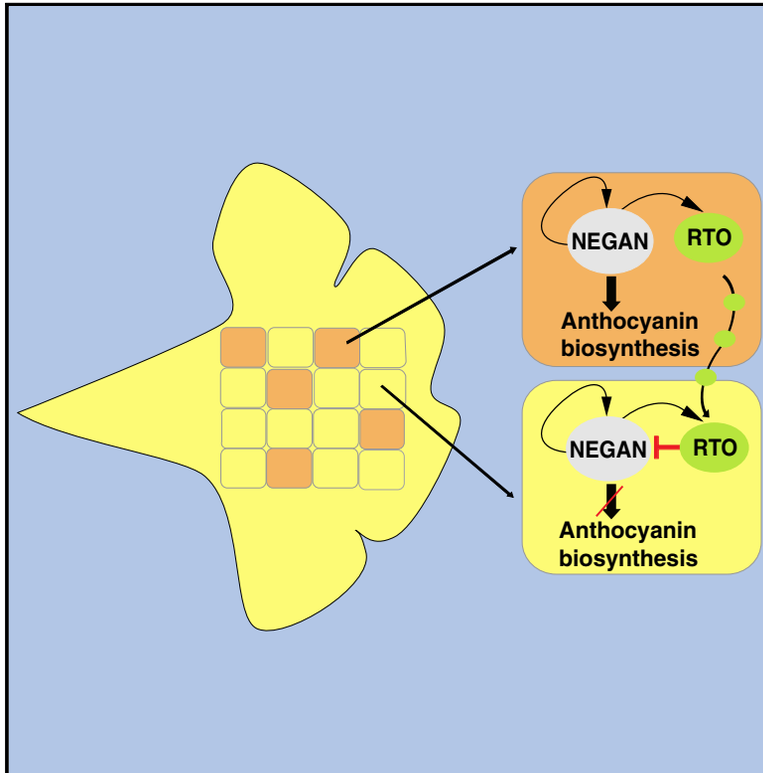


Current Biology

Two MYB Proteins in a Self-Organizing Activator-Inhibitor System Produce Spotted Pigmentation Patterns

Graphical Abstract



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In Brief

Many species exhibit visually striking spotted or striped pigmentation patterns. By studying mutants that alter the red spots that dapple monkeyflower petals, Ding et al. uncover a simple activator-inhibitor system that conforms to long-standing theoretical predictions for how such patterns self-organize during development.

Highlights

- An activator-inhibitor system produces spotted pigment patterning in *Mimulus* flowers
- Expression of the inhibitor, RTO, is promoted by the activator, NEGAN
- RTO protein is mobile and inhibits NEGAN function in neighboring cells
- Modulating this nectar guide patterning system influences bumblebee visitation



Two MYB Proteins in a Self-Organizing Activator-Inhibitor System Produce Spotted Pigmentation Patterns

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SUMMARY

Many organisms exhibit visually striking spotted or striped pigmentation patterns. Developmental models predict that such spatial patterns can form when a local autocatalytic feedback loop and a long-range inhibitory feedback loop interact. At its simplest, this self-organizing network only requires one self-activating activator that also activates a repressor, which inhibits the activator and diffuses to neighboring cells. However, the molecular activators and inhibitors fully fitting this versatile model remain elusive in pigmentation systems. Here, we characterize an R2R3-MYB activator and an R3-MYB repressor in monkeyflowers (*Mimulus*). Through experimental perturbation and mathematical modeling, we demonstrate that the properties of these two proteins correspond to an activator-inhibitor pair in a two-component, reaction-diffusion system, explaining the formation of dispersed anthocyanin spots in monkeyflower petals. Notably, disrupting this pattern impacts pollinator visitation. Thus, subtle changes in simple activator-inhibitor systems are likely essential contributors to the evolution of the remarkable diversity of pigmentation patterns in flowers.

INTRODUCTION

Pigmentation patterns such as the stripes of zebras, patches of giraffes, and petal spots of many flowers have fascinated biologists and mathematicians for centuries. These patterns often function as visual cues in mate choice and interspecies

communications, adapting organisms to their environments and potentially contributing to reproductive isolation among species [1–4]. Unlike previously well-characterized pigmentation patterns in insect wings and some flowers that are spatially organized by coinciding expression domains of regulatory genes that serve as position-specific signals [5–11], our understanding of how dispersed pigmentation patterns emerge in concert with or entirely untethered from such positional information is primarily theoretical [12–15]. The reaction-diffusion (RD) model, initially proposed by Turing [16] and then independently developed and elaborated by Gierer and Meinhardt [17–19] and further extended by others [12, 20–22], postulates that local activation of pattern differentiation factors combined with long-range inhibition of the activity of those factors can produce defined, regularly spaced, tissue-level spot and stripe patterns. In its simplest form, the RD model for self-organizing pattern formation only requires two interacting components: an activator that activates itself as well as a repressor, which diffuses to neighboring cells faster than the activator and inhibits the activator's function [12, 17, 18]. However, when the RD interacting network contains more than two nodes, pattern formation does not necessarily require differential diffusivity of the activator and inhibitor as in the two-component RD system [22]. Candidate activators and inhibitors that fit the RD model have been advanced for several developmental patterns ([15, 23–27]; but see [13, 28, 29] for alternative mechanisms not reliant on molecular players). However, the molecular identities and dynamics of the activation and inhibition factors that contribute to self-organized pigmentation patterning have remained elusive, particularly in relevant animal systems where mutant alleles often have deleterious pleiotropic consequences or critical embryonic developmental stages are challenging to study experimentally.

Anthocyanin spots in flower petals provide an excellent empirical system to reveal the molecular basis for the formation and evolution of self-organized pigmentation patterns. These



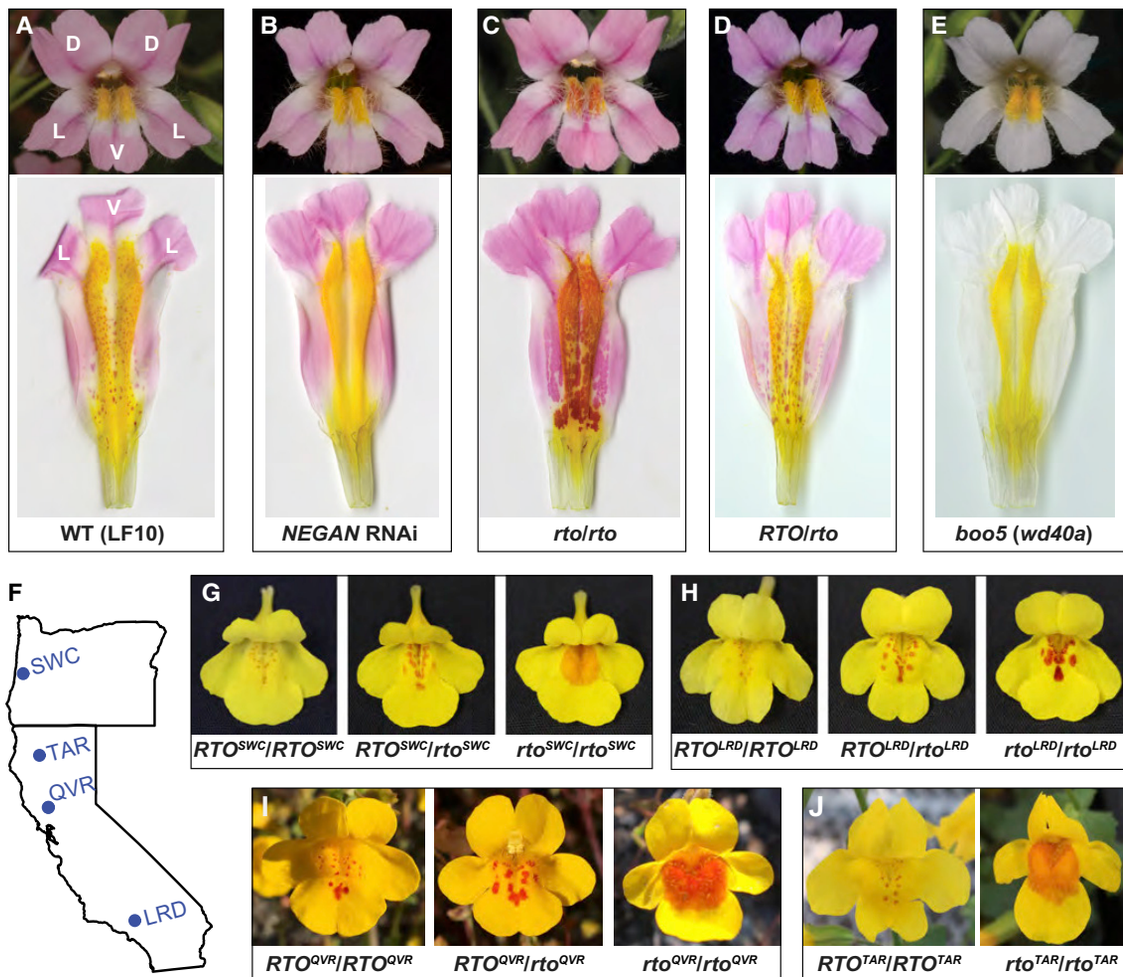


Figure 1. Dispersed Anthocyanin Spots in *M. lewisii* and *M. guttatus*

(A) Anthocyanin spots on the yellow nectar guides of the wild-type (WT) *M. lewisii* (LF10). D, dorsal; L, lateral; V, ventral.
 (B) *NEGAN* RNAi line lacking anthocyanins in the nectar guides.
 (C) Homozygous *rto* mutant with expanded anthocyanin pigmentation.
 (D) Heterozygous *rto* mutant with intermediate phenotype.
 (E) Homozygous *wd40a* mutant lacking anthocyanins in both petal lobes and nectar guides.
 (F–J) Anthocyanin spot patterns of *M. guttatus* variants segregating in natural populations in Oregon and California. Sampling locations (F) and wild-type homozygote, heterozygote, and *rto*-like homozygote phenotypes are shown for the SWC (G), LRD (H), QVR (I), and TAR (J) populations.
 See also [Figures S1](#) and [S2](#).

patterns, which are highly diverse in angiosperms even among different varieties of the same species [30, 31], are known to serve as critical cues in plant-pollinator interactions [32–34], and the genetic network controlling anthocyanin pigment production is otherwise well described [35, 36]. Although some factors involved in the regulation and natural variation of floral pigment patterning have been identified, these factors act solely through pre-patterning by underlying developmental programs and do not yield periodic or randomly dispersed spatial patterns [7–9]. To reveal developmental mechanisms that contribute to dispersed floral pigment patterns, we have focused on the anthocyanin spots that dot the nectar guides of many species in the monkeyflower genus *Mimulus*. The presence of these spots facilitates effective pollinator visitation [32], and genomic resources and methods for genetic transformation that foster

rigorous genetic and developmental studies have been established for multiple *Mimulus* species [37–39].

Previously, we identified a self-activating R2R3-MYB transcription factor, NECTAR GUIDE ANTHOCYANIN (*NEGAN*), that activates anthocyanin spot formation in the monkeyflower species *Mimulus lewisii* [40] (Figure 1A). Downregulation of *NEGAN* expression abolishes anthocyanin production in the nectar guides (i.e., the two yellow ridges of the ventral petal; Figure 1B). We hypothesized that the dispersed anthocyanin spots could be explained by a simple two-component RD model, which predicts (1) there should be an inhibitor corresponding to the anthocyanin activator *NEGAN*, and loss-of-function mutations in the presumptive inhibitor gene should cause expansion of the spots; (2) the inhibitor is activated by the activator (i.e., *NEGAN*); (3) the inhibitor represses the function of the activator;

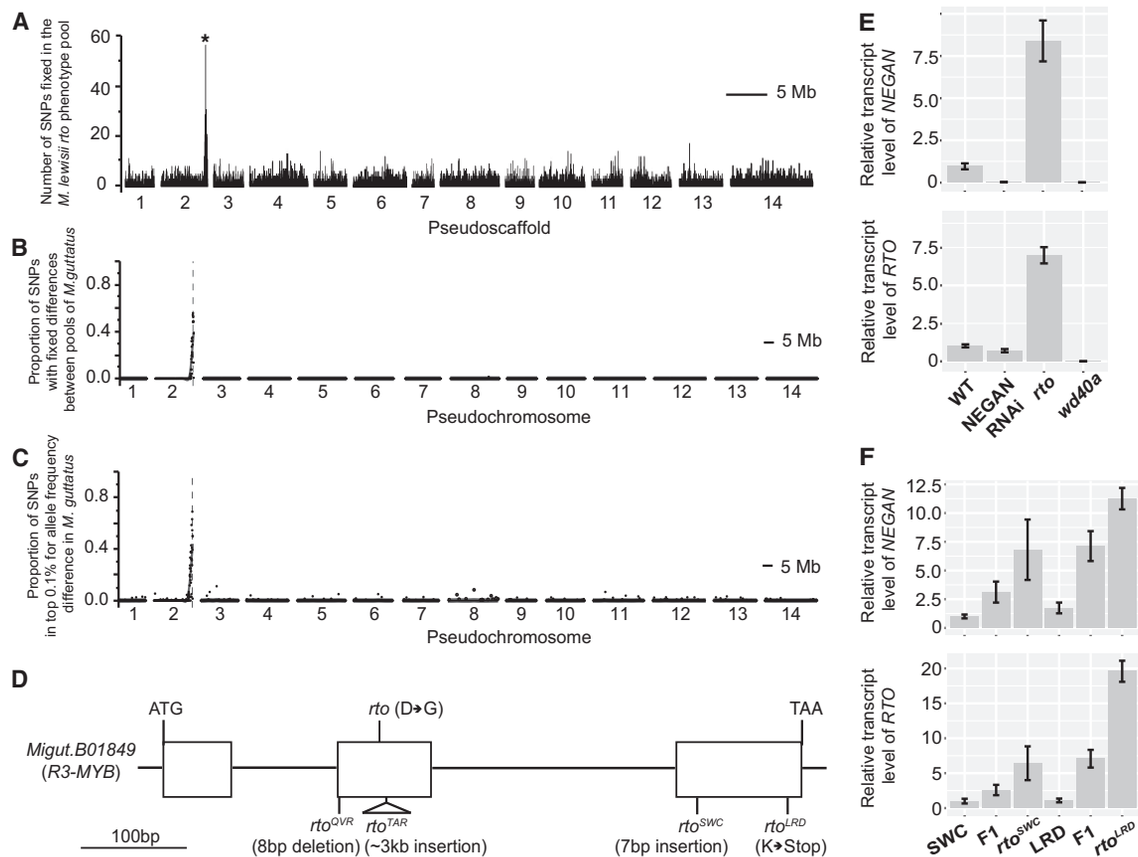


Figure 2. Identification of the RTO Gene in *M. lewisii* and *M. guttatus* and Its Relative Expression in Various Mutant and Transgenic Lines (A–C) Bulk segregant analyses of *rto* in *M. lewisii* (A) and *rto*^{SWC} (B) and *rto*^{LRD} (C) in *M. guttatus* narrowed RTO down to the same genomic interval. (D) Schematic of the RTO (*R3-MYB*) gene showing the molecular lesions of the five mutant alleles. (E and F) Relative transcript level of NEGAN (upper) and RTO (lower) in *M. lewisii* (E) and *M. guttatus* (F) as measured by qRT-PCR, standardized to the corresponding wild type (LF10 for *M. lewisii*; SWC for *M. guttatus*). Error bars represent 1 SD from three biological replicates. See also Figures S3 and S4 and Tables S4 and S6.

and (4) the inhibitor can move from cell to cell. Here, through genetic mapping of a chemically induced mutant and natural variants with expanded anthocyanin spots followed by manipulative experiments and computer simulations, we identify a second molecular factor involved in anthocyanin spot patterning in *Mimulus* petals, reveal how this protein functions as a long-range inhibitor of NEGAN to regulate spot formation, and evaluate the evolutionary and ecological impacts of modulating this self-organizing patterning mechanism.

RESULTS

The R3-MYB Gene RED TONGUE Is a Negative Regulator of Anthocyanin Spot Patterning

To search for the putative inhibitor involved in anthocyanin spot formation, we carried out an ethyl methanesulfonate mutagenesis screen using the *M. lewisii* inbred line LF10 and isolated one candidate mutant, named *red tongue* (*rto*). Homozygous *rto* mutants develop a near-uniform patch of anthocyanin color across the entire nectar guide region (Figure 1C). Coincidentally, we discovered naturally occurring mutants with *rto*-like phenotypes segregating in several geographically distinct populations

of the congener *Mimulus guttatus* (Figures 1F–1J). Complementation crosses among these populations indicate that these natural *M. guttatus* variants represent different alleles of the same locus (Figure S1A). The *rto* mutant and *rto*-like natural alleles behave semi-dominantly; F₁ plants derived by crossing these alleles to corresponding wild-type individuals exhibit intermediate anthocyanin spot number and area phenotypes (Figures 1D, 1G–1I, and S2).

Although *M. lewisii* and *M. guttatus* diverged about 20 million years ago [41], the similarity and specificity of the altered pigmentation phenotypes led us to suspect that all these mutant phenotypes are caused by lesions in the same gene. To identify the gene responsible for the *rto* phenotype in each species, we performed bulked segregant analyses by whole-genome sequencing on F₂ mapping panels segregating for the *M. lewisii* mutant and the two *rto*-like *M. guttatus* alleles found at Sweetwater Creek (SWC) in Oregon (Figure 1G) and Litterlock Dam (LRD) in Southern California (Figure 1H), i.e., *rto*^{SWC} and *rto*^{LRD}. As expected, the independent mapping experiments all pinpointed the same causal genomic interval (Figures 2A–2C). Inspection of the bulked segregant reads aligning to this ~60-kb region in the wild-type *M. lewisii* (LF10) genome revealed

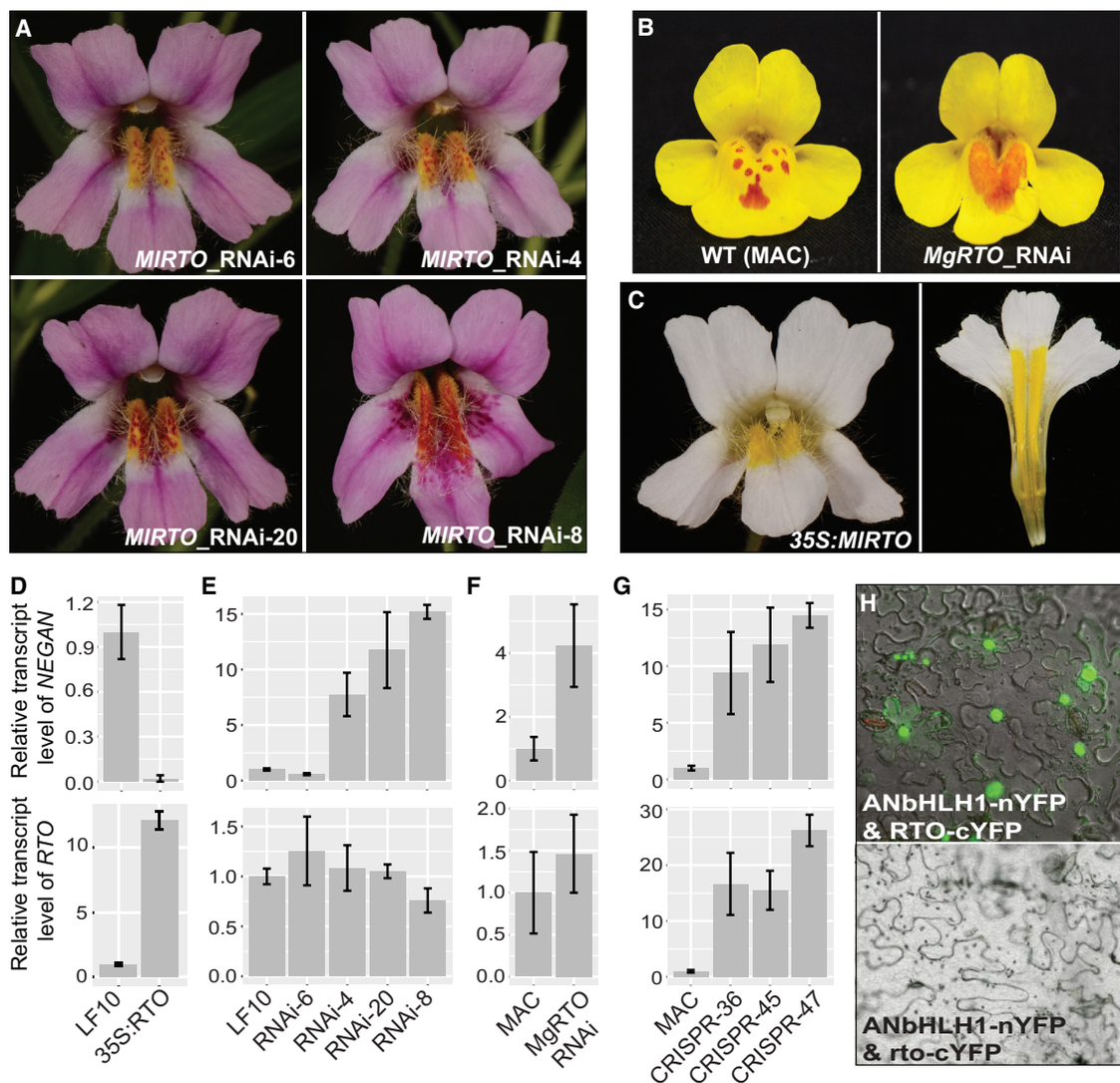


Figure 3. Functional Characterization of RTO in *M. lewisii* and *M. guttatus*

(A) RNAi of RTO in *M. lewisii* generates a range of anthocyanin spot patterns.

(B) RNAi of MgRTO in *M. guttatus* recapitulates the *rto*-like phenotype.

(C) Overexpression of RTO in *M. lewisii* abolishes anthocyanin production throughout the corolla.

(D–G) Relative expression of NEGAN and RTO in *M. lewisii* RTO overexpression lines (D), *M. lewisii* RTO RNAi lines (E), *M. guttatus* RTO RNAi lines (F), and *M. guttatus* CRISPR/Cas9-mediated knockout lines (G). All relative transcript levels are measured by qRT-PCR, standardized to the corresponding wild type (LF10 for *M. lewisii*; Mackville Road, CA [MAC] for *M. guttatus*). Error bars represent 1 SD from three biological replicates.

(H) Bimolecular fluorescence complementation (BiFC) assay shows that the wild-type RTO protein interacts with ANbHLH1, whereas the D>G amino acid replacement in the mutant *rto* protein abolishes or attenuates the interaction.

See also Figures S2 and S4–S6 and Tables S5 and S6.

only one mutation, which causes an aspartic acid-to-glycine replacement at a highly conserved site of a small R3-MYB protein (the ortholog of Migut.B01849) that is 91 amino acids long (Figures 2D and S3A). Furthermore, fine mapping resolved the *M. guttatus rto*^{SWC} allele to an interval spanning four genes (Figure S3B), including the same R3-MYB gene *Migut.B01849*. Sanger sequencing of this gene revealed at least one putative loss-of-function mutation in each of the four *M. guttatus rto*-like alleles (Figure 2D). Notably, this R3-MYB is closely related to a group of R3-MYBs that are known to repress anthocyanin biosynthesis (Figure S3A), including ROSE INTENSITY1 in

M. lewisii [42], MYBx in *Petunia* [43], and CAPRICE in *Arabidopsis* [44]. Taken together, these results strongly suggest this R3-MYB gene is the causal gene underlying the RTO locus in both *M. lewisii* and *M. guttatus*.

To verify the function of this R3-MYB gene, we knocked down its expression level in *M. lewisii* and *M. guttatus* by RNA interference (RNAi). We obtained multiple independent transgenic lines from both species with severe phenotypes that recapitulate the mutant phenotypes (Figures 3A, 3B, and S2A). Some lines in the *M. lewisii* background have an even stronger phenotype than *rto* toward the throat of the corolla

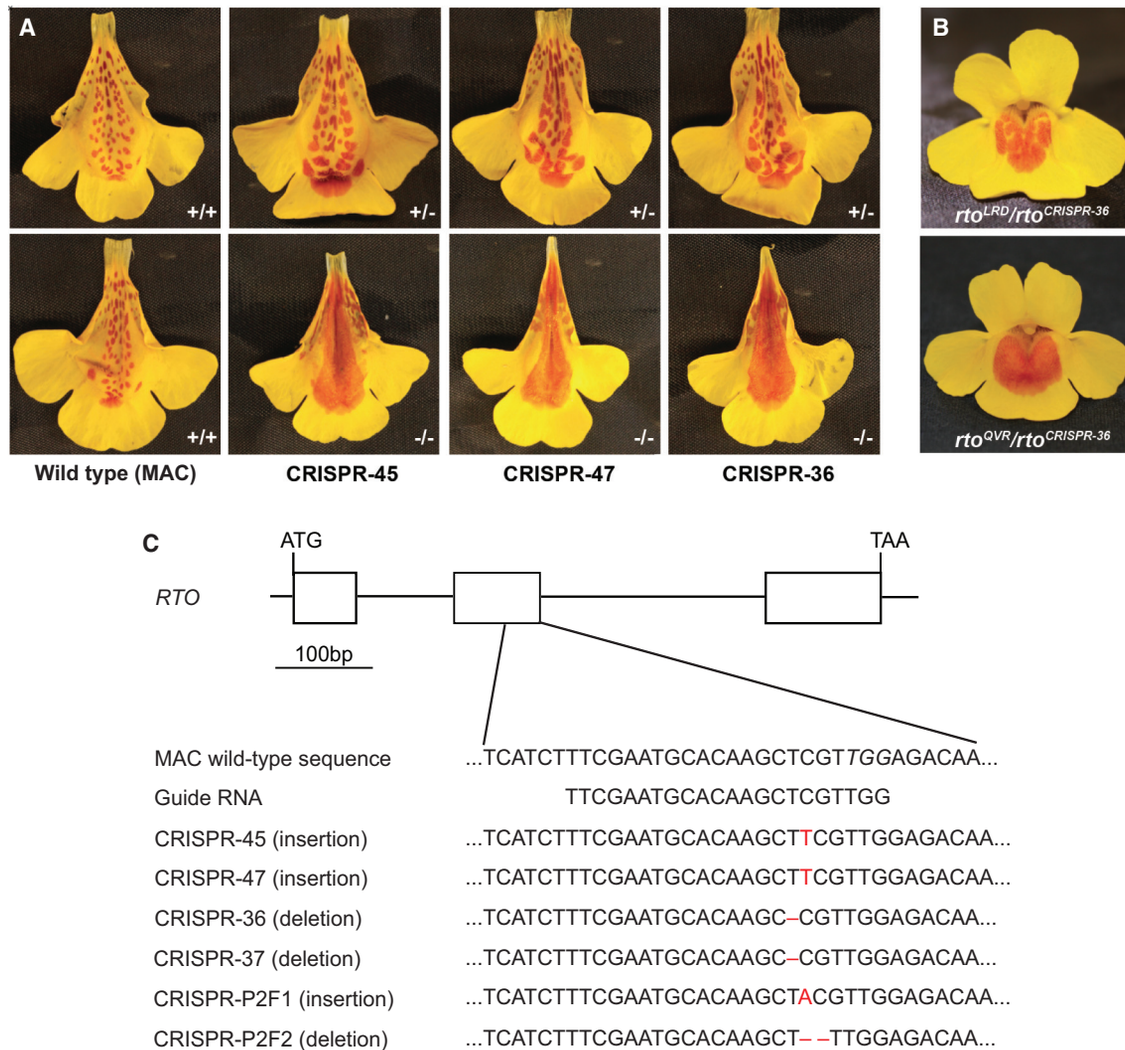


Figure 4. CRISPR/Cas9-Mediated Knockout of *MgRTO* in *M. guttatus*

(A) Flower phenotypes of three independent CRISPR knockout lines in their heterozygous (+/-) and homozygous (-/-) conditions, compared to the wild type (MAC; +/+).

(B) Complementation crosses between a CRISPR allele ($rto^{CRISPR-36}$) and two natural mutant alleles (rto^{LDR} and rto^{QVR}).

(C) Examples of six independent CRISPR alleles relative to the guide RNA (gRNA) position in *MgRTO*. The primer pair 5'-GTCTTCATATATTTCCCATCTCTT-3' and 5'-CGTGCTCGGTGTAAGTAACG-3' was used for amplifying and sequencing the target region.

See also Figure S2 and Table S5.

(e.g., RNAi line 8; Figure 3A), suggesting that *rto* is likely hypomorphic (partially functional) rather than a null allele. Notably, CRISPR/Cas9-mediated frameshift mutations in *MgRTO* also recapitulate the *rto* phenotype (Figures 4A, 4C, and S2B) and the targeted mutation does not complement natural variants (Figure 4B), confirming that they are allelic. Together, the association of expanded pigmentation with five independent lesions in *RTO* and with RNAi knockdown and/or targeted disruptions of *RTO* in two congeneric species strongly indicates that *RTO* functions as an inhibitor of anthocyanin spot patterning. As such, we expected that overexpression of *RTO* driven by the CaMV 35S promoter would attenuate or completely abolish the anthocyanin spots. Indeed, the corollas of strong 35S:*MIRTO* lines in *M. lewisii* are completely white and lack anthocyanin spots in the nectar

guide region (Figure 3C), confirming that *RTO* acts as a potent inhibitor of anthocyanin pigmentation.

NEGAN Promotes *RTO* Expression

Having identified the putative inhibitor, we then tested whether *NEGAN* and *RTO* function as a local activator/lateral inhibitor pair consistent with the simple two-component RD model [17, 18]. In particular, we sought to test whether the self-activating activator *NEGAN* activates the inhibitor (i.e., *RTO*), which then diffuses laterally and inhibits the function of the activator along the diffusion path. To test the first prediction that the local activator also activates the inhibitor, we compared *RTO* gene expression levels between wild-type and *NEGAN* RNAi plants as well as the *M. lewisii* *wd40a* mutant. The *wd40a* mutant is relevant because in all angiosperms characterized

to date, the anthocyanin-activating R2R3-MYB transcription factor (e.g., *NEGAN*) functions in an obligatory protein complex that also contains a bHLH protein and a WD40 protein [35, 36]. In *M. lewisii* flowers, the bHLH and WD40 components of the complex (*MIANbHLH1* and *MIWD40a*, respectively) are expressed in the entire corolla. The R2R3-MYB component is supplied by two paralogs, *NEGAN* and *PETAL LOBE ANTHOCYANIN (PELAN)*, that are expressed in nectar guides and petal lobes, respectively [40]. The *wd40a* mutants completely lack anthocyanins in both the petal lobes and the nectar guides (Figure 1E). Downregulation of *NEGAN* by RNAi abolishes the anthocyanin spots in the nectar guides but does not affect the petal lobe color (Figure 1B), whereas the loss-of-function *pelan* mutant and *PELAN* RNAi lines lose all the anthocyanin pigments in the petal lobe while the nectar guide spots remain unperturbed [40] (Figure S4A). If nectar guide expression of *RTO* is activated by *NEGAN* acting in a complex with *MIANbHLH1* and *MIWD40a*, we would expect reduced *RTO* expression in *NEGAN* RNAi lines and the *wd40a* mutant compared to wild-type *M. lewisii* plants.

Indeed, for whole corollas sampled from 10-mm-stage flower buds, the developmental stage when both *NEGAN* and *RTO* show peak expression level and the anthocyanin spots just start appearing (Figures S4B, S4D, and S4F), the *RTO* transcript level decreased to an undetectable level in the *wd40a* mutant compared to the wild type (Figure 2E). *NEGAN* expression is also abolished in the *wd40a* mutant (Figure 2E), as previously shown [40], because the ability of *NEGAN* to self-activate also requires the complete *NEGAN-MIANbHLH1-MIWD40a* protein complex. In the 10-mm-stage whole corollas of *M. lewisii* *NEGAN* RNAi lines, however, the *RTO* transcript level only slightly decreased compared to the wild type (Figure 2E). We predicted this finding resulted from activation of *RTO* expression in the petal lobe by *PELAN*. Indeed, when examined in dissected petal lobes and nectar guides at this stage, petal lobe *RTO* expression is unaffected in the *NEGAN* RNAi lines but reduced to an undetectable level in the *pelan* mutant, whereas nectar guide *RTO* expression is unaffected in the *pelan* mutant but clearly reduced in the *NEGAN* RNAi lines (Figure S4C). These results suggest that *MIRTO* expression is activated by *PELAN* and *NEGAN* in the petal lobes and nectar guides, respectively, and requires the same WD40 component in both tissues. That *PELAN* cannot self-activate [40] may explain the generally uniform anthocyanin pigmentation in the petal lobes, compared to the dispersed spots in the nectar guides. Similarly, when *MgNEGAN* expression was knocked down in *M. guttatus*, the anthocyanin spots completely disappeared from nectar guides of strong RNAi lines, and *MgRTO* expression levels decreased accordingly (Figure S5). Taken together, our results confirm that the activator, *NEGAN*, activates the expression of the inhibitor, *RTO*, in the nectar guides of both *Mimulus* species.

RTO Is a Negative Regulator of NEGAN Function

To test whether the inhibitor represses the activator's function, we assayed relative *NEGAN* expression levels in the *rto* mutant and *rto*-like natural variants as well as various *RTO* transgenic lines. Because *NEGAN* expression is self-activated, if *RTO* inhibits the activity of *NEGAN*, loss of *RTO* function should

increase *NEGAN* expression, and overexpression of *RTO* should do the opposite. Consistent with those predictions, *NEGAN* transcript abundance increased 6- to 11-fold in *rto* and the *rto*-like natural variants (Figures 2E and 2F) and decreased >10-fold in the strong 35S:*MIRTO* lines (Figure 3D). In the *M. lewisii* *RTO* RNAi lines, *NEGAN* expression increased as much as 15-fold, and *NEGAN* expression level is positively correlated with the severity of the anthocyanin phenotype (Figures 3A and 3E). Likewise, *MgNEGAN* expression is also upregulated in the *MgRTO* RNAi and targeted knockout lines (Figures 3F and 3G). These results strongly indicate that *RTO* inhibits *NEGAN* function.

That the transcript level of *RTO* itself in the *RTO* RNAi lines is not lower than in the wild-type plants (Figures 3E and 3F) seems surprising at first glance. However, this finding is consistent with the anticipated dynamic interactions between *NEGAN* and *RTO*. As shown above, knockdown of *RTO* transcript levels by RNAi resulted in upregulation of *NEGAN*, which in turn would upregulate *RTO*, but the increased *RTO* transcript level would then be counterbalanced by RNAi. Consequently, unlike *NEGAN*, *RTO* transcript level showed no clear correlation with the severity of the anthocyanin phenotype in the *RTO* RNAi lines. Notably, in *MgRTO* knockout lines, where similar feedback is not expected, *RTO* levels are conspicuously elevated relative to wild-type plants (Figure 3G).

The mechanism by which R3-MYB repressors inhibit R2R3-MYB activator function is well understood for their homologous proteins in *Arabidopsis* and *Petunia* [43–45]. These R3-MYB proteins have neither DNA-binding nor activation domains, but they compete with the R2R3-MYB activator for the limited supply of bHLH proteins, sequestering the bHLH proteins into inactive complexes. Like *NEGAN*, *RTO* also contains the bHLH-interacting motif in its R3 domain. The aspartic acid-to-glycine replacement in the mutant *rto* protein is expected to disrupt the bHLH-interacting motif (Figure S3A). Indeed, in bimolecular fluorescence complementation assays, pairing wild-type *MIRTO* protein and *MIANbHLH1* showed reconstitution of fluorescence but pairing mutant *rto* protein and *MIANbHLH1* yielded no detectable fluorescence signal (Figures 3H and S6A), indicating that the mutation abolishes or attenuates the interaction between R3-MYB and the bHLH protein.

RTO Exhibits Intercellular Movement

To test whether *RTO* exhibits long-range lateral inhibition (i.e., diffusible to neighboring cells), we compared the spatial pattern of *RTO* transcription with the *RTO* protein distribution [46]. If *RTO* moves between cells, *RTO* proteins should be observable over a broader domain of cells than where *RTO* is transcribed. To make this comparison, we first examined the spatial pattern of *RTO* transcription by transforming the wild-type *M. lewisii* (LF10) with a construct that expresses cyan fluorescent protein (CFP) with an endoplasmic reticulum (ER) retention signal driven by the *RTO* promoter. The CFP signal marks the cells with *RTO* promoter activity, as the CFP-ER product cannot move from the source cell to neighboring cells. Multiple independent *RTO_{pro}:CFP-ER* lines showed the same pattern; the CFP fluorescence signal co-localizes with the anthocyanin spots (Figure 5A), consistent with the expectation that both *RTO* and the anthocyanin biosynthetic genes are regulated by the same transcriptional activator (i.e., *NEGAN*).

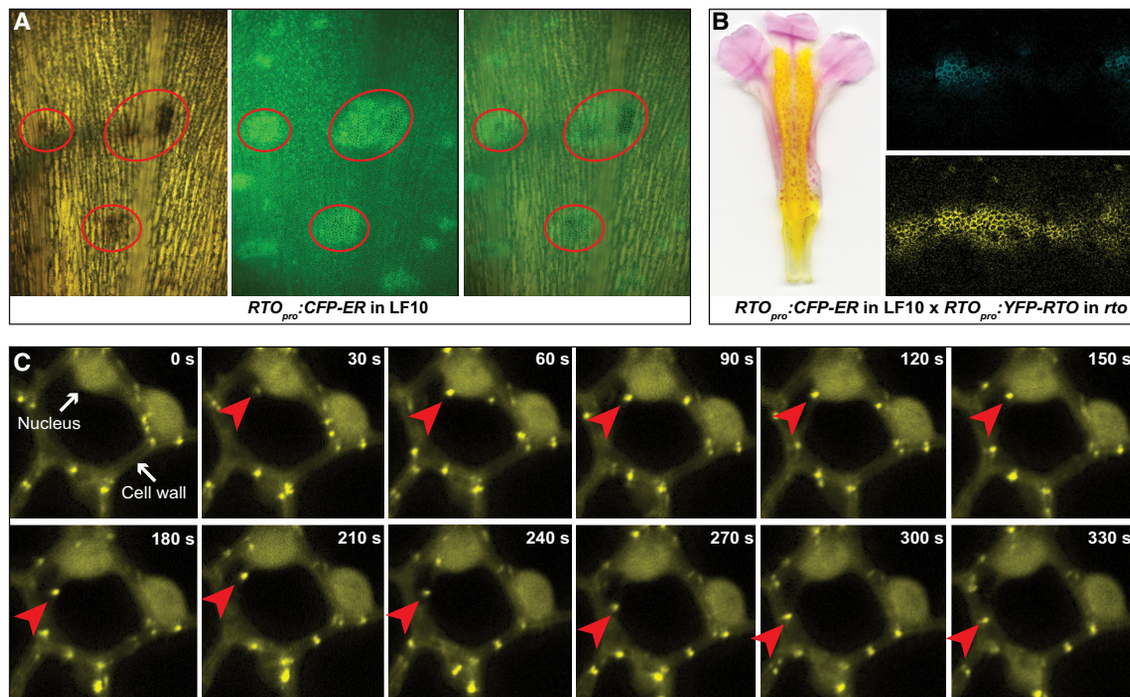


Figure 5. Intercellular Movement of RTO in *M. lewisii*

(A) Spatial pattern of *RTO* promoter activity revealed by the *RTO_{pro}:CFP-ER* construct. The same nectar guide area was imaged under bright light (left) and the green channel (center). The right image is an overlay between the left and the center. The yellow background under bright light is due to carotenoid pigments, and the anthocyanin spots are red. CFP fluorescence signal (i.e., the green spots in the green channel) foreshadows and co-localizes with anthocyanin production (red ovals).

(B) F₁ hybrids between *RTO_{pro}:CFP-ER* line 25 (in the wild-type background) and *RTO_{pro}:YFP-RTO* line 22 (in the *rto* background) bear flowers similar in anthocyanin pigmentation to the non-transgenic *RTO/rto* heterozygote and reveal a broader spatial distribution of RTO protein (yellow) than *RTO* promoter activity (blue).

(C) A series of still images from a time-lapse video taken every 30 s to track the movement of YFP-RTO from the nucleus to the cytoplasm in *35S:YFP-RTO* petal epidermal cells.

See also [Figure S6](#), [Table S5](#), and [Video S1](#).

Next, to reveal the spatial distribution of RTO proteins, we expressed a yellow fluorescent protein (YFP)-RTO fusion protein driven by the same *RTO* promoter in the *rto* background. In the majority of the *RTO_{pro}:YFP-RTO* transformants, the nectar guide anthocyanin pigmentation was reduced and the extent of this “rescue” of the *rto* phenotype was positively correlated with the transgene expression level ([Figures S6B](#) and [S6C](#)). Thus, the YFP-RTO fusion protein retains the function of RTO as an anthocyanin repressor *in vivo*. The complete elimination of nectar guide anthocyanin rather than restoration to the wild-type pattern for a substantial number of *RTO_{pro}:YFP-RTO* lines in the *rto* background was unexpected ([Figure S6B](#)). One possible explanation is that a recessive epistatic interaction between the hypomorphic *rto* allele and the transgene shifts the dynamics of the NEGAN-RTO interaction. However, when we crossed representative *RTO_{pro}:CFP-ER* and *RTO_{pro}:YFP-RTO* lines to bring the two transgenes together in an *RTO/rto* heterozygous background, we observed typical *RTO/rto* anthocyanin patterning rather than the “over-rescue” phenotype ([Figure 5B](#)), indicating wild-type NEGAN-RTO dynamics are re-established. In this background, YFP-RTO signal was detected not only in the source cells of the *RTO* promoter activity, as reflected by the CFP-ER signal, but also in neighboring cells

([Figures 5B](#), [S6D](#), and [S6E](#)). Taking these transcript and protein localization results together, we conclude that the *RTO* gene is transcribed in the anthocyanin spot cells but the RTO protein moves to adjacent cells.

To gain further insight into the mechanism of RTO movement, we generated *35S:YFP-RTO* transformants in *M. lewisii*. Like the strong *35S:RTO* lines, strong *35S:YFP-RTO* transgenic lines produce completely white flowers, confirming again that the YFP tag does not interfere with RTO function. We found YFP-RTO localized to both the nucleus and cytoplasm. We also found many punctate YFP-RTO signals, mainly in the cytoplasm. These puncta travel from the nucleus to the membrane ([Figure 5C](#); [Video S1](#)), reminiscent of the RTO homolog in *Arabidopsis*, CAPRICE, and other mobile proteins such as SHORT ROOT, which are known to interact with protein transport partners and travel through the endosome system in *Arabidopsis* roots [47]. These similarities suggest that the same mechanism might be involved in RTO intercellular movement.

The NEGAN/RTO Pair Fits the Simplest Two-Component Reaction-Diffusion Model

The properties we have described for NEGAN and RTO conform to a two-component activator-inhibitor model, one of the

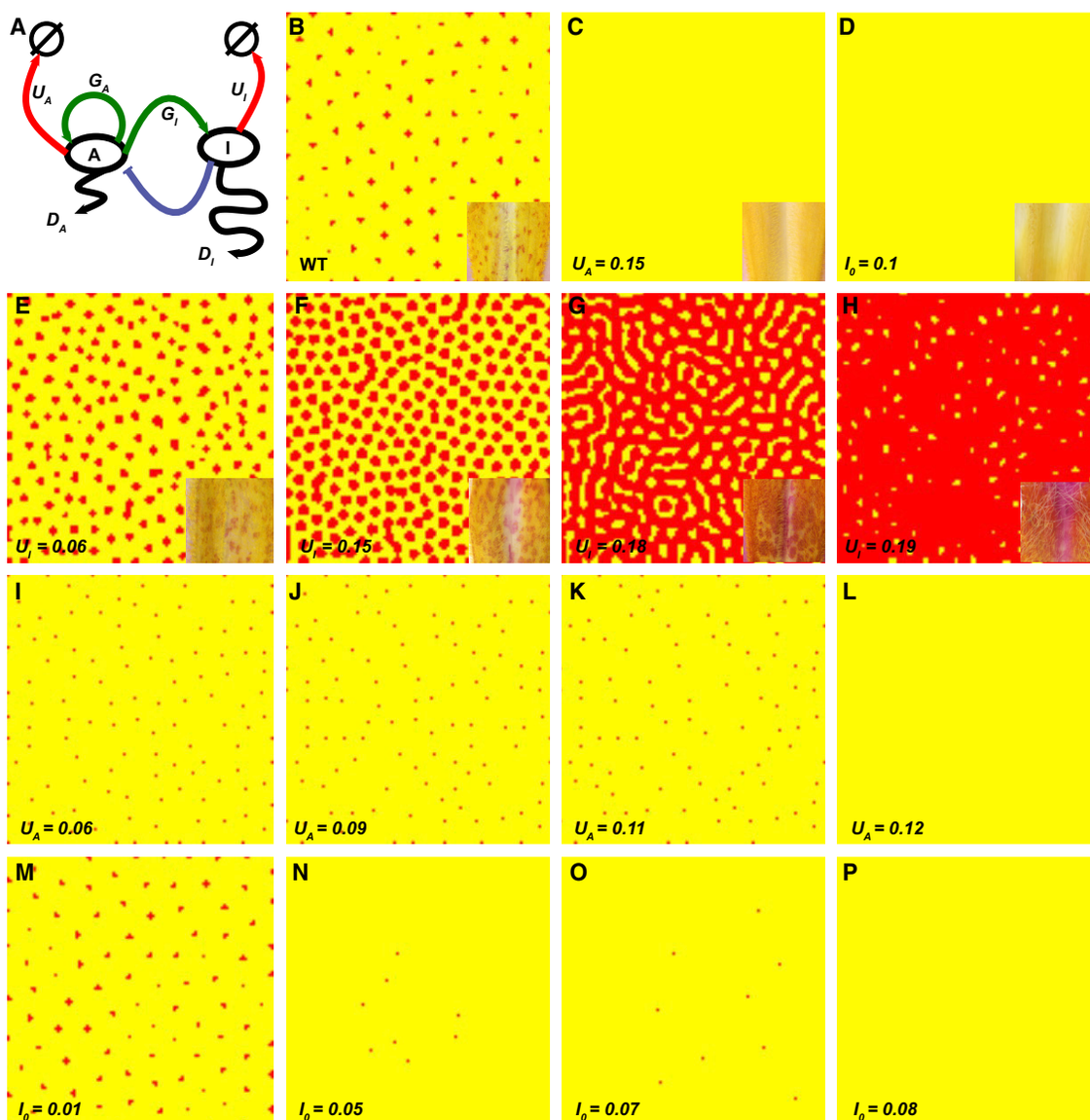


Figure 6. Computer Simulation of the Anthocyanin Spot Patterning

(A) Schematic illustration of a two-component activator-inhibitor model. *A*, activator; *I*, inhibitor; D_A and D_I , diffusion coefficient of the activator and inhibitor, respectively; U_A and U_I , degradation rates; A_0 and I_0 , background production rates; G_A and G_I , potency of the activation by the activator. The specific equations are described in STAR Methods.

(B) Simulated pattern resembling the wild-type *M. lewisii*, with the following parameter values: $D_A = 0.01$; $D_I = 0.5$; $U_A = 0.03$; $U_I = 0.03$; $A_0 = 0.01$; $I_0 = 0$; $G_A = 0.08$; $G_I = 0.12$.

(C–P) Simulated patterns that mimic *NEGAN* RNAi (C and I–L), *RTO* overexpression (D and M–P), and *RTO* RNAi lines (E–H). All parameter values are the same as in the wild-type condition (B), except for one modification for each perturbation as shown in the lower left of each panel. Real flower images of the center of the nectar guides from the corresponding lines are shown in the lower right corner of (B)–(H).

See also Figure S7.

simplest Gierer-Meinhardt topologies underlying pattern formation [17, 18]. We therefore simulated this Gierer-Meinhardt activator-inhibitor model (Figure 6A) with Virtual Cell software [48] to investigate whether tinkering with the kinetics of the two molecules *in silico* can recapitulate the phenotypes that were produced by experimental perturbations. We first determined the parameter values that generated a sparsely spotted pattern resembling wild-type *M. lewisii* (Figure 6B), guided by previous studies simulating spot pattern formation with similar parameter

values that were considered biologically realistic [12, 20, 49, 50]. We then increased the degradation rate of the activator *NEGAN* (U_A), analogous to RNAi-mediated knockdown of *NEGAN* expression. Consistent with the experimental results (Figures 1B and S5), a 4-fold increase of activator degradation rate is sufficient to abolish the spots in the simulation (Figures 6C and 6I–6L). Similarly, increasing the baseline production rate of the inhibitor *RTO* (I_0), akin to the overexpression of *RTO*, resulted in the loss of spots (Figures 6D and 6M–6P). Finally, gradually

Table 1. Anthocyanin Spot Patterning Affects Bumblebee Visitation

Genotype	Observed Visits	Pooled G Value	df	p Value	Heterogeneity G Value	df	p Value
<i>RTO/RTO</i>	246	30.99	1	<0.001	83.34	9	<0.001
<i>rto/rto</i>	393						
<i>RTO/RTO</i>	450	17.92	1	<0.001	37.01	9	<0.001
<i>RTO/rto</i>	566						
<i>RTO/rto</i>	404	1.09	1	0.3	14.18	9	0.12
<i>rto/rto</i>	420						

Pooled goodness-of-fit test over ten trials per pairwise *RTO* genotype combination. Significant heterogeneity G values indicate variation in bee preference among individual trials; nonetheless, in all cases, the effects of genotype in significant individual trials were in the same direction as the pooled test. df, degree(s) of freedom. See also [Figure S1](#) and [Tables S1–S3](#).

increasing the degradation rate of the inhibitor *RTO* (U), mimicking the different degrees of *RTO* knockdown by RNAi, produced patterns resembling the actual RNAi phenotypes in *M. lewisii* ([Figures 6E–6H](#)). Notably, the seemingly counterintuitive experimental observation that *RTO* transcript level did not decrease in the *RTO* RNAi lines ([Figures 3E and 3F](#)) is actually predicted by the reaction kinetics of the Gierer-Meinhardt model. Computer simulations showed that the increase in inhibitor degradation rate resulted in not only higher average activator level, as expected, but also slightly higher average inhibitor level ([Figure S7A](#)). Simulations with another commonly used activator-inhibitor RD model [20, 50], as implemented in the software simRDj [12], also generated similar patterns ([Figure S7B](#)). Taken together, these simulation results further suggest that the anthocyanin spot patterning in *Mimulus* is controlled by a simple two-component activator-inhibitor system.

Alleles Disrupting *MgRTO* Occur at Non-trivial Frequencies in Natural Populations and Impact Bumblebee Attraction

Having demonstrated that *NEGAN* and *RTO* function as an activator-inhibitor pair that patterns anthocyanin spots in the nectar guides, we next explored how variants that alter the *NEGAN*-*RTO* interaction dynamics are distributed in natural populations of *M. guttatus* ([Figures 1F–1J](#)) and their potential impacts on plant performance. In two of three population surveys for *SWC* and *LRD*, we found that *M. guttatus* *rto*-like alleles segregate at low but non-trivial frequencies in the wild (2011 survey: *rto*^{SWC} frequency 0.1, $n = 10$ maternal individuals; 2017 surveys: *rto*^{LRD} frequency 0.02, $n = 51$ and *rto*^{SWC} frequency 0, $n = 49$). Moreover, *rto*-like plants have been seen in at least two additional populations, each >200 km from either *SWC* or *LRD* ([Figure 1F](#)). In one of these populations, *rto*-like plants carry an 8-bp deletion that removes a splice acceptor site and causes a frameshift at the start of the second exon of *MgRTO* ([Figure 2D](#)), and we estimate this allele is present at a frequency of ~ 0.11 in the population based on phenotype (2018 survey: $n = 200$ individuals). In a second population, *rto*-like plants have an ~ 3 -kb insertion that disrupts the second exon and is present at a frequency of ~ 0.07 (2018 survey: $n = 15$ maternal individuals).

The non-trivial frequencies of *rto*-like alleles in several *M. guttatus* populations and the fact that these anthocyanin spots serve as important nectar guides in the closely related *Mimulus luteus* [32] motivated us to test whether natural

variation in the anthocyanin spot patterns could impact pollinator visitation. We carried out a series of bumblebee attraction experiments in a laboratory setting using *Bombus impatiens* and using the *rto*^{SWC} allele because the heterozygous and homozygous genotypes form a broad continuum of spot expansion ([Figure 1G](#)). Although this commercially obtainable species is not native to *M. guttatus* habitats, *B. impatiens* worker bees are comparable in size to those of native congeners (e.g., *Bombus vosnesenskii* and *Bombus californicus*), pollinate *M. guttatus* effectively, and recognize variation in and exert selection on *M. guttatus* floral characters [51–56]. In a direct comparison between genotypes, naive bumblebees preferentially visited *rto*^{SWC} homozygote flowers over wild-type flowers, and bumblebees also preferred heterozygote to wild-type flowers ([Tables 1 and S1–S3](#)). Although we detected significant heterogeneity in attraction among trials for both genotypic contrasts, all trials where one genotype was visited preferentially showed that the *rto*^{SWC}-containing genotype was favored. This heterogeneity may indicate that individual variation in preference segregates within bumblebee populations. Bees did not discriminate between heterozygote flowers and *rto*^{SWC} homozygote flowers, however. Given that corolla UV pigmentation and cell shape do not differ between full-sib *RTO*^{SWC} homozygous and *rto*^{SWC} carrier flowers ([Figures S1B–S1D](#)), we conclude that altered anthocyanin pigmentation pattern is responsible for preferred visitation to *rto*^{SWC} homozygotes and *RTO*^{SWC}/*rto*^{SWC} heterozygotes. Determining whether the *rto*-like alleles represent nascent but incomplete selective sweeps due to positive impacts on pollination or other functions, are drifting neutrally, or are polymorphisms maintained by frequency-dependent selection or at mutation-selection balance due to costs (e.g., florivore attraction) will require further study in field conditions, particularly as pollinator learning can diminish the influence of innate preferences and thus how the polymorphism impacts individual fitness in natural populations. Nonetheless, these findings demonstrate that, in principle, changes in nectar guide patterning that arise by modulating the underlying patterning system can influence fundamental plant-pollinator interactions.

DISCUSSION

In this study, we have shown that a pair of MYB proteins functions as a simple activator-inhibitor system underlying the formation of the dispersed anthocyanin spots that dot the nectar

guides of *Mimulus* flowers. The two-component activator-inhibitor model [12, 17, 18, 20] predicts that the activator self-activates and activates the inhibitor; the inhibitor represses the activator function; and the inhibitor can move from cell to cell. Our experimental evidence is fully consistent with these theoretical predictions. First, the R2R3-MYB activator *NEGAN* activates its own expression [40]. Second, *NEGAN* also activates the expression of the R3-MYB inhibitor *RTO*; when *NEGAN* or its interacting partner *WD40a* is knocked down/out by RNAi or chemical mutagenesis, *RTO* expression is also reduced (Figures 2E and S5). Third, the inhibitor *RTO* represses the activity of *NEGAN*. When *RTO* function is impaired by RNAi or mutations, *NEGAN* expression level increases (Figures 2E, 2F, and 3E–3G), and when *RTO* is overexpressed, *NEGAN* expression is abolished (Figure 3D). Finally, *RTO* shows intercellular movement, as demonstrated by comparing the spatial pattern of *RTO* transcription with the *RTO* protein distribution (Figures 5B, S6D, and S6E). To our best knowledge, these findings are the first to reveal a developmental mechanism that generates dispersed pigment spots in plants. In addition, through genetic mapping and knockdown or targeted knockout experiments of *MgRTO*, our work represents the first instance in which the identity and molecular impact of a specific causal variant underlying natural phenotypic variation have been confirmed by functional perturbation in the evolutionary model system *M. guttatus*, and the first successful instance of gene editing in the genus *Mimulus* as a whole.

Computer simulations based on simple activator-inhibitor models [12, 17, 18, 20] provided further support that *NEGAN* and *RTO* correspond to an activator-inhibitor pair in a two-component RD system. Gradual increase of the activator degradation rate (U_A), inhibitor degradation rate (U_I), and background inhibitor production rate (I_0) in simulations can recapitulate patterns generated by *NEGAN* RNAi, *RTO* RNAi, and *RTO* overexpression, respectively (Figures 6 and S7B). Furthermore, the simulation results demonstrated that the seemingly surprising observation that *RTO* transcript level did not decrease in the *RTO* RNAi lines (Figures 3E and 3F) is actually a predictable outcome of the activator-inhibitor reaction kinetics (Figure S7A). These mathematical models will serve as an important guide for future studies that aim to experimentally determine the relative diffusion rates, molecular half-lives, and potency of activation and inhibition of this protein pair. By extending the tools for studying gene and protein function we have newly applied in *Mimulus* and adopting additional technologies (e.g., light-sheet fluorescence microscopy), we anticipate that these difficult tasks will be achievable.

Although our results strongly suggest that the RD mechanism plays a critical role in the formation of dispersed anthocyanin spots in *Mimulus*, we cannot rule out the possibility that other mechanisms may also contribute to the pattern formation in this system. An alternative mechanism often invoked to explain spatial patterning during development is the “positional information” (PI) model [14, 57–59]. In contrast to the self-organizing nature of the RD model, the PI model involves fine orchestration of a hierarchical genetic cascade during development that leads to gradual refinement of discrete boundaries within tissues. The source of the positional information is often associated with anatomical landmarks (e.g., veins of butterfly wings or

flower petals). Whereas historically the PI model and RD model were usually viewed as mutually exclusive [14, 58], recent studies suggest that these two mechanisms may act together in pattern formation [25, 27, 57]. The restriction of spot formation and *NEGAN* expression to the ventral (lower) petal in *Mimulus* indicates the existence of upstream positional information that sets the initial conditions for the activator. Future experiments seeking to identify upstream regulators of *NEGAN* will be necessary to elucidate the putative positional information.

We have shown that an array of anthocyanin spot patterns can be generated by modulating the interaction of a local activator and a lateral inhibitor through experimental manipulations or natural mutations, and altering the spot patterns can impact bumblebee visitation. Thus, subtle changes in this simple interacting network are likely essential contributors to the remarkable diversification of pigmentation patterns in nature, particularly when overlaid on different positional information that imposes unique initial conditions or boundary conditions for the formation of the self-organized patterns. For instance, many *Mimulus* and other angiosperm species display spotted, blotched, or striped anthocyanin patterns in their nectar guides that may influence interactions with pollinator communities, potentially facilitating reproductive isolation and speciation [7, 32, 33, 60]. A similar R2R3- and R3-MYB pair has been implicated in the periodic patterning of leaf trichomes and root hairs in *Arabidopsis* [23, 61–63], although the R2R3-MYB in that case does not self-activate. The evolution of novel R3-MYB repressors through duplication and then degeneration of R2R3-MYB sequences also has been observed [64]. Thus, the origin of two-component RD systems involving an R3-MYB inhibitor that is evolutionarily derived from the R2R3-MYB activator may be a particularly common route to the evolution of pigmentation patterns in flowering plants. We expect that molecular circuits with similar properties do explain various pigmentation patterns in other systems as first envisioned by Turing [16] and Gierer and Meinhardt [17] several decades ago.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at <https://doi.org/10.1016/j.cub.2019.12.067>.

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AUTHOR CONTRIBUTIONS

Y.-W.Y., B.K.B., and H.D.B. conceptualized and guided the project. B.D., J.L., L.E.S., F.P., H.D.B., and Y.-W.Y. conducted the experiments in *M. lewisii*. E.L.P., S.V.H., G.A.J., A.B.G., and B.K.B. conducted the experiments in *M. guttatus*. B.D., E.L.P., S.V.H., B.K.B., F.P., and Y.-W.Y. analyzed the data. M.L.B. performed mathematical modeling and computer simulations. B.D., B.K.B., and Y.-W.Y. wrote the manuscript with input from all authors.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Bacterial and Virus Strains		
<i>Agrobacterium tumefaciens</i> strains EHA105 and GV3101	Widely distributed	EHA105, GV3101
Biological Samples		
SWC wt x SWC <i>rto</i> -like <i>M. guttatus</i> F ₂ seed	This Paper	N/A
LRD wt x LRD <i>rto</i> -like <i>M. guttatus</i> F ₂ seed	This Paper	N/A
<i>rto</i> x SL9 <i>M. lewisii</i> F ₂ seed	This Paper	N/A
Chemicals, Peptides, and Recombinant Proteins		
1-naphthalene acetic acid	PhytoTechnology Labs	Cat#N5999
2-N-morpholino ethane sulfonic acid	Sigma-Aldrich	Cat#M2933
Acetosyringone (4'-Hydroxy-3',5'-dimethoxyacetophenone)	Sigma-Aldrich	Cat#D134406
Calcium-D-gluconate	Acros Organics	Cat#299-28-5
Cefotaxime	GoldBio	Cat#C-104-25
Cetyl Trimethyl Ammonium Bromide (CTAB)	Sigma-Aldrich	Cat#H5882
Gelrite (Gellan gum)	Scotts Labs	Cat#4900-1890
Kanamycin	GoldBio	Cat#K-120-10
Meta-topolin	PhytoTechnology Labs	Cat#T-841
Murashige and Skoog salts with Gamborg B5 vitamins	PhytoTechnology Labs	Cat#M404
N-(2-Chloro-4-pyridyl)-N'-phenylurea (CPPU)	PhytoTechnologyLabs	Cat#C279
Phosphinothricin	GoldBio	Cat#P-165-250
Rifampicin	GoldBio	Cat#R-120-1
Timentin Ticarcillin / Clavulanate	GoldBio	Cat#T-104-250
<i>S. pyogenes</i> Cas9 nuclease	New England Biolabs	Cat#M0386S
Critical Commercial Assays		
AMPure XP beads	Beckman Coulter	Cat#A63880
Quant-iT dsDNA Broad Range Assay Kit	ThermoFisher	Cat#Q33130
NEBNext Ultra II DNA Library Prep Kit for Illumina	New England Biolabs	Cat#E7645S
Multiplex Oligos for Illumina Kit	New England Biolabs	Cat#E7335S
Phusion High Fidelity PCR Kit	New England Biolabs	Cat#E0553S
Spectrum Plant Total RNA Kit	Sigma-Aldrich	Cat#STRN50-1KT
Zymo Direct-Zol RNA kit	Zymo Research	Cat#R2055
SuperScript III First-Strand Synthesis System	Invitrogen	Cat#18080051
DNase I	Invitrogen	Cat#18068015
DNase I Set	Zymo Research	Cat#E1010
GoScript Reverse Transcriptase System	Promega	Cat#A5000
PowerUp SYBR Green Master Mix	Applied Biosystems	Cat#A25741
iQ SYBR Green Supermix	Bio-Rad	Cat#1708880
Deposited Data		
Genomic sequence reads generated for bulked segregant analyses	NCBI Sequence Read Archive	SRA: BioProject PRJNA481753
<i>Mimulus guttatus</i> v2.0 reference genome	Phytozome	https://phytozome.jgi.doe.gov/pz/portal.html#info?alias=Org_Mguttatus

(Continued on next page)

Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
Experimental Models: Organisms/Strains		
<i>Mimulus lewisii</i>	Inbred line LF10	LF10
<i>Mimulus guttatus</i>	Sweet Creek Road (OR), Littlerock Dam (CA), Trinity Alps Road (CA), Quarry Valley Road (CA), Mackville Road (CA)	SWC, LRD, TAR, QVR, MAC
<i>Nicotiana benthamiana</i>	Widely distributed	N/A
<i>Bombus impatiens</i>	Koppert Biological Systems	Naturpol
Oligonucleotides		
Primers for cloning, sequencing, genotyping, qPCR	This paper (Tables S4–S6)	N/A
Recombinant DNA		
pFGC5941	<i>Arabidopsis</i> Biological Resource Center	CD3-447
pMCS:GW	<i>Arabidopsis</i> Biological Resource Center	CD3-1933
<i>ER-cb</i>	<i>Arabidopsis</i> Biological Resource Center	CD3-954
pMCS:YFP-GW	<i>Arabidopsis</i> Biological Resource Center	CD3-1934
pEarleyGate 100	<i>Arabidopsis</i> Biological Resource Center	CD3-724
pEarleyGate 104	<i>Arabidopsis</i> Biological Resource Center	CD3-686
pUC119-gRNA	Addgene	52255
pFGC-pcoCas9	Addgene	52256
pUBC-nYFP-Dest	[65]	N/A
pUBC-cYFP-Dest	[65]	N/A
pENTR/D-TOPO	Invitrogen	Cat#K240020
AS-121P-CR3	Alex Schultink, Innovative Genomics Institute	N/A
MIRTO RNAi	This paper	N/A
MgRTO RNAi	This paper	N/A
MINEGAN RNAi	This paper	N/A
MgNEGAN RNAi	This paper	N/A
RTO sgRNA ⁺ Cas9	This paper	N/A
RTO _{pro} :CFP-ER	This paper	N/A
RTO _{pro} :YFP-RTO	This paper	N/A
35S:MIRTO	This paper	N/A
35S:YFP-MIRTO	This paper	N/A
pUBC-cYFP-ANbHLH1	This paper	N/A
pUBC-nYFP-RTO	This paper	N/A
pUBC-nYFP-rto	This paper	N/A
Software and Algorithms		
bwa	[66]	https://github.com/lh3/bwa
samtools	[67]	https://github.com/samtools
ImageJ	[68]	https://imagej.nih.gov/ij/
VCell	[48]	http://vcell.org
RDsimJ.jar	[12]	https://science.sciencemag.org/content/suppl/2010/09/22/329.5999.1616.DC1
JMP Pro 13	SAS Institute Inc.	https://www.jmp.com/
E-CRISP	[69]	http://www.e-crisp.org/E-CRISP/
Custom Scripts	This Paper	https://github.com/BlackmanLabUCB/Bulked-Segregant-Analysis

LEAD CONTACT AND MATERIALS AVAILABILITY

Further information and requests for *M. lewisii* and *M. guttatus* resources should be directed to and will be fulfilled by Yao-Wu Yuan (yaowu.yuan@uconn.edu) and Benjamin K. Blackman (bkblackman@berkeley.edu), respectively.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Plants

Mimulus lewisii plants were grown in the University of Connecticut research greenhouses under natural light supplemented with sodium vapor lamps, ensuring a 16h day length. EMS mutagenesis was performed using the *Mimulus lewisii* Pursh inbred line LF10 as previously described [70].

Mimulus guttatus individuals with fully or partially expanded nectar guide spots were observed in the selfed offspring of wild plants collected along Sweet Creek Road in Mapleton, OR (SWC; 44°00'51"N, 123°51'36"W) and along Emma Road east of Cheeseboro Road near Littlerock Dam (LRD; 34°30'23"N, 118°01'39"W). SWC plants, F₁s, F₂s, and 2011 maternal families were grown in Fafard 4P soil in 2.5 square-inch pots in the Duke University greenhouses (18h 21°C days: 6h 18°C nights). After cold stratification (7d at 4°C), plants were misted and bottom watered twice daily on flood benches for 10d and then just bottom watered for the remainder of the experiment. LRD plants, F₁s, F₂s, and the LRD and SWC 2017 maternal families were grown in Sunshine Mix #1 in 2.5 square-inch pots in the UC Berkeley Oxford Tract Facility greenhouses (16h 24°C: 8h 14°C). Seeds were cold stratified (7d at 4°C), then watered on a mist bench for two weeks as they germinated, and then moved to a standard bench and bottom-watered daily. For both mapping populations, all pots were thinned to one plant per pot. Once plants began flowering, we recorded nectar guide phenotypes and flash froze bud tissue, which was maintained at –80°C until DNA extraction for fine mapping. We also observed *rto*-like plants in the field or in families grown in the greenhouse from field-collected seed for *M. guttatus* populations at Quarry Valley Road in the Donald and Sylvia McLaughlin Natural Reserve (QVR; 38°51'37"N, 122°25'15"W) and on CA-3 at Trinity Alps Road (TAR; 40°51'09"N, 122°51'56"W).

Animals

A Natupol hive of *Bombus impatiens* was purchased from Koppert Biological Systems (Howell, MI). To conduct pollinator choice trials, we constructed a 3' x 4' x 2' flight cage underneath four Reptisun fluorescent bulbs that provided visible and UV light. We connected the hive to the flight cage with a small tube and a series of gates, which allowed us to let an individual bee into the flight cage.

METHOD DETAILS

Bulked Segregant Analysis in *Mimulus lewisii*

To identify the causal gene underlying *rto*, we performed bulked segregant analysis following Yuan et al. [37]. Briefly, the homozygous *rto* mutant (in the LF10 background) was crossed with the mapping line SL9. A single F₁ individual was then selfed to generate the F₂ mapping population (n = 468). DNA samples from 100 F₂ segregants displaying the mutant phenotype were pooled with equal representation. A whole genome sequencing library (insert size of ~320 bp) was prepared for the pooled sample, and 100-bp paired-end reads were generated by an Illumina HiSeq 2000 (Illumina, San Diego, CA, USA). The resulting ~196 million short reads were mapped to the SL9 pseudoscaffolds, and the resulting raw SNPs were filtered by depth of coverage, tendency of clustering, and variant frequency. The SL9 pseudoscaffolds were then scanned for regions enriched with homozygous SNPs to determine the candidate gene interval (Figure 2A).

Bulked Segregant Analyses in *Mimulus guttatus*

For SWC, an individual with a large anthocyanin patch was crossed to an individual with typical nectar guide patterning descending from a maternal family that bred true for that phenotype. A single F₁ individual was selfed to generate the F₂ mapping population (n = 452 individuals). Similarly, we crossed an LRD individual with expanded anthocyanin spots to a wild-type individual from another maternal line and selfed an F₁ descendant to generate an F₂ mapping population (n = 631).

For bulked segregant analysis of each mapping panel, we randomly selected 100 wild-type and 100 *rto*-like plants derived from the corresponding F₂ mapping population, and we then collected three equal-sized buds from each plant into one of four 15 mL tubes (25 plants per tube) for each phenotype. We extracted DNA using a modified CTAB protocol [71], including a final cleanup with 2M NaCl, 4% PEG solution. The resulting DNA was quantified using a Quant-iT dsDNA Broad Range Assay Kit (ThermoFisher), and 250 ng from each of the four tubes was pooled for a total of 1 µg per phenotype in 50 µL water. The pooled samples were sonicated for 65 s using a Covaris M220 in order to retrieve ~350-bp fragments. AMPure Purification beads were used to remove fragments smaller than 200 bp, and remaining fragments were end-repaired and A-tailed before adaptor ligation using the NEBNext Ultra II DNA Library Prep Kit for Illumina (New England Biolabs). For adaptors, we used the Multiplex Oligos for Illumina Kit (New England Biolabs), and, once indexed, we again size selected for 200–400-bp fragments by bead purification. The libraries were enriched using the Phusion High Fidelity PCR Kit (New England Biolabs), and bead purified once more to remove the smallest fragments. The final whole genome sequencing libraries were run on an Agilent Bioanalyzer to determine fragment size and concentration and then

combined in equimolar concentration into one sample. Samples were submitted either to the Institute for Integrative Genome Biology Genomics Core at University of California, Riverside for one lane of 100SR sequencing on an Illumina HiSeq2500 platform or to the QB3 Vincent J. Coates Genome Sequencing Laboratory at University of California, Berkeley for one lane of 100SR sequencing on an Illumina HiSeq4000 platform.

The LRD wild-type and *rto*-like pools yielded ~184 million and ~172 million reads, respectively. Because the initial sequence data obtained for the two pooled SWC libraries yielded a low number of assignable reads, we constructed new libraries from two pools of individual DNAs from 100 wild-type and from 100 *rto*-like phenotype plants. Library construction and sequencing were completed as above. Data from both SWC sequencing runs were then combined for analysis (~59 and ~76 million total reads for the wild-type and *rto*-like pools, respectively).

Fine Mapping of *rto*^{SWC} in *Mimulus guttatus*

We identified recombinants in the candidate genomic interval resulted from the bulked segregant analysis by first genotyping all individuals in the SWC F₂ population for four EPIC (exon-primed, intron-capture) markers spanning the region through fragment analysis on the University of Virginia Department of Biology Genomics Core Applied Biosystems 3130 instrument. We then developed additional microsatellite or CAPS (cleaved, amplified polymorphic sequence) markers from polymorphisms distinguishing the parental alleles using the bulked segregant re-sequencing data. These markers were selectively scored on the recombinant individuals using the ABI 3130 or by gel electrophoresis as the region containing the causal locus was iteratively refined. Finally, we confirmed that pigmentation phenotype perfectly co-segregated with the *rto*^{SWC} frameshift mutation (Figure 2D) or a 723-bp insertion in the *RTO* 3'UTR in the SWC and LRD panels, respectively, by gel electrophoresis. Primer sequences for all markers are listed in Table S4.

Plasmid Construction

RNAi plasmids

We built *RTO* RNAi constructs by cloning fragments of the *M. lewisii* or *M. guttatus* *RTO* gene into the pFGC5941 vector ([72]; ABRC stock no. CD3-447) in both the sense and antisense directions separated by an intronic spacer. This fragment, and every 12-bp block within it, matched only a single region of the *M. lewisii* or *M. guttatus* gene (100% identity), indicating target specificity. Likewise, we built *MgNEGAN* RNAi constructs following the same procedure.

CRISPR-Cas9 plasmid

We also undertook a targeted genome editing approach using the Cas9-sgRNA system to recapitulate the *RTO* loss of function phenotype in *M. guttatus*. The synthetic guide-RNA (sgRNA) comprised an optimized *trans*-activating CRISPR RNA (tracrRNA) and a CRISPR RNA (crRNA) that contains a 20-bp guide proximal to an NGG-site recognized by *S. pyogenes* Cas9. Several sgRNA guides were designed *in silico* (<http://www.e-crisp.org/E-CRISP/>) to target an exonic region of *MgRTO*. The 20-bp guide sequences were verified for its specificity by using a blast algorithm to scan the *M. guttatus* v2.0 reference genome (<https://phytozome.jgi.doe.gov>). The guides were pre-tested for their cleavage efficiency with an *in vitro* assay system using *S. pyogenes* Cas9 nuclease (New England Biolabs) to edit a PCR-amplified double-stranded DNA fragment of *RTO*. A binary plasmid construct expressing plant-codon-optimized Cas9 (pFGC-pcoCas9; Addgene#52256) was modified to express a *RTO* sgRNA (TTTCGAATGCACAAGCTCGT) that was successful *in vitro*. We synthesized the cassette of AtU6-polymerase III promoter driving sgRNA and poly-T terminator sequences by PCR. The AtU6-polymerase promoter was amplified from a plasmid pUC119-gRNA (Addgene: 52255). The tracrRNA and crRNA sequence with the poly-T terminator was amplified from plasmid AS-121P-CR3 (provided by A. Schultink, Integrative Genomic Institute, UC Berkeley). The cassette (AtU6p:sgRNA:terminator) for the expression of sgRNA was inserted into plasmid pFGC-pcoCas9 using *AscI* and *Pacl* restriction sites. Insertion of the sgRNA cassette was confirmed by Sanger sequencing.

*RTO*_{pro}:CFP-ER and *RTO*_{pro}:YFP-*RTO* plasmids

To monitor the spatial expression pattern of *RTO* in the wild-type *M. lewisii* (LF10), we constructed a plasmid with the *RTO* promoter (a ~1.7-kb DNA fragment upstream of the translation initiation site) driving CFP with an ER retention signal peptide. To this end, we first cloned the promoter fragment into the *pMCS:GW* vector ([73]; ABRC stock no. CD3-1933) using the *AatII* and *XhoI* restriction enzyme sites and obtained an intermediate vector *RTO*_{pro}:*GW*. We then amplified the *CFP-ER* fragment from the vector *ER-cb* ([74]; ABRC stock no. CD3-954) and recombined it into the intermediate vector by LR recombination, resulting in the final plasmid *RTO*_{pro}:*CFP-ER*. To monitor the *RTO* protein distribution, we constructed the *RTO*_{pro}:*YFP-RTO* plasmid using a similar approach. We first cloned the same promoter fragment into *pMCS:YFP-GW* (ABRC stock no. CD3-1934) and then recombined the *RTO* CDS into the intermediate vector through LR recombination, resulting in the final *RTO*_{pro}:*YFP-RTO* plasmid.

35S:*RTO* and 35S:*YFP-RTO* plasmids

To generate the 35S:*RTO* and 35S:*YFP-RTO* plasmids, we first amplified the full-length CDS of *RTO* from the wild-type *M. lewisii* (LF10) cDNA using the Phusion enzyme (NEB, Ipswich, MA). The amplified fragment was cloned into the pENTR/D-TOPO vector (Invitrogen), and then a linear fragment containing the CDS flanked by the *attL1* and *attL2* sites was amplified using M13 primers. This linear fragment was subsequently recombined into the Gateway vector pEarleyGate 100 ([75]; ABRC stock no. CD3-724) or pEarleyGate 104 (CD3-686), to generate the final 35S:*RTO* and 35S:*YFP-RTO* plasmids, respectively.

BiFC plasmids

To test the interaction between the wild-type *RTO* or mutant *rto* protein with ANbHLH1, bimolecular fluorescence complementation (BiFC) plasmids were constructed by recombining the *ANbHLH1* CDS into the destination vector pUBC-nYFP-Dest and the *RTO*

or *rto* CDS into pUBC-cYFP-Dest [65], following the same procedure as described above for constructing the 35S:*RTO* plasmid. Primers used for plasmid constructions are listed in Table S5.

Plant Transformation

Mimulus lewisii transformation followed Yuan et al. [42]. For *M. guttatus*, we introduced the *RTO* and *NEGAN* RNAi constructs and the *RTO* targeting Cas9-sgRNA construct into the plant through *Agrobacterium* mediated transformation of callus tissue in tissue culture. To this end, we first transformed the RNAi or Cas9-sgRNA construct into *Agrobacterium tumefaciens* strain EHA105. Overnight cultures in yeast extract peptone medium supplemented with kanamycin (50 mg/L) and rifampicin (40 mg/L) were resuspended to OD600 = 0.2 with half-strength MS salts supplemented with acetosyringone (200 μ M). The resuspended cultures were shaken at 50 rpm in darkness for 4h for virulence induction.

All plant cultures were grown on MS growth medium (Murashige and Skoog salts with Gamborg B5 vitamins, 2% sucrose, 1.3 g/L calcium gluconate, 0.25 g/L 2-N-morpholino ethane sulfonic acid, and 0.25% gelrite), pH 5.6. Plant cultures were incubated under fluorescent lights (200 μ mol/m²/s) on 14h light: 10h dark photoperiod, at 21 \pm 1°C during transformation and tissue culture. We excised fully expanded leaves with petioles intact from *M. guttatus* plants, grown *in vitro* from seeds from inbred lines of the MAC population (Mackville Road, 1-2 mi N of Liberty Road, Sacramento County, CA; 38°14'56"N, 121°05'26"W). Petioles were dipped in the *Agrobacterium* cultures and co-cultivated on MS growth medium supplemented with 1 mg/L N-(2-Chloro-4-pyridyl)-N'-phenylurea (CPPU) and 100 μ M acetosyringone. After 3d of co-cultivation in darkness, the explants were washed with 100 mg/L timentin and 50 mg/L cefotaxime and maintained on the callus induction MS growth medium supplemented with 1 mg/L CPPU, antibiotics, and 6 mg/L phosphinothricin (PPT). We sub-cultured callus tissue that survived selection for 25-28 days, and then transferred callus exhibiting shoot organogenesis to shoot induction MS growth medium supplemented with 0.1 mg/L meta-topolin. We further sub-cultured shoots > 1 cm long on half-strength MS growth medium supplemented with 0.1 mg/L 1-naphthalene acetic acid and PPT. Rooted plantlets were transplanted to pots filled with soil-less potting medium and hardened on a mist-bench in the green house for four weeks. We confirmed *RTO* and *NEGAN* RNAi lines by genotyping and RT-PCR in at least 10 T₀ plants per construct and then selfed these individuals to generate T₁ full-sib families for phenotyping and expression analyses. For the *RTO* targeting Cas9-sgRNA experiments, transgenic T₀ and T₁ plants showing phenotypes resembling *RTO/rto* heterozygotes were genotyped by Sanger sequencing to verify the mutations in sgRNA target region.

Expression Analysis by RT-qPCR

For *M. lewisii*, total RNA was isolated from whole corolla or dissected petal lobes and nectar guides of 10 mm flower buds (unless noted otherwise) using the Spectrum Plant Total RNA Kit (Sigma-Aldrich). For each sample we treated 1 μ g of total RNA with amplification grade DNaseI (Invitrogen) before cDNA synthesis using the SuperScript III First-Strand Synthesis System (Invitrogen). cDNA samples were diluted 10-fold before quantitative reverse transcriptase PCR (qRT-PCR). All qRT-PCRs were performed using iQ SYBR Green Supermix (Bio-Rad) in a CFX96 Touch Real-Time PCR Detection System (Bio-Rad). Samples were amplified for 40 cycles of 95°C for 15 s and 60°C for 30 s. Amplification efficiencies for each primer pair (Table S6) were determined using critical threshold values obtained from a dilution series (1:4, 1:8, 1:16, 1:32) of pooled cDNA. *MIUBC* was used as a reference gene to normalize expression levels following the delta-delta Ct method.

For *M. guttatus*, lower lobes of buds measuring approximately 2-3mm (SWC and MAC populations) and 4-5mm (LRD population) were harvested under magnification lenses (Figure S4E). Pools of tissue from least 12-14 buds per each of four biological replicates were frozen in liquid nitrogen and stored in -80°C until processing for RNA extraction. Total RNA was extracted using the Zymo Research Direct-Zol RNA kit and DNase treated with the Zymo Research DNase I Set following the manufacturer protocols. First strand cDNA was synthesized using 500ng of RNA with oligo d(T)15 primer using GoScript™ Reverse Transcriptase System manufacturer kit (Promega Corp. Madison, WI, USA) and diluted 4-fold before qPCR. Three technical replicates per qPCR amplification were conducted in 10 μ L reactions using PowerUp™ SYBR Green Master Mix (Applied Biosystems). All reactions were run on a 7500 Fast Real-Time PCR System (Applied Biosystems) for 40 cycles of 95°C for 3 s and 60°C for 30 s with initial UDG activation at 50°C for 2min and polymerase activation at 95°C for 2 min. Amplification efficiencies for each primer pair (Table S6) were determined using critical threshold values obtained from a dilution series (1, 1:2, 1:4, 1:16, 1:64, 1:256) of pooled cDNA. The specificity of each primer pair for its target gene was verified visually by the melt curves obtained with default instrument settings and by Sanger sequencing. The mean Ct values for the three technical replicates per sample-target combination were used to calculate relative expression values. *MgUBQ* was used as a reference gene to normalize expression levels following the delta-delta Ct method.

BiFC Assay

We transformed the *pUBC:ANbHLH1-nYFP*, *pUBC:RTO-cYFP*, and *pUBC:rto-cYFP* plasmids separately into *Agrobacterium* strain GV3101. Equal volumes of *pUBC:ANbHLH1-nYFP* and *pUBC:RTO-cYFP* cultures or *pUBC:ANbHLH1-nYFP* and *pUBC:rto-cYFP* cultures were infiltrated into the abaxial side of *Nicotiana benthamiana* leaves, following the procedure described previously [76]. The YFP signal was imaged 4-6 days after agroinfiltration. To ensure that the constructs are properly expressed in the infiltrated tobacco leaf tissue, we excised leaf tissue from half of the infiltrated area and stored it in liquid nitrogen until RNA extraction, and we excised the other half for fluorescence imaging. Relative expression levels of *ANbHLH1* and *RTO* were assayed with semiquantitative RT-PCR. The *Nicotiana benthamiana* endogenous *ACTIN* gene was used as the reference gene.

Fluorescence Imaging

For the $RTO_{pro}:CFP-ER$ lines, ventral petals (dissected from 10 mm corolla) were imaged using a Zeiss Axioskop2 microscope with a 5X objective (FLUAR 5X/0.25) under the bright-field for anthocyanin spots (Figure 5A, left) and green channel for CFP signal (Figure 5A, center).

For the F_1 hybrids with both $RTO_{pro}:CFP-ER$ and $RTO_{pro}:YFP-RTO$ transgenes, a NIKON A1R confocal laser scanning microscope with a 10X objective was used to image CFP and YFP signals with the excitation wavelength of 457 nm and 514 nm, respectively (Figures 5B, S6D, and S6E).

For the BiFC assays, a NIKON A1R confocal laser scanning microscope equipped with a 20X objective (S Plan Fluor ELWD 20X) was used to acquire the images (Figures 3H and S6A) under the transmitted light and the green channel (excitation wavelength 514 nm).

To track the intracellular movement of the YFP-RTO fusion proteins in the $35S:YFP-RTO$ lines, YFP-RTO fluorescence signals were captured automatically every 30 s within a time span of 15 min using the same confocal laser scanning microscope with the green channel (Figure 5C; Video S1).

UV Photography

We imaged UV-absorbent nectar guides using a Canon T2i Rebel camera converted by LifePixel (Mukilteo, WA) to replace the internal hot mirror filter with a high quality custom manufactured UV band pass filter.

Scanning Electron Microscopy

We dissected fully opened *M. guttatus* flowers to retain only the ventral petal, fixed tissue with glutaraldehyde, and dehydrated the fixed samples through a graded ethanol series. Following dehydration, the samples were further trimmed to retain the pigmented area of the petal to fit the mounting space, and imaging was completed using the Zeiss Sigma VP HD Field SEM instrument at the University of Virginia School of Medicine Advance Microscope Facility.

Mathematical Modeling

The co-localization of RTO (inhibitor) promoter activity with the anthocyanin spots (Figure 5A) suggests that the activator and the inhibitor have expression patterns that are in-phase, consistent with one of the two simplest Gierer-Meinhardt topologies — the two-component activator-inhibitor model [17, 18]. Therefore, we implemented a modified Gierer-Meinhardt model (based on Equation 2.1 in [18]) between an autocatalytic activator A (NEGAN) and a rapidly diffusing inhibitor I (RTO) (Figure 6A). A diffuses with a diffusion coefficient D_A , degrades with a rate U_A . A has a background production rate A_0 and is self-activated with a potency G_A . The activator production is slowed down by the inhibitor I . A residual value ($k = 0.001$) was added to the denominator to avoid the situation where the denominator becomes zero.

$$\frac{dA}{dt} = D_A \cdot \nabla^2 A + G_A \frac{A^2 + A_0}{I + k} - U_A \cdot A$$

Similarly, I diffuses with a diffusion coefficient D_I , degrades with a rate U_I , has a background production rate I_0 and is activated by the activator with a potency G_I .

$$\frac{dI}{dt} = D_I \cdot \nabla^2 I + G_I \cdot A^2 + I_0 - U_I \cdot I$$

Computer Simulations

We implemented the modified Gierer-Meinhardt model in the Virtual Cell modeling and simulation platform [48, 77] and performed a number of two-dimensional numerical simulations to explore the pattern formation for different transgenic lines. This required integration of the partial differential equations described above. We used fully implicit, adaptive time step finite volume method. The finite volume method represents partial differential equations as algebraic discretization equations which exactly preserves conservation laws. Values are calculated at discrete places on a meshed geometry. This method employs Sundials stiff solver CVODE for time stepping. 2D discretization is the only source of solution error. According to this error, the time-step is resized during the simulation. All simulations were conducted with a domain size of 100×100 , a spatial step size of 1.0, an activator initiation concentration of 1.0 randomly distributed across the whole domain, an inhibitor initiation concentration of 0.0, and a zero flux boundary condition.

For simulating patterns resembling the wild-type *M. lewisii*, we used the following parameter values: $D_A = 0.01$, $D_I = 0.5$, $U_A = 0.03$, $U_I = 0.03$, $G_A = 0.08$, $G_I = 0.12$, $A_0 = 0.01$, and $I_0 = 0$ (Figure 6B). These parameter values were largely adapted from previous studies employing similar two-component activator-inhibitor models [12, 20, 49, 50], and were considered biologically realistic. We then investigated additional cases corresponding to three different types of transgenic manipulations. All parameter values used in these cases were the same as those in the simulation for the wild-type pattern, except one modification for each perturbation, as follows:

- (i) *NEGAN* RNAi, with gradually increased activator degradation rate ($U_A = 0.06, 0.09, 0.11, 0.12$, and 0.15 ; Figures 6C and 6I–6L).
- (ii) *RTO* overexpression, with gradually increased background production rate for the inhibitor ($I_0 = 0.01, 0.05, 0.07, 0.08$, and 0.1 ; Figures 6D and 6M–6P).

(iii) *RTO* RNAi, with gradually increased inhibitor degradation rate ($U_A = 0.06, 0.15, 0.18, \text{ and } 0.19$; Figures 6E–6H).

We used the activator level (A) as the pattern readout [12, 20, 49, 50], and set a threshold value for pattern to appear, with the assumption that the activator should be above a certain threshold level for pigment production. All patterns shown in Figure 6 are based on the threshold value 3.0.

The Virtual Cell model is available in the VCell software (<http://vcell.org>). After logging in as a guest, search for “Monkeyflower_pigmentation_v2” in VCellDB - > BioModels panel in the left bottom part of the VCell interface. Double click on the model name in the bottom left panel (under Public BioModels) to launch it. Simulation results are available by double clicking on “Pattern_formation” under Applications in the left column and selection of the Simulations tab. To retrieve the simulation results, select the name of a simulation (e.g., WT, *RTO*_RNAi_0.06), and click the “Native Simulation Results” icon. To view patterns, select “Pattern_threshold_3.0” under “Variables” on the left, and move time slider to see pattern development over time. Similarly, one can see individual variables (e.g., A, I) changes over time. To match simulation results with a pattern visible in plants, we exported VCell images in monochrome Nearly Raw Raster Data (*.nrrd) format into ImageJ, and applied LookUp Table with initial entry yellow (255,255,0) and final entry red (255,0,0).

Besides the VCell simulations, we also performed simulations with another commonly used two-component, activator-inhibitor model ([20]; Figure S7B), as implemented in the java program RDSimJ.jar [12]. The parameter values used to simulate the wild-type (WT) pattern are shown in the screenshot of the graphic interface of RDSimJ.jar (Figure S7). The patterns corresponding to the three types of transgenic manipulations (i.e., *NEGAN* RNAi, *RTO* overexpression, and *RTO* RNAi) are shown in Figure S7B, with the single altered parameter value (compared to the WT values) shown below each simulated pattern.

Allele Frequency Surveys

Mature fruits were collected from 51 maternal plants at the LRD population in May 2017, from 10 maternal plants at the SWC population in June 2011, and from 48 maternal plants at the SWC population in June 2017. Three to seventeen (2011 survey) or sixteen (both 2017 surveys) progeny from each maternal family were then raised in the Duke University greenhouses or the UC Berkeley Oxford Tract Facility as described above. Heterozygosity for the premature stop mutation was confirmed for individuals with nectar guide phenotypes resembling the *RTO*^{LRD}/*rto*^{LRD} heterozygous phenotype (Figure 1) by amplifying, cloning, and Sanger sequencing the PCR products generated with the *Migut.B01849* fine mapping primers (Table S4). The QVR population (Quarry Valley Road, Donald and Sylvia McLaughlin Natural Reserve) was phenotypically surveyed in the field. Counts of wild-type, intermediate, and *rto*-like plants were tallied in 8 patches along a transect through the population (25 randomly selected plants per patch, $n = 200$ individuals total). We confirmed by Sanger sequencing that *rto*-like plants ($n = 4$) carry a QVR-specific 8-bp deletion in *RTO* not observed in the wild-type.

Pollinator Preference Experiments

To determine whether bumblebees preferred flowers of *rto*^{SWC} variant carriers to flowers with wild-type nectar guides, we conducted pollinator cage experiments with a commercially obtained hive of *Bombus impatiens*. Before adding plants to the cage, we counted the total number of open flowers on each plant. We then placed six plants from the SWC F₂ mapping population – three of one *RTO* genotype and three of another *RTO* genotype – in a regularly-spaced checkerboard pattern based on phenotype such that each plant’s direct neighbors were of a different genotype. We conducted 10 independent trials for each pairwise combination of *RTO*^{SWC} genotypes.

For each trial, we allowed one bee to enter and forage for 20 minutes. Any bee that did not start foraging within five minutes was removed. To ensure all bees were naive, any bee which entered the cage, whether or not they successfully foraged, was marked upon removal and returned to the hive. We did not collect any further foraging data from marked bees. If a bee did start foraging, we recorded each visit with the plant that was visited, the time of landing on a flower, the orientation of the visit, and the time of departure. After each successful trial, we replaced the plants. The effect of *RTO* genotype on pollinator visits was analyzed using G-tests of goodness of fit [78], conducting separate analyses for each pairwise combination of *RTO*^{SWC} genotypes. Expected visit values were calculated based on the proportion of open flowers of each genotype included in the pollinator cage for each trial.

QUANTIFICATION AND STATISTICAL ANALYSIS

Quantification of Anthocyanin Spots

We quantified the anthocyanin spots (number of spots and relative area in the nectar guides) for both species, including the various mutants and transgenic lines (Figure S2). For *M. lewisii*, the ventral (lower) petals were scanned using an Epson WorkForce Pro WP-4530 printer. The spots were manually counted for 10 flowers per line from the scanned images with ImageJ. To quantify the relative area of anthocyanin spots in the nectar guides (i.e., the ratio of the areas pigmented by anthocyanin to the total area of the yellow background), we analyzed five scanned flower images for each line. A customized Python script with OpenCV tools was used to perform the analysis. Briefly, we first used the “cv2.findContours” function to isolate nectar guide area from the rest of flower petal images based on color contrast. Then we used “cv2.findContours” again with different argument settings to separate red spots from the yellow background. We also calculated nectar guide area and spots area with the “cv2.contourArea” function.

For *M. guttatus*, the ventral petals were stretched out to remove the creases, and a transparent tape was used to fix the petal on a sheet of white color paper. Color photographs of the petal were taken along with a ruler to calibrate area estimates. The calibrating ruler was placed close to the petal to minimize errors caused by the distortion of the camera lens. A Canon Rebel T2i digital single-lens reflex (DSLR) camera mounted on a tripod was used for photography. The pictures were imported to ImageJ2, and the scale of the image was set using the metrics on the ruler to obtain area estimates in square centimeters. The free hand tool was used to trace along the petal to measure the area of the whole petal, and individual spots were outlined in a similar approach to measure the area covered by the spots. The ratio of the total area of spots to the whole ventral petal area was calculated. The number of anthocyanin spots was manually counted from each picture. Ten to thirteen plants per genotype were sampled.

Bulked Segregant Analyses for *M. guttatus*

For each mapping population, sequencing data for each pool were aligned to the *M. guttatus* v2.0 reference genome (<https://phytozome.jgi.doe.gov>) with the `aln` and `samse` commands in BWA [66] and converted to sorted BAM files with `samtools` [67]. A merged BAM file was then converted to `mpileup` format, and a list of SNPs was called using a custom script that removed indel variants, sites covered by < 4 reads or covered by > 200 reads, sites that segregated for > 2 alleles, and sites where one or both alleles were supported by < 2 reads with Illumina base quality scores < 4. Pileup files were then generated from the sorted BAM files for each pool, and these were then parsed with a custom script and the SNP list to yield a table listing the allele counts and frequencies for each bulk and the allele frequency difference between the two bulks at each SNP site. SNP sites were filtered for those where each bulk had > 10x read coverage and the total read coverage was $\leq 160x$. For the SWC panel, the proportion of SNPs with fixed differences between the two bulk pools was calculated for 50-kb intervals (median = 94 SNPs/interval) in JMP Pro 13 (SAS Institute Inc.). For the LRD panel, the proportion of SNPs in the top 0.1% allele frequency difference (> 0.66) was calculated for 50-kb intervals (median = 60 SNPs/interval). Different metrics were calculated and plotted for each panel due to the difficulty of visually distinguishing the LRD panel heterozygote phenotype from the LRD wild-type homozygote phenotype. As a consequence of this ambiguity, heterozygous individuals were included in the wild-type pattern LRD bulked pool at low frequency, leading to a lack of fixed differences overall between the wild-type and *rto-like* pools.

DATA AND CODE AVAILABILITY

The accession number for the sequence data reported in this paper is NCBI SRA: PRJNA481753. Code utilized in this paper can be found on GitHub: <https://github.com/BlackmanLabUCB/Bulked-Segregant-Analysis>. Many protocols used in this study are available on protocols.io: <https://www.protocols.io/groups/mimulus/publications>.