

A novel R3 MYB transcriptional repressor associated with the loss of floral pigmentation in *Lochroma*

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Summary

- Losses of floral pigmentation represent one of the most common evolutionary transitions in flower color, yet the genetic basis for these changes has been elucidated in only a handful of cases.
- Here we used crossing studies, bulk-segregant RNA sequencing, phylogenetic analyses and functional tests to identify the gene(s) responsible for the transition to white flowers in *Lochroma loxense*.
- Crosses between *L. loxense* and its blue-flowered sister species, *L. cyaneum*, suggested that a single locus controls the flower color difference and that the white allele causes a nearly complete loss of pigmentation. Examining sequence variation across phenotypic pools from the crosses, we found that alleles at a novel R3 MYB transcription factor were tightly associated with flower color variation. This gene, which we term *MYBL1*, falls into a class of MYB transcriptional repressors and, accordingly, higher expression of this gene is associated with downregulation of multiple anthocyanin pigment pathway genes. We confirmed the repressive function of *MYBL1* through stable transformation of *Nicotiana*.
- The mechanism underlying the evolution of white flowers in *L. loxense* differs from that uncovered in previous studies, pointing to multiple mechanisms for achieving fixed transitions in flower color intensity.

Introduction

From a genetic and biochemical perspective, the biosynthesis of anthocyanin pigment represents one of the best studied metabolic pathways in plants. The red, purple and blue anthocyanins are the products of a series of enzymatic steps that are highly conserved across plants, and they constitute the most common pigments responsible for flower and fruit coloration (Winkel-Shirley, 2001; Grotewold, 2006). The regulation of anthocyanin production varies in different taxa, but members of the WD-repeat (WDR), basic helix–loop–helix (bHLH) and MYB families are commonly important regulators of this pathway (Feller *et al.*, 2011; Davies *et al.*, 2012). Often, these transcription factors form a complex (the MYB–bHLH–WDR or MBW complex) that can coordinately activate or repress multiple steps in the pathway to modulate pigment production (Albert *et al.*, 2014).

This deep understanding of anthocyanin pathway function and regulation has made it possible to identify the genetic basis of pigment variation in many nonmodel plants. These studies have begun to reveal predictable patterns in pigment evolution (Streisfeld & Rausher, 2011; Sobel & Streisfeld, 2013). For example, evolutionary transitions to white flowers via losses of floral anthocyanin production have consistently involved

loss-of-function mutations in R2R3 MYB transcriptional activators (Quattrocchio *et al.*, 1999; Schwinn *et al.*, 2006; Hoballah *et al.*, 2007). Given that a wide range of pathway mutations can give rise to white flowers (de Vlaming *et al.*, 1984; van Houwelingen *et al.*, 1998), this pattern has been attributed to preferential fixation of these R2R3 MYB mutations (Streisfeld & Rausher, 2011). R2R3 MYB transcription factors are among the largest gene families in plants, and each member is highly specialized in terms of spatial expression, timing of expression and target genes (Kranz *et al.*, 1998; Ramsay & Glover, 2005). Thus, mutations in R2R3 MYBs have the potential to change anthocyanin gene expression only in the flower, with minimal pleiotropic consequences for the production of anthocyanins or related flavonoids in other tissues (Wessinger & Rausher, 2012). Nonetheless, this apparent preferential fixation is based on studies of floral pigment loss in two genera (*Antirrhinum* and *Petunia*) and remains to be validated more broadly.

Here we examine the genetic basis for the transition to white flowers in the Andean genus *Lochroma* (Solanaceae). *Lochroma* is a relatively small genus of *c.* 25 species, but presents a wide range of flower colors, including red, blue, green, white, yellow and orange (Smith & Baum, 2006). The ancestral state in the genus is blue flowers, which derive their color from delphinidin-based

anthocyanins (Smith & Baum, 2007; Berardi *et al.*, 2016). From this ancestral state, the clade has experienced multiple transitions, leading to a variety of derived color phenotypes, including at least three to white or yellow flowers due to losses of floral anthocyanins (Smith & Goldberg, 2015). The present study focuses on the most recent of these transitions, along the branch leading to the white-flowered *I. loxense*. Its sister species, *I. cyaneum*, is a horticulturally important plant in which the anthocyanin pathway and anthocyanin production have been previously characterized (Smith & Rausher, 2011; Berardi *et al.*, 2016). Taking advantage of the crossability of these two taxa, segregating backcross populations were created to identify genes associated with flower color variation. Given the large number of potential candidate genes, including the many R2R3 MYBs, we adopted a bulk-segregant RNA-Seq approach to identify potentially causative loci and determine their effects on pathway gene expression. The results implicate a novel class of R3 MYB transcriptional repressors, which appear to influence flower color via the same developmental changes as the previously identified R2R3 loss-of-function mutations.

Materials and Methods

Crossing and phenotyping

We created a cross between the blue-flowered *I. cyaneum* and the white-flowered *I. loxense* using two cultivated accessions. The parental *I. cyaneum* individual was derived from an accession cultivated at the Missouri Botanical Garden by W. G. D'Arcy, and this accession was used to generate a low-coverage genome in previous work (Gates *et al.*, 2016). The parental *I. loxense* individual was derived from material from the University of Loja Botanical Gardens near Loja, Ecuador. Voucher specimens for each accession (Smith 265 and 235, respectively) are deposited at the University of Wisconsin–Madison Herbarium and the Missouri Botanical Garden. Previous biochemical studies confirm that *I. cyaneum* produces over 97% blue delphinidin-based anthocyanin pigments, while *I. loxense* produces no detectable floral anthocyanins (Berardi *et al.*, 2016). A single F₁ individual was backcrossed to each parental accession to create segregating backcross populations. Backcross individuals were grown in glasshouse conditions with natural light. Flowers from each backcross individual were photographed, and individuals were visually grouped into three phenotypic classes (blue, white and F₁ phenotype). Floral anthocyanins were quantified by eluting pigments from fresh corolla tissue in 500 µl methanolic HCl (1%) and measuring absorbance at 530 nm. Reflectance spectra were collected from the corolla tube and lobe of each individual using a JAZ spectrometer as described by Ng & Smith (2016). Spectral variation was examined by principal component analysis of the combined corolla tube and lobe data in the visible spectrum (400–700 nm).

Association mapping

Bulk transcriptome sequencing of each parent and phenotypic pool was used to search for allelic variants associated with flower

color. Floral bud RNA was extracted following Coburn *et al.* (2015) from the blue and white parents (two biological replicates each); 20 blue and 15 F₁ phenotype individuals from the backcross to the blue parent; and eight white and nine F₁ phenotype individuals from the backcross to the white parent. Two lanes of Illumina 100-base-pair paired-end RNAseq of Tru-Seq libraries (Illumina, Inc., San Diego, CA, USA) were generated for these six pools of RNA. We created a transcriptome index from the *I. cyaneum* transcriptome (Gates *et al.*, 2016) with BOWTIE 2.02 (Langmead *et al.*, 2009) with default settings and mapped reads from each pool using TOPHAT 2.1.0 (Kim *et al.*, 2013) allowing five mismatches per read (-N flag) and a final read edit distance of 8. This TOPHAT mapping for each pool generated a BAM file to be used in both association and expression analyses. In addition to each TOPHAT BAM mapping file, downstream expression and variant calling analyses also require a gtf mapping file. Without a prespecified gtf from an annotated genome assembly, TOPHAT makes run-specific gtf mapping files along with each BAM mapping but can exclude low expression genes of interest (e.g. anthocyanin pathway genes in white-flowered pools) in these gtf files. To avoid potentially missing genes in the gtf file, we used BWA-MEM (Li & Durbin, 2009) to construct a pan-gtf file by mapping reads from all libraries to the *I. cyaneum* transcriptome. This mapping was passed to TOPHAT for gtf file construction and the resulting gtf file contained all gene models with reads that were mapped in any pool. We then used SAMTOOLS (Li *et al.*, 2009) to generate variant calls in each of the mappings. Before association mapping, we removed low-quality variants with < 10-fold coverage and with a quality score of < 50.

Two approaches were used to test for associations between genes and color phenotypes using the mapped reads. First, we conducted a transcriptome-wide search for genes with patterns and frequencies of single nucleotide polymorphism (SNP) variation consistent with the phenotypic pools. We focused on genes that differed between the parents by three or more SNPs with > 200× coverage to increase confidence in the assignment of alleles to the *I. cyaneum* or *I. loxense* parents. We filtered this initial list of genes, saving those which showed no mapped SNPs in the blue backcross pool (i.e. all reads match the blue *I. cyaneum* parent). This follows from the expectation that blue-flowered individuals should be homozygous for the blue parental allele at the causative locus. We further reduced the pool by applying a second filter, keeping only genes with 35–65% blue variants in the F₁ phenotypic pools and < 35% blue variants in the white phenotypic pool. With precise phenotyping, we would expect 0% blue variants in the white pool; however, distinguishing the two phenotypes was difficult in the backcross to the white parent. We expected the F₁ phenotypic pool to have intermediate allele frequency, but not precisely 50% blue variants because of the difficulty in phenotyping as well as any *cis*-regulatory effects.

Next, we compared loci recovered from this transcriptome-wide pipeline to SNP variation at candidate loci in the anthocyanin pathway. We included seven structural genes (*Chs*, *Chi*, *F3h*, *F3' h*, *F3' 5' h*, *Dfr*, *Ans*) and six known transcription factors (the R2R3 MYB activator *AN2* and repressor *MYB27*, the bHLH genes *jaf13* and *ANI*, the WDR *AN11* and the R3 MYB

repressor *MYBx*) (Koes *et al.*, 2005; Albert *et al.*, 2014). We refer to the names of the loci in *Petunia hybrida* for six transcription factors (*AN1*, *AN2*, etc.) for ease of comparison with existing literature. The *Ichroma* orthologs for these pathway genes have been characterized in previous studies (Smith & Rausher, 2011; Coburn *et al.*, 2015; Gates *et al.*, 2016). For each locus, we calculated the proportion of reads matching the blue *I. cyaneum* parent allele in each phenotypic pool.

Phylogenetic analyses of *MYBL1*

A combination of bioinformatic and phylogenetic analyses was used to elucidate the evolutionary history of *MYBL1*, an R3 MYB gene identified by our pipeline as having a strong association with flower color. Potential orthologs in related taxa were retrieved through a combination of BLAST searches and gene prediction using Augustus (Stanke & Morgenstern, 2005). Genome assemblies for tomato (*Solanum lycopersicum*), potato (*S. tuberosum*), *S. pennellii*, pepper (*Capsicum annuum*), tobacco (*Nicotiana tabacum*), petunia (*P. hybrida*) and coffee (*Coffea canephora*) were accessed through the Sol Genomics Network (solgenomics.net) and the Coffee Genome Hub (coffee-genome.org). Protein sequences for *MYBL1* were aligned to similar Solanaceae MYB sequences along with representatives of the major MYB subgroups in *Arabidopsis* (Stracke *et al.*, 2001; Dubos *et al.*, 2010) for phylogenetic analysis. The sequences were aligned using Clustal in GENEIOUS 6.05 (Biomatters Inc., Auckland, New Zealand) with default settings. The phylogeny was estimated in MRBAYES 3.2.5 (Ronquist & Huelsenbeck, 2003) with two independent runs, each of 1 million generations, and a mixed prior on models. Convergence was judged by potential scale reduction factor (PSRF) values approaching 1.0, estimated sample sizes (ESS) exceeding 300 and similarity of the consensus trees from the two runs.

Expression analyses

Expression levels of anthocyanin transcription factors and regulatory genes in each of our six RNA pools (the two parents and the two phenotypes in each backcross population) were measured by mapping reads to the *I. cyaneum* transcriptome as described above. After mapping we calculated fragments per kilobase per million (FPKM) and tested for differential expression using CUFFDIFF 1.1.2 (Trapnell *et al.*, 2010) with our custom .gtf file. CUFFDIFF uses an optimization routine to obtain a maximum *a priori* estimate of the FPKM for a given fragment (typically a gene). From this sampling routine, the program also calculates a 95% confidence interval based upon the variance–covariance relationship of high likelihood samples that are also weighted by an information index (as a means of ensuring lower coverage fragments will have appropriate levels of uncertainty). We used the FPKM estimates and accompanying confidence intervals to compare expression levels across pools for each gene by using a two-sample *t*-test where the standard error for the expression of each gene at each pool is calculated as the confidence interval divided by 1.96. We also conducted quantitative real-time PCR (qPCR)

to confirm patterns of expression across the phenotypic pools. Three individuals were randomly selected from the four backcross pools. We completed two replicate qPCRs for each individual and three for each parent for seven structural genes and four transcription factors (see Supporting Information Table S1 for primers). Two of the six transcription factors included in the association analysis (*MYB27* and *AN1*) were excluded from the qPCR experiment because of their low expression across all pools in the RNASeq experiment. Protocols for cDNA synthesis, qPCR conditions and data analysis follow Coburn *et al.* (2015).

Functional testing

The function of *ILMYBL1* was tested through stable transformation of *N. tabacum*. First, the complete coding sequence of *MYBL1* from *I. loxense* was synthesized (GenScript Corp., Piscataway, NJ, USA) and cloned into the pRTL2 vector (Carrington & Freed, 1990), resulting in constitutive expression regulated by the enhanced cauliflower mosaic virus 35S (e35S) promoter with the tobacco etch virus (TEV) translational enhancer element. The resultant expression cassette was subsequently subcloned into the binary vector pXP212 (Hajdukiewicz *et al.*, 1994). Tobacco leaf disks were transformed with this vector following the protocol outlined by Clemente (2006), and 20 of the resulting transformants were grown from seed in the glasshouse. Floral pigmentation was quantified from both transgenic and wild-type plants by eluting anthocyanins from petal disks with 500 µl methanolic HCl (1%) and measuring absorbance at 530 nm.

Results

Patterns of phenotypic variation in crosses suggest a single underlying locus

Crosses between the white-flowered *I. loxense* and the blue-flowered *I. cyaneum* resulted in a pale (nearly white) phenotype. The F₁ has a white floral tube with slight anthocyanin pigmentation near the mouth of the corolla (Fig. 1a). Such patterning is not apparent in the parents, which are uniformly colored throughout the tube and the mouth. The highly reduced level of floral pigmentation in the F₁ suggests nearly complete dominance of the allele (or alleles) from the white-flowered *I. loxense*.

Backcrosses of the F₁ to *I. loxense* and *I. cyaneum* recovered the three parental phenotypes, consistent with a single major locus determining the flower color difference. Both blue and F₁ phenotypes segregated in the backcross to the blue *I. cyaneum* parent, and F₁ and white phenotypes in the backcross to the white *I. loxense* parent (Fig. 1a). As expected for a single gene controlling the phenotype, the numbers of individuals with each phenotype did not differ significantly from 1 : 1 (backcross to blue, 35 blue: 38 F₁ phenotype; backcross to white, 8 white: 11 F₁ phenotype, *P* = 0.8, Fisher's exact test). Fewer individuals were scored in the backcross to white because of low viability in that cross (possibly due to incipient incompatibility between the sister species).

Quantitative comparisons of pigment concentration showed a clear split between the blue phenotype and the other two

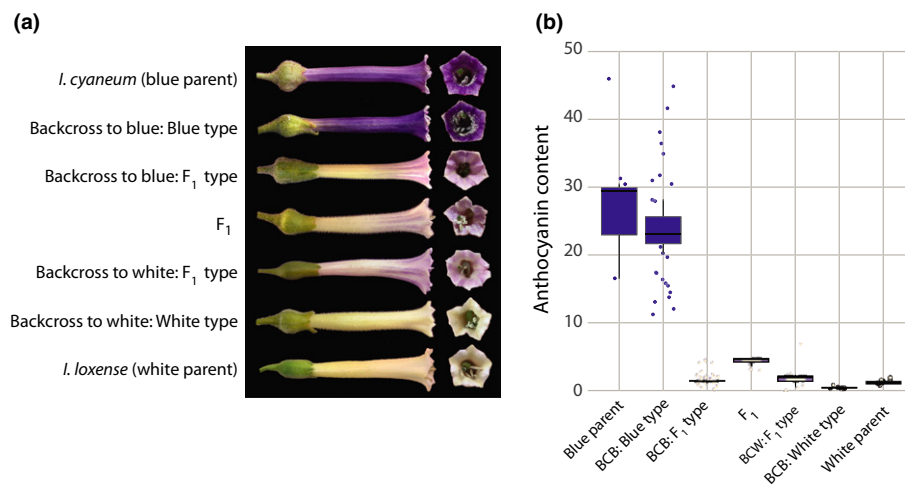


Fig. 1 Phenotypic variation in parents and backcrosses. (a) Flowers and corolla mouths of parental species (*Ipomoea cyaneum* and *I. loxense*), their F_1 and example phenotypes from the backcrosses. (b) Boxplots of anthocyanin content in corolla tissue, measured as absorbance at 530 nm g^{-1} fresh tissue. Order of the phenotypes along the x-axis follows (a). Boxes encompass the upper and lower quartiles and the horizontal line represents the median.

phenotypes. The blue-flowered parent and the blue-flowered individuals from the backcross to the blue parent produced about five times the amount of anthocyanin pigment as the white or F_1 phenotype individuals, with no overlap in values (Fig. 1b). Although the white-flowered backcross individuals had lower average pigment content than the F_1 phenotype individuals, there was significant overlap among the pools (Fig. 1b).

A similar pattern was observed in a principal component (PC) analysis of floral color spectra, which sharply divided the blue-flowered individuals and the rest. The first principal component axis captured 84% of the variation and split the individuals into two clusters, *I. cyaneum* and the blue-flowered individuals from the backcross to *I. cyaneum* vs the others (Fig. S1a). This axis corresponds to brightness (the total amount of light reflected), as each of the wavelengths of light has roughly equal loading (Fig. S1b). As expected, blue flowers have low brightness, while the white and F_1 phenotype flowers have high brightness (Fig. S1a). The other minor PC axes incorporate relative reflectance across different wavelengths (floral hue), and here indicate different reflectance in green/yellow wavelengths vs red/blue (Fig. S1b).

Bulk-segregant mapping points to an R3 MYB associated with color

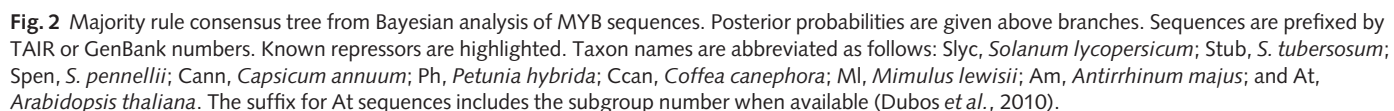
We used a bulk-segregant RNA-Seq approach to identify genes associated with the phenotypic classes from the crosses. Although the white and F_1 phenotypes overlap in pigment concentration and reflectance spectra, we sequenced each of the pools separately in order to detect expression differences that could account for their distinct color patterns (Fig. 1a). Our first filtering of the transcriptomic pools for associated genes (requiring three confidently called SNPs between the parents and no variants called between *I. cyaneum* and the blue-flowered pool) resulted in 35 candidates (Table S2). Among these candidates, only one belonged to a class of genes (MYB transcription factors) known to regulate anthocyanin production. Although the gene appeared to be a single repeat R3 MYB, the closest BLAST hit in tomato was an R2R3 MYB with uncharacterized function (Table S2). We next filtered the list of 35 candidates by SNP frequency in the remaining pools, retaining genes with 35–65% reads matching

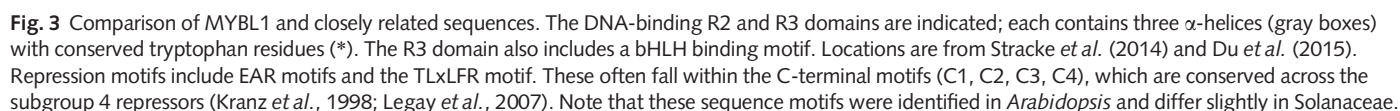
I. cyaneum in the F_1 phenotype pools and <35% in the white pool. The apparent R3 MYB sequence was the only gene to pass this second filter, and given its phylogenetic proximity to *A. thaliana* MYBL2 (see phylogenetic analysis in next Results section), this locus was designated *MYBlike-1* or *MYBL1*. The *MYBL1* sequences for *I. loxense* and *I. cyaneum* have been uploaded to GenBank as KY658468 and KY658469.

We compared these results to the known structural and regulatory genes of the anthocyanin pathway, none of which emerged from the transcriptome-wide search. All of the structural genes were segregating both blue and white parental alleles in the blue-flowered pool, indicating that none of these comprises the causative locus. The same pattern was observed for four of the six transcription factors (the R2R3 MYB *AN2*, the bHLH *jaf13*, the WDR *AN11* and the R3MYB *MYBx*). The *AN1* bHLH ortholog presented nearly undetectable floral expression (Table S3), and thus no SNPs could be confidently called across the pools. This result suggests that, between *jaf13* and *AN1*, the former is probably the principal bHLH partner in the anthocyanin regulatory complex in *Ipomoea* flowers. The final candidate locus, the R2R3 MYB repressor *MYB27*, did present zero called variants in the blue-flowered pool, as would be predicted for a causal locus. However, this gene was not recovered in the transcriptome-wide search because of its low expression (Table S3) and accordingly low coverage. Such low floral expression is consistent with studies in *Petunia*, suggesting that its primary role is repression of anthocyanins in vegetative tissue (Albert *et al.*, 2011). Thus, the lack of called variants for *MYB27* in the blue pool is probably due to linkage with the causative locus. Indeed, additional bioinformatic analyses (see gene annotation analyses in next Results section) suggest that the *MYBL1* gene detected in the transcriptome-wide search is closely linked to the *MYB27* ortholog in *Ipomoea*.

Novel R3 MYB belongs to a clade of R2R3 MYB repressors

The similarity of *Ipomoea* *MYBL1* to an R2R3 MYB in tomato raised the possibility that this R3 MYB might be derived from a past duplication of an R2R3 MYB. Indeed, additional searches of nearby tomato scaffolds from chromosome 5 uncovered a highly similar sequence in an unannotated region that was 10 kb from





Phylogenetic analyses of *MYBL1* along with other R2R3 and R3 MYBs are consistent with a recent origin via duplication within the clade of R2R3 MYB repressors. The *MYBL1* sequences from *Iochroma*, *Capsicum* and *Solanum* form a well-supported clade (100% posterior probability, Fig. 2) that is closely related to a group of ‘MYB3-like’ Solanaceae R2R3 genes, including the tomato R2R3 originally uncovered in the BLAST search. These

Members of the *MYBL1* clade share several changes in their coding sequence that may relate to their functional evolution. First, these MYBs present large deletions in the R2 domain, comparable to *AtMYBL2* and other R3 MYBs (Fig. 3). These deletions have resulted in the loss of the helix–turn–helix motifs

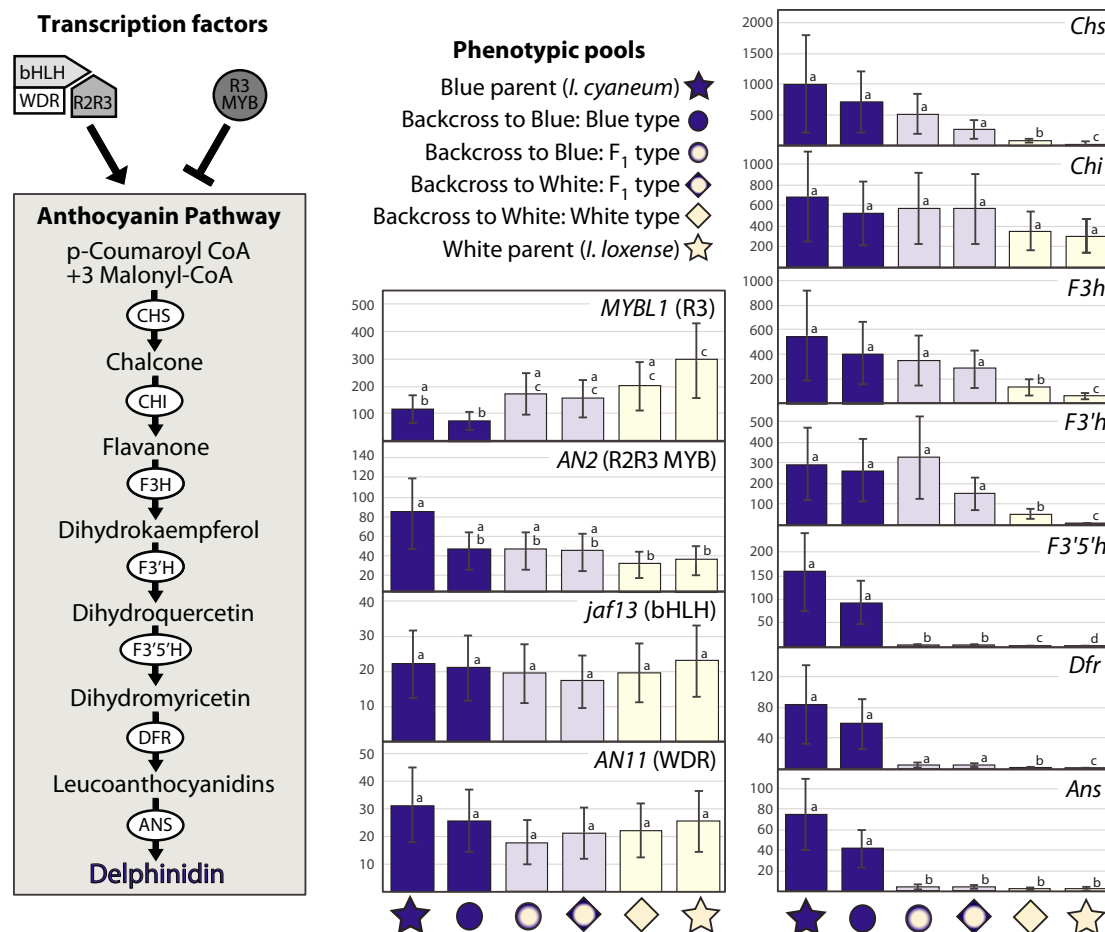


Fig. 4 Variation in pathway gene expression across phenotypic pools. A simplified pathway is depicted on the left comprising the enzymes chalcone synthase (CHS), chalcone isomerase (CHI), flavonone 3-hydroxylase (F3H), flavonoid 3'-hydroxylase (F3'H), flavonoid 3',5'-hydroxylase (F3'5'H), dihydroflavonol 4-reductase (DFR) and anthocyanidin synthase (ANS). Branches of the pathway leading to pelargonidin and cyanidin pigments are not shown as *I. cyaneum* produces only delphinidin. Note that in *Lochroma*, F3'5'H has little 3' activity and thus appears not to act on dihydrokaempferol (Smith & Rausher, 2011). Elements of the pathway are regulated by bHLH, WDR, and R2R3 and R3 MYB proteins. Expression levels are given for each of these loci in fragments per kilobase per million (FPKM). Error bars denote 95% confidence intervals, and lower-case letters indicate significantly different values.

required for DNA-binding and functionality of the domain (Ogata *et al.*, 1994; Williams & Grotewold, 1997). Second, the *MYBL1* sequences have lost the C-terminal active repression motif DLN_x2P present in the MYB3-like R2R3 sequences and in *Petunia's* *MYB27*. Nonetheless, they have gained a novel repression motif (LxLxL) near the end of the R3 domain (Fig. 3). Both of these repression motifs fall into the class of ethylene-responsive element binding factor-associated amphiphilic repression or EAR motifs, which are found through plants (Kagale & Rozwadowski, 2011). Thus, it appears that the repressive activity of *MYBL1* was probably retained despite a *c.* 40% reduction in the length of the protein. Neither the *MYBL1* clade nor *AtMYBL2* contain the WxM motif required for cell-to-cell movement (Kurata *et al.*, 2005), suggesting they do not diffuse like the smaller R3 MYBs of the CPC clade (Fig. 2).

Differential expression associated with color phenotypes

Comparison of gene expression levels across the backcross pools points to upregulation of *MYBL1* associated with loss of

pigmentation, consistent with its putative role as a pigment repressor. In the backcross-to-blue, the F₁ phenotype individuals have roughly twice the *MYBL1* expression of the blue-flowered individuals (Fig. 4). Given that the *trans*-regulatory environment should be homogenized across these two pools, this difference would be best explained by a *cis*-regulatory change at *MYBL1*. Expression levels for *MYBL1* were slightly elevated in the white backcross individuals compared to those with the F₁ phenotype, but the difference was not significant (Fig. 4). A marked difference in expression of *MYBL1* was also observed between the parents, although the *I. cyaneum* expression level overlapped with multiple backcross pools. These qPCR experiments produced very similar patterns, with the strongest differences between the two parents and between the two phenotypic pools in the backcross to blue (Fig. S2). Given the relatively subtle differences in expression, additional experiments would be useful to better quantify allele-specific *MYBL1* expression across the individuals in these segregating populations.

By contrast with *MYBL1*, the other transcription factors showed no evidence of differential expression in association with

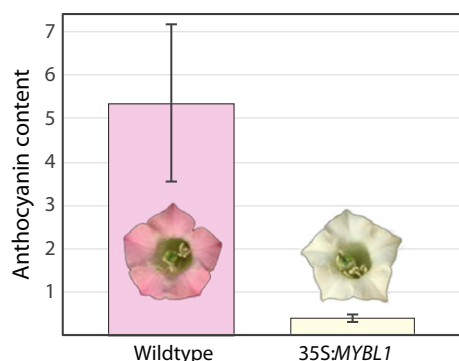


Fig. 5 Effects of *MYBL1* expression in *Nicotiana tabacum*. Anthocyanin content is measured as absorbance at 530 nm g⁻¹ fresh tissue. Error bars show ± 1 SD. Representative flowers are shown for each treatment.

the color phenotypes. The three transcription factors with quantifiable floral expression (*AN2*, *jaf13* and *AN11*) showed statistically indistinguishable levels of expression across the backcross pools in both the transcriptomic analysis and the qPCR experiment (Figs 4, S2). The lack of variation in expression at these loci suggests that *MYBL1* does not regulate these transcription factors, and instead controls flower pigmentation through regulation of the structural genes of the pathway.

Consistent with this hypothesis, we observed marked variation in the expression of structural genes. In particular, *F3'5'h*, *Dfr* and *Ans* showed an almost complete loss of expression in both the white-flowered and the F₁ phenotype pools, suggesting that their expression is strongly inhibited by the increase in *MYBL1* expression. The significant downregulation of these three loci was also recovered in the qPCR experiment (Fig. S2). Two of the early genes, *Chs* and *F3h*, also showed a trend toward lower expression with decreasing pigmentation, but these patterns were not consistently significant (Figs 4, S2). By contrast with the other structural genes, *Chi* did not exhibit lowered expression in the less pigmented pools. This gene may be controlled by different transcriptional regulators, as implicated by studies in several other systems (Dubos *et al.*, 2008; Yi *et al.*, 2010; Koehler *et al.*, 2012).

R3 MYB acts as a functional floral anthocyanin repressor in *Nicotiana*

Expression of *MYBL1* from *I. loxense* in *N. tabacum* (tobacco) resulted in a nearly complete loss of floral anthocyanins. All of the transformed lines presented the white-flowered phenotype, with 11–20 times lower absorption across them and 14 times lower absorbance on average (Fig. 5). *N. tabacum*, another member of the Solanaceae, is estimated to be diverged from *Iochroma* 32 Ma (De-Silva *et al.*, 2017), indicating that the R3 MYB has the ability to repress anthocyanins in distantly related taxa. Also, the primary pigment in *N. tabacum* is cyanidin (McCarthy *et al.*, 2017), as opposed to delphinidin, which is in the blue-flowered *Iochroma*. Given that cyanidin, a derivative of dihydroquercetin, does not require F3'5'H activity, this result suggests that the level of downregulation at the other pathway genes (Fig. 4) is probably sufficient to abolish pigment production.

Discussion

Through a combination of crossing studies and bulk-segregant RNA-sequencing, our results indicate that a novel R3 MYB transcription factor, *MYBL1*, controls the intensity of floral pigmentation in *Iochroma*. The *MYBL1* allele from the white-flowered *I. loxense* acts in a nearly dominant fashion, with the F₁ hybrid between *I. loxense* and the blue-flowered *I. cyaneum* presenting only traces of anthocyanin pigmentation towards the mouth of the corolla. Patterns of gene expression suggest that this loss of pigmentation is due to downregulation of structural genes in the pathway, particularly the late anthocyanin-specific elements. Although we cannot exclude the possibility of causative coding sequence mutations at *MYBL1*, the elevated expression of this gene in the F₁ phenotype and white-flowered pools suggests a *cis*-regulatory change as the underlying mechanism, analogous to the R3 MYB transcription factor controlling flower color in *Mimulus* (Yuan *et al.*, 2013). Under this scenario, the transition from the ancestral state of blue flowers to the derived state of white flowers along the *I. loxense* lineage could be explained by the increased expression of the *MYBL1* regulator leading to downregulation of anthocyanin pathway genes and suppression of floral pigment production.

While the structure of *MYBL1* resembles known R3MYB repressors, phylogenetic analyses demonstrate that *MYBL1* is evolutionarily distinct. Most of the previously described R3 MYB repressors, such as *CAPRICE* (*CPC*) in *Arabidopsis* and *MYBx* in *Petunia*, fall into a single clade that is distantly related to *MYBL1* (Fig. 2). Similar to *MYBL1*, these genes lack an R2 domain and contribute to the regulation of anthocyanin production (Zhu *et al.*, 2009) along with other aspects of epidermal cell differentiation (Wada *et al.*, 1997; Schnittger *et al.*, 1999; Serna, 2008; Wang & Chen, 2014). While the closest relatives of the CPC clade of R3 MYBs remain unclear, the *MYBL1* clade is well resolved as a part of the subgroup 4 R2R3 repressors (Fig. 2). All of the characterized subgroup 4 genes in *Arabidopsis* (*AtMYB4*, *AtMYB7* and *AtMYB2*) regulate elements of the phenylpropanoid pathway, which includes anthocyanins (Jin *et al.*, 2000; Preston *et al.*, 2004; Fornale *et al.*, 2014). *Petunia MYB27* also belongs to this subgroup 4 clade and modulates anthocyanin production (Albert *et al.*, 2011). These patterns suggest that anthocyanin regulation is probably the ancestral state for the *MYBL1* clade as well as the other post-duplication copy, *MYB3like*. Given the physical proximity of the *MYBL1* genes to the *MYB3like* R2R3 genes in the Solanoid genomes (*c.* 10 kb), we hypothesize that these R3 MYBs arose by tandem duplication followed by loss of the R2 domain.

The molecular evolution of the *MYBL1* clade within Solanaceae presents a striking parallel to the evolution of the *AtMYB2* gene in *Arabidopsis*. Both of these single-domain R3 MYB repressors arose from clades of R2R3 MYB repressors, and both show a dynamic history of loss and gain of repression motifs. *AtMYB2* lost the EAR motif shared among the subgroup 4 R2R3 MYB repressors (Liu *et al.*, 2015) but gained a novel TLxLFR repression motif at the C-terminus (Matsui *et al.*, 2008). Similarly, *MYBL1* lost the EAR motif characteristic of the

MYB3like genes and *Petunia MYB27* (Albert *et al.*, 2011), but acquired a new EAR motif near the end of the R3 domain (Fig. 3). Like *AtMYBL2*, the *MYBL1* clade has retained the conserved bHLH binding motif in the R3 domain (Zimmermann *et al.*, 2004) and thus is likely to bind with the bHLH transcription factors and act as part of an MBW regulatory complex.

The presence of the EAR repression motif in *MYBL1* suggests that it functions as an active transcriptional repressor, although some degree of passive repression is possible. Given the structural similarity to *AtMYBL2*, we predict that *MYBL1* binds the bHLH factor and possibly also to the R2R3 partner (Albert *et al.*, 2014). The presence of the R2R3 MYB (*AN2* homolog), which normally acts as an activator, allows the complex to identify the transcriptional targets, while the R3 MYB induces epigenetic silencing via the EAR motif (Kagale & Rozwadowski, 2011). Thus, the replacement of one of the R2R3 MYB partners in the MBW complex with an R3 MYB could transform the complex from an activator of anthocyanin gene transcription to a repressor (Albert *et al.*, 2014) (Fig. 6). Depending on its abundance relative to the R2R3 activators, it is also possible that *MYBL1* could passively repress anthocyanin gene expression by titrating bHLH factors, similarly to the CPC clade of R3 MYBs that lack the EAR active repression motifs (Mitsuda & Ohme-Takagi, 2009). Both mechanisms are consistent with the pattern seen in the backcrosses, where intermediate levels of *MYBL1* expression (as in the F_1) allow some pigment production (presumably because some MBW complexes contain only R2R3 MYB activators). When R3 MYB expression is high, as in the white-flowered pools and parent, most MBW complexes contain at least one R3 MYB, and thus function as repressors. Additional studies would be needed to confirm this model for *MYBL1* function.

The mode of action of *MYBL1* may have important consequences for understanding the evolutionary trajectory that led to the transition to white flowers in *I. loxense*. Judging from expression levels in the phenotypic pools (Fig. 4), a 30% increase in *MYBL1* expression might be sufficient to convert a blue flower into a nearly white flower. Even a small increase might be sufficient to produce a pale phenotype that could be targeted by selection, whether by biotic factors, like pollinators, or abiotic factors, such as climatic conditions (Rausher, 2008). We do not presently know the mutation(s) that caused the observed expression change in *MYBL1*, but if it were accomplished with a single mutation, the resulting allele could spread quickly through the population given its nearly dominant action (Haldane, 1924, 1927). This stands in contrast to most of the previously identified genetic changes associated with transitions to white flowers, which involve loss-of-function mutations in the R2R3 MYB activators (Quattrocchio *et al.*, 1999; Schwinn *et al.*, 2006; Hoballah *et al.*, 2007). These alleles are recessive and thus would experience a lower probability of fixation, a phenomenon known as ‘Haldane’s Sieve’ (Turner, 1981; Charlesworth, 1992).

Although loss-of-function R2R3 MYB mutations and gain-of-function R3 MYB mutations are expected to experience different rates of fixation, their ultimate effect on development is likely to be similar. Because the R3 MYB acts by partnering with the bHLH and using the R2R3 MYB to identify targets, the set of

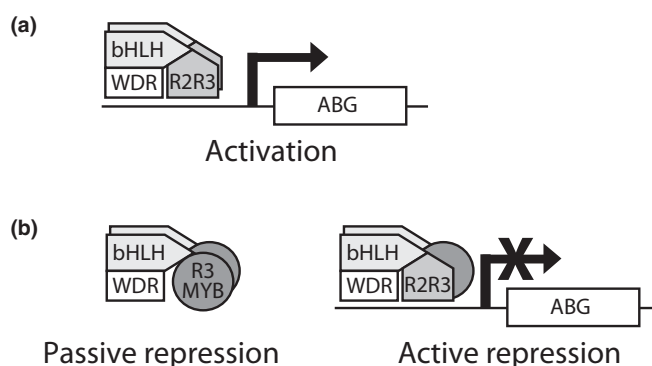


Fig. 6 Model for R3 MYB repression of anthocyanin production. (a) Activation of anthocyanin biosynthetic genes (ABG) by the MBW complex based on the model of Albert *et al.* (2014). The MYB and bHLH proteins are thought to be present as dimers within the complex, with MYBs being bridged through their binding to the bHLH factors. (b) The R3 MYB (dark circle) could block anthocyanin production by titrating bHLH factors (and their bound WD40 partners) as passive repressors (left) or by partnering with R2R3 MYB activators as part of MBW complexes and triggering active (transcriptional) repression (right).

anthocyanin genes which experience downregulation are expected to be the same. In *Iochroma* and other asterids, the MBW complex, whether acting as an activator or repressor, most strongly targets the late genes of the pathway. The isolation of effects to the late genes may be important in allowing the suppression of pigment production but maintenance of other pathway products, such as flavones and flavonols (Streisfeld & Rausher, 2011; Wessinger & Rausher, 2012). Similarly, the apparent bias towards fixation of regulatory mutations as opposed to coding mutations in structural genes may reflect selection to maintain the intact pathway for pigment production in other tissues and developmental stages (Wessinger & Rausher, 2012; Ho & Smith, 2016). Overall, this work together with studies in other flowering plants suggests that convergence at the level of individual genes and mutations may not be expected when the underlying mechanisms present multiple, developmentally similar, pathways to the same phenotype.

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Author contributions

S.D.S. and D.J.G. designed and carried out the experiments and analyzed the data; B.J.S.C.O. collaborated on the bioinformatic analysis; T.E.C. contributed materials and expertise to the functional test in *Nicotiana*. D.J.G. and S.D.S. wrote the paper, and all authors contributed to revision of the manuscript.

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Supporting Information

Additional Supporting Information may be found online in the Supporting Information tab for this article:

Fig. S1 Principal components analysis of floral color spectra.

Fig. S2 Differences in gene expression based on qualitative real-time PCR.

Table S1 Primer pairs for qPCR

Table S2 Thirty-five candidate genes from first bioinformatics filter

Table S3 SNP frequencies and expression levels for anthocyanin pathway genes

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