

1 **Dissecting the basis of novel trait evolution in a radiation with widespread**
2 **phylogenetic discordance**

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17 RRH: Trait evolution under widespread discordance

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23 **ABSTRACT**

24 Phylogenetic analyses of trait evolution can provide insight into the evolutionary
25 processes that initiate and drive phenotypic diversification. However, recent
26 phylogenomic studies have revealed extensive gene tree-species tree discordance, which
27 can lead to incorrect inferences of trait evolution if only a single species tree is used for
28 analysis. This phenomenon—dubbed “hemiplasy”—is particularly important to consider
29 during analyses of character evolution in rapidly radiating groups, where discordance is
30 widespread. Here we generate whole-transcriptome data for a phylogenetic analysis of 14
31 species in the plant genus *Jaltomata* (the sister clade to *Solanum*), which has experienced
32 rapid, recent trait evolution, including in fruit and nectar color, and flower size and shape.
33 Consistent with other radiations, we find evidence for rampant gene tree discordance due
34 to incomplete lineage sorting (ILS) and several introgression events among the well-
35 supported subclades. Since both ILS and introgression increase the probability of
36 hemiplasy, we perform several analyses that take discordance into account while
37 identifying genes that might contribute to phenotypic evolution. Despite discordance, the
38 history of fruit color evolution in *Jaltomata* can be inferred with high confidence, and we
39 find evidence of *de novo* adaptive evolution at individual genes associated with fruit
40 color variation. In contrast, hemiplasy appears to strongly affect inferences about floral
41 character transitions in *Jaltomata*, and we identify candidate loci that could arise either
42 from multiple lineage-specific substitutions or standing ancestral polymorphisms. Our
43 analysis provides a generalizable example of how to manage discordance when
44 identifying loci associated with trait evolution in a radiating lineage.

45

46 *Key words:* phylogenomics, rapid radiation, hemiplasy, convergence, *Jaltomata*, *Solanum*

47

48 **INTRODUCTION**

49 Phylogenies contribute to our understanding of the evolutionary history of traits
50 (Felsenstein, 1985). When the patterns of relationship among species is known, robust
51 inferences about character state evolution can be made, including the number of times a
52 character evolved, the direction of character evolution, and the most likely ancestral
53 character state. Phylogenies can also reveal whether lineages with similar phenotypic
54 traits have evolved these via independent evolution (convergence or parallelism) or
55 whether a single origin is more likely (Wake et al., 2011). The recent use of whole
56 genomes or transcriptomes to make phylogenetic inferences from thousands to millions
57 of sites (“phylogenomics”), has succeeded in its aim of generating species trees with high
58 levels of statistical support. However, other genome-wide analyses have begun to reveal
59 unexpected complexities in the evolutionary history of rapidly radiating lineages—
60 including widespread gene tree discordance due to incomplete lineage sorting and/or
61 introgression (Degnan & Rosenberg, 2009). This frequent discordance among individual
62 gene trees can amplify incorrect inferences of trait evolution on even well-supported
63 species trees. In particular, when a trait is determined by genes whose topologies do not
64 match the species topology, incorrect inferences of homoplasy (independent evolution of
65 the same character state) are substantially elevated—a phenomenon known as
66 ‘hemiplasy’ (Avice & Robinson, 2008; Hahn & Nakhleh, 2016; Storz, 2016). Because
67 understanding trait evolution—including the underlying genetic changes—is of particular
68 interest in species radiations, extra care must be taken to consider and account for the
69 influence of hemiplasy in these cases.

70 The fraction of the genome affected by hemiplasy will depend upon the amount
71 and sources of gene tree discordance in a clade. In rapidly radiating species groups,
72 widespread discordance has been attributed to the effects of both incomplete lineage
73 sorting (ILS) and introgression between lineages (Degnan & Rosenberg, 2009). ILS
74 affects gene tree topologies when segregating ancestral variation is maintained through
75 consecutive speciation events (Maddison, 1997). Because the effect of ILS is proportional
76 to ancestral population size, and inversely proportional to the time between speciation
77 events (Pamilo & Nei, 1988), ILS is expected to be particularly exaggerated in radiations
78 where a diverse ancestral population undergoes rapid speciation. Indeed, gene tree
79 discordance has been noted for a substantial fraction of the genome in rapidly radiating
80 groups, including the *Drosophila simulans* sub-clade (Garrigan et al., 2012), African
81 cichlid fishes (Brawand et al., 2014), wild tomatoes (Pease et al., 2016), and the genus
82 *Arabidopsis* (Novikova et al., 2016). When there is introgression, discordance emerges
83 because genes that are introgressed among lineages will show historical patterns of
84 relatedness that differ from the loci in the genome into which they are introduced.
85 Substantial introgression has also been identified among rapidly radiating lineages
86 through genome-wide analysis, including in *Xiphophorus* fishes (Cui et al., 2013),
87 *Heliconius* butterflies (Martin et al., 2013), Darwin's finches (Lamichhaney et al., 2015),
88 and *Anopheles* mosquitoes (Fontaine et al., 2015).

89 Both ILS and introgression contribute to hemiplasy because they cause a
90 proportion of gene trees to disagree with the species tree (Avice & Robinson, 2008; Hahn
91 & Nakhleh, 2016; Storz, 2016). Specifically, the probability of hemiplasy is expected to
92 be 1) proportional to the fraction of gene trees that are discordant with the species tree;

93 and 2) negatively correlated with the branch length leading to clades with similar
94 phenotypes (Hahn & Nakhleh, 2016). A higher proportion of discordant gene trees
95 increases the probability that a character of interest is underpinned by genes that have a
96 tree topology that differs from the species tree; shorter branch lengths increase the chance
97 of incorrectly inferring homoplasy, as they leave relatively little time for convergent
98 evolution to happen (Hahn & Nakhleh, 2016). Both conditions are expected to be
99 exaggerated specifically in rapidly diversifying groups. Therefore, in these cases mapping
100 characters onto a single species tree has a substantially elevated risk of incorrectly
101 inferring the number of times a trait has evolved and the timing of trait changes (Avisé &
102 Robinson, 2008; Hahn & Nakhleh, 2016; Storz, 2016). Hemiplasy also affects inferences
103 about the specific loci inferred to underlie trait transitions because, when ILS or
104 introgression are common, the substitutions underlying trait transitions may occur on
105 gene trees that are discordant with the species tree (Mendes et al., 2016). Accordingly,
106 genome-wide analyses must take into account the extent and distribution of ILS and
107 introgression if they are to accurately infer the number and timing of evolutionary
108 changes in specific traits, and the genes underlying these changes.

109 In this study, we used genome-wide data to investigate the morphologically and
110 ecologically diverse plant genus *Jaltomata*, in which several key trait transitions appear
111 to have occurred in parallel (Miller et al., 2011), and have been inferred to be
112 independent convergent responses to similar selective pressures. However, because trait
113 diversification has occurred in a relatively short period in this group, the probability of
114 hemiplasy is also expected to be elevated. Our main goals were to assess the timing of
115 lineage and trait diversification in the group, and to identify sources of genetic variation

116 that potentially contribute to rapid trait diversification in *Jaltomata*, while taking into
117 account the potential for hemiplasy. To do so, we generated a clade-wide whole-
118 transcriptome dataset and explicitly evaluated alternative scenarios to explain trait
119 evolution by: 1) reconstructing phylogenetic relationships among target species, and
120 evaluating the extent of discordance with the resulting inferred species tree; 2) evaluating
121 patterns of trait variation and evolution in key reproductive (flower and fruit) characters,
122 in the context of best and least supported nodes in this tree; and, 3) evaluating specific
123 scenarios of the genetic changes associated with this trait evolution, in order to identify
124 candidate loci that might be causally responsible. Our results imply two different
125 scenarios of trait evolution for fruit color versus floral traits, reflecting the different
126 amounts of hemiplasy associated with the two traits. While fruit color evolution in
127 *Jaltomata* could be confidently inferred—along with potential *de novo* molecular changes
128 on the relevant evolutionary branches—inferring the history of floral trait evolution and
129 the potential contributing loci requires more careful treatment that considers the high
130 probability of hemiplasy.

131

132 **MATERIALS AND METHODS**

133 Study system

134 The plant genus *Jaltomata* includes approximately 60-80 species, distributed from
135 the southwestern United States through to the Andes of South America (Mione, 1992;
136 Mione et al., 2015) (Figure 1). It is the sister genus to *Solanum*, the largest and most
137 economically important genus in the family Solanaceae (Olmstead et al., 2008; Särkinen
138 et al., 2013). Species of *Jaltomata* live in a wide range of habitats, and are phenotypically

139 diverse in vegetative, floral, and other reproductive traits (Mione 1992; Kostyun & Moyle
140 2017). Floral diversity is particularly pronounced in *Jaltomata*. In comparison to closely
141 related clades (including *Solanum*, *Capsicum*, and *Lycianthes*) which predominately have
142 ‘flattened’ rotate corollas (petals) (Knapp, 2010), *Jaltomata* species exhibit a variety of
143 corolla shapes, including rotate, as well as campanulate and tubular (Miller et al., 2011).
144 All *Jaltomata* species also produce at least some nectar, including noticeably red- or
145 orange-colored nectar in some lineages, while nectar is not produced by species in
146 *Solanum*.

147 Species also differ in fruit color, and fruit color variation appears to characterize
148 major subgroups within the genus as separate dark purple, red, and orange-fruited clades
149 (Miller et al., 2011; Särkinen et al., 2013). Several species also have green fruit at
150 maturity, although these lineages appear to be distributed across the three major
151 *Jaltomata* clades, suggesting multiple convergent losses of fruit pigment (Miller et al.,
152 2011). The first molecular phylogeny of this genus (Miller et al., 2011) was inferred from
153 a single gene (*waxy*), and indicates that the lineage of species with red fruits is sister to
154 the rest of the genus. However, a more recent study using seven loci (5 plastid and 2
155 nuclear, including *waxy*) showed a conflicting topology, with purple-fruited lineages
156 sister to the remaining groups, and red-fruited lineages more closely related to lineages
157 with orange fruits (Särkinen et al., 2013). The inconsistency between the two studies
158 might be the result of using few loci, or of reconstructions performed with loci that have
159 different evolutionary histories.

160

161 *RNA preparation, sequencing, and transcript assembly*

162 We chose 14 target *Jaltomata* species that are distributed across the three
163 previously identified major clades (Miller et al., 2011), and that span representative floral
164 diversity within the genus (Figure 2A, Table S1). Tissues for RNA extraction included
165 seven reproductive tissues (ranging from early bud, to mature pollinated flower, to early
166 fruit) and four vegetative tissues (roots, early leaf buds, and young and mature leaves),
167 from a single representative individual of each target species (see Supplementary text).
168 All sampled individuals were housed at the Indiana University research greenhouse,
169 under standardized temperature (15-20°C), watering (twice daily), and lighting (16-hour
170 days) conditions.

171 Tissue collection and RNA extraction followed Pease et al. (2016): briefly, tissue
172 was collected into pre-chilled tubes under liquid nitrogen, each sample was individually
173 ground under liquid nitrogen, and RNA was extracted from <100mg ground tissue using
174 the Qiagen Plant RNeasy kit. RNA quality/quantity was checked via Nanodrop (Thermo
175 Fisher Scientific); qualified samples of >50ng/uL with 260/280 and 260/230 between 1.8-
176 2.0 were brought to the IU Center for Genomics and Bioinformatics (CGB) for library
177 preparation. Separate reproductive and vegetative libraries for RNA-seq were prepared
178 by pooling equi-molar RNA samples from all reproductive tissues, and all vegetative
179 tissues, respectively, for each species. Both reproductive and vegetative libraries were
180 prepared for all species except for *J. grandibaccata*, for which only vegetative RNA
181 could be obtained.

182 Libraries were sequenced using 100-bp paired-end reads in a single lane of
183 Illumina Hi-seq 2000 (San Diego, CA, USA). Raw paired-end reads were filtered for
184 quality using the program Shear (<https://github.com/jbpease/shear>) by removing low

185 quality reads and ambiguous bases, and trimming adapter ends (see Supplementary text).
186 The retained reads (length >50 bps) from vegetative and reproductive transcriptomes of
187 the same species were combined prior to assembly using Trinity with the default settings
188 (Grabherr et al., 2011). The open reading frame of each assembled transcript was
189 predicted using TransDecoder v.2.0.1 with default settings (Haas et al., 2013). All the
190 predicted protein-coding sequences within each *Jaltomata* species transcriptome were
191 reduced using CD-HIT v4.6 with -c 0.99 -n 10 (Fu et al., 2012). Each sequence in the
192 assembled transcriptome was presented as a haploid representative of particular transcript.
193 To include domestic tomato (*Solanum lycopersicum*) as the outgroup in the following
194 analyses, we also downloaded the annotated tomato protein-coding sequences from
195 SolGenomics (<ftp://ftp.solgenomic.net>).

196

197 Protein-coding gene ortholog identification

198 To infer orthologous gene clusters, we followed a pipeline designed for
199 transcriptome data in non-model species, that begins with an all-by-all BLAST search
200 followed by several steps that iteratively split sub-clusters of homologs at long internal
201 branches, until the subtree with the highest number of non-repeating/non-redundant taxa
202 is obtained (Yang et al., 2015; Yang & Smith, 2014) (see Supplementary text, Figure S1).
203 For the primary analyses, our homologous clusters were required to include a *S.*
204 *lycopersicum* (tomato) homolog in each cluster. For one of our downstream analyses
205 (molecular evolution on the basal branch leading to *Jaltomata*; see below) we also used
206 *Capsicum annuum* (pepper) sequence data. To do so, we added the *C. annuum* sequences
207 if the tomato sequence in the orthologous cluster had an identified 1-to-1 ortholog in a

208 pepper gene model (<http://peppersequence.genomics.cn>).

209 We prepared multiple sequence alignments of orthologous genes using the
210 program GUIDANCE v.2.0 (Sela et al., 2015) with PRANK v.150803 (Löytynoja &
211 Goldman, 2005) as the alignment algorithm, with codons enforced and ten bootstrap
212 replicates. As a final quality check, we further removed poorly aligned regions using a
213 sliding window approach that masked any 15-bp window from alignment if it had more
214 than three mismatches (indels/gaps were not counted) between ingroup sequences, or
215 had more than five/seven mismatches when tomato/pepper sequences were included.
216 After this process, any alignment with more than 20% of its sequence masked was
217 removed from the analysis. The resulting sequence alignments were converted to the
218 Multisample Variant Format (MVF), and then genetic distances were computed in all
219 possible pairs of species using the program MVFtools (Pease & Rosenzweig, 2015).

220

221 Estimating the amount of shared variation

222 To quantify the amount of variation shared among species and subclades in
223 *Jaltomata*, the reads from all 14 species were mapped to the reference tomato genome
224 (The Tomato Genome Consortium, 2012) using STAR v2.5.2 (Dobin et al., 2013). SAM
225 files generated were converted to sorted BAM files using SAMtools v. 0.1.19 (Li et al.,
226 2009). SAMtools *mpileup* was then used to call alleles from the BAM files for all
227 lineages. VCF files were processed into MVF files using *vcf2mvf* from the MVFtools
228 package (Pease & Rosenzweig, 2015), requiring non-reference allele calls to have Phred
229 scores ≥ 30 and mapped read coverage ≥ 10 . Based on the MVF files, the numbers of
230 variant sites shared between different subclades of *Jaltomata* species were counted.

231

232 *Phylogenetic analysis*

233 We used four different, but complementary, inference approaches to perform
234 phylogenetic reconstruction: 1) maximum likelihood applied to concatenated alignments;
235 2) consensus of gene trees; 3) quartet-based gene tree reconciliation; and 4) Bayesian
236 concordance of gene trees. Because these four approaches use different methods to
237 generate a phylogeny, we applied all four to evaluate the extent to which they generated
238 phylogenies that disagreed, as well as to identify the specific nodes and branches that
239 were robust to all methods of phylogenetic reconstruction. For the concatenation
240 approach, we first aligned all orthologous genes (n=6431), and then used those
241 alignments to build a supermatrix of sequences (6,223,350 sites in total). The species tree
242 was then inferred by maximum likelihood using the GTRCAT model in RAxML v8.23
243 with 100 bootstraps (Stamatakis, 2006). We also inferred chromosome-concatenated
244 phylogenies with this method. The other three methods (i.e. consensus, quartet-based, and
245 Bayesian concordance) infer species relationships based on gene trees. First, we inferred
246 the majority rule consensus tree with internode certainty (IC) and internode certainty all
247 (ICA) support scores using RAxML with the option for Majority Rule Extended
248 (Salichos & Rokas, 2013). Second, we inferred a quartet-based estimation of the species
249 tree by using the program ASTRAL v.4.10.9 with 100 bootstraps (Mirarab & Warnow,
250 2015). All 6431 RAxML gene trees were used as input in the consensus and quartet-
251 based approaches. Finally, the Bayesian primary concordance tree and associated
252 concordance factors (CFs; indicative of the posterior probability of gene trees supporting
253 a node) at each internode of the primary concordance tree was computed in the program

254 BUCKy v1.4.4 (Larget et al., 2010). Because BUCKy is computationally intensive for a
255 large number of input gene trees, we only used orthologous gene sets that were
256 potentially informative for resolving gene trees in these analyses. Specifically, we utilized
257 1517 genes that showed average bootstrap values >50 across the RAxML-inferred gene
258 trees. (These are generally the loci with sufficient genetic variation across the tree to
259 provide information about branch support.) The input of a posterior distribution of gene
260 trees was generated from an analysis with MrBayes v3.2 (Huelsenbeck & Ronquist,
261 2001). We ran MrBayes for one million Markov chain Monte Carlo (MCMC) generations,
262 and every 1000th tree was sampled. After discarding the first half of the 1000 resulting
263 trees from MrBayes as burnin, BUCKy was performed for one million generations with
264 the default prior probability that two randomly sampled gene trees share the same tree
265 topology is 50% ($\alpha = 1$) (Larget et al., 2010).

266 All inferred species trees were plotted using the R package “phytools” (Revell, 2012).
267 To estimate dates of divergence, we used the function “chronos” in the R package “ape”
268 (Paradis et al., 2004) to fit a chronogram to the RAxML genome-wide concatenated
269 phylogeny by using penalized likelihood and maximum likelihood methods implemented
270 in chronos. Times were calibrated using a previous estimate of the divergence time
271 between *Solanum* and *Jaltomata* at ~17 Ma (Särkinen et al., 2013). To visualize gene tree
272 discordance, a “cloudogram” of 183 gene trees with average node bootstrap values
273 greater than 70 was prepared using DensiTree v 2.2.1 (Bouckaert, 2010).

274

275 Ancestral state reconstruction

276 The number of sampled species (14) is small compared to the size of this clade (60-
277 80 species), and sparse taxon sampling is known to affect the reconstruction of ancestral

278 character states (Heath et al., 2008). Nonetheless, to assess whether our confidence in the
279 number and placement of transitions generally differs among different traits in our clade,
280 we reconstructed ancestral states for fruit color, nectar color, nectar volume and corolla
281 shape. We used the ultrametric species tree inferred from the RAxML genome-wide
282 concatenated phylogeny and the distribution of traits at the tips of phylogeny (Figure 2A)
283 as input. While nectar volume is a quantitative trait, the other three traits are categorical
284 (fruit color: purple/red/orange/green; nectar color: red/clear; corolla shape:
285 rotate/campanulate/tubular). Ancestral character states were inferred using the standard
286 maximum likelihood method with the equal rates model “ER” within the phytools
287 package (Revell, 2012), which models the evolution of discrete-valued traits using a
288 Markov chain, and the evolution of continuous-valued traits using Brownian motion.

289

290 Testing for introgression

291 We searched for evidence of post-speciation gene flow, or introgression, using the
292 ABBA-BABA test (Durand et al., 2011; Green et al., 2010) on the concatenated
293 orthologous sequence alignment (n=6431 and 6,223,350 sites in total). The ABBA-
294 BABA test detects introgression by comparing the frequency of alternate ancestral (“A”)
295 and derived (“B”) allele patterns among four taxa. In the absence of gene flow, the
296 alternate patterns ABBA and BABA should be approximately equally frequent, given the
297 equal chance of either coalescence pattern under incomplete lineage sorting (ILS). An
298 excess of either ABBA or BABA patterns is indicative of gene flow. Because of the low
299 resolution of many recent branches within major clades of the phylogenetic tree (see
300 Results), evidence for introgression was only evaluated between four well-supported

301 major subclades, each of which is characterized by a distinct fruit color in our specific
302 dataset (i.e. the purple-, red-, and orange-fruited major clades, and a two-species clade of
303 green-fruited taxa; see Results). Patterson's D -statistic was calculated for all four-taxon
304 combinations including one taxon from the green-fruit lineage, one from red or orange-
305 fruit lineage, one from purple-fruit lineages, and one from tomato as the outgroup.
306 Patterson's D -statistic is calculated as $(ABBA-BABA) / (ABBA+BABA)$ for biallelic
307 sites in the multiple sequence alignment (Durand et al., 2011; Green et al., 2010). To
308 further investigate the taxa involved in and the direction of introgression specifically
309 involving purple-fruited lineages, we also used a symmetric five-taxon phylogeny
310 method 'D-foil' test (Pease & Hahn, 2015) on the transcriptome-wide concatenated
311 dataset, however the results of these analyses were inconclusive (see Supplementary text).

312

313 Identifying genetic variation associated with trait evolution

314 We used two general strategies to identify loci that might contribute to important
315 phenotypic trait (fruit and floral) transitions within *Jaltomata*. First, to identify loci that
316 have experienced lineage-specific *de novo* adaptive molecular evolution, we evaluated
317 loci for patterns of molecular evolution indicative of positive selection on specific
318 phylogenetic branches (i.e. $d_N/d_S > 1$). Second, to identify variants that might have been
319 selected from segregating ancestral variation, we identified genetic variants that had
320 polyphyletic topologies that grouped lineages according to shared trait variation rather
321 than phylogenetic relationships ('PhyloGWAS'; Pease et al., 2016).

322

323 *1) Lineage-specific de novo evolution associated with trait variation:* We identified loci

324 with signatures of *de novo* adaptive molecular evolution (i.e. significantly elevated rates
325 of non-synonymous substitution) across each available locus in our transcriptome
326 (sometimes called ‘reverse ecology’; Li et al., 2008) as well as in a set of *a priori*
327 candidate loci identified based on known or putative functional roles associated with
328 floral or fruit trait variation (Krizek & Anderson, 2013; Rausher, 2008; Specht &
329 Howarth, 2015) (see supplementary text). Tests were only performed on the four best-
330 supported branches within the phylogeny (see Results). For each locus (group of
331 orthologs), we inferred putative adaptive evolution (i.e. $d_N/d_S > 1$) using PAML v4.4
332 branch-site model (model = 2 and NS sites = 2) on the target branches (Yang, 2007). In
333 each analysis, a likelihood ratio test (LRT) was used to determine whether the alternative
334 test model (fixed_omega = 0) was significantly better than the null model (fixed_omega
335 = 1). In addition, because PAML uses a tree-based d_N/d_S model to reconstruct ancestral
336 states and lineage-specific substitutions, and because high levels of incongruence of gene
337 trees caused by ILS and introgression can produce misleading results when gene trees do
338 not match the assumed species tree (Mendes et al., 2016; Pease et al., 2016), we limited
339 our tests of molecular evolution to the subset of genes for which 1) the RAxML gene tree
340 contained the target ancestral branch (that is, the target branch was supported by the
341 genealogy of the tested/target locus); and 2) there was at least one non-synonymous
342 substitution that could be unambiguously assigned to this branch.

343 Prior to testing individual loci, we further filtered our data to ensure that poorly
344 aligned and/or error-rich regions were excluded from our alignments (as these tests are
345 particularly sensitive to alignment errors that generate spurious non-synonymous
346 changes). To do so, we used the program SWAMP v1.0 to remove regions from

347 alignments when they showed higher than expected non-synonymous substitutions (i.e.
348 more than five non-synonymous substitutions in 15 codons; the second sliding-window
349 sequence alignment check, different from above) and a minimum sequence length of 50
350 codons (Harrison et al., 2014). We also required that each alignment must contain a
351 tomato sequence and orthologous sequences from all investigated *Jaltomata* species
352 (except for *J. grandibaccata* because reproductive tissues were not sampled from this
353 species). The resulting sequence alignments were converted to codon-based MVF file
354 format (Pease & Rosenzweig, 2015), prior to performing branch-site tests. We first
355 defined putative genes showing positive selection by using the uncorrected p -value <0.01
356 as cutoff. The false discovery rate (FDR; (Benjamini & Hochberg, 1995) was then
357 calculated for the PAML p -values in each branch-specific test.

358 For the PAML analyses of *a priori* candidates, we used a slightly less stringent
359 statistical cutoff for consideration and required only that each cluster of orthologous
360 sequences had a minimum representation of species from each of the major clades (see
361 supplementary text); this allowed us to evaluate more of these loci while still testing
362 molecular evolution only on the four well-supported branches. For *a priori* genes
363 showing a significant signature of positive selection ($p <0.05$) and for genes identified by
364 the genome-wide unbiased analyses (FDR <0.1), we manually checked the sequence
365 alignments to examine whether they contain putative multi-nucleotide mutations
366 (MNM), which can cause false inferences of positive selection in the PAML branch-site
367 test (Venkat et al., 2017). Here we assigned an MNM in cases where we observed that a
368 single codon had 2 or 3 substitutions on the selected branch.

369 To determine the putative functional categories of genes with elevated per site

370 non-synonymous substitution rates, and to assess whether these were enriched for
371 particular functional categories, genes with uncorrected p -value <0.05 from the
372 transcriptome-wide analysis were also examined using Gene Ontology (GO) terms. GO
373 term reference was obtained from the Gene Ontology project (www.geneontology.org).
374 GO terms for each gene were obtained from SolGenomics (<ftp://www.solgenomics.net>).
375 GO term enrichment analysis was performed with ONTOLOGIZER v2.0 using the
376 parent-child analysis (Bauer et al., 2008).

377

378 2) *Ancestral genetic variation associated with trait variation*: To identify shared ancestral
379 variants that were associated with trait variation across lineages, we used a ‘PhyloGWAS’
380 approach (Pease et al., 2016) in which we searched for SNPs that were shared by current
381 accessions that share the same character state, regardless of their phylogenetic relatedness.
382 This approach is only informative in cases where trait variation is not confounded with
383 phylogenetic relationships, which in *Jaltomata* applies to floral shape transitions from the
384 ancestral ‘rotate’ form to the two derived forms (see Results). For our analysis, we treated
385 both campanulate and tubular corolla as the derived state, and rotate corolla as the
386 ancestral state (Figure S2). These categories of floral shape are also perfectly associated
387 with nectar color variation; species with rotate corollas have small amounts of clear/very
388 lightly colored nectar (ancestral), whereas species with campanulate or tubular corollas
389 have larger amounts of darkly colored red or orange nectar (derived). To assess whether
390 the number of nonsynonymous variants found to be associated with our defined groups of
391 floral traits (see Results) was greater than expected by chance, we generated a null
392 distribution due to ILS alone by simulating datasets over the species tree (Figure 2A)

393 using the program *ms* (Hudson, 2002). An associated *p* value was determined by the
394 proportion of simulated datasets that have a greater number of genes perfectly associated
395 with the floral trait distribution than our observed value (see Supplementary text).

396

397 **RESULTS**

398 *Transcriptome assembly and ortholog inference identified >6000 orthologs*

399 In assembled transcriptomes from both reproductive and vegetative tissues for
400 each of 14 *Jaltomata* species (except for *J. grandibaccata*, which only included
401 vegetative tissues), the number of transcripts per lineage ranged from 46,841-132,050,
402 and mean transcript length ranged from 736-925 bp (Table S2). Based on our criteria for
403 ortholog identification (see Methods, Figure S1), we ultimately identified 6431 one-to-
404 one orthologous genes for which we had sequences from all 14 investigated *Jaltomata*
405 species and a unique tomato annotated coding sequence. All of these 6431 genes were
406 used in the concatenation, majority rule, or quartet-based phylogeny reconstructions.
407 From this dataset, we used 1517 genes in the BUCKy reconstruction (see Methods).
408 Since we did not sample RNA from the reproductive tissues of *J. grandibaccata*, we
409 excluded this species from analyses of locus-specific adaptive evolution; this resulted in a
410 slightly larger number of orthologs, including those expressed solely in flowers. The
411 resulting dataset had 6765 alignments of orthologous coding sequences, each containing
412 sequences from the remaining 13 *Jaltomata* species (with *J. grandibaccata* excluded) and
413 tomato. Among them, 4248 genes also have *C. annuum* orthologs, thus could also be used
414 to test for positive selection on the ancestral branch leading to *Jaltomata*.

415

416 *Phylogenomic reconstruction of Jaltomata lineages supports several major clades*

417 All four phylogenetic inference methods (concatenation, majority rule, quartet-
418 based, and Bayesian concordance) generated a nearly identical species tree topology
419 (Figure 2A, S3). In all trees, the first split in the species tree produces a well-supported
420 clade that includes three north- and central-American species (*J. procumbens*, *J.*
421 *repandidentata*, and *J. darciana*) that all share floral traits (rotate corollas and light
422 nectar) and make dark purple/purple fruit (CF=87). The remaining 11 species, that are
423 found exclusively in South America and vary in floral traits and fruit colors, form a single
424 moderately supported clade (CF=66). Based on sequence divergence at synonymous sites
425 (Table S3), lineages within the non-purple-fruited clade have pairwise distance of 0.26%
426 to 0.53%, and differ from the purple-fruited lineages by 0.95% to 1.29%. Within the non-
427 purple-fruited group, our reconstruction indicates that the red-fruited species *J. auriculata*
428 is sister to the remaining species. The remaining species are split into a moderately
429 supported clade (CF=67) of two species (*J. calliantha* and *J. quipuscoae*) that share floral
430 traits and green fruits, and a relatively poorly supported clade (CF=19) consisting of the
431 remaining eight species that vary extensively in floral traits but all produce orange fruit.
432 Although quantitative support for different clades varied based on inference method
433 (Figure S3), all trees produced the same topology, with the exception of inferred
434 relationships within the orange-fruited group among *J. biflora*, *J. sinuosa*, *J. aijana* and *J.*
435 *umbellata*. In that case, all concatenation, quartet-based, and concordance trees supported
436 *J. sinuosa* as more closely related to *J. aijana* and *J. umbellata* (6% of gene trees support
437 this grouping), while consensus trees placed *J. sinuosa* as sister to *J. biflora* (7% of gene
438 trees support this grouping).

439

440 *Segregating variation is broadly shared among species in different subclades*

441 To quantify how much variation shared among present subclades—presumably
442 because of either shared ancestral variation or ongoing introgression—we mapped RNA-
443 seq reads from each species to the tomato reference genome and called high-quality
444 variants from ~8 million sites with more than 10X sequencing depth for all investigated
445 species. We identified a large number of sites that are sorting the same allele among
446 different subclades. Among them, 4303 variant sites are sorting in all four subclades
447 (Figure 3A). We also quantified how many sites that are heterozygous in one lineage
448 (accession) have the same two alleles sorting in other subclades. Within each lineage, the
449 proportion of heterozygous sites range from 0.02%~0.16% (Table S4), which is
450 comparable to the level of heterozygosity observed in self-compatible tomato species
451 (Pease et al., 2016). For 13.57% to 64.35% of heterozygous sites in one species (Figure
452 3A, Table S4), both alleles could also be identified in other subclades, again indicative of
453 a large amount of shared allelic variation.

454

455 *Phylogenomic discordance accompanies rapid diversification*

456 As expected given the large number of genes ($n=6431$ and 6,223,350 sites in
457 total) used for phylogenetic inference, our resulting species trees had very strong
458 bootstrap support for almost all nodes (Figure S3A and S3C). Despite this,
459 reconstructions also revealed evidence of extensive gene tree discordance consistent with
460 rapid consecutive lineage-splitting events in this group (Figure 2B; S3B and S3D). For
461 instance, the 6431 genes inferred 6431 different topologies, none of which matched the

462 topology of the inferred species tree (Figure 2A). The concatenation tree has many
463 extremely short internal branches, where gene trees show high levels of phylogenetic
464 discordance. We detected a strong correlation between the internal branch length and
465 levels of discordance ($P = 0.0001$, Figure S4), consistent with both ILS and introgression.
466 Short branch lengths and extensive discordance were also detected for trees built
467 individually for each of the 12 chromosomes (Figure S5; supplementary text).

468 Only three branches within *Jaltomata* are supported with relatively little
469 discordance, i.e., with Bayesian CFs greater than 50 (Figure S3D): the branch leading to
470 the purple-fruited clade, the branch uniting all non-purple-fruit *Jaltomata* lineages, and
471 the branch leading to the two green-fruited lineages (Figure 2A). Along with the ancestral
472 *Jaltomata* branch, these were the four branches on which most of our subsequent
473 analyses were performed.

474

475 *Introgression after speciation among major clades of Jaltomata lineages*

476 Given the apparent high level of phylogenetic discordance among our examined
477 species, we tested for evidence of introgression on the background of presumed ILS. To
478 do so, we calculated genome-wide D -statistics using the ABBA-BABA test. We only
479 examined introgressions across well-supported subclades. In particular, we compared the
480 distribution of trees in which one of two sister taxa (here, a species from either the red-,
481 green-, or orange-fruited lineage) is closer to more distantly related species (in the purple-
482 fruited clade) than the other. We found several such cases (Figure 3B). For example,
483 there was a significant excess of sites that grouped the red-fruited lineage (*J. auriculata*)
484 with a purple-fruited lineage, relative to the number of sites that grouped the green-

485 fruited lineage with the purple-fruited lineage, indicative of detectable gene flow between
486 the red-fruited and purple-fruited lineages since their split (Figure 3B and Table S5). We
487 also inferred putative introgression, in at least two separate events, involving six species
488 in the orange-fruited clade with the purple-fruited clade (Figure 3B and Table S5). First,
489 we inferred a shared introgression event between the purple-fruited group and three of the
490 orange-fruited species (*J. grandibaccata*, *J. dendroidea*, and *J. incahuasina*); this excess
491 includes shared specific sites that support the same alternative tree topology for each of
492 these three ingroup species, suggesting that it likely involved the common ancestor of all
493 three contemporary orange-fruited species (Table S6). Second, we detected evidence for
494 gene flow between the remaining orange-fruited species (*J. yungayensis*, *J. biflora* and *J.*
495 *sinuosa*) and the purple-fruited lineage, in the form of significant genome-wide *D*-
496 statistics (Table S5). Because we did not observe an excess of shared specific sites
497 supporting the same alternative tree topology among these three orange-fruited species
498 (Table S6), these patterns are suggestive of three putative independent introgression
499 events. However, given very low resolution of patterns of relatedness among orange-
500 fruited species, the specific timing of these events is hard to resolve.

501

502 Ancestral state reconstruction suggests different histories for fruit color and floral trait
503 evolution

504 Based on the inferred species tree (Figure 2A), we reconstructed the ancestral
505 states of fruit and floral traits (Figure 4; S6). The four subclades of *Jaltomata* species
506 were inferred to have evolved different fruit colors at their corresponding common
507 ancestors (Figure 4A). Our reconstruction suggests that the derived nectar traits

508 (orange/red nectar color, and increased nectar volume) probably evolved at the common
509 ancestor of the green/orange-fruited clade (Figure S6A and S6B), with two subsequent
510 reversions to ancestral conditions within this clade. The evolution of the two derived
511 corolla shapes in *Jaltomata* (campanulate and tubular) appears to be more complex
512 (Figure 4B). At the majority of internodes within the non-purple-fruited lineages, all three
513 corolla shapes (i.e. rotate (ancestral), campanulate and tubular) show $\geq 10\%$ probability of
514 being the ancestral state, making specific inferences about corolla shape evolution within
515 this clade uncertain. Consistent with these patterns, we found that concordance factors
516 were very low at almost all internodes within the radiating subgroup that displays the
517 derived floral traits (i.e., the non-purple-fruited lineages) (Figure 2A; S3D), whereas they
518 were considerably higher on branches associated with fruit color evolution (including the
519 branch uniting the two green-fruited species analyzed).

520 These analyses suggest alternative evolutionary and genetic histories for our traits
521 of interest. In particular, strong associations between fruit color transitions and specific
522 branches/clades within *Jaltomata* suggests that the underlying genetic changes are more
523 likely due to conventional lineage-specific *de novo* evolution along the relevant branches.
524 In contrast, the distribution of floral trait variation produces an ambiguous reconstruction
525 of trait transitions, especially for floral shape, such that the distribution of ancestral
526 versus derived floral shape variation is unassociated with phylogenetic relationships in
527 the non-purple-fruited *Jaltomata* clade (Figure 4B). While strictly *de novo* evolution
528 occurring multiple times is not excluded as an explanation of floral evolutionary
529 transitions, one alternative is that these trait transitions drew upon shared variation
530 segregating in the ancestor of these lineages. Accordingly, in the next sections we

531 evaluate both lineage-specific *de novo* evolution and selection from standing ancestral
532 variation when searching for genetic variants that might have contributed to floral trait
533 evolution. The lineage-specific *de novo* evolution analysis alone is used to identify
534 potential candidates for fruit color evolution, since the approach we use to identify
535 standing ancestral variation is only informative in cases where trait variation is not
536 confounded with phylogenetic relationship.

537

538 *Loci with patterns of positive selection associated with lineage-specific trait evolution*

539 We performed tests of molecular evolution for all orthologous clusters that
540 contained a sequence from every *Jaltomata* accession and an ortholog from the tomato
541 outgroup. Depending upon the specific branch being tested, we detected evidence for
542 positive selection in ~1-2% of loci in our dataset, based on whether the locus had
543 significantly elevated d_N/d_S ratios ($d_N/d_S > 1$; $p < 0.01$). This included 1.88% of genes (67
544 out of 3556 testable genes; Table S7) on the *Jaltomata* ancestral branch, 1.58% in the
545 purple-fruited group (48 out of 3033 testable genes; Table S8), 2.61% in the red-fruited
546 group (70 out of 2686 testable genes; Table S9), 1.96% in the green-fruited group (30 out
547 of 1531 testable genes; Table S10), and 0.74% in the non-purple-fruited *Jaltomata*
548 lineages (15 out of 2039 testable genes; Table S11). Many of the genes showing elevated
549 d_N/d_S appear to have general molecular functions (e.g. transcription, protein synthesis, or
550 signaling), including numerous genes involved in various stress responses, such as heavy
551 metal tolerance, sugar starvation response, protection from ultraviolet (UV) radiation and
552 extreme temperature, and herbivore and pathogen resistance (Table S7-11). Our
553 positively selected loci contain genes functionally associated with photosynthesis,

554 fatty/lipid biosynthesis and transportation, and sugar signal transduction (observed in the
555 GO enrichment analysis; Table S12-16), as well as loci with unknown functions.

556 After controlling for multiple tests using an FDR <0.1 on each branch tested, only
557 three genes on the *Jaltomata* ancestral branch, one gene on the purple-fruited ancestral
558 branch, and four genes on the red ancestral branch, remained significant for $d_N/d_S > 1$.
559 Interestingly, for 5 of these 8 loci, the inference of positive selection appears to be due to
560 the presence of a multi-nucleotide mutation (MNM) specifically on the target branch, a
561 mutational pattern known to produce spurious inferences of positive selection in PAML's
562 branch-site test (Venkat et al., 2017). With one exception, all of these potential false
563 positives were found on the ancestral *Jaltomata* or purple-fruited clade branches, the
564 longest *Jaltomata*-specific branches in our analyses; as MNMs are more likely to appear
565 on long branches (they are relatively rare mutations) these are expected to be enriched for
566 these spurious inferences in the branch-site test (Venkat et al., 2017). Our three remaining
567 loci, with positive selection on the red-fruited branch, include a gene (*BANYULS*;
568 ortholog to *Solyc03g031470*) with functional roles in pigmentation (see Discussion).

569 We also detected several instances where slightly less stringent criteria ($d_N/d_S > 1$;
570 $p < 0.05$) revealed lineage-specific adaptive evolution of our *a priori* candidate genes
571 occurring on a branch that is also inferred to be associated with the evolution of derived
572 traits (Table S17). Most notably, we found evidence of positive selection on candidate
573 loci that are likely to be involved in fruit color, including a gene encoding ζ -carotene
574 isomerase (*Z-ISO*; ortholog to *Solyc12g098710*) on the red-fruited lineage, and the two
575 other genes significant on the ancestral branch of the green-fruited lineages encoding
576 carotenoid cleavage enzyme 1A (*CCDIA*; ortholog to *Solyc01g087250*) and zeaxanthin

577 epoxidase (*ZEP*; ortholog to *Solyc02g090890*) (Figure 5, see Discussion). We detected
578 signatures of positive selection on fewer of the genes involved in floral development,
579 mostly notably in the MADS-box gene *APETALA3* (*AP3/DEF*, ortholog to
580 *Solyc04g081000*) on the ancestral branch to the purple-fruited lineage. Overall, we note
581 that many of our loci (including *a priori* candidates) did not meet the requirements to be
582 tested for positive selection (Table S7); in particular, gene trees for many loci lacked the
583 required support for a specific internal branch, either because of incongruence or limited
584 phylogenetic signal, especially within the rapidly diverging orange-fruited clade (Table
585 S17).

586

587 *Loci potentially associated with trait evolution from standing ancestral variation*

588 To investigate whether ancestral variants are potentially associated with floral
589 trait diversification, we performed a “PhyloGWAS” analysis (Pease et al., 2016). Such
590 variants will have differentially fixed among descendant lineages, leading to genes that
591 cluster species together based on floral traits regardless of their overall phylogenetic
592 relationships. We found 31 genes with nonsynonymous variants perfectly associated with
593 the derived floral traits (Table S18), which was significantly more than the number of
594 loci expected by chance to have segregation patterns that exactly match the tip states ($p <$
595 9.3×10^{-5}). Most of these genes are characterized by only one or few nucleotide
596 differences, which is an expected pattern for variants recently selected from standing
597 ancestral variation (Pease et al., 2016). These results suggest that one or few molecular
598 variants present in ancestral populations could contribute to the multiple apparent
599 transitions to derived floral shapes in *Jaltomata*. Among the loci identified by our

600 approach, some genes are potentially functionally related to petal development, including
601 *ARGONAUTE1* (*AGO1*) and xyloglucan endotransglucosylase/hydrolase 2 (*DcXTH2*)
602 (see Discussion).

603

604 **DISCUSSION**

605 Within rapidly radiating groups, the patterns of genetic relatedness among
606 lineages provide essential data for determining the pace and location of important trait
607 transitions, and their underlying genes; both are critical for understanding the drivers of
608 rapid diversification and speciation. Our phylogenomic analyses of the 14 investigated
609 *Jaltomata* species revealed genome-wide gene tree discordance, and a highly complex
610 history of genetic relatedness among contemporary lineages, consistent with other studies
611 of recently radiating groups (Brawand et al., 2014; Garrigan et al., 2012; Novikova et al.,
612 2016; Pease et al., 2016). We identified substantial ILS and shared ancestral
613 polymorphism, as well as evidence of putative introgressions among the subclades of
614 *Jaltomata* species, as the sources of this observed complex genome-wide history. This
615 complexity was also reflected in inferences about the evolution of major trait transitions
616 within the group. We found differences in the patterns of fruit versus floral character
617 evolution and in our inferred confidence in the reconstruction of these patterns, including
618 their likely risk of hemiplasy. Given this, we used several strategies to identify loci that
619 might contribute to the evolution of these traits, including examining lineage-specific *de*
620 *novo* adaptive evolution along well-supported branches and identifying variants that
621 might have been selected from standing ancestral variation. Overall, by combining
622 evidence from molecular evolution with data on trait variation across a clade—and a

623 more direct accounting for the risk of hemiplasy—we generated more conservative, but
624 credible, inferences of candidate genes responsible for the evolution of ecologically
625 important phenotypic traits. Specifically, we identified several functionally relevant
626 candidate genes for our target trait transitions, and different potential sources of adaptive
627 evolution fueling changes in target floral versus fruit traits.

628

629 *Extensive ILS and several introgression events produce a complex genome-wide history*
630 *during rapid diversification*

631 Our reconstruction of phylogenetic relationships based on a transcriptome-wide
632 dataset agrees with previous studies that identified three major *Jaltomata* sub-clades
633 primarily distinguished from each other by their fruit colors (Miller et al., 2011; Särkinen
634 et al., 2013), including the previous inference that the clade of purple-fruited species is
635 sister to the rest of the genus (Särkinen et al., 2013). However, the relationships among
636 species within each subclade show high levels of discordance, and the distribution of
637 genetic variation strongly indicates a rapid and recent evolutionary origin; the internal
638 branch lengths within each subclade are short, especially in the non-purple-fruited
639 lineages (Figure 2A). Accordingly, the history of these species—including the
640 orange/green-fruited lineages that show the most floral trait diversity among *Jaltomata*
641 species—is expected to be strongly affected by ILS and introgression.

642 Indeed our genome-wide reconstruction indicated that, although concatenation or
643 quartet-based approaches generated a species tree with high bootstrap support values
644 (commonly observed when inferring trees from large amounts of data (Kubatko &
645 Degnan, 2007; Salichos & Rokas, 2013), gene tree discordance was rampant, and

646 individual gene trees showed highly variable support for the specific placement of
647 individual species (Figure 2B) especially at short branches (Figure 2A). Our finding of
648 extensive genome-wide ILS in the genus *Jaltomata* agrees with other recent studies on
649 contemporary (Novikova et al., 2016; Pease et al., 2016) and relatively ancient adaptive
650 radiations in plant species (Wickett et al., 2014; Yang et al., 2015), and is emerging as a
651 universal signal of rapid radiation in comparative genome-wide datasets.

652 Resolution of phylogenetic relationships among the species within each *Jaltomata*
653 subclade was insufficiently clear to investigate introgression within subgroups. However,
654 across major sub-clades we identified at least two clear introgression events that involved
655 either orange and red-fruited lineages with the purple-fruited lineages, similar to the
656 detection of introgression events in other recent genome-wide studies on closely related
657 plant species (Eaton & Ree, 2013; Novikova et al., 2016; Owens et al., 2016; Pease et al.,
658 2016). Interestingly, in one case an excess of sites supported a shared introgression
659 pattern between three orange-fruited species (i.e. *J. grandibaccata*, *J. dendroidea*, and *J.*
660 *incahuasina*) and the purple-fruited clade, consistent with a scenario in which
661 introgression involved the recent common ancestor of these three orange-fruited species.
662 Moreover, in this case, the inference of a single shared introgression event itself provided
663 more confidence in this specific ancestral branch within the orange-fruited clade. Overall,
664 as with ILS, post-speciation introgression is another inference frequently emerging from
665 contemporary phylogenomic studies of radiations.

666

667 *Inferring the history of trait evolution and the contributing loci in the presence of*
668 *rampant discordance*

669 The complex history of genomic divergence in this clade has clear consequences
670 for inferences of trait and gene evolution. When ILS or introgression can plausibly
671 explain the discordant distribution of traits, it might be impossible to infer trait evolution
672 with any certainty in the absence of additional independent information about target traits,
673 such as their genetic basis (Hahn & Nakhleh, 2016). In contrast, the evolutionary
674 transitions of some traits might be confidently inferred as long as the relevant branches
675 and resulting relationships are associated with higher levels of concordance (Hahn &
676 Nakhleh, 2016). Species in *Jaltomata* exhibit extensive trait diversity, most notably in
677 fruit color, corolla shape, and nectar volume and color (Figure 2A) (Miller et al., 2011),
678 and one of the main goals in this study was to better understand the evolutionary history
679 of these trait transitions, including the genetic basis of the traits associated with rapid
680 phenotypic diversification. A previous phylogenetic study based on a single locus
681 suggested that floral traits (including corolla shape and nectar color) might have evolved
682 multiple times independently in *Jaltomata* species (Miller et al., 2011). However, the
683 presence of rampant discordance, and abundant evidence of shared ancestral variation,
684 makes inferring the history of trait transitions and their genetic basis especially
685 challenging in this group. Indeed, our analyses indicated that different classes of trait
686 transition—most notably fruit color versus floral shape variation—were differently
687 susceptible to hemiplasy. For floral shape evolution in *Jaltomata*, a lack of resolution and
688 high gene tree discordance at key nodes within the phylogeny, including within the clade
689 displaying the greatest phenotypic diversity (Figure 2A and Figure 4B), mean that
690 hemiplasy is a plausible explanation of the discordant distribution of similar floral
691 traits—rather than multiple independent evolutionary events (i.e. homoplasy). In contrast,

692 we showed that the history of fruit color evolution could be confidently inferred, as these
693 trait transitions occurred on branches with higher levels of concordance and therefore
694 lower risks of hemiplasy (Figure 2A and Figure 4A).

695 Accounting for the potential influence of hemiplasy is also critical when
696 generating hypotheses about the loci that could have contributed to trait transitions. In
697 general, incorrect reconstructions of trait history will suggest incorrect candidates
698 involved in the evolution of those traits. Moreover, tests of molecular evolution can be
699 specifically misled if trait transitions occur on discordant gene trees (Mendes et al., 2016).
700 Accordingly, to identify loci that might be responsible for any particular trait transitions,
701 different approaches will be appropriate depending upon the confidence with which
702 hemiplasy can be excluded or not. For traits evolving on branches where discordance is
703 low, confidence is high, and hemiplasy is unlikely, it is reasonable to expect that lineage-
704 specific *de novo* substitutions are a substantial contributor to relevant trait evolution.
705 Given a high risk of hemiplasy, genetic variation underpinning trait evolution could
706 potentially come from additional sources, including recruitment of ancestral
707 polymorphisms and/or introgression. These differences are exemplified in our study by
708 the alternative histories, and different genetic hypotheses, generated for fruit color versus
709 floral shape traits.

710

711 **Floral shape evolution** – Based on the inferred species tree (Figure 2A), the 14
712 investigated *Jaltomata* species are not related according to their floral shape traits (i.e.
713 floral shape is distributed paraphyletically). Moreover, the branch lengths leading to
714 lineages with derived character states are uniformly short with high levels of gene tree

715 discordance (Figure 2A), so that the probability of hemiplasy is expected (Hahn &
716 Nakhleh, 2016) to be very high. Indeed, when we reconstruct the evolution of corolla
717 shape (Figure 4B), the three alternative corolla morphs were inferred to be almost equally
718 likely at the common ancestor of non-purple-fruited lineages. It is possible that this lack
719 of resolution is due to introgression. For example, the ancestral floral character states (i.e.
720 rotate corolla shapes and clear nectar) found in *J. yungayensis* and *J. sinuosa* within the
721 orange-fruited clade could be due to alleles introgressed from purple-fruited species, as
722 we identified putative introgression events between those lineages (Figure 3B). However,
723 the lack of a reference genome for *Jaltomata* precluded us from more directly
724 investigating evidence (for example, locus-specific patterns of introgression) that
725 introgression might contribute to the distribution of ancestral floral traits within this sub-
726 clade. Instead, because of the high risk of hemiplasy and low resolution of ancestral states,
727 we used several approaches to identify the genetic variants associated with the two other
728 potential sources of trait variation.

729 First, if the paraphyletic distribution of derived traits is due to hemiplasy among
730 species, the relevant nucleotide differences should be at the same sites in all lineages that
731 share derived traits (Hahn & Nakhleh, 2016). Using this rationale, Pease et al. (2016)
732 identified tens to hundreds of genetic variants among wild tomato lineages that were
733 exclusively associated with each of three ecological factors, variants that are candidate
734 targets of parallel ecological selection on standing ancestral variation in that group. Here,
735 we used an analogous approach to look for variants associated with phenotypic (floral)
736 trait variation in *Jaltomata*, and identified 31 candidate genes with nonsynonymous
737 variants that are completely correlated with the distribution of floral trait variation

738 (derived vs. ancestral; Figure S2). Among them, the gene *AGO1* (ortholog to
739 *Solyc02g069260*) is known to be necessary in *Arabidopsis* floral stem cell termination
740 and might act through *CUC1* and *CUC2* (*a priori* candidate genes), which redundantly
741 specify boundaries of floral meristem (Ji et al., 2011; Kidner & Martienssen, 2005).
742 Another gene *DcEXPA2* (ortholog to *Solyc02g091920*) is known to be markedly up-
743 regulated in the petals of carnation (*Diathathus caryophyllus*), and is potentially
744 associated with the petal growth and development (Harada et al., 2010). We also note that,
745 although these shared hemiplasious variants could be due to post-speciation introgression
746 rather than sorting from ancestral variation, given the often allopatric geographical
747 distribution of our lineages (Figure 1) and evidence of substantial allelic sharing among
748 contemporary subclades (Figure 3A), we infer that these variants were more likely sorted
749 from ancestral variation.

750 Second, because our reconstruction of floral trait evolution could not exclude the
751 role of lineage-specific *de novo* mutation, we also examined our transcriptomes for genes
752 showing lineage-specific evolution associated with the derived floral traits. However, we
753 found that only a small number of genes were testable due to the extensive gene tree
754 discordance specifically at the branches leading to subclades with the derived floral
755 characters. Moreover, none of our *a priori* candidate genes involved in floral
756 development showed suggestive patterns of molecular evolution on internal branches of
757 *Jaltomata* (Table S7), nor did we find any other functionally suggestive (floral
758 development-related) genes adaptively evolving on branches leading to specific
759 subclades within *Jaltomata*, with the exception of *APETALA3* (*AP3*)—a gene associated
760 with the formation of petals and stamens in flowering plants, including

761 *Arabidopsis* (Wuest et al., 2012; Specht & Howarth, 2015)—although this was detected
762 on the branch leading to the purple-fruited clade, within which all species retain the
763 ancestral rotate corolla form.

764

765 **Fruit color evolution** – In contrast to floral traits, ancestral state reconstruction in the 14
766 investigated *Jaltomata* species suggested that fruit color transitions follow phylogenetic
767 relationships. The ancestral states of fruit colors at most relevant internodes can be
768 inferred with high confidence (Figure 4A), and the internal branches leading to fruit trait
769 transitions are well-supported, indicating a low probability of hemiplasy (Figure 2A).

770 Accordingly, we identified a set of loci showing adaptive evolution specifically on these
771 branches, provided that each tested gene tree 1) supported the internal branch being
772 tested, and 2) showed at least one clade-specific nonsynonymous substitution on that
773 branch (Mendes et al., 2016; Pease et al., 2016), to avoid potential inference errors

774 associated with examining genes that have topologies discordant with the species tree
775 (Mendes et al., 2016). Our analyses revealed multiple adaptively evolving candidate loci
776 with clear functional relevance to these trait transitions, including several *a priori*

777 candidate genes involved in the carotenoid pathway (Yuan et al., 2015) (Figure 7). First,
778 *Z-ISO* (ortholog to *Solyc12g098710*), a key enzyme in the production of red-colored
779 lycopene in the carotenoid biosynthetic pathway (Chen et al., 2010), was positively
780 selected on the branch leading to our one species with bright red fruit, *J. auriculata*.

781 Interestingly, this gene was also previously found to show adaptive evolution specifically
782 on the branch leading to the red-fruited (Esculentum) group in wild tomatoes (Pease et
783 al., 2016). Second, on the branch leading to our two green-fruited species, we found

784 significantly elevated d_N/d_S ratios for both *CCDIA* (ortholog to *Solyc01g087250*), a gene
785 whose product participates in the conversion of carotenoid pigments to isoprenoid
786 volatiles (Ilg et al., 2014), and for *ZEP* (ortholog to *Solyc02g090890*), which converts
787 zeaxanthin to violaxanthin (Marin et al., 1996). *CCDI* has previously been identified in
788 tomato fruits as responsible for generating flavor volatiles (Auldridge et al., 2006; Simkin
789 et al., 2004). This functional observation from a closely related group is intriguing
790 because fruits of our two green-fruited species (*J. quipuscoae* and *J. calliantha*) appear to
791 produce the strongest scent within the 14 *Jaltomata* species analyzed here (J. Kostyun,
792 unpubl. data). This apparent increase in fragrance—presumably due to changed volatile
793 organic compounds—might play a role in attracting vertebrate frugivores for seed
794 dispersal.

795 In addition to carotenoids, among *a priori* candidate genes involved in the
796 biosynthesis pathway of water-soluble vacuolar anthocyanin pigments, we detected
797 *BANYULS* (ortholog to *Solyc03g031470*) selected on the red-fruited branch. (Note that
798 PAML also indicates adaptive molecular evolution of this locus on the purple-fruited
799 branch, but this appears to be due to the presence of a MNM; see results). In addition to *a*
800 *priori* candidates, genome-wide unbiased analyses also identified that multiple genes
801 belonging to *R2R3MYB*, *BHLH* and *WD40*-repeats classes of loci were under positive
802 selection in purple-fruited lineages and the red-fruited lineage. The *MYB-BHLH-WD40*
803 TF complexes are known to regulate cellular differentiation pathways, including of the
804 epidermis, as well as transcription of anthocyanin structural genes (Gonzalez et al., 2008;
805 Jaakola, 2013; Ramsay & Glover, 2005).

806

807 *Implications for the inference of phenotypic trait evolution and causal genetic variation*
808 *in rapid radiating lineages*

809 As highlighted here and in numerous recent phylogenomic studies (Eaton & Ree,
810 2013; Novikova et al., 2016; Owens et al., 2016; Pease et al., 2016), both ILS and
811 introgression contribute to the history of diversification within radiating clades, so that
812 evolution in these groups is more complex than can be represented by a simple
813 bifurcating species tree. This complexity has important implications for empirical
814 inferences about historical relationships and trait evolution, because assuming resolved
815 relationships without taking into account incongruence can fundamentally mislead
816 inferences in both these cases (Hahn & Nakhleh, 2016). It also must be accounted for
817 when considering the genetic changes that might have fueled diversification; when a trait
818 has several possible alternative evolutionary histories, it is necessary to investigate the
819 range of alternative sources of genetic variation—including *de novo* lineage-specific
820 evolution and selection from ancestral variation—that could fuel this trait evolution.
821 Here, we provided a genome-wide analysis of the recently diversified plant genus
822 *Jaltomata* in which we consider the relative risk of hemiplasy while identifying
823 candidates for the specific loci underlying trait evolution. Our analysis highlights a
824 growing appreciation that rapid radiations can and likely do draw on multiple sources of
825 genetic variation (Hedrick, 2013; Pease et al., 2016; Richards & Martin, 2017). Indeed,
826 while independently originating variants could explain the recurrent evolution of
827 phenotypic similarity—a frequent observation in adaptive radiations—it is clear that
828 shared ancestral genetic variation, or alleles introgressed from other lineages, have also
829 made substantial contributions (Elmer & Meyer, 2011; Stern, 2013). Going forward, it

830 will be necessary to distinguish between these alternative scenarios to understand how
831 different evolutionary paths contribute to phenotypic convergence and differentiation
832 (Martin & Orgogozo, 2013; Stern, 2013) and to identify the specific variants responsible.

833

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838

839 **AUTHOR CONTRIBUTIONS**

840 L.C.M, M.W., M.W.H., and J.L.K. designed the experiments; J.L.K generated the
841 experimental materials; M.W. conducted the bioinformatics analyses; and M.W. and
842 L.C.M. wrote the paper with contributions from M.W.H. and J.L.K.

843

844 **DATA AVAILABILITY**

845 Raw reads (FASTQ files) for generating the 14 species transcriptomes are
846 deposited in the NCBI SRA (BioProject: PRJNA380644). Multiple sequence alignments
847 used for phylogenetic tree reconstruction and molecular evolution analyses are available
848 at Dryad (doi: XXX). All commands and scripts used for analyses in this study can be
849 found in our project directory on GitHub (<https://github.com/wum5/JaltPhylo>).

850

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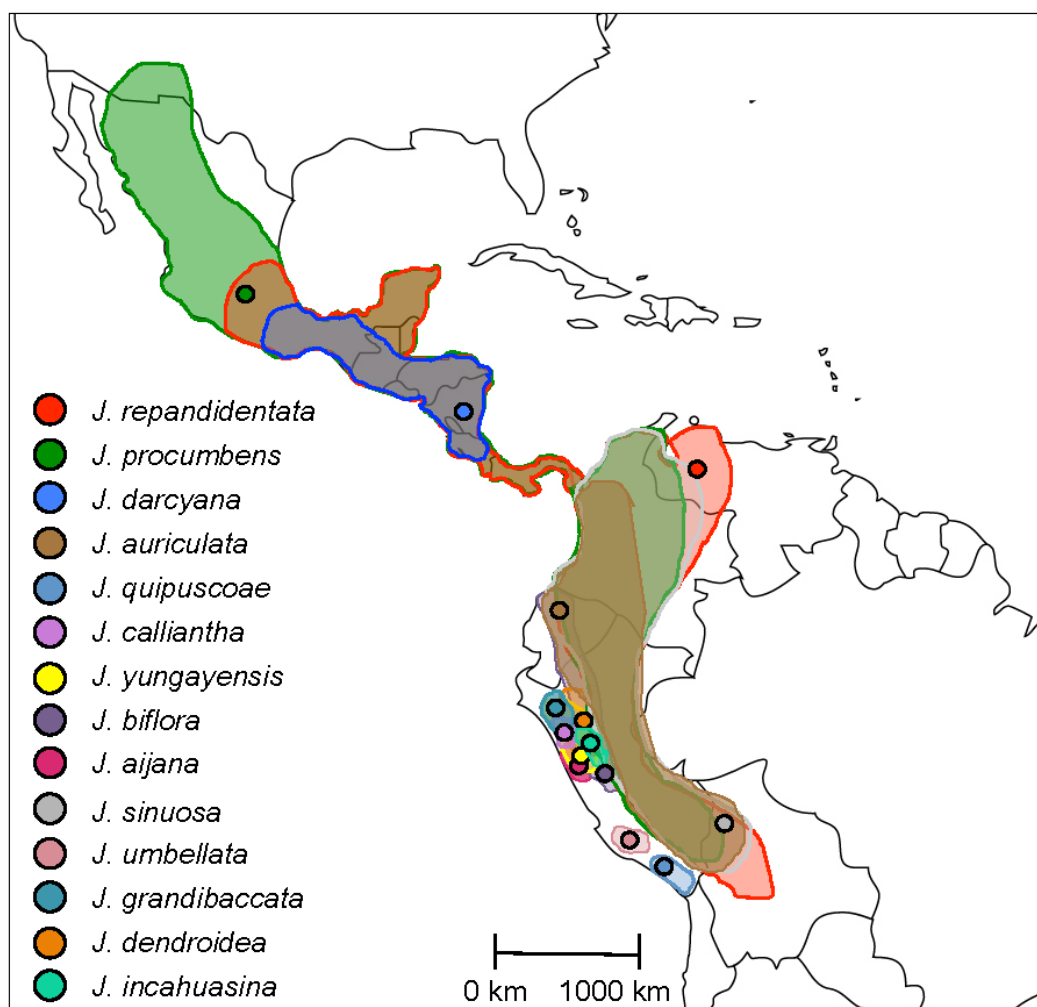


Fig 1. Geographic distribution of investigated *Jaltomata* species. For each species, the sample location is labeled. Ranges estimated from herbarium specimens (T. Mione and S. Leiva G., pers. comm.; J. L. Kostyun, unpub.).

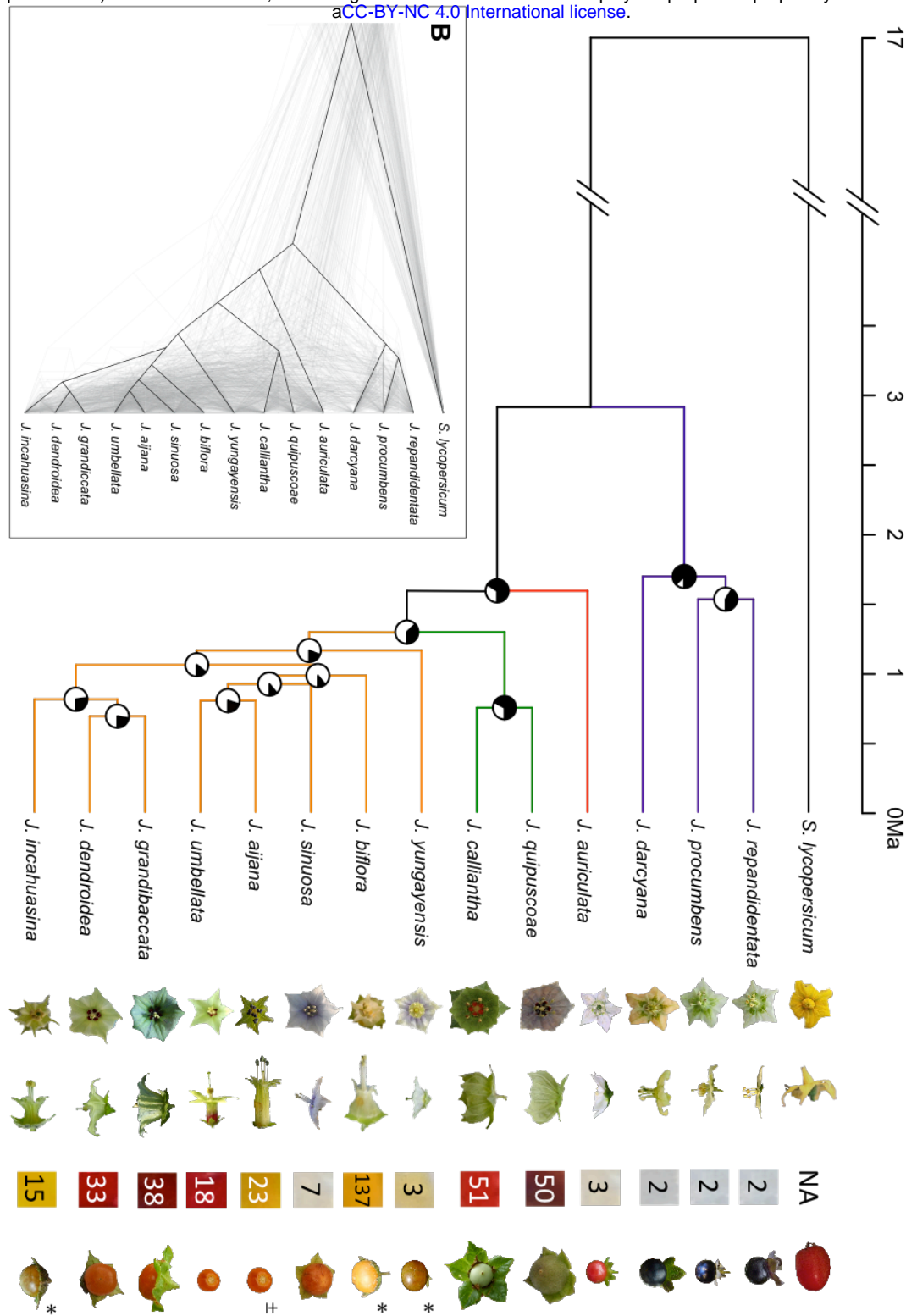


Fig 2. The phylogeny of investigated *Jaltomata* species. **(A)** A whole-transcriptome concatenated phylogeny of *Jaltomata* species with *Solanum lycopersicum* as outgroup. The Pie chart on each internode shows the concordance factor estimated from BUCKy, with the amount of black representing the degree of concordance. Divergence times estimated in *ape* with 17MYA *Jaltomata-Solanum* calibration from (Särkinen et al., 2013). Representative flower and fruit images to the right of species names: front view of flower, lateral view of flower, nectar color and volume (uL) per flower, and ripe fruit. Image with ± indicates that fruit from a similar species is shown, and * indicates contributed by Dr. Thomas Mione at Central Connecticut State University.

(B) A ‘cloudogram’ of 183 gene trees whose average bootstrap values are larger than 70 across the nodes. For contrast, the concatenated tree (Fig 2A) is shown in black.

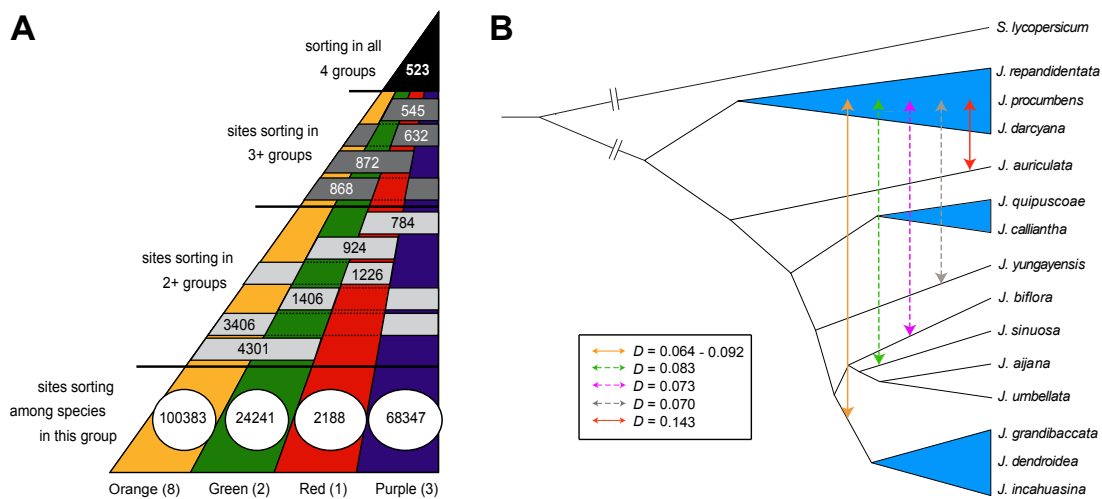


Fig 3. (A) Allele sorting, with the number of genetic variants within or shared between each *Jaltomata* subclade. (B) The introgression pattern among *Jaltomata* lineages. The solid lines indicate strong evidence of introgression between two lineages or sub-clades, while the dashed lines indicate putative introgression. The corresponding Patterson's *D* statistic value is labeled for each putative introgression event.

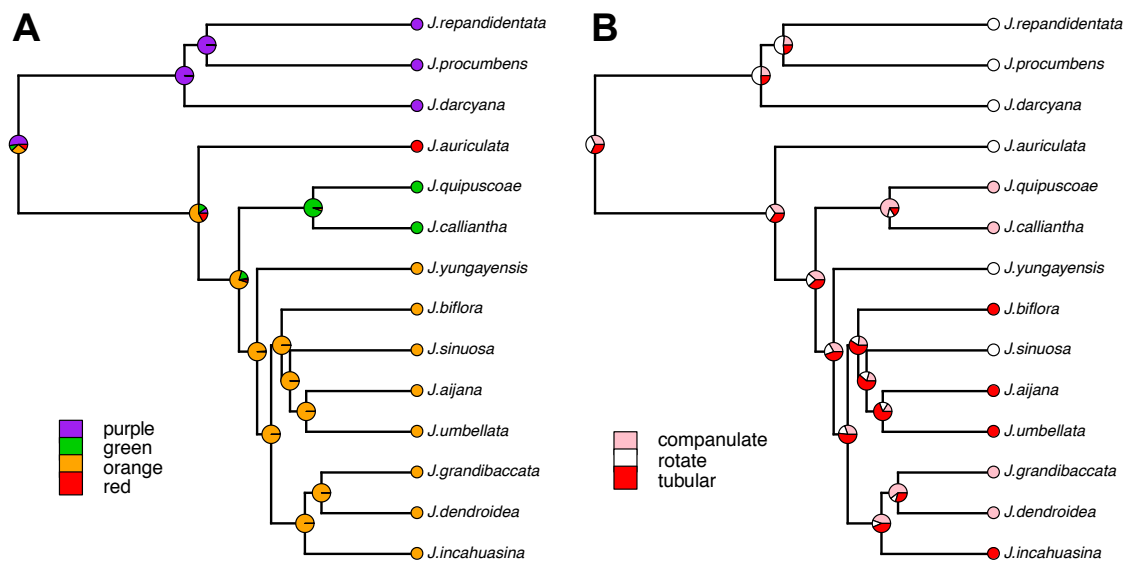


Fig 4. Ancestral character-state reconstruction of (A) fruit color, (B) corolla shape in investigated *Jaltomata* using maximum likelihood.

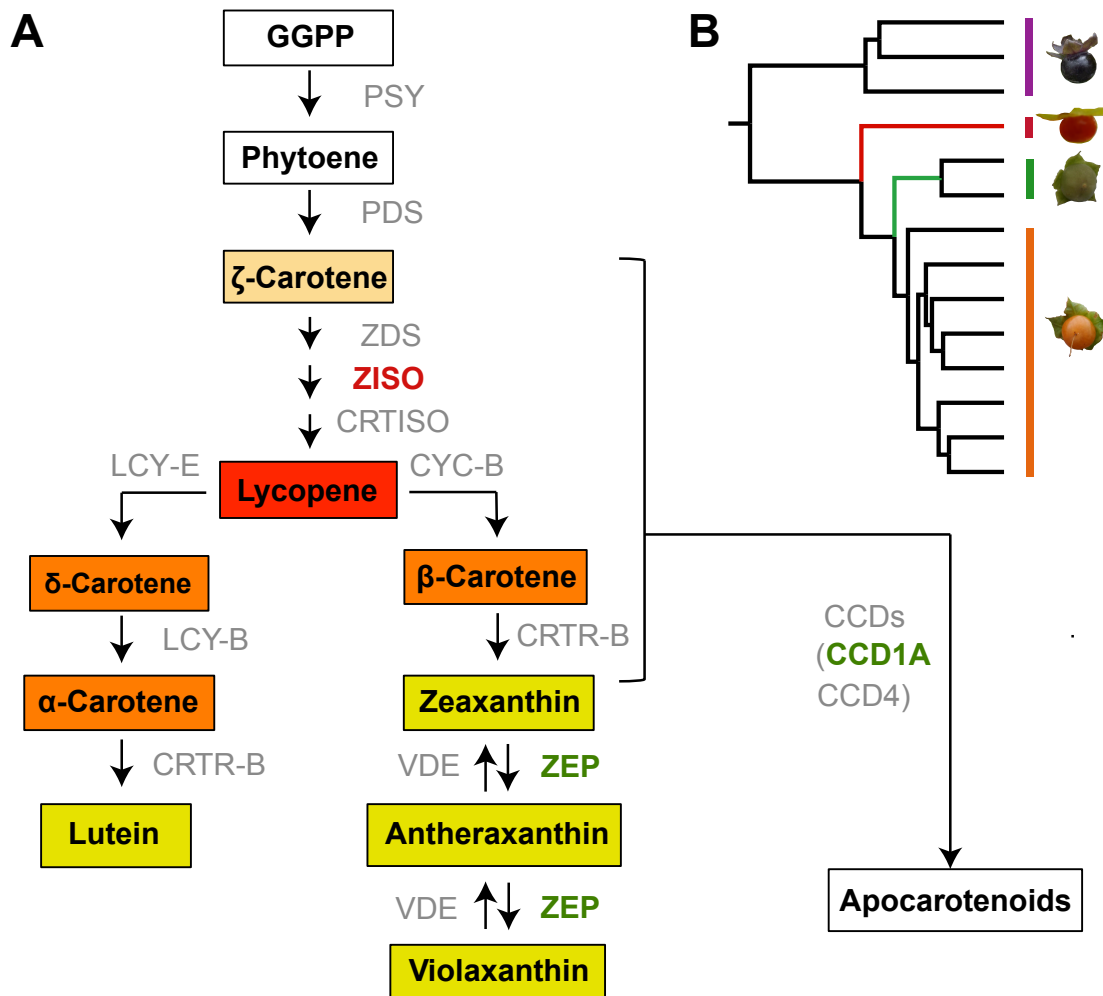


Fig 5. Genes under adaptive evolution in the carotenoid biosynthesis pathway. **(A)** Simplified carotenoid biosynthesis pathway modified from (Yuan et al. 2015). Genes under adaptive evolution are indicated by their names highlighted in colors corresponding to particular branches (see panel B). *PSY*, phytoene synthase; *PDS*, phytoene desaturase; *ZDS*, ζ-carotene desaturase; *ZISO*, ζ-carotene isomerase; *CRTISO*, carotenoid isomerase; *LCY-E*, lycopene ε-cyclase; *LYC-B*, lycopene β-cyclase; *CRTR-B*, β-ring hydroxylase; *CYC-B*, chromoplast specific lycopene β-cyclase; *ZEP*, zeaxanthin epoxidase; *VDE*, violaxanthin de-epoxidase; *CCD*, carotenoid cleavage dioxygenase. Metabolites are boxed and colored according to their compound colors, whereas white boxes indicate no color. **(B)** Positive selection signatures of genes on different branches are indicated by different colors: red-fruited lineages (Red), and green-fruited lineages (Green).