The regulatory network for petal anthocyanin pigmentation is shaped by the MYB5a/NEGAN transcription factor in *Mimulus*

Xingyu Zheng,1,2 Kuenzang Om,3 Kimmy A. Stanton,3 Daniel Thomas,3 Philip A. Cheng,3 Allison Eggert,3 Emily Simmons,1 Yao-Wu Yuan,3,4 Gregory D. Conradi Smith,1 Joshua R. Puzey,1,*† and Arielle M. Cooley3,*,†

1Departments of Biology and Applied Science, William & Mary, Williamsburg, VA 23185, USA
2School of Biological Sciences, Cold Spring Harbor Laboratory, Cold Spring Harbor, NY 11724, USA
3Department of Biology, Whitman College, Walla Walla, WA 99362, USA
4Department of Ecology and Evolutionary Biology, University of Connecticut, Storrs, CT 06269, USA

*These authors contributed equally to this work.

†These authors contributed equally to this work.

*Corresponding authors: puzeyjr@wm.edu. (J.R.P.); cooleya@whitman.edu (A.M.C.)

**Abstract**

Much of the visual diversity of angiosperms is due to the frequent evolution of novel pigmentation patterns in flowers. The gene network responsible for anthocyanin pigmentation, in particular, has become a model for investigating how genetic changes give rise to phenotypic innovation. In the monkeyflower genus *Mimulus*, an evolutionarily recent gain of petal lobe anthocyanin pigmentation in *M. luteus* var. *variegatus* was previously mapped to genomic region *pla2*. Here, we use sequence and expression analysis, followed by transgenic manipulation of gene expression, to identify MYB5a—orthologous to the NEGAN transcriptional activator from *M. lewisii*—as the gene responsible for the transition to anthocyanin-pigmented petals in *M. l. variegatus*. In other monkeyflower taxa, MYB5a/NEGAN is part of a reaction-diffusion network that produces semi-repeating spotting patterns, such as the array of spots in the nectar guides of both *M. lewisii* and *M. guttatus*. Its co-option for the evolution of an apparently non-patterned trait—the solid petal lobe pigmentation of *M. l. variegatus*—illustrates how reaction-diffusion can contribute to evolutionary novelty in non-obvious ways. Transcriptome sequencing of a MYB5a RNAi line of *M. j. variegatus* reveals that this genetically simple change, which we hypothesize to be a regulatory mutation in cis to MYB5a, has cascading effects on gene expression, not only on the enzyme-encoding genes traditionally thought of as the targets of MYB5a but also on all of its known partners in the anthocyanin regulatory network.

**Keywords:** anthocyanin; color patterning; differential expression; reaction-diffusion; Turing network; feedback loops; monkeyflower; Mendelian trait; evolution of novelty

**Introduction**

Anthocyanins, the red and purple pigments that color many plant tissues, are largely responsible for the tremendous visual diversity of floral pigmentation in angiosperms. The evolutionary lability of anthocyanin pigmentation, combined with its ecological and agricultural importance, has helped anthocyanin biosynthesis and regulation become a model for investigating the genetic mechanisms for trait evolution.

In dicots, the anthocyanin biosynthetic pathway is functionally divided into the early biosynthetic genes (EBGs) and the late biosynthetic genes (LBGs) (Dubos et al. 2010). The LBGs are coordinately activated by an MBW complex consisting of three kinds of proteins: a subgroup-6 R2R3 MYB protein (M) that directly binds the promoters of the enzyme-encoding genes, and bHLH (B) and WD40 (W) co-factors that physically interact with the MYB (Martin and Gerats 1993; Lepiniec et al. 2006; Petroni and Tonelli 2011; Xu et al. 2014; Zhu et al. 2015; Lloyd et al. 2017). An R3 MYB protein represses anthocyanin biosynthesis by sequestering the bHLH proteins and preventing them from taking part in MBW formation (Zhu et al. 2009; Yuan et al. 2013; Albert et al. 2014).

A growing body of empirical work has demonstrated that the anthocyanin regulatory network is highly interconnected. For example, the MBW complex triggers increased transcription of its own bHLH component, as well as increasing the expression of the R3 MYB repressor (Xu et al. 2015). In some cases, the MBW complex further upregulates the R2R3 MYB activator gene (Yuan et al. 2014). This positive feedback loop is essential to generating spatial patterns via a self-organizing series of molecular interactions described as a reaction-diffusion system.

In the 1950s, Alan Turing proposed such a reaction-diffusion model to explain the developmental origins of semi-regularly repeating patterns in biological organisms, such as the spots on a cheetah or the stripes on a zebra (Turing 1952), and the idea has been further developed by others (Gierer and Meinhardt 1972;...
Within the monkeyflower genus *Mimulus* (synonym *Erythranthe*; Barker et al. 2012; Lowry et al. 2019), floral anthocyanin traits have evolved repeatedly (Grossenbacher and Whittall 2011) and both types of patterning appear to be involved. The developmental genetics of floral anthocyanins have been studied in the greatest detail in *M. lewisii* and *M. guttatus*. In both of these species, the R2R3 MYB activator gene Negan/MYB5 is part of a reaction-diffusion system that creates spots of red anthocyanin pigment in the nectar guide region (Ding et al. 2020). A paralagous R2R3 MYB gene, PELAN, similarly operates as part of an MBW complex in the petal lobe region of *M. lewisii*, with one important difference: PELAN lacks autocatalytic ability, and thus cannot initiate reaction-diffusion dynamics. Consistent with this finding, PELAN generates a continuous field of pigment within the positionally specified petal lobe region, rather than a repeated spotting pattern, and RNAi against PELAN causes a uniform lightening of pigmentation rather than the altered spatial pattern expected under a reaction-diffusion model (Yuan et al. 2014).

At first glance, the solid purple pigmentation of *Mimulus luteus* var. *variegatus* ([Figure 1A](#)) would appear to be a classic case of positional specification. This evolutionarily recent, Mendelian trait shows no spotting, striping, or any other kind of the repetition that is thought to be the hallmark of reaction-diffusion traits. Surprisingly, when *M. l. variegatus* is crossed to its yellow-petaled conspecific, *M. l. luteus*, the offspring of the cross feature multiple spots in their petal lobes ([Figure 1B](#)). This altered spatial pattern suggests that the gain of petal lobe anthocyanin in *M. l. variegatus* is in fact a cryptic reaction-diffusion trait. And indeed, while solid petal color across the genus *Mimulus* maps to the genomic region containing the non-autocatalytic gene PELAN ([Figure 1A](#)), petal lobe pigmentation in *M. l. variegatus* maps to the region containing the reaction-diffusion activator gene Negan ([Figure 1A](#)).

In this work, we explore the gain of an apparently unpatterned trait via a reaction-diffusion activator gene that, in other *Mimulus* species, creates spotting patterns. We find that MYB5a, the ortholog of Negan, has been co-opted for petal lobe anthocyanin pigmentation while also retaining its ancestral function of activating anthocyanin spot patterns in the nectar guide. We report the first evidence of a positive correlation in expression between the R2R3 MYB activator and its WD40 co-factor, suggesting a possible new feedback loop within the MBW transcription-factor complex. Using transcriptome data in combination with mathematical modeling, we model the interaction between the R2R3 MYB activator, MYB5a, and the R3 MYB repressor, RTO.

The results highlight how a reaction-diffusion network, and its two resulting phenotypes of nectar guide and petal lobe pigmentation, can respond in a dramatic and coordinated fashion to a relatively modest change in the expression of a single component part. The estimates of interaction strengths will help to inform and constrain future mathematical models of the evolutionary and developmental systems that give rise to the color pattern diversity of flowering plants. Finally, we note that petal lobe anthocyanin in *M. l. variegatus*, which looks to all appearances like a traditional positionally specified feature, is in fact controlled by a reaction-diffusion network. The existence of such “cryptic” reaction-diffusion traits suggests that the importance of this mechanism to evolutionary and developmental biology may be substantially greater than previously supposed.

### Materials and methods

#### Plant sources and growth conditions

Seeds were originally collected from natural populations in central Chile in 2004, and were inbred for multiple generations prior to the work described here ([Table 1](#)). Seeds were sown in 2-in pots onto wet Black Gold potting soil (SunGro Horticulture, Agawam, MA, USA) or Miracle-Gro potting soil (Scotts Miracle-Gro Co., Marysville, OH, USA), and were misted daily until germination. For transgenic experiments, seedlings were transplanted into 6" pots to promote large size. Plants were maintained in the Whitman College greenhouse under 16-hour days and were watered daily by an automatic misting system. After expansion of the first true leaves, fertilizer (Grow Big, FoxFarm Soil & Fertilizer Co., Arcata, CA, USA, N:P:K = 6:4:4; Open Sesame, FoxFarm Soil & Fertilizer Co., Arcata, CA, USA, N:P:K = 5:45:19; and/or Miracle-Gro Bloom Booster, Scotts Miracle-Gro Co., Marysville, OH, USA, N:P:K = 1:3:2) was applied two to three times weekly to promote growth and flowering.

---

**Figure 1** Petal anthocyanin pigmentation is evolutionarily labile within and between species of *Mimulus*. (A) Shifts in petal color in *Mimulus* have primarily been tracked to genomic region *pla1*. The causal gene in the *pla1* genomic region has been identified as the R2R3 MYB gene MaMYB2 in *M. aurantiacus* (Streisfeld et al. 2013), its ortholog PELAN in *M. lewisii* (Yuan et al. 2014), and one or more R2R3 MYB genes at the orthologous genomic region in *M. guttatus* (Lowry et al. 2012), *M. cupreus* (Cooley et al. 2011), and *M. naiandinus* (A. Cooley, unpublished data). Genomic region *pla2* contains MaMYB1 in *M. aurantiacus*, its ortholog Negan in *M. lewisii*, and MYB5 in *M. guttatus*, *M. cupreus*, *M. l. variegatus*, and *M. naiandinus*. The two varieties of *M. luteus* are the purple-flowered *M. l. variegatus* (left) and the yellow-flowered *M. l. luteus* (right). Photo credits: *M. aurantiacus* courtesy of Matt Streisfeld, *M. naiandinus* courtesy of Melia Matthews. Divergence times not to scale. (B) Crosses between the two varieties of *M. luteus* reveal a spotting mechanism underlying the apparently solid purple pigmentation of *M. l. variegatus*, consistent with Turing-style reaction-diffusion.
Expression analyses of candidate anthocyanin-activating genes

Two candidate anthocyanin-activating genes, MYB4 and MYB5, had been previously identified in M. l. variegatus by genetic mapping and sequence characterization (Cooley et al. 2011). Because members of the luteus complex are allotetraploids (Mukherjee and Vickery 1962; Vallejo-Marín et al. 2015; Edger et al. 2017), many genes have a homeologous copy located on a different chromosome, which could inflate expression estimates for a target gene if care is not taken to distinguish between the copies. In the process of investigating MYB5 in the sequenced M. l. luteus genome, we discovered its apparent homeolog, MYB5b (Figure 2A).

**Table 1** Seed sources

| Taxon               | Line ID   | Generations inbred | Population       | Location               |
|---------------------|-----------|--------------------|------------------|------------------------|
| M. luteus var. luteus | Ml-EY7    | 12                 | El Yeso          | 33.4°S, 70.0°W (2600 m) |
| M. luteus var. variegatus | Mlv-RC6   | 11                 | Rio Cipreses     | 34.2°S, 70.3°W (1200 m) |

Locations from which seeds were collected are given as latitude, longitude (meters above sea level).

**Figure 2** Sequence and expression of MYB5a/NEGAN. (A) Alignment of MYB5a, MYB5b, and MYB4. Amino acid sequences were inferred by translating DNA sequences, obtained from the M. l. luteus genome and transcriptome (Edger et al. 2017; Stanton et al. 2017) and from PCR and Sanger sequencing of M. l. variegatus. Black rectangles identify putative Subgroup 6 motifs. Note that this motif, which strongly predicts ability to activate anthocyanin production, is missing from M. l. variegatus exon 3. Gray boxes denote exon boundaries for MYB5a and MYB5b. Created in Geneious 9.1 by Biomatters (www.geneious.com). (B) qRT-PCR estimates of MYB5a/NEGAN and MYB5b expression in M. l. variegatus and M. l. luteus. Each point indicates the mean of three technical replicates, ±95% CI, relative to the reference gene Actin and a reference pool of cDNA.
MYB5 will therefore be referred to in this work as MYB5a, to distinguish it from its homeolog. In the M. l. luteus reference genome, the three exons that make up MYB5b are identical in their coding sequence to the first three exons of MYB5a. However, substantial divergence in the 5'-UTR (as well as downstream of the third exon) enabled the development of copy-specific primers, allowing us to disentangle the expression of MYB5a from its homeolog. We were not able to identify any homeolog of MYB4 in the M. l. luteus genome.

The gene responsible for activating anthocyanin biosynthesis in M. l. variegatus petal lobes is expected to be expressed in developing floral buds, during or just prior to the accumulation of visible anthocyanin pigment. Developmentally, this occurs approximately 72 hours before anthesis (Cooley et al. 2011), before the bud has emerged from the calyx. Endpoint and quantitative RT-PCR were used to investigate patterns of expression for each candidate gene, as well as for MYB5b, in developing floral buds and also young leaves of the purple-lobed M. l. variegatus compared to the yellow-lobed M. l. luteus. Primers were designed based on the sequenced genome of M. l. luteus (Supplementary Table S1), and their functionality in M. l. variegatus was confirmed by PCR prior to their use in expression analyses.

Although M. l. variegatus and M. l. luteus have striking differences in petal lobe pigmentation, with abundant anthocyanin production in the former and none in the latter, they share the trait of anthocyanin spotting in the nectar guide region of the corolla. Petals of developing flower buds were therefore dissected into lobe vs nectar guide components and the two tissue types were analyzed separately.

RNA was extracted using the Agilent Plant RNA Isolation Kit (Santa Clara, CA, USA). cDNA was synthesized using the ProtoScript First Strand cDNA Synthesis Kit from New England Biolabs, Inc. (Ipswich, MA, USA). Endpoint (qualitative) RT-PCR was performed on cDNA, with gDNA template as a control for primer efficacy. In addition to the MYB5a primers, either Actin or GAPDH primers were included as a control for successful cDNA synthesis (Supplementary Table S1). PCRs were performed with 0.2 mM dNTPs, 0.2 μM of each primer, 0.2 μL of G-Biosciences Taq DNA Polymerase per 25-μL reaction, and 1x G-Biosciences buffer. Amplification was performed with temperature settings of 95°C for 3 min, followed by 30 cycles of (95°C for 30 s, variable temperature for 30 s, 72°C for 1 min per kb of product), and then 72°C for 10 min. The annealing temperature was set to 0°C lower than the lowest melting temperature of the set of primers used in the PCR, and the minimum extension time used was 30 s.

To test for expression patterns consistent with the activation of petal lobe anthocyanin synthesis, quantitative RT-PCR was performed on MYB5a and MYB5b using SYBR-Green Brilliant III Ultra-Fast reagents (Agilent Technologies, USA), in optical 96-well plates (Greiner Bio-One, Belgium), on an MxPro3000p analyzer (Stratagene, USA). Four biological replicates were used for each taxon and tissue type, collected from different plants of the same inbred line. Each biological replicate was run in triplicate and the average of the three technical replicates was used for statistical analyses. Samples were amplified for 40 cycles of 95°C for 10 s and 51°C for 20 s. Melt curves were obtained by heating from 51°C to 95°C with a ramp speed of 0.01°C per second. The Actin gene was used as a normalizer. Raw qPCR fluorescence data were collected and analyzed by the default settings of the MxPro software v.4.10 (Agilent Technologies, USA). Amplification efficiencies for each primer pair were determined using the Ct (threshold) values obtained from a 1/4 dilution series (1:4, 1:16, 1:64, 1:256, and 1:1024).

Cloning and sequencing MYB5a splice variants

Two splice variants of M. l. variegatus MYB5a, containing exons 1-2-3 and 1-2-4 respectively, were PCR-amplified using primers MYB5a-10F and MYB5a-53R (Supplementary Table S1). PCR products were purified using a Genomic DNA Clean and Concentrate Kit (Zymo Research, Irvine, CA, USA), then cloned using the pGEM T-Easy kit and manufacturer’s protocol (Promega, Madison, WI, USA). pGEM vector containing the desired PCR product was transformed into competent JM109 E. coli cells (Promega, Madison, WI, USA). Colonies were evaluated using blue-white screening and PCR with M13 universal primers. Inserts of the expected length were sent to Eton Bioscience (San Diego, CA, USA) for Sanger sequencing.

Transgene construction

A previously built overexpression construct (Yuan et al. 2014), containing the complete NEGAN coding sequence from M. lewisii, driven by a 35S promoter, was used to test the sufficiency of MYB5a for activating petal lobe anthocyanin biosynthesis in M. l. luteus, a variety that normally lacks petal lobe anthocyanin pigmentation (Figure 1A).

Two RNAi constructs were built, targeting M. l. variegatus MYB5a-specific sequences in the 5'-UTR and fourth exon respectively, using the Gateway method (Earley et al. 2006; Katzen 2007). In brief, each fragment of interest was PCR-amplified using a forward primer with a 5'-CACC tag. The tag permitted the fragment to be directionally cloned into a pENTR entry vector (Thermo Fisher Scientific, Waltham, MA, USA); attL/attR recombination was used to transfer the correctly oriented fragment into the Gateway-compatible destination vector pB7GWIWG2(I) (VIB, Ghent, Belgium). M. l. variegatus genomic DNA for PCR was extracted using the E.Z.N.A. SP Plant DNA Kit (Omega Bio-Tek, Norcross, GA, USA). PCR amplification was performed using Phusion High Fidelity DNA Polymerase (New England Biolabs). The inserts and primers used for RNAi transgene construction are summarized in Table 2.

Stable transformation

Agrobacterium-mediated plant transformation was performed using floral spray and vacuum infiltration as described in Yuan et al. 2014. Two transgenic lines were obtained, one of which was chosen for further studies.

Table 2 Construction of MYB5a/NEGAN RNAi transgenes for use in M. l. variegatus

| Insert                  | Insert length (bp) | Primers                                                                 |
|-------------------------|--------------------|-------------------------------------------------------------------------|
| M. l. variegatus 5'-UTR fragment | 307                | 24 F: caccCAAAAATTTTGGTTTTTGCCATTT 37 R: TCGGTCAATTTAAATGCACA          |
| M. l. variegatus exon 4 fragment   | 377                | 60 F: caccGAGGAAACGGCCAAAGCAC 56 R: GCAGCTTCCCCGTTTCA                  |

Both inserts were cloned into the pB7GWIWG2(I) RNAi vector. Primers are labeled with a number and F for Forward or R for Reverse. Their sequences are shown 5’ to 3’. Nucleotides added to primers for purposes of directional cloning or restriction enzyme digests are underlined. bp, base pairs. Lowercase cacc- represents nucleotides added to primers to enable cloning into the pENTR entry vector.
et al. (2013). Each experiment utilized approximately 12 robustly budding 2- to 6-month-old plants growing in 6” pots. The commercial *Agrobacterium tumefaciens* strain LBA4404 (ThermoFisher Scientific) produced only two transformed seedlings from the infiltration of 67 plants (K. Om, senior thesis 2017). Subsequent experiments were performed using strain GV3101, and this yielded higher rates of transformed seedlings.

The concentration of Silwet-L77 (Lehle Seeds, Round Rock, TX, USA) used by Yuan et al. (2013) was 0.1%, or 1 mL per L of *Agrobacterium* culture. This unusually high concentration of Silwet was previously found to increase transformation success in *Mimulus* (Y.-W. Yuan, unpublished data). Acetosyringone was added to a final concentration of 0.1 M as in Yuan et al. (2013).

The predominantly outcrossing taxa *M. luteus* and *M. l. variegatus* were manually self-pollinated for 2 weeks after the first post-infiltration flower opened. Seeds were collected and densely sown in 25 cm × 50 cm flats. As soon as germination was observed, flats were sprayed daily with 1:1000 Finale (Farnam Companies, Inc., Phoenix, AZ, USA) to eliminate non-transgenic plants. Transgenic seedlings were grown to flowering and photographed. cDNA was extracted and the presence of the transgene was verified by PCR, using a forward primer targeted to the transgene and a reverse primer targeted to the atrR2 region of the transgenic plasmid (Table 2 and Supplementary Table S1).

RNA extractions, cDNA synthesis, and endpoint RT-PCR were performed as described in the Expression Analyses section, to evaluate the expression of the target gene relative to an untransformed control plant. Two different primer pairs targeting MYBSa were used, 33 F-40R and 44 F-45R, along with primers to reference gene GAPDH (Stanton et al. 2017) as a positive control (Supplementary Table S1).

**RNA extraction and library preparation for RNAseq**

Transcriptome analysis was used to confirm the specific knockdown of target gene MYBSa by RNAi, and to explore the effects of this knockdown on other anthocyanin-related genes. Petal tissue for transcriptome sequencing was collected separately from three individuals of our highly inbred wild-type line of *M. l. variegatus*, RC6, and from three white-flowered offspring (Vrna1.1, 1.3, and 1.5, Supplementary Figure S2) of the white-flowered RNAi line Vrna1, and *M. naidinus* as an outgroup. As expected, the samples clustered by treatment with the outgroup being an outlier (Supplementary Figure S2).

After performing a shrinkage estimation for dispersion to address the inaccuracy introduced by the small sample size and reduce the false positive rate (Love et al. 2014), the logarithmic fold change between RNAi treatment and control samples was used to evaluate differential expression. Following a false discovery rate control using the Benjamini-Hochberg Correction method, transcripts that were log-2-fold up- or down-regulated with a F-value < 0.05 were considered to be significantly differentially expressed. Transcript expression profiles were normalized to reads per kilobase per million (RPKM) for further analysis and for plotting (Supplementary Table S3).

To annotate differentially expressed genes, all gene sequences from the *M. l. luteus* reference genome were translated to protein sequences using EMBoss (6.5.7) “transeq” command and searched against Arabidopsis thaliana (TAIR) protein database (www.arabidopsis.org) using the “BLASTp” query with an e-value cut-off < 10^-6. For each coding sequence from *M. l. luteus*, the best-hit *A. thaliana* gene was used to annotate the transcript with a gene name and Gene Ontology (GO) annotation terms.

These annotations were then applied to the *M. l. variegatus* transcriptome data. GO enrichment analysis was conducted, for genes that were differentially expressed between wild-type *M. l. variegatus* and the MYBSa RNAi line Vrna1, using topGO (Version 2.36.0) with the “org.At.tair.db” database in R. Pathway enrichment tests were done using the KEGG (the Kyoto Encyclopedia of Genes and Genomes, https://www.genome.jp/kegg/; Release 93.0) pathway assignments for Arabidopsis with the KEGGREST Bioconductor package (Kanehisa et al., 2016).

**Tree construction**

A gene tree was constructed to determine the relationship of *M. l. luteus* 12200, the candidate MYBSa gene, to previously studied unaware mapping approach to assemble the *M. l. variegatus* transcriptomes to the *M. l. luteus* genome (Stein 2011; Williams et al. 2014). Sequencing quality control was performed by plotting the sequence nucleotide distribution and sequencing quality scores for all samples. Transcriptome libraries were aligned using Bowtie2 (Version 2.3.5.1) under -very-sensitive-local mode (Langmead and Salzberg 2012) for best results in distinguishing homolog expression, given that the *M. luteus*-group species are putative tetraploids (Mukherjee and Vickery 1962; Vickery 1995). Overall alignment rates were greater than 90% for all 7 samples. From the sequence alignment maps we counted reads per gene for all samples using the exon coordinates included in the published luteus GFF (general feature format) file. Read count was performed with software HT-seq (Version 0.11.2) (Anders et al. 2014) in Python. For detailed alignment documents, see the Supplementary Data. Pipeline code, alignment documents, and results are accessible through the GitHub repository (https://github.com/cici-xingyu-zheng/Luteus-RNA-seq) and for raw read count, see Supplementary Table S2.

**Transcriptome analysis and functional annotation**

Analyses of differential gene expression were conducted in DESeq2 (Version 1.26.0) in the R/Bioconductor environment (Love et al. 2014) (R Version 3.6.2; Bioconductor Release 3.10). After normalizing each gene by sequencing depth, we performed a principal component analysis plot of the seven transcriptome samples (three wild-type *M. l. variegatus*, three *M. l. variegatus* from RNAi line Vrna1, and *M. naidinus* as an outgroup). As expected, the samples clustered by treatment with the outgroup being an outlier (Supplementary Figure S2).

Because the expression of anthocyanin-producing genes in Mimulus flowers is highest early in bud development, just before the flower emerges from the calyx. Anthers were removed and the remaining petal tissue (including both lobe and nectar guide regions) was snap frozen in liquid nitrogen. RNA was extracted from each of the six samples using the Agilent Plant RNA Isolation Kit (Wilmington, DE, USA). A stranded RNA-Seq plus Ribo-Zero library preparation, followed by one lane of Illumina HiSeq 4000 50-bp single-read sequencing, was performed by the Duke University sequencing core facility (Durham, NC, USA). The FASTQ format RNA libraries were deposited in the NCBI Sequence Read Archive (SRA) repository under accession number PRJNA629107.

**Transcriptome alignment**

With a published *M. l. luteus* genome and gene feature annotation file available (Edger et al. 2017), we chose the genome splice-
Mimulus MYBs. Sequences used to construct the gene tree in Figure 2 of Yuan et al. (2014) were downloaded from GenBank. A full list of GenBank sequence IDs are available in the Supplementary Table S4. Amino acid sequences were aligned with Clustal Omega and an ML gene tree was constructed using PhyML using a WAG substitution model and 100 bootstrap replicates. After initial tree construction, a subset of sequences that formed the core clade of MYBs of interest was realigned using Clustal Omega. The alignment cleaned manually to remove poorly aligned regions, and a tree was constructed using PhyML. The full gene tree from this analysis is available in the Supplementary Figure S3.

**Regulatory and pathway analysis**

In order to test for an effect of MYB5a down-regulation on the anthocyanin biosynthetic pathway, we examined the expression of genes corresponding to six core pathway enzymes that produce the cyanidin pigment found in M. l. variegatus (Cooley et al. 2008). These are: chalcone synthase (CHS), chalcone isomerase (CHI), flavonoid-3-hydroxylase (F3H), dihydroflavonol-4-reductase (DFR), anthocyanidin synthase (ANS), and UDP-flavonoid-3-glucosyl-transferase (UFGT) (Holton and Cornish 1995; Tanaka et al. 2008). A list of M. l. variegatus transcripts with RPKM >1 and annotated with descriptions matching these enzymatic activities was obtained. Genes were considered truly orthogonal if they were the best hit in a reciprocal BLAST or had the same high BLAST score as a putative homeolog.

In addition to the enzyme-encoding genes, we were also interested in what other regulatory factors might be affected when we knock down the MYB5a transcriptional activator. In all species that have been examined, the regulatory complexes of the anthocyanin pathway include members of the MYB, basic helix-loop-helix (bHLH), and WD40 repeat families (Dooner et al. 1991; Koes et al. 2005). Therefore, we subset from the differentially expressed genes a list of genes annotated as members of these three transcriptional regulator families based on comparison to A. thaliana. Within this subset, we used reciprocal best 267 BLAST hits to identify orthologs to the genes that reported to be involved in 268 anthocyanin regulation in M. guttatus and M. lewisii.

**Data availability**

Seeds, bacterial strains, and plasmids are available upon request. Individual MYB4, MYB5a, and MYB5b coding sequences are located in GenBank. Whole-genome sequence for M. l. luteus was released by Edgar et al. (2017) along with the draft genome and are located at https://datadryad.org/stash/dataset/doi:10.5061/dry-ad.d4vr0. Raw transcriptome data for M. l. luteus are published in Stanton et al. (2017) and are located at http://dx.doi.org/10.5061/dryad.84655. The M. l. variegatus transcriptome raw reads are available on the Sequence Read Archive (https://www.ncbi.nlm.nih.gov/bioproject/PRJNA629107/). Transcriptome analysis code and alignment documents are accessible through the GitHub repository (https://github.com/cici-xingyu-zheng/Luteus-RNA-seq).

Supplementary material is available at figshare DOI: https://doi.org/10.25386/genetics.13350800.

**Results**

Expression of an alternately spliced R2R3 MYB, MYB5a/NEGAN, covaries with petal anthocyanin

In M. l. variegatus, only two anthocyanin-related genes were found within the genetically mapped petal anthocyanin locus pla2 (Cooley et al. 2011). The two genes, MYB4 and MYB5a, show phylogenetic similarity to the anthocyanin-activating Subgroup 6 of the R2R3 MYB gene family (Stracke et al. 2001; Cooley et al. 2011). Reciprocal BLAST searches of the M. lewisii genome indicate that MYB5a is orthologous to NEGAN, a gene shown to be responsible for nectar guide anthocyanin spots in M. lewisii (Yuan et al. 2014). NEGAN has previously been shown to be orthologous to MYB5a from M. guttatus (Yuan et al. 2014).

MYB4, while somewhat similar in sequence to MYB5a/NEGAN, has a nonconservative amino acid substitution in the functionally critical DNA-binding motif that is a hallmark of Subgroup 6 transcription factors (Figure 2A) and has a top BLAST hit on NCBI to M. guttatus LOC105953416, which encodes a GL1-like trichome differentiation protein. Additionally, unsuccessful efforts to amplify MYB4 out of M. l. variegatus floral bud cDNA, using primers that successfully amplify the gene from gDNA, indicate that this gene is not detectably expressed in developing M. l. variegatus buds (Supplementary Figure S1). Consistent with its similarity to a trichome differentiation gene, we did detect expression of MYB4 in leaf tissue (Supplementary Figure S1).

In contrast, quantitative RT-PCR revealed that MYB5a/NEGAN is strongly and specifically expressed in the anthocyanin-pigmented petal lobes of M. l. variegatus, while showing little to no expression in the non-anthocyanin-pigmented petal lobes of M. l. luteus (Figure 2B). Expression of MYB5a/NEGAN was modest in the anthocyanin-pigmented nectar guide tissue of both taxa, and undetectable in their leaf tissue (Figure 2B). The homeologous gene copy, MYB5b, showed consistently low expression across both taxa and all tissue types, at levels less than 10% of that observed in M. l. variegatus petal lobes (Figure 2B).

The anthocyanin-activating subgroup 6 R2R3 MYBs generally have three exons. The diagnostic Subgroup 6 amino acid motif is KRPR[S/T]F in Arabidopsis, and it is located in the third exon (Stracke et al. 2001). In M. l. luteus and M. l. variegatus, however, MYB5a genomic sequence data revealed the presence of a fourth exon that is extremely similar in sequence to the third exon. In both taxa, this fourth exon appears to be intact and potentially functional, including the Subgroup 6 motif. The fourth exon differs between M. l. luteus and M. l. variegatus by 7 amino acid substitutions and one in-frame indel (Figure 2A). The third exon, although apparently intact in M. l. luteus, has a 26-bp frameshift-inducing deletion in M. l. variegatus that eliminates the critical Subgroup 6 domain, alters 35 amino acids, and creates a premature stop codon (Figure 2A).

MYB5a/NEGAN is sufficient and necessary for activating petal anthocyanin in M. luteus

A pilot experiment showed that NEGAN from M. lewisii, when overexpressed with the 3SS promoter, was sufficient to activate anthocyanin production in the petal lobe, as well as the nectar...
guide region and leaf tissue, of the normally yellow-petaled M. l. luteus. Encouraged by this result, we used a 35S-driven RNAi transgene to test the hypothesis that the exon 1-2-4 transcript of MYB5a is necessary for petal lobe anthocyanin pigmentation in the purple-petaled M. l. variegatus. In the strongest RNAi line, “Vrnai1,” anthocyanin pigmentation was completely abolished in both the petal lobe and the nectar guide (Figure 3B). Anthocyanin pigmentation in the stems, the leaves, and the adaxial side of the corolla appeared to be unaffected, indicating that the effect of MYB5a knockdown is spatially specific. Successful knockdown of MYB5a in line Vrnai1 was demonstrated via qualitative RT-PCR (Figure 3C), and confirmed through transcriptome analysis (next section).

Considerable variation in the severity of the RNAi phenotype was observed across different transgenic lines, as is typical for this technique (Figure 4). From 12 wild-type M. l. variegatus transformed with RNAi targeting exon 4 of MYB5a, six putatively transgenic seedlings were obtained. Four of these (Vrnai1, Vrnai3, Vrnai5, Vrnai7) had a visible reduction in petal lobe anthocyanin pigmentation. From an additional 12 wild-type M. l. variegatus transformed with RNAi targeting the 5’-UTR of MYB5a, two putatively transgenic seedlings were obtained. One of these (Vrnai6) had a visible reduction in petal lobe anthocyanin pigmentation. Genotyping with a MYB5a forward primer and the attR2 transgene-specific reverse primer confirmed the presence of the RNAi transgene in all eight lines. In line Vrnai6, the RNAi

---

**Figure 3** MYB5a/NEGAN is sufficient and necessary for activation of petal lobe anthocyanins. (A) Overexpression of the coding sequence of MYB5a/NEGAN from M. lewisii in the normally yellow-petaled M. l. luteus activates anthocyanin biosynthesis in both leaf and petal tissue. (B) RNAi targeting MYB5a exon 4 in the normally purple-petaled M. l. variegatus eliminates anthocyanin biosynthesis in the petal lobes and nectar guides, but not elsewhere in the plant. The strongest of 8 RNAi lines, Vrnai1, is shown here. See Supplementary Data for images of the other RNAi lines. (C) Qualitative (end-point) RT-PCR on cDNA from developing petal lobes of wild type (W) and transgenic (T) M. l. variegatus reveals a reduction in MYB5a expression in the Vrnai1 transgenic line. Reference gene GAPDH was used as a positive control.
phenotype declined over the lifetime of the plant (Figure 4). This phenomenon has been shown to be caused by methylation and epigenetic silencing, particularly of the 35S promoter, in both *A. thaliana* and the tobacco species *Nicotiana attenuata* (Daxinger et al. 2008; Mlotshwa et al. 2010; Weinhold et al. 2013).

The Vrnai1 plant was self-fertilized, and eight of the resulting seeds were planted and grown to flowering. As expected for a single-copy (heterozygous) insertion of a dominantly acting transgene in the Vrnai1 plant, we observed a 3:1 ratio of six white-flowered and two wild-type plants among the offspring (Supplementary Figure S2). The six white-flowered offspring exhibited slight anthocyanin mottling on some flowers (Supplementary Figure S2).

Transcriptomic differences between wild type and MYB5a RNAi lines of *M. l. variegatus*

We found a total of 632 genes that were significantly differentially expressed between wild-type and Vrnai1 lines of *M. l. variegatus*, with 346 genes down-regulated and 290 up-regulated in the Vrnai1 line (Supplementary Table S5). The differentially expressed genes are enriched in a variety of functions including response to UV-B, anthocyanin-containing compound biosynthesis, pollen exine formation, and phenylpropanoid biosynthetic process (P-value <0.001) (Supplementary Table S6). A pathway enrichment analysis identified two significantly enriched pathways (Supplementary Table S7): 1. cutin, suberine and wax biosynthesis, and 2. flavonoid biosynthesis, which includes the anthocyanin biosynthetic pathway (Holton and Cornish 1995).

The *M. l. variegatus* MYB5a DNA sequence, which had previously been determined by PCR and Sanger sequencing, had best hits to three coding sequences in the *M. l. luteus* reference genome: Mlu 12200, Mlu 12207, and Mlu 42095. In the *M. l. variegatus* transcriptomes, the latter two transcripts were not detectably expressed in developing petal tissue, in either wild-type or RNAi lines of *M. l. variegatus*, eliminating them from further consideration as the petal anthocyanin activator. In contrast, Mlu 12200 showed expression patterns consistent with our expectation for MYB5a: it was robustly expressed in all six libraries, with three-fold higher expression in the wild-type compared to Vrnai1 (Figure 5A). Both the Mlu 12200 and the Mlu 42095 transcripts from *M. l. variegatus* grouped phylogenetically with *M. guttatus* MYB5 and *M. lewisii* NEGAN (Figure 6).

To check for off-target effects of the RNAi transgene, we asked whether any other Subgroup 6 R2R3 MYB genes were significantly down-regulated in the RNAi line. Five MYBs were identified among the down-regulated genes (Mlu 24690,
Mlu 05348, Mlu 27563, Mlu 17841, and Mlu 00921), but none of them contained a Subgroup 6 motif. We conclude that the loss-of-pigment phenotype observed in line Vrnai1 is due to the reduction in MYB5a expression alone, and was not caused by incidental down-regulation of another anthocyanin-activating MYB gene.

RNAi knockdown of MYB5a reduces transcription of the late anthocyanin biosynthesis genes

We identified a total of 30 genes that are annotated to have enzymatic functions corresponding to the six core enzymes in the cyanidin pathway (Holton and Cornish 1995) (Supplementary Table S8). The EBGs (CHS, CHI, and F3H) are particularly enriched in copy number, each having more than the two homeologous copies expected in a tetraploid.

In eudicots, the anthocyanin biosynthetic genes are usually divided into two groups, the early (EBGs) and late (LBGs) biosynthetic genes, with the latter being tightly regulated by an MBW complex (Martin and Gerats 1993; Lepiniec et al. 2006; Dubos et al. 2010; Petroni and Tonelli 2011; Xu et al. 2014; Lloyd et al. 2017). In Arabidopsis, CHS, CHI, and F3H belong to the EBGs, producing the precursors of not only flavonoid pigments but other flavonol compounds. DFR, ANS, and UF3GT comprise the LBGs. Conserved as the enzymatic pathway is, though, the break point of regulation between early and late genes can vary across species (Streisfeld and Rausher 2009).

We utilized our transcriptomic data to determine where the breakpoint occurs in M. l. variegatus. Consistent with expectations from the literature, the anthocyanin biosynthetic genes in M. l. variegatus show a...
RNAi knockdown of MYB5a reduces transcription of other anthocyanin regulators

Within the list of differentially expressed genes, we identified 20 genes that are annotated to be transcription factors with MYB or helix-loop-helix (HLH) domains, or that encode a WD40 protein that could potentially belong to a MYB–bHLH–WD40 regulatory complex (Supplementary Table S9). Since different MBW genes regulate a variety of traits besides anthocyanin synthesis and gene functions might have diverged between *Mimulus* and *Arabidopsis*, we further compared this subset of genes to the MBW regulatory genes within *M. lewisii*, the *Mimulus* species with the most thoroughly characterized anthocyanin regulatory network. One WD40 gene (labeled MlutWD40a) was the best hit to *M. lewisii* M1W40a, a gene that co-activates anthocyanin expression in the corolla. Two bHLH genes (MlutbHLH1 and MlutbHLH2) have the highest hit to the same bHLH gene in *M. lewisii*, ANbHLH3, indicating they are duplicates of each other. Interestingly, in *M. lewisii*, ANbHLH3 is not detectably expressed in the petal lobes, suggesting a functional diversification in the bHLH co-factors as well as in MYB5a/NEGAN (Yuan et al. 2014).

Recent studies show that single-repeat R3 MYBs are activated by the R2R3 MYB member of the MBW complex, and inhibit anthocyanin biosynthesis by directly interacting with the bHLH component of the same complex. In two *Mimulus* species, *M. lewisii* and *M. guttatus*, the same R3 MYB (RTO) is reported to down-regulate floral anthocyanin production. We found that two R3 MYB genes, Mlu 13044 (MlutR3MYB1) and Mlu 33990 (MlutR3MYB2), both have the highest BLAST hit to this RTO gene and are strongly and significantly down-regulated in the MYB5a RNAi line, Vrnai1. Their down-regulation in Vrnai1 is consistent with the model that the R2R3 MYB, MYB5a, is an activator of its own inhibitor, although the impact of MYB5a on two apparently homeologous gene copies is unique. The activating properties of the anthocyanin-activating R2R3 MYB on itself, its R3 MYB inhibitor, and its bHLH cofactor, have been previously shown in other taxa (Yuan et al. 2014; Xu et al. 2015). We hypothesized that in *M. l. variegatus* RNAi lines, the bHLH and RTO candidate anthocyanin regulators that we identified would show the same directional expression change as MYB5a (Li 2014). In fact, our expression results showed that down-regulating MYB5a in the RNAi line of *M. l. variegatus* led to a down-regulation in all the putative regulatory genes (Supplementary Table S10), surprisingly including the WD40 genes (Figure 5, C and D). The involvement of the WD40 genes in a positive feedback loop has not, to our knowledge, been reported previously. The activators of anthocyanin biosynthesis—including R2R3 MYB, bHLH, and WD40 transcription factors—were all two- to threefold down-regulated in the RNAi line. The two R3 MYB inhibitors showed much more dramatic effects: they were 19- and 29-fold down-regulated (Figure 5, C and D).

**Discussion**

Here we demonstrate that MYB5a is both sufficient and necessary for the evolutionarily recent, single-locus gain of petal lobe anthocyanin within *M. luteus*. We identify an unusual four-exon structure to this Subgroup 6 R2R3 MYB, along with evidence for alternative splicing. While the role of coding vs cis-regulatory evolution (Hoekstra and Coyne 2007; Stern and Orgogozo 2008) was not explicitly examined in this paper, we found that patterns of expression of MYB5a corresponded to pigmentation: the gene was strongly expressed in the heavily anthocyanin-pigmented petal...
lobes of *M. l. variegatus*; not detectably expressed in the petal lobes of *M. l. luteus*, a conspecific which lacks petal lobe anthocyanin; and modestly expressed in the partly pigmented nectar guides of both taxa. Further supporting the importance of expression change rather than coding change is that the orthologous coding sequence from the evolutionarily distant *M. lewisii*, which is approximately 20 MY divergent from *M. variegatus* (Nie et al. 2006), was highly effective at activating anthocyanin in the normally yellow-flowered *M. l. luteus*. We therefore predict that the causal mutation is either cis to *MYB5a*, or located in a trans-regulator of *MYB5a* that is closely linked to *MYB5a* within the mapped *pla2* genomic region. Additional transgenic experiments, such as coding or promoter swaps, will help to clarify the extent and nature of *MYB5a* functional divergence in *M. l. variegatus*.

The adaptive significance of this evolutionary change is not yet fully understood. Flower color differences in *Mimulus* can have dramatic effects on pollinator behavior (Schemske and Bradshaw 1999). However, in wild populations in Chile, both *M. l. luteus* and *M. l. variegatus* are pollinated by the native Chilean bumblebee *Bombus dahlbomii*, which did not exhibit either collective or individual flower color preferences in a mixed-phenotype swarm of *Mimulus* hybrids (Cooley et al. 2008). Anthocyanins also protect plants against a variety of environmental stressors including extremes of heat, cold, drought, and UV radiation (Chaikiner-Scott 1999). The closely related *M. cupreus*, which recently gained petal lobe anthocyanins independently of *M. l. variegatus* (Figure 1A), does show greater drought tolerance than the yellow-flowered *M. l. luteus* (Stanton et al. 2016), but this work has not yet been extended to *M. l. variegatus*.

Our RNAi results indicate that *MYB5a* is responsible for anthocyanin pigmentation in two distinct regions of the *M. l. variegatus* corolla: the nectar guide and the petal lobe. The role of *MYB5a/NEGAN* in nectar guide spotting appears to be conserved across much of the genus. Its deployment to the petal lobes is unique in diverse species including *M. lewisii*, *M. aurantiacus*, *M. cupreus*, and *M. naiandinus* (Cooley and Willis 2009; Cooley et al. 2011; Streisfeld et al. 2013; Yuan et al. 2014), petal lobe anthocyanin pigmentation is controlled by ancient paralogs of *MYB5a/NEGAN*, located at the functionally similar but genetically unlinked genomic region *pla1* (Figure 1).

The R2R3 MYB gene PELAN, which is responsible for petal lobe pigmentation in *M. lewisii*, differs fundamentally from *MYB5a/NEGAN* in that it does not appear to function as part of a reaction-diffusion network (Yuan et al. 2014). Several observations indicate that *MYB5a* in *M. l. variegatus* is operating via reaction-diffusion, despite the apparently unpatterned nature of its petal lobe anthocyanin phenotype. (1) Crossing *M. l. variegatus* to the conspecific variety *M. l. luteus* reveals a pattern of anthocyanin spots in the petal lobes of the F1 hybrids, rather than a spatially uniform reduction in pigment intensity. (2) Knocking down *MYB5a* in *M. l. variegatus* eliminates the nectar guide spots, indicating that *MYB5a* has retained this ancestral ability which relies on reaction-diffusion (Ding et al. 2020). Our transcriptome data show that no other Subgroup 6 R2R3 MYB was significantly down-regulated in the *MYB5a* RNAi line, confirming that the loss of nectar guide spots is unlikely to be caused by an off-target effect of the RNAi transgene silencing a paralogous MYB gene. (3) The RNAi lines with weaker phenotypes did not show a spatially uniform reduction in pigment intensity, as has been observed for RNAi against the non-autocatalytic R2R3 MYB PELAN (Yuan et al. 2014). Rather, pigment was completely lost in portions of the petal, and the remaining pigment was sometimes expressed in a spotted pattern (i.e., line Vrnai6; Figure 4).

The genomic region harboring PELAN appears to have been used for petal pigmentation across *Mimulus* more often than the genomic region harboring *MYB5a/NEGAN* (Figure 1). More work is required to determine whether the inability of PELAN to self-activate is a recently derived trait within *M. lewisii*, or a conserved and evolutionarily ancient feature. It is interesting to note that the orthologous (*pla1*) genomic region in *M. guttatus* has been found to create semi-stochastically distributed red dots of anthocyanin pigment on the calyx (Lowry et al. 2012), suggesting that the causal gene in this instance is capable of reaction-diffusion-type patterning. While circumstantial, this discovery suggests that the lack of self-activation in PELAN in *M. lewisii* may be the exception rather than the rule for anthocyanin regulatory networks in *Mimulus*.

The repeated gains and losses of anthocyanin pigmentation across *Mimulus* (Grossenbacher and Whittall 2011), combined with the genomic and molecular resources that characterize the genus (Sobel and Streisfeld 2013; Twyford et al. 2015; Yuan 2019), create fertile ground for investigating how two major developmental approaches to trait formation—reaction-diffusion and positional specification—are utilized in the evolution of biological diversity. One challenge in studying anthocyanin pigmentation by either mechanism is that the number of interactions in the regulatory network makes it difficult to precisely predict the effects of perturbing any one component. Yet, the ability to make such predictions is central to generating and testing mechanistic hypotheses about how the system functions.

Mathematical modeling provides a potentially powerful solution to this problem. Yet, empirical estimates for many of the necessary parameters—such as the strength of the interactions among the different regulatory proteins—are lacking. Discovering the values of such parameters is essential in order to constrain a wide universe of possible models to a narrower range of biologically plausible ones.

One striking result from our transcriptome data was that an RNAi knockdown of *MYB5a/NEGAN* had a much stronger effect on the RTo repressor than on *MYB5a* itself (Figure 7). To further explore this result, we used mathematical modeling to identify conditions that could give rise to the observed patterns of gene expression change between wild-type and Vrnai1 lines of *M. l. variegatus*. We found that the experimental observation that a knock-down of *MYB5a/NEGAN* decreases the RTo concentration is consistent with activator-inhibitor systems that serve as minimal models of the MBW regulatory network (see Supplementary File S1). Working with such a minimal model, we found it straightforward to identify parameter values that demonstrate how a knock-down of *MYB5a/NEGAN* can result in a relatively larger fold change of the inhibitor concentration. Additionally, by changing parameters, we are able to demonstrate a wide variety of theoretical behaviors consistent with previous experimental results (Ding et al. 2020). According to this model, a knock-down of activator will always cause a decrease in inhibitor concentration; the relative size of the inhibitor change will vary depending on the parameter values.

Overall, our transcriptome analyses and models highlight the inter-relatedness of expression patterns across the network of anthocyanin regulatory and biosynthetic genes (Figure 7). These results illustrate how the network can respond dynamically to expression changes of a single network component, creating an avenue for a relatively modest expression change—such as that caused by a single cis-regulatory mutation—to have a major effect on the transcriptome as well as on the ultimate phenotype.
Acknowledgments
The authors gratefully acknowledge assistance with molecular work from Baoqing Ding, Lauren Stanley, Rachel Eguia, Johanna Au, Jonah Rodewald, and Jeanette Schwensen. We thank Larry North for construction of a vacuum infiltration apparatus, Carter Cooke and Bella Rivera for greenhouse assistance, and Amy LaFountain and Qiaoshan Lin for assistance in obtaining transgenic seeds.

Funding
This material is based upon work supported by the Murdock College Research Program for Life Sciences Award 2015270, National Science Foundation microMORPH training grant, and National Science Foundation - Division of Environmental Biology award 1655311 to A.M.C.; and National Science Foundation - Division of Mathematical Sciences award 1951646 and National Institutes of Health - National Center for Complementary and Integrative Health award R01AT01816 to G.D.C.S.

Conflicts of interest
None declared.

Literature cited
Albert NW, Davies KM, Lewis DH, Zhang H, Montefiori M, et al. 2014. A conserved network of transcriptional activators and repressors regulates anthocyanin pigmentation in eudicots. Plant Cell. 26:962–980.
Anders S, Theodor PT, Huber WH. 2015. HTSeq—a Python framework to work with high-throughput sequencing data. Bioinformatics 31:166–169.
Ball P. 2015. Forging patterns and making waves from biology to geology: a commentary on Turing (1952) ‘the chemical basis of morphogenesis’. Philos Trans R Soc B. 370:20140218.
Barker W, Nesom G, Beardsley PM, Fraga NS. 2012. A taxonomic conspectus of Phrymaceae: a narrowed circumscription for Mimulus, new and resurrected genera, and new names and combinations. Phytoneuron 39:1–60.
Chalker-Scott L. 1999. Environmental significance of anthocyanins in plant stress responses. Photochem Photobiol. 70:1–9.
Cooley A, Carvallo G, Willis J. 2008. Is floral diversification associated with pollinator divergence? Flower shape, flower colour and pollinator preference in Chilean Mimulus. Ann Bot. 101:641–650.
Cooley AM, Modliszewski JL, Rommel ML, Willis JH. 2011. Gene duplication in Mimulus underlies parallel floral evolution via independent trans-regulatory changes. Curr Biol. 21:700–704.
Cooley AM, Willis JH. 2009. Genetic divergence causes parallel evolution of flower color in Chilean Mimulus. New Phytol. 183:729–739.
Daxinger L, Hunter B, Sheikh M, Jauvion V, Gascioli V, et al. 2008. Unexpected silencing effects from t-DNA tags in Arabidopsis. Trends Plant Sci. 13:4–6.
Ding B, Patterson EL, Holalu SV, Li J, Johnson GA, et al. 2020. Two MYB proteins in a self-organizing activator-inhibitor system produce spotted pigmentation patterns. Curr Biol. 30:802–814.
Dooner HK, Robbins TP, Jorgensen RA. 1991. Genetic and developmental control of anthocyanin biosynthesis. Annu Rev Genet. 25:173–199.
Dubos C, Stracke R, Grotewold E, Weisshaar B, Martin C, et al. 2010. MYB transcription factors in Arabidopsis. Trends Plant Sci. 15:573–581.
Earley KW, Haag JR, Pontes O, Opper K, Juehne T, et al. 2006. Gateway-compatible vectors for plant functional genomics and proteomics. Plant J. 45:616–629.
Edger PP, Smith R, McKain MR, Cooley AM, Vallejo-Marín M, et al. 2017. Subgenome dominance in an interspecific hybrid, synthetic
allopolyplploid, and a 140-year-old naturally established neo-allopolyplploid monkeyflower. Plant Cell. 29:2150–2167.
Gierer A, Meinhardt H. 1972. A theory of biological pattern formation. Kybernetik. 12:30–39.
Gompel N, Prud’homme B, Wittkopp PJ, Kassner VA, Carroll SB. 2005. Chance caught on the wing: cis-regulatory evolution and the origin of pigment patterns in Drosophila. Nature. 433:481–487.
Grossenbacher DL, Whitall JB. 2011. Increased floral divergence in sympatric monkeyflowers. Evolution. 65:2712–2718.
Hoekstra HE, Coyne JA. 2007. The locus of evolution: evo devo and the genetics of adaptation. Evolution. 61:995–1016.
Holton TA, Cornish EC. 1995. Genetics and biochemistry of anthocyanin biosynthesis. Plant Cell. 7:1071.
Kanehisa M, Sato Y, Kawashima M, Furumichi M, Tanabe M. 2016. Kegg as a reference resource for gene and protein. Nucleic Acids Res. 44:D457–D462.
Katzen F. 2007. Gateway™ recombinational cloning: a biological operating system. Exp Opin Drug Discov. 2:571–589.
Koes R, Verweij J, Quattrocchio F. 2005. Flavonoids: a colorful introduction to the genetics of adaptation. Evolution. 61:995–1016.
Koester S, Pruss GJ, Gao Z, Mgutshini NL, Li J, et al. 2010. Transmissional silencing induced by Arabidopsis t-DNA mutants is associated with 35S promoter siRNAs and requires genes involved in siRNA-mediated chromatin silencing. Plant J. 64:699–704.
Love MI, Huber W, Anders S. 2014. Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biol. 15:550.
Lowry DB, Sheng CC, Lasky JR, Willis JH. 2012. Five anthocyanin polymorphisms are associated with an R2R3-MYB cluster in Mimulus guttatus (Phrymaceae). Am J Bot. 99:82–91.
Lowry DB, Sobel JM, Angert AL, Ashman T-L, Baker RL, et al. 2019. The case for the continued use of the genus name Mimulus for all allopolyploid species and variegates. Am J Bot. 106:30–39.
Martin TR, Jiang P, Rausher MD. 2017. How petals change their spots: cis-regulatory re-wiring in Clarkia (Onagraceae). New Phytol. 216:510–518.
Mihalich H. 2012. Turing’s theory of morphogenesis of 1952 and the subsequent discovery of the crucial role of local self-enhancement and long-range inhibition. Interface Focus. 2:407–416.
Meinhardt H, Gierer A. 2000. Pattern formation by local self-activation and lateral inhibition. Bioessays. 22:753–760.
Mokshwala S, Pruss GJ, Gao Z, Mtutshini NL, Li J, et al. 2010. Transcriptional silencing induced by Arabidopsis t-DNA mutants is associated with 35S promoter siRNAs and requires genes involved in siRNA-mediated chromatin silencing. Plant J. 64:699–704.
Mukherjee BB, Vickery RK. 1962. Chromosome counts in the section simiolus of the genus Mimulus (Scrophulariaceae). v. the chromosomal homologies of M. guttatus and its allied species and varieties. Madrono. 16:141–155.
Nie Z-L, Sun H, Beardsley PM, Olmstead RG, Wen J. 2006. Evolution of biogeographic disjunction between Eastern Asia and Eastern North America in Phryma (Phrymaceae). Am J Bot. 93:1343–1356.
Nijhout H. 1991. Pattern formation in biological systems. In: Lectures in Complex Systems, SFI Studies in the Sciences of Complexity. Vol. IV. L, Nadel, D, Stein, editors. p. 159–187. Addison Wesley Publishing Company.
Petrone K, Tonelli C. 2011. Recent advances on the regulation of anthocyanin synthesis in reproductive organs. Plant Sci. 181:219–229.
Schaafsma DW, Bradshaw H. 1999. Pollinator preference and the evolution of floral traits in monkeyflowers (Mimulus). Proc Natl Acad Sci U S A. 96:11910–11915.
Sobel JM, Streisfeld MA. 2013. Flower color as a model system for studies of plant evo-devo. Front Plant Sci. 4:321.
Stanton K, Valentin CM, Wijnen ME, Stustman S, Palacios JJ, et al. 2016. Absence of postmatting barriers between a selfing vs. outcrossing Chilean Mimulus species pair. Am J Bot. 103:1030–1040.
Stanton KA, Edger PP, Puzyr JK, Kinser T, Cheng P, et al. 2017. A whole-transcriptome approach to evaluating reference genes for quantitative gene expression studies: a case study in Mimulus. G3 (Bethesda). 7:1085–1099.
Stein LD. 2011. An introduction to the informatics of “next-generation” sequencing. Curr Protoc Bioinformatics. 36:11.
Stern DL, Orgogozo V. 2008. The loci of evolution: how predictable is genetic evolution? Evolution. 62:2155–2177.
Stracke R, Werber M, Weisshaar B. 2001. The R2R3-MYB gene family in Arabidopsis thaliana. Curr Opin Plant Biol. 4:447–456.
Streisfeld MA, Rausher MD. 2009. Altered trans-regulatory control of gene expression in multiple anthocyanin genes contributes to adaptive flower color evolution in Mimulus aurantiacus. Mol Biol Evol. 26:433–444.
Streisfeld MA, Young WN, Sobel JM. 2013. Divergent selection drives genetic differentiation in an R2R3-MYB transcription factor that contributes to incipient speciation in Mimulus aurantiacus. PLoS Genet. 9:e1003385.
Tanaka Y, Sasaki N, Ohmiya A. 2008. Biosynthesis of plant pigments: anthocyanins, betalains and carotenoids. Plant J. 54:733–749.
Turing A. 1952. The chemical theory of morphogenesis. Philos Trans R. Soc. 13:5–72.
Twyford AD, Streisfeld MA, Lowry DB, Friedman J. 2015. Genomic studies on the nature of species: adaptation and speciation in Mimulus Mol Ecol. 24:2601–2609.
Vallejo-Marín M, Buggs RJ, Cooley AM, Puzyr JK. 2015. Speciation by genome duplication: repeated origins and genomic composition of the recently formed allopolyploid species Mimulus peregrinus. Evolution. 69:1487–1500.
Vickery RK. 1995. Speciation by aneuploidy and polyploidy in Mimulus (Scrophulariaceae). Great Basin Nat. 55:174–176.
Weinhold A, Kallenbach M, Baldwin IT. 2013. Progressive 35S promoter methylation increases rapidly during vegetative development in transgenic Nicotiana attenuata plants. BMC Plant Biol. 13:99.
Werner T, Koshikawa S, Williams TM, Carroll SB. 2010. Generation of a novel wing colour pattern by the Wingless morphogen. Nature. 464:1143–1148.
Williams AG, Thomas S, Wyman SK, Holloway AK. 2014. RNA-seq data: challenges in and recommendations for experimental design and analysis. Curr Protoc Hum Genet. 83:11.13.1–11.13.20.
Xu W, Dubos C, Lepiniec L. 2015. Transcriptional control of flavonoid biosynthesis by MYB–bHLH–WDR complexes. Trends Plant Sci. 20:176–185.
Xu W, Grain D, Bobet S, Gourrierec JL, Thévenin J, et al. 2014. Complexity and robustness of the flavonoid transcriptional regulatory network revealed by comprehensive analyses of MYB–bHLH–WDR complexes and their targets in Arabidopsis seed. New Phytol. 202:132–144.

Yuan Y-W. 2019. Monkeyflowers (Mimulus): new model for plant developmental genetics and evo-devo. New Phytol. 222:694–700.

Yuan Y-W, Sagawa JM, Frost L, Vela JP, Bradshaw HD, Jr. 2014. Transcriptional control of floral anthocyanin pigmentation in monkeyflowers (Mimulus). New Phytol. 204:1013–1027.

Yuan Y-W, Sagawa JM, Young RC, Christensen BJ, Bradshaw HD. 2013. Genetic dissection of a major anthocyanin qtl contributing to pollinator-mediated reproductive isolation between sister species of Mimulus. Genetics. 194:255–263.

Zhu H-F, Fitzsimmons K, Khandelwal A, Kranz RG. 2009. Cpc, a single-repeat R3 MYB, is a negative regulator of anthocyanin biosynthesis in Arabidopsis. Mol Plant. 2:790–802.

Zhu Z, Wang H, Wang Y, Guan S, Wang F, et al. 2015. Characterization of the cis elements in the proximal promoter regions of the anthocyanin pathway genes reveals a common regulatory logic that governs pathway regulation. J Exp Bot. 66:3775–3789.

Communicating editor: K. Bomblies