1. Introduction

Sex has long been a popular topic of research among evolutionary biologists. Our personal fascination with the subject is related to the variation that is seen in sexual behavior. This includes the different roles that make up mating rituals, such as courtship traits or preference for the traits, and the variation of these behaviors observed both within and between species. Understanding the biological basis of mating behavior is not only interesting, it is also important for our understanding of evolution as it can shed light on how species boundaries are formed and maintained. Different mating behaviors of closely related species can act as an isolating barrier that stops gene flow between two interbreeding populations. This usually results from closely related species having diverse mating signals: one or both of the sexes fail to identify the other as a suitable mate [1–4]. For example, males of some species court conspecific females more often or with more vigor than heterospecific females [5] and females mate more readily with conspecific than heterospecific males [6, 7]. The impact of Drosophila in this area of research has been pronounced primarily because many obstacles can be bypassed in this system. First, the stereotypical mating behavior observed in this genus is relatively easy to score [8–10], there are genetic tools available to allow manipulation of the development and physiology of mating behavior [11], and there is relative ease in housing large numbers of individuals in a uniform environment. Second, many Drosophila sister species are only partially isolated in a lab setting, producing viable and fertile hybrids [2]. Females of most Drosophila species are usually the sex that determines whether copulation occurs [9]. Males preferentially court conspecific females with larger body sizes, which is a good indicator of female fecundity [12], and in some species (e.g., D. virilis) males are able to discriminate against heterospecific females [5]. However, it is more often found that males readily court heterospecific females [13], while females discriminate against heterospecifics males [6]. Females easily prevent unwanted copulations by flying away from the courting male or extruding her ovipositor [8]. Furthermore, mating behavior has been found to be cyclic.
with alternating bouts of high mating activity and low mating activity; with the use of arrhythmic mutants, it has been shown that females determine when mating occurs [14]. Therefore, in order to understand what isolates species from each other, attention should be focused on female mating behavior.

2. The Evolution of Genes for Female Mating Behavior

The majority of research in behavioral isolation has been influenced by the Modern Synthesis [15], which is a general account of speciation and evolution. The tenants of the Modern Synthesis state that a population contains genetic variation which is apparent at both the gene level (with multiple alleles produced by random mutation) and at the chromosomal level (with different combinations of alleles within individuals produced by recombination). A population's gene frequencies and allele combinations can change over time through multiple processes, including natural selection and genetic drift. While the contribution of natural selection has been wellsupported, the impact of genetic drift has been debated within the literature. For example, a computer model exploring the development of behavioral isolation via sexual selection [16] and research that employed extreme bottlenecking [17] both showed that genetic drift can rapidly lead to some level of sexual isolation. On the other hand, speciation by genetic drift has been shown to be unlikely to occur [18, 19] because genes for mating behavior are most likely either pleiotropic and directly under natural selection, or are closely linked to genes that are under selection and therefore would not simply be fixed by a random process [20].

If a population is divided, the newlyformed subpopulations can potentially become genetically differentiated from each other. Over time, the genomes of each subpopulation can diverge from each other due to the different distribution of alleles that make up the founder population, the different selective pressures on these alleles, and new genetic mutations that arise. As with other traits, the genetic variants that contribute to male and female mating behaviors may cause a difference in phenotype between individuals of the two subpopulations (for review, see [21]). Differences in mating signals can influence female mate choice, which can subsequently act to reduce gene flow between the groups if they come into contact.

Secondary contact between diverging populations can, however, produce hybrids between species in nature. If these hybrids have a relatively high fitness, it is possible that enough gene flow can occur between these two species to cause them to merge back into one. In contrast, if hybrids have a low fitness, a selective pressure to assortatively mate within both populations can act directly on the genes for mating behavior, favoring alleles that differentiate courtship behaviors and enhance preferences for traits that distinguish potential mates of the two groups. This phenomenon, known as reinforcement, has been observed in nature where two closely related species, for example, Drosophila pseudoobscura and D. persimilis, have partially overlapping regions. In response to the selective pressure to avoid heterospecific matings, populations from the sympatric region have a greater level of behavioral isolation compared to those from allopatric populations [22, 23]. However, reinforcement's role in speciation was historically disputed as alternative theories could explain the increased level of behavioral isolation [24], controlled experiments on the topic were generally lacking [25], and some experiments failed to support the theory of reinforcement. For example, the presence of reinforcement within D. mojavensis and D. arizonensis was tested with the use of two groups: one with the traditional rearing substrate of banana agar food, and the other with fermenting cactus—the natural food of these two species. Although behavioral isolation was still found between the species, and the general pattern of reinforcement was still present, the sympatric population was not significantly more behaviorally isolated than the allopatric population [26]. Additionally, reinforcement is not required for differences in mating behavior to arise. For example, a population of Drosophila was subdivided into three groups within the lab: one group remained on the ancestral food source, while two other subpopulations were reared on novel food sources for multiple generations. Afterwards, female mating preferences were tested and were found to be changed in parallel with population divergence [27].

Recently, however, strong empirical data in support of reinforcement has surfaced. Lab investigations have shown that reinforcement can strengthen behavioral isolation between two closely related species [28], and once the selective pressure for species discrimination is removed, the likelihood of interspecific mating has been found to increase [29]. A meta-analysis also found evidence to support the previouslyuntested predictions of reinforcement, such as concordant isolation asymmetries (because reinforcement potentially evolves due to unfit hybrids, pre- and postzygotic isolation should evolve in the same direction) and rare-female effect (females from the smaller population would encounter more heterospecific males and therefore have a stronger selective pressure to choose conspecific males) [30]. Thus, separated populations can evolve divergent behaviors, and these behaviors can potentially be enhanced when the populations are once again in contact.

3. Intraspecific Sexual Selection versus Interspecific Female Mate Choice

The relationship between within- and between-species mating preferences is not fully understood, but they are often considered extreme ends of the same continuum. With time, two populations of a species are thought to slowly slide from assortative mating to heterospecific discrimination, by sexual selection either directly acting on genes that influence mating behavior or indirectly acting on genes that enhance survival. Blows and Allan [1] argued that if species isolation was produced by sexual selection, then the traits involved in species isolation should be the same traits used by both sexes during within-species mate choice. To test this hypothesis, they investigated the behavioral isolation between D. serrata...
and *D. birchii*. These two species have overlapping geographic regions along the east coast of Australia. Although morphologically very similar, there is strong behavioral isolation between the two species [31]. They showed that the two species have different cuticular hydrocarbon (CHC) profiles, which are used as sexual pheromones. By performing perfuming experiments, which transferred CHCs from one species onto another, these researchers determined that the same mechanism (olfaction sensation of CHCs) is used for within-species mate choice (sexual selection) and between-species female mate preference (behavioral isolation).

Although this shows that variation in the same trait can be used for both within- and between-species female mate choice, it does not necessarily mean that they have the same genetic basis. The assumption is that there is a set of genes that control female mating behavior in the ancestral population, and once the population is divided, those genes accumulate mutations in the new populations which cause changes in the behavior. The genes that control intraspecific behavioral variation, however, may not be the same genes that are important in interspecific behavioral variation. Although, for example, genes for olfactory system development used to detect different CHC profiles could be important for normal female mating behavior in both species, the genetic basis for the interpretation of variation in the CHC profile may vary between species.

Investigation into this question led to a series of studies that showed the relationship between interspecific hybridization and intraspecific receptivity. Carracedo et al. [32] proposed that if intraspecific and interspecific mating behaviors have the same genetic basis, females that are slower to accept conspecific males may also be more reluctant to accept heterospecific males. In other words, high level of general within-species receptivity would be selected against due to its pleiotropic effect on high interspecies hybridization. In the lab, when a high level of interspecies hybridization (reduced choosiness) in females was selected for, a decrease in time to start copulation with conspecific males was also found [33]. This was interpreted as a linked increase in intra- and interspecific receptivity, giving support to the notion that both types of mating behavior have the same genetic underpinning. However, when interspecific mating was directly tested by placing females in a choice assay with conspecific and heterospecific flies of the opposite sex, almost no heterospecific matings were observed, showing that selection for heterospecific mating is unlikely to influence within species mate choice in nature, where multiple males are available [34]. The ultimate test of whether the genetic bases of intra- and interspecific mating behavior are under the same genetic control would be to determine and compare the genetic basis of both systems. Unfortunately, no gene has been identified to be involved with interspecific female mate choice. However, a few studies that have identified the regions that most likely contain genes that isolate species do not seem to overlap those regions that contain genes known to influence within-species mating behavior [6, 7, 35].

More unexpected results came from the female mating behavior of island populations. When migrants populate a new island, it is likely that the least choosy females will propagate the most offspring since the most choosy females may not find a high-quality male and therefore will not reproduce [36, 37]. Assuming that low intraspecific choosiness results in high hybridization rates, we would then expect isolated island species to have high levels of hybridization. Although we do find this relationship in the North American and Bogota strains of *D. pseudoobscura* [38], we see the opposite trend in many other species pairs [19]. For example, *D. mauritiana* and *D. sechellia* females, both from island populations, are more choosy against males from the closely related mainland species, *D. simulans*, than mainland females are against island males.

4. Genes for Interspecific Female Mating Behavior

Mating behaviors in *D. simulans* usually have a genetic basis (e.g., of an exception, see [39]). The genetic information that one inherits can predispose a female to behave a specific way: which partner she chooses to accept. These genetic factors can influence both behavioral variation within a species and behavioral differences between species. The latter of these two is critical for our understanding of the genetic basis of species isolation, as it is thought that these behavioral differences are the first barrier to arise in species isolation [40]. By identifying the genetic variants that cause interspecific differences in mating behavior, we can determine which mutations and alterations in the genetic material cause the differences in behavior between two isolated species, and thus may underlie the speciation process itself.

Despite its importance for species isolation, the genetic basis of behavioral isolation is not well understood. This is primarily due to the most commonly used method in genetics for locating genes that contribute to variation in a quantitative trait, namely, recombination mapping. This method necessitates crossing two divergent lines and producing fertile offspring. However, by definition, separate species usually do not produce either fertile or viable offspring. Second, identifying the genetic basis of a behavior requires the location of multiple genes with different effect sizes [41], necessitating a repeatable measure of the behavior, large sample sizes, and the availability of powerful genetic tools such as readily available single gene mutant lines [42].

Despite these obstacles, the genetic basis of mating behavior has been studied in different species of animals and plants. The genetic basis of floral scent production in *Petunia axillaris* (*Petunia*) has been found to play an important role in pollinator attraction and thus contributes to isolation between related species of plants [43]. Research on butterfly mating behavior has found a consistent relationship between wing color and mate preference [44] and both traits may be caused by the same gene (*wingless*) or multiple genes linked to *wingless* [45]. Male cichlids in Lake Victoria have divergent species-specific coloration which has been shown to be driven by female choice [46] and this interspecific female mating preference for conspecific coloration has been found to be heritable in cichlids, with only a few loci responsible [47]. Although butterfly and cichlid coloration...
and preference have provided insight into the genetic basis of behavioral isolation, these systems are limited in that they do not have the powerful genetic tools that are available in *Drosophila*, a well-developed genetic model system.

Using mutagenesis studies, multiple genes have been identified in *D. melanogaster* that influence within-species mating behaviors for both males and females. Male behavior has traditionally taken the spotlight in genetic studies on mating behavior. Through mutagenesis studies, approximately 55 genes have been identified to influence within species male mating behavior, while only a handful of genes have been identified that act within a female to increase or reduce her receptivity (see Supplementary Table 1 available online at doi:10.1155/2012/328392).

These studies are of great importance as they provided crucial information into both the sensory system used in *Drosophila* mating and the types of genes that can influence the construction of mating behavior. However, these studies eliminate the gene’s function in order to test whether it affects a behavior. While this demonstrates that the gene is important for creation of the behavior, it does not necessarily tell us anything about the naturally occurring genetic variation that contributes to the differences seen within or between species. For example, genes identified during mutagenesis for normal male mating behavior were not found to contribute to variation seen in courtship [48], did not contribute to variation between low and high mating male lines [41], and did not vary in expression in a natural population of *D. melanogaster* [49]. The genes important for normal female mating behavior were also not found to vary in expression between courted and naive same-age virgin females [50]. The genes identified through mutagenesis consistently do not appear to influence the variation in mating behavior within a species, and, therefore, may also not contribute to the variation observed between species [51].

Although no individual genes for behavioral isolation have been identified, recombination mapping studies have located regions of the genome that influence behavioral isolation, which do not include genes identified through mutagenesis (see below). However, since the preliminary observations of interspecific female mating behavior do not resemble the expectations set out by prevailing theory, it is difficult to determine strong candidate genes for interspecific female receptivity within these regions [52, 53]. In order to identify which genes are candidates for influencing interspecific female mating behavior, we could first evaluate which signals females are basing their choice upon.

5. The Modes of *Drosophila* Male Signaling during Courtship

The variability we see in female preference, both within and between species, is most likely dictated by the integration of the auditory and olfactory systems [54]. To complicate investigation of these two systems, the amount that females of each species rely on one system over the other is most likely species specific [3, 4, 55, 56]. A gene for interspecific female preference is most likely going to be associated with the signaling pathway of the auditory system used to recognize differences in male courtship song characteristics [3, 57], the olfactory system used to recognize CHC pheromone profiles [1], or both systems via the organization of the part of the brain that receives and interprets signals from both pathways [54, 58]. This is because both modes of signaling are used during *Drosophila* courtship [8–10] and vary between species [1, 3, 56, 59]. A candidate region for such integration in the brain is the mushroom body, which receives signals from many sensory systems in *Drosophila* [60], including the olfaction system [61], and has been linked to sexual behavior [62, 63], specifically female receptivity [64].

There are two main elements to the courtship song—the sine song and the interpulse interval—and males of different species usually differ from each other on both accounts [53]. A female’s ability to identify conspecific song over heterospecifics can lead to behavioral isolation [3]. For females in the melanogaster group of *Drosophila*, the most important element of courtship song is the interpulse interval (IPI) which differs among the males, and preference for variants of IPI seems to differ among females [65]. The most famous gene to influence courtship song is the *period (per)* gene. Mutations in this gene influence IPI [66], and transgenic *D. melanogaster* flies with *D. simulans* produced *D. simulans*-typical rhythm [57]. Instead of a species difference reflecting a complex genetic basis, the species differences in song rhythm reflect just a small number of amino acid changes [57]. Females from this same transgenic line showed associated preference for the transgenic male’s IPI [67], and a later study also showed evidence of assortative mating with a different *per*-transgenic line [68]. Although the genetic basis of this preference is not straightforward, it is clear that females may be using the variations in song between species in determining mate choice. Females can detect male song and male movement with use of the receptors in the antenna; neurons from the antenna project to the dorsal brain, which requires feminization in order for females to be receptive (for review, see [58]).

In addition to song, females also use pheromonal cues to distinguish mates. Each species of *Drosophila* has cuticular hydrocarbons (CHCs) on the outer surface of their body that act as a protective barrier to desiccation and most likely evolved as an adaptation to dry climates [69]. These compounds also are important in mating behavior [70] and are used during mate selection as pheromones that both allow males to distinguish females [71] and affect female receptivity [72]. The majority of CHCs are nonvolatile compounds that are detected by both males and females, most likely through touch (gustation) at close proximity, rather than smell at long distances [70]. Detection of the CHC profile occurs through a large family of odorant receptors that send information about the environment via odorant sensory neurons to the antennal lobe, which is analogous to the olfactory bulb in mammals (for a review, see [58]).

Billeter et al. [71] used a Gal4-UAS system to block the development of oenocytes, which are cells specialized to produce the cuticular hydrocarbons. Flies without working oenocytes (oe−) were completely devoid of all CHCs but behave normally. However, female response towards oe−
males was significantly altered: wildtype females were significantly less receptive to oe− males and oe− males took significantly longer to achieve copulation. Therefore, CHCs not only enhance within species female receptivity [71], but they can also potentially be used to deter females from heterospecific matings [1]. Furthermore, it has been shown that males’ CHC profiles respond more easily to lab-induced natural and sexual selection than the females’ CHC profile [73], indicating that the male profile could be a more likely avenue by which selection acts in nature.

Although there are more than 20 different CHC molecules on the cuticle of the fly, only the predominant hydrocarbons have received much examination and have been primarily studied within the melanogaster subgroup of Drosophila [74]. D. simulans and D. mauritiana have a monomorphic CHC profile, with the main hydrocarbon of both males and females being the same 23-carbon chain compound, cis 7-trisose (7-T). However, D. melanogaster and D. sechellia are dimorphic: the males have large amounts of 7-T, but females lack this hydrocarbon and instead have large amounts of a 27-carbon molecule, cis, cis-7,11-heptacosadiene (7,11-HD) [75]. Most Drosophila species have males that predominately produce 7-T as their main CHC and also share multiple minor compounds as well. However, the ratio between the different CHCs is slightly altered between species, creating unique pheromone “blends” [70].

Through mutagenesis studies, genes have been identified to affect CHC production, such as dsat1 and dsat2 [76], Enhancer of zest [77], Ddc [78], nerd [79], seven pentacosene, and smoq [80], as well as some sex determination genes, such as doublesex [81]. However, only the genetic basis of the main CHC components (7-T and 7,11-HD) have been examined. Additionally, it is unclear if variation in these genes produces the variation that is seen in CHC production between populations of the same species, or variation in production between species [56, 82, 83].

From the research dedicated to identifying the genetic basis of CHC variation between species and courtship song variation between males of different species, we can comfortably deduce that different species have different CHC profiles, different courtship songs, and females preferentially mate with conspecific males based at least partially on both signals.

6. Genetic Basis of Female Behavioral Isolation for Different Species Pairs

To date, no individual genes have been identified as influencing intra- or interspecific female preference in Drosophila, although the trait has a clear heritable basis [8]. Due to the requirement of fertile hybrids for traditional recombination mapping, the majority of studies seeking to address this question have been done in Drosophila species other than D. melanogaster (Table 1), since D. melanogaster does not produce fertile offspring with any of its sibling species[2, 3, 6, 7, 59, 84]. The majority of studies that have examined the behavioral isolation between D. melanogaster and D. simulans have done so in a limited way, showing that specific chromosome arms influence behavioral isolation, and until recently these attempts have not come close to isolating individual genetic variants that affect behavioral isolation [85–87]. However, the genomes of 12 different species of Drosophila have now been sequenced [88], and recently the powerful genetic tools available in D. melanogaster, such as the Gal4-UAS system (used to manipulate gene expression) and transposon vectors (for use in mutagenesis studies), have now been modified for other species of Drosophila [89]. Despite the previous limitations, various genomic regions have been identified that contribute to behavioral isolation in multiple species of Drosophila, and the expansion of the available tools makes further refinement of these studies now possible.

6.1. D. pseudoobscura and D. persimilis. Drosophila pseudoobscura are found across much of Western North America and are located both in sympatry and in allopatry with D. persimilis [123]. The initial genetic basis of isolation between these species, termed basal isolation, was found to be caused by only two regions in the genome: one on the left arm of the X chromosome (which is homologous to the X in D. melanogaster) and one on the second chromosome (homologous to the right arm of chromosome 3, called 3R, in D. melanogaster), within an interspecific inversion that differentiates D. pseudoobscura and D. persimilis [84].

Female D. pseudoobscura from sympatric regions hybridize less with male D. persimilis than females from allopatric regions without D. persimilis, which has made this a model system for studying reinforcement [22]. Ortiz-Barrioneto et al. [109] investigated the genetic basis of the increased discrimination of sympatric D. pseudoobscura females. By introgressing (crossing) pieces of the sympatric D. pseudoobscura genome into an allopatric D. pseudoobscura background, they mapped the increase in behavioral isolation to two allelics of strong effect, one on the right arm of the X chromosome (called Coy-1; homologous to 3L in D. melanogaster) and one on the fourth chromosome (called Coy-2; homologous to 2L in D. melanogaster). However, Barnwell and Noor [124] used six pairs of different inbred strains in a quantitative trait locus (QTL) mapping study to try to replicate the previous identification of Coy-1 and Coy-2. They could not, and therefore determined that Coy-1 and Coy-2, although they may be important, are not the primary loci causing increased behavioral isolation in sympatric versus allopatric populations. These alleles may be present at low frequencies in natural populations and therefore would not be present in most inbred laboratory lines.

Although they may not underlie species-wide discrimination, an examination of the two loci could still provide important insight into the genetic basis of reinforcement. To this end, each of the D. pseudoobscura sympatric and allopatric Coy2 alleles was introgressed into a D. persimilis background (creating perCoy2sym and perCoy2allo lines) [110]. If the reinforced behavioral isolation was caused by an increased receptivity for D. pseudoobscura (conspecifics) by the D. pseudoobscura sympatric population, the expected results would be that perCoy2sym females are more likely to
Table 1: Summary of existing genetic analyses of *Drosophila* species pairs that are behaviorally isolated. The current mode of isolation, trait studied, experimental design (E D), and number of loci potentially affecting behavioral isolation are listed. E D’s are chromosome substitution (C), deficiency complementation mapping (D), complementation mapping of single genes (G), homozygous for a mutation (H), introgression (I), microarray (M), quantitative trait locus mapping (Q), and recombination mapping (R).

| Species pair                      | Isolation       | Trait                                    | E D     | Number of loci |
|-----------------------------------|-----------------|------------------------------------------|---------|----------------|
| *D. melanogaster* (two “races”)  | Allopatric      | Male prezygotic isolation [90–92]        | C, I    | ≥5             |
|                                   |                 | Female prezygotic isolation [90–93]      | C, I, M | ≥4             |
|                                   |                 | Female pheromone production [94]         | R       | 1              |
| *D. melanogaster* and *D. simulans* | Sympatric      | Female pheromone production [95]         | D       | ≥5             |
| *D. simulans* and *D. sechellia* | Allopatric      | Female pheromone production [56, 74]     | Q       | ≥11            |
|                                   |                 | Male prezygotic isolation [59]           | Q       | ≥1             |
|                                   |                 | Male copulation duration [59]            | Q       | ≥1             |
|                                   |                 | Male genital morphology [101]            | Q       | ≥9             |
|                                   |                 | Male sex comb tooth number [96]          | Q       | ≥4             |
|                                   |                 | Male pheromone production [59, 97, 98]  | Q, C    | ≥1–5           |
| *D. simulans* and *D. mauritiana* | Allopatric      | Female prezygotic isolation [2]          | C       | ≥2             |
|                                   |                 | Male courtship song [99]                | Q       | ≥6             |
| *D. mauritiana* and *D. sechellia* | Allopatric      | Female pheromone production [104]        | R       | ≥6             |
| *D. mojavensis* (different populations) | Allopatric   | Male courtship success [105]             | Q       | ≥1             |
|                                   |                 | Male copulation latency [105]            | Q       | ≥3             |
| *D. mojavensis* and *D. arizonae* | Sympatric       | Male prezygotic isolation [106]          | C       | ≥2             |
|                                   |                 | Female prezygotic isolation [106]        | C       | ≥2             |
| *D. heteroneura* and *D. silvestris* | Sympatric     | Male head shape [107, 108]               | C       | ≥9–10          |
|                                   |                 | Female prezygotic isolation and reinforcement [109, 110] | Q, I | ≥4 |
| *D. pseudoobscura* and *D. persimilis* | Sympatric  | Male prezygotic isolation [111, 112]     | C, R    | ≥3             |
|                                   |                 | Male courtship song [113]                | Q       | ≥2–3           |
|                                   |                 | Female prezygotic isolation [84]         | I       | ≥2             |
|                                   |                 | Pheromone production [114]              | C       | ≥2             |
| *D. virilis* and *D. littoralis*  | Sympatric       | Male song production [115]               | C       | ≥3             |
| *D. virilis* and *D. lummei*     | Sympatric       | Male courtship song [116]                | C       | ≥4             |
|                                   |                 | Male pheromone productions [117]        | C       | ≥5             |
| *D. virilis* and *D. a. texana*  | Sympatric       | Male prezygotic isolation [118]          | Q       | ≥1             |
| *D. virilis* and *D. novamexicana* | Sympatric      | Male prezygotic isolation [118]          | Q       | ≥1             |
| *D. auraria* and *D. biuraria*   | Sympatric       | Male courtship song [55]                 | C       | ≥2             |
| *D. montana* (different strains) | Sympatric       | Male pheromone production [119]          | Q       | ≥9             |
| *D. santomea* and *D. yakuba*    | Sympatric       | Female prezygotic isolation [7]          | Q       | ≥3             |
|                                   |                 | Male trait [7]                          | Q       | ≥3             |
| *D. ananassae* (different populations) | Sympatric  | Assortative mating [120]                 | H       | ≥1             |
| *D. ananassae* and *D. pallidosa* | Sympatric       | Female prezygotic isolation [3, 121]     | C, I, R | ≥2 |
|                                   |                 | Male song production [122]              | C       | ≥2             |
mate with D. pseudoobscura than perCoy2allo, but instead they found the opposite: perCoy2sym females were less likely to mate with D. pseudoobscura than perCoy2allo. This suggests that an allele for reduced interspecific mating within a species (Coy2sym) can cause the same reduction in interspecific mating when placed within another species [110]. The explanation provided by Ortiz-Barrientos and Noor is that Coy-2 may be a “One-Allele” mating locus. This theory suggests that one allele (Coy-2) can exist in both the sympatric population of D. pseudoobscura and in D. persimilis population, and aids in the reinforced behavioral isolation between those populations, but not in the basal behavioral isolation. In other words, the same allele causes females of both species to have an increased discrimination against heterospecifics. This is possible if, for example, the gene encodes for increased odor sensitivity or reduced dispersal [125]. This theory would explain why perCoy2sym females were less likely to mate with D. pseudoobscura than perCoy2allo.

6.2. D. ananassae and D. pallidosa. Drosophila ananassae and D. pallidosa are present in overlapping pan-tropical geographic regions. Males of both species court females of both species, but there is strong female interspecific female preference that reduces the gene flow between the two. The genetic basis of this behavior was first explored with female F1 hybrids, which were found to prefer D. ananassae males over D. pallidosa males [3]. This suggests that D. ananassae genes for interspecific female choice must be dominant over those from D. pallidosa. The same study created introgression lines to locate the genomic regions responsible for this behavior. A region on the left arm of the second chromosome (homologous to 3R in D. melanogaster) near the Delta locus was identified to play a role in female species mate choice: females that were almost entirely D. pallidosa except for a small region near the Delta locus mated significantly more with D. ananassae males and significantly less with D. pallidosa males [3]. In other words, this locus both increased intraspecific mating in D. ananassae and decreased interspecific mating between D. ananassae females with heterospecific males. This region was later confirmed by a study that found 2L (3R in D. melanogaster) to be important for the willingness of D. pallidosa females to mate with D. ananassae males, and XL, 2L, and 3R (X, 3R, and 2L in D. melanogaster, resp.) for D. ananassae female’s willingness to mate with D. pallidosa males. All of the identified regions had species specific inversions [121], suggesting that regions of the genome with reduced recombination between the species may be more likely to harbor behavioral isolation loci.

6.3. D. santomea and D. yakuba. Drosophila santomea and D. yakuba diverged approximately 400,000 years ago [126]. D. yakuba is widespread across Africa, including some of the islands off the coast. On one of these islands, D. santomea are found [127]. Although this species pair has a small overlapping geographic region, no reinforcement has been observed [128]. Male courtship behavior may contribute to the behavioral isolation between these two species as D. santomea males do not court heterospecific females with any vigor. To investigate the genetic basis behind the female interspecific mating, a QTL map was created for female rejection of heterospecific males [7]. Three QTLs were identified for D. santomea female discrimination against D. yakuba males: two on the X chromosome (homologous to X in D. melanogaster) and one on the third chromosome (3R in D. melanogaster).

6.4. D. simulans and D. sechellia. Drosophila simulans is a cosmopolitan species, while its closely related sibling species D. sechellia is only found on the Seychelles Islands in the Indian Ocean. There is an asymmetrical behavioral isolation between D. simulans and D. sechellia: D. simulans females are less choosy against D. sechellia males than D. sechellia females are against D. simulans males [2]. Hybrids have an intermediate level of D. simulans rejection when paired with D. simulans males, suggesting an additive genetic basis. Further backcrossing of these F1 hybrids to D. simulans males, and pairing the female offspring with D. simulans males, revealed no isolation, and therefore locating the genes for behavioral isolation in D. sechellia females is not possible with this technique. When the F1 hybrids are backcrossed to D. sechellia males, and the resulting females were assayed with D. simulans males, the second and third chromosomes (2 and 3 in D. melanogaster) were found to have a moderate and strong effect, respectively [2].

6.5. D. simulans and D. mauritiana. D. simulans is a cosmopolitan species and D. mauritiana is only found on the island of Mauritius in the Indian Ocean. It is thought that D. mauritiana resulted from colonization by a recent common ancestor with D. simulans about 250,000 years ago [129]. Females of these species are almost identical, and the males are only distinguishable by the shape of their genital arch [130]. Asymmetrical species isolation is present, with D. simulans being the less choosy of the two courted females. Although D. simulans females are not choosy and readily mate with D. mauritiana males, matings between these two species are abnormally short and result in no or limited sperm transfer, decreasing the number of hybrid offspring [2].

The absence of heterospecific mating by D. mauritiana females is due to the rejection of males by these females, since females of both species are courted vigorously by males of both species [13]. Hybrids produced by D. mauritiana males and D. simulans females mate readily with D. simulans males, and thus the genes for interspecific mate discrimination in D. mauritiana females must be recessive [2, 13]. By backcrossing the hybrids to D. mauritiana males, Coyne was able to assess each D. mauritiana chromosome’s effect on decreasing mating with D. simulans males [13]. He found each of the main autosomes has very large effects with the effect of X being very small [13]. Further dissections of the second chromosome determined that each arm of the second chromosome contains at least one gene for reducing D. mauritiana female mating with D. simulans males (2R and 2L in D. melanogaster); this method of uncovering
recessive *D. mauritiana* genes also possibly removed *D. simulans* genes for conspecific mate preference—these genes may or may not be one in the same. When the same pairings were examined with a more refined map, seven QTL were identified that contribute to *D. mauritiana* discrimination against *D. simulans* males: two on the X chromosome, two on the second chromosome, and three on the third chromosome (X, 2, and 3 in *D. melanogaster*, resp.) [6].

6.6. *D. simulans* and *D. melanogaster*. *Drosophila melanogaster* and *D. simulans* are both cosmopolitan species found worldwide and have broad overlapping geographic distribution. Although both females show some behavioral isolation, *D. simulans* females are far more choosy [131, 132]: interspecific crosses with *D. melanogaster* females are produced with relative ease in the lab, but the reciprocal interspecific cross with *D. simulans* females very rarely occurs [133]. F1 hybrids made from *D. melanogaster* females are all sterile females, and from the reciprocal cross are all sterile males. Due to the complete sterility of hybrids, the conventional method of QTL mapping is not possible as this would require an F2 generation, typically through backcrossing to one of the parental species. Therefore, other methods used to determine the genetic basis of behavioral isolation between these two species have been employed.

Using chromosomal substitution, a genomic region was identified on the third chromosome for *D. melanogaster* female receptivity, and genomic regions on all three major chromosomes were identified for rejection of *D. simulans* males by *D. melanogaster* females [85]. Although there is some evidence that male *D. simulans* may have reduced courtship of interspecific females, and thus contribute to the behavioral isolation [132], there is no such evidence for discrimination by *D. melanogaster* males [134]. Therefore, the strong behavioral isolation demonstrated by *D. simulans* females is largely due to rejection of heterospecific (*D. melanogaster*) males.

To investigate whether there is genetic variation for *D. simulans* female preference, different lab strains of *D. simulans* females [86, 135] and *D. melanogaster* males [86] were compared for their rate of interspecific mating. Crossability, the ability for the parental strains or species to successfully produce offspring, varied among strains for both *D. melanogaster* males and *D. simulans* females [86, 87], but were still highly correlated [135]. When strains of *D. simulans* were crossed, the pure species F1 females were then crossed *D. melanogaster* males and the crossability was compared to the two parental strains. Mixed results were found: while one study found that F1 females always showed greater levels of hybridization [87], another study found that in most cases F1 females showed significantly lower levels of hybridization [86], making it unclear whether increased discrimination within *D. simulans* against heterospecifics is dominant or not. Further inconsistencies include one study that found that X and the third chromosome act additively to contribute to the rejection of *D. melanogaster* males by *D. simulans* females [87], while another study found that the X and the left arm of the second chromosome influenced the trait [133]. These results may be due to the low genetic variability within inbred laboratory lines, and may support the hypothesis that the genetic basis of behavioral isolation varies among populations of the same species. Recently, the right arm of the third chromosome (3R) was mapped using deficiency mapping, revealing five regions (all in areas of low recombination) that contribute to the rejection behavior of *D. simulans* females towards a courting *D. melanogaster* male [35]. While a list of candidate genes in these regions was generated, fine mapping of these regions to the individual gene level remains.

6.7. *M* and *Z* Forms of *D. melanogaster*. *Drosophila melanogaster* are found all over the world, usually commensally with humans, and it was once thought that there was gene flow between populations, including those found spread across large continents [136]. However, a Zimbabwe population was found to have twice the amount of genetic variation compared to North American populations, with certain variants only present in Zimbabwe [137]. Females from these Zimbabwe lines (Z) show behavioral isolation against males from cosmopolitan regions (M): when they have the choice, Z females prefer to mate with Z males, but show no postzygotic isolation (hybrid sterility or inviability) when they are mated with M males. Females from cosmopolitan regions also show behavioral isolation with Z males, but it is weaker than that seen in Z females [90]. The genetic basis for this strong preference in Z females was mapped to all three major chromosomes, with the largest effect being contributed by the third chromosome [91]. With the use of recombinant lines and visible markers (dominant mutations to identify which homologous chromosome was inherited from which parental species), the genetic basis of the female preference in Z females for Z males was mapped to four regions: a region of large effect and a region of minor effect on the left arm of the third chromosome (3L) and a region that most likely houses two loci on the right arm of the third chromosome (3R) [92].

7. Conclusions

In the quest to identify the genetic basis of behavioral isolation, genomic regions have been mapped for interspecific female receptivity in a variety of species pairs. These efforts have yielded maps that vary in refinement from whole chromosomes, chromosomal arms, subchromosomal regions, to specific QTLs. Although the genetic basis of female discrimination may be species pair specific [135], one common attribute of these loci is their location in the genome: most of these loci fall within areas of low recombination, such as species inversion polymorphisms, regions near the centromere, and regions near the telomere. Behavioral isolation loci between *D. santomea* and *D. yakuba* were found near the centromere on 3R [7], and near the telomere for both the *D. simulans* and *D. mauritiana* species pair [6] and the M and Z forms of *D. melanogaster* [92]. Loci responsible for the behavioral isolation between *D. ananassae* and *D. pallidosa* [121], and the isolation between *D. pseudoobscura* and *D. persimilis* [84] all fell within interspecific inversion polymorphisms. Although this was
not true for the regions responsible for increased behavioral isolation caused by reinforcement in the latter species pair [109], these loci for reinforcement were not confirmed by further studies [124].

Inversions have also been shown to play a role in within-species assortative mating. Unlike other species of Drosophila, D. ananassae males have spontaneous meiotic recombination which contributes to the entire species having a high degree of inversion polymorphisms. One inversion, called “alpha,” is a large paracentric inversion covering the majority of 2L (3R in D. melanogaster). To investigate whether this inversion could contribute to behavioral isolation within this species, Nanda and Singh [120] created karyotypically different strains homozygous for one of three naturally occurring inversions. Through mate choice assays, they found a preference for homogamic matings in all three populations.

Genomic rearrangements, centromeric, and telomeric areas can act as an island of low recombination between two potentially interbreeding populations, allowing for the creation and maintenance of population-specific gene complexes (genes inherited together). Over time, new mutations can occur within these complexes and, due to reduced recombination [138], can create a population-typical phenotype if the complexes contain variants for local adaptation [139]. Therefore, even in the face of gene flow between the two groups, a new population identity can be created.

While it has been shown that similar sensory systems may be used for both intra- and interspecific mate discrimination, it is unknown whether these two levels of discriminatory behavior have the same genetic basis. Genomic regions identified as influencing species-specific female preference could contain genes that affect either the auditory or olfactory system, as both are used in mate discrimination, or the brain where this information is processed. If these genes could tolerate a genetic variant causing a slight change in function, selection could then act directly on a new allele, or on other genes within this genetic island, to cause different alleles to reach a high frequency in different populations, causing a slight difference in female mating preference between them. If mutations that occur within these regions cause a change in female preference by influencing assortative mating within species [120], these areas can influence behavioral isolation between species, and thus potentially induce a speciation event [84, 140].

The genetic basis of interspecific female preference is a significant component necessary for understanding the genetic basis of species isolation. While many broad-scale mapping studies have allowed for a solid understanding of the genetic architecture underlying female preference—the number and relative location of genomic regions contributing to female discrimination—to date, no individual genes for this trait have been identified. This limits the ability to assess the mechanism by which females process and evaluate heterospecific mating signals, and thus maintain species isolation. As the genetic tools available in D. melanogaster become more widely available in other systems, and as new mapping techniques are developed, refined genetic dissection of this trait is becoming more tenable. By identifying the genetic mutations that cause interspecific variation in mating behavior, we can start to understand the biological basis species isolation, and better our understanding on the definition of a species. Perhaps the most interesting aspect, however, is that we can finally begin to understand the molecular basis of sex.

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References

[1] M. W. Blows and R. A. Allan, “Levels of mate recognition within and between Drosophila species and their hybrids,” American Naturalist, vol. 152, no. 6, pp. 826–837, 1998.
[2] J. A. Coyne, “Genetics of sexual isolation in females of the Drosophila simulans species complex,” Heredity, vol. 60, no. 1, pp. 25–31, 1992.
[3] M. Doi, M. Matsuda, M. Tomaru, H. Matsubayashi, and Y. Oguma, “A locus for female discrimination behavior causing sexual isolation in Drosophila,” Proceedings of the National Academy of Sciences of the United States of America, vol. 98, no. 12, pp. 6714–6719, 2001.
[4] B. Moulin, T. Aubin, and J. M. Jallon, “Why is there a one-way crossability between D. melanogaster and D. simulans? An ontogenic explanation,” Genetics, vol. 120, no. 1–3, pp. 285–292, 2004.
[5] D. Nickel and A. Civetta, “An X chromosome effect responsible for asymmetric reproductive isolation between male Drosophila virilis and heterospecific females,” Genome, vol. 52, no. 1, pp. 49–56, 2009.
[6] A. J. Moehring, J. Li, M. D. Schug et al., “Quantitative trait loci for sexual isolation between Drosophila simulans and D. mauritiana,” Genetica, vol. 167, no. 3, pp. 1265–1274, 2004.
[7] A. J. Moehring, A. Llopart, S. Elwyn, J. A. Coyne, and T. F. C. Mackay, “The genetic basis of prezygotic reproductive isolation between Drosophila santomea and D. yakuba due to mating preference,” Genetics, vol. 173, no. 1, pp. 215–223, 2006.
[8] J. C. Hall, “The mating of a fly,” Science, vol. 264, no. 5166, pp. 1702–1714, 1994.
[9] R. J. Greenspan, “Understanding the genetic construction of behavior,” Scientific American, vol. 272, no. 4, pp. 74–79, 1995.
[10] R. J. Greenspan and J. F. Ferveur, “Courtship in Drosophila,” Annual Review of Genetics, vol. 34, pp. 205–232, 2000.
[11] K. A. Matthews, T. C. Kaufman, and W. M. Gelbart, “Research resources for Drosophila: the expanding universe,” Nature Reviews Genetics, vol. 6, no. 3, pp. 179–193, 2005.
[12] P. G. Byrne and W. R. Rice, “Evidence for adaptive male mate choice in the fruit fly Drosophila melanogaster,” Proceedings of the Royal Society B, vol. 273, no. 1589, pp. 917–922, 2006.
[13] J. A. Coyne, “Genetics of sexual isolation between two sibling species, Drosophila simulans and Drosophila mauritiana,” Proceedings of the National Academy of Sciences of the United States of America, vol. 86, no. 14, pp. 5464–5468, 1989.
[14] T. Sakai and N. Ishida, “Circadian rhythms of female mating activity governed by clock genes in Drosophila,” Proceedings of the National Academy of Sciences of the United States of America, vol. 98, no. 16, pp. 9221–9225, 2001.

[15] T. Dobzhansky, “Speciation as a stage in evolutionary divergence,” American Naturalist, vol. 74, no. 753, pp. 312–321, 1940.

[16] J. C. Uyeda, S. J. Arnold, P. A. Hohenlohe, and L. S. Mead, “Drift promotes speciation by sexual selection,” Evolution, vol. 63, no. 3, pp. 583–594, 2009.

[17] P. Nanda and B. N. Singh, "Origin of sexual isolation in Drosophila ananassae due to founder effects," Genetica, vol. 139, no. 6, pp. 779–787, 2011.

[18] M. Turelli, N. H. Barton, and J. A. Coyne, “Theory and speciation,” Trends in Ecology and Evolution, vol. 16, no. 7, pp. 330–343, 2001.

[19] J. A. Coyne and A. Orr, Speciation, Sinauer and Associates, Sunderland, UK, 2004.

[20] J. M. Sobel, G. F. Chen, L. R. Watt, and D. W. Schemske, “The biology of speciation,” Evolution, vol. 64, no. 2, pp. 295–315, 2010.

[21] P. Nanda and B. N. Singh, "Behavioural reproductive isolation and speciation in Drosophila," Journal of Biosciences, vol. 37, no. 2, pp. 359–374, 2012.

[22] M. A. Noor, “Speciation driven by natural selection in Drosophila,” Nature, vol. 375, no. 6533, pp. 674–675, 1995.

[23] E. Zouros and C. J. d’Entremont, “Sexual isolation among populations of Drosophila mojavensis; response to pressure from a related species,” Evolution, vol. 34, no. 3, pp. 421–430, 1980.

[24] A. R. Templeton, “Mechanisms of speciation—a population genetic approach,” Annual Review of Ecology, Evolution, and Systematics, vol. 12, pp. 23–48, 1981.

[25] M. A. F. Noor, “Reinforcement and other consequences of sympathy,” Heredity, vol. 83, no. 5, pp. 503–508, 1999.

[26] J. H. Jennings and W. J. Etges, “Species hybrids in the laboratory but not in nature: a reanalysis of premating isolation between Drosophila arizonae and D. mojavensis,” Evolution, vol. 64, no. 2, pp. 587–598, 2010.

[27] H. D. Rundle, S. F. Chenoweth, P. Doughty, and M. W. Blows, “Divergent selection and the evolution of signal traits and mating preferences,” PLoS Biology, vol. 3, no. 11, article e368, 2005.

[28] D. R. Matute, “Reinforcement can overcome gene flow during speciation in Drosophila," Current Biology, vol. 20, no. 24, pp. 2229–2233, 2010.

[29] E. M. Myers and W. A. Frankino, “Time in a bottle: the evolutionary fate of species discrimination in sibling Drosophila species,” PLoS ONE, vol. 7, no. 2, Article ID e31759, 2012.

[30] R. Yukilevich, “Asymmetrical patterns of speciation uniquely support reinforcement in Drosophila," Evolution, vol. 66, no. 5, pp. 1430–1446, 2012.

[31] R. J. Ayala, "Relative fitness of populations of Drosophila serrata and Drosophila birchii," Genetics, vol. 51, pp. 527–544, 1965.

[32] M. C. Carracedo, L. Garcia-Florez, and E. San Miguel, "Sexual maturation in Drosophila melanogaster and hybridization with D. simulans males: a study of inheritance modes," Journal of Heredity, vol. 80, no. 2, pp. 157–158, 1989.

[33] R. Pineiro, M. C. Carracedo, J. I. Izquierdo, and P. Casares, "Bidirectional selection for female receptivity in Drosophila melanogaster," Behavior Genetics, vol. 23, no. 1, pp. 77–83, 1993.

[34] J. I. Izquierdo, M. C. Carracedo, R. Pineiro, and P. Casares, "Response to selection for increased hybridization between Drosophila melanogaster females and D. simulans males," Journal of Heredity, vol. 83, no. 2, pp. 100–104, 1992.

[35] M. Laturney and A. J. Moehring, "Fine-scale genetic analysis of species-specific female preference in Drosophila simulans," Journal of Evolutionary Biology, vol. 25, pp. 1718–1731, 2012.

[36] K. Y. Kaneshiro, "Ethological isolation and phylogeny in the Planitia subgroup of Hawaiian Drosophila," Evolution, vol. 30, no. 4, pp. 740–745, 1976.

[37] K. Y. Kaneshiro, "Sexual selection and direction of evolution in the biosystematics of Hawaiian Drosophilidae," Annual Review of Entomology, vol. 28, pp. 161–178, 1983.

[38] Y. K. Kim, M. Ruiz-Garcia, D. Alvarez, D. R. Phillips, and W. W. Anderson, "Sexual isolation between North American and Bogota strains of Drosophila pseudoobscura," Behavior Genetics, vol. 42, no. 3, pp. 472–482, 2012.

[39] J. Ringo, G. Sharon, and D. Segal, "Bacteria-induced sexual isolation in Drosophila," Fly, vol. 5, no. 4, pp. 310–315, 2011.

[40] J. A. Coyne and H. A. Orr, "Patterns of speciation in Drosophila revisited," Evolution, vol. 51, no. 1, pp. 295–303, 1997.

[41] A. I. Moehring and T. F. C. Mackay, "The quantitative genetic basis of male mating behavior in Drosophila melanogaster," Genetics, vol. 167, no. 3, pp. 1249–1263, 2004.

[42] R. R. H. Anholt and T. F. C. Mackay, "Quantitative genetic analyses of complex behaviours in Drosophila," Nature Reviews Genetics, vol. 5, no. 11, pp. 838–849, 2004.

[43] U. Klahre, A. Gurba, K. Hermann et al., "Pollinator choice in Petunia depends on two major genetic loci for floral scent production," Current Biology, vol. 21, no. 9, pp. 730–739, 2011.

[44] N. I. Morehouse and R. L. Rutowski, "In the eyes of the beholders: female choice and avian predation risk associated with an exaggerated male butterfly color," American Naturalist, vol. 176, no. 6, pp. 768–784, 2010.

[45] M. R. Kronforst, L. G. Young, D. D. Kapan, C. McNelly, R. J. O’Neill, and L. E. Gilbert, "Linkage of butterfly mate preference and wing color preference cue at the genomic location of wingless," Proceedings of the National Academy of Sciences of the United States of America, vol. 103, no. 17, pp. 6575–6580, 2006.

[46] M. E. Maan, O. Seehausen, L. Söderberg et al., "Intraspecific sexual selection on a speciation trait, male coloration, in the Lake Victoria cichlid Pundamilia nyererei," Proceedings of the Royal Society B, vol. 271, no. 1556, pp. 2445–2452, 2004.

[47] M. P. Haesler and O. Seehausen, "Inheritance of female mating preference in a sympatric sibling species pair of Lake Victoria cichlids: implications for speciation," Proceedings of the Royal Society B, vol. 272, no. 1560, pp. 237–245, 2005.

[48] G. E. Carney, "A rapid genome-wide response to Drosophila melanogaster social interactions," BMC Genomics, vol. 8, article 288, 2007.

[49] E. A. Ruedi and K. A. Hughes, "Age, but not experience, affects courtship gene expression in male Drosophila melanogaster," PLoS ONE, vol. 4, no. 7, Article ID e6150, 2009.

[50] M. K. N. Lawiczak and D. J. Begun, "A genome-wide analysis of courting and mating responses in Drosophila melanogaster," Behavior Genetics, vol. 23, no. 1, pp. 77–83, 1993.
[51] T. P. C. Mackay, S. L. Heinsohn, R. F. Lyman, A. J. Moehring, T. J. Morgan, and S. M. Rollmann, “Genetics and genomics of Drosophila mating behavior,” Proceedings of the National Academy of Sciences of the United States of America, vol. 102, no. 1, pp. 6622–6629, 2005.

[52] R. Butlin and M. G. Ritchie, “Evolutionary biology: searching for speciation genes,” Nature, vol. 412, no. 6842, pp. 31–33, 2001.

[53] C. P. Kryiakou, “Single gene mutations in Drosophila: what can they tell us about the evolution of sexual behaviour?” Genetica, vol. 116, no. 2–3, pp. 197–203, 2002.

[54] D. Yamamoto and Y. Nakano, “Sexual behavior mutants revisited: molecular and cellular basis of Drosophila mating,” Cellular and Molecular Life Sciences, vol. 56, no. 7–8, pp. 634–646, 1999.

[55] M. Tomaru, H. Yamada, and Y. Oguma, “Female mate recognition and sexual isolation depending on courtship song in Drosophila sechellia and its siblings,” Genes and Genetic Systems, vol. 79, no. 3, pp. 145–150, 2004.

[56] J. M. Gleason, J. M. Jallon, J. D. Rouault, and M. G. Ritchie, “Quantitative trait loci for cuticular hydrocarbons associated with sexual isolation between Drosophila simulans and D. sechellia,” Genetics, vol. 171, no. 4, pp. 1789–1798, 2005.

[57] D. A. Wheeler, C. P. Kryiakou, M. L. Greenacre et al., “Molecular transfer of a species-specific behavior from Drosophila simulans to Drosophila melanogaster,” Science, vol. 251, no. 4997, pp. 1082–1085, 1991.

[58] B. J. Dickson, “Wired for sex: the neurobiology of Drosophila mating decisions,” Science, vol. 322, no. 5903, pp. 904–909, 2008.

[59] A. Civetta and E. J. F. Cantor, “The genetics of mating recognition between Drosophila simulans and D. sechellia,” Genetical Research, vol. 82, no. 2, pp. 117–126, 2003.

[60] R. L. Davis, “Traces of Drosophila memory,” Neuron, vol. 14, no. 1, pp. 8–19, 2011.

[61] J. S. de Belle and M. Heisenberg, “Associative odor learning in Drosophila abolished by chemical ablation of mushroom bodies,” Science, vol. 263, no. 5147, pp. 692–695, 1994.

[62] M. Balakireva, R. F. Stocker, N. Gendre, and J. F. Ferveur, “Volia, a new Drosophila courtship variant that affects the nervous system: behavioral, neural, and genetic characterization,” Journal of Neuroscience, vol. 18, no. 11, pp. 4335–4343, 1998.

[63] K. M. C. O’Dell, J. D. Armstrong, M. Y. Yang, and K. Kalser, “Functional dissection of the Drosophila mushroom bodies by selective feminization of genetically defined subcompartments,” Neuron, vol. 15, no. 1, pp. 55–61, 1995.

[64] W. S. Neckameyer, “Dopamine modulates female sexual receptivity in Drosophila melanogaster,” Journal of Neurogenetics, vol. 12, no. 2, pp. 101–114, 1998.

[65] J. M. Gleason, “Mutations and natural genetic variation in the courtship song of Drosophila,” Behavior Genetics, vol. 35, no. 3, pp. 265–277, 2005.

[66] C. P. Kryiakou and J. C. Hall, “Circadian rhythm mutations in Drosophila melanogaster affect short-term fluctuations in the male’s courtship song,” Proceedings of the National Academy of Sciences of the United States of America, vol. 77, no. 11, pp. 6729–6733, 1980.

[67] M. G. Ritchie and C. P. Kryiakou, “Reproductive isolation and the period gene of Drosophila,” Molecular Ecology, vol. 3, no. 6, pp. 595–599, 1994.

[68] T. Sakai and N. Ishida, “Time, love and species,” Neuroendocrinology Letters, vol. 22, no. 4, pp. 222–228, 2001.

[69] J. D. Rouault, C. Marican, C. Wicker-Thomas, and J. M. Jallon, “Relations between cuticular hydrocarbon (HC) polymorphism, resistance against desiccation and breeding temperature; a model for HC evolution in D. melanogaster and D. simulans,” Genetica, vol. 120, no. 1–3, pp. 195–212, 2004.

[70] J. F. Ferveur, “Cuticular hydrocarbons: their evolution and roles in Drosophila pheromonal communication,” Behavior Genetics, vol. 35, no. 3, pp. 279–295, 2005.

[71] J. C. Biller, J. Atallah, J. J. Krupp, J. G. Millar, and J. D. Levine, “Specialized cells tag sexual and species identity in Drosophila melanogaster,” Nature, vol. 461, no. 7266, pp. 987–991, 2009.

[72] D. Scott, “Genetic variation for female mate discrimination in Drosophila melanogaster,” Evolution, vol. 48, no. 1, pp. 112–121, 1994.

[73] M. D. Sharma, J. Hunt, and D. J. Hosken, “Antagonistic responses to natural and sexual selection and the sex-specific evolution of cuticular hydrocarbons in Drosophila simulans,” Evolution, vol. 66, no. 3, pp. 665–677, 2012.

[74] J. M. Gleason, R. A. James, C. Wicker-Thomas, and M. G. Ritchie, “Identification of quantitative trait loci function through analysis of multiple cuticular hydrocarbons different between Drosophila simulans and Drosophila sechellia females,” Heredity, vol. 103, no. 5, pp. 416–424, 2009.

[75] J. F. Ferveur, M. Cobb, H. Boukella, and J. M. Jallon, “Worldwide variation in Drosophila melanogaster sex pheromone: behavioural effects, genetic bases and potential evolutionary consequences,” Genetica, vol. 97, no. 1, pp. 73–80, 1996.

[76] R. Dallerac, C. Labeur, J. M. Jallon, D. C. Knipple, W. L. Roelofs, and C. Wicker-Thomas, “A δ desaturase gene with a different substrate specificity is responsible for the cuticular diene hydrocarbon polymorphism in Drosophila melanogaster,” Proceedings of the National Academy of Sciences of the United States of America, vol. 97, no. 17, pp. 9449–9454, 2000.

[77] C. Wicker-Thomas and J. M. Jallon, “Control of female pheromones in Drosophila melanogaster by homeotic genes,” Genetical Research, vol. 78, no. 3, pp. 235–242, 2001.

[78] C. Marican, L. Duportets, S. Birman, and J. M. Jallon, “Female-specific regulation of cuticular hydrocarbon biosynthesis by dopamine in Drosophila melanogaster,” Insect Biochemistry and Molecular Biology, vol. 34, no. 8, pp. 823–830, 2004.

[79] J. F. Ferveur and J. M. Jallon, “Nerd, a locus on chromosome III, affects male reproductive behavior in Drosophila melanogaster,” Naturwissenschaften, vol. 80, no. 10, pp. 474–475, 1993.

[80] J. F. Ferveur and J. M. Jallon, “Genetic control of male cuticular hydrocarbons in Drosophila melanogaster,” Genetical Research, vol. 67, no. 3, pp. 211–218, 1996.

[81] J. M. Jallon, G. Lauge, L. Orsaud, and C. Antony, “Female pheromones in Drosophila melanogaster are controlled by the doublesex locus,” Genetica, vol. 51, no. 1, pp. 17–22, 1988.

[82] J. A. Coyne, C. Wicker-Thomas, and J. M. Jallon, “A gene responsible for a cuticular hydrocarbon polymorphism in Drosophila melanogaster,” Genetical Research, vol. 73, no. 3, pp. 189–203, 1999.

[83] M. G. Ritchie and M. A. Noor, “Evolutionary genetics: gene replacement and the genetics of speciation,” Heredity, vol. 93, no. 1, pp. 1–2, 2004.
[84] M. A. F. Noor, K. L. Gratos, L. A. Bertucci, and J. Reiland, “Chromosomal inversions and the reproductive isolation of species,” Proceedings of the National Academy of Sciences of the United States of America, vol. 98, no. 21, pp. 12084–12088, 2001.

[85] M. C. Carracedo, R. Pineiro, and P. Casares, “Chromosomal substitution analysis of receptivity and sexual isolation in Drosophila melanogaster females,” Heredity, vol. 75, no. 5, pp. 541–546, 1995.

[86] M. C. Carracedo, A. Suarez, A. Asenjo, and P. Casares, “Genetics of hybridization between Drosophila simulans females and D. melanogaster males,” Heredity, vol. 80, no. 1, pp. 17–24, 1998.

[87] T. Ueno and Y. Inoue, “Genetic studies on premating isolation in Drosophila simulans. I. A D. simulans line highly crossable with D. melanogaster,” Japanese Journal of Genetics, vol. 70, no. 3, pp. 365–371, 1995.

[88] Drosophila 12 Genomes Consortium, “Evolution of genes and genomes on the Drosophila phylogeny,” Nature, vol. 450, no. 7167, pp. 203–218, 2007.

[89] S. Holtzman, D. Miller, R. Eisman, H. Kuwayama, T. Niimi, and T. Kaufman, “Transgenic tools for members of the genus Drosophila with sequenced genomes,” Fly, vol. 4, no. 4, pp. 349–362, 2010.

[90] C. I. Wu, H. Hollocher, D. J. Begun, C. F. Aquadro, Y. Xu, and M. L. Wu, “Sexual isolation in Drosophila melanogaster: a possible case of incipient speciation,” Proceedings of the National Academy of Sciences of the United States of America, vol. 92, no. 7, pp. 2519–2523, 1995.

[91] H. Hollocher, C. T. Ting, M. L. Wu, and C. I. Wu, “Incipient speciation by sexual isolation in Drosophila melanogaster: extensive genetic divergence without reinforcement,” Genetics, vol. 147, no. 3, pp. 1191–1201, 1997.

[92] C. T. Ting, A. Takahashi, and C. I. Wu, “Incipient speciation by sexual isolation in Drosophila: concurrent evolution at multiple loci,” Proceedings of the National Academy of Sciences of the United States of America, vol. 98, no. 12, pp. 6709–6713, 2001.

[93] R. I. Bailey, P. Innocenti, E. H. Morrow, U. Friberg, and A. Qvarnström, “Female Drosophila melanogaster gene expression and mate choice: the X chromosome harbours candidate genes underlying sexual isolation,” PLoS ONE, vol. 6, no. 2, Article ID e17358, 2011.

[94] A. Takahashi, S. C. Tsaur, J. A. Coyne, and C. I. Wu, “The nucleotide changes governing cuticular hydrocarbon variation and their evolution in Drosophila melanogaster,” Proceedings of the National Academy of Sciences of the United States of America, vol. 98, no. 7, pp. 3920–3925, 2001.

[95] J. A. Coyne, “Genetics of differences in pheromonal hydrocarbons between Drosophila melanogaster and D. simulans,” Genetics, vol. 143, no. 1, pp. 353–364, 1996.

[96] S. J. Macdonald and D. B. Goldstein, "A quantitative genetic analysis of male sexual traits distinguishing the sibling species Drosophila simulans and D. sechellia," Genetics, vol. 153, no. 4, pp. 1683–1699, 1999.

[97] J. A. Coyne, A. P. Crittenden, and K. Mah, “Genetics of a pheromonal difference contributing to reproductive isolation in Drosophila,” Science, vol. 265, no. 5177, pp. 1461–1464, 1994.

[98] J. A. Coyne, “Genetics of a difference in male cuticular hydrocarbons between two sibling species, Drosophila simulans and D. sechellia,” Genetics, vol. 143, no. 4, pp. 1689–1698, 1996.

[99] J. M. Gleason and M. G. Ritchie, “Do quantitative trait loci (QTL) for a courtship song difference between Drosophila simulans and D. sechellia coincide with candidate genes for intraspecific QTL? ” Genetics, vol. 166, no. 3, pp. 1303–1311, 2004.

[100] J. A. Coyne, “The genetics of an isolating mechanism between two sibling species of Drosophila,” Evolution, vol. 47, no. 3, pp. 778–788, 1993.

[101] J. R. True, J. Liu, L. F. Stam, Z. B. Zeng, and C. C. Laurie, “Quantitative genetic analysis of divergence in male secondary sexual traits between Drosophila simulans and Drosophila mauritiana,” Evolution, vol. 51, no. 3, pp. 816–832, 1997.

[102] Z. B. Zeng, J. Liu, L. F. Stam, C. H. Kao, J. M. Mercer, and C. C. Laurie, “Genetic architecture of a morphological shape difference between two Drosophila species,” Genetics, vol. 154, no. 1, pp. 299–310, 2000.

[103] J. A. Coyne, “Genetics of sexual isolation in male hybrids of Drosophila simulans and D. mauritiana,” Genetical Research, vol. 58, no. 3, pp. 211–220, 1996.

[104] J. A. Coyne and B. Charlesworth, “Genetics of a pheromonal difference affecting sexual isolation between Drosophila mauritiana and D. sechellia,” Genetics, vol. 145, no. 4, pp. 1015–1030, 1997.

[105] W. J. Eges, C. C. de Oliveira, E. Gragg, D. Ortiz-Barrientos, M. A. F. Noor, and M. G. Ritchie, “Genetics of incipient speciation in Drosophila mojavensis. I. Male courtship song, mating success, and genotype X environment interactions,” Evolution, vol. 61, no. 5, pp. 1106–1119, 2007.

[106] E. Zouros, “The chromosomal basis of sexual isolation in two sibling species of Drosophila: D. arizonensis and D. mojavensis,” Genetics, vol. 97, no. 3–4, pp. 703–718, 1981.

[107] A. R. Templeton, “Analysis of head shape differences between two interfertile species of Hawaiian Drosophila,” Evolution, vol. 31, no. 3, pp. 630–641, 1977.

[108] F. C. Val, “Genetic analysis of the morphological differences between two interfertile species of Hawaiian Drosophila,” Evolution, vol. 31, no. 3, pp. 611–629, 1977.

[109] D. Ortiz-Barrientos, B. A. Counterman, and M. A. F. Noor, “The genetics of speciation by reinforcement,” PLoS Biology, vol. 2, no. 12, article e146, 2004.

[110] D. Ortiz-Barrientos and M. A. F. Noor, “Evolution: evidence for a one-allele assortative mating locus,” Science, vol. 310, no. 5753, article 1467, 2005.

[111] M. A. F. Noor, “Genetics of sexual isolation and courtship dysfunction in male hybrids of Drosophila pseudoobscura and Drosophila persimilis,” Evolution, vol. 51, no. 3, pp. 809–815, 1997.

[112] M. A. F. Noor, K. L. Grams, L. A. Bertucci, Y. Almendarez, J. Reiland, and K. R. Smith, “The genetics of reproductive isolation and the potential for gene exchange between Drosophila pseudoobscura and D. Persimilis via backcross hybrid males,” Evolution, vol. 55, no. 3, pp. 512–521, 2001.

[113] M. A. Williams, A. G. Blouin, and M. A. F. Noor, “Courtship songs of Drosophila pseudoobscura and D. persimilis. II. Genetics of species differences,” Heredity, vol. 86, no. 1, pp. 68–77, 2001.

[114] M. A. F. Noor and J. A. Coyne, “Genetics of a difference in cuticular hydrocarbons between Drosophila pseudoobscura and D. persimilis,” Genetical Research, vol. 68, no. 2, pp. 117–123, 1996.

[115] A. Hoikkala, S. Päälysaho, J. Aspi, and J. Lumme, “Localization of genes affecting species differences in male courtship song between Drosophila viridis and D. littoralis,” Genetical Research, vol. 75, no. 1, pp. 37–45, 2000.
[116] A. Hoikkala and J. Lumme, “Genetic control of the difference in male courtship sound between Drosophila virilis and D. lummei,” Behavior Genetics, vol. 14, no. 3, pp. 257–268, 1984.

[117] J. O. Limatainen and J. M. Jallon, “Genetic analysis of cuticular hydrocarbons and their effect on courtship in Drosophila virilis and D. lummei,” Behavior Genetics, vol. 37, no. 5, pp. 713–725, 2007.

[118] T. R. Shirangi, H. D. Dufour, T. M. Williams, and S. B. Carroll, “Rapid evolution of sex pheromone-producing enzyme expression in Drosophila,” PLoS Biology, vol. 7, no. 8, Article ID e1000168, 2009.

[119] W. J. Etges, C. C. de Oliveira, M. A. F. Noor, and M. G. Ritchie, “Genetics of incipient speciation in Drosophila mojavensis. III. Life-history divergence in allopatry and reproductive isolation,” Evolution, vol. 64, no. 12, pp. 3549–3569, 2010.

[120] P. Nanda and B. N. Singh, “Effect of chromosome arrangements on mate recognition system leading to behavioral isolation in Drosophila ananassae,” Genetica, vol. 139, no. 2, pp. 273–279, 2011.

[121] K. Sawamura, H. Zhi, K. Setoguchi et al., “Genetic analysis of female mating recognition between Drosophila ananassae and Drosophila pallidosa: application of interspecific mosaic genome lines,” Genetica, vol. 133, no. 2, pp. 179–185, 2008.

[122] H. Yamada, M. Matsuda, and Y. Oguma, “Genetics of sexual isolation based on courtship song between two sympatric species: Drosophila ananassae and D. pallidosa,” Genetica, vol. 116, no. 2-3, pp. 225–237, 2002.

[123] S. E. McGaugh and M. A. Noor, “Genomic impacts of chromosomal inversions in parapatric Drosophila species,” Philosophical Transactions of the Royal Society B, vol. 367, no. 1587, pp. 422–429, 2012.

[124] C. V. Barnwell and M. A. F. Noor, “Failure to replicate two mate preference QTLs across multiple strains of Drosophila pseudoobscura,” Journal of Heredity, vol. 99, no. 6, pp. 653–656, 2008.

[125] M. R. Servedio and M. A. F. Noor, “The role of reinforcement in speciation: theory and data,” Annual Review of Ecology, Evolution, and Systematics, vol. 34, pp. 339–364, 2003.

[126] A. Llopart, S. Elwyn, D. Lachaise, and J. A. Coyne, “Genetics of a difference in pigmentation between Drosophila yakuba and Drosophila santomea,” Evolution, vol. 56, no. 11, pp. 2262–2277, 2002.

[127] D. Lachaise, M. Harry, M. Solignac, F. Lemeunier, V. Benassi, and M. L. Cariou, “Evolutionary novelties in islands: Drosophila santomea, a new melanogaster sister species from São Tomé,” Proceedings of the Royal Society B, vol. 267, no. 1452, pp. 1487–1495, 2000.

[128] J. A. Coyne, S. Y. Kim, A. S. Chang, D. Lachaise, and S. Elwyn, “Sexual isolation between two sibling species with overlapping ranges: Drosophila santomea and Drosophila yakuba,” Evolution, vol. 56, no. 12, pp. 2424–2434, 2002.

[129] R. M. Kliman, P. Andolfatto, J. A. Coyne et al., “The population genetics of the origin and divergence of the Drosophila simulans complex species,” Genetics, vol. 156, no. 4, pp. 1913–1931, 2000.

[130] L. Tsacas and J. David, “Drosophila mauritiana n. sp. du groupe melanogaster de l’île Maurice,” Bulletin de la Société Entomologique de France, vol. 79, pp. 42–46, 1974.

[131] D. Wood, J. M. Ringo, and L. L. Johnson, “Analysis of courtship sequences of the hybrids between Drosophila melanogaster and Drosophila simulans,” Behavior Genetics, vol. 10, no. 5, pp. 459–466, 1980.

[132] M. C. Carracedo, A. Asenjo, and P. Casares, “Genetics of Drosophila simulans male mating discrimination in crosses with D. melanogaster,” Heredity, vol. 91, no. 3, pp. 202–207, 2003.

[133] M. C. Carracedo, A. Asenjo, and P. Casares, “Inheritance mode of Drosophila simulans female mating propensity with D. melanogaster males,” Journal of Heredity, vol. 89, no. 1, pp. 102–104, 1998.

[134] J. A. Jamart, M. C. Carracedo, and P. Casares, “Sexual isolation between Drosophila melanogaster females and D. simulans males. Male mating propensities versus success in hybridization,” Experientia, vol. 49, no. 6-7, pp. 596–598, 1993.

[135] M. C. Carracedo, C. Suarez, and P. Casares, “Sexual isolation between Drosophila melanogaster, D. simulans and D. mauritiana: sex and species specific discrimination,” Genetica, vol. 108, no. 2, pp. 155–162, 2000.

[136] M. Kreitman and M. Aguade, “Genetic uniformity in two populations of Drosophila melanogaster as revealed by filter hybridization of four-nucleotide-recognizing restriction enzyme digests,” Proceedings of the National Academy of Sciences of the United States of America, vol. 83, no. 10, pp. 3562–3566, 1986.

[137] D. J. Begun and C. F. Aquadro, “African and North American populations of Drosophila melanogaster are very different at the DNA level,” Nature, vol. 365, no. 6446, pp. 548–550, 1993.

[138] L. S. Stevison, K. B. Hoehn, and M. A. Noor, “Effects of inversions on within- and between-species recombination and divergence,” Genome Biology and Evolution, vol. 3, pp. 830–841, 2011.

[139] J. L. Feder, R. Gejji, T. H. Q. Powell, and P. Nosil, “Adaptive chromosomal divergence driven by mixed geographic mode of evolution,” Evolution, vol. 65, no. 8, pp. 2157–2170, 2011.

[140] D. B. Lowry and J. H. Willis, “A widespread chromosomal inversion polymorphism contributes to a major life-history transition, local adaptation, and reproductive isolation,” PLoS Biology, vol. 8, no. 9, Article ID e1000500, 2010.
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