how ICD codes map to U-codes) and the average age and gender of respondents from HODaR and MEPS were used to calculate a disease decrement for each U-code, based on quality of life norms from the general population. These U-code disease decrements can then be applied to the age and gender distribution of each U-code, based on information from GBD about the prevalence and age distribution of each - using information about the incidence of sequelae associated with them (as described in Section 4.2.3) and information about the durations of disease (see Table C. 22 Appendix C).
    ${ }^{72}$ For example, the evidence about quality of life from HODaR and MEPS suggests that the impact of U037 on quality of life is greater than indicated by DALY disability weights. The quality of life effects of U141, although still very significant, are lower than indicated by DALY disability weights.

[^25]:    ${ }^{73}$ Information about the size and age and gender distribution is only available at U-code level. Therefore U-code ratios are applied to all the ICD codes that contribute to a particular U-code. Note that, unlike ICD codes, U-codes do not map directly to PBCs so some ICDs in different PBCs may belong to the same U-code and therefore have the same U-code ratio. Some ICDs are not included in the U-code classification of disease. Most of these are procedural codes where we do not assign life year and QALY effects anyway (any health effects would be evident in other ICD codes), so it was not necessary to impute ratios for them (84 out of 1562). Of the others most were associated with PBC16 with a zero outcome elasticity so did not require imputation either (186 out of 1562). Imputation based on the median ratio across the ICDs within the PBC was required for the remaining (482 out of 1562). Eighty eight of these cannot be mapped into U-codes. The remaining 394 were associated with U-codes where the ratio was undefined because the denominator (YLL) was zero. In both these cases, values were imputed based on the median ratio across the ICDs within the PBC. Since the distribution of ratios within a PBC tend to be highly positively skewed, imputation based on the median is likely to be conservative with respect to health effects and especially in the latter case where mortality effects appear to be a much less important aspect of the disease.
    ${ }^{74}$ It is important to note that it would be inappropriate to calculate an average of the ratios within a PBC and then apply this 'average ratio' to life year effects at PBC level, rather than calculate QALY effects at ICD level by applying the relevant ratio. The results, however, can be presented as an implied PBC ratio (i.e., a ratio of averages), see Table C. 43 in Appendix C.

[^26]:    ${ }^{77}$ Recall that in Section 4.3 each life year gained could be assumed to be lived in full health, lived in a quality of life that reflects age and gender norms of the general population or lived in a quality of life that reflects the original disease state. Using measures of QALY burden means that disease decrements are assigned to any life years gained during the duration of disease, i.e., insofar as more patients survive as a consequence of increased expenditure they do so in the diseased state for the remaining duration of the disease. Quality of life norms are assigned for any life

[^27]:    ${ }^{89}$ The cost per life year threshold in Table 4.9 can be interpreted as cost per QALY thresholds conditional on the assumption that all life years are lived in full health and the quality of life with disease is zero (equivalent to death).
    ${ }^{90}$ Note that the proportionate difference between the estimates in column 3 and columns 1 and 2 are greater in lines 1 and 2, reflecting the additional health effects from considering the likely impact of changes in expenditure on quality of life during disease. These differences are less marked in line 3 because the effects in those PBCs where an outcome elasticity can be estimated are extrapolated to the other PBCs using proportionate effect on QALY burden and measures of QALY burden in these other PBCs (see the discussion in Section 4.4.2 for a more details).

[^28]:    ${ }^{91}$ It would be inappropriate to assign all the change in GMS expenditure to the estimate of cost per QALY based only on the 11 PBCs with outcome elasticities because it would imply that GMS only contributes to these PBCs. Restricting attention to the 11 PBCs with outcome elasticities but allocating part of the change in GMS expenditure to them based on their proportional share of changes in overall expenditure would yield a slightly higher cost per QALY than reported in line 2.

[^29]:    ${ }^{92}$ The cost per life year threshold in column 1 can be interpreted as cost per QALY thresholds conditional on the assumption that all life years gained or lost are lived in full health but the quality of life with disease is zero (equivalent to death).
    ${ }^{93}$ The cost per life year adjusted for quality of life in column 2 can be interpreted as cost per QALY threshold conditional on the assumption that the quality of life with disease is zero (equivalent to death); effectively ignoring any effects on those who survive with disease.

[^30]:    ${ }^{108}$ Rather than solve for this type of 'certainty equivalent' a probabilistic analysis of the cost-effectiveness of a technology which integrated the uncertainty associated with the cost per QALY threshold as well, would take account of these issues, i.e., the technology would be cost-effective if it offered the highest expected net benefit when averaged over all Monte Carlo simulations, including sampling from the distribution of the cost per QALY threshold.
    ${ }^{109}$ Although health benefits can be expressed in terms of consumption (in money) using some consumption value of the health effects (willingness to pay), NHS costs must be first converted into health forgone, using an uncertain estimate of the threshold, before these are also expressed in consumption (money terms) using the same consumption value of health, i.e., the non linear effect of the threshold remains unavoidable. Failure to account for the threshold and the implications of its uncertainty would only be reasonable in a heath care system where expenditure was not constrained and/or all costs fell on private consumption.

[^31]:    ${ }^{110}$ What can be estimated is the health effect over the observed variation in expenditure. This will also be the 'true' marginal effect (tangency at a budget of B1) if health returns to expenditure diminish at a constant rate (the second derivative is constant) as illustrated in Figure 5.3. Since nothing is 'truly' marginal the important question is how the threshold changes with the sign and scale of the non marginal budget impact associated with approval of a new technology.

[^32]:    ${ }^{111}$ Due to the diminishing marginal returns illustrated in Figure 5.4 (see section 5.5 for further explanation).

[^33]:    ${ }^{1122008}$ expenditure expressed in 2007 NHS prices based on $3.9 \%$ NHS inflation from the HCHS index - see Section B11.5 in Appendix B.

[^34]:    ${ }^{113}$ See Table C55 and C82 in Appendix C for a summary of outcome and expenditure elasticities and total expenditure by PBC in 2007 and 2008. Also compare Table C80 in Appendix C to Table 5.2 above for an indication of these net effects on the share of health effects and changes in expenditure.
    ${ }^{114}$ If the growth rate in the nominal threshold between 2007 and 2008 was applied the current 2012 threshold would be expected to be $£ 16,895$
    ${ }^{115}$ See above

[^35]:    ${ }^{119}$ This is implicit in the estimates of outcome elasticities presented in Chapter 3. Recall that although 3 years of mortality data are used in the analysis of each year of expenditure, these are averaged to an annual value prior to estimating outcome elasticities. Therefore, the estimated outcome elasticities represent the proportionate effect on

[^36]:    ${ }^{124}$ These effects will be picked up in the cross sectional variation, at least partially, so long as there is some variation in the health effects achieved and scale of simultaneous investment or disinvestment across PCTs.
    ${ }^{125}$ This would be particularly interesting when re-considering the subgroup analysis in Section 5.5 with panel data. ${ }^{126}$ Recall that we have taken account of competing risks or counterfactual deaths (which might appear in any of the PBCs) in our calculation of net years of like lost - see Section 4.2.3)

[^37]:    ${ }^{127}$ Recall that no health effects where attributed to changes in GMS expenditure (PBC23).
    ${ }^{128}$ The vast majority of hip and knee replacements are for osteoarthritis which is included in PBC15

[^38]:    ${ }^{1}$ This is the aim of the new Value Based Pricing approach currently under development by the Department of Health [10, 44]

[^39]:    ${ }^{2}$ In fact in the 2004 NICE Methods Guide [30] noted that "the threshold will change over time as the budget for healthcare changes" (p. 33). However, there is no clear reference to this change in the 2008 Methods Guide [31].

[^40]:    ${ }^{1}$ This study builds on previous work that was undertaken as part of the Quest for Quality and Improved Performance, a five-year initiative of the Health Foundation.

[^41]:    ${ }^{2}$ Strictly speaking, these local health authorities are Primary Care Organisations (PCOs) but the vast majority of these are 'Trusts' and we retain this terminology throughout.
    ${ }^{3}$ In April 2010 two PCTs (East \& North Hertfordshire (5P3) and West Hertfordshire (5P4)) merged to form a single organisation (Hertfordshire PCT (5QV)) so that, since this date, there have been 151 PCTs. At the same time Blackburn and Darwen PCT (5CC) became Blackburn and Darwen Teaching Care Trust Plus (TAP). In April 2011 Solihull Care Trust (TAM) became a PCT (5QW).

[^42]:    ${ }^{4}$ Some commentators have suggested that some of the within programme variation in expenditure observed across PCTs reflects different accounting conventions or unknown local factors. One way of reducing the impact of such unobserved heterogeneity is to construct a longitudinal data set with expenditure and mortality for each PCT for several years. With the availability of several years of data for both expenditure and mortality, we wanted to estimate a panel data model. However, most of the instruments employed here are based on the 2001 Census and thus estimation of a panel model will not be possible until these too become time variant; this should occur later this year with release of the 2011 Census data at PCT level. The same difficulty arises with the estimation of an incremental model.
    ${ }^{5}$ This figure ignores intra-category changes (for example, where an ICD10 code is re-allocated from category 1A to 1B) and only counts cross-category changes (for example, where the code is switched from category 1 to category 2). ${ }^{6}$ This expert review also led to the introduction of 40 additional sub-categories including 10 sub-categories for the cancer and tumour programme.

[^43]:    ${ }^{7}$ Expenditure on, for example, community care, A\&E, ambulance services, and outpatients can be difficult to attribute a particular PBC. Critical care, rehabilitation, and specialised commissioning across care settings will also be difficult to attribute to a particular programme.

[^44]:    ${ }^{8}$ This cost adjustment reflects the fact that health economy input prices vary considerably across the country and, for some inputs, are up to $40 \%$ higher in London and the south east of England than elsewhere. We have used a weighted average of the three Market Forces Factor Indices (MFFs) for HCHS, for prescribing, and for GMS/PMS to adjust the raw expenditure figures in Table 4.2 for local input prices (see Department of Health, 2009).
    ${ }^{9}$ This needs adjustment incorporates the AREA resource allocation formula for HCHS (see Department of Health, 2005c).

[^45]:    ${ }^{11}$ One exception to this is the mortality rate for the trauma and injuries programme where initially only SMRs were available.
    ${ }^{12}$ The NHS IC reports mortality rates using deaths pooled over a three year period because the relatively small number of annual deaths in some disease categories might lead to large year-on-year fluctuations in death rates at PCT level.

[^46]:    Note: these statistics are unweighted across PCTs and reflect the values for these variables as available for the regression analysis of PB expenditure data for $2007 / 8$ and for $2008 / 9$.
    Sources: Population Census 2001, Department of Health (2009), NHS Information Centre website.

[^47]:    ${ }^{13}$ However, we do experiment with replacing and supplementing this all service measure of need with more programme specific measures where these are available (e.g., using the diabetes and epilepsy prevalence rates).
    ${ }^{14}$ Whilst need is a function of mortality/morbidity in the resource allocation formula, the relationship is not sufficiently strong enough for us to be concerned about the endogeneity of the need in any individual care programme.
    ${ }^{15}$ When estimating expenditure equations using PB data for 2005/6 for cancer and circulatory disease we persevere (for continuity with previous studies) with the use of the circulatory disease SYLLR as the proxy for other programme need in the cancer programme, and we use the cancer SYLLR as the proxy for other programme need in the circulatory disease programme (see Martin, Rice and Smith, 2008a \& 2012).

[^48]:    ${ }^{16}$ The IV procedure involves the estimation of the second-stage expenditure equation as specified in equation 6.1 and the estimation of a first-stage expenditure equation associated with equation 6.2. The same variable might have different coefficients in these two equations because the equations will have different sets of covariates.

[^49]:    ${ }^{17}$ For the case of a single endogenous regressor and three excluded instruments, Stock and Yogo (2002) critical values are as follows in term of the bias of 2SLS relative to bias of OLS as follows: relative bias $5 \%$ critical value $=$ 13.9; relative bias $10 \%$, critical value $=9.08$; relative bias $20 \%$, critical value $=6.46$; relative bias $30 \%$, critical value $=5.39$.
    ${ }^{18}$ The OLS version of Ramsey's reset test was invoked using Stata's -ovtest- command, and the IV equivalent was invoked using -ivreset-.
    ${ }^{19}$ As all PCTs face the same prescribing costs, the prescribing MFF is 1 for all PCTs.

[^50]:    ${ }^{20}$ The 'big four' programmes are the cancer, circulatory disease, respiratory problems, and gastro-intestinal problems programmes. They are 'big' programmes in terms of the number of deaths associated with each programme.

[^51]:    ${ }^{0.111 *}$
    0.158]

[^52]:    ${ }^{21}$ The programme specific cost per life and life year estimates presented here will underestimate the true programme specific costs because not all PCT expenditure can be allocated to a specific programme (for example, all GMS expenditure is allocated to PBC23 rather than being split between cancer, circulatory disease, respiratory problems, etc). However, this more generic expenditure is incorporated into the calculation of the cost of a life year when this calculation is undertaken across all programmes.
    ${ }^{22}$ These are the 'big four' programmes in terms of the number of lives (or life years) lost.

[^53]:    ${ }^{26}$ Note that implied need=unified weighted population/(CARAN MFF* raw population).

[^54]:    Notes: (a) iAREA need $1=$ AREA unified weighted population/HCHS MFP

[^55]:    Notes: (a) iAREA need $1=$ AREA unified weighted population/HCHS MFF

[^56]:    Notes: (a) iAREA need $1=$ AREA unified weighted population/HCHS MFF

[^57]:    ${ }^{27}$ Ideally, the test F statistic should equal to or greater than ten.

[^58]:    ${ }^{28}$ Clearly, some expenditure in year $t$ will have an effect on mortality beyond $t+2$ but we have no mortality data that would allow us to include this in our modelling work. We must assume that, for expenditure that affects mortality beyond $t+2$, PCTs have reached some sort of equilibrium position in terms of their expenditure choices and the outcomes secured.

[^59]:    ${ }^{29}$ When re-estimating the all PCT model for 'high spenders' and then for 'low spenders' no attempt was made to adjust the estimating equation for any implied model mis-specifiaction.
    ${ }^{30}$ The cost of a life year estimates presented in Table 8.28 are not adjusted for the mismatch in the ICD10 coverage of the expenditure and mortality data because such an adjustment would not affect our conclusions.

[^60]:    Note: 'high spending' PCTs are those whose predicted spend per person is greater than the average predicted spend per person (ceteris paribus), and

[^61]:    Note that for those over target, the average amount (percentage) is $£ 13.415 \mathrm{~m}(3.6 \%)$; for those under target, the average amount (percentage) is $£ 10.575 \mathrm{~m}(2.6 \%)$

[^62]:    Note that the adjustment for the coverage of the Y1L data relative to the spend data uses deaths under age 75 in England in 2008.

[^63]:    ${ }^{31}$ See column 1 of Table 8.19 for the estimated IV cancer outcome model.
    ${ }^{32}$ We used a symmetric distribution about zero because we have no priors about the signs of the coefficients on the instruments. The use of a uniform distribution is arbitrary but of no significance.

[^64]:    ${ }^{33}$ The outcome model for circulatory disease reported in Table B8.19 (using PB expenditure for 2006/7 and mortality data for $2006 / 7 / 8$ ) contains four instruments. The application of the sensitivity analysis described in this section is considerably easier to implement if only two instruments are present and re-estimation of the outcome model for circulatory disease without the two least significant instruments generates very similar results to those obtained with all four instruments (for example, the coefficient on expenditure declines marginally from - 1.434 to 1.427). Therefore the sensitivity analysis reported here uses the outcome model containing only two instruments.

[^65]:    ${ }^{34}$ The Kleibergen-Paap F statistic is very close to the target value of ten for both the genitor-urinary and infectious diseases outcome models.

[^66]:    (ii) the addition of unpaid carers as an instrument for the endoctine outcome model generates a Hansen-Sargen test statistic of 0.372 ( p -value 0.5418 ) and the coefficient on expenditure is -0.423 .

[^67]:    Note: robust standard errors in brackets, *** $\mathrm{P}<0.01,{ }^{* *} \mathrm{p}<0.05,{ }^{*} \mathrm{p}<0.1$

[^68]:    Note: robust standard errors in brackets, ${ }^{* * *} \mathrm{p}<0.01,{ }^{* *} \mathrm{p}<0.05,{ }^{*} \mathrm{p}<0.1$

[^69]:    ${ }^{35}$ Note that the apparent outcome elasticities -- 0.465 and 0.414 -- shown in the calculations (11.1) and (11.2) are not pure outcome elasticities but incorporate both the expenditure and outcome elasticities. For pure outcome elasticities see row 13 of Table B11.5.
    ${ }^{36}$ Expenditure on, for example, community care, $\mathrm{A} \& E$, ambulance services, and outpatients can be difficult to attribute a particular PBC. Critical care, rehabilitation, and specialised commissioning across care settings will also be difficult to attribute to a particular programme.

[^70]:    ${ }^{37}$ With the index for $1987 / 8$ set equal to 100 , then $2005 / 6=240.9,2006 / 7=249.8,2007 / 8=257.0$, and 2008/9=267.0 (Curtis, 2011, p209).

[^71]:    Note: these are the first-stage regressions for the IV models reported in Table B10.1. Robust standard errors in brackets, ${ }^{* * *} \mathrm{p}<0.01,{ }^{* *} \mathrm{p}<0.05,{ }^{*} \mathrm{p}<0.1$

[^72]:    Note: these are the first-stage regressions for the models reported in Table B10.2. Robust standard errors in brackets, ${ }^{* * *} \mathrm{p}<0.01$, ${ }^{* *} \mathrm{p}<0.05$, ${ }^{*} \mathrm{p}<0.1$

[^73]:    Note: these are the first-stage regressions for the IV models reported in Table B11.1. Robust standard errors in brackets, *** $\mathrm{p}<0.01, * * \mathrm{p}<0.05$, , $\mathrm{p}<0.1$

[^74]:    ${ }^{2}$ The YLL available from NHS IC represented all deaths from maternity and all deaths under 28 days across PBCs. The coverage factor ( 0.68 ) adjusts this YLL to represent maternity and all deaths $<1$ year across PBCs. The calculation is described in Appendix B, footnote (v) of Table B5.1.
    ${ }^{3}$ Figures for England, from http://www.ons.gov.uk/ons/rel/subnational-health4/life-expec-at-birth-age-65/2004-06-to-2008-10/statistical-bulletin.html\#tab-National-life-expectancy.

[^75]:    ${ }^{4}$ Note that the outcome elasticities are based on PBC mortality that is sensitive to changes in expenditure (i.e., is avoidable) at the margin so no assumptions about how much of the PBC mortality is avoidable is required.

[^76]:    5 Although this research was not funded to purchase access to GPRD data we were able to examine a sample of it which comprised of $22,313,086$ rows/patient-ICD10 events (3 digit) representing 4,229,910 patients with data on new diagnosis of diseases observed between 1 Jan 2006 and 24 June 2011 (see Addendum 1). Although GPRD data could, in principle, provide this type of information the difficulties of reliability, face validity and interpretation of the sample data in the form available to us meant that it was not directly useful.
    ${ }^{6}$ We are aware that the 2000-2002 WHO GBD study and the update which was published in 2008 using 2004 data has itself recently been updated. However, the report and tools where not publically available at the time this research was conducted.
    ${ }^{7}$ Throughout the analyses in this Appendix, mortality, life years and QALY were not assigned to procedural ICD codes (i.e. those in ICD chapter Z, Factors influencing health status and contact with health services). Health effects from increased spending on these ICD codes would either be non-existant or would be evident in other ICD codes related to the procedure.

[^77]:    ${ }^{9}$ This information is also used in Section C.2.3.

[^78]:    ${ }^{10}$ Indeed, since some of the variation in mortality in 1st year that is not sustained to the 3rd year will nevertheless be sustained for 1 or 2 years, 2 life years per death averted represents somewhat less than the minimum, consistent with restricting live years gain to the observed mortality data.

[^79]:    ${ }^{12}$ Note that this is the ratio of total change in health to total change in expenditure across these PBC (rather than an average ratio) and the contribution that each of these PBCs make to these total effects on health and expenditure depends on the estimated expenditure as well as outcome elasticities.
    ${ }^{13}$ Indeed, applying the absolute health effect of expenditure from the 11 PBCs with outcome elasticities implies different (higher) proportionate effects in the other PBCs

[^80]:    ${ }^{14}$ Note that the proportionate difference between the estimates in column 3 and columns 1 and 2 are greater in lines 1 and 2, reflecting the additional health effects from considering the likely impact of changes in expenditure on quality of life during disease. These differences are less marked in line 3 because the effects in those PBCs where an outcome elasticity can be estimated are extrapolated to the other PBCs using proportionate effect on QALY burden and measures of QALY burden in these other.

[^81]:    ${ }^{15}$ Which are determined by the estimated expenditure elasticities (the proportionate change in PBC expenditure due to a change in overall expenditure) and total PBC expenditure (see Chapter 3 and section B11 in Appendix B).

[^82]:    ${ }^{16}$ Which are determined by the outcome elasticities (the proportionate effects on mortality and YLL of a proportionate change in PBC expenditure (see section C.2.3 for details of how these estimates can be applied to measures of QALY burden in all PBCs).
    ${ }^{17}$ See section C.2.3 for how PBC level effects can be allocated to the contributing ICD codes and how measures of QALY burden for each ICD code can be established

[^83]:    ${ }^{18}$ Insofar as measures of contribution to variance based on HES data (see Section C.2.3.2) will tend to introduce a bias against those ICD codes where costs are more likely to be recorded in primary care and community services (e.g., more common mental health problems such as depression) then the potential underestimation of health effects is likely to be greater (since these interventions appear more cost effective) and the likelihood that the overall threshold of $£, 15,701$ is overestimated will tend to be greater

[^84]:    ${ }^{19}$ Prof. Craig Currie and Sara Jenkins-Jones
    ${ }^{20}$ This represents six fewer than the incidence data as in these instances the end dates for the disease were beyond the end of the data collection period.

[^85]:    ${ }^{21}$ Mapping algorithms were provided by the NHS Connecting for Health group, see
    http://www.connectingforhealth.nhs.uk/systemsandservices/data/clinicalcoding/crossmap for more details
    ${ }^{22}$ Representing instantaneous, one month, one year, five years and life-long
    ${ }^{23}$ For more information on access to the Toolkit see
    http:/ /www.who.int/healthinfo/global_burden_disease/tools_nbd_toolkit/en/index.html

[^86]:    ${ }^{24}$ For more information on the surveys and the data they collect see http://www.dh.gov.uk/en/Publicationsandstatistics/PublishedSurvey/HealthSurveyForEngland/index.h tm

[^87]:    ${ }^{25}$ from http://www.yhpho.org.uk/resource/view.aspx?RID=49488

[^88]:    ${ }_{26}$ Available at http://www.rightcare.nhs.uk/atlas/index.html

[^89]:    ${ }^{27}$ This contrast was informed by our clinical representative
    ${ }_{28}$ Available at http://www.nhs.uk/Pages/HomePage.aspx accessed on 10/10/2011

[^90]:    ${ }^{29}$ See: http://www.dispensingdoctor.org/content.php?id=1335 accessed 03/05/2012
    ${ }^{30}$ Estimate by Tim Kendal

[^91]:    ${ }^{31}$ This view was informed by our clinical advisors
    ${ }^{32}$ This view was informed by our clinical advisors

