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Chapter 2

Seven questions on the chemical ecology and neurogenetics of resource-mediated speciation

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Abstract

Adaptation to different environments can result in reproductive isolation between populations and the formation of new species. Food resources are among the most important environmental factors shaping local adaptation. The chemosensory system, the most ubiquitous sensory channel in the animal kingdom, not only detects food resources and their chemical composition, but also mediates sexual communication and reproductive isolation in many taxa. Chemosensory divergence may thus play a crucial role in resource-mediated adaptation and speciation. Understanding how the chemosensory system can facilitate resource-mediated ecological speciation requires integrating mechanistic studies of the chemosensory system with ecological studies, to link the genetics and physiology of chemosensory properties to divergent adaptation. In this review, we use examples of insect research to present seven key questions that can be used to understand how the chemosensory system can facilitate resource-mediated ecological speciation in consumer populations.

Keywords: Ecological speciation, resource heterogeneity, food, adaptation, reproductive isolation
**Introduction**

Ecological speciation has been recognized as a major mechanism in generating biodiversity (Rundle and Nosil, 2005; Schluter, 2009; Nosil, 2012). Populations occurring in heterogeneous environments may experience divergent selection, resulting in reduced gene flow, which may eventually lead to the evolution of distinct species (Rundle and Nosil, 2005; Schluter, 2009; Nosil, 2012). Classical examples of ecological speciation in animals include the Darwin’s finches on the Galapagos islands, three-spined sticklebacks in limnetic or benthic habitats and host-plant races in herbivorous insects (reviewed in Schluter, 2009; Nosil, 2012). All animals rely on their sensory systems to sense and respond to their local environments. Among the different environmental factors, food resources are among the most important ones shaping local adaptation. The chemosensory system used to find food resources and avoid toxic substances are important targets of divergent selection (Smadja and Butlin, 2009; Hansson and Stensmyr, 2011; Stevens, 2013). Here, we present seven key questions that can be used to understand how the chemosensory system can facilitate resource-mediated ecological speciation in consumer populations (Figure 1). We specifically focus on insects, as they provide several model systems at the forefront of ecological speciation research and present striking evidence of speciation driven by host plant heterogeneity (Matsubayashi et al., 2010). In addition, the insect chemosensory system has been explored using state-of-the-art methods in the *Drosophila* genus for several decades, providing valuable examples on how insects adapt to their chemical environment.

We start by identifying which aspects of food resources are heterogeneous in a given environment (Question 1), as this is the starting point of divergent selection. We then discuss which aspects of food exert selection on consumers (Question 2), and explore how consumers detect (Question 3), exploit (Question 4) and adapt to these resources (Question 5) to understand how local adaptation proceeds. Finally, we discuss whether successful exploitation of new food resources is genetically inherited or acquired during an individual’s lifetime (Question 6) and review the mechanisms that reduce gene flow between individuals that specialized on alternative resources (Question 7). Rather than providing a comprehensive review, we aim to show how answering these seven questions can explain chemosensory-mediated speciation, by integrating approaches traditionally employed by different scientific disciplines, such as chemical ecology, evolutionary biology and neurobiology.
Chapter 2

Figure 1. An overview of seven key questions that can be used to explore the processes leading up to resource-mediated ecological speciation, with methods at the bottom that can be used to study each question. To dissect resource-mediated ecological speciation, we first need to understand the different forms of food resource heterogeneity which could be variation in spatial location, temporal availability, and chemical composition of different food resources like the yellow and red fruits (Question 1). Then, it is important to identify which aspects of the food resources exert selection on consumers (Question 2). It could be because of the toxicity of the food resources. For example, flies which feed on yellow fruit but do not tolerate the associated toxins will die. To understand how consumers interact with food resources, it is necessary to explore how consumers detect (Question 3), exploit (Question 4) and adapt to heterogeneous resources (Question 5). Consumers can detect food resources by food odors (represented by yellow dots), exploit food resources by tolerating defensive compounds (represented by yellow proboscis) and adapt to food resources by the coupling of detection and digestion (represented by yellow odor dots and abdomen). Apart from genetic variation, phenotypic plasticity also modulates consumers' exploitation of a resource (Question 6). Phenotypic plasticity determines whether successful exploitation can be learned and modified during an individual’s lifetime (represented by life cycles of flies). Finally, we come to the mechanisms preventing gene flow between individuals that specialized on alternative resources and those that did not (Question 7). When mating takes place at a specific food resource, while alternative resources are available, assortative mating can emerge as a by-product of divergence in foraging behaviour. For example, the yellow flies are isolated from red flies. The approaches presented by the color at the bottom of each panel from left to right are: field observation (navy blue), analytical chemistry (light blue), toxicology experiment (light green), behavioural analysis (dark red), neurogenetics (pink), chemosensory analysis (dark blue), population genetics (yellow), experimental evolution (dark green) and developmental biology (ochre).
Question 1: Which aspects of the food resource are heterogeneous?

In nature, several aspects of a food resource can be heterogeneous at the same time and induce divergent selection. For example, spatial and temporal variation in food availability may influence the chances of finding mates (Singer and Parmesan, 2010; Fei et al., 2014; Kambach et al., 2016), chemical composition may affect resource perception and feeding (Caillaud and Via, 2000; Webster and Cardé, 2017), and variation in nutritional quality may lead to changes in growth, survival and reproduction (Wetzel et al., 2016; Moreau et al., 2017; Verschut and Hambäck, 2018). An example in which several forms of resource heterogeneity simultaneously act on consumers can be found in the desert fly, Drosophila mojavensis (Diptera: Drosophilidae) (Figure 2), which exploits spatially heterogeneous host cacti. These cacti occur in the Sonoran Desert of North America and are distributed along isolated sites including the Baja California Peninsula, the mainland Sonoran Desert, the Mojave Desert, and the Santa Catalina Island (Heed, 1978). They are susceptible to microbe-induced necroses, which serve as food sources for D. mojavensis (Fogleman and Danielson, 2001). Different necroses from these cacti occur at different times of the year and last for periods of variable duration (Breitmeyer and Markow, 1998), constituting a temporal heterogeneous food resource. An additional, but temporally overlooked form of resource heterogeneity, is microbe-induced heterogeneity. The microbes associated with cacti fermentate the carbohydrates in the cactus and produce specific volatile profiles which are important cues for host detection and exploitation (Becher et al., 2012; Date et al., 2013). These microbes also metabolize the cactus tissues and affect the cactus secondary metabolites which are mainly toxins including triterpene glycosides, medium chain fatty acids, alkaloids, and sterols diols, increasing or decreasing the toxicity of the cactus and thus changing the availability of the nutrients (Fogleman and Danielson, 2001). Together, these different sources of heterogeneity including spatial, temporal and microbe-induced heterogeneity impose divergent selection on D. mojavensis, targeting the flies’ abilities to recognize the odors of its native host, locate the resource and tolerate its secondary metabolites.

Apart from natural variation, human-mediated resource manipulation has become a major driver of consumer adaptation by dramatically changing resource abundance and distribution. A classic example is the introduction and domestication of apple varieties in North America in the mid-1800s. As a result, the local apple maggot fly, Rhagoletis pomonella (Diptera: Tephritidae), which originally fed on hawthorn, started to feed on apples and rapidly spread through North America (Bush, 1969; Feder and Filchak, 1999; Feder et al., 1994). The subsequent introduction of sweeter and earlier ripening apples contributed to a larger diet breadth and longer developmental time for the flies, leading to the formation of a specialized apple race (Feder and Filchak, 1999; Filchak et al., 2000; Hood et al., 2020). An even more recent example is the spread of the spotted wing Drosophila, Drosophila suzukii (Diptera: Drosophilidae) on commercially grown soft fruits like cherries, blueberries, raspberries and grapes. Because this species is attracted to ripening fruit odors and is equipped with an enlarged ovipositor for piercing ripening fruits (Keesey et al., 2015; Revadi et al., 2015; Karageorgi et al., 2017), it utilizes resources at a much earlier stage than other Drosophilids.
and rapidly invaded North America and Europe from Japan (Lee et al., 2011; Cloonan et al., 2018). Food and beverages for humans resulting from yeast fermentation like beer, wine and bread also offer novel food resources for many Drosophila species that naturally feed on yeast and enable them to expand their niches to breweries, wineries and bakeries ( Günther and Goddard, 2019). Consequently, feeding on human-mediated resources may cause consumer populations to become morphologically and genetically distinct from their ancestors, potentially leading to ecological speciation (Capy et al., 2000; Haerty et al., 2003).

Figure 2. Phylogenetic tree of several Drosophila species with their hosts and the key host volatiles. Drosophila sechellia can tolerate the octanoic acid emitted by its host Morinda citrifolia. The seasonal specialist Drosophila erecta is attracted to its host Pandanus by the volatile 3-methyl-2-butenyl acetate. Drosophila mojavensis is attracted by the propyl propionate emitted from its host cactus. Drosophila images are from Nicolas Gompel; host images were taken from https://pixabay.com/.

Question 2: Which aspects of the food resource exert selection on consumers?

To understand how resource heterogeneity underlies divergent selection and speciation in consumer organisms, it is necessary to identify which aspect of the resource is actually exerting selection. This is not trivial, as several aspects of a resource may be heterogeneous at the same time, while adaptation may be primarily driven by one critical factor. For example, the phenology of introduced apple varieties is the trait that initially generated selection on the apple maggot fly to shift their developmental and diapause times (Bush, 1969; Feder and Filchak, 1999; Filchak et al., 2000). Subsequently, the apple race became allochronically isolated from the hawthorn race (Filchak et al., 2000). Among the aspects of the food resource exerting selection on consumers, the chemical composition is most obvious because different nutritional compounds or toxins require different adaptive responses in consumers. For example, because of differences in the chemical composition of pea aphid’s host plants, clover and alfalfa, different host plant races of pea aphid reject their alternative host plant and mainly feed and mate on their native plants (Caillaud and Via, 2000; Campo et al., 2003).
Similarly, differences in volatile profile and secondary metabolites between cactus species are responsible for host specialization and population differentiation among *D. mojavensis* populations (Fogleman and Danielson, 2001).

Toxins are a major driver of food specialization for many herbivorous and frugivorous insects as species that tolerate these toxins can escape competition (Karageorgi et al., 2019; Auer et al., 2020). For example, the noni fruit, *Morinda citrifolia* (L.) (Rubiaceae), produces hexanoic and octanoic acids that are toxic to most insects (Legal et al., 1994; Amlou et al., 1998). However, *Drosophila sechellia* (Diptera: Drosophilidae), a species endemic to the Seychelles islands, tolerates and even prefers it for feeding and oviposition (R'Kha et al., 1991; Legal et al., 1992; Matsuo et al., 2007) (Figure 2). The precursor 3,4-dihydroxyphenylalanine (l-DOPA) in the noni fruit increases female fertility and egg volume in *D. sechellia*, facilitating the egg survival in the toxic environment, enhancing the fitness of adults and offspring to develop on noni fruit and helping *D. sechellia* escape competition with its co-occurring sibling species *Drosophila simulans* (Diptera: Drosophilidae) (Lavista-Llanos et al., 2014). Similarly, the monarch butterfly, *Danaus plexippus* (Lepidoptera: Nymphalidae) feeds on various glycoside-containing milkweeds of the Apocynaceae family that are toxic and repel other insect species. The larvae and adults of the monarch butterfly are physiologically insensitive to the glycoside that otherwise affect an insect’s cardiac physiology and functioning (Holzinger et al., 1992; Karageorgi et al., 2019). To study how different aspects of the food resource exert selection on consumer organisms, integration between toxicology, analytical chemistry, and neurogenetics is needed to identify the toxic effects (Legal et al., 1992), to find chemical cues influencing behaviour (Legal et al., 1994), and to test the organisms’ behavioural, neurological, and physiological responses, for example by using sensory mutants (Auer et al., 2020).

**Question 3: How are consumer chemosensory systems tuned to detect and identify a novel food resource?**

The chemosensory system and its associated neurons generally form the first line of contact between consumers and their food resources (Dicke, 2000; Bruce et al., 2005; Missbach et al., 2014). The ability to detect novel stimuli is pivotal to accessing new resources and changing foraging habits, making the mechanism of resource localization a critical step in resource-mediated ecological speciation. In insects, the chemosensory receptors involved in these processes are distributed in three principal families that have distinct functions in resource detection: the gustatory receptors (GRs) for taste and contact cues (Montell, 2013; Scott, 2018), the odorant receptors (ORs) for olfaction (Vosshall et al., 1999; Liang and Luo, 2010; Joseph and Carlson, 2015), and the ionotropic receptors (IRs) for olfaction, taste, thermosensation and hygrosensation (Benton, 2008; Silbering and Benton, 2010; Enjin et al., 2016; Rimal and Lee, 2018). Genes encoding classical olfactory receptors and their associated sensory neurons are subject to divergent selection in the *Drosophila* genus (Anholt, 2020). For example, the central African *Drosophila erecta* (Diptera: Drosophilidae) locates the fruits of *Pandanus candelabrum* (P. Beauv.) (Pandanaceae) via an increased number of...
olfactory sensory neurons that respond to 3-methyl-2-butenyl acetate (Rio et al., 1983; Linz et al., 2013) (Figure 2). Wild African *D. melanogaster* detects and uses its host marula fruit *Sclerocarya birrea* (A. Rich.) (Anacardiaceae) through the strong response of its Or22a-expressing neurons to the marula volatile ethyl isovalerate (Mansourian et al., 2018). Likewise, the preference of *D. suzukii* for ripening fruits correlates with the duplication of an OR gene and positive selection for ORs tuned to fresh fruit aromas (Revadi et al., 2015; Ramasamy et al., 2016). The leaf-mining fruit fly, *Scaptomyza flava* (Diptera: Drosophilidae), has receptors tuned to volatiles from Brassicaceae leaves rather than yeasts (Goldman-Huertas et al., 2015; Matsunaga et al., 2019). In the mosquito *Aedes aegypti* (Diptera: Culicidae), the transition from forest mammal to human hosts in urban centers coincided with an increased expression of its odorant receptor *AegOr4*, which detects the human body odorant sulcatone (McBride et al., 2014).

Of all *Drosophila* species, the mechanisms by which chemosensory genes contribute to resource adaptation have been especially well characterized for *D. sechellia*. The olfactory response of *D. sechellia* to noni fruits maps to one distinct glomerulus in the antennal lobes (AL, Figure 3), the first olfactory processing center in the brain, receiving input from the odorant receptor Or22a, specific to methyl esters (a noni fruit volatile) (Dekker et al., 2006; Auer et al., 2020). Sequence comparison and functional tests showed that the higher attraction of *D. sechellia* to methyl esters compared to that of *D. simulans* and *D. melanogaster* is determined by three amino acids in the N-terminus of Or22a, exemplifying that a molecular difference in Or22a orthologues can contribute to species-specific olfactory behaviours. Olfactory receptor genes are, however, not the only chemosensory genes contributing to adaptation of *D. sechellia* to noni fruits. A single amino acid change in the ionotropical receptor gene Ir75b is sufficient to provoke the attraction of *D. sechellia* to hexanoic acid (Prieto-Godino et al., 2017). Another change in the chemoreceptor repertoire of *D. sechellia* (compared to its sibling species) involves the odorant-binding proteins (OBP) *Obp57d* and *Obp57e* (Matsuo et al., 2007). OBPs are located in the lymph of sensilla (Larter et al., 2016) and are hypothesized to bind and transfer odorants to the OSNs (Vogt et al., 1991; Leal, 2013; Pelosi et al., 2014) (Figure 3). A mutation in the promoter region of *D. sechellia’s Obp57e* gene prevents its expression, removing taste avoidance towards hexanoic and octanoic acid and reducing repulsion towards noni fruit (Matsuo et al., 2007; Tomioka et al., 2012). When *Obp57e* and *Obp57d* were deleted in *D. melanogaster*, or when hybrids between the sibling species were produced, the behavioural repulsion to the acids also changed (Matsuo et al., 2007), showing that molecular differences in OBP orthologues can contribute to the species-specific olfactory behaviours underlying ecological differences.
Figure 3. A simplified overview showing how odors are detected by the olfactory pathway. Odors are detected by hair-like sensilla on the antennae and the maxillary palps. Panel (A) exemplifies how the antennae of *Drosophila* are covered with the basiconic (blue), trichoid (magenta) and coeloconic (green) sensilla types. These sensilla contain olfactory receptor neurons (ORN) that express olfactory receptors (ORs) and ionotropic receptors (IRs). Panel (B) exemplifies a trichoid sensilla in which the dendrite of the ORN (magenta) is covered with ORs (purple). In *Drosophila*, the ORNs can express one to four olfactory receptors, but as a simplification only one is shown. The odorant binding proteins (OBP - blue) are secreted by the accessory cells (dark grey) into the sensillar lymph and carry the odorants (orange) from the pores to the dendrites. Panel (C) shows a simplified antennal lobe, found in the insect brain, in which odors picked up by the ORN activate specific glomeruli (magenta) that it converges into. The glomeruli that were not activated by the odor are in dark grey.

Consistent with their role in mediating contact between animals and their environment, the genes encoding chemosensory proteins evolve fast (Arguello et al., 2016). Selection operates on standing variation at chemosensory loci. This variation may be particularly easy to maintain because individual chemosensory proteins rarely have large phenotypic effects. These chemosensory genes properties may facilitate rapid evolutionary change when confronted with a new ecological niche.

Question 4: Which trait allows the consumer to successfully exploit the food resource?

While the detection and identification of resources are important steps towards exploiting them, many food resources possess a wide range of toxic defenses or antifeedant secondary metabolites to protect their tissues (Moraes et al., 2001; Kessler and Baldwin, 2001; Ballaré, 2011; Mithöfer and Boland, 2012). The interaction between noni fruits and *D. sechellia* offers a good example of a consumer that not only has chemosensory adaptations to detect its resource but also possesses the ability to develop on the resource. Genetic studies identified a chromosomal region with strong effects on fatty acid octanoic acid (OA) tolerance (Jones, 1998), which mapped to a 170 kb region on the third chromosome, harbouring several odorant-binding proteins and *Osiris* genes (a family of putative transmembrane proteins of unknown function) (Hungate et al., 2013). To establish a causal relationship between the
genes identified by Hungate et al. (2013) and OA tolerance, Andrade López et al. (2017) used RNAi in *D. melanogaster* to knock down each of these 18 genes and test the phenotypic effects. They observed that three genes in the *Osiris* family (*Osi6, Osi7, Osi8*) significantly influenced OA resistance. They also found that *Osi6* and *Osi7* were expressed at significantly higher levels in *D. sechellia* than in intolerant sibling species, indicating that these two genes in particular affect OA resistance. Lanno et al. (2017) subsequently implicated *Osi6*, but also identified many other differentially expressed genes through RNA seq (104 genes in total). Thus, there is a complex genetic architecture to tolerance, which also involves genetic loci not involved in chemosensory perception.

Similar to *D. sechellia*, the leaf-mining fruit fly *S. flava* has a specialized olfactory system alongside a tolerance to compounds that are otherwise lethal to Drosophilids. This species stands out from other Drosophilids by ovipositing and developing on leaves of the Brassicaceae family (Goldman-Huertas et al., 2015; Matsunaga et al., 2019). Gloss et al. (2019) suggested that the shift from ancestral yeast feeding *Drosophila* to the herbivorous *Scaptomyza* is the result of higher turnover rates of the heme-binding cytochrome P450 monoxygenases (CYPs) enzymes that are involved in the detoxification of harmful environmental chemicals. For two Cyp genes, *cyp6g1* and *GstE5-8*, there is an increase in the number of copies in the *S. flava* genome compared to its ancestors. *Cyp6g1* is involved in the evolution of resistance in other insect species through cis-regularly mutations and gene duplication events (Goff and Hilliou, 2017), and *GstE5-8* underlies the production of enzymes that detoxify mustard oils (isothiocyanates). As mustard oils are the main toxic compound in plants of the Brassicaceae family, the tolerance for these toxins may have allowed ancestral *Scaptomyza* species to evade competition for fruits and fungi with other *Drosophila* species by adapting to a novel leaf mining niche. Interestingly, another Cyp gene, *Cyp6as*, has been suggested to facilitate the shift from carnivory to florivory in florivorous bees from the *Apidae* family compared to carnivorous species from the *Vespidae* family, by allowing the metabolism of flavonols in nectar and honey (Johnson et al., 2018). Together, these patterns suggest that detoxification genes can play a crucial role in the successful exploitation of food resources.

**Question 5: Are exploitation and sensory detection genetically linked?**

An organism can only successfully adapt to a novel food resource when the ability to digest the food coincides with the ability to detect it (or even the tendency to be attracted to it). This implies that the genetic architecture of these traits, and how tightly they are linked, strongly influence the likelihood and speed of resource-mediated divergence. Specifically, a shared genetic basis or a genetic correlation between detection and digestion would greatly facilitate niche expansion, and possibly lead to ecological speciation (Lande, 1979; Jaenike, 1990; Fry et al., 1996; Grosjean et al., 2011; Janz, 2011). As reviewed above, both attraction and tolerance have been extensively studied in *D. sechellia*. The introgression lines between *D. sechellia* and *D. simulans* that Hungate et al. (2013) used to map OA tolerance, were also used to explore genetic linkage: flies were subjected to behavioural tests to quantify both
tolerance and preference. Contrary to expectation, genotypes that scored high on tolerance did not also express preference for noni fruit over control food. Thus, the genetic region that confers tolerance does not include loci influencing preference - in fact, other studies suggest that factors influencing preference are located on another chromosome (Yassin et al., 2016). A similar situation is observed in the cactophilic D. mojavensis, which has evolved metabolic pathways to detoxify the toxic compounds in cactus necroses. Different populations express different detoxification genes to utilize their own cactus hosts. For instance, Baja California flies express high levels of the detoxification gene Cyp6g1 for detoxifying Agria, while mainland Sonoran flies mostly depend on the Ugt86Dd gene for detoxification (Matzkin, 2012). The same goes for attraction preferences. All four geographically isolated D. mojavensis populations use volatiles emitted from fermenting cactus necroses to identify their host plant and differ in preferences and sensitivity to the volatiles of these four cactus species (Newby and Etges, 1998; Date et al., 2013; Crowley-Gall et al., 2016). For example, the Mojave population which inhabits Santa Catalina Island displays increased olfactory responses to aromatics (Date et al., 2013), the main volatile of its host plant, barrel cactus, and upregulates two chemosensory genes Or67b and Or71a to detect aromatics (Crowley-Gall et al., 2016). Yet, even though both gene families for attraction and detoxification are paramount in cactus host adaptation, there is no direct genetic linkage between attraction and detoxification (Matzkin et al., 2013; Matzkin, 2014; Allan and Matzkin, 2019).

The evidence presented above, albeit coming from a limited number of species, suggests that physical genetic linkage between resource-specific digestive abilities (including toxin tolerance) and sensory responses towards that same resource (i.e., detection and attraction) does not actually dominate in nature. Instead, the two traits tend to evolve sequentially. Successful exploitation of a new resource may start with a single evolutionary innovation (e.g., the ability to detect a particular chemical compound), which is only maintained if it is quickly followed by the second (e.g., the ability to tolerate an associated toxin). This raises the question as to what comes first in general, tolerance or preference? Island populations of D. yakuba, which recently specialized on M. citrifolia as well, carry the same genetic changes that underlie octanoic acid tolerance in D. sechellia – while their preference for the toxic fruit appears to be largely mediated by other loci (Yassin et al., 2016). These findings suggest that attraction preceded tolerance. This is because once one of the two innovations has evolved, the second one is expected to emerge very rapidly by strong selection on the existing genetic variation within the population. Given that a substantial proportion of genetic variation will be shared between the ancestral D. sechellia and the derived D. yakuba, this leads to the rapid fixation of the same tolerance-conferring alleles (Yassin et al., 2016). However, we need more data from other species to see how consistent this novel food adaptation pattern is, and whether genetic linkage between sensory detection and exploitation is a major determiner of speciation dynamics.
Question 6: How does phenotypic plasticity contribute to the successful exploitation of a novel food resource?

The previous sections illustrate that adaptation to a food resource relies on a chemosensory system that is tuned to locate the resource and on detoxifying enzymes that neutralize toxic compounds (Scriber and Slansky, 1981; Gloss et al., 2016; Webster and Cardé, 2017). The important role of genetically encoded proteins that directly interact with the food resource suggests that these traits may be hard wired during early development. Evolution thus proceeds through classical selection of alleles that confer adaptation. However, some degree of plasticity may allow consumers to adjust their responses through experience. Neural plasticity occurs throughout the olfactory pathway, from olfactory receptor neurons to higher-order neurons, undergoing experience-based changes in synaptic efficacy to neurogenesis and apoptosis (Wilson et al., 2004; Gadenne et al., 2016). This plasticity might promote niche expansion, exposing organisms to new selective pressures that actually favour specialization. On the other hand, olfactory plasticity may interfere with the evolution of resource specialization: if plasticity allows successful detection and exploitation of a (novel) resource, then there is no selective advantage for alleles conferring certain specialist abilities, nor for assortative mating (i.e., individuals mating more often with individuals possessing similar phenotypes). Understanding how preference for certain resources and the ability to exploit it develop, is thus critical to understanding how divergent resource specialization evolves.

In the example of D. sechellia, chemosensory genes are obviously important for the attraction to noni fruit, but they do not solely account for the behavioural differences with its sibling species D. simulans. First, while D. sechellia develops twice as many Or22a-expressing neurons during pupal development compared to its sibling species, this higher number of neurons is not sufficient to explain host attraction. Auer et al. (2020) showed that substituting D. sechellia’s Or22a receptors with those from D. melanogaster, while maintaining their higher number of neurons, does not result in the same level of attraction to noni fruit as wild-type flies. Second, additional differences are found in D. sechellia’s lateral horn, a higher-order brain region that processes innate responses to olfactory cues (Jefferis et al., 2007), in which projection neurons connecting to Or22a-expressing neurons have a prominent branch innervating an area that is not targeted by the homologous neurons in its sibling species (Auer et al., 2020). These differences in projection neuron axon innervations suggest that changes in central-circuit connectivity also allowed for the adaptation to noni fruit. These examples illustrate that although chemosensory gene families evolve fast, additional traits shaping detoxifying enzymes, central-circuit connectivity and projection neuron innervations also account for the evolution of resource detection.

For many insects, resource detection and exploitation are subject to behavioural plasticity resulting from feeding and mating. Hunger generally increases sensitivity towards food odors and decreases sexual receptivity in many insect species (Edgecomb et al., 1994; Lightle et al., 2010; Luo et al., 2013 ; Ko et al., 2015; Lebreton et al., 2017). In fruit flies, this is
controlled through the release of insulin, which leads to sensitization of odorant receptor neurons tuned to juvenile food sources like yeasts (Ribeiro and Dickson, 2010; Root et al., 2011). The increased sensitivity to yeast may, in turn, increase the sexual receptivity of these mated females, leading them to remate at the juvenile food resources (Gorter et al., 2016). In addition, feeding status and food-derived odors influence female responses to male pheromones (Lebreton et al., 2015; Das et al., 2017). Thus, food-induced behavioural changes may indirectly affect mating opportunities. Many species also show behavioural plasticity after mating. The so-called ‘post-mating switch’ lowers responses to pheromones, while increasing responses to host odors or oviposition-sites (Anderson and Anton, 2014; Gadenne et al., 2016). For example, mated female fruit flies show a strong preference for yeast (Ribeiro and Dickson, 2010) and need to locate yeast that allows for the development of their offspring (Becher et al., 2012). Hussain et al. (2016) showed that mated females show an increased expression in the olfactory system of a receptor of myoinhibitory peptides, resulting in a higher sensitivity to the smell of polyamines, which are indicative of substrates that enhance larval development. Both sexes of the cotton leafworm *Spodoptera littoralis* (Lepidoptera: Noctuidae) also exhibit a post-mating response to sex pheromones and host odors. In males, the post-mating switch results in reduced responses to sex pheromones and host leaf odors signalling mating sites, but not diminished attraction to odors signaling adult food (Kromann et al., 2015). In females, mating abolishes attraction to lilac flowers *Syringa vulgaris* (L.) (Oleaceae), but causes attraction to green-leaf odor of the larval host plant cotton, *Gossypium hirsutum* (L.) (Malvaceae) (Saveer et al., 2012). Thus, behavioural plasticity enables consumers to interact with food resources in a way that matches their physiological states and needs.

**Question 7: How do individuals that adapted to an alternative food resource become reproductively isolated?**

A critical step in any ecological speciation scenario is the evolution of some level of reproductive isolation, which can be mediated by various evolutionary mechanisms (Kirkpatrick and Ravigné, 2002). In some of these scenarios divergent adaptation immediately causes reproductive isolation as a by-product. More specifically, when mating takes place at a specific resource, while alternative resources are available, assortative mating can emerge as a by-product of divergence in foraging behaviour. This is one of the most straightforward routes to reproductive isolation, as it does not require the evolution of additional changes in traits involved in mate selection (Kirkpatrick and Ravigné, 2002; Servedio and Boughman, 2017; Hood et al., 2020). This mechanism contributed to the origin of the apple maggot fly, where colonization of apple trees reduced gene flow through spatiotemporal segregation from the ancestral population (Bush, 1969; Dambroski et al., 2005; Hood et al., 2020). Similar spatial or temporal segregation occurs in many other species, including several of the examples mentioned in the sections above, making it one of the major drivers of species diversity in insects (Matsubayashi et al., 2010). Establishing the contribution of spatiotemporal isolation to speciation requires quantifying the degree of
overlap in resource distribution, as well as behavioural studies into the dispersal tendencies and abilities of the consumer organism.

Answers to many of the questions outlined in this review exemplify that divergent adaptation to alternative resources often involves changes in chemosensory perception. Given the importance of chemical signals in sexual communication, such changes may also affect mating probabilities between individuals exploiting different resources. This can constitute another fairly direct route to reproductive isolation, but it requires alignment between the sensory adaptations involved in resource detection, identification and processing, together with adaptations in sexual communication. For several insect species, such alignment occurs when dietary compounds influence the cuticular hydrocarbons (CHC) involved in mate recognition (Etges et al., 2006; Smadja and Butlin, 2009; Sharon et al., 2010; Fedina et al., 2012). When these changes in CHC profiles coincide with changes in mate selection, resource-based assortative mating can evolve, as seen for the cactophilic D. mojavensis (Etges, 1992; Brazner and Etges, 1993; Khallaf et al., 2020). Additionally, gut microbiota obtained by fruit flies through feeding can strongly affect their mate choice, as they select individuals harboring specific microbiota (Sharon et al., 2010, 2011; Brucker and Bordenstein, 2012; Engel and Moran, 2013; Shropshire and Bordenstein, 2016). Consequently, selection for microbiota can be expected to lead to the divergence of host lineages that feed on different substrates, ultimately leading to speciation. It is important to note, however, that these changes are often phenotypically plastic, which modulates their impact on reproductive isolation and species divergence (Otte et al., 2018).

Another caveat concerns the association between signal divergence and mate selection, as changes in chemical signals must align with changes in mating preferences in order to generate reproductive isolation. This is easily achieved when mating depends on the similarity of chemical signals (i.e., ‘self-referent phenotype matching’ (Weddle et al., 2013)), but when mate selection targets specific trait values, assortative mating requires that preferences change in accordance with the chemical signals. This is not always the case, and several examples of chemical mismatches have been documented in Drosophila (e.g., Kwan and Rundle, 2010; Xue et al., 2018). Even without changes in chemical signals themselves, the combination with different food odors might modulate sexual communication. As discussed above, female responses to male pheromones can be affected by food-derived odors (e.g., (Das et al., 2017)). To establish whether such effects could impact the extent of assortative mating, future research should explore whether synergistic effects of food-derived and fly-derived compounds are resource-specific.

In the absence of direct effects of divergent adaptation on sexual communication, assortative mating may evolve as a consequence of selection for local adaptation. A special case concerns ‘reinforcement’, where assortative mating between individuals with the same adaptations evolves to avoid the production of maladapted hybrids (Craig et al., 1997; Caillaud and Via, 2000). In addition to reducing net gene flow, this generates selection for mechanisms that increase the probability of assortative mating, as observed for several
Drosophilids (Yukilevich, 2012). For example, pre-mating isolation in several *Drosophila* sibling species is based on male responses to female CHC profiles (Coyne et al., 1994; Billeter et al., 2009; Shahandeh et al., 2018; Seeholzer et al., 2018). Based on quantitative trait locus (QTL) studies dissecting female CHC profiles (Gleason et al., 2005, 2009) and male courtship preferences (Shahandeh et al., 2018; Shahandeh and Turner, 2020), the most parsimonious scenario is that female CHC profiles diverged due to differences in microclimatic conditions, followed by divergence in male courtship preferences to avoid costly interspecific courtship and mating (Rundle and Dyer, 2015; Shahandeh and Turner, 2020).

Adaptation-assortative mating may also result from mating preferences for traits reflecting the adaptation of prospective mates to the local circumstances. In the most straightforward case, individuals prefer to mate with partners in good physical condition - thereby selecting those individuals that are best adapted. Somewhat more complex is the evolution of novel mate preferences, targeting traits that specifically reflect local performance. Sexual preferences for mates in good conditions is a well-documented phenomenon in behavioural ecology (Davies et al., 2012). In insects, chemical signals play a role in mate choice for unrelated or older mates (Kuo et al., 2012), but in the context of host race formation evidence for the contribution of sexual selection for local adaptation during divergence is scarce. Rundle et al. (2009) observed that experimental selection of *Drosophila serrata* (Diptera: Drosophilidae) on different food resources generated changes in both CHC profiles and CHC-based mating preferences, but failed to accelerate divergent adaptation through sexual selection. Currently, it seems unclear whether chemically-mediated preference for locally adapted mating partners is an important contributor to assortative mating during resource-driven speciation in insects (Rundle et al., 2009).

**Discussion**

In this review, we presented seven questions to explore the processes leading up to resource-mediated ecological speciation, with a particular focus on chemosensory interactions in insects. For each of the seven questions, major insights have been generated using various taxa and approaches, documenting the critical resource properties that drive chemosensory adaptations in consumers, the molecular basis of these adaptations and the characterization of the behavioural responses that contribute to reproductive isolation. Our analysis also revealed important gaps in our knowledge. Notably, different approaches are needed for different questions: we need field ecology and chemical analyses to characterize environmental heterogeneity and its effects on consumer populations, neurobiology and physiology to examine how individuals detect and exploit food resources, developmental biology and genetics to assess the molecular basis of these traits, behavioural observation to document mechanisms underlying reproductive isolation, and evolutionary biology to link selective regimes to evolutionary change. There is rarely one model system in which all seven questions have been thoroughly answered; different organisms are generally employed to answer different questions. For example, *D. mojavensis* is used to explore resource
heterogeneity and population differentiation, while laboratory strains of *D. melanogaster* are used to study receptors, olfactory pathways, neutral circuits and genes involved in resource detection and exploitation. Niche specialists such as *D. sechellia* and *D. mojavensis* are used to examine host specialization and its underlying molecular determinants, while pea aphids and apple maggot flies are used to study the evolutionary ecology of incipient sympatric speciation. In order to extrapolate and generalize, it will be important to incorporate insights from one model system into studies of others. Certainly, all ‘model organism’ and methods have their limitations. *Drosophila* has proven to be a powerful model in neurogenetics and evolutionary genetics because of its unique genetic tools (Bellen et al., 2010; Kazama, 2015; Castillo and Barbash, 2017). These tools however require the use of genetically homogenous or inbred strains so that the control and experimental groups only differ at the level of the mechanism studied, limiting “noise” coming from individual genetic variation. This is essential for data interpretation. However, these inbred *Drosophila* laboratory strains do not represent the diversity and complexity that ecologists aim to explain (Zuk et al., 2014) because ecological speciation relies on the existence of natural genetic variation between individuals, which can then be selected from. It is therefore important to combine the mechanistic insights acquired from the laboratory inbred strains with the study of the natural variants found in nature (Alfred and Baldwin, 2015).

One major knowledge gap concerns the nature of the heterogeneity that exerts divergent selection (Question 1). Environments are heterogeneous in many ways, but it is often not clear what kind of heterogeneity, or which aspect of a particular resource, will generate divergence in consumers. This restricts our predictive ability and hampers our understanding of the relative importance of intrinsic and extrinsic factors in speciation. As noted above, chemosensory genes evolve relatively fast, and this may be true for other sensory genes as well (Stevens, 2013). This may be explained by their specific genomic architectures and functioning, leading to high levels of redundancy and little evolutionary constraint. We may speculate that it is because of these properties that ecological speciation often involves divergence in sensory traits, which would suggest a major role for intrinsic organismal factors in explaining patterns of speciation. Alternatively, it is primarily the environment that governs the likelihood of speciation, and the nature and extent of environmental heterogeneity that determine the strength of divergent selection. Of course, both are important, but a major challenge for speciation research is to establish whether there are general rules across taxa in the contributions of specific intrinsic and extrinsic factors to explaining variation in speciation propensity.

Obviously, additional questions can be formulated. For instance, while we provided several examples of how the adaptation to a food resource relies on a chemosensory system that is tuned to specific substances, the incapacity to detect compounds that are repellent to closely related species may also be coupled to niche specialization (Bernays, 2001). It is also important to explore what happens after speciation has been initiated and some level of reproductive isolation has been achieved. The long-term persistence of new species will
depend on the maintenance of environmental heterogeneity, the geographic setting, the dimension of selection and the genetic changes involved in divergence. Generally, in plant-insect interactions, the diversity and abundance of plants directly affect speciation and extinction in associated insect taxa (Nyman et al., 2012). For instance, the collapse of the host plants drives the extinction of specialized insect species (Wagner and Driesche, 2010). When host plants are segregated in space, persistence of two new herbivore species will be easily maintained because of the absence of gene flow (Nosil, 2012). In line with this, phylogenetic analyses reveal that geographic isolation is more important than host plant heterogeneity in maintaining new species (sawflies: Nyman, 2010; aphids: Jousselin, 2013). In addition to environmental heterogeneity itself, its dimensionality may affect the maintenance of incipient species. Divergence in multiple niche dimensions may not only promote diversification but also the maintenance of differentiation (Nosil, 2012). Finally, the genetic changes involved in divergence may influence the long-term persistence of incipient species. For example, the evolution of reproductive incompatibilities may take a long time (Coyne and Orr, 2004) but once present, greatly facilitates species persistence. It is still an open question whether the factors that promote speciation are also involved in maintaining diversity (Butlin et al., 2012).

One potentially important avenue for future research concerns the role of the consumer gut microbes in resource-mediated speciation. In addition to the microbes present in the substrate, consumer gut microbes also play a role in resource-mediated speciation by mediating resource exploitation ability (Brucker and Bordenstein, 2012; Shropshire and Bordenstein, 2016). For example, the elimination of an endosymbiont in natural Japanese populations of pea aphid A. pisum significantly reduced populations’ exploitation of white clover Trifolium repens (L.) (Fabaceae) (Tsuchida et al., 2004). Gut microbiota may also influence mate choice of hosts. For example, D. melanogaster that were reared on different media acquired different microbiotas, which in turn influenced their cuticular hydrocarbon compositions and led to assortative mating among the flies reared on the same medium (Sharon et al., 2011; Heys et al., 2018). Therefore, microbes in the gut could be a promising direction for future research on resource-mediated ecological speciation.

Theoretical and methodological advances in both speciation biology and chemical ecology make today an exciting time for multidisciplinary speciation research. Speciation biologists have a wealth of methodological tools at their disposal that are increasingly applicable to other taxa beyond the traditional laboratory model organisms, making more taxa available for future comparative studies. We hope that these tools, together with a multidisciplinary perspective and the seven-question framework presented here, will contribute to a more integrated and complete understanding of resource-mediated speciation.
Chapter 2

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