Chromosome-Scale Genome Assembly of *Gilia yorkii* Enables Genetic Mapping of Floral Traits in an Interspecies Cross

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Abstract

Substantial morphological variation in land plants remains inaccessible to genetic analysis because current models lack variation in important ecological and agronomic traits. The genus *Gilia* was historically a model for biosystematics studies and includes variation in morphological traits that are poorly understood at the genetic level. We assembled a chromosome-scale reference genome of *G. yorkii* and used it to investigate genome evolution in the Polemoniaceae. We performed QTL (quantitative trait loci) mapping in a *G. yorkii*/*C. capitata* interspecific population for traits related to inflorescence architecture and flower color. The genome assembly spans 2.75 Gb of the estimated 2.80-Gb genome, with 96.7% of the sequence contained in the nine largest chromosome-scale scaffolds matching the haploid chromosome number. *Gilia yorkii* experienced at least one round of whole-genome duplication shared with other Polemoniaceae after the eudicot paleohexaploidization event. We identified QTL linked to variation in inflorescence architecture and petal color, including a candidate for the major flower color QTL—a tandem duplication of flavanol 3',5'-hydroxylase. Our results demonstrate the utility of *Gilia* as a forward genetic model for dissecting the evolution of development in plants including the causal loci underlying inflorescence architecture transitions.

Key words: evolution of development, genome assembly, *Gilia*, inflorescence architecture, QTL mapping, whole-genome duplication.

Significance

Forward genetics with model species has made substantial contributions to our understanding of the genes regulating plant growth and development; however, these model species represent only a small fraction of existing plant diversity. Here, we report the development of genetic and genomic resources in *Gilia yorkii*, including a high-quality genome assembly and interspecies mapping population. These resources will facilitate the identification of genes underlying diversity in important plant traits, including floral and inflorescence architecture.

Introduction

The broad adoption of a few model organisms in genetics and molecular biology has been highly productive for diverse fields of biology. In plants, sustained focus on all aspects of *Arabidopsis thaliana* (*Arabidopsis*) informs research in a diversity of species and problems beyond that narrow model (Woodward and Bartel 2018). However, *Arabidopsis* lacks important agronomic and ecological traits. Even apparently conserved traits can diverge at the genetic level via developmental system drift (True and Haag 2001), underscoring the need for comparative work outside of these established models. Judiciously
expanding the suite of plant species used for genetic studies remains imperative, particularly in fields like evolution of development (evo–devo), where understanding the emergence and change of developmental networks requires comparative work (Wagner 2014). Despite efforts to develop phylogenetically diverse land plant models for evo–devo (Rensing et al. 2008; Kramer 2009; Banks et al. 2011; Bowman et al. 2017; Yuan 2019), considerable plant morphological diversity remains inaccessible because no species polymorphic for these traits are currently subjects of genetic analysis.

Historically, genome size limited potential model species, but advances in high-throughput sequencing are making assembly of large complex genomes routine (Zimin et al. 2017; Hufford et al. 2021), opening a much larger suite of possible species. An important remaining constraint is the ability to employ forward genetics to move beyond the limitations of candidate gene studies. In the context of evo–devo, a forward genetic approach involves crossing closely related species that are polymorphic for interesting morphological traits to identify quantitative trait loci (QTL) and eventually the causative polymorphisms underlying morphological evolution (Colosimo et al. 2005; Doebley et al. 2006; Jeong et al. 2008). Harnessing the power of forward genetics in wild species requires that they not only be polymorphic for traits of interest, but also capable of producing a fertile hybrid so that segregating mapping populations can be generated.

The ability to make wide hybrids is common in many plant lineages, and pioneering work examining the genetic basis of ecological and morphological differences among closely related species (Clausen and Hiesey 1958; Grant 1981) can now be revisited with the increased resynthesis of modern genomic and molecular approaches. Among these early studies, work on the leafy-stemmed gilias (Gilìa section Gilìa, Polemoniaceae) stands out (Grant and Grant 1956). Leafy-stemmed gilias are mostly diploid annuals native to western North and South America, grow readily in greenhouse conditions, hybridize easily, and are polymorphic for a diverse array of morphological traits (Grant 1954). We have chosen two closely related leafy-stemmed Gilìa species that form a fertile hybrid for forward genetic analysis, the broadly dispersed G. capitata and the narrow endemic G. yorkii. These species differ in multiple morphological features including flower color and inflorescence architecture.

Here, we propose G. yorkii and G. capitata as a promising new model for the evolution of development. We have generated a chromosome-scale genome assembly for G. yorkii which facilitated an analysis of whole-genome duplication (WGD) history in Ericales and Polemoniaceae and identification of QTL for morphological traits that distinguish these species.

## Results

### Genome Assembly and Annotation

A chromosome squash revealed that metaphase-stage mitotic cells of G. yorkii contain 18 chromosomes (2n = 2x = 18) (fig. 1a and supplementary fig. S1, Supplementary Material online). Using a kmer analysis, we estimated the G. yorkii genome to be 2.801 Gb, with 75% estimated to be repetitive sequences. The sequenced plant exhibited low heterozygosity (0.133%; supplementary fig. S2, Supplementary Material online). The genome was sequenced with PacBio to approximately 67× coverage, and sequencing reads were assembled into 3,947 contigs (table 1). The contig assembly was polished with the PacBio and Illumina reads to correct SNP and InDel errors and then combined into 2,043 scaffolds (table 1) using Hi-C sequencing reads. The scaffolded assembly is chromosome scale, with 96.7% of the total sequence length assembled into the nine largest scaffolds (fig. 1b), corresponding to the haploid chromosome number of G. yorkii. The pattern of chromosomal interactions in the ordered Hi-C heatmap was characteristic of full-length chromosomes in which arms of a given chromosome interact with each other (supplementary fig. S3, Supplementary Material online) in the previously described Rabl conformation (Cowan et al. 2001).

In addition, seven of the nine chromosome-scale scaffolds have the canonical telomeric repeat sequences (TTTAGGG) in at least one end of the chromosome, with three containing the telomeric sequences at both ends, indicative of a largely complete chromosome-scale assembly (fig. 1b). A benchmarking of universal, single-copy orthologs (BUSCO) analysis also indicated that the assembly is largely complete, with 97.6% of BUSCOs being identified in the scaffolded assembly and 96.8% identified as complete (table 1). As expected for a diploid, most BUSCOs (92.4%) were identified as single copy. Given these results, we refer to the largest nine scaffolds hereafter as chromosomes, numbered from largest (Gy1) to smallest (Gy9).

In total, 75.60% of the assembly was annotated as repetitive (supplementary table S2, Supplementary Material online), in agreement with the kmer-based prediction. Long-terminal repeat (LTR) retrotransposons of the Class I transposable elements are the most common repetitive sequences in the genome, accounting for 45.81% of the entire genome assembly, with Copia LTRs alone representing 29.43% of the genome. A large proportion of the genome (21.92%) was annotated as unknown repetitive sequences, suggesting that many of the repetitive sequences in G. yorkii are family-, genus-, or species-specific.

Genes were annotated in the G. yorkii assembly using transcript evidence from PacBio Iso-Seq data assembled into a final mapped transcriptome data set consisting of 96,691 full-length transcripts. An analysis of BUSCO genes indicated that the transcriptome assembly was largely complete, with only slightly fewer BUSCOs identified in the transcriptome
Chromosome-Scale Genome Assembly of G. yorkii

43,969 genes. Final gene models were strongly supported by the evidence used for annotation, with 80% of the genes having annotation edit distance values less than 0.4 (supplementary fig. S4, Supplementary Material online). Genes are distributed throughout all nine chromosomes, with high density near the ends of chromosomes that generally decreases with increasing distance from the ends (fig. 1b).

Whole-Genome Duplication in G. yorkii

Duplicated, collinear genes are distributed among all chromosomes in the G. yorkii genome, in many cases occurring in long stretches of collinear genes (fig. 2a). To assess whether these duplications were consistent with ancient WGDs in G. yorkii, we analyzed the age distribution of gene duplications. The mixture model supported a single peak consistent with WGD around $K_s \approx 1.07$ (named GILIx, fig. 2b, supplementary table S3, Supplementary Material online). Comparison of this gene age distribution to transcriptomes of two additional members of the temperate subfamily of the Polemoniaceae, Saltugilia latimeri and Phlox cuspidata (Larson et al. 2020), revealed that both species show a similar peak of duplication at $K_s \approx 1$ (fig. 2b). The ortholog divergences among S. latimeri, P. cuspidata, and G. yorkii, were $K_s = 0.2–0.7$ (fig. 2c and supplementary table S4, Supplementary Material online). This suggests that the ancient WGD is likely shared by at least the temperate subfamily of Polemoniaceae. We then compared G. yorkii with the more distantly related species Diospyros oleifera (Ebenaceae) (Suo et al. 2020) in the same order (Ericales). The D. oleifera genome has a peak of duplication at $K_s \approx 0.68$ (fig. 2b), but G. yorkii and D. oleifera diverged before this duplication event with a median ortholog divergence of $K_s = 0.94$ (fig. 2c). This suggests that the ancient WGD in Polemoniaceae may not be shared with Ebenaceae or other more distantly related families of Ericales.

To test the hypothesized phylogenetic placement of this ancient WGD, we used the Multi-Taxon Paleopolyploidy Search (MAPS) approach (Li et al. 2018) with the G. yorkii genome; the S. latimeri and P. cuspidata transcriptomes, and transcriptomes or genomes of Fouquiéra macdougalii (Leebens-Mack et al. 2019), D. oleifera, Cornus floridana (Leebens-Mack et al. 2019), and Solanum lycopersicum (Hosmani et al. 2019) as outgroups. We found one significant burst of shared gene duplication that was statistically consistent with our simulations of WGD in the ancestry of the three Polemoniaceae species (fig. 2d and supplementary table S5, Supplementary Material online). However, we do not find any significant episodes of gene duplication at other nodes in the phylogeny.

To confirm that G. yorkii has undergone a single round of WGD after the ancient eudicot hexaploidy event, we compared the syntenic depth ratio between the genomes of G. yorkii and Vitis vinifera. We observed an overall two-
Table 1
Statistics of the Gilia yorkii Contig, Scaffold, and Transcriptome Assemblies

|                       | Genome Contig Assembly | Genome Scaffold Assembly | Transcriptome Assembly |
|-----------------------|------------------------|--------------------------|------------------------|
| Length (Mb)           | 2,753                  | 2,754                    | 192.63                 |
| % of estimated genome size | 98.29                | 98.32                    | NA                     |
| Contigs/scaffolds     | 3,947                  | 2,043                    | 96.691                 |
| Contig/scaffold N50 (Mb) | 2.54                | 285.77                   | 0.0022                 |
| Contig/scaffold L50   | 317                    | 5                        | 31,532                 |
| Longest contig/scaffold (Mb) | 18.30                | 374.98                   | 0.0099                 |
| Complete BUSCO genes (%) | 1,330 (96.7)          | 1,331 (96.8)             | 1,322 (96.1)           |
| Single copy (%)       | 1,264 (91.9)           | 1,271 (92.4)             | 233 (16.9)             |
| Complete, duplicate (%) | 66 (4.8)              | 60 (4.4)                 | 1,089 (79.2)           |
| Fragmented BUSCO genes (%) | 14 (1.0)             | 11 (0.8)                 | 12 (0.9)               |
| Missing BUSCO genes (%) | 31 (2.3)              | 33 (2.4)                 | 41 (3.0)               |
| Total BUSCO genes searched | 1,375                 | 1,375                    | 1,375                  |

To—one syntenic depth ratio between G. yorkii and V. vinifera (supplementary fig. 5S, Supplementary Material online). We further compared the G. yorkii genome with the other Ericales genomes from Actinidia eriantha (Tang et al. 2019), C. sinensis (Xia et al. 2020), D. oleifera, and Rhododendron simsii (Yang et al. 2020). We found a two-to-two syntenic depth ratio to C. sinensis, D. oleifera, and R. simsii (fig. 2e and supplementary fig. 5S, Supplementary Material online), but a two-to-four ratio with A. eriantha (fig. 2f), consistent with Actinidia having experienced an additional round of ancient WGD compared with other Ericales genomes (Shi et al. 2010; Huang et al. 2013; Wu et al. 2019). For comparison to G. yorkii, we also analyzed the syntenic depth ratios between V. vinifera and A. eriantha, C. sinensis, D. oleifera, or R. simsii. In A. eriantha, we found a four-to-one syntenic depth ratio to V. vinifera (supplementary fig. 5S, Supplementary Material online). In C. sinensis, D. oleifera, and R. simsii, we recovered a two-to-one ratio to V. vinifera (supplementary fig. 5S, Supplementary Material online). Combined with other results above, the syntenic evidence confirms a single round of ancient WGD in G. yorkii after the eudicot ancient hexaploidization event.

Gilia yorkii and G. capitata Are Morphologically Distinct Species that Form a Fertile F1 Hybrid

We used the chromosome-scale G. yorkii genome assembly to investigate the genetic basis of morphological traits that are polymorphic with G. capitata. Gilia yorkii is a narrow endemic in the Monarch Wilderness (CA) (Shevock and Day 1998). The inflorescence of G. yorkii is an open panicle with white flowers, typically producing only one solitary lateral flower before the production of a terminal flower (fig. 3). Unlike G. yorkii, G. capitata is a morphologically diverse species, with eight described subspecies distributed throughout the Pacific coast of North America (Grant 1950). The tight capitate inflorescence composed of up to 80 blue flowers forms a robust landing pad on which larger pollinating insects can alight (Grant and Grant 1965). The G. capitata inflorescence is a compound raceme which can produce ≥30 solitary lateral flowers before the apical meristem is consumed in a terminal flower (fig. 3). The tight capitate structure of this raceme results in part from the failure of internodes and pedicels to elongate (fig. 3b and c).

The phylogenetic relationships of leafy-stemmed gilias are poorly resolved, but current work suggests G. yorkii and the morphologically distinct Gilia achilleifolia are closely allied to a subset of G. capitata which is likely a polyphyletic taxon (Johnson 2007; Johnson and Porter 2017). We crossed G. yorkii to G. capitata from three different populations spanning multiple subspecies (see methods). All crosses produced vigorous hybrid F1 progeny, but two of these crosses resulted in sterile hybrids. One isolate, G. capitata (Cow Canyon), produced a vigorous and fertile F1 progeny. Nine chromosome pairs (2n = 2x = 18) were observed in meiotic pollen from these fertile hybrids (supplementary fig. S6, Supplementary Material online). No obvious lagging or bridge chromosomes were observed.

In addition to inflorescence architecture and flower color, G. capitata (Cow Canyon) was polymorphic with G. yorkii for many additional morphological features including leaf shape, floral shape (corolla size and shape, stamen exsertion), and calyx pubescence (wooly vs. stipitate glandular, not shown), as well as breeding system, with G. capitata being self-incompatible whereas G. yorkii is self-compatible. The F1 hybrid was self-compatible, and largely intermediate in all polymorphic characters (fig. 3).

QTL Mapping Reveals Candidate Genes for Major Effect Flower Color Locus and Complex Inheritance of Inflorescence Architecture

A population of 189 F2 individuals was generated from the selfing of a single F1 G. yorkii × G. capitata hybrid. Petal color (PC) was scored on a scale of 1–7, with one having no color (white) and seven having saturated blue/purple petals. All G. yorkii individuals displayed a score of 1, whereas all G. capitata individuals displayed a score of 7. PC of six F1
individuals was 5, intermediate between the parents. The F2 distribution of PC had distinct peaks at 1 and 5 (fig. 4a). This bimodal distribution is consistent with one to a few major loci controlling PC, and in combination with the intermediate F1 phenotype suggests that the *G. capitata* blue allele(s) are semidominant to the white *G. yorkii*.

Solitary flower number (SFN) refers to the number of solitary flowers below the terminal flower of the inflorescence and is a quantitative measure of inflorescence type with races having a high SFN and panicles a low SFN. Given the presence of both terminal and lateral inflorescences in all parental and F2 progeny, we scored SFN only for the terminal inflorescence in all individuals. In the inflorescence of *G. yorkii*, SFN was always 1, but ranged between 20 and 70 in *G. capitata*, with an average of 36. The F1 was intermediate between the parents, though strongly skewed toward *G. yorkii*, with a SFN average of 8. The F2 distribution was normal and centered around 8 or 9, with a range from 1 to 15 (fig. 4b). No F1 or F2 individuals approached the average SFN for *G. capitata*. The normal distribution of phenotypes suggests that SFN is controlled by multiple loci of smaller effect.

We used all 189 F2 individuals to generate a linkage map comprised 5,335 SNP markers on nine linkage groups. Positions of markers on the genetic map were highly colinear with their physical positions (supplementary fig. S7a, Supplementary Material online). A single marker on Gy8 was incongruent between the physical and genetic maps and was removed from further analyses. Although the majority of the physical map was represented in the genetic map, large regions of Gy3, Gy4, and Gy5 had no corresponding markers on the genetic map (supplementary fig. S7b, Supplementary Material online). The nonrandom filtering of these linked markers could be consistent with segregation distortion associated with either the self-incompatibility system of *G. capitata* or other genetic incompatibilities generated by the wide cross.

We performed a preliminary single-QTL analysis for the traits PC and SFN. PC is inherited as a relatively simple trait where candidate genes from the anthocyanin biosynthesis pathway are well characterized (Saigo et al. 2020). On the other hand, SFN, a quantitative measure of inflorescence type (panicle vs. raceme), correlates with an important morphological transition that is less well studied and likely involves a more complex set of loci.

With a threshold LOD score of 4.16 (*P* = 0.05) PC showed a single strong peak on Gy1 (LOD 30.0, *P* = 4.3e-32), with a minor peak on Gy4 (LOD 5.4, *P* = 3.2e-7), suggesting a single major QTL with a possible modest effect modifier (fig. 4c). The presence of a single major QTL regulating PC is consistent with the bimodal distribution of PC in the F2 population. An investigation of phenotypes conditioned on the most significant markers for these two peaks showed no evidence of epistasis (supplementary fig. S8, Supplementary Material online). We investigated annotated genes near the large-effect
QTL on Gy1 and found a tandem duplication of a flavanoid 3',5'-hydroxylase (F3'5'H) gene, an essential enzyme in the anthocyanin biosynthesis pathway shown previously to regulate flower color in other species (Holton et al. 1993; Zabala and Vodkin 2007; Hopkins and Rausher 2011; Smith and Rausher 2011; Moreau et al. 2012; Wessinger and Rausher 2015). No obvious deleterious mutations were observed in these genes, although a promoter or other regulatory mutation could not be ruled out. Although the causative polymorphism is still unclear, one or both members of this F3'5'H duplication are strong candidates for the loss of anthocyanin in G. yorkii.

In contrast to PC, SFN had a much more complex inheritance pattern, with significant peaks on several chromosomes, including Gy1, Gy3, Gy5, and Gy9 (fig. 4d), suggesting that the shift in inflorescence architecture is likely to be controlled by multiple, smaller effect QTL. Further analysis will be needed to determine relative effect sizes of these loci and to identify candidate genes. Nevertheless, these results confirm that a forward genetic approach is capable of identifying loci necessary to dramatically alter inflorescence architecture and thus highlight the promise of Gilia as a model for evo–devo.

**Discussion**

**Genome Assembly and WGDs**

The G. yorkii genome assembly reported here is the first for Polemoniaceae and adds to a growing number of assemblies within Ericales. Like many flowering plants, evidence indicates
that WGDs occurred throughout the history of the Ericales. Putative WGD peaks in gene age distributions are found in all Ericales transcriptomes (Shi et al. 2010; Leebens-Mack et al. 2019; Larson et al. 2020), and multiple ancient WGDs have been confirmed by syntenic analyses (Hosmani et al. 2019; Tang et al. 2019; Larson et al. 2020; Suo et al. 2020; Xia et al. 2020; Yang et al. 2020). Recent phylogenomic analyses also suggest a WGD occurred near the origin of Ericales, although the exact timing remains unsolved (Leebens-Mack et al. 2019; Larson et al. 2020; Wang et al. 2021). Resolving WGDs in Ericales has been difficult due to substantial rate heterogeneity and rapid radiation (Leebens-Mack et al. 2019; Larson et al. 2020; Wang et al. 2021). Using our G. yorkii genome, we found evidence for one WGD based on synteny, Ks plot, and phylogenomic approaches. Our syntenic evidence for WGD in the Polemoniaceae confirms the putative WGD previously found in Polemoniaceae based on transcriptomes (Leebens-Mack et al. 2019; Larson et al. 2020). Although a recent phylogenomic study suggests an ancestral Ericales WGD (Wang et al. 2021), our syntenic and phylogenomic analyses indicate that the Polemoniaceae WGD was not shared with other Ericales and the similar Ks peaks in Polemoniaceae, Fouquieriaceae, and Ebenaceae may be three independent WGDs. This conflicting evidence suggests that the exact phylogenetic placement of a single ancestral Ericales WGD or multiple independent ancient WGDs remains to be resolved and will require additional genome assemblies from key families, such as from Primulaceae and Sapotaceae.

Gilia as a Promising Evo–Devo Model

The chromosome-scale genome assembly for G. yorkii provides a crucial reference for functional dissection of polymorphic traits in Gilia. Although the Polemoniaceae includes some popular horticultural species such as Phlox, it lacks any food or forage species which likely accounts for the previous lack of attention at the genomic level. The diverse ecology and morphology represented by members of the genus Gilia have made it an attractive model for multiple evolutionary studies (Schoen 1982, 1983; Morrell and Rieseberg 1998; Morrell et al. 2000; Takebayashi et al. 2006), and the leafy-stemmed gilias in particular were a classic model for Verne and Alva Grant’s pioneering work on plant evolution and speciation (Grant 1950, 1952, 1953, 1954, 1956, 1966, 1981; Grant and Grant 1956). The complete Gilia yorkii genome reference will facilitate re-analysis of multiple questions in this group with modern genetic approaches. Additionally, Gilia yorkii has many traits commonly sought in a model genetic system including relatively short annual life cycle (~3 months), self-compatibility, simple crossing, copious seed production, and minimal culturing requirements. We show that despite striking morphological divergence, G. yorkii and G. capitata form a fertile F1 hybrid and F2 progeny that segregate the traits of their parents. This positions Gilia alongside other plant evo–devo models (e.g., Erythranthe, formerly Mimulus, Wu et al. 2008; Aquilegia, Kramer 2009; Penstemon, Wessinger et al. 2014; Petunia, Klahre et al. 2011) in which a forward genetic QTL approach can be used to identify the molecular causes of species differences. Similar approaches have proven especially informative in animal evo–devo (Shapiro et al. 2004; Jeong et al. 2008). Plants hybridize much more frequently and promiscuously than animals, opening up a host of diverse morphologies to dissect at the genetic level. With the growing ease of genome assembly and high-throughput genotyping, many other morphologically divergent but interfertile species pairs are similarly poised for development.
Preliminary Mapping of QTL for Flower Color and Inflorescence Architecture

Although hybridization is common, not all hybrids are fertile and among those that are, structural rearrangements can inhibit crossing over and thus limit the resolution of genetic mapping (Fishman and Sweigart 2018). Our preliminary interspecies genetic map recovered the majority of the physical map of G. yorkii. This result, in combination with the apparently normal segregation of meiotic chromosomes during microsporogenesis, suggests that crossing over is largely undisrupted between these genomes, although we cannot rule out local interference due to smaller scale rearrangements. Even with a small mapping population, the major PC locus on Gy1 mapped precisely, allowing the identification of a strong pair of candidate genes. Despite the high resolution of most chromosomes, large regions of Gy3, Gy4, and Gy5 were absent from our genetic map. These regions contained a large proportion of markers with segregation distortion that were filtered before creating the genetic map. There are many possible causes of segregation distortion (Hall and Willis 2005) and it is still unclear which of these are contributing, although loci linked to the self-incompatibility locus (or loci) from G. capitata were possibly selected against. In addition to distorted markers, many markers were filtered because they did not cosegregate consistently with adjacent markers. Considering the wide cross and highly repetitive nature of the G. yorkii genome, a genotype by sequencing (GBS) approach to identify polymorphic markers may not have been ideal. The random genome reduction resulting from GBS is necessarily biased to intergenic and repetitive regions, and thus many of the polymorphic SNPs are likely to be from repetitive regions and may not be homologous across these species. A different marker technology, such as targeted capture (Grover et al. 2012) of genes across the G. yorkii genome may help alleviate these concerns in future QTL mapping experiments.

We identified a tandem gene duplication as candidates for the major PC QTL. It is still unclear if this duplication is present in both G. capitata and G. yorkii, and the lack of obvious deleterious mutations makes it difficult to speculate how they may cause the loss of anthocyanin in G. yorkii. Thus, either gene, or perhaps even both genes together, are potentially causative. These genes encode orthologs of F3′5′H that catalyzes hydroxylation of dihydroeinorcinol (DHQ) to dihydroevincinidin (DHM) required for the formation of blue/purple anthocyanidins. In multiple studies of PC change in wild species, loss of F3′5′H activity results in a blue to red color shift (Hopkins and Rausher 2011; Smith and Rausher 2011; Wessinger and Rausher 2015), unlike the shift from blue to white seen in G. yorkii. It is not immediately clear why loss of F3′5′H would create a white as opposed to red flower in Gilia, but the PC response to mutations in the anthocyanin biosynthesis pathway can be complex (Berardi et al. 2021) and an F5′3′H ortholog in soybean mediates a purple to white transition (Zabala and Vodkin 2007). More work will be necessary to confirm the molecular nature of the flower color change. Beyond this specific cross, blue/white flower polymorphism occurs in native accessions of G. capitata (Grant 1950), as well as more broadly in the leafy-stemmed gillas (Grant 1954) representing an attractive system to investigate molecular parallelism versus convergence in flower color.

Unlike flower color polymorphism, which has been addressed in multiple wild species, inflorescence architecture has received little attention (Prusinkiewicz et al. 2007; Lippman et al. 2008). Angiosperm inflorescence architecture is highly diverse (Rickett 1944; Stebbins 1973; Weberling 1989) and contributes to fecundity, making it important both ecologically and agriculturally (Wyatt 1982; Fishbein and Venable 1996; Friedman and Harder 2004; Jiao et al. 2010; Bommert et al. 2013; Gao et al. 2015). Models of inflorescence architecture (Lijegren et al. 1999; Prusinkiewicz et al. 2007; Park et al. 2012) involve competitive interaction of genes that promote or delay acquisition of floral meristem identity and are based on induced mutations or natural variants in crop species that typically generate quantitative as opposed to qualitative shifts in inflorescence type. The inflorescence shift from G. capitata to G. yorkii is remarkable in that it involves changes from a raceme to a panicle, which along with cymes characterize the majority of inflorescence diversity in the angiosperms (Prusinkiewicz et al. 2007). Developmental modeling and surveys of inflorescence architecture in the angiosperms show that transitions from racemes to panicles to cymes are particularly rare, whereas transitions from racemes to panicles appear to be more common (Prusinkiewicz et al. 2007; Harder and Prusinkiewicz 2013). Unlike cymes, both panicles and racemes have an indeterminate apical meristem that produces multiple lateral meristems. Although definitions of raceme and panicle are not uniform (Rickett 1944; Prenner et al. 2009; Endress 2010), a simplified distinction can be reduced to the fate of lateral meristems. Specifically, in a raceme the lateral meristems take on a determinate floral fate immediately, whereas in a panicle, the lateral meristems maintain an indeterminate inflorescence meristem identity, only producing a determinate lateral flower after several rounds of branching. Developmental modeling indicates that the raceme panicle transition is quantitative rather than qualitative with multiple intermediate states (Prusinkiewicz et al. 2007; Harder and Prusinkiewicz 2013). In Gilia, SFN is a robust measure of the quantitative raceme-panicle continuum and is normally distributed in F2 segregating progeny showing a range of intermediate morphologies. SFN is regulated by at least four QTL, suggesting that multiple loci of smaller effect mediate the shift from panicle to raceme. Identifying these genes will require the development of new genetic mapping populations such as recombinant inbred and near isogenic lines.

Beyond inflorescence architecture, G. yorkii and G. capitata differ in other morphological traits that could be dissected...
with a QTL-cloning approach including leaf, floral organ, and trichome morphology, and presence or absence of a self-incompatibility system. Many of these traits are also polymorphic more broadly in the leafy-stemmed gilias as well. The *G. yorkii* genome will provide a platform to dissect the genetic basis of these traits in future studies.

**Materials and Methods**

**Plant Material and Mapping Populations**

*Gilia yorkii* Shevock and A.G. Day was grown from seed collected from an herbarium sheet (BRY-614948). Plants were self-fertilized and an inbred line produced by single-seed descent for eight generations. *Gilia capitata* Sims was grown from seed collected in Cow Canyon near Mt. Baldy, San Bernardino Co., CA (BRY-642206), and propagated by sibling crossing. F1 hybrids were generated by emasculating and crossing *G. yorkii* × *G. capitata*. A single seed source of *G. yorkii* was used to which *G. capitata* from three different populations (*G. capitata* 9228, *G. capitata* Baja, and *G. capitata* Cow Canyon) were crossed. F1 hybrid progeny (a minimum of three) for each *G. yorkii* × *G. capitata* cross were grown to maturity. Although all F1 hybrids were vigorous, only the cross with *G. capitata* Cow Canyon produced a fertile hybrid, and these F1 plants were self-fertilized to generate an F2 population. PC was determined by averaging by eye more than five flowers from each plant and using a printed scale of graded white to purple shades as a standard. SFN was determined for only the cross with *G. capitata* Cow Canyon produced a fertile hybrid, and these F1 plants were self-fertilized to generate an F2 population. Flower buds, and roots. One SMRT cell was used to generate 198 Gb of continuous long read (CLR) sequencing data.

**Sequencing**

Illumina sequencing of *G. yorkii* was performed on HiSeq X by Novogene using 550-bp insert DNA libraries prepared at Brigham Young University’s DNA Sequencing Center (BYUDNASC), generating a total of 161.5 Gb of 150-bp paired-end sequencing data. PacBio library preparation and sequencing of HWM *G. yorkii* DNA was performed on PacBio Sequel by the BYUDNASC. Three SMRT cells were used to generate 198 Gb of continuous long read (CLR) sequencing data. PacBio Iso-Seq library preparation and sequencing were performed at the BYUDNASC using a single RNA sample created by pooling equal amounts of RNA extracted individually from leaf primordia, mature leaves, shoot apex, mature flowers, flower buds, and roots. One SMRT cell was used to generate 2,341,411 circular consensus sequencing reads (193.3 Mb of data).

**Whole-Genome Assembly and Annotation**

Illumina sequencing reads were trimmed using Trimmomatic to remove leading and trailing bases with a quality score below 20 or average per-base quality of 20 over a four-nucleotide sliding window; trimmed reads shorter than 75 nucleotides in length were removed. The frequency of kmers in the sequencing data was calculated using Jellyfish (Marçais and Kingsford 2011), and the genome size was estimated based on kmer frequency using GenomeScope (Ranallo-Benavidez et al. 2020) with kmer length = 21, read length = 150, and kmer max coverage = 1,000,000. PacBio CLR reads were assembled into contigs using Canu v1.9 (Koren et al. 2017) with coMhapSensitivity = normal, corOutCoverage = 40, and ovlMerThreshold = 500. SNP and Indel correction were performed with the PacBio CLR reads in two successive rounds using the Arrow algorithm in the
PacBio SMRT Tools package v7.0, and additional polishing was performed with the Illumina reads (Indel correction only) in one round using Pilon v1.9 (Walker et al. 2014). The assembly was scaffolded using in vivo Hi-C data by Phase Genomics as previously described (Lightfoot et al. 2017), using 228,955,144 pairs of Illumina sequencing reads. Juicebox (Rao et al. 2014; Durand et al. 2016) was used to manually correct scaffolding errors.

PacBio Iso-Seq reads were assembled into a transcriptome using the IsoSeq v3 pipeline in the PacBio SMRT Tools package, including using pbmm2 to align the Iso-Seq reads to the scaffolded genome and IsoSeq v3 to collapse the transcripts. Completeness of the Hi-C scaffolded genome and transcriptome assembly were assessed using BUSCO v4 (Simão et al. 2015) with the embryophytaodb10 data set.

Repeat sequences were identified and classified using RepeatModeler (Smit and Hubley 2008) and RepeatMasker (Smit et al. 2013), respectively, using RepBase database version 20181026. Genes were annotated using MAKER2 (Holt and Yandell 2011) using the Iso-Seq assembly as transcript evidence. Alternative EST and protein evidence were provided by sequences from Actinidia chinensis (Wu et al. 2019) and Camellia sinensis (Wei et al. 2018). The uniprot-sprot database (downloaded September 25, 2018) was also used as alternative protein evidence. An initial gene annotation was performed using Gila-specific AUGUSTUS (Stanke et al. 2004) gene prediction models and Arabidopsis thaliana SNAP (Korf 2004) gene models provided to MAKER, and tRNA genes were predicted with tRNAscan-SE (Lowe and Eddy 1997). Telomeres were identified by performing a BLAST search of four repeats of the canonical telomere sequence (TTAGGG) (Richards and Ausubel 1988). The chromosomal positions of BLAST hits with 100% identity, as well as the chromosomal positions of annotated genes, were plotted using karyoplotR (Gel and Serra 2017).

**Analysis of Ancient WGDs**

The DupPipe pipeline was used to construct gene families and estimate the age distribution of gene duplications (Barker et al. 2008, 2010). DNA sequences were translated and open reading frames (ORFs) identified by comparing the GeneWise (Birney et al. 2004) alignment to the best-hit protein from a collection of proteins from 25 plant genomes from Phytozome (Goodstein et al. 2012). Protein-guided DNA alignments were used to align nucleic acid sequences while maintaining the ORFs. Synonymous divergence (Ks) was estimated using PAML with the F3X4 model (Yang 2007) for nodes in the gene trees. Mixture modeling was used to identify peaks consistent with a potential WGD and estimate their median paralog Ks values. Significant peaks were identified using a likelihood ratio test in the boot.comp function of the package mixtools in R (Benaglia et al. 2009).

**Estimating Orthologous Divergence**

Synonymous divergence of orthologs was estimated among pairs of species that may share a WGD based on their phylogenetic position and evidence from the within-species Ks plots. The mean and median synonymous divergence of orthologs were estimated with The RBH Orthologue pipeline (Barker et al. 2010) and compared with the synonymous divergence of inferred paleopolyploid peaks. Orthologs were identified as reciprocal best BLAST hits in pairs of transcriptomes. Using protein-guided DNA alignments, the pairwise synonymous divergence for each pair of orthologs was estimated using PAML with the F3X4 model (Yang 2007).

**Genome Synteny Analyses**

Genomic collinearity blocks for intra- and interspecies comparisons for G. yorkii and other genomes were identified by MCScan (Tang et al. 2008). All-against-all LAST (Kiebsa et al. 2011) hits with a distance cutoff of ten genes were chained, requiring ≥ five gene pairs per syntenic block. The syntenic “depth” function in MCScan was applied to estimate the duplication history in respective genomes. Genomic synteny was visualized by the Python version of MCScan (Tang et al. 2008).

**Multi-tAxon Paleopolyploidy Search Analyses**

Multi-tAxon Paleopolyploidy Search (Li et al. 2015, 2018) was used to determine the timing of WGD in G. yorkii. Orthologous groups were obtained from OrthoFinder (Emms and Kelly 2015) using default parameters, retaining only gene families that with ≥ 1 gene copy from each taxon. Trees for 8,093 gene families constructed by PASTA (Miranab et al. 2015) were analyzed by MAPS. Both null and positive simulations of the background gene birth and death rates were performed to compare with the observed number of duplications at each node, using WGDgc (Rabier et al. 2014) to generate gene birth (λ) and death (μ) rates for null simulations. Gene count data of each gene family were obtained from OrthoFinder (Emms and Kelly 2015). The estimated parameters (λ = 0.007; μ = 0.014) were configured in MAPS, and the gene trees were simulated within the species tree using “GuestTreeGen” from GenPhyloData (Sjöstrand et al. 2013). For each species tree, 3,000 gene trees were simulated with at least one tip per species: 1,000 gene trees respectively at 0.5x, 1x, and 3x the estimated λ and μ, according to the 1KP analyses (Leebens-Mack et al. 2019; Li and Barker 2020). Random resampling of 1,000 trees without replacement from the total pool of gene trees 100 times provided a measure of uncertainty on the percentage of subtrees at each node. Fisher’s exact test was used to identify significant increases in gene duplication compared with a null simulation. For positive simulations, gene trees were simulated using the same methods described above. However, WGDs
were incorporated at the location in the MAPS phylogeny resulting in significantly larger numbers of gene duplications at this phylogenetic position compared with the null simulation. At least 20% of the genes were retained following the simulated WGD (Leebens-Mack et al. 2019; Li and Barker 2020).

**Genotyping-by-Sequencing**

Approximately 30 mg of fresh leaf tissue was lyophilized for G. yorkii, G. capitata, a newly generated F1 hybrid, and each of the 189 F2 progeny in the G. yorkii × G. capitata population. Samples were sent to Freedom Markers, and genotyped using the modified TGBS protocol (Ott et al. 2017). Libraries were prepared with the restriction enzyme Bsp1286I and sequenced using Illumina HiSeq X, generating a total of 377,694,355 pairs of reads. Reads were trimmed by removing bases with PHRED quality scores less than 15 (out of 40) using Lucy (Chou and Holmes 2001). Trimmed reads were mapped to the polished, scaffolded reference G. yorkii genome using GSNAP (Wu and Nacu 2010). Reads with more than two mismatches every 36 bp or more than five mismatches every 75 bp were removed prior to genotype calling.

**Genetic Map**

Polymorphic sites were identified as any site that differed from the reference in at least one sample. Genotypes at each site were identified using the following criteria: if the most common allele was supported by ≥80% of aligned reads and ≥5 unique reads, the SNP was classified as homozygous. SNPs were considered heterozygous if each of the two most common alleles were supported by ≥30% of aligned reads and ≥5 unique reads. Polymorphisms in the first and last 3 bp of reads were ignored, and a PHRED quality score of ≥20 was required for each SNP.

These SNPs were then filtered accordingly: minimum calling rate ≥50%, allele number = 2, number of genotypes ≥2, minor allele frequency ≥10%, and heterozygosity rate range of 10–65%, resulting in 73,482 SNPs used to create a linkage map. Linkage groups were formed using the R/qtl package in R version 3.6.2 (Broman et al. 2003; Broman and Sen 2009). The 5,335-marker data set (supplementary table S1, Supplementary Material online) was used for single-QTL analyses using an extended Haley–Knott regression analysis in RStudio using R/qtl in R version 3.6.2. The LOD threshold of 4.16 was determined using the 95th percentile of LOD scores for all markers across all chromosomes, with 100,000 permutations.

**QTL Analysis**

The 5,335-marker data set (supplementary table S1, Supplementary Material online) was used for single-QTL analyses using an extended Haley–Knott regression analysis in RStudio using R/qtl in R version 3.6.2. The LOD threshold of 4.16 was determined using the 95th percentile of LOD scores for all markers across all chromosomes, with 100,000 permutations.

**Supplementary Material**

Supplementary data are available at Genome Biology and Evolution online.

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**Data Availability**

The data that support the findings of this study are openly available in GenBank at https://www.ncbi.nlm.nih.gov/genbank/ (last accessed March 3, 2022), BioProject PRJNA729797. Additional information is provided in supplementary table S6, Supplementary Material online. The Gila yorkii genome assembly and annotation are available on CoGe (https://genomevolution.org/coge/, last accessed March 3, 2022) under Genome ID 62042.

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