Genetic analysis of a wild-caught hybrid between non-sister *Heliconius* butterfly species

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Interspecific hybridization occurs regularly in wild Heliconius butterflies, although hybrid individuals are usually very rare. However, hybridization generally occurs only between the most closely related species. We report a rare naturally occurring hybrid between non-sister species and carry out the first genetic analysis of such distant hybridization. Mitochondrial and nuclear genes indicate that the specimen is an F₁ hybrid between a female Heliconius ethilla and a male Heliconius melpomene, originating from a group of 13 species estimated to have diverged over 2.5 Myr ago. The presence of such distant natural hybrids, together with evidence for backcrossing, suggests that gene flow across species boundaries can take place long after speciation. Adaptive genes such as those involved in wing coloration could thus be widely shared among members of this highly mimetic genus.

Keywords: *Heliconius ethilla*; *Heliconius melpomene*; hybridization

1. INTRODUCTION

Evolutionary biologists generally accept the biological species concept, and this has led to hybridization between species being regarded as uncommon and unimportant (Mayr 1963). Although hybrids are, almost by definition, very rare on a per individual basis, recent surveys have shown that many species do in fact hybridize (Coyne & Orr 2004; Mallet 2005). On average, 10% of animal species and 25% of plant species are known to hybridize in the wild, and in some particularly charismatic groups, such as ducks, birds of paradise and North American *Papilio* butterflies, 40–75% of species are known to hybridize (Mallet 2005).

Many wild-caught interspecific hybrid specimens between *Heliconiina* butterflies are known. Hybrids occur in 35% of the 46 species in the genus *Heliconius* and 27% of the 73 species in the larger sub-tribe Heliconiinae (Mallet *et al.* 2007). These butterflies have been well studied owing to their bright wing coloration and extensive variation in wing pattern

Electronic supplementary material is available at http://dx.doi.org/ 10.1098/rsbl.2007.0401 or via http://www.journals.royalsoc.ac.uk. morphology both between species and among geographical races within species. The bright wing colours act as a warning of their unpalatability to potential predators. Many species share similar patterns with unrelated Heliconiinae and Ithomiinae, leading to impressive Müllerian mimicry rings (Bates 1862; Turner 1981; Beccaloni 1997; Joron & Mallet 1998). Although hybrids are rare within most *Heliconius* populations, the existence of naturally occurring hybrids is intriguing because it suggests that gene flow will be possible between species. Hence, this could lead to the transfer of adaptive genes between species (Gilbert 2003).

Hybridization in *Heliconius* has hitherto been examined genetically only in a few cases that involve closely related species: *Heliconius erato* and *Heliconius himera*, or *Heliconius melpomene* and *Heliconius cydno* (Jiggins *et al.* 1997; Bull *et al.* 2006; Kronforst *et al.* 2006). In all other cases, inferences of hybridization have been made on the basis of phenotypes of wildcaught specimens. In contrast, the extremely rare hybrids between *H. melpomene* and the more distant species in the 'silvaniform' *Heliconius* (Mallet *et al.* 2007) have never been examined genetically. In this paper, we describe a new rare hybrid formed between two distantly related *Heliconius* species in the *H. melpomene* group, recently caught in Peru, and use molecular markers to determine the parental types.

2. MATERIAL AND METHODS

We are engaged in an extensive study of heliconiine and ithomiine butterflies in the Departamento de San Martín, Peru (Mallet & Barton 1989; Joron *et al.* 2001; Whinnett *et al.* 2005). In addition to the pure species, subspecies and many interracial hybrids sampled, we recently caught a peculiar specimen, a putative interspecific hybrid from the *melpomene*-silvaniform group (specimen ID 06-921; table 1) near Moyobamba in Northern Peru. The hybrid specimen will be deposited in the collection at the Museo de Historia Natural, Universidad Nacional Mayor de San Marcos (UNMSM), Lima, Peru. The remaining specimens analysed (table 1) are of common species, held at University College London and vouchered at UNMSM.

DNA was extracted from a single leg of the hybrid specimen using a QIAamp DNA Micro Kit (QIAGEN), and from one-third of the thorax of all other specimens using the DNeasy Blood and Tissue Kit (QIAGEN). Approximately 2200 bp of mtDNA comprising CoI and 5' end of CoII genes were amplified by PCR in three sections for the hybrid individual, representatives of Heliconius melpomene amaryllis, Heliconius elevatus and each of the four sympatric silvaniform species: Heliconius numata; Heliconius ethilla; Heliconius pardalinus; and Heliconius hecale. In addition, five nuclear loci (Tektin, Rpl5, Tpi, Mpi and inv) were amplified in the hybrid, representatives of H. melpomene and H. ethilla. PCR products were cleaned and cycle sequenced with the PCR primers using the Big DYE TERMINATOR v. 3.1 Cycle Sequencing Kit (Applied Biosystems). Table S1 of the electronic supplementary material contains details of the primers used, PCR conditions and sequence accession numbers.

Apart from mtDNA and *Tektin*, all other loci spanned one or more intron regions. Indels in the intronic regions often resulted in the amplification of alleles with different sizes from a single individual. Unless sequence quality was low, the indel could be readily identified and the two alleles deconvoluted using information from the double-peak signals following the indel (Flot *et al.* 2006). See the electronic supplementary material for details of this procedure.

3. RESULTS

The putative hybrid (figure 1) is a male, and its wing pattern is unlike that of other local *melpomene*-silvaniform group species. Instead, the markings appear to combine those of *H. m. amaryllis* (the local 'postman'-patterned *melpomene* race) and of a local

specimen ID	species	collection locality	latitude	longitude
06-921	Heliconius hybrid	Rumiyacu, near Moyobamba	06°05′23″ S	076°58′09″ W
04-286	Heliconius melpomene	Bosque von Humboldt	08°49′48″ S	075°03′28″ W
04-288	Heliconius melpomene	Bosque von Humboldt	08°49′48″ S	075°03′28″ W
02-366	Heliconius melpomene	Davidcillo	06°14′47″ S	076°15′58″ W
02-944	Heliconius melpomene	Puente Serranayacu	05°40′48″ S	077°40′50″ W
02-1839	Heliconius melpomene	Boca Toma Río Shilcayo	06°27′20″ S	076°20'40″ W
02-1850	Heliconius melpomene	Shapaja	06°34'29″ S	079°16'48″ W
02-1882	Heliconius melpomene	Chumia	06°36'57″ S	076°11'07″ W
02-1894	Heliconius melpomene	km 30, Tarapoto-Yurimaguas	06°24'33″ S	076°18'24″ W
02-2060	Heliconius melpomene	km 26, Yurimaguas-Tarapoto	05°58'30″ S	076°14'15″ W
02-3	Heliconius ethilla	Boca Toma Río Shilcayo	06°27'20″ S	076°20'40″ W
02-975	Heliconius ethilla	km 10, Tarapoto-Yurimaguas	06°27'18″ S	076°17'46″ W
02-1483	Heliconius ethilla	La Antena, km 16, Tarapoto-Yurimaguas	06°27'19″ S	076°17'54″ W
02-2037	Heliconius numata	km 26, Yurimaguas-Tarapoto	05°58'30″ S	076°14'15″ W
02-364	Heliconius elevatus	Davidcillo	06°14'47″ S	076°15'58″ W
05-1196	Heliconius pardalinus	Urahuasha	06°27'43″ S	076°19'36″ W
02-1330	Heliconius hecale	km 7.2, Pongo-Barranquita	06°17'41″ S	076°13'53″ W

Table 1. Details of samples and collection localities.



Figure 1. (a) Dorsal and (b) ventral wing colour patterns of the hybrid specimen 06-921. Dorsal wing colour patterns of the putative parent species (c) H. ethilla aerotome and (d) H. melpomene amaryllis.

silvaniform species: H. numata; ethilla; hecale; or pardalinus. Compared with H. m. amaryllis, the forewing crimson colours have become burnt orange and these orange markings extend over both fore- and hindwings, rather than being restricted to a forewing bar. The yellow hindwing bar of the latter species has also become orange on the upperside, although the yellow is almost fully expressed on the underside. Compared with Heliconius ethilla aerotome (identified as the other parent, see below), the orange markings are much reduced, and on the hindwings narrowed into 'rays' reminiscent of those found in races of H. melpomene such as Heliconius melpomene aglaope, which occurs over the mountain range in the Amazonian lowlands to the northeast of the capture site. The spotty melanic markings of H. ethilla are also broadened in the putative hybrid to form black smears, particularly in the central part of the forewing. Apart from the underside-expressed yellow hindwing bar, the underside and upperside patterns are similar. This hybrid is similar to a specimen in the Natural History Museum, London, originally named as a separate species '*Heliconius hippola*' Hewitson (Mallet *et al.* 2007).

A BLAST search of the mtDNA sequence revealed 99.5% similarity to *H. ethilla*. Subsequent comparison with mtDNA sequences obtained from locally caught specimens of the four potential silvaniform species showed unambiguously that the hybrid possessed a *H. ethilla* mitochondrial sequence (figure 2; table S2, electronic supplementary material). Diagnostic sequence differences between *H. melpomene* and *H. ethilla* were not



Figure 2. Neighbour-joining trees for six loci showing the relationship between alleles found in the hybrid and other melpomene–cydno–silvaniform group species. Nodes with 50% or greater bootstrap support are labelled. Bold vertical lines indicate diagnostic indels between *H. melpomene* and *H. ethilla*. Scale bars represent percentage sequence divergence. Asterisks, additional information in the electronic supplementary material.

found at *Rpl5*, but were present at the other four nuclear loci. The hybrid individual was found to possess both a *melpomene-* and an *ethilla-*type allele at each of these four diagnostic nuclear loci (figure 2).

An F_1 hybrid should be heterozygous at all nuclear genes, bearing alleles of each parental type. In contrast, a backcross hybrid should only be heterozygous at half its nuclear genes and homozygous for one parental type over its remaining nuclear genes. As the hybrid is heterozygous at four diagnostic nuclear loci, this indicates that the individual is unlikely to be a backcross or F_2 hybrid (binomial $p=0.5^4=0.0625$); an F_1 hybrid between *H. ethilla* and *H. m. amaryllis* is 16 times more likely than any second-generation hybrid.

4. DISCUSSION

Although interspecific hybridization is a common phenomenon among heliconiine butterflies, at the individual level, hybridization is rare, usually comprising less than 1 in 1000 wild individuals (Mallet *et al.* 2007). Most wild hybrids are either intraspecific

(between different wing pattern races) or between closely related, usually sister taxa (Mallet *et al.* 2007). Only 10 putative hybrid specimens have been documented between *melpomene* and silvaniform species, and only four are putatively between *H. melpomene* and *H. ethilla* (Mallet *et al.* 2007). However, no molecular verification of these rare and distant hybrids has hitherto been carried out, so the identity of the parents and whether they are in fact hybrids or aberrations is in doubt (Mallet *et al.* 2007). Here, genetic evidence for natural hybridization between such distant non-sister *Heliconius* species (*H. melpomene* and *H. ethilla*) has been obtained for the first time.

Heliconius ethilla and H. melpomene are approximately 5% different at the mtDNA studied. Assuming that this gene evolves in a clock-like manner at 2% Myr⁻¹ (Brower 1994), this suggests hybridization events occurring *ca* 2.5 Myr after speciation has occurred. The occurrence of a wild adult F_1 hybrid between these species indicates that such hybrids can develop normally and survive in the wild. Some putative wild hybrids between *melpomene* and silvaniform group butterflies are considered to be backcrosses

(Mallet et al. 2007), providing evidence that some such hybrids may be fertile and capable of reproduction. Two of the four putative hybrids between H. melpomene and *H. ethilla* known from Colombia are clearly not F_1 hybrids, and are presumably backcrosses to H. ethilla (Mallet et al. 2007). Although those putative hybrids have not been analysed genetically, the more ethilla-like patterns, the strong expression of yellow coloration in the forewing (which is recessive in hybrids) and strongly ethilla-like hindwing pattern all provide clear evidence of backcrossing to ethilla in nature. The other two hybrids are *H. hippola*-like and are presumably F_1 hybrids (Mallet et al. 2007). Backcrosses via male F_1 hybrids in a complex cross involving *H. melpomene* and the silvaniforms H. hecale and H. atthis (the latter close to H. ethilla) have been obtained in captivity, although female hybrids were reported to be sterile (Mallet et al. 2007), indicating that such backcrossing is possible. This potential for backcrossing may result in transfer of genes between species.

As hybridization is regular, species boundaries in Heliconius have the potential to be porous. Within the melpomene-cydno group, hybridization and backcrossing has led to interspecific introgression at some, but not all, genomic regions (Bull et al. 2006; Kronforst et al. 2006), and has apparently produced at least one hybrid species (Mavárez et al. 2006). If extensive hybridization among these two closely related species can cause adaptive genes to introgress as a result of hybridization, rarer hybridization between more distant species, such as between the silvaniform and the *melpomene-cydno* group species, as here, may also play a role. Closely related Heliconius species are often members of different colour pattern mimicry rings, and very similar and apparently homoplasious mimetic patterns are often found between related non-sister species; for instance, H. elevatus (a silvaniform) shares almost identical ray patterns with some races of H. melpomene and Heliconius timareta. Heliconius besckei (also a silvaniform), in contrast, shares a melpomene amaryllis-like postman pattern with races of H. melpomene and H. timareta. Our data therefore contribute evidence for the intriguing possibility that mimetic wing patterns may be shared via introgression among distant as well as closely related species (Gilbert 2003; Mallet et al. 2007).

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