Mimics without models: causes and consequences of allopatry in Batesian mimicry complexes

David W. Pfennig* and Sean P. Mullen

Batesian mimicry evolves when a palatable species (the ‘mimic’) co-opts a warning signal from a dangerous species (the ‘model’) and thereby deceives its potential predators. Longstanding theory predicts that this protection from predation should break down where the model is absent. Thus, mimics are expected to only co-occur with their model. Yet, many mimics violate this prediction and occur in areas where their model is absent. Here, we discuss the causes and consequences of such allopatric mimics. We also describe how these ‘rule-bending’ mimics provide critical insights into diverse topics ranging from how Batesian mimicry evolves to its possible role in speciation.

Keywords: mimicry; predation; speciation; gene flow; species ranges

1. INTRODUCTION

Batesian mimicry evolves when individuals of a palatable species gain the selective advantage of reduced predation because they resemble a toxic species that predators avoid (Ruxton et al. 2004). This idea traces to Bates (1862), who regarded convergent evolution between a palatable species (the ‘mimic’) and an unpalatable one (the ‘model’) as, ‘a most powerful proof of the theory of natural selection’ (Bates 1862, p. 511). Darwin and Wallace, the co-discoverers of natural selection, agreed. Indeed, Darwin wrote to Bates that his paper was, ‘one of the most remarkable and admirable papers I ever read’ (letter from Darwin to Bates, dated 20 November 1862; cited in Burkhardt et al. (2008)). Even today, Batesian mimicry is used in textbooks (e.g. Campbell & Reece 2005; Zimmer 2010) and popular books (e.g. Carroll 2009) as a prime example of natural selection’s efficacy in promoting adaptation.

Although evolutionary biologists have long known about Batesian mimicry, many aspects of its evolution remain unclear (reviewed in Sherratt 2002; Brodie & Brodie 2004; Ruxton et al. 2004). One issue requiring clarification is whether and how a mimic can occur in the absence of its model (Pfennig et al. 2001; Ruxton et al. 2004; Joron 2008). As we describe in detail below, the occurrence of mimics outside the range of their model potentially undercuts the notion that selection to avoid predation drives convergence of models and their putative mimics. Consequently, it has long been postulated that mimics should only occur with their model (Ruxton et al. 2004). Indeed, as Wallace (1867, p. 8) put it, ‘The first law is, that in an overwhelming majority of cases of mimicry, the animals (or the groups) which resemble each other inhabit the same country, the same district, and in most cases are to be found together on the very same spot’.

Here we describe how, unexpectedly, many Batesian mimics occur in the absence of their model. We also consider the various factors that promote such paradoxical species distributions and explore the consequences of allopatric mimics. Finally, we highlight how the allopatric occurrence of Batesian mimics provides fertile testing ground for mimicry theory.

2. MIMICS WITHOUT MODELS?

Classical Batesian mimicry theory predicts that mimics should occur only in sympathy with their model (reviewed in Ruxton et al. 2004). Specifically, this theory predicts that protection from predation should be inversely frequency dependent, such that its effectiveness decreases as the model becomes less abundant (in both a relative and absolute sense; Ruxton et al. 2004). Protection should break down completely when the model is entirely absent (as in allopatry), because predators that do not co-occur with the model will not be under selection to recognize it, or any other species that resemble it, as dangerous (Waldbauer & Sternburg 1987; Pfennig et al. 2001). Moreover, because mimics resemble models that are typically aposematic and, thus, conspicuous to potential predators (Ruxton et al. 2004), predation on these more apparent—but unprotected—allopatric mimics should be particularly intense.

Recent field experiments have substantiated these theoretical expectations. For example, free-ranging carnivore mammals attacked replicas of non-venomous scarlet kingsnakes (Lampropeltis elapsoides), which mimic eastern coral snakes (Micrurus fulvius), and Sonoran mountain kingsnakes (Lampropeltis pyromelana), which mimic Sonoran coral snakes (Micruroides euryxanthus), more frequently in allopatry than in sympathy with
their model (Pfennig et al. 2001). Moreover, within allopatry, replicas exhibiting the mimetic pattern of \textit{L. elapsoides} were attacked more frequently than replicas exhibiting a non-mimetic pattern, suggesting that allopatric mimics do indeed suffer enhanced predation (Pfennig et al. 2007). Thus, theoretical and empirical studies predict that the mimic’s distribution should fall entirely within that of its model. Yet, do the geographical distributions of Batesian mimics support this prediction?

As expected, the geographical distributions of Batesian mimics and their models always overlap; there are no known situations in which a Batesian mimic occurs solely in allopatry (Ruxton et al. 2004). Contrary to expectation, however, the distributions of mimics often do not fall entirely within the distributions of their models. In particular, there are numerous, well-studied Batesian mimicry complexes in which mimics are found in both sympatry and allopatry with their model. Indeed, mimics can occur hundreds of kilometres outside the range of their model. For example, the coral snake mimics \textit{L. elapsoides} (figure 1) and \textit{L. pyromelana} (Stebbins 2003) extend 650 and 800 km, respectively, beyond the range of their coral snake models. Even more remarkable is the case of the (probably) coral snake mimic, \textit{L. zonata}, which occurs 1500 km beyond the range of its model, \textit{M. euryanthus} (Stebbins 2003). Moreover, a majority of a mimic’s range may lie in allopatry. For example, for \textit{L. elapsoides}, a remarkable 60 per cent of the total area of its range is in allopatry.

Figure 1. An allopatric mimic. \textit{(a)} (i) The highly venomous eastern coral snake, \textit{Micrurus fulvius}, and (ii) its non-venomous mimic, \textit{Lampropeltis elapsoides}. \textit{(b)} Geographical distributions of model and mimic in the southeastern United States (green shade, allopatry (only mimic present); yellow shade, sympatry (model and mimic present). Scale bar, 200 km.
Table 1. Representative butterfly and snake Batesian mimicry complexes in the United States and Canada, showing instances in which the mimic occurs entirely outside the geographical range of its model and instances in which the mimic occurs at least partly outside the geographical range of its model.

<table>
<thead>
<tr>
<th>mimicry complex (mimic species, model species)</th>
<th>does the mimic’s range extend beyond its model’s range?</th>
<th>reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>butterfly mimicry complexes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Limenitis arthemis astyanax, Battus philenor</td>
<td>yes</td>
<td>Ries &amp; Mullen (2008)</td>
</tr>
<tr>
<td>Limenitis lorgani Adelpha bredowii</td>
<td>yes</td>
<td>Prudic et al. (2002)</td>
</tr>
<tr>
<td>Papilio glaucus, B. philenor</td>
<td>yes</td>
<td>Brower &amp; Brower (1962)</td>
</tr>
<tr>
<td>Speyeria diana, B. philenor</td>
<td>no</td>
<td>Brower &amp; Brower (1962)</td>
</tr>
<tr>
<td>Snake mimicry complexes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lampornpeltis elapoides, Micrurus fulvius</td>
<td>yes</td>
<td>Pfennig et al. (2001)</td>
</tr>
<tr>
<td>Lampornpeltis triangulum, M. fulvius</td>
<td>yes</td>
<td>Conant &amp; Collins (1998)</td>
</tr>
<tr>
<td>Cenophora coccinez, M. fulvius</td>
<td>yes</td>
<td>Conant &amp; Collins (1998)</td>
</tr>
<tr>
<td>Lampornpeltis pyromelana, Micrurita euryxanthus</td>
<td>yes</td>
<td>Pfennig et al. (2001)</td>
</tr>
<tr>
<td>Lampornpeltis triangulum, M. euryxanthus</td>
<td>yes</td>
<td>Stebbins (2003)</td>
</tr>
<tr>
<td>Chionactis occipitalis, M. euryxanthus</td>
<td>yes</td>
<td>Stebbins (2003)</td>
</tr>
<tr>
<td>Chionactis palostris, M. euryxanthus</td>
<td>no</td>
<td>Stebbins (2003)</td>
</tr>
</tbody>
</table>

(area of allopatric range = 322 km²; area of sympatric range = 217 km²; figure 1).

The existence of allopatric mimics poses a paradox for Batesian mimicry theory. Although some might contend that such a situation is atypical and requires no special explanation, allopatry in Batesian mimicry complexes may be more common than is often assumed. For example, two groups of organisms in which Batesian mimicry has been especially well studied are butterflies and snakes (e.g. Brower 1958; Wallace 1867; Wickler 1968; Greene & McDirmid 1981; Pough 1988; Savage & Slowinski 1992; Brodie & Brodie 2004). We carried out an informal survey of representative butterfly and snake Batesian mimicry complexes in the United States and Canada and found that mimics occur in allopatry in a clear majority of cases (table 1). Thus, the true extent of allopatric occurrence of Batesian mimics may be greater than is generally supposed.

Regardless of the incidence of the phenomenon, clarifying the causes of allopatry in Batesian mimicry complexes is important for at least two reasons. First, the occurrence of mimics outside the range of their models has often been used to dispute whether such systems represent true Batesian mimicry complexes. For example, before recent field experiments confirmed the basic prediction that coral snake mimics are protected only in sympathy with their model (see above), the existence of these mimics in allopatry was used as prima facie evidence against the mimicry hypothesis (Brattstrom 1955; Grobman 1978). Second, as we describe below, the occurrence of mimics in allopatry with their model can have important evolutionary consequences. Before exploring these consequences, however, we first consider the important issue of why mimics occur outside the geographical range of their model.

3. CAUSES OF ALLOPATRIC MIMICS

Three non-mutually exclusive processes can explain the occurrence of allopatric mimics: selection, range contraction/expansion, and gene flow. Below, we discuss each process in turn.

(a) Selection

Selection might favour mimetic patterns even in the absence of the model through at least five major routes. Even when these routes do not actually favour a mimetic phenotype over an alternative non-mimetic phenotype, they could render the mimetic phenotype less costly in allopatry, thereby possibly either maintaining allopatric mimics in the presence of gene flow or maintaining them for longer periods of time following range contraction/expansion.

In the first route, both the model and the mimic are toxic, and what is regarded as a Batesian mimicry complex may actually be an unrecognized case of Müllerian mimicry. Each species would be protected, even in allopatry, because each is toxic and predators associate this noxiousness with a distinctive phenotype on which both species have converged independently. For instance, Vice-royst butterflies (Limenitis archippus) closely resemble several distasteful species of milkweed feeding butterflies in the genus Danaus. Once considered a classic example of defensive Batesian mimicry (Brower 1958), this relationship is now considered to represent quasi-Batesian (Speed 1999) or Müllerian mimicry, because subsequent research has demonstrated that L. archippus sequester phenolic glycosides (Prudic et al. 2007), which are toxic to predators (Ritland 1991, 1995). Of course, if a putative Batesian mimic is found to be toxic, we no longer face a conundrum, even if it occurs in allopatry with its putative model: the putative mimic is protected from predation, even when it occurs alone. It is also possible that a species might be a Müllerian mimic only during certain times or places: a species’ level of toxicity might vary temporally and spatially (especially in species that acquire their toxins from the environment; e.g. Saporito et al. 2007).

In the second route, selection might maintain mimetic phenotypes in allopatry if predators possess an unlearned, generalized avoidance of certain phenotypes, such as those with bright, contrasting colours (Lindstrom 1999). For example, experiments have demonstrated that two species of birds, turquoise-browed motmots (Eumomota superciliosa) and great kiskadees (Pitangus
show unlearned avoidance responses towards dowels painted with red-yellow-black rings (Smith 1975, 1977), which are diagnostic of coral snakes and their mimics (Greene & McDiarmid 1981). Such unlearned avoidance responses might even be expressed in allopatry (but see Rubinoff & Kropach 1970). For instance, naive striated herons (Butorides striatus) exhibit unlearned avoidance responses towards highly venomous yellow-bellied sea snakes (Pelamis platurus), but not towards harmless snakes or eels, even when these herons are derived from populations where sea snakes do not occur (Caldwell & Rubinoff 1983). Other experiments have demonstrated that naive avian predators avoid black and yellow stripes (Schuler & Hesse 1985; Lindstrom 1999), bright red (Roper 1990), and red and yellow prey items (Mastrotta & Mench 1995). More generally, many predators appear to harbour unlearned avoidance of novel stimuli (Coppinger 1970). Thus, many predators might possess an unlearned, generalized avoidance of aposematic signals (Exnerova et al. 2007). Even a slight tendency towards avoiding such phenotypes would reduce the costs of expressing a mimetic phenotype in allopatry.

In the third route, a noxious species and its presumed mimic might have converged independently on a similar phenotype that deters predation, but for different selective reasons (Gadow 1911). A phenotype that acts as a warning signal might, for instance, in some environmental contexts actually enhance crypsis. For example, some researchers have suggested that the brightly coloured, ringed patterns that characterize coral snakes and their mimics (see figure 1) might be cryptic on certain backgrounds or in certain contexts. In particular, on rough backgrounds, such patterns might break up the form of an elongate body, thereby generating a disruptive effect (Brattstrom 1955). Moreover, to some visual systems, these patterns might cause a moving snake to blur into a solid colour and make the snake appear motionless, thereby aiding its escape from predators (Pough 1976). Thus, depending upon the environmental context, a phenotype might function both as a warning and cryptic signal, thereby selectively maintaining it in both sympatry and allopatry.

In each of the above three routes, multiple species might have independently converged on the same phenotype that predators avoid. However, a fourth route by which mimetic phenotypes are selectively maintained in allopatry occurs when the model and mimic have independently converged on a phenotype that is favoured for selective reasons other than deterring predation (Malcolm 1990). For example, if competitors (or prey) are distracted by conspicuous signals, or if they harbour an unlearned, generalized fear of them (see above), then mimetic individuals might gain an advantage over non-mimetic individuals in resource competition, even in allopatry (Rashed & Sherratt 2007; Cheney 2010). Moreover, although mimetic phenotypes might be opposed by natural selection in allopatry, sexual selection might maintain them in such regions (e.g. Summers et al. 1999; Maan & Cummings 2008). Additionally, conspicuous phenotypes might possess non-signalling functions, such as aiding thermoregulation (Lindstedt et al. 2009).

Thus, phenotypic similarity between a model and mimic might reflect convergent evolution in response to similar selective pressures other than predation. Moreover, the mimetic phenotype might have originally evolved because of the benefits of mimicry, and only later acquired additional functions (or vice versa). Regardless of how such phenotypes evolved, a mimetic phenotype might provide its bearer with multiple selective benefits, some of which might maintain such phenotypes in allopatry.

Finally, a fifth route by which mimetic phenotypes might be selectively maintained in allopatry occurs when predators have large home ranges or when they migrate between areas where models are present to those where models are absent. This idea, first proposed by Poulton (1909), is plausible if predators with either innate (Smith 1975, 1977) or learned avoidance, owing to co-occurrence with a toxic model, subsequently avoid aposematic signals when they encounter a palatable look-alike in areas where the model does not occur. For instance, it has been suggested that certain coral snake mimics occur in allopatry (see above), in part because predatory birds migrate seasonally from sympathy (where they are at risk of encountering coral snakes) to allopatry (Groban 1978).

In the fifth route described above, expansion of the predator’s geographical range maintains mimics in allopatry. In §3b, we describe how changes in the model’s or mimic’s geographical range could maintain mimics in allopatry.

(b) Range contraction

A second major cause of allopatric mimics is the differential extinction of model populations. Specifically, the mimetic phenotype might persist in allopatry if the model (but not the mimic) has undergone a range contraction. In such situations, predators might continue to avoid mimics in the newly created allopatric region if they had previously learned or evolved innate tendencies to avoid the dangerous model in what was formerly sympatry. Even if predators fail to avoid mimics in allopatry, there may not have been sufficient time for the mimic to undergo a similar range contraction or to evolve a non-mimetic pattern (see below).

Such range contraction might have contributed to allopatric mimics in a coral snake mimicry complex. In the eastern United States, non-venomous scarlet kingsnakes (L. elapsoides) and milksnakes (Lampropeltis triangulum) closely resemble highly venomous coral snakes (M. fulvius), but they range well outside the latter’s current distribution (Conant & Collins 1998). Micrurus fulvius are now restricted to the southeastern corner of the United States. However, fossils from the Middle Miocene (11.5–16 Myr ago) of Nebraska reveal that they may have formerly occurred much further north and west of their present-day distribution (Holman 2000); i.e. in areas where mimics currently occur in allopatry. The model, therefore, might have experienced a much greater range contraction than did their mimics, which might have contributed to the occurrence of allopatric mimics in this system.

(c) Range expansion and gene flow

Range expansion and/or gene flow by mimics from sympathy into allopatry might maintain mimetic phenotypes in allopatry. Range expansion might create allopatric mimics if models do not undergo a similar range
expansion. For example, Batesian mimicry involving herbivorous insects (e.g. butterflies) may be strongly influenced by ecological limitations on the ranges of their respective host plants. If a mimic’s host plant expands its range more rapidly in response to changing environmental conditions than its model’s host, then dispersal and colonization by the mimetic herbivore could lead to persistence of the mimetic phenotype, even in areas where the model is locally rare or absent. Similarly, recent evidence suggest that the coral snake mimic, *L. elapoides* (figure 1), underwent a range expansion into allopatry within the past 10,000 years (Harper & Pfennig 2008). The lack of a similar range expansion by its model (which actually appears to have undergone an earlier range contraction; see above) might explain the occurrence of allopatric mimics in this system.

Gene flow might also promote the maintenance of allopatric mimics. In particular, allopatric mimics may arise as a consequence of gene flow between mimetic individuals, sympatric with the model, and non-mimetic conspecifics that occur outside the model’s range. Hybridization between mimetic and non-mimetic subspecies of the polytypic butterfly *Limenitis arthemis astyanax* complex may represent an example of this phenomenon (Mullen et al. 2008), and recent work suggests that the position of this hybrid zone is maintained by frequency dependent selection on wing pattern (Ries & Mullen 2008). Ongoing gene flow between sympatry and allopatry may also account for the occurrence of allopatric mimics in the *L. elapoides-M. fulvius* coral snake mimicry complex (Harper & Pfennig 2008).

In some cases, range expansions by mimics into allopatry might have an anthropogenic cause. For example, many species of hoverflies (Diptera: Syrphidae) are Batesian mimics of bees and wasps (reviewed in Gilbert 2005). Yet, in Europe, some mimetic species frequently occur in areas where their models are absent (Howarth & Edmunds 2000). Azmeh et al. (1998) suggested that human-induced changes to the habitat might explain why these mimics occur in allopatry. They suggested that the shift in present-day allopatry from ancient forest to agricultural or urban habitat might have greatly increased the abundance of the hoverfly’s aphid prey, which could account for the mimetic hoverflies’ expansion into allopatry.

To summarize, three processes—selection, range contraction/expansion and gene flow—can promote the allopatric occurrence of Batesian mimics. Although these different processes need not act mutually exclusively, each could, in theory, act alone and thereby account for allopatric mimics.

4. APOSTATIC PREDATION AND THE MAINTENANCE OF ALLOPATRIC MIMICS

Once one or more of the above factors (see §3) promotes the allopatric occurrence of Batesian mimics, even if selection normally disfavours allopatric mimics (Pfennig et al. 2007), it might not act strongly against such mimics if they are rare. Indeed, in a polymorphic population consisting of mimetic and non-mimetic phenotypes, selection acting in allopatry might actually *favour* rare mimetic phenotypes over more common non-mimetic phenotypes, if predators engage in frequency-dependent (i.e. apostatic) predation (Punzalan et al. 2005; Merilaita & Ruxton 2009). With apostatic predation, predation on common prey phenotypes is higher than expected (i.e. based on their actual abundance within the population), if predators develop through learning (or evolve) preferences for those prey phenotypes that they encounter frequently (search image formation is a common explanation offered for apostatic predation; reviewed in Endler 1991). By contrast, predation on rare phenotypes is lower than expected. Although not all predators engage in an apostatic fashion (indeed, some are antiapostatic; see Lindstrom et al. 2001; Endler & Mappes 2004), a recent empirical test suggests that predation on allopatric coral snake mimics in the eastern United States might be apostatic (Pfennig et al. 2007). With apostatic predation, mimics might persist in allopatry as long as they are rare, because predator-mediated selection against rare mimetic phenotypes is weak or non-existent. Consistent with this hypothesis, mimetic phenotypes are much rarer than non-mimetic phenotypes is allopatric populations of coral snake mimics in the eastern United States (Williams 1978).

5. CONSEQUENCES OF ALLOPATRIC MIMICS

Evaluating which of the factor(s) listed in §§3 and 4 contribute to allopatric distributions of mimics will require more research. These studies are important, not only to clarify why Batesian mimics occur in allopatry, but also because the existence of such allopatric mimics can have important evolutionary ramifications. Below, we describe three such consequences of allopatric mimics.

(a) Variable selection for mimicry

The sympathy/allopatry dichotomy that we have presented up to now oversimplifies the true nature of the distributions and abundances of models and mimics in many natural systems. In most cases, the abundance of the model probably declines gradually as one approaches the sympathy/allopatry boundary. Consequently, the fitness benefits of mimicry may attenuate long before reaching this boundary. Thus, selection for mimicry should vary spatially, even within sympathy.

To see how variation in model abundance can lead to variation in selection for mimicry (Endler & Mappes 2004) consider that in areas where the probability of encountering the model is likely to be high, such as in deep sympathy, predators would be under strong selection to avoid any species that remotely resembles the model, especially if the model is highly noxious (Getty 1985). Therefore, in such areas, even imprecise mimics may be protected from predation (Sherratt 2002). By contrast, in areas where the probability of encountering the model is likely to be low, such as on the sympathy–allopatry boundary, selection to avoid the model (and any look-alikes) should be weak (Huheey 1964; Oaten et al. 1975; Getty 1985). In such situations, only those mimics that most closely resemble the model should receive any protection, and natural selection should favour only the best mimics; i.e. those that are the most precise replicas of their model. Indeed, if the model becomes very rare near the boundary, even perfect mimicry might not be favoured (especially if the model is not
particular noxious). In these cases, an alternative (e.g. cryptic) phenotype might be favoured over a phenotype that perfectly matches a (rare) model.

Recent studies of geographical variation in model–mimic resemblances in a coral snake mimicry complex have substantiated these predictions (Harper & Pfennig 2007). Thus, counter intuitively, the best mimics might occur on the sympathy–allopatry boundary rather than in deep sympathy. Consequently, during gene flow, alleles for the best mimics would not have to travel far to reach allopatry.

In summary, when the range of a Batesian mimic extends beyond that of its model, predators in different populations will necessarily vary in their likelihood of encountering the model. Consequently, selection for mimicry will also vary spatially.

(b) Degradation of the mimetic phenotype
A possible ramification of allopatry in Batesian mimicry complexes is the breakdown of the mimetic phenotype by selection. Although mimics are generally protected from predation in sympathy (Pfennig et al. 2001), they might often experience higher than random predation in allopatry (Pfennig et al. 2007). Accordingly, allopatric mimics might often either go extinct or evolve a non-mimetic phenotype (Harper & Pfennig 2008; Joron 2008).

Selection to break down maladaptive Batesian mimicry has been demonstrated in butterflies (Ries & Mullen 2008) and snakes (Harper & Pfennig 2008). In butterflies, selection acts against the introgression of mimetic alleles into a previously non-mimetic population (Limenitis a. arthemis) that has secondarily come into contact with a mimetic subspecies (L. a. astyanax; Mullen et al. 2008; Savage & Mullen 2009). In snakes, range expansion by the coral snake mimic L. elapoides has lead to selection against mimicry in allopatry and erosion of the mimetic phenotype despite ongoing gene flow from sympatry (Harper & Pfennig 2008). Thus, in the absence of an alternative form of selection favouring the mimetic phenotype (see §§3a and 4), allopatric mimics are unlikely to persist.

(c) Speciation
Selection should always favour the maintenance, and even enhancement, of the mimetic phenotype in sympathy. By contrast, selection might often favour the breakdown of this phenotype in allopatry (Pfennig et al. 2007; Harper & Pfennig 2008; Joron 2008). In such situations, allopatric and sympatric populations would experience a divergent pattern of natural selection. Indeed, the transition between these two selective environments might be abrupt. If mimics are selected against in allopatry, but if the best mimics occur on the sympathy–allopatry boundary (see §5a), then there will be a sharp transition between these two contrasting selective environments.

As a consequence of this divergent selection, individuals that select mates from the opposite selective environment would produce offspring that are poorly adapted for either selective environment. These offspring would presumably be intermediate in phenotype and therefore neither mimetic (favoured in the sympatric selective environment) nor cryptic (favoured in the allopatric selective environment). Consequently, selection should favour individuals that cryptic phenotype that perfectly matches a (rare) model. Such assortative mating could, over time, lead to reduced gene flow between selective environments, thereby possibly resulting in the evolution of reproductive isolation between sympatric and allopatric populations.

Presently, there are no known examples in which allopatric mimics have lead to speciation. However, populations in sympathy and allopatry should experience strong divergent selection, and speciation is a possible outcome of such divergent selection (Jiggins et al. 2001, 2004; Naisbit et al. 2001; Servedio 2004).

6. ALLOPATRIC MIMICS AS TESTING GROUND FOR MIMICRY THEORY
The above discussion rests on the assumption that the system of interest is actually a Batesian mimicry complex. Allopatric mimics are valuable because they enable tests of such fundamental assumptions. Indeed, the occurrence of mimics in allopatry allows for tests of the critical expectation that protection from predators should break down where the unpalatable form is not present. Consequently, such tests can be used to establish whether or not a putative mimic is, in fact, a true Batesian mimic (Pfennig et al. 2001).

More generally, allopatric mimics provide an ideal setting for testing mimicry theory, as illustrated by three additional examples. First, allopatric mimics enable tests of how Batesian mimicry evolves in the first place. Evolutionary biologists have long debated how such resemblances can arise gradually (Charlesworth & Charlesworth 1975). Indeed, it is unclear how a population can transition from a cryptic ancestral phenotype to a derived mimetic one if the population must pass through a phase in which it expresses a phenotype intermediate between these two extremes. Generally, such intermediates should be disfavoured because they would be neither cryptic nor mimetic (Charlesworth & Charlesworth 1975). However, intermediate phenotypes might be favoured in areas where models are abundant, such as in deep sympathy (see §5a). Essentially, high model abundance may facilitate the gradual evolution of Batesian mimicry. A recent empirical test, using a Batesian mimicry complex in which mimics occur in both sympathy and allopatry, supports this hypothesis (Kituchi & Pfennig 2010).

Second, such systems can be used to explore mimic–model coevolution. Because models may suffer increased predation as mimics become more numerous, it has long been postulated that selection should cause models to evolve away from their mimics (Gavrilets & Hastings 1998; Speed & Turner 1999; Rowland et al. 2007). However, this ‘chase-away’ selection should be strong only in areas where models are rare relative to mimics, such as at the sympathy–allopatry boundary. Moreover, such selection might be overcome by gene flow from models in deep sympathy, where any mimicry ‘burden’ should be weak (because models are more abundant, many mimics in deep sympathy are likely to be imprecise replicas of the model; Harper & Pfennig 2007; see also §5a).
Tests of this hypothesis, using systems in which mimics
occur in both sympatry and allopatry, could help resolve this issue.

Finally, the occurrence within the same species of both good mimics in sympatry and poor mimics in allopatry (see §5b) presents an ideal opportunity for investigating the genetic basis of mimicry. Specifically, by evaluating the phenotypes produced by the offspring of crosses between good and poor mimics, one can gain insights into the genetic architecture of the mimetic pattern. For example, one could use such data to evaluate whether mimicry involves just a few mutations of large effect, as has been postulated (Joron 2008).

7. CONCLUDING REMARKS

Longstanding theory predicts that the geographical ranges of Batesian mimics should always fit entirely within that of their model (Wallace 1867; Ruxton et al. 2004). Yet, in direct opposition to this longstanding theoretical expectation, mimics often occur outside the range of their model (figure 1 and table 1). Clarifying the processes that account for allopatric mimics (see §§3 and 4) is imperative, because the allopatric occurrence of Batesian mimics can have important evolutionary consequences (see §5). Moreover, the occurrence of Batesian mimics in allopatry offers an ideal setting in which to test mimicry theory (see §6).

A number of critical issues remain to be explored. Here, we highlight five topics that are promising. First, is allopatry more prevalent among certain taxa or when mimicking certain types of models? For example, allopatric mimics may be associated with models that are highly toxic more often than with those that are less toxic. Second, are there any examples of the reverse—models occurring where mimics are absent—and, if so, do allopatric and sympatric models tend to differ? If selection generally favours models that evolve away from their range of their model (figure 1 and table 1)? Clarifying which processes that account for allopatric mimics (see §§3 and 4) is imperative, because the allopatric occurrence of Batesian mimics can have important evolutionary consequences (see §5). Moreover, the occurrence of Batesian mimics in allopatry offers an ideal setting in which to test mimicry theory (see §6).

In short, studies into the causes and consequences of rule-bending allopatric mimics promise to offer fundamental insights into the evolution of mimicry and thereby continue to provide, ‘a most powerful proof of the theory of natural selection’.

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REFERENCES


