Disruptive sexual selection against hybrids contributes to speciation between Heliconius cydno and Heliconius melpomene

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Understanding the fate of hybrids in wild populations is fundamental to understanding speciation. Here we provide evidence for disruptive sexual selection against hybrids between Heliconius cydno and Heliconius melpomene. The two species are sympatric across most of Central and Andean South America, and coexist despite a low level of hybridization. No-choice mating experiments show strong assortative mating between the species. Hybrids mate readily with one another, but both sexes show a reduction in mating success of over 50% with the parental species. Mating preference is associated with a shift in the adult colour pattern, which is involved in predator defence through Müllerian mimicry, but also strongly affects male courtship probability. The hybrids, which lie outside the curve of protection afforded by mimetic resemblance to the parental species, are also largely outside the curves of parental mating preference. Disruptive sexual selection against F1 hybrids therefore forms an additional post-mating barrier to gene flow, blurring the distinction between pre-mating and post-mating isolation, and helping to maintain the distinctness of these hybridizing species.

Keywords: Lepidoptera; Nymphalidae; hybridization; mate choice; post-mating isolation; pre-mating isolation

1. INTRODUCTION

Studies of recently diverged species are increasingly producing examples of sympatric species that hybridize in the wild yet remain distinct (Grant & Grant 1992; Mallet et al. 1998). Understanding the fate of hybrids is therefore critical to our understanding of the nature of the species boundary and of the forces that drive speciation.

Given incomplete assortative mating, the fitness of hybrids will determine the extent to which gene flow occurs between species. Hybrid fitness can be reduced by (i) sterility or inviability (‘genomic’ incompatibility), (ii) disruptive ecological selection, or (iii) disruptive sexual selection. Most attention has been focused on the first effect, hybrid sterility and inviability, with the majority of studies using Drosophila. That work has yielded a number of consistent generalizations about speciation, for instance Haldane’s rule, which states that where one sex of F1 hybrid is absent, rare or sterile, it is the heterogametic sex (Haldane 1922; Coyne 1992; Sperling 1993). However, these studies of genomic incompatibility arguably tell us little about the early stages of speciation: a growing body of work is demonstrating that speciation can proceed in the absence of this form of post-mating isolation (Feder et al. 1994; Bradshaw et al. 1995; Grant & Grant 1997; McMillan et al. 1997; Seehausen et al. 1997; Hatfield & Schluter 1999). More recently, there has been an upsurge in theoretical and empirical interest in the second source of selection against hybrids: ecological forms of disruptive selection (McMillan et al. 1997; Schluter 1998; Dieckmann & Doebeli 1999; Kondrashov & Kondrashov 1999).

Rather less experimental work has investigated mate choice during speciation and the possibility of the third type of selection against hybrids: disruptive sexual selection. Recent theory suggests that disruptive sexual selection could be extremely important in speciation (Payne & Krakauer 1997; Higashi et al. 1999). Several recent examples have shown a reduced mating probability of F1 hybrids between a pair of butterfly species (Davies et al. 1997), preference for conspecific over F1 hybrid males in sticklebacks (Vamosi & Schluter 1999), strong mating discrimination against hybrids in lacewings (Wells & Henry 1998) and almost complete behavioural sterility of both sexes of F1 hybrid in wolf spiders (Stratton & Uetz 1986). This form of selection against hybrids provides an additional element of speciation for which Haldane’s rule might hold, as it has in the effects on vigour or choosiness of female hybrids seen in Anartia butterflies (Davies et al. 1997). Sexual selection will limit the extent to which introgression is possible following hybridization and, if mating asymmetries exist, they will influence the direction in which gene flow might proceed.

Three processes are likely to contribute to divergence in mating preference: (i) pleiotropic or otherwise genetically correlated effects of ecological selection, (ii) disruptive sexual selection, and (iii) reinforcement. First, assortative mating can arise as a by-product of disruptive natural selection (Schluter 1998). Assortment will result if a trait under ecological selection also forms the basis of mate choice or through the recruitment of other traits involved in mate choice by the build-up of linkage disequilibrium with the ecological character (Dieckmann & Doebeli 1999; Kondrashov & Kondrashov 1999). This will generate populations with divergent ecology that are also reproductively isolated and, therefore, that are able to coexist. Second, recent theory suggests that the genetic
correlations between preferences and traits generated purely by disruptive sexual selection can also drive sympatric speciation (Turner & Burrows 1995; Payne & Krakauer 1997, 2000; Higashi et al. 1999). However, the daughter populations from such a process may be able to coexist genetically, but without adaptive differentiation they will not be able to coexist ecologically (but see Lande & Kirkpatrick 1988; Van Doorn et al. 1998). Reinforcement provides a third process that can strengthen pre-mating isolation, however produced, between differentiated populations through selection for assortative mating if hybridization produces offspring with low fitness (Liou & Price 1994). Interestingly, once mating divergence occurs, hybrids are potentially left in an adaptive valley between the ranges of mating preference of the two parental species, which itself can promote additional divergence by further reinforcement.

*Heliconius cydno* and *Heliconius melpomene* (*Lepidoptera: Nymphalidae*) are sister species that are sympatric across much of Central and Andean South America below 1500 m (Brown 1979; Brower 1996). Both are unpalatable and warningly coloured, and speciation has accompanied a shift in their Müllerian mimicry. *Heliconius melpomene* is black with red and yellow markings and mimics *Heliconius erato*, while *H. cydno* is black with yellow or white markings and usually mimics members of the *Heliconius sapho/elencchia* clade (Linares 1997; Jiggins et al. 2001). The two species have diverged in habitat use, with *H. melpomene* in second growth and *H. cydno* in forest understorey, which matches the distribution of their co-mimetic species (Smiley 1978a; Waage et al. 1981; Mallet & Gilbert 1995). However, there is considerable overlap and the two species can be found flying together. They also differ in their degree of host plant specialization within the *Passiflora* (Smiley 1978a,b) and female *F₁* hybrids are sterile (Linares 1989). Despite these differences the two species do hybridize in the wild and *F₁* and backcross hybrids are known from across their range, although they probably form less than 0.1% of natural populations (Mallet et al. 1998).

In this paper, we investigate mating interactions between these two species and estimate the extent of sexual selection against hybrids.

2. METHODS

The work was carried out with *Heliconius melpomene rosina* and *Heliconius cydno chineus* and their *F₁* hybrids. Crosses and mating trials were performed in outdoor insectaries in Gamboa, Republic of Panama, between October 1999 and March 2000. This area lies close to Pipeline Road in Soberania National Park where stocks of both species were collected. The two species will cross in the insectary although rarely, and one direction of cross, *H. melpomene* female × *H. cydno* male, could not be obtained to supply hybrids for these experiments: males of *H. cydno* are consistently more reluctant to mate with heterospecific than are *H. melpomene* males. Although the *H. melpomene* female × *H. cydno* male cross has been produced on two earlier occasions, in both cases the female died before laying eggs.

Mating behaviour was analysed using no-choice trials. Each female was left to eclose in a 1 m × 1 m × 2 m insectary in the presence of a single male and the pair observed at 30 min intervals during the daytime hours for a minimum of 2 days. Males were allowed to mature for at least 5 days beforehand with access to pollen sources (*Psiguria* and *Lantana*) and artificial nectar (10% sugar solution) (McMillan et al. 1997). All individuals were reared without access to the opposite sex and were used once only. Pairs stayed coupled for at least 1 h and so all matings were observed. In addition, females were dissected in order to check for the presence of a spermatophore. The pair were disturbed at each observation period if perching and, once they had come into contact, male behaviour such as chasing the female, fluttering courtship or attempted mating was recorded. This no-choice experimental design was adopted in order to mimic the natural situation, where males patrol larval host plants and mate teneral females soon after eclosion (Mallet 1986). Male choice is almost certainly the primary determinant of mating probability at this stage. Females use their wings for fending off unwanted males, but the wings of teneral females are too soft to be used in this way. A female can use the sperm from a single spermatophore to fertilize several eggs each day over her 6 month lifespan, although a fraction of females do remate in the wild (Bogs 1979).

Mating probabilities were estimated using likelihood in order to test between hypotheses differing in complexity, as well as obtaining measures of reliability (McMillan et al. 1997). A binomial mating probability *P*(_i,j_) was obtained for each combination of i-type female and j-type male, thereby maximizing the expression for log-likelihood given by

\[
\log P_{i,j} + \log (1 - P_{i,j})
\]

where *m* and *n* are the numbers of trials in which the pair mated or remained unmated, respectively. The log-likelihoods for the *P*(_i,j_) values were maximized using the SOLVER algorithm supplied with Microsoft Excel. Support limits for *P*(_i,j_) which are asymptotically equivalent to 95% confidence intervals, were obtained at the parameter values that led to a decrease in the log-likelihood of two units (Edwards 1972). In the case of the parameters *a*, *b*, *c* and *e*, which are multiplicatively combined in the final model (see below), support limits were obtained while maximizing the likelihoods for the other parameters. Fitting models with different numbers of parameters allowed a test for differences in the mating probability across trials using a likelihood ratio test with *G* = 2 *L* log *L₀*, which asymptotically follows a *χ²*-distribution (Edwards 1972).

The likelihood model was fitted in a stepwise manner by adding the parameters to an initial null model with a single mating probability (*a* = *b* = *c* = *d* in table 2) across all trials. Estimating the mating probability separately for interspecific trials (parameter *c* versus *a* = *b* = *d* for the rest) gave a significant improvement (*G* = 30.12, d.f. = 1 and *p* < 0.01). Further improvement was achieved in a three-parameter model estimating separate probabilities for trials among individuals of like genotype (parameter *a* in table 2), hybrid × parental (*b* = *d*) and interspecific mating (parameter *c*) (*G* = 10.14, d.f. = 1 and *p* < 0.01). Males of *H. cydno* displayed a consistently poor performance throughout the period of the experiment, with reduced courtship effort and mating success in all trials including those with a conspecific female. This was not the case in a different set of experiments performed the year before (Jiggins et al. 2001). It was apparent in the offspring of several wild-caught females and seemed to reflect poor adaptation of *H. cydno* to the cage environment. Here we account for the poor performance of *H. cydno* males by estimating an extra parameter (parameter *e*) (*G* = 14.25, d.f. = 1 and *p* < 0.01). Separate mating probabilities

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3. RESULTS

There was very strong assortative mating between the species, with no interspecific mating occurring in 30 trials (table 1). Hybrid males had a high mating success with hybrid females, but both sexes of hybrids had considerably reduced mating probabilities with the parental species (table 1). Male courtship in the first 2 h after female eclosion was a good predictor of male mating success in the trials (figure 1): male mating probability is highly correlated with male courtship probability ($r^2 = 0.65$ and $p < 0.01$).

Interpretation of the likelihood model is complicated by the poor performance of the $H. cydno$ males in both courtship effort and mating success (figure 1 and parameter $e$ in table 2). Since courtship appears to determine male mating performance, it is reasonable to treat the performance parameter (parameter $e$) as an adjustment for the effect of the poor performance by male $H. cydno$. We believe this to be an artefact of this experiment, since a different study with the same pair of species showed similar conspecific mating probabilities within $H. cydno$ and within $H. melpomene$ (Jiggins et al. 2001). This variation between experiments places doubt on the precision of some of the parameter estimates, but the relationships between parameters hold up for the males of each pure species (for example $a > b > c$), regardless of whether the performance correction for $H. cydno$ is used. We can then compare the four mating parameters (parameters $a$–$d$ in table 2). There is very strong assortative mating of the parental species (test of $a = c$, $G = 44.33$, d.f. = 1 and $p < 0.001$) and there is good evidence of discrimination against female hybrids (test of $a = b$, $G = 6.41$, d.f. = 1 and $p < 0.05$). Males of the parental species are therefore most likely to mate with conspecific females, less so with hybrids, and are extremely unlikely to pair with heterospecifics. For hybrid males, pairing with hybrid females is more likely than with either parental species (test of $a = d$, $G = 15.79$, d.f. = 1 and $p < 0.01$): the probability of mating between hybrid males and parental females is estimated to be less than half that of parental $×$ parental or hybrid $×$ hybrid matings (table 2). As noted previously, the mating success of male and female hybrids with parental partners does not differ significantly (test of $b = d$, $G = 1.29$, d.f. = 1 and $p > 0.05$).

4. DISCUSSION

There is extremely strong assortative mating between $H. cydno$ and $H. melpomene$ and reduced mating success of $F_1$ hybrids of both sexes with the parental species. The probability of mating is strongly correlated with the probability of courtship, suggesting that, while female receptivity may play a role, male choice is the primary determinant of mating probability. Although hybrids were not tested in earlier work, a similar correlation was obtained in a study of male courtship and mate choice in $H. cydno$ from Panama and $H. melpomene$ from Panama and French Guiana: in both cases, mating success seems mainly a result of male courtship interest largely due to the males’ colour pattern preference (Jiggins et al. 2001). Male choosiness is not unexpected since the spermatophore of Heliconius represents a considerable nutrient investment, providing the female with amino acids used in egg production (Boggs 1979). In the experiments described here, the mating probabilities for male and female parentals are intermediate when tested with hybrid individuals (conspecific $>$ hybrid $>$ heterospecific) and hybrid success is greatest with hybrids (hybrid $>$ parental). Although the precise nature of mating cues is uncertain, the hybrid male preferences and hybrid female signals from our data seem to be intermediate between those of the two parental species, suggesting an approximately additive genetic basis of both mating cue and response. Similar cases of intermediate signals and preferences of hybrids exist in other species, for example in tree frogs (Doherty & Gerhardt 1983) and lacewings (Wells & Henry 1998), but there are also instances where the mating success of hybrids is similar to that of the

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Table 1. Results from the no-choice mating trials.

(The expected number of matings were estimated from the five-parameter model using the parameters in the final column.)

<table>
<thead>
<tr>
<th>Female</th>
<th>Male</th>
<th>number of trials occurring</th>
<th>total number of trials</th>
<th>expected number of matings</th>
<th>parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>$H. melpomene$</td>
<td>$H. melpomene$</td>
<td>15</td>
<td>17</td>
<td>13.4</td>
<td>$a$</td>
</tr>
<tr>
<td>$F_1$</td>
<td>$F_1$</td>
<td>5</td>
<td>14</td>
<td>4.2</td>
<td>$d$</td>
</tr>
<tr>
<td>$H. melpomene$</td>
<td>$H. cydno$</td>
<td>0</td>
<td>10</td>
<td>0.0</td>
<td>$c \times e$</td>
</tr>
<tr>
<td>$F_1 \times H. melpomene$</td>
<td>8</td>
<td>18</td>
<td>8.2</td>
<td>$b$</td>
<td></td>
</tr>
<tr>
<td>$F_1 \times F_1$</td>
<td>11</td>
<td>16</td>
<td>12.6</td>
<td>$a$</td>
<td></td>
</tr>
<tr>
<td>$F_1 \times H. cydno$</td>
<td>3</td>
<td>16</td>
<td>2.7</td>
<td>$b \times e$</td>
<td></td>
</tr>
<tr>
<td>$H. cydno \times H. melpomene$</td>
<td>0</td>
<td>20</td>
<td>0.0</td>
<td>$c$</td>
<td></td>
</tr>
<tr>
<td>$H. cydno \times F_1$</td>
<td>4</td>
<td>16</td>
<td>4.8</td>
<td>$d$</td>
<td></td>
</tr>
<tr>
<td>$H. cydno \times H. cydno$</td>
<td>5</td>
<td>18</td>
<td>5.3</td>
<td>$a \times e$</td>
<td></td>
</tr>
</tbody>
</table>
parents (Davies et al. 1997; McMillan et al. 1997). In only one published case that we could find, namely that of a spider, was hybrid mating success non-additive, being very low with all genotypes (Stratton & Uetz 1986).

The concept of the ‘shape’ of mating preference provides a unifying theme in the study of sexual selection and species recognition (Ryan & Rand 1993; Ritchie 1996). The shape of mating preference is the relationship between the values of a trait and their probability of acceptance by a mating partner (Ritchie 1996). It determines how sexual selection acts within a population, as well as the probability of hybridization between populations. The distribution of trait values within a population and the shape of preference are expected to coevolve closely, although recognition may be elicited by trait values beyond the normal range of the population, as found for syllable number in the song of the cricket Ephippiger ephippiger (Ritchie 1996). In the initial stages of divergence the preference distributions of two incipient species will overlap. The trait values found in hybrids will depend on the genetic architecture of those traits, but if a number of cues combine additively to form the basis for recognition, hybrids will lie at some intermediate point. Hybrids may therefore be recognized as potential mates by both species and could even have superior mating success to parents in mixed populations (dashed line in figure 2a). Another pair of sister species, Heliconius himera and H. erato, illustrate a possible intermediate stage of divergence as in figure 2a, with strong assortative mating but little discrimination against hybrids (McMillan et al. 1997). At this point, mate choice on its own merely provides stabilizing selection, thereby opposing further divergence. However, if hybrids are selected against, reinforcement can lead to a narrowing of the preference function of the parental species, which might reduce the extent to which hybrids are accepted as mates (figure 2b).

**Table 2.** Multiplicative mating probability parameters from the five-parameter model with support limits.

<table>
<thead>
<tr>
<th>type of parameter</th>
<th>parameter</th>
<th>maximum likelihood value</th>
<th>support limits</th>
</tr>
</thead>
<tbody>
<tr>
<td>pure x pure and hybrid x hybrid mating probability</td>
<td>a</td>
<td>0.786</td>
<td>0.625–0.903</td>
</tr>
<tr>
<td>pure male x hybrid female mating probability</td>
<td>b</td>
<td>0.455</td>
<td>0.257–0.672</td>
</tr>
<tr>
<td>interspecific mating probability</td>
<td>c</td>
<td>0.000</td>
<td>0.000–0.081</td>
</tr>
<tr>
<td>pure female x hybrid male mating probability</td>
<td>d</td>
<td>0.300</td>
<td>0.155–0.479</td>
</tr>
<tr>
<td>H. cydno male mating performance parameter</td>
<td>e</td>
<td>0.371</td>
<td>0.176–0.648</td>
</tr>
</tbody>
</table>

**Figure 1.** Courtship and mating in the no-choice trials. Each cell shows the percentage of males displaying courtship interest towards the female during four observations in the first 2 h after eclosion and the percentage mating over the whole trial period. Courtship was taken to include chasing the female, fluttering courtship above her and attempted or actual mating.

**Table 2.** Multiplicative mating probability parameters from the five-parameter model with support limits.
Alternatively, discrimination may result from further trait divergence of the parental species and coevolution of their mating preference (figure 2c). In either case, disruptive sexual selection against hybrids is generated. This is exactly what we see in *H. melpomene* and *H. cydno*, with an additive pattern of hybrid mating, very strong assortative mating and a reduction in the mating success of hybrids with the parental species of over 50%. This sexual selection may only delay mating for hybrid females, but should result in a strong reduction in lifetime reproductive success for male hybrids. The strength of this sexual selection depends on an Allee effect (Allee et al. 1949) for hybrids: male mating opportunities are only scarce because female hybrids are rare. Hybrids between *H. melpomene* and *H. cydno* are always at extremely low density (less than one in 1000 individuals) (Mallet et al. 1998), so they are unlikely to encounter other hybrids as compatible mates. This creates what is in effect a novel form of post-mating isolation due to poor hybrid mating success.

Ecological adaptation in this pair of species initiated a chain of divergence that led to speciation. Changes in habitat use exposed populations to different suites of potential Müllerian co-mimics (Waage et al. 1981; Mallet & Gilbert 1995), thereby selecting for a shift in mimetic allegiance. This switch in mimicry then led to pleiotropic changes in mate choice, as assortative mating coevolved with colour pattern (Jiggins et al. 2001). Reduced production of hybrids created further disruptive predator selection, leading to low fitness of non-mimetic hybrids, probably in the order of 50% as seen in *H. erato* hybrid zones (Mallet & Barton 1989). Selection within each nascent species after separation almost certainly led to the acquisition of female hybrid sterility (Linares 1989), presumably via Muller’s (1940) classical pleiotropic route for the evolution of genomic incompatibility between populations. These ecological and genomic post-mating barriers created the conditions necessary for reinforcement, which appears to have strengthened the assortative mating in sympatric populations: allopatric *H. melpomene* court and mate with *H. cydno* more readily than the same species pair in sympathy (Jiggins et al. 2001). This greater choosiness in sympathy very probably resulted in still stronger disruptive sexual selection against hybrids. Thus, it is possible to see how simple ecological divergence can trigger a cascade of further changes that lead to full speciation. Each step in divergence we have documented leads to conditions that promote further divergence by reducing gene flow and creating additional disruptive selection pressures in a series of examples of positive feedback in the speciation process. Of course not every species is mimetic or has such clear selective forces at work, but mimicry is a particularly good example where an ecological change that has pleiotropic effects on both pre-mating isolation and post-mating isolation may ultimately cause speciation. Many other examples of similar pleiotropy probably exist.

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