Pollination syndromes are sets of floral traits that are adapted to different guilds of animal pollinators (Faegri 1979; Fenster et al. 2004). The floral traits that constitute a particular pollination syndrome can be separated into three categories: traits that attract the pollinator, such as floral color and scent; traits that reward the pollinator, such as nectar volume; and efficiency traits, such as the positioning of reproductive organs that affect pollen removal and deposition (Bradshaw et al. 1995). Although it is debated whether pollination syndromes can accurately predict the type of pollinating visitor to a flower, the pollination syndrome concept is a useful way to try to understand floral diversification and to frame questions about evolutionary transitions (Waser et al. 1996; Kingston and Mc Quillan 2000; Fenster et al. 2004; Olerton et al. 2009; Danieli-Silva et al. 2012). Pollination syndromes can be considered as a spectrum, with generalist species that produce flowers that appeal to many pollinators at one end and flowers that require pollination by specific pollinators at the other end (Fenster et al. 2004).

The force behind the convergent evolution of pollination syndromes across plant families is thought to be selection pressure exerted by the particular guilds of pollinators (van der Pijl 1960; Faegri 1979; Fenster et al. 2004). Animals have an innate preference for certain floral attributes and/or learn to associate the flower with providing a food source, such as nectar or pollen (Riffell 2013). In cases in which a variant arises that is more attractive to a particular pollinator, this will lead to a higher number of pollinator visits and may result in an increase in the fitness of the plant. This increase in fitness can cause the trait to spread as the variant allele increases in frequency in the population.

Floral constancy shown by the pollinator causing assortative mating may then lead to prezygotic isolation and possibly even speciation. This process has traditionally been thought of as coevolutionary, with plants also exerting selection on pollinators (Feinsinger 1983). However, more recent work in traits of attraction, color and scent has suggested that the plants have adapted their traits to sensory capabilities already present in the pollinators (Schiestl and Dötterl 2012), although there may be some fine-tuning of pollinator sensory preferences (Chittka and Menzel 1992; Raine and Chittka 2007).

Switches among syndromes are widespread. For instance, in the Solanaceae, hummingbird pollination is thought to have arisen at least 10 times (Knapp 2010). This raises the question of the genetic basis of the evolution of pollination syndromes. If each of the individual traits is encoded by multiple genes that must coordinately evolve, how is it possible that switches among syndromes happen frequently? To answer this question, it is necessary to identify the underlying plant genes that are under selection by animal pollinators. In their seminal study, Bradshaw et al. (1995, 1998) investigated the genetic bases for multiple divergent pollination syndrome traits in the monkey flower Mimulus. In its natural habitat, Mimulus lewisii is pollinated by bumblebees, whereas M. cardinalis is pollinated by birds. The two species live in habitats that are altitudinally separated except for a hybrid zone in which they overlap. Although the species are cross-compatible, they are almost completely reproductively isolated by the different pollinators that they attract (Schemske and Bradshaw 1999; Ramsey et al. 2003). To investigate the genetics defining the differences in pollination syndromes between these two species, a quantitative trait locus (QTL) analysis was performed. It was found that all of the pollination syndrome traits investigated (which included traits of pollinator attraction, reward, and efficiency) could be explained by one to six QTLS each (Bradshaw et al. 1995). This implies that there may only be a limited number of genes that determine
differences in individual pollination traits and that genes of large effect may have a central role in the pollinator-determined prezygotic isolation of plant species.

Quantitative trait loci analysis provides rough map positions, and each of the identified loci may still consist of multiple linked genes. Thus, the next logical step is to identify the genes underlying the QTL. The search for “pollination syndrome genes” in plants has been in full progress during the last 10 years. Once isolated, it is possible to determine the number and nature of the genetic changes required to change an individual floral trait. Do genetic changes preferably occur in structural or in regulatory genes and is there evidence for selection? Interspecific crosses followed by reciprocal introgressions of single loci also provide attractive material for pollinator choice studies, both under controlled laboratory conditions and in the natural habitat. They make it possible to determine the extent to which differences in single traits/loci condition differences in pollinator visitation and their effect on female and male fitness in the plant. Ultimately, transgenic lines can determine the effect of single genes.

It is important here to make the distinction between gene polymorphisms that have arisen in nature (natural variants) versus those that have arisen in the laboratory (mutants) or by breeding (varieties). Most mutants and varieties will have reduced overall fitness and are unlikely to survive in the wild. Thus, we focus here on natural color variants and the genes that determine these.

So far, the molecular basis of natural variation is known for only two pollination syndrome traits: petal color, specifically anthocyanin-based color, and scent production. We discuss the types of genes that have been found, their role in the attraction of different classes of pollinators, and the similarities among species. We focus specifically on anthocyanin-based petal color and volatile benzenoid-derived scent, both of which are derived from phenylalanine (see Fig. 1). We base our discussion around the genus *Petunia*, in which the biochemistry and molecular genetics of the two pathways have been worked out in considerable detail.

The genus *Petunia* comprises 14 species, 11 of which show a typical bee pollination syndrome (Stehmann et al. 2009). *Petunia integrifolia*, a representative of the bee clade, has a purple corolla with a wide tube that gives bees access to the nectar, and stamens and stigma within the length of the tube (Fig. 2A) (Stuurman et al. 2004). *Petunia axillaris* possesses a hawkmoth pollination syndrome (Fig. 2B), with white flowers that produce a strong scent at dusk. Hawkmoths reach the abundant dilute nectar by extending their proboscis through the long, narrow floral tubes (Stuurman et al. 2004). *P. integrifolia* conforms to its pollination syndrome, with visits primarily by

![Figure 1](symposium.cshlp.org)
bee species (Gübitz et al. 2009), whereas P. axillaris is pollinated by both nocturnal hawkmoths and diurnal bees (Hoballah et al. 2007). Petunia exserta has bright red petals that are reflexed, with the reproductive organs exserted from the floral tube (Fig. 2C) (Gübitz et al. 2009). These traits are typical for the hummingbird pollination syndrome, but detailed field observations of this rare species are scarce (Lorenz-Lemke et al. 2006).

**FLORAL COLOR**

**Floral Color Can Influence Pollinator Behavior in Different Ways**

Particular floral colors characterize different pollination syndromes and are associated with particular classes of pollinators. This must be conditioned in part by the perceptive abilities of the pollinators. All insects can see into the ultraviolet (UV) but have limited ability to see into the red end of the light wavelength spectrum (Menzel and Backhaus 1991). Pollinating birds have four photoreceptors with peak sensitivities in the UV, blue, green, and red light ranges (Herrera et al. 2008). The association of floral colors with particular pollinators is also likely to be affected by innate color preferences held by the pollinators. Innate preferences have been shown in many species to initially attract pollinators or aid recognition of flowers (Giurfa et al. 1995; Kelber 1997; Weiss 1997; Gumbert 2000). For instance, nectar guides are patterns of contrasting colors, often comprising “bulls-eye” spots at the mouth of the floral tube. Nectar guides increase pollinator efficiency by helping insects to orient on the flower and guide them to the food source more quickly than noninnately preferred colors (Giurfa et al. 1995). Color patterns can also influence the attractiveness of the flower and increase the efficiency of pollination. For example, nectar guides are patterns of contrasting colors, often comprising “bulls-eye” spots at the mouth of the floral tube. Nectar guides increase pollinator efficiency by helping insects to orient on the flower and guide them to the food source more quickly than they would without (Waser and Price 1981, 1983; Dinkel and Lunau 2001).

The question of whether an alteration in floral color alone can affect pollinator attraction was addressed in further work on *Mimulus* (Schemske and Bradford 1999; Bradshaw and Schemske 2003). The bumblebee-pollinated *M. lewisii* possesses pink flowers, whereas hummingbird-pollinated *M. cardinalis* has red flowers. The difference in color can be attributed to one locus, *yellow upper* (*yup*), from which the dominant allele from *M. lewisii* prevents carotenoid accumulation, whereas the homozygous recessive *yup* allele from *M. cardinalis* enables carotenoid accumulation (Bradshaw and Schemske 2003).

Reciprocal introgressions of the *yup* locus into the parental lines resulted in a yellow-orange *M. lewisii* line and a dark-pink *M. cardinalis* line. Small differences in six other traits important for pollinator visitation were present but were probably not functionally significant (Bradshaw and Schemske 2003). The differences in color

![Figure 2](https://example.com/figure2.png) Closely related *Petunia* species attract different animal pollinators: *Petunia integrifolia* with the solitary bee species *Callonychium petuniae* (A), *Petunia axillaris* ssp. axillaris with the hawkmoth *Hyles lineata*, Uruguay 2008 (B), *Petunia exserta* lab-line with the hummingbird *Hylocharis chrysura*, Uruguay 2009 (C). Photographs: Alexandre Dell’Olivo.
altered the preferences of the pollinators, decreasing the attraction of the species to its original pollinator and increasing the attractiveness to its noncognate pollinator. This shows that color likely has an important role in maintaining the reproductive isolation of these two Mimulus species. It also indicates that a change at a single locus can alter the pollinator class attracted to a plant. Although at this time the underlying gene (or genes) has not been identified, it seems likely that the yup locus has a role in the regulation or biosynthesis of carotenoid compounds.

Can a change in pollinator preference between two species be pinpointed to a difference in one gene? To study the role of flower color in pollinator attraction, we have used Petunia as a fitting model system. In Petunia, the contrasting color differences in the pollination syndromes are characterized by anthocyanin-based petal colors. The anthocyanin pathway is one of the best-characterized metabolic pathways in plants and much of the biochemical and molecular-genetic studies were performed in Petunia (Holton and Cornish 1995; Koes et al. 2005; Tornielli et al. 2009).

The Anthocyanin Pathway Is a Useful Model in which to Study Differences in Floral Color

Anthocyanins, a class of flavonoid, are the predominant floral pigment in angiosperms (Winkel-Shirley 2001). Other flavonoids also serve as floral pigments: aurones produce yellow flowers in some species (e.g., see Schwinn et al. 2006), whereas flavonols are important for absorption in the UV light range and can act as copigments in the presence of anthocyanins. Other components, such as the pH of the vacuoles in which the anthocyanins are stored, can also alter the light-absorption properties of anthocyanin compounds (Grotewold 2006).

The anthocyanin biosynthetic pathway is depicted in Figure 3. Excellent reviews have been written on anthocyanin biosynthesis and its regulation in Petunia and other species (Koes et al. 2005; Grotewold 2006; Quattrocchio et al. 2008; Tornielli et al. 2009). Here, we give a brief overview of the pathway and focus on the main genes that influence color. The early enzymes of the pathway convert phenylaniline to dihydroflavonols (dihydrokaempferol, dihydroquercetin, and dihydromyricetin), the precursors for both anthocyanins and UV-absorbing flavonols. Flavonol synthase (FLS) specifically catalyzes the synthesis of the flavonols, whereas dihydroflavonol reductase (DFR) is the first dedicated enzyme in the anthocyanin branch of the pathway. Anthocyanidin synthase (ANS) establishes the conjugated double-bond structure that makes the molecule appear colored to the human eye. The three main classes of anthocyanidins are orange-red pelargonidins, red-blue cyanidins, and blue-purple delphinidins. Further modification occurs by glucosyltransferases, acyltransferases, and rhamnosyltransferases, and methyltransferases produce “redder” compounds: peonidins from cyanidins, and petunidins and malvidins from delphinidins.

The type of anthocyanins produced can be influenced by the presence/absence as well as substrate specificities of the biosynthetic enzymes. P. integrifolia has highly decorated anthocyanins, such as malvidin-3-(p-coumaroyl)-rutinoside-5-glucoside (Fig. 3), whereas P. exserta produces simple anthocyanins, such as delphinidin-3-rutinoside and cyanidin-3-rutinoside (Fig. 3) (Ando et al. 1999). The low affinity of P. hybrida DFR for dihydrokaempferol and dihydromyricetin results in the production of mainly delphinidin-based anthocyanins (Gerats et al. 1982; Tornielli et al. 2009). This is likely attributable to the specific amino acid sequence in the putative substrate-binding region of DFR (Johnson et al. 2001). Competition for substrates may occur between the different branches of the anthocyanin pathway and/or between the anthocyanin and other branches of the flavonoid pathway. For instance, studies have shown that when a dominant flavonol (fl) locus is present in P. hybrida, the anthocyanin biosynthetic genes likely cannot compete against the flavonol biosynthetic gene FLS for the common substrate dihydroquercetin, resulting in a lower production of cyanidin-derived anthocyanins (Gerats et al. 1982).

Regulation of the Anthocyanin Pathway Is by a Conserved Group of Transcription Factors

Floral pigmentation mutants in Petunia have revealed a number of genes that activate anthocyanin structural genes. AN2, AN1, and AN11 code for a R2R3-MYB transcription factor (R2R3-MYB), a basic-helix-loop-helix (bHLH) transcription factor, and a WD40-repeat (WDR) protein, respectively. These proteins act in a complex (MYB-bHLH-WDR) to coordinately up-regulate the expression of late anthocyanin biosynthetic genes, from DFR onward (Beld et al. 1989; Quattrocchio et al. 1993). Another, as-yet-unidentified group of regulators must be responsible for expression of the early biosynthetic genes because they are expressed independently (Tornielli et al. 2009). It is proposed that this separate regulation provides a mechanism for the diversification of color while allowing other classes of flavonoids, with other roles in the plant, to be produced in tissues in which anthocyanin pigments are not present (Quattrocchio et al. 2008).

A number of closely related R2R3-MYBs confer pigmentation in specific tissues. The R2R3-MYB AN2 is expressed mainly in the petal limb in which it is the major regulator of anthocyanin production, whereas AN4 is primarily responsible for pigmentation in the anthers and floral tube (Quattrocchio et al. 1998, 1999; Kroon 2004). DEEPURPLE (DPL)/MYBB expression is correlated with flower tube venation, whereas another R2R3-MYB, PURPLEHAZE (PHZ), regulates anthocyanin production in light-exposed tissue in floral buds (Albert et al. 2011). A number of experiments that use ectopic and heterologous expression in mutants show that, in many cases, these R2R3-MYBs can compensate for the loss of function of native R2R3-MYBs, producing pigmentation in tissues in which they are expressed (Quattrocchio et al. 1998;
Figure 3. The flavonoid pathway leading to the production of anthocyanins and flavonols in *Petunia hybrida*. Enzymes are shown in red. Light-red enzyme names and gray arrows leading to the pelargonidins and myricetin indicate enzymatic reactions that do not occur in *Petunia hybrida*, owing to an inability of the enzyme to use the substrate. (3'AMT and 3'5'AMT) Anthocyanidin 3' and 3'5' O-methyltransferase, (3GT and 5GT) anthocyanidin 3 and 5 glucosyltransferase, (4CL) 4-coumarate CoA-ligase, (AAT) anthocyanidin 3-rutinoside acyltransferase, (ANS) anthocyanidin synthase, (ART) anthocyanidin 3-glucoside rhamnosyltransferase, (C4H) cinnamate 4-hydroxylase, (CHI) chalcone isomerase, (CHS) chalcone synthase, (DFR) dihydroflavonol reductase, (F3'H and F3'5'H) flavonoid 3' and 3'5' hydroxylase, (F3H) flavone 3-hydroxylase, (FLS) flavonol synthase, (Glc) glucose, (PAL) phenylalanine ammonia lyase, (Rha) rhamnose.
Spelt et al. 2000; Albert et al. 2011). Specific spatial expression of the R2R3-MYBs thus determines the production of anthocyanin compounds in only certain floral tissues (Tornielli et al. 2009).

In contrast, bHLH and WDR genes have wider expression profiles and also affect other processes in the plant. In Petunia, AN1 is expressed in pigmented tissues and is required for anthocyanin production, but it also has other functions besides pigmentation that influence vacuolar pH and morphology of the seed coat (Spelt et al. 2000, 2002). Another bHLH gene, JAF13, also has a role in anthocyanin pigmentation, although its exact role is unclear (Spelt et al. 2000). The WDR gene AN11 is expressed in all tissues, even in those that are nonpigmented (de Vetten et al. 1997). Its role has not been precisely characterized but it also may regulate vacuolar pH, and plants deficient in AN11 have defects in the epidermal cells of seeds and flowers as well as pigmentation loss in both tissues (Quattrocchio 1994; Zenoni et al. 2011). Repressors of anthocyanin production have also been characterized. The R3-MYB transcription factor MYBx is known to repress anthocyanin biosynthesis, possibly by competing with the R2R3-MYBs for the promoters of target genes (Kroon 2004).

The MYB-bHLH-WDR complex comprises the main classes of proteins involved in anthocyanin regulation across all species studied thus far (Koes et al. 2005; Grotewold 2006). Readers are referred to recent reviews for tables of these homologous regulators in different species (Koes et al. 2005; Petroni and Tonelli 2011; Davies et al. 2012). In dicotyledonous plants, the MYB-bHLH-WDR complex up-regulates late anthocyanin biosynthetic genes; however, the starting point of the “late” genes differs among species. In Petunia, F3′5′H is regulated along with DFR and subsequent genes (Brugliera et al. 1999), whereas in Ipomoea, the early biosynthetic genes CHS, CHI, and F3H may be partly regulated by the MYB-bHLH-WDR complex but are also independently regulated (Morita et al. 2006). This regulatory separation contrasts with the monocotyledonous plant maize, in which all anthocyanin biosynthetic genes are regulated together by the MYB-bHLH-WDR complex (Mol et al. 1998).

In other species, it is also likely that bHLH and WDR proteins are involved in additional processes unrelated to pigmentation while interacting with R2R3-MYBs that provide specific functions (Koes et al. 2005). In Antirrhinum and Ipomoea, it has also been shown that different MYB and vegetative tissues (Morita et al. 2006; Schwinn et al. 2006). In Arabidopsis, four highly related R2R3-MYB sequences similarly regulate late anthocyanin biosynthetic genes. These genes are not involved in floral anthocyanin production because Arabidopsis has white flowers and reproduces by self-fertilization; one of these R2R3-MYBs, PAP1, is necessary for anthocyanin production in young seedlings, whereas the function of the other genes is unknown (Gonzalez et al. 2008). This clade of Arabidopsis genes is classified as subfamily 6 of the R2R3-MYB transcription factor family (Stracke et al. 2001). Anthocyanin-implicated R2R3-MYBs from many species, including AN2 from Petunia, show homology with this subfamily (see Fig. 3 in Cooley et al. 2011).

Regulatory and Structural Genes Contribute to Natural Variation in Color

In the previous two sections, we discussed the major biosynthetic and regulatory genes in anthocyanin production. Natural variation in these genes may form the basis for color differences between species and variants. Most of these genes have been inactivated by mutations in P. hybrida. However, for some of these, mutational changes will not only result in shifts in petal color but also affect other pathways and, thereby, may negatively affect overall fitness (Coberly and Rausher 2003). So, which of these genes are likely targets of natural selection during shifts in pollination syndromes? This section gives an overview of the kind of molecular changes that have been attributed to causing color changes in natural populations of plant species. It also considers whether color variants that have arisen and spread in the population could be attributed to pollinator behavior. This information is summarized in Table 1. Examples are given in which a color polymorphism has been linked to a specific gene and the investigators have shown this by transgenic experiments or by genetic studies showing cosegregation of the gene with flower color. Studies of particular interest are discussed in greater detail below.

The first gene shown to be involved in a naturally occurring color transition was the R2R3-MYB transcription factor AN2 (Wijsman 1983; Quattrocchio et al. 1999). As described above, AN2 regulates the later part of the anthocyanin biosynthetic pathway. The gene is active in the purple flowers of P. integrifolia but is inactive in the white flowers of P. axillaris (Quattrocchio et al. 1999; Hoballah et al. 2007). The role of AN2 and color in pollinator attraction was shown by transforming a functional AN2 allele into the P. axillaris genetic background that produced a P. axillaris line with anthocyanin-pigmented flowers. Pollinator choice tests showed that the hawkmoth Manduca sexta (one of P. axillaris’ natural pollinators) preferred the white P. axillaris parent, whereas the bumblebee pollinator Bombus terrestris showed a preference for the transgenic line (Hoballah et al. 2007). This experiment showed that the modification of a single gene can have a major effect on pollinator behavior. The loss of function of AN2, and thus color, has happened multiple times, as shown by different mutations in populations of P. axillaris (Hoballah et al. 2007). Although it is debated whether the loss of AN2 function represented the first step in the divergence of these two species (Hoballah et al. 2007; A. Lorenz-Lemke, pers. comm.), the loss of color constitutes a necessary step in the transition from a bee pollination syndrome to a moth pollination syndrome in P. axillaris.

The genus Mimulus contains a yellow, carotenoid-colored clade, in which red, anthocyanin-colored flowers are thought to have independently arisen five times (Cooley and Willis 2009). Two of these gains of red color have been elucidated and have each been attributed to a single,
Table 1. Genetic polymorphisms associated with natural variants in floral anthocyanin-derived color and benzenoid floral volatiles in species that are pollinated by animal pollinators

<table>
<thead>
<tr>
<th>Taxon</th>
<th>Gene/s (locus)</th>
<th>Gene family Mutation</th>
<th>Molecular polymorphism</th>
<th>Role in pollinator preference</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antirrhinum</td>
<td>ROSEA1, ROSEA2 (rosea); VENOSA (venosa)</td>
<td>R2R3-MYB (subfamily 6)</td>
<td>R</td>
<td>Causal polymorphisms not yet identified in natural accessions. Differences in expression and target specificity of these genes contribute to variation in at least six species of <em>Antirrhinum</em>.</td>
<td>Schwinn et al. 2006; Whibley et al. 2006; Shang et al. 2011</td>
</tr>
<tr>
<td>Iochroma</td>
<td>F3(^5)H</td>
<td>CYP75A</td>
<td>S</td>
<td>Deletion of a F3(^5)H gene in <em>I. gesnerioides</em> compared to <em>I. cyaneum</em> leads to pelargonidin-derived anthocyanins instead of cyanidin-derived anthocyanins.</td>
<td>Smith et al. 2008; Smith and Rausher 2011</td>
</tr>
<tr>
<td>Ipomoea purpurea</td>
<td>F3(^5)H (P)</td>
<td>CYP75B</td>
<td>S</td>
<td>A loss-of-function mutation in the F3(^5)H gene causes <em>pp</em> individuals to be unable to produce cyanidin-derived anthocyanins.</td>
<td>Zufall and Rausher 2003</td>
</tr>
<tr>
<td>Ipomoea purpurea</td>
<td>CHS-D (A)</td>
<td>CHS</td>
<td>S</td>
<td>A loss-of-function mutation in the CHS-D gene causes <em>aa</em> individuals to be unable to produce anthocyanins in flowers and also affects flavonoid production in other tissues.</td>
<td>Coberly and Rausher 2003; Fehr and Rausher 2004</td>
</tr>
<tr>
<td>Ipomoea purpurea</td>
<td>IpMYB1 (W)</td>
<td>R2R3-MYB (subfamily 6)</td>
<td>R</td>
<td>A loss-of-function mutation in IpMYB1 is correlated with the down-regulation of multiple anthocyanin biosynthetic genes in <em>ww</em> individuals and loss of pigmentation.</td>
<td>Epperson and Clegg 1987; Tiffin et al. 1998; Chang et al. 2005</td>
</tr>
<tr>
<td>Ipomoea quamoclit</td>
<td>F3(^5)H</td>
<td>CYP75B</td>
<td>R</td>
<td>Causal polymorphism not yet identified. A <em>cis</em>-regulatory change to F3(^5)H affects expression of the gene and influences flux between cyanidin-derived anthocyanins and pelargonidin-derived anthocyanins.</td>
<td>Des Marais and Rausher 2010</td>
</tr>
<tr>
<td>Mimulus aurantiacus</td>
<td>MYB</td>
<td>R2R3-MYB (subfamily 6)</td>
<td>R</td>
<td>Causal polymorphism not yet identified. A <em>cis</em>- or <em>trans</em>-regulatory change in R2R3-MYB influences color in a complex fashion in yellow- and red-flowered races of <em>M. aurantiacus</em>.</td>
<td>Streisfeld and Kohn 2005, 2007; Streisfeld and Rausher 2009a</td>
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(Continued)
### Table 1. (Continued)

<table>
<thead>
<tr>
<th>Taxon</th>
<th>Gene/s (locus)</th>
<th>Gene family</th>
<th>Mutation</th>
<th>Molecular polymorphism</th>
<th>Role in pollinator preference</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Mimulus cupreus</em></td>
<td>McMYB1, McMYB2, McMYB3 (pla1)</td>
<td>R2R3-MYB (subfamily 6)</td>
<td>R</td>
<td>Causal polymorphism not yet identified. The <em>M. cupreus</em> <em>pla1</em> locus and, in particular, McMYB2 expression are correlated with higher expression of anthocyanin biosynthetic genes compared to alleles from the yellow morph of <em>M. cupreus.</em></td>
<td>Pollinator visitation rate by <em>Bombus dahlbomii</em> to <em>M. cupreus</em> in the population of central Chile was low compared to other <em>Mimulus</em> species at the same study site. Discrimination was not associated with flower color. Population at this site is thought to be maintained by selfing.</td>
<td>Cooley et al. 2008, 2011</td>
</tr>
<tr>
<td><em>Mimulus luteus</em></td>
<td>MvMYB4, MvMYB5 (pla2)</td>
<td>R2R3-MYB (subfamily 6)</td>
<td>R</td>
<td>Causal polymorphism not yet identified. The <em>M. l. variegatus</em> allele of MvMYB5 correlated with higher expression of anthocyanin biosynthetic genes compared to the allele from yellow <em>M. l. luteus.</em></td>
<td>Pollinator visitation by single, generalist pollinator in the population in central Chile did not show discrimination between red <em>M. l. variegatus</em> and yellow <em>M. l. luteus.</em> Some assortative mating may occur owing to preferences of individual pollinators.</td>
<td>Cooley et al. 2008, 2011</td>
</tr>
<tr>
<td><em>Petunia</em></td>
<td>AN2</td>
<td>R2R3-MYB (subfamily 6)</td>
<td>R</td>
<td>Functional gene regulates production of anthocyanins in the corolla of <em>P. integrifolia.</em> At least five independent loss-of-function mutations in the coding region contribute to down-regulation of anthocyanin biosynthetic genes in the corolla of <em>P. axillaris.</em></td>
<td>Introduction of functional AN2 into white <em>P. axillaris</em> partially restored anthocyanin production. The pigmented, transgenic line showed fourfold less feedings by <em>Manduca sexta</em> and threefold more visits by <em>Bombus terrestris</em> compared to wild type in pollinator-choice assays in controlled conditions.</td>
<td>Quattrocchio et al. 1999; Hoballah et al. 2007</td>
</tr>
<tr>
<td><em>Petunia</em></td>
<td>ODO1</td>
<td>R2R3-MYB (subfamily undefined)</td>
<td>R</td>
<td>Causal polymorphisms not yet identified. A cis-regulatory change is suggested to cause expression differences.</td>
<td>NILs that differed only in floral scent production were bred into the genetic backgrounds of <em>P. axillaris</em> and <em>P. exserta.</em> <em>Manduca sexta</em> moths showed a significant preference for scented lines over nonscented lines. No preference was displayed when confronted with conflicting visual and olfactory cues.</td>
<td>Klahre et al. 2011</td>
</tr>
<tr>
<td><em>Phlox drummondii</em></td>
<td>MYB</td>
<td>R2R3-MYB (subfamily 6)</td>
<td>R</td>
<td>Causal polymorphisms not yet identified. A cis-regulatory change affects expression of the <em>MYB</em>, which is correlated with the expression level of anthocyanin biosynthetic genes.</td>
<td>Sympatric populations of <em>P. drummondii</em> with <em>P. cuspidata</em> show dark red flower color; allopatric populations are light blue. Common garden-field experiment showed reduced hybridization between light- and dark-colored plants caused by color-intensity constancy by pollinators.</td>
<td>Hopkins and Rausher 2011, 2012; Hopkins et al. 2011</td>
</tr>
<tr>
<td></td>
<td>C3'5'H</td>
<td>CYP75A</td>
<td>R</td>
<td>A cis-regulatory change affects expression of <em>F3'5'H</em> and influences color hue.</td>
<td>No evidence for discrimination by color hue, despite previous study showing selection on the “red” allele.</td>
<td>Lile 2007; Lile et al. 2009</td>
</tr>
</tbody>
</table>

The Mutation column refers to the type of mutation that has occurred: either affecting the coding region of a structural (S) gene or affecting regulatory (R) sequences (either a cis-regulatory element or the coding sequence of a transcription factor). Where available, information has been included about the role of the genetic polymorphism in pollinator preference and/or selection by pollinators.

(AN2) Anthocyanin2, (CHS) chalcone synthase, (CYP) cytochrome P450 monoxygenase, (DFR) dihydroflavonol reductase, (F3’H and F3’5’H) flavonoid 3’ and 3’5’ hydroxylase, (NIL) near isogenic line, (ODO1) odorant1.
POLLINATION SYNDROME GENES

dominant gain-of-function mutation in loci containing R2R3-MYBs of subfamily 6. The locus, petal lobe anthocyanin 1 (pla1), is associated with red color in the species M. cupreus, whereas the locus pla2 produces red pigmentation in M. luteus variegatus (Cooley and Willis 2009; Cooley et al. 2011).

The pla1 locus may also be associated with the production of anthocyanin-pigmented corolla banding and calyx spotting in M. guttatus (Lowry et al. 2012). Each locus contains duplicated homologous R2R3-MYBs and the loci themselves are likely to be related by a previous duplication event (Cooley et al. 2011). For both loci, it was shown that the level of anthocyanins is correlated with the coordinate up-regulation of both early and late anthocyanin biosynthetic genes, indicating that the R2R3-MYBs function as regulators of the anthocyanin pathway (albeit there may be some differences to other dicotyledonous species). The exact role of each R2R3-MYB in influencing the difference in color between the yellow ancestors and derived red species has not yet been elucidated. However, this example of parallel evolution shows the central role of R2R3-MYB factors in determining color differences in Mimulus. Similarly, two trans-acting regulatory loci are implicated in determining color differences between parapatrically distributed red- and yellow-flowered races of M. aurantiacus, a species from a separate Mimulus clade (Streisfeld and Rausher 2009a). These races are likely under selection by their hawkmoth and hummingbird pollinators who show strong preferences for flower color (Streisfeld and Kohn 2007). Conversely, little evidence was found to suggest that flower color in M. cupreus, M. l. variegatus, and a yellow-flowered species M. l. luteus influences pollinator behavior (Cooley et al. 2008). An alternative explanation for the occurrence of red-flowered species is that selection on other traits has caused anthocyanin pigmentation to arise and it may be a pleiotropic effect of the biosynthetic pathways controlling pigmentation (Cooley and Willis 2009).

A member of the R2R3-MYB subfamily 6 has also been shown to be responsible for a gain in red color in Phlox. In this example, the change in the R2R3-MYB works in concert with a cis-regulatory change to the F3′5′H gene. Phlox drummondii has light blue flowers similar to another species, P. cuspidata. Where these species are found in sympatry, P. drummondii produces red flowers. The level of anthocyanins produced by the red flowered P. drummondii is higher than that in P. cuspidata, and this is correlated with a higher expression of R2R3-MYB in P. drummondii (Hopkins and Rausher 2011). The up-regulation of R2R3-MYB is caused by an as-yet-unidentified change in a cis-regulatory element of the P. drummondii allele. The change in hue is attributed also to a cis-regulatory difference acting on F3′5′H, meaning that flux in the pathway is redirected from the malvidin branch of the pathway to the cyanidin branch (Hopkins and Rausher 2011). Field experiments showed that the “color intensity” R2R3-MYB locus maintains reproductive isolation of the red-flowered P. drummondii from P. cuspidata by decreasing interspecific pollen flow. Surprisingly, the F3′5′H locus affecting hue does not seem to be important, with current knowledge, for maintaining the species (Hopkins and Rausher 2012), despite the previous finding of a selective sweep on the allele causing red color (Hopkins et al. 2011). This highlights the necessity to compare population genetics studies with tests of pollinator preference and fitness differences in the field.

In Ipomoea, a different mechanism has been shown to be responsible for the production of red, bird pollinated flowers. The ancestral color state is believed to be blue-purple flowers which are pollinated by bees, with red-flowered hummingbird pollinated species having arisen four times independently (Streisfeld and Rausher 2009b). Ipomoea quamoclit represents one member of a clade of red-flowered species. In this species, Des Marais and Rausher (2010) showed that downregulation of the biosynthetic gene, F3′H, at least partly as a result of mutation in a cis-regulatory element, resulted in flux in the pathway being directed from cyanidin-derived anthocyanins to pelargonidin-derived anthocyanins. This regulation is tissue-specific, occurring in the floral tissue but not in the vegetative tissue. The investigators speculate that this change characterizes the other species in the I. quamoclit clade, and that it is similar to what has occurred in another red-flowered lineage of Ipomoea, I. horsfalliae. In this species, F3′H is also downregulated, but the causal mechanism has not been found (Streisfeld and Rausher 2009b). The investigators postulate that F3′H could be a common target of mutations to transition from bee pollination to hummingbird pollination; many red-flowered angiosperm species which have gone through a transition from blue-purple flowers are colored by pelargonidin-derived anthocyanins (Rausher 2008; Streisfeld and Rausher 2009b). It remains to be established whether the transition to red flowers in Ipomoea is as a result of selection from pollinators.

In all the above examples, color changes have involved changes to the regulation of the anthocyanin pathway—either regulatory genes (trans-regulatory changes) or the targets of regulatory genes, regulatory elements (cis-regulatory changes). However, examples of changes to color caused by mutations in structural genes are also present. The common morning glory Ipomoea purpurea has populations that are highly polymorphic for color. All of these polymorphisms have been attributed to variation in four loci (Epperson and Clegg 1988). The genetic basis for three of these loci has been elucidated and two of them result from mutations to biosynthetic genes. For the P locus, a mutation in the coding region of the F3′H gene produces a truncated protein that cannot use dihydroquercetin and, thus, cannot produce cyanidin-derived anthocyanins (Zufall and Rausher 2003). Flux is redirected from the cyanidin branch of the biosynthetic pathway to the pelargonidin branch, thereby producing red flowers in a similar way as to that described for I. quamoclit. The A locus produces a class of white-colored variants of I. purpurea that cannot produce anthocyanins owing to a transposon insertion in the CHS-D gene (Johzuka-Hisatomi et al. 1999). This loss-of-function mutation in the CHS-D gene means that neither anthocyanins nor other flavonoid compounds can be produced throughout the whole plant. These mutants show decreased fitness and
this is likely attributable to a deficiency in flavonoid compounds that function to ameliorate heat stress (Coberly and Rausher 2003). The third locus represents a regulatory change controlled by IpMYB1, another member of the R2R3-MYB subfamily 6, and is also responsible for producing a class of white-flowered variants (Chang et al. 2005). In this example, white-flowered variants do not appear to be under selection by pollinators (for review, see Clegg and Durbin 2003).

The Role of Ultraviolet Color May Be Linked to Anthocyanin Pigmentation

Insects, pollinating birds, and bats have photoreceptors for UV light (Menzel and Backhaus 1991; Winter et al. 2003; Herrera et al. 2008), and many flowering plants absorb in the UV. UV-absorbing nectar guides that help to orient insects with regard to the source of nectar have been known for some time (Daumer 1958; Jones and Buchmann 1974; Penny 1983). Recent work in Mimulus has shown that UV-absorbing petals can be important for the attraction of bees and other pollinators in a natural habitat and also reaffirmed that these petals can be important for bee orientation (Rae and Vamosi 2012).

UV-absorbing floral pigments may be useful for detection of flowers by night-active insects. Night-active hawkmoths generally visit white-flowered plants, such as Petunia axillaris. These flowers’ maximal reflectance seems optimal for the flower to be perceived at low-light intensity. However, most white flowers absorb UV light and, thus, do not reflect all wavelengths perceived by the insect’s photoreceptors and do not take advantage of all available photons (Kevan et al. 1996). Interestingly, hawkmoths have been shown to display true color vision at light intensities as low as dim starlight (Kelber et al. 2002). Therefore, hawkmoths probably use color vision to detect UV-absorbing flowers against a background of “dull” foliage that reflects to a low degree across all wavelengths (Kevan et al. 1996).

In Petunia, flavonols are the main pigments contributing to the UV absorbance of the flower. The absence of anthocyanins and the presence of flavonols in the P. axillaris flower may be attributable to selection by hawkmoths, but alternative roles of flavonols in plant defense against pathogens or abiotic stresses should also be considered (Gould and Lister 2005). Another possibility is that the presence of flavonols is a consequence of selection for loss of anthocyanin biosynthesis, because down-regulation of DFR would redirect metabolic flux toward FLS and flavonol biosynthesis (Fig. 1). Molecular genetics in combination with behavioral experiments can test such hypotheses.

FLORAL SCENT

Floral Volatiles Act as Species-Specific Long-Distance Signals to Attract Pollinators

Floral scent bouquets may consist of as many as 20–60 different volatiles (Knudsen et al. 2006), synthesized from diverse primary metabolites (Dudareva et al. 2006). Most floral volatiles belong to fatty acid derivatives, terpenoids, or benzenoids. The number and concentration of compounds provide species-specific olfactory cues (Raguso et al. 2003; Hoballah et al. 2005; Raguso 2008), and the combination with visual stimuli allows searching pollinators to use different sensory channels to identify target flowers.

Bats, bees, butterflies, hummingbirds, and moths are able to sense floral scents and have been shown to discriminate among odor differences (von Helversen et al. 2000; Kunze and Gumbert 2001; Wright et al. 2002, 2005; Ōmura and Honda 2005; Kessler et al. 2008; Zvi et al. 2012). Recently, it was shown that the hawkmoth Manduca sexta processes olfactory stimuli through two olfactory channels, allowing it to coordinate an innate preference with olfactory learning (Riffell et al. 2012). Hawkmoths use odor cues to locate target plants over long distances (Raguso and Willis 2002a), which may explain the high odor emission rates in flowers displaying moth-pollination syndromes. For instance, moth-pollinated Clarkia breweri flowers emit strong floral scents compared to flowers of the closely related bee-pollinated C. concinna (Raguso and Pichersky 1995). However, the importance of olfactory and visual stimuli for the attraction of specific pollinator guilds differs (Raguso and Willis 2005; Goyret et al. 2007; Klahre et al. 2011).

The Biosynthesis of Volatile Benzenoid Compounds

Petunia x hybrida cv Mitchell is a popular model system to elucidate the biosynthesis of floral volatile benzenoids, and many of the structural and regulatory genes have been identified in Petunia. For other odor compounds, the reader can refer to excellent reviews of Pichersky and Gershenzon 2002; Dudareva et al. 2006.

Phenylalanine-derived volatiles come in three groups that are identified by the number of carbon atoms in their side chains (Fig. 4; all volatile benzenoid compounds are highlighted in purple). In the C$_6$–C$_2$ group, the key enzyme phenylacetalddehyde synthase (PAAS) catalyzes the decarboxylation of phenylalanine to phenylacetaldehyde, which can subsequently be reduced, oxidized, or esterified.

The C$_6$–C$_3$ and C$_6$–C$_1$ pathways both start from trans-cinnamic acid, which is synthesized from phenylalanine by the activity of phenylalanine ammonia lyase (PAL). Entry into the C$_6$–C$_1$ pathway requires shortening of the C$_3$ side chain of trans-cinnamic acid by two carbons, which can occur via both a CoA-dependent β-oxidative pathway analogous to the catalolism of fatty acids (Hertweck et al. 2001; Moerkercke et al. 2009; Klempien et al. 2012; Qualley et al. 2012) and a CoA-independent non-β-oxidative pathway (Boatright et al. 2004; Orlova et al. 2006).

The conversion of trans-cinnamic acid into p-coumaric acid leads into the C$_6$–C$_3$ branch. The specific reaction that gives rise to coniferyl alcohol is as-yet unknown.
Figure 4. The benzenoid pathway leading to the production of volatile compounds in scented Petunia. Volatile compounds emitted by scented Petunia are highlighted in purple. Enzymes shown in red indicate known steps in Petunia; unknown enzymatic steps are indicated by dashed arrows. Multiple dashed arrows indicate multiple unknown steps. Transcriptional regulators are highlighted in yellow. (4CL) 4-Coumarate-CoA ligase, (ADT) arogenate dehydratase, (BALDH) benzaldehyde dehydrogenase, (BSMT) (S-adenosyl-L-methionine) benzoic acid/salicylic acid carboxyl methyltransferase, (BPBT) benzoyl-CoA-benzylalcohol/2-phenylethanol benzoyltransferase, (C4H) cinnamate 4-hydroxylase, (CFAT) coniferyl alcohol acyltransferase, (CHD) cinnamoyl-CoA hydratase-dehydrogenase, (CM) chorismate mutase, (CNL) cinnamate CoA-ligase, (DAHPS) 3-deoxy-D-arabino-heptulosonate-7-phosphate synthase, (EGS) eugenol synthase, (EPSPS) 5-enolpyruvylshikimate-3-phosphate synthase, (IGS) isoeugenol synthase, (KAT1) 3-ketoacyl-CoA thiolase, (PAAS) phenylacetaldehyde synthase, (PAL) phenylalanine ammonia lyase.
Coniferyl alcohol acyltransferase (CFAT) acetylates coniferyl alcohol to produce coniferyl acetate. This serves as the substrate for isoeugenol synthase (IGS) and eugenol synthase (EGS), which catalyze the formation of isoeugenol and eugenol, respectively (Clark 2009). Some \textit{P. axillaris} emit neither eugenol nor isoeugenol but produce high levels of dihydroconiferyl acetate. In \textit{P. a. paradox} \textit{S7}, EGS is functional, but the gene encoding IGS contains a frameshift mutation leading to its inactivity. Unexpectedly, this does not lead to higher eugenol levels (Koeduka et al. 2009).

Because all benzenoid volatiles derive from phenylalanine, substrate competition may regulate the fluxes through the $C_6-C_2$, $C_6-C_1$, and $C_6-C_3$ pathways. Such substrate competition may be determined by the relative abundance and enzymatic properties of the branch-specific enzymes. Indeed, there is evidence that a higher affinity for phenylalanine by PAL compared to PAAS causes a higher flux through the $C_6-C_1$ and $C_6-C_3$ branches of the pathway (Maeda et al. 2010).

The Regulation of Floral Scent Production

The emission of floral volatile benzenoids from the epidermal cells of \textit{P. axillaris} and \textit{P. hybrida} cv Mitchell petals begins at anthesis and ceases after pollination (Negre et al. 2003; Underwood et al. 2005). Emission follows a circadian rhythm that peaks at dusk (Verdonk et al. 2005; Clark 2009). Together, this indicates precise spatial (Van Moerkercke et al. 2012), developmental, and temporal regulation of the pathway. So far, three transcriptional regulators of scent production, ODO1, EMISSION OF BENZENOIDS II (EOBII), and PhMYB4, have been identified. Expression of the \textit{R2R3-MYB} \textit{ODO1} peaks before the onset of volatile emission and its transcript levels are low the next morning. ODO1 activates the promoter of the gene encoding EPSPS, an enzyme of the shikimate pathway, suggesting that ODO1 is involved in the petal-specific up-regulation of phenylalanine as the precursor for volatile production (Verdonk et al. 2005). Accordingly, down-regulation of \textit{ODO1} in transgenic \textit{P. hybrida} cv Mitchell plants strongly reduced benzenoid emission (Verdonk et al. 2005). The expression levels of genes encoding DAHPS, EPSPS, PAL, and CM were decreased, but, interestingly, the late-acting genes encoding BSMT and BPBT were up-regulated. Expression of the structural genes BSMT, BPBT, together with PAAS and IGS, precedes volatile emission and has been detected at high levels during the light period and low levels during the dark period (Colquhoun et al. 2010). However, the rhythmic transcription accumulation of at least BSMT and PAAS seems not to cause the rhythmic emission of the floral volatiles, based on constant activities detected over a 24-h period (Kolosova et al. 2001). Instead, oscillations of early precursor pools and \textit{ODO1} expression have been suggested to control the rhythmic emission of floral fragrance.

The second \textit{R2R3-MYB} transcription factor, EOBII, binds to a specific MYB-binding site located in an upstream enhancer region to activate the expression of \textit{ODO1} (Van Moerkercke et al. 2011). EOBII also activates \textit{PAL} and \textit{IGS} promoters (Spitzer-Rimon et al. 2010). Silencing of \textit{EOBII} down-regulates \textit{ODO1} and genes of the shikimate and phenylpropanoid pathways. As with \textit{ODO1}, it has a rhythmic expression pattern, but the transcript levels peak in the early morning hours and are lowest during the late afternoon. The difference in phasing of the \textit{ODO1} and \textit{EOBII} rhythms might be explained by time-delayed activation of \textit{ODO1} expression. But \textit{EOBII} is expressed in early flower development when \textit{ODO1} is not. Functional analyses of \textit{ODO1} promoter fragments have provided evidence that at least one additional factor is needed for high \textit{ODO1} promoter activity (Van Moerkercke et al. 2012). This as-yet-unidentified factor might partly explain the interaction of \textit{EOBII} and \textit{ODO1} despite their different expression patterns and the finding that overexpression of \textit{EOBII} did not result in significant higher \textit{ODO1} levels (Spitzer-Rimon et al. 2010).

The third identified transcription factor, PhMYB4, is also an \textit{R2R3-MYB} transcription factor. It negatively regulates C4H (Colquhoun and Clark 2011; Colquhoun et al. 2011), indirectly controlling the emission of isoeugenol and eugenol by precursor supply. It was hypothesized to “fine-tune” the bouquet of floral fragrance by affecting the balance between the $C_6-C_1$ and $C_6-C_3$ branches of the pathway.

Interaction Between Color and Scent at the Biochemical Level

Interaction between the branches of the benzenoid and flavonoid pathways might occur owing to pleiotropic regulation or to metabolic competition for the common precursor, phenylalanine. Redirective metabolic flux from one branch to the other has been shown in \textit{Dianthus caryophyllus} (Zuker et al. 2002). These investigators showed that suppression of \textit{F3’H} led to white-flowered mutants that produced higher amounts of methylbenzoate compared to colored control plants. In contrast, the introduction of the \textit{Arabidopsis} \textit{R2R3-MYB} transcription factor \textit{PRODUCTION OF ANTHOCYANIN PIGMENT 1} (PAP1; a member of the \textit{R2R3-MYB} subfamily 6) into \textit{P. hybrida} enhanced both branches of the phenylpropanoid pathway, leading to the production of color and scent. This argues against metabolic competition but supports the notion that PAP1 has a pleiotropic effect on both pathways (Zvi et al. 2006, 2008). Similarly, \textit{35S:PAP1} transgenic \textit{Rosa hybrida} showed increased pigmentation as well as eugenol accumulation (Zvi et al. 2012). A more general argument against metabolic competition is the temporal separation of the two pathways. Flavonoids are produced during growth and development of the flower, whereas benzenoids are only produced when the flowers open (Verdonk et al. 2005).

However, multiple examples contradict the idea of a pleiotropic regulator of scent and color. The introduction of the key regulator of anthocyanin biosynthesis, AN2,
into *Petunia axillaris* altered flower color but had no effect on scent production (Hoballah et al. 2007), suggesting that this *Petunia* PAP1 homolog has no effect on volatile emission. Silencing *ODO1* and thus decreasing shikimate precursor compounds for benzenoid production did not lead to decreased flavonoid production (Verdonk et al. 2005). In addition, *EOBII* expression in transgenic *Nicotiana tabacum* increased PAL transcript levels but *CHS* and *F3H* expression levels and anthocyanin levels were not affected (Spitzer-Rimon et al. 2010). Because pleiotropic effects have been observed when foreign transgenes were expressed from a strong viral promoter, the evidence in favor of regulatory interactions between the two pathways should be interpreted with caution.

### Behavioral Experiments Test the Importance of Color and Scent in Pollinator Attraction

Classical behavioral studies have tried to disentangle the importance of olfactory and visual stimuli for pollinator attraction using artificial flowers and cues (Raguso and Willis 2002b; Balkenius and Kelber 2006; Balkenius et al. 2006; Goyret et al. 2007). To disentangle visual and olfactory cues in complex flowers, Hirota et al. (2012) generated F₁ and F₂ hybrids between the butterfly-pollinated daylily and the hawkmoth-pollinated nightlily, each differing in floral color and scent. In an F₂ progeny, segregating for both floral traits, neither butterflies nor hawkmoths showed significant preferences for overall scent emission. However, butterflies preferentially visited red- or orange-colored flowers, resembling the daylily color phenotype, whereas moths preferred the yellow-colored flowers, representing the nightlily color phenotype. Thus, the visual stimuli were more important for pollinator attraction. It should be noted, however, that additional floral traits might have been segregating in a F₂ population that were not considered in this study.

In *Petunia*, we set out to construct near isogenic lines (NILs) through repeated backcrossing of hybrids among species with different pollination syndromes. The aim was to obtain plants that differ genetically only in single relevant loci, and phenotypically only in single traits, for instance, petal color or scent production. This makes it possible to dissect the interactions between visual and olfactory stimuli in complex flowers. Such plants can be studied under controlled conditions but can also be introduced into the natural habitats. They can thus be exposed to the variety of natural pollinators visiting during the season, and fitness across generations can be determined. In addition, NILs are attractive genetic material for the isolation of the genes underlying shifts in pollination syndromes.

Using this approach, we crossed the red-flowered, nonscented *P. exserta* with the white, scented *P. axillaris* and mapped two major QTL for scent production in the F₂ population to chromosomes II and VII. The scent QTL on chromosome VII covered the position of *ODO1* (Stuurman et al. 2004; Klahre et al. 2011). *ODO1* is highly expressed in *P. axillaris* but is present at low levels in *P. exserta*. In the F₁ hybrid, most of the *ODO1* transcript was produced from the *P. axillaris* allele, indicating that the difference in expression level of the two alleles is a cis-acting effect (Klahre et al. 2011). The actual polymorphism, potentially a sequence change in the *ODO1* promoter, has not yet been determined. The gene(s) underlying the chromosome II QTL are presently unknown. However, repeated backcrossing resulted in an unscented *P. axillaris* NIL as well as a scented *P. exserta* NIL and made it possible to study hawkmoth attraction to scent in otherwise indistinguishable genetic backgrounds. Both in the *P. axillaris* and *P. exserta* backgrounds, the moths had a strong preference for the scented lines over the nonscented lines. Interestingly, when the insects were given the choice between the red, scented *P. exserta* NIL and the white, nonscented *P. axillaris* NIL, a situation in which they had to choose between conflicting visual and olfactory cues, they displayed no preference. This suggests that, at least under the artificial conditions of a wind tunnel experiment, color and scent appear to be equivalent cues (Klahre et al. 2011).

### CONCLUSIONS

Pollination syndromes consist of specific traits, such as floral color, scent, floral morphology, and nectar volume and composition. The correlation between floral traits and pollinator classes across the angiosperms suggests that this convergent evolution is driven by pollinator-mediated selection (Fenster et al. 2004). However, other contributing factors must also be considered. Rausher (2008) discusses this in depth, pointing out that the few examples investigated thus far show selection on color changes but that the selective agent cannot necessarily be attributed to pollinators.

To attract a new pollinator, coordinated changes in multiple traits are required. This raises important questions about the presumably complex molecular genetics of shifts in pollination syndromes. This review focuses on the molecular basis of the pollination syndrome traits, color and scent. For both traits, substantial knowledge has been acquired about the biochemical pathways and transcriptional regulators. A central question in the ecology and evolution of pollination syndromes is which of the trait-determining genes were selected in nature to achieve a shift in pollinator attraction. Altogether, how many genes were involved and how did they change in a coordinated fashion? For color and scent, both structural and regulatory genes contribute to major phenotypic changes, although more regulatory changes are represented in the limited number of examples found so far (Table 1). Genes of small effect would in most cases have escaped detection, but the fact that major genes were found at all is of great significance to the debate about the role of major mutations in adaptive evolution (Bradshaw and Schemske 2003).

The first transcription factor involved in natural variation in petal color was AN2 and *R2R3-MYBs* are the only regulatory genes represented so far (Table 1). Subsequent discovery of genes underlying natural variation has relied on candidate gene approaches that are inevitably biased...
toward R2R3-MYBs. We note that no genes causing natural variation in pollination syndromes have yet been identified de novo. Changes consist of alterations in the coding region of the R2R3-MYBs themselves or changes in cis-regulatory elements. Other regulatory changes concern putative cis changes or changes to the coding sequences of biosynthetic genes, for instance, F3′H or F3′5′H genes. Such changes have effects on the types of anthocyanins produced by adjusting flux through branches of the biosynthetic pathways. Duplication also had a role in the diversification of color, with many of the described examples involving duplication of R2R3-MYBs. Duplication is thought to be a significant factor influencing phenotypic novelty in plants (Flagel and Wendel 2009).

R2R3-MYBs are also involved in the regulation of volatile benzenoid biosynthesis. The R2R3-MYBs EOBII and ODO1 regulate scent biosynthesis in different species and there is evidence that ODO1 underlies natural variation in scent production. However, both ODO1 and EOBII primarily affect early steps in the pathway and alterations to these genes might be expected to have pleiotropic effects. It seems likely, therefore, that additional regulators of natural variation in scent remain to be discovered.

Coordinated changes in multiple traits could be brought about by genetic changes in master genes that affect multiple traits or by genetic linkage of multiple regulators. There is little convincing evidence thus far for the involvement of coregulation of scent and color in natural variants despite regulators being found that influence early, shared steps of the phenylpropanoid pathway. Linkage of genes in the genome may have occurred in Mimulus and Aquilegia (Bradshaw et al. 1995; Hodges et al. 2002), but much more detailed genetic and physical maps, as well as a better understanding of the genetic and biochemical basis of the traits, will be necessary to address this issue.

Undoubtedly, the rapid advances in whole-genome sequencing will accelerate the identification of the genes that cause natural variation. Functional analysis of such genes, at the level of the gene, trait, and population, will provide new insight not only into plant–pollinator interactions but also in reproductive isolation and ultimately speciation.

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