

1 TITLE: Target enrichment and extensive population sampling help untangle the recent, rapid
2 radiation of *Oenothera* sect. *Calylophus*

3
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32 ABSTRACT

33 *Oenothera* sect. *Calylophus* is a North American group of 13 recognized taxa in the
34 evening primrose family (Onagraceae) with an evolutionary history that may include
35 independent origins of bee pollination, edaphic endemism, and permanent translocation
36 heterozygosity. Like other groups that radiated relatively recently and rapidly, taxon boundaries
37 within *Oenothera* sect. *Calylophus* have remained challenging to circumscribe. In this study, we
38 used target enrichment, flanking non-coding regions, summary coalescent methods, tests for
39 gene flow modified for target-enrichment data, and morphometric analysis to reconstruct
40 phylogenetic hypotheses, evaluate current taxon circumscriptions, and examine character
41 evolution in *Oenothera* sect. *Calylophus*. Because sect. *Calylophus* comprises a clade with a
42 relatively restricted geographic range, we were able to extensively sample across the range of
43 geographic and morphological diversity in the group. We found that the combination of exons
44 and flanking non-coding regions led to improved support for species relationships. We
45 reconstructed potential hybrid origins of some accessions and note that if processes such as
46 hybridization are not taken into account, the number of inferred evolutionary transitions may be
47 artificially inflated. We recovered strong evidence for multiple origins of the evolution of bee
48 pollination from ancestral hawkmoth pollination, the evolution of edaphic specialization on
49 gypsum, and permanent translocation heterozygosity. This study applies newly emerging
50 techniques alongside dense infraspecific sampling and morphological analyses to effectively
51 address a relatively common but recalcitrant problem in evolutionary biology.

52

53 *Keywords.*– Gypsum Endemism, Onagraceae, *Oenothera* sect. *Calylophus*, Pollinator Shift,
54 Recent Radiation, Phylogenomics, Target Enrichment

55

56 INTRODUCTION

57 The challenges of reconstructing species histories for groups that arose through recent,
58 rapid radiations are well established. Phylogenetic signal can be obscured by processes such as
59 incomplete lineage sorting (ILS) and gene flow (Maddison and Knowles 2006; Knowles and
60 Chan 2008; Christie and Knowles 2015), resulting in short branch lengths and conflicting gene
61 tree topologies. Consequently, approaches that use few loci or concatenation may fail to resolve
62 the most accurate species tree (Eckert and Carstens 2008; Leaché et al. 2014; Xi et al. 2014;
63 Giarla and Esselstyn 2015). This may be particularly common in plants that are thought to have
64 experienced rapid or recent radiation with ongoing hybridization and high levels of ILS. The
65 application of target enrichment methods that efficiently sequence hundreds of nuclear loci,
66 coalescent-based phylogenetic methods that account for ILS and gene flow, and extensive
67 sampling of morphologically diverse populations across the geographic range should allow for
68 more accurate representations of phylogenetic relationships (Maddison and Knowles 2006;
69 Knowles and Chan 2008; Knowles 2009; Mamanova et al. 2010; Lemmon et al. 2012; Straub et
70 al. 2012; Bryson et al. 2014; Weitemier et al. 2014; Mandel et al. 2014; Stephens et al. 2015;
71 Johnson et al. 2016).

72 *Oenothera* sect. *Calylophus* currently comprises seven species (thirteen taxa) with a
73 center of diversity in western Texas, southern New Mexico, and north-central Mexico (Fig. 1;
74 Towner 1977; Turner and Moore 2014; Wagner 2021). Previous analyses suggest that *Oenothera*
75 sect. *Calylophus* forms a well-supported, morphologically coherent clade with a relatively
76 restricted geographic range (Towner 1977; Levin et al. 2004; Wagner et al. 2007; Turner and
77 Moore 2014; Wagner 2021). However, as with other groups that have experienced rapid

78 radiations, taxon boundaries within *Oenothera* sect. *Calylophus* have been challenging to define,
 79 likely due to phenomena such as overlapping morphological boundaries, ongoing introgression,
 80 and incomplete lineage sorting.

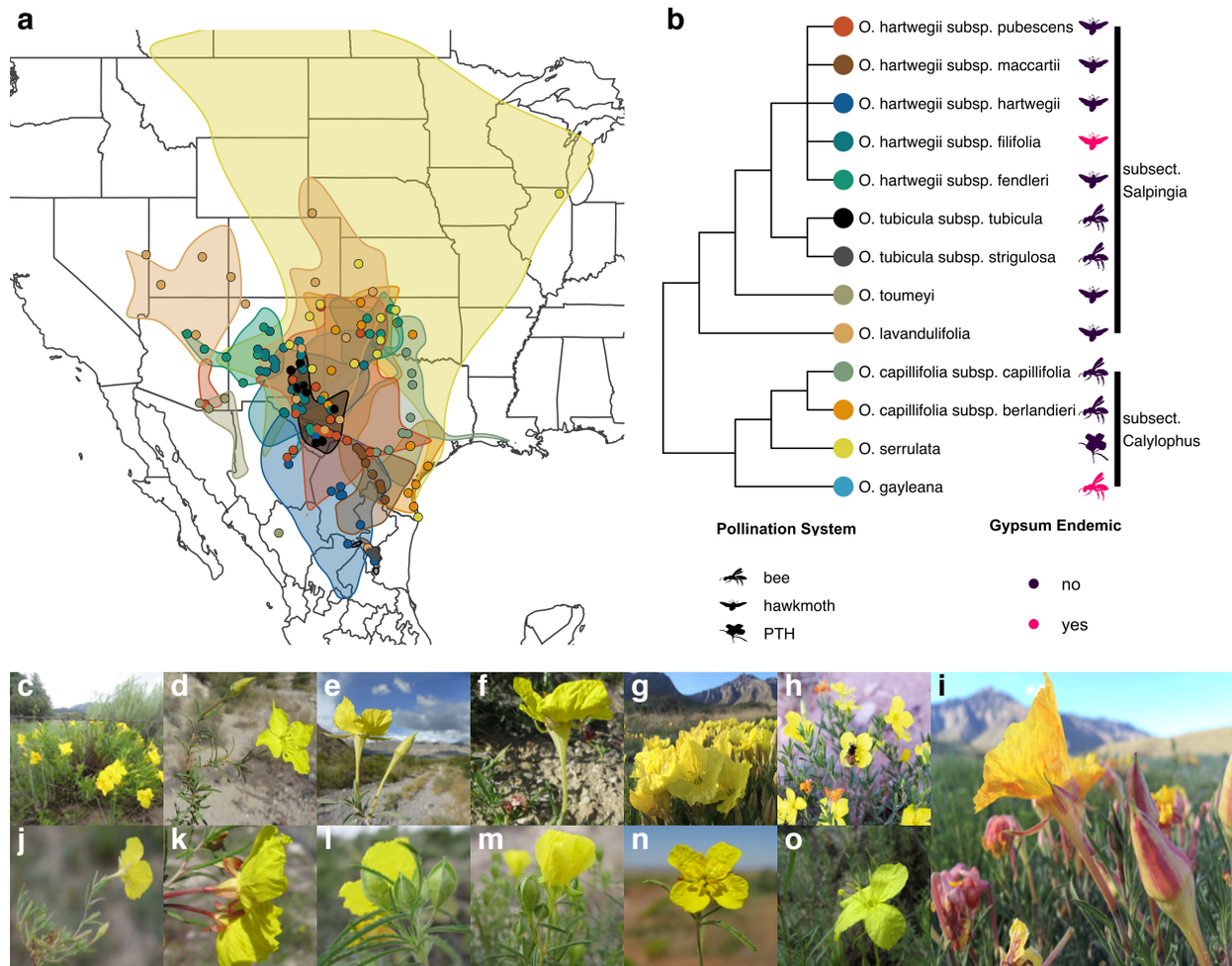


Figure 1

81 In the most comprehensive study of the group to date, Towner (1977) circumscribed taxa
 82 using morphology, breeding system, geography, and ecology, but it was noted (and our field
 83 observations confirm) that characters often overlap among taxa (Towner 1977). Taxa within
 84 *Oenothera* sect. *Calylophus* are divided into two easily recognizable subsections: subsect.
 85 *Salpingia* and subsect. *Calylophus* (Towner 1977; Wagner et al. 2007). Pollination varies

86 between the two subsections; flowers of subsect. *Salpingia* are adapted to vespertine pollination
87 by hawkmoths, except for *O. tubicula*, which opens in the morning and is primarily pollinated by
88 bees (Towner 1977), while taxa in subsect. *Calylophus* are predominantly bee-pollinated
89 (Towner 1977). Taxa in both subsections have geographic ranges that partially (or even largely)
90 overlap, resulting in occasional morphologically intermediate populations (Towner 1977).
91 Although confounding for morphological-based analyses, this observed pattern of reticulation is
92 consistent with a recent, rapid radiation occurring in parallel with climatic fluctuations and
93 increasing aridity in the region since the Pleistocene (Raven 1964; Towner 1977; Nason et al.
94 2002; Katinas et al. 2004). Hawkmoth pollination, which is ancestral in Onagraceae and common
95 in *Oenothera* sect. *Calylophus*, is known to result in long-distance pollen movement (Stockhouse
96 1973; Skogen et al. 2019). Therefore, gene flow may have been extensive over the evolutionary
97 history of hawkmoth-pollinated taxa, increasing the chances that processes such as historical
98 introgression may obscure phylogenetic signal in extant plants (Elrich and Raven 1969). With a
99 phylogenomic approach that samples hundreds of nuclear loci, we may better illuminate both the
100 history of these species and the key evolutionary processes related to speciation in this group.

101 Understanding speciation in angiosperms remains a fundamental question in evolutionary
102 biology (Barrett et al. 1996; Rajakaruna 2004; van der Niet et al. 2006; Wilson et al. 2007;
103 Crepet and Niklas 2009; Peakall et al. 2010; Xu et al. 2011; Van der Niet and Johnson 2012;
104 Boberg et al. 2014). *Oenothera* section *Calylophus* has an evolutionary history that likely
105 involves changes in reproductive system (pollinator functional group, breeding system) and
106 edaphic endemism. For example, there are thought to be two independent shifts between
107 pollinators from hawkmoth to bee pollination (Towner 1977; Fig. 1b), despite many studies in
108 other plant groups showing a directional bias in shifts from bee to hummingbird or hawkmoth

109 pollination (Barrett et al. 1996; Wilson et al. 2007; Thomson and Wilson 2008; Tripp and Manos
110 2008; Barrett 2013). However, pollinator shifts that do not follow this sequence may be more
111 likely when the extent of trait divergence and specialization does not completely inhibit
112 secondary pollinators such as bees, as has been suggested in *Oenothera* sect. *Calylophus*
113 (Stebbins 1970; Tripp and Manos 2008; Van Der Niet et al. 2014). Shifts to autogamy are also
114 frequent across angiosperms and in Onagraceae alone there are an estimated 353 shifts to modal
115 autogamy (i.e. mostly autogamous; Raven 1979). *Oenothera* sect. *Calylophus* also includes at
116 least one autogamous species, *O. serrulata*, which exhibits permanent translocation
117 heterozygosity, a phenomenon in which all chromosomes are translocationally heterozygous
118 (PTH; Towner 1977) (Fig. 1b). While the evolution of PTH has been assessed in molecular
119 phylogenetic analyses across Onagraceae (Johnson et al. 2009; Hollister et al. 2019), no study to
120 date has examined this transition in a well-sampled clade with extensive population sampling.
121 Lastly, abiotic ecological factors such as edaphic specialization are also known to drive
122 speciation in some groups (Rajakaruna 2004; van der Niet et al. 2006), including in *Oenothera*
123 sect. *Pachylophus* (Patsis et al. in press). For example, serpentine endemics represent ~10% of
124 the endemic flora in California even though serpentine soils account for about 1% of terrestrial
125 habitat in the state (Brady et al. 2005). Similarly, the Chihuahuan Desert is comprised of
126 numerous isolated islands of gypsum outcrops and current estimates suggest that at least 235 taxa
127 from 36 different plant families are gypsum endemics (Moore and Jansen 2007; Moore et al.
128 2014). It is suspected that gypsum endemism has also evolved independently in *Oenothera* sect.
129 *Calylophus* at least twice (Towner 1977; Turner and Moore 2014; Fig. 1b). Ultimately, to
130 understand the role that these transitions have played in shaping the diversity of *Oenothera* sect.
131 *Calylophus*, a robust phylogeny is required.

132 Here, we use target enrichment, summary coalescent methods, and morphometric
133 analyses to reconstruct a phylogenetic relationships, leveraging this information to re-examine
134 previous taxonomic concepts, test for instances of hybridization, and resolve the history of
135 pollinator shifts, PTH, and gypsum endemism in *Oenothera* sect. *Calylophus*. Target enrichment
136 is a cost-effective method for sequencing hundreds of loci across a high volume of samples,
137 producing highly informative datasets for phylogenetics (Lemmon et al. 2012; Straub et al. 2012;
138 Mandel et al. 2014; Weitemier et al. 2014; Heyduk et al. 2015; Stephens et al. 2015; Johnson et
139 al. 2016). While target enrichment is generally designed to capture coding regions, a significant
140 proportion of flanking non-coding regions can be recovered (the “splash-zone”; Weitemier et al.
141 2014). The inclusion of non-coding regions may be particularly informative for recent radiations,
142 since these regions are less constrained by selective pressures and may contain on average more
143 informative sites at shallower time scales (Folk et al. 2015). We included these flanking non-
144 coding regions in our sequence alignments to evaluate their impact on reconstructing lower-order
145 relationships. Importantly, we sampled extensively, including individuals from numerous
146 populations across the geographic and morphological ranges of all thirteen taxa in the section
147 (Fig. 1a). This study presents an example of how combining these molecular techniques with
148 dense sampling and morphological analysis can be used to effectively address a common
149 problem in evolutionary biology.

150

151 MATERIALS AND METHODS

152 A total of 194 individuals spanning the geographic, morphological, and ecological ranges
153 of all 13 recognized taxa in *Oenothera* sect. *Calylophus* [following Towner (1977) and Turner
154 and Moore (2014)] were included in this study (Fig. 1a, S1, S2) along with 8 outgroups

155 representing other major sections of *Oenothera* (*Eremia*, *Gaura*, *Kneiffia*, *Lavauxia*, *Oenothera*,
156 *Pachylophus*, and *Ravenia*) and other genera (*Chylismia* and *Eulobus*) in Onagraceae (S1, S2).
157 DNA was extracted from fresh, silica-dried leaf tissue (S3). PTH status was confirmed or
158 reassessed for individuals in subsect. *Calylophus* by assessing pollen fertility, when flowers were
159 present, using a modified Alexander stain (Alexander 1969, 1980; S3).

160 Target nuclear loci for enrichment were determined by clustering transcriptome
161 assemblies of *Oenothera serrulata* (1KP accession *SJAN*) and *Oenothera capillifolia* subsp.
162 *berlandieri* (1KP accession *EQYT*). Starting with the 956 phylogenetically informative
163 *Arabidopsis* loci identified by Duarte et al. (2010; S3), we identified 322 homologous, single-
164 copy loci in our clusters and used these in the probe design process. Libraries were enriched for
165 these loci using the MyBaits protocol (Arbor Biosciences, Ann Arbor, MI, USA) and sequenced
166 on an Illumina MiSeq (2 x 300 cycles, v3 chemistry; Illumina, Inc., San Diego, California,
167 USA). Raw reads have been deposited at the NCBI Sequence Read Archive (BioProject
168 PRJNA544074; See S3 for details). Reads were trimmed using Trimmomatic (Bolger et al. 2014;
169 S3) and trimmed, quality-filtered reads were assembled using HybPiper (Johnson et al. 2016).
170 From the assembled loci, we produced two datasets: “**exons**”, consisting of exon-only
171 alignments, and “**supercontig**”, consisting of alignments containing both the exon alignment and
172 flanking non-coding regions (the “splash-zone” per Weitemier 2014 and reconstructed using
173 supercontigs produced by HybPiper). We used these two datasets to test the most recent
174 taxonomic circumscription of the group with several methods: (1) phylogenetic inference of
175 concatenated alignments (two analyses: exons and supercontigs) using RAxML (Stamatakis
176 2014), (2) ASTRAL-II (Mirarab and Warnow 2015; Sayyari and Mirarab 2016) species tree
177 inference (two analyses: exons and supercontigs), (3) SVD Quartets (Chifman and Kubatko

178 2014, 2015) species tree inference (one analysis: supercontigs), (4) Phyparts (Smith et al. 2015;
179 one analysis: supercontigs), (5) IQtree (Minh et al. 2018) with both gene and site concordance
180 factors (one analysis: supercontigs).

181 We used HyDe (Blischak et al. 2018) to test for putative hybrid origins of selected taxa
182 and accessions by calculating D-Statistics (Green et al. 2010) for a set of hypotheses (S3). To
183 further characterize population-level processes or genetic structure within sect. *Calylophus*, we
184 extracted and filtered SNPs by mapping individual reads against reference supercontigs (see
185 <https://github.com/lindsawi/HybSeq-SNP-Extraction>) and used Discriminant Analysis of
186 Principal Components (Jombart et al. 2010) as implemented in the R package *adeigenet* (Jombart
187 2008) and the *snmf* function in the LEA package (Frichot and François 2015) in R (R Core
188 Team, 2020; S3).

189 We evaluated current taxonomic concepts and patterns of morphological variation by
190 measuring character states for vegetative and floral structures that have been used historically to
191 discriminate taxa in *Oenothera* sect. *Calylophus* (Towner 1977): plant height, leaf length (distal),
192 leaf width (distal), leaf length/width ratio (distal), leaf length (basal), leaf width (basal), leaf
193 length/width (basal), sepal length, and sepal tip length (S3). Measurements were made for 125 of
194 the sequenced samples (S11); we were unable to measure all traits for 73 samples because we did
195 not have access to the herbarium vouchers, or the trait of interest was not captured on the
196 voucher, therefore some samples were dropped from the analysis due to missing values. Finally,
197 the number of transitions and inferred ancestral conditions of reproductive system were mapped
198 onto an ASTRAL species tree, with individuals grouped into species, using the stochastic
199 mapping function in the R package *phangorn* version 2.5.5 (Schliep 2011; S3).

200

201 RESULTS AND DISCUSSION

202 *Sequencing and Phylogenetic Results*

203 Sequencing resulted in a total of 80,273,296 pairs of 300-bp reads with an average of
204 625,323 reads per sample. Following quality filtering, assembly and alignment, we recovered
205 204 loci that were present in at least 70% of the samples. Across all datasets and analyses,
206 *Oenothera* sect. *Calylophus* was monophyletic. At the subsection level there was strong
207 agreement in topology between concatenation and coalescent-based trees. For example, subsect.
208 *Calylophus* was recovered as sister to *Oenothera* subsect. *Salpingia* (minus *O. toumeyi*) with
209 strong support across all analyses and *O. toumeyi* [considered by Towner (1977) to be in subsect.
210 *Salpingia*; Fig. 2, S4-7] was recovered as sister to subsect. *Calylophus* across all trees, with
211 strong bootstrap support. Within subsect. *Calylophus* there was poor resolution for currently
212 recognized taxa in all analyses, whereas taxon relationships were better resolved in subsect.
213 *Salpingia* (Fig. 2, S4-7). With coalescent-based tree reconstruction, most taxa *sensu* Towner
214 (1977) were recovered as monophyletic with moderate to strong support (Fig. 2, S6-8). In
215 contrast, both the exon and supercontig concatenation trees recovered most currently recognized
216 taxa as non-monophyletic (S4, S5). Given that concatenation has been shown to produce
217 incorrect topologies in the presence of high ILS (Roch and Steel 2015) and that *Oenothera* sect.
218 *Calylophus* underwent recent radiation, we believe the paraphyly of taxa in both concatenation
219 trees might be artifactual. We therefore interpret relationships based on our coalescent-based
220 trees, which comprise the focus for the remainder of the paper (Fig. 2, S6-8).

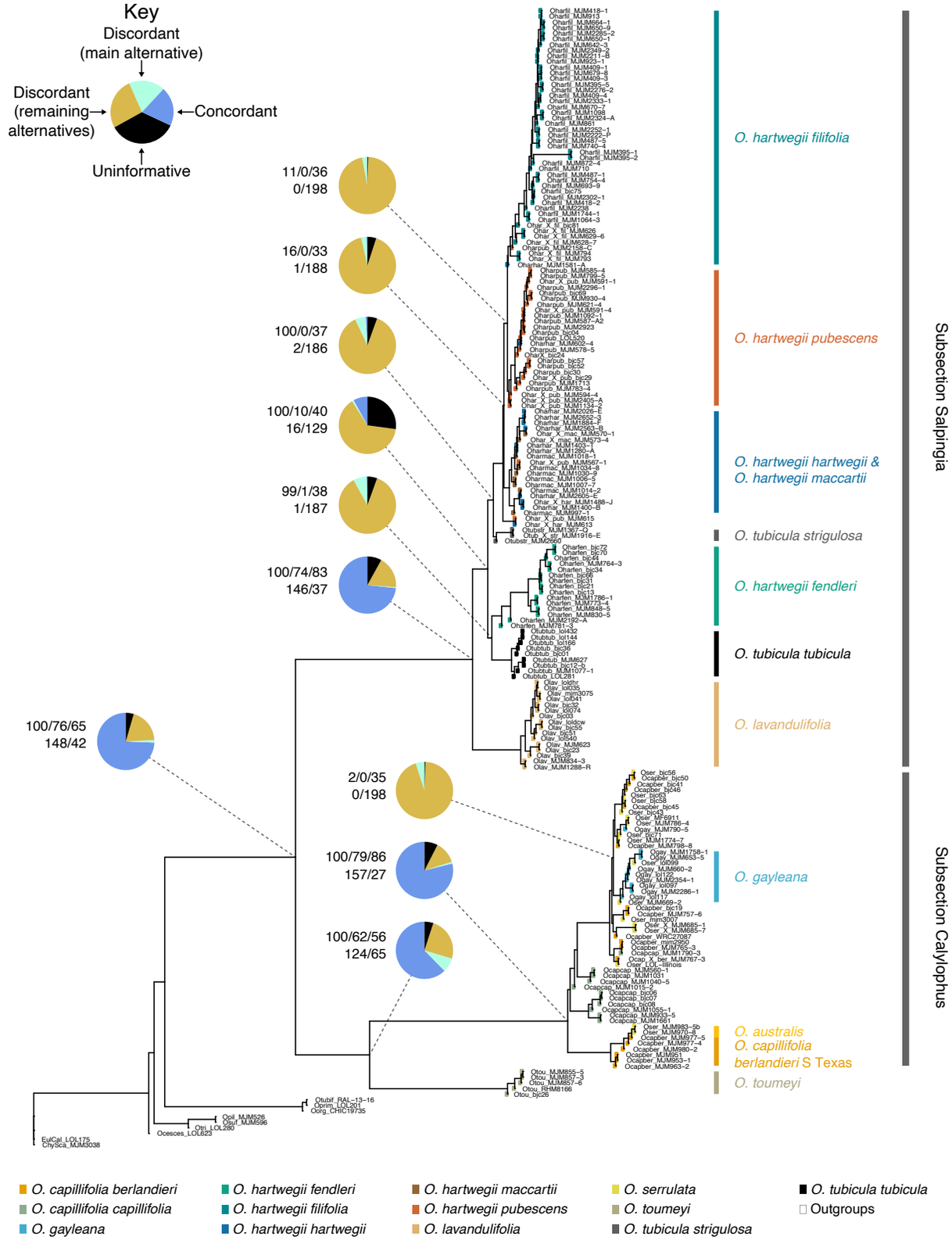


Figure 2

221 To understand whether summary coalescent relationships display a consistent signal
222 across the genome, we quantified gene tree and site concordance using Phyparts and IQtree. We
223 found that gene tree concordance was highest at the deepest nodes at the species level where we
224 expected less ILS and more time between speciation events (Fig. 2, S8, S9). Correspondingly,
225 gene tree concordance was lowest at the subspecies level where increased sharing of ancestral
226 alleles and ongoing gene flow are more likely (Fig. 2, S8, S9). For example, within *O. hartwegii*
227 less than 1% of genes were concordant for bifurcations representing all currently recognized taxa
228 at the subspecies level (Fig. 2, S8, S9). For species-level nodes with high support and high gene
229 tree concordance, site concordance was also high; for example, *O. lavandulifolia* had high
230 bootstrap support in summary coalescent trees (BS = 100), high gene tree concordance (Phyparts
231 = 94% concordance, gCF = 74), and high site concordance (sCF = 83; Fig. 2, S9h). For
232 subspecies with high support, but low gene tree concordance, site concordance was moderate.
233 For example *O. hartwegii* subsp. *fendleri* had high bootstrap support (BS = 99), low gene tree
234 concordance (Phyparts = <1% concordance, gCF = 1), and moderate site concordance (sCF = 38;
235 Fig. 2, S9b). For subspecies that were monophyletic in our coalescent-based trees, but that had
236 low support and low gene tree concordance, site concordance was moderate with an average of
237 35% of sites in agreement for taxa at these nodes (S9[c,f,g,v]). For example *O. hartwegii* subsp.
238 *filifolia* had low bootstrap support, low gene tree concordance (Phyparts = <1% concordance,
239 gCF = 0), and moderate site concordance (sCF = 36; Fig. 2, S9c). This is an important finding
240 because while species tree relationships can be obscured by ILS, site concordance factors, which
241 may be less constrained and less subject to ILS at shallower evolutionary timescales, provide a
242 key alternative method of support (Minh et al. 2018).

243 In general, topologies of exon and supercontig datasets were similar, with no major
244 differences in clade membership, but the inclusion of the flanking non-coding regions increased
245 support at shallow nodes in our trees. However, this trend was not universal. For example, in
246 subsect. *Salpingia*, using the supercontig dataset decreased support slightly for one taxon (*O.*
247 *hartwegii* subsp. *filifolia*), and in subsect. *Calylophus* it led to paraphyly of another (*O.*
248 *capillifolia* subsp. *capillifolia*). For six other taxa, our results showed that using supercontigs
249 increased bootstrap support. Therefore, these results demonstrated a net benefit of including
250 flanking non-coding regions for resolving relationships among closely related taxa.

251

252 *Hybridization and Geneflow*

253 Using concatenated loci from the supercontig dataset, we used HyDe (Blischak et al.
254 2018) to test for signals of hybridization. We used 552,521 sites and tested 22 hypotheses for
255 either individuals or groups suspected to be of hybrid origin based on field observations of
256 morphological intermediacy, geographic location, and topological position in our coalescent-
257 based trees and found evidence of hybridization in three individuals representing two taxa, both
258 in subsect. *Salpingia*. The highest signal of hybridization, with a gamma value ($\hat{\gamma}$) of 0.947
259 suggesting more historic gene-flow, was observed in one of three sampled individuals of *O.*
260 *tubicula* subsp. *strigulosa* (*MJM1916.E*). This involved admixture between *O. tubicula* subsp.
261 *tubicula* and the clade consisting of *O. hartwegii* subsp. *hartwegii* and *O. hartwegii* subsp.
262 *maccartii* (Z-score = 5.585, p-value = 0.000, $\hat{\gamma}$ = 0.947; Fig. 3a, S10). We also detected
263 significant levels of hybridization, with $\hat{\gamma}$ ranging from 0.332 to 0.338 suggesting more
264 contemporary gene-flow, in two individuals in *O. hartwegii* subsp. *pubescens*, *BJC29* (Z-score =
265 2.378, p-value = 0.009, $\hat{\gamma}$ = 0.338) and *MJM594* (Z-score = 2.094, p-value = 0.018, $\hat{\gamma}$ = 0.332).

266 This more recent gene flow involved admixture between *O. hartwegii* subsp. *pubescens* and the
267 clade consisting of *O. hartwegii* subsp. *hartwegii* and *O. hartwegii* subsp. *maccartii* (Fig. 3a,
268 S10).

269 The finding that one individual of *O. tubicula* subsp. *strigulosa* may be of hybrid origin is
270 consistent with gene flow between *O. tubicula* subsp. *strigulosa* and its sister taxon *O. tubicula*
271 subsp. *tubicula* (sensu Towner 1977). In our coalescent-based analyses, the two subspecies of *O.*
272 *tubicula* were not recovered as sister taxa, and this relationship was strongly supported (S9[j-k]).
273 If the two *O. tubicula* taxa arose independently, this would support the hypothesis that bee
274 pollination arose in *Oenothera* sect. *Calylophus* independently three times. However, while the
275 summary coalescent analyses we utilized to estimate phylogenies accounted for ILS in tree
276 estimation, they did not account for gene flow (Meng and Kubatko 2009; Gerard et al. 2011;
277 Kubatko and Chifman 2019). Our HyDe results may support the hypothesis that *O. tubicula*
278 subsp. *strigulosa* has experienced gene flow from two closely related taxa, and may have hybrid
279 origins resulting from crossing between *O. tubicula* subsp. *tubicula* and *O. hartwegii* subsp.
280 *hartwegii* (Fig. 3a, S10). This is consistent with Towner's interpretation of this taxon, which he
281 hypothesized may represent a stabilized derivative of introgression between *O. tubicula* subsp.
282 *tubicula* and *O. hartwegii* subsp. *hartwegii* (Towner 1977). Therefore, the placement of *O.*
283 *tubicula* subsp. *strigulosa* as sister to the rest of the *O. hartwegii* species complex in our trees
284 may result from past gene flow and hence may not represent independent origins of bee
285 pollination in subsect. *Salpingia*. These results underscore the importance of explicitly including
286 tests for hybridization in phylogenetic studies. In the case of these data, estimating a species tree
287 given a set of gene trees within a coalescent framework without considering other non-ILS
288 sources of signal conflict could artificially inflate the number of inferred evolutionary transitions.

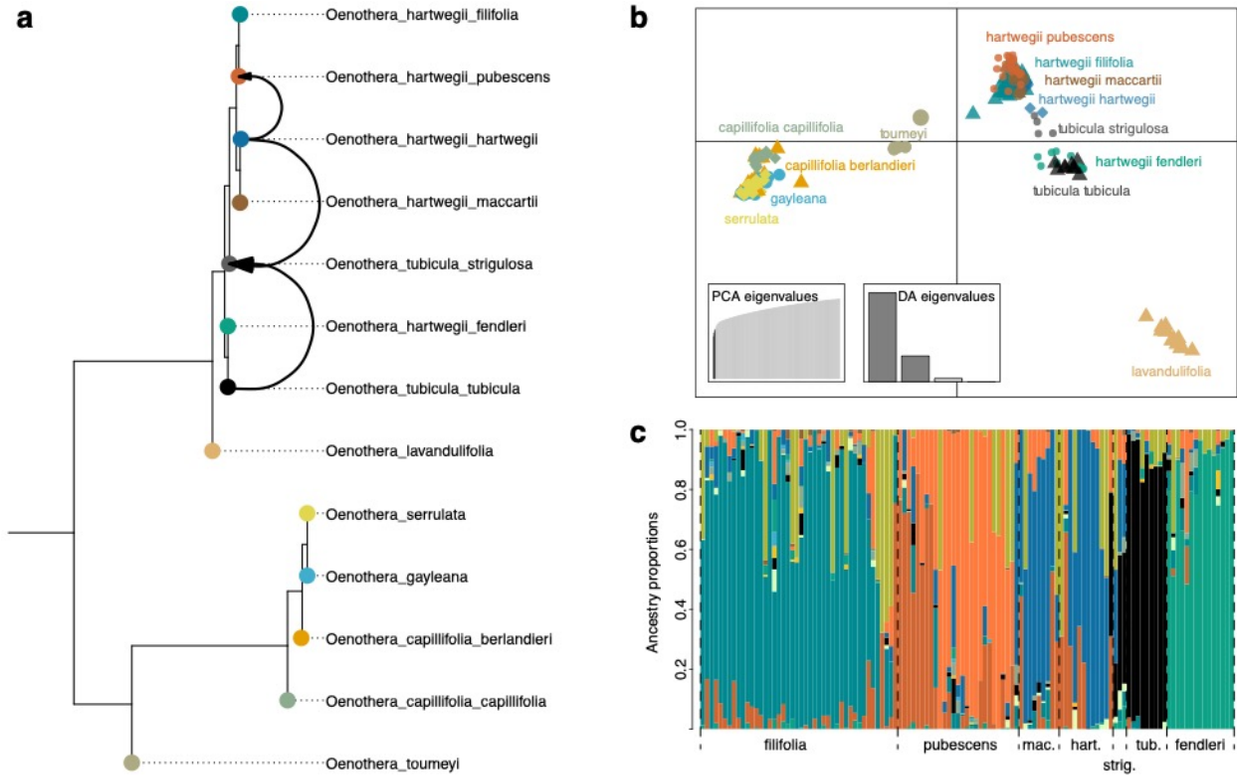


Figure 3

289 Our HyDe results also suggest that at least some of the morphological intermediacy and
 290 overlap among taxa in the group is due to continued, or at least recent, gene flow. For example,
 291 both *O. hartwegii* subsp. *pubescens* individuals that are inferred to have significant levels of
 292 admixture were collected from morphologically intermediate populations of *O. hartwegii* subsp.
 293 *pubescens* and *O. hartwegii* subsp. *hartwegii*. In addition, *O. hartwegii* subsp. *hartwegii* was a
 294 parent in all three hybridization events (S10). Thus, gene flow may explain this taxon's non-
 295 monophyly in our summary coalescent results. However, despite the often confounding patterns
 296 of overlapping morphological variation among closely related taxa in subset. *Salpingia*, this
 297 pattern does not necessarily appear to be the result of admixture, as many of the tests for
 298 hybridization based on field observations were not significant (Fig 3a, S10). What is also clear
 299 from these results is that much like collecting hundreds of nuclear genes provides a more

300 nuanced picture of phylogenetic signal and taxon relationships, our results show that collecting
301 multiple individuals from across the geographic and morphological ranges is necessary for a
302 more complete picture of relationships among closely related taxa.

303 After filtering, we extracted a set of 9,728 single nucleotide polymorphisms (SNPs) from
304 both coding and non-coding regions. A Discriminant Analysis of Principal Components (DAPC;
305 Fig. 3b) using these data clearly distinguishes *Oenothera* subsect. *Salpingia* from *O.* subsect.
306 *Calylophus*, with *O. toumeyi* intermediate between the two, which is consistent with the
307 phylogenetic results presented here. Additionally, the DAPC identifies *O. lavandulifolia* as a
308 distinct genetic cluster from the remaining taxa in subsection *Salpingia*. The overlap between
309 taxa, for example between the remaining taxa in subsection *Salpingia*, is consistent with the high
310 levels of gene tree discordance identified by PhyParts (Fig. 3a). For this latter group of taxa, we
311 computed estimates of ancestry coefficients using snmf, which suggests a substantial amount of
312 shared ancestral polymorphisms while also showing some evidence of clear genetic structure
313 among taxa (Fig. 3c). Consistent with the phylogenetic analyses, there does not appear to be any
314 clear genetic distinction between *O. hartwegii* subsp. *hartwegii* and *O. hartwegii* subsp.
315 *maccartii*, whereas *O. hartwegii* subsp. *fendleri*, *O. tubicula* subsp. *tubicula*, and *O. hartwegii*
316 subsp. *filifolia* appear to be largely distinct.

317

318 *Morphological Analysis*

319 We conducted morphometric Principal Components Analysis (PCA) to determine if
320 morphological patterns were consistent with phylogenetic results and to examine if specific
321 characters could be used to diagnose taxa as circumscribed by our phylogenetic analysis. Towner
322 (1977) observed overlapping and confounding patterns of morphological variation among taxa

323 within subsections, particularly within the *O. hartwegii* species complex. Despite this, because
 324 some taxa (e.g., *O. hartwegii* subsp. *fendleri*) were strongly supported by our summary
 325 coalescent trees we expected that they would be well distinguished in morphometric analysis.

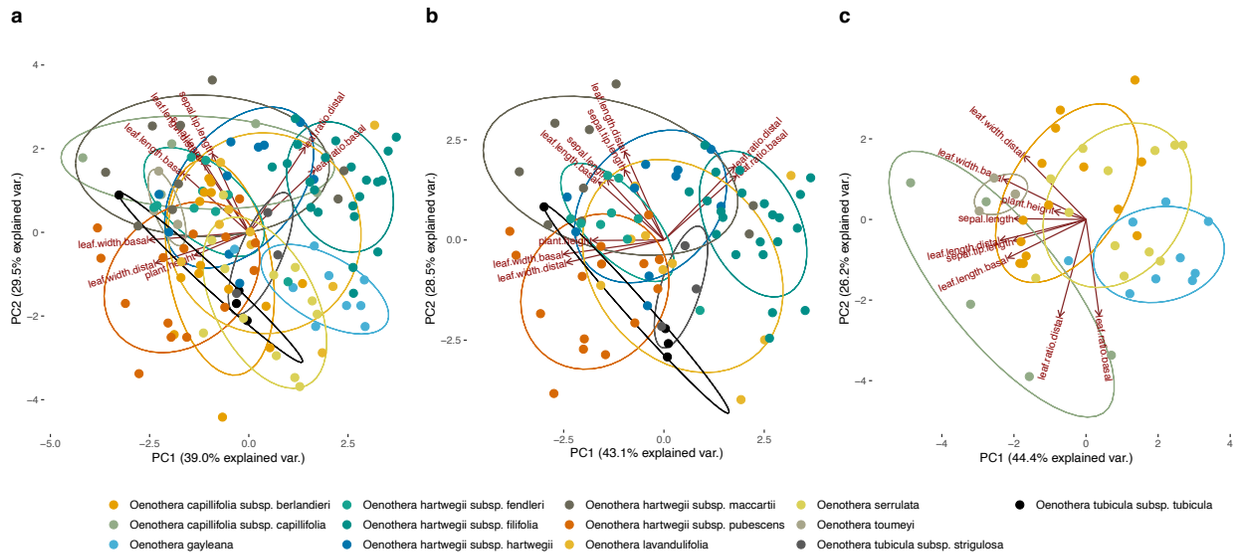


Figure 4

326 The main traits that separated taxa in subsect. *Salpingia* were leaf traits and plant size, while
 327 in subsect. *Calylophus* the main traits that separated taxa were sepal traits. In subsect. *Salpingia*,
 328 PC1 accounted for 43.1% of variance in PCA, while PC2 accounted for 28.5% (Fig. 4b).
 329 Morphological characters most associated with PC1 were leaf width (distal and basal), plant
 330 height, and leaf length/width ratio (distal and basal). Those associated with PC2 were leaf length
 331 (distal and basal), sepal length, and sepal tip length (Fig. 4b). In subsect. *Calylophus*, PC1
 332 accounted for 44.4% of explained variance and PC2 accounted for 26.2% (Fig. 4c). The
 333 characters most associated with PC1 in subsect. *Calylophus* include leaf length (distal and basal),
 334 sepal length, and sepal tip length. Those most associated with PC2 were leaf length/width ratio
 335 (distal and basal) and distal leaf width (Fig. 4c).

336 Our results support Towner's understanding of taxon boundaries by underscoring previous
337 difficulties in identifying individuals in this group based on morphology (Towner 1977). We
338 found substantial overlapping morphological variation among currently recognized taxa in both
339 subsections, though some taxa exhibited better grouping than others. The amount of overlap
340 between taxa was not a function of the strength of tree support for a given taxon in our summary
341 coalescent results. For example, *O. hartwegii* subsp. *fendleri*, a taxon that was well supported in
342 our summary coalescent trees, exhibited some of the highest degree of overlap with other taxa in
343 PCA space. Conversely, both *O. hartwegii* subsp. *filifolia* and *O. hartwegii* subsp. *pubescens*,
344 two taxa that formed poorly supported clades in our trees, formed clusters on the outer edges of
345 PCA space and had less overlap than other taxa (Fig. 4b). Interestingly, *O. hartwegii* subsp.
346 *hartwegii*, the taxon that was identified as a parent in all three instances of admixture in our
347 HyDe analysis, also overlaps morphologically with most other taxa in subsect. *Salpingia* (Fig.
348 4b). This is not surprising given that it is widely distributed in northern Mexico and western
349 Texas and frequently comes into contact with related taxa resulting in sympatric populations and
350 occasional morphologically intermediate populations.

351

352 *Implications for Reproductive Systems and Edaphic Endemism*

353 Our results show that shifts from hawkmoth to bee pollination likely occurred twice in
354 sect. *Calylophus* (S3) and thus may be more common in *Oenothera* than previously thought. The
355 strongly supported sister relationship of *O. toumeyi* to remaining subsect. *Calylophus* in our
356 summary coalescent results is consistent with two independent shifts to bee pollination, once in
357 the ancestor of subsect. *Calylophus*, and another in subsect. *Salpingia* on the branch leading to
358 *O. tubicula* (Fig. 2). Independent shifts to bee pollination from hawkmoth pollination are perhaps

359 not surprising considering that within sect. *Calyophus*, hawkmoth-pollinated floral forms exhibit
360 variation between populations in hypanthium length and diameter and do not prevent occasional
361 pollination by bees (Lewis 2015; Towner 1977). Hawkmoth-pollinated taxa in sect *Calylophus*
362 exhibit vespertine anthesis, which separates them temporally from diurnal bees, but variation in
363 the timing of anthesis is also common between populations (Towner 1977), and hawkmoths are
364 documented to vary greatly in abundance spatiotemporally (Miller 1981; Campbell et al. 1997;
365 Artz et al. 2010). In cases of pollen limitation, night-blooming plants benefit from bimodal
366 pollination between moths and bees by acquiring pollinator assurance against yearly variation or
367 local extinction of specific pollinators, as shown for other *Oenothera* species (Barthell and
368 Knops 1997; Artz et al. 2010), *Lonicera japonica* (Miyake and Yahara 1998) and for night-
369 blooming *Ancistrophora* cacti (Schlumpberger et al. 2009).

370 Additionally, it has been shown that florivore-mediated selection drives floral trait shifts
371 in sect. *Calylophus* towards bee-pollinated floral forms (Jogesh et al. 2017; Bruzzese et al. 2019).
372 Variation in reproductive traits that allows some continued pollination by bees provides an
373 alternative mode of pollen transfer and may represent a mechanism for ensuring pollination.
374 While studies have shown that pre mating barriers contribute greatly to reproductive isolation
375 (Stanton et al. 2016), our results show that multiple, independent shifts from hawkmoth to bee
376 pollination and associated morphological changes, such as the shorter floral tube length of bee
377 pollinated flowers, may occur in sect. *Calylophus*, and hence may not be a particularly reliable
378 character for diagnosing taxa in this group.

379 Stochastic mapping (supplemental) suggests that there are multiple origins of permanent
380 translocation heterozygosity (PTH) in sect. *Calylophus* While ring chromosomes are common
381 and found in all taxa in sect. *Calylophus*, PTH is currently known from only one taxon, *O.*

382 *serrulata*. Because neighboring populations of *O. serrulata* and its putative progenitor *O.*
383 *capillifolia* subsp. *berlandieri* often resemble each other phenetically, Towner (1977)
384 hypothesized that *O. serrulata* may have originated multiple times through independent origins
385 of translocation heterozygosity in different geographic regions, and may be best recognized as “a
386 complex assemblage of populations having a common breeding system.” However, this has
387 never before been explored in a phylogenetic context, nor has it been clearly demonstrated with
388 phylogenetic studies in Onagraceae. In our summary coalescent trees, all currently recognized
389 taxa in subsect. *Calylophus* were paraphyletic and *O. serrulata* was scattered throughout the
390 subsection (Fig. 2, S6, S7). Although support values are not always high for the positions of
391 various individuals of *O. serrulata*, there is at least one well defined, well supported split among
392 populations of *O. serrulata*. In our summary coalescent trees, the two *O. serrulata* accessions
393 from south Texas (*MJM970* & *MJM983*) grouped with other south Texas populations of *O.*
394 *capillifolia* subsp. *berlandieri* with generally strong support (Fig. 2, S6, S7). This relationship
395 was supported in PCA space as well, where *MJM983* was morphologically more similar to the
396 south Texas *O. capillifolia* subsp. *berlandieri* accessions than to other *O. serrulata* (Fig 4C). Our
397 results are therefore consistent with an independent origin of PTH in coastal Texas populations
398 of *O. serrulata*, demonstrating at minimum two origins of PTH (see *Taxonomic Implications*
399 below). However we cannot rule out other independent origins of PTH. Given the prevalence of
400 translocations among partial sets (i.e. not all seven) of homologous chromosomes in *O.* section
401 *Calylophus* (Towner 1977), and if this could be considered an intermediate step towards
402 “complete” PTH, perhaps it is not surprising to reconstruct multiple origins. Our results suggest
403 that a more complete assessment of the extent and distribution of this phenomenon in section
404 *Calylophus* is needed.

405 Independent origins of gypsum endemism in sect. *Calylophus* are also supported by our
406 analyses (supplemental). Edaphic specialization is a fundamental driver of speciation in plants
407 and contributes greatly to endemism and species diversity in areas with geologically distinct
408 substrates such as gypsum and serpentine outcrops (Kruckeberg 1984; Anacker et al. 2011;
409 Cacho and Strauss 2014; Moore et al. 2014). To date, two gypsum endemic taxa have been
410 described in sect. *Calylophus*, one in each subsection: *O. hartwegii* subsp. *filifolia*, which is
411 relatively widespread on gypsum in New Mexico and trans-Pecos Texas and only rarely
412 sympatric with other taxa, and the recently described *O. gayleana*, which is found in
413 southeastern New Mexico and adjacent western Texas, with disjunct populations in northern
414 Texas and western Oklahoma (Turner and Moore 2014). Despite low support and low gene tree
415 congruence in our analyses, the two gypsum endemic taxa had moderate sCF support (Fig. 2,
416 S9), much like other taxa with similarly low support and high levels of discordance. In
417 addition, while both gypsum endemics overlapped with other taxa in the morphometric analysis,
418 they occupied morphological extremes in PCA space (Fig. 4). Given that other well-supported
419 taxa also overlap morphologically, it is perhaps not surprising that the two gypsum endemic taxa
420 are not more differentiated from other taxa morphologically. Perhaps the strongest evidence in
421 our data for their recognition as distinct taxa is that we found no evidence of admixture between
422 either of these gypsum endemic taxa and other closely related taxa (Fig. 3a, S10).

423

424 *Taxonomic Implications*

425 The most consequential taxonomic result that arises from our analyses is the position of
426 *O. toumeyi*, a member of subsect. *Salpingia* as circumscribed by Towner (1977). In the present
427 study, this species is resolved as sister to subsect. *Calylophus* with strong support, rendering

428 subsect. *Salpingia* paraphyletic (Fig. 2, S9). Towner (1977) grouped *O. toumeyi* with *O.*
429 *hartwegii* due to similar floral and bud characters including large flowers and long floral tubes
430 suggestive of hawkmoth pollination, and rounded buds with long, free sepal-tips. Because the
431 breeding system is a defining difference in the current circumscription between the two
432 subsections in sect. *Calylophus*, this result supports abandoning subsections altogether in sect.
433 *Calylophus*.

434 Within subsect. *Salpingia* our results also suggest the need for revision. While our
435 phylogenetic analyses strongly support the current circumscription of *O. lavandulifolia* (sensu
436 Towner 1977) as a distinct species within subsect. *Salpingia* (Fig. 2, S9), the relationships of the
437 other two species *O. hartwegii* and *O. tubicula* are less clear. Towner (1977) differentiated these
438 two species by the breeding system and grouped the five subspecies of *O. hartwegii* together
439 based on a pattern of reticulate and intergrading variation in which taxa were distinguished from
440 one another by often slight differences in pubescence and leaf shape. Our morphometric analysis
441 confirmed this pattern; however, our phylogenetic results indicated that one taxon, *O. hartwegii*
442 subsp. *fendleri*, shares a closer relationship with the bee pollinated *O. tubicula* subsp. *tubicula*
443 than other taxa in the hawkmoth pollinated *O. hartwegii* species complex (Fig. 2, S6, S7). This
444 relationship was strongly supported and renders *O. hartwegii*, according to the current
445 circumscription, paraphyletic (Towner 1977). Based on strong phylogenetic support for this
446 clade, and its strong morphological distinctiveness as described by Towner (1977), we suggest
447 that *O. hartwegii* subsp. *fendleri* be elevated to the species rank along with both subspecies of *O.*
448 *tubicula* which were equally well supported in phylogenetic analysis and are geographically
449 isolated. Furthermore, our results support a the possible elevation to species rank for *O.*
450 *hartwegii* subsp. *filifolia*. While this taxon was poorly supported in our summary coalescent trees

451 (Fig. 2, S6, S7), we found no evidence of hybridization between this taxon and other closely
452 related taxa. In addition, *O. hartwegii* subsp. *fillifolia* is restricted to gypsum. Therefore, we
453 believe that the ecological distinctiveness and lack of gene flow of *O. hartwegii* subsp. *fillifolia*
454 with other taxa in the *O. hartwegii* species complex warrants its elevation as a distinct species. In
455 light of these changes, and to maintain consistency in classification in the subsection, we feel
456 that despite the evidence of hybridization of *O. hartwegii* subsp. *pubescens* with *O. hartwegii*
457 subsp. *hartwegii*, it possesses a morphological distinctiveness that is supported by our
458 phylogenetic results. We therefore recommend *O. hartwegii* subsp. *pubescens* be elevated to the
459 species level, while *O. hartwegii* subsp. *hartwegii* and *O. hartwegii* subsp. *maccartii* be retained
460 as is, forming a polytypic species with two subspecies.

461 In contrast to the relatively clear divisions among taxa in subsect. *Salpingia*, none of the
462 four currently recognized taxa in the subsect. *Calylophus* were consistently recovered as
463 monophyletic. For example, *O. capillifolia* subsp. *capillifolia* was monophyletic in our exon-
464 only summary coalescent tree, but not in the “supercontig” tree, and *O. capillifolia* subsp.
465 *berlandieri* and *O. serrulata* were scattered throughout sect. *Calylophus* in both trees, perhaps
466 suggesting widespread gene flow and/or multiple origins of PTH (Fig. 2, S6). Importantly, our
467 results suggest that the circumscription of *O. gayleana* sensu Turner and Moore (2014) should be
468 amended. Specifically, we find that the populations of subsect. *Calylophus* from northern Texas
469 and western Oklahoma that were assigned to *O. gayleana* by Turner and Moore (2014; *MJM790-*
470 *5, BJC71*) may instead may belong to *O. serrulata* based on both their phylogenetic positions
471 (Fig. 2, S6, S7) and reduced pollen fertility (S12). These north Texas/western Oklahoma
472 populations seem to represent slightly narrower-leaved individuals of *O. serrulata*, which is a

473 common inhabitant of the extensive gypsum outcrops of this area (although it is not restricted to
474 gypsum there).

475 Finally, our results highlight an unrecognized cryptic taxon within *O. capillifolia* formed
476 by southern Texas coastal populations currently recognized as *O. capillifolia* subsp. *berlandieri*.
477 Towner (1977) described *O. capillifolia* as a polytypic species with two well-differentiated
478 morphological races. Though he noted the geographic and cytological distinction of the southern
479 Texas coastal populations of *O. capillifolia* subsp. *berlandieri*, these populations were included
480 in *O. capillifolia* subsp. *berlandieri* primarily because of completely overlapping morphological
481 variation. In our results, this cryptic clade of southern Texas coastal populations of *O. capillifolia*
482 subsp. *berlandieri* is the most phylogenetically well supported clade in subsect. *Calylophus* and
483 therefore may warrant taxonomic distinction based on our data (Fig. 2, S9r). Similarly, the
484 southern Texas coastal populations of *O. serrulata*, which is likely an independent origin of PTH
485 derived from this cryptic southern Texas coastal clade of *O. capillifolia* subsp. *berlandieri*, are
486 ecologically distinctive and geographically disjunct from other *O. serrulata* (occurring in coastal
487 dunes, unlike other populations in western Texas, Oklahoma, and northern Texas). In the past
488 they were considered distinctive enough to be described as a species, *Calylophus australis*
489 (Towner & Raven 1970). However, Towner (1977) later combined this species with *O. serrulata*
490 based on his decision to treat all PTH populations as *O. serrulata*. Combined with our results
491 here and the ecogeographic distinctiveness consistent with an independent origin of PTH in
492 coastal Texas, we believe this taxon also warrants recognition as a second PTH species in
493 *Oenothera* sect. *Calylophus*.

494

495 CONCLUSIONS

496 Here we describe a robust example of resolving a recent, rapid radiation using multiple
497 sources of evidence: (1) extensive sampling from populations throughout the geographic and
498 morphological range, (2) target enrichment for hundreds of nuclear genes, (3) the inclusion of
499 flanking non-coding regions, (4) gene tree-based hybridization inference, (5) SNPs extracted
500 from target enrichment data, and (6) morphometrics. Our results indicate that in recently radiated
501 species complexes with low sequence divergence and/or high levels of ILS that could be an
502 intractable problem with traditional loci, the use of targeted enrichment in addition to flanking
503 non-coding regions provides a net benefit and is essential to recover species-level resolution. Our
504 results also underscore the importance of summary coalescent methods and evaluating gene tree
505 discordance for resolving historical relationships in recalcitrant groups. By explicitly testing for
506 hybridization using gene tree approaches, we also demonstrate that the estimated number of
507 character state transitions may be artifactually inflated if hybridization is not taken into account.
508 This, in combination with morphometrics, provided key evolutionary insights where
509 relationships in summary coalescent methods may be obscured by gene flow. Importantly, our
510 study uncovers strong evidence for multiple origins of biologically important phenomena,
511 including the evolution of bee pollination, PTH, and edaphic specialization. Consequently,
512 *Oenothera* sect. *Calylophus* might represent a powerful system for understanding these
513 phenomena, especially with future genome sequence data.

514

515 DATA AVAILABILITY

516 Illumina reads generated for this study are available at the NCBI Sequence Read Archive under
517 BioProject PRJNA544074. Assembled exon and supercontig sequences, multiple sequence
518 alignments, and SNP files are available at [http://dx.doi.org/10.5061/dryad.\[NNNN\]](http://dx.doi.org/10.5061/dryad.[NNNN]).

519

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543
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752 FIGURE LEGENDS

753

754 **Figure 1.** (a) Range map of all taxa in *Oenothera* sect. *Calylophus* (based on Towner 1977).

755 Sampling locations of leaf tissue samples (points; color corresponds to taxa in cladogram to the

756 right [Figure 1b]) and estimated taxon ranges (polygons; colors correspond to Figure 1b)

757 proposed by Towner (1977) and Turner and Moore (2014). (b) Estimated cladogram of

758 *Oenothera* sect. *Calylophus* sensu Towner (1977) and Turner and Moore (2014). Symbols to the

759 right of tip labels signify pollination system (bee, hawkmoth, or Permanent Translocation

760 Heterozygosity [PTH]) and the symbol color specifies whether a given taxon is a gypsum

761 endemic (purple = no, pink = yes). Photo panels: (c) *Oenothera hartwegii* subsp. *fendleri* (d)

762 *Oenothera hartwegii* subsp. *filifolia* (e) *Oenothera hartwegii* subsp. *hartwegii* (f) *Oenothera*

763 *hartwegii* subsp. *maccartii* (g) *Oenothera hartwegii* subsp. *pubescens* (h) *Oenothera tubicula*

764 subsp. *tubicula* (i) *Oenothera lavandulifolia* (j) *Oenothera tubicula* subsp. *strigulosa* (k)

765 *Oenothera capillifolia* subsp. *berlandieri* (l) *Oenothera capillifolia* subsp. *capillifolia* (m)

766 *Oenothera gayleana* (n) *Oenothera serrulata* (o) *Oenothera toumeyii*

767

768 **Figure 2.** ASTRAL-II summary coalescent tree constructed using the “Supercontig” dataset with
769 100 bootstraps. At relevant nodes, piecharts represent Phyparts analysis (blue = concordant,
770 green = most frequent conflict, yellow = all other conflict, black = uninformative gene trees), top
771 row of support values are bootstrap values from ASTRAL-II, and gCF and sCF from IQTree
772 (BS/gCF/sCF), bottom two support values are number of concordant gene trees for the node and
773 total number of gene trees minus the number of concordant gene trees at that node
774 (concord/discord). Colored tip points correspond to taxon designation.

775

776 **Figure 3.** (a) Summary of HyDe Analysis annotated on ASTRAL-III species tree constructed
777 using the “Supercontig” dataset; black arrows represent direction of admixture detected by HyDe
778 analysis. (b) Discriminant Analysis of Principal Components based on a filtered set of SNPs
779 extracted from the entire supercontig dataset, and (c) sNMF plot of inferred ancestry coefficients
780 using the same set of filtered SNPs but limited to subsection *Salpingia* (minus *O. lavandulifolia*)
781

781

782 **Figure 4.** Morphometric Principal Components Analysis (PCA) using 9 morphological
783 characters (plant height, leaf length [basal and distal], leaf width [basal and distal], leaf
784 length/width ratio (basal and distal), and sepal tip length) for (a) Section *Calylophus* (b)
785 Subsection *Salpingia* without *O. toumeyi*, and (c) Subsection *Calylophus* with *O. toumeyi*
786 included.

787 SUPPLEMENTAL MATERIAL

788 S1.

789 See excel table “S1 Accessions and Seq Stats”

790

791 S2.

792 Number of leaf tissue accessions sequenced from each taxon

Taxon	No. of Accessions ⁷⁹³	
<i>O. lavandulifolia</i>	16	794
<i>O. tubicula</i> subsp. <i>strigulosa</i>	3	795
<i>O. tubicula</i> subsp. <i>tubicula</i>	9	796
<i>O. hartwegii</i> subsp. <i>fendleri</i>	15	797
<i>O. hartwegii</i> subsp. <i>filifolia</i>	44	798
<i>O. hartwegii</i> subsp. <i>hartwegii</i>	12	799
<i>O. hartwegii</i> subsp. <i>maccartii</i>	9	800
<i>O. hartwegii</i> subsp. <i>pubescens</i>	26	801
<i>O. hartwegii</i>	1	802
<i>O. toumeyi</i>	5	803
<i>O. capillifolia</i> subsp. <i>berlandieri</i>	17	804
<i>O. capillifolia</i> subsp. <i>capillifolia</i>	11	805
<i>O. gayleana</i>	9	806
<i>O. serrulata</i>	17	807
Outgroups:		808
<i>O. pilosella</i>	1	809
<i>O. organensis</i>	1	810
<i>O. primiveris</i>	1	811
<i>O. tubifera</i>	1	812
<i>O. triloba</i>	1	813
<i>O. cespitosa</i> subsp. <i>cespitosa</i>	1	814
<i>O. suffrutescens</i>	1	815
<i>Chylismia scapoidea</i> subsp. <i>scapoidea</i>	1	816
<i>Eulobus californicus</i>	1	817
Total	203	817
		818

819 S3.

820 MATERIALS AND METHODS

821 *Taxon Sampling, DNA Extraction, and Determination of PTH*

822 A total of 194 individuals spanning the geographic, morphological, and ecological ranges
 823 of all 13 recognized taxa in *Oenothera* sect. *Calylophus* [following Towner (1977) and Turner
 824 and Moore (2014)] were included in this study (Fig. 1a, S1, S2) along with eight outgroups
 825 representing other major sections of *Oenothera* (*Eremia*, *Gaura*, *Kneiffia*, *Lavauxia*, *Oenothera*,
 826 *Pachylophus*, and *Ravenia*) and other genera (*Chylismia* and *Eulobus*) in Onagraceae (S1, S2).
 827 All leaf tissue samples were collected from individuals in the field between 2007 and 2015 and
 828 voucher specimens were deposited at the United States National Herbarium (US), with

829 duplicates in most cases at either the Nancy Rich Poole Herbarium (CHIC) or the George T.
830 Jones Herbarium at Oberlin College (OC; S1). DNA was extracted from fresh, silica-dried leaf
831 tissue using either (1) a modified CTAB protocol (Doyle 1987), (2) the Nucleon PhytoPure DNA
832 extraction kit (GE Healthcare Life Sciences, Pittsburgh, Pennsylvania, USA), or (3) a modified
833 CTAB and silicon dioxide purification protocol (Doyle 1987; Sharma and Purohit 2012; See S13
834 for details) followed by passing any extractions retaining a brown or yellow coloration through a
835 Qiagen Qiaquick PCR spin column for additional purification according to the manufacturer's
836 protocol (Qiagen, Venlo, Netherlands). The third DNA extraction method was used for difficult
837 to extract, polysaccharide-rich leaf tissue samples that yielded gooey, discolored DNA following
838 initial extraction. PTH status was determined for individuals in subsect. *Calylophus* using floral
839 morphology and/or when flowers were present by assessing pollen fertility using a modified
840 Alexander stain, as PTH taxa have a demonstrated 50% reduction in pollen fertility (Towner
841 1977). For accessions identified as either *O. capillifolia* subsp. *berlandieri* or *O. serrulata* that
842 had sufficient pollen available, pollen was removed from flowers and stained using a modified
843 Alexander stain (Alexander 1969, 1980). Accessions with less than 50% viable pollen were
844 assigned to *O. serrulata*, the only currently recognized PTH taxon in subsect. *Calylophus*. Pollen
845 count data are provided in Supplement 12 (See S12 for details).

846

847 *Bait Design, Library Construction, Target Enrichment, and Sequencing*

848 We targeted 322 orthologous, low-copy nuclear loci determined by clustering
849 transcriptomes of *Oenothera serrulata* (1KP accession *SJAN*) and *Oenothera berlandieri* (1KP
850 accession *EQYT*) to select a subset of the 956 phylogenetically informative *Arabidopsis* loci
851 identified by Duarte et al. (2010). Transcriptomes of two *Oenothera* species, *O. serrulata* and *O.*
852 *berlandieri*, were assembled and optimal isoforms were filtered for the longest reading frame
853 using custom Perl scripts. Assembled transcripts were aligned as amino acids to the 956 TAIR
854 loci of *Arabidopsis* in TranslatorX (Abascal et al. 2010). This alignment identified 956
855 orthologous sequences, from which 322 loci were randomly selected. BLAST searches of amino
856 acid sequences from these loci were carried out to ensure orthology between the transcript loci
857 and the *Arabidopsis* TAIR locus. The bait set was designed from these 322 loci, which were
858 selected from both *O. serrulata* and *O. berlandieri* sequences. A set of 19,994 120-bp baits tiled
859 across each locus with a 60 base overlap (2x tiling) was manufactured by Arbor Biosciences
860 (formerly MYcroarray, Ann Arbor, Michigan, USA). Sequencing libraries for 67 samples were
861 prepared with the Illumina TruSeq Nano HT DNA Library Preparation Kit (San Diego,
862 California, USA) following the manufacturer's protocol, except using half volumes beginning
863 with the second addition of Dynabeads MyOne Streptavidin C1 magnetic beads (Invitrogen,
864 Carlsbad, CA, USA). DNA samples were sheared using a Covaris M220 Focused-Ultrasonicator
865 (Covaris, Woburn, Maryland, USA) to a fragment length of ~550 bp (for an average insert size
866 of ~420 bp). The remaining 134 libraries were constructed by Rapid Genomics (Gainesville,
867 Florida, USA), with custom adapters. The Illumina i5 and i7 barcodes were used for all libraries.

868 Libraries were enriched for these loci using the MyBaits protocol (ArborBiosciences
869 2016) with combined pools of libraries totaling 1.2 µg of DNA (12 libraries/pool at 100
870 ng/library). Libraries with less than 100 ng of total recovered DNA were pooled together in
871 equimolar concentrations using available product, resulting in some pools with less than 1.2 µg
872 of DNA. The smallest successful pool contained four samples with 6 ng of library each.
873 Hybridization was performed at 65°C for approximately 18 hours. The enriched libraries were
874 reamplified with 14 to 18 PCR cycles and a final cleanup was performed using a Qiagen

875 QiaQuick PCR cleanup kit following the manufacturer's protocol to remove bead contamination
876 (Qiagen, Venlo, Netherlands). DNA concentrations were measured using a Qubit 2.0
877 Fluorometer (Life Technologies, Carlsbad, California, USA) and molarity was measured on an
878 Agilent 2100 Bioanalyzer (Agilent Technologies, Santa Clara, California, USA). A final
879 cleaning step using Dynabeads MyOne Streptavidin C1 magnetic beads was performed on pools
880 with adapter contamination as detected on the Bioanalyzer. Pools were sequenced in four runs at
881 equimolar ratios (4 nM), on an Illumina MiSeq (2 x 300 cycles, v3 chemistry; Illumina, Inc., San
882 Diego, California, USA) at the Pritzker DNA lab (Field Museum, Chicago, IL, USA). This
883 produced 80,273,296 pairs of 300-bp reads. Reads were demultiplexed and adapters trimmed
884 automatically by Illumina Basespace (Illumina 2016). Raw reads have been deposited at the
885 NCBI Sequence Read Archive (BioProject PRJNA544074).

886

887 *Quality Filtering, Assembly and Alignment*

888 A summary of read quality from each sample was produced using FastQC
889 (<http://www.bioinformatics.babraham.ac.uk/people.html>), which revealed read-through adapter
890 contamination in many of the poorer quality samples. To remove read-through contamination
891 and filter for quality, reads were trimmed for known Illumina adapters using Trimmomatic
892 (Bolger et al. 2014) with the following settings:

893 ILLUMINACLIP:<illumina_adapters.fasta>:2:30:10 LEADING:10 TRAILING:10
894 SLIDINGWINDOW:4:20 MINLEN:20. Trimmed, quality-filtered reads were assembled using
895 the HybPiper pipeline (Johnson et al. 2016) with default settings, followed by the intronrate.py
896 script, to produce both exons and the “splash zone” flanking non-coding region-containing
897 supercontigs. Only pairs with both mates surviving trimming and quality filtering were used for
898 HybPiper.

899 To compare the influence of “splash-zone” non-coding regions, two sets of alignments
900 were created: (1) exons alone, and (2) coding sequences plus the “splash-zone” (Hereby referred
901 to as supercontigs). For multiple sequence alignments of exons alone, protein and nucleotide
902 sequences assembled in HybPiper were gathered into fasta files by gene. For protein sequences
903 only, stop codons were changed to “X” using a sed command-line regular expression to facilitate
904 alignment, and sequences were aligned using MAFFT with settings: --auto --adjustdirection --
905 maxiterate 1000 (Katoh et al. 2002). Aligned protein sequences were then used to fit unaligned
906 nucleotide sequences into coding frame alignments using pal2nal with default settings (Suyama
907 et al. 2006). In-frame, aligned DNA sequences were trimmed to remove low-coverage positions
908 and sequences composed only of gaps using TrimAl with the automated setting, which is
909 optimized for maximum likelihood analyses (Gutierrez et al. 2009). For supercontigs, nucleotide
910 sequences assembled using HybPiper were gathered into fasta files by gene, gene names were
911 removed from fasta headers using a command-line regular expression, and sequences were
912 aligned in MAFFT with settings: --auto --adjustdirection --maxiterate 1000 (Katoh et al. 2002).
913 Reverse compliment tags (“_R_”) were removed from taxon names using a command-line
914 regular expression, and sequences were trimmed using TrimAl with previously listed settings
915 optimized for maximum likelihood analyses (Gutierrez et al. 2009). To minimize the effects of
916 missing data on phylogenetic analyses, accessions with < 50% of loci passing quality filtering
917 were removed, and genes that were recovered across < 70% of the total remaining samples were
918 also removed. Following quality filtering, we recovered 204 high quality loci (present in at least
919 70% of samples) and an average of 625,323 reads per sample (S1). Final assemblies are available
920 at Dryad ([http://dx.doi.org/10.5061/dryad.\[NNNN\]](http://dx.doi.org/10.5061/dryad.[NNNN])). Bioinformatics pipelines and analyses were

921 run at the high-performance computing cluster (“Treubia”) at the Chicago Botanic Garden unless
922 otherwise specified.

923

924 *Phylogenetic Reconstruction*

925 We conducted phylogenetic analyses using two strategies for each set of alignments.
926 Alignments were concatenated and analyzed using maximum likelihood (ML) in RAxML
927 (Stamatakis 2014; hereafter referred to as “concatenation”), whereas coalescent-based analyses
928 were conducted using ML gene trees in ASTRAL-II (Mirarab and Warnow 2015; Sayyari and
929 Mirarab 2016) and using unlinked SNPs in SVDquartets (Chifman and Kubatko 2014, 2015)
930 implemented in PAUP* beta version 4.0a168 (Swofford 2003; S3). In concatenation analyses,
931 after aligning each gene separately in MAFFT, genes were concatenated, partitioned, and
932 maximum likelihood trees were reconstructed in RAxML Version 8 (Stamatakis 2014) using the
933 GTRCAT model with 100 “rapid-boostrapping” psuedoreplicates and *Chylismia scapoidea* as
934 the outgroup, on the CIPRES Science Gateway computing cluster (Miller et al. 2010). For
935 coalescent analyses, individual gene trees were first estimated using RAxML Version 8
936 (Stamatakis 2014), with 100 “rapid-boostrapping” psuedo-replicates and settings: -p 12345 -x
937 12345 -N 100 -c 25 -f a -m GTRCAT -s, and *Chylismia scapoidea* as the outgroup. Gene trees
938 based on supercontigs were not partitioned by codon position. Coalescent-based analyses of
939 accessions were conducted in ASTRAL-II (Mirarab and Warnow 2015; Sayyari and Mirarab
940 2016) with default settings using the best RAxML gene trees and their associated bootstrap files
941 as input, and in SVD Quartets (Chifman and Kubatko 2014, 2015) with default settings using an
942 inframe aligned supermatrix with all 204 loci and supercontigs. ASTRAL-II and SVD Quartets
943 analysis was performed with 100 multi-locus bootstraps.

944 To assess concordance among gene trees and provide additional support complementary
945 to bootstrap values, we conducted two additional analyses. First, we assessed raw gene tree
946 concordance using Phyparts (Smith et al. 2015). Prior to running Phyparts, nodes with < 33%
947 support in the supercontig RAxML gene trees were collapsed using the sumtrees command in
948 Dendropy (Sukumaran 2010). These gene trees were then re-rooted using *Chylismia scapoidea*
949 as the outgroup and ASTRAL-II was rerun using these collapsed, re-rooted gene trees as the
950 input files. Pie charts showing gene tree discordance were generated and overlaid on the
951 resulting ASTRAL-II tree using the PhypartsPiecharts script
952 (<https://github.com/mossmatters/phyloscripts/tree/master/phypartspiecharts>). Phyparts piecharts
953 and gene tree concordance values were also added to Figure 1 by importing the two data files
954 produced by the Phypartspiecharts.py into R and manually matching them to key nodes on our
955 ASTRAL-II supercontig tree using *ggtree* version 1.14.6 (G Yu, DK Smith, H Zhu, Y Guan
956 2017). We also generated gene and site concordance factors for our ASTRAL-II tree constructed
957 using supercontigs in IQTree v1.7-beta16 (Minh et al. 2018). IQtree calculates the gene
958 concordance factor (gCF) and accounts for incomplete taxon coverage among gene trees and
959 therefore may provide a more accurate representation of agreement among gene trees than other
960 methods. In addition to gCF, IQTree calculates the site concordance factor (sCF), which is
961 defined as the percentage of decisive nucleotide sites supporting a specific node (Minh et al.
962 2018). We used the RAxML gene trees produced for the ASTRAL-II supercontig analysis, and
963 supercontig alignments themselves, as the inputs for IQtree. For computing sCF, we randomly
964 sampled 100 quartets around each internal node. Finally, we mapped gCF and sCF values to the
965 ASTRAL-II supercontig tree produced in our previous summary coalescent analysis (Fig. 2). All

966 phylogenetic trees, with the exception of the full Phyparts picharts tree, were visualized using the
967 R package *ggtree* version 1.14.6 (G Yu, DK Smith, H Zhu, Y Guan 2017).

968

969 *Ancestral State Reconstruction*

970 To infer ancestral conditions and the number of transitions in reproductive system, we
971 used the *phangorn* (Schliep 2011) package in R. First, a Coalescent-based species tree with
972 accessions grouped into taxa using a mapping file was estimated in ASTRAL-III (Zhang et al.
973 2018) with default settings using the best RAxML gene trees and their associated bootstrap files,
974 from the supercontig alignments, as input. Next we time calibrated the ASTRAL-III species tree
975 to 1 million years based on estimates of other taxa in the genus (Evans et al. 2009) using the
976 *makeChronosCalib* function in the *ape* (Paradis et al. 2004) package in R, and estimated an
977 ultrametric tree using the *chronos* function in *ape* (Paradis et al. 2004) with settings: $\lambda = 1$,
978 $\text{model} = \text{"relaxed"}$. Finally, we performed marginal reconstruction of ancestral character states
979 using the maximum likelihood method using the *optim.pml* and *ancestral.pml* functions in the
980 *phangorn* (Schliep 2011) package in R.

981

982 *Testing for Hybrid Origins with HyDe*

983 To test for putative hybrid origins of selected taxa, we used HyDe (Blischak et al. 2018)
984 to calculate D-Statistics (Green et al. 2010) for a set of hypotheses (S10). Briefly, HyDe
985 considers a four-taxon network of an outgroup and a triplet of ingroup populations to detect
986 hybridization from phylogenetic invariants that arise under the coalescent model with
987 hybridization. Introgression between P3 and either P1 or P2 influences the relative frequencies of
988 ABBA and BABA, and the D-statistic measures the imbalance between these frequencies. We
989 tested the triplets in (S10) and set *Chylismia scapoidea* as the outgroup. We considered
990 hypothesis tests significant at an overall $\alpha < 0.05$ level with estimates of γ between 0 and 1. Z-
991 scores greater than 3 are generally interpreted as strong evidence of introgression.

992

993 *Population-level Analysis*

994 To further characterize population-level processes or genetic structure within sect.
995 *Calylophus*, we extracted and filtered SNPs by mapping individual reads against reference
996 supercontigs (see <https://github.com/lindsawi/HybSeq-SNP-Extraction>). To account for
997 duplicates arising from PCR during HybSeq in SNP calling and filtering, first we selected the
998 sample with the highest target recovery rate and sequencing depth as a target reference sequence
999 (*Oenothera capillifolia berlandieri_bjc19*) and gathered supercontigs for this individual into a
1000 single target FASTA file. We then ran BWA (Li and Durbin 2009) to align sequences, Samtools
1001 'index' (Danecek et al. 2011) and GATK CreateSequenceDictionary (Poplin et al. 2017),
1002 respectively, on the resulting target FASTA file. Next we ran a custom script
1003 "variant_workflow.sh" using both read files from each *Calylophus* sample as input to create a
1004 vcf file for each sample. SNP's were called for each individual using GATK (Poplin et al. 2017)
1005 and the vcf file from each sample as input. The resulting vcf file created in the previous step was
1006 filtered to remove indels using GATK and the original target FASTA file as input, and then
1007 filtered again based on read mapping and quality with GATK VariantFiltration with settings: --
1008 filterExpression "QD < 5.0 || FS > 60.0 || MQ < 40.0 || MQRankSum < -12.5 || ReadPosRankSum
1009 < -8.0" (Poplin et al. 2017). Finally, we generated a reduced SNP file in FastStructure format
1010 using PLINK (Purcell et al. 2007) to remove SNPs that did pass filter using the command: plink -
1011 -vcf-filter --vcf Pachylophus.filtered.snps.vcf --const-fid --allow-extra-chr --geno --make-bed --

1012 recode structure. Finally we used Discriminant Analysis of Principal Components (Jombart et al.
1013 2010) as implemented in the R package *adegenet* (Jombart 2008) and the *snmf* function in the
1014 LEA package (Frichot and François 2015) in R (R Core Team, 2020).

1015

1016 *Morphological Measurements and Analysis*

1017 To assess taxon boundaries and patterns of morphological variation, we measured
1018 character states for the following key morphological structures that have been used historically to
1019 discriminate taxa in sect. *Calylophus* (Towner 1977): plant height, leaf length (distal), leaf width
1020 (distal), leaf length/width ratio (distal), leaf length (basal), leaf width (basal), leaf length/width
1021 (basal), sepal length, and sepal tip length. Measurements were made with digital calipers when
1022 possible, or with a standard metric ruler and dissecting scope, from voucher specimens of nearly
1023 all sampled populations of sect. *Calylophus* included in our molecular phylogenetic analyses.
1024 Measurements are provided in S11 and have been deposited at Dryad
1025 ([http://dx.doi.org/10.5061/dryad.\[NNNN\]](http://dx.doi.org/10.5061/dryad.[NNNN])). Morphological measurements were log transformed
1026 using the R base function ‘log’ (R Core Team 2018) prior to Principal Components Analysis
1027 (PCA), which was conducted in R using the *stats* package version 3.7 and the function
1028 ‘prcomp’ (R Core Team 2018). All ‘NA’ values were omitted from analysis. Plots of PCA results
1029 were visualized using the *ggplot2* package in R (Wickham 2016).

1030

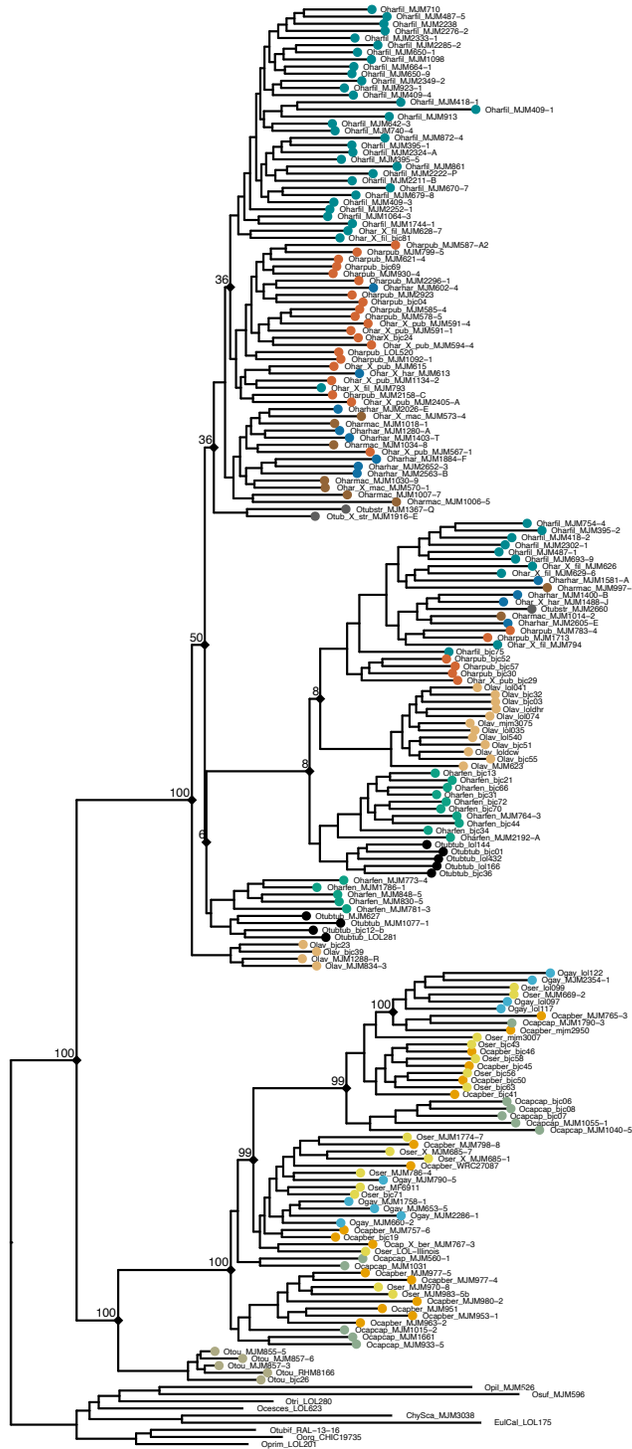
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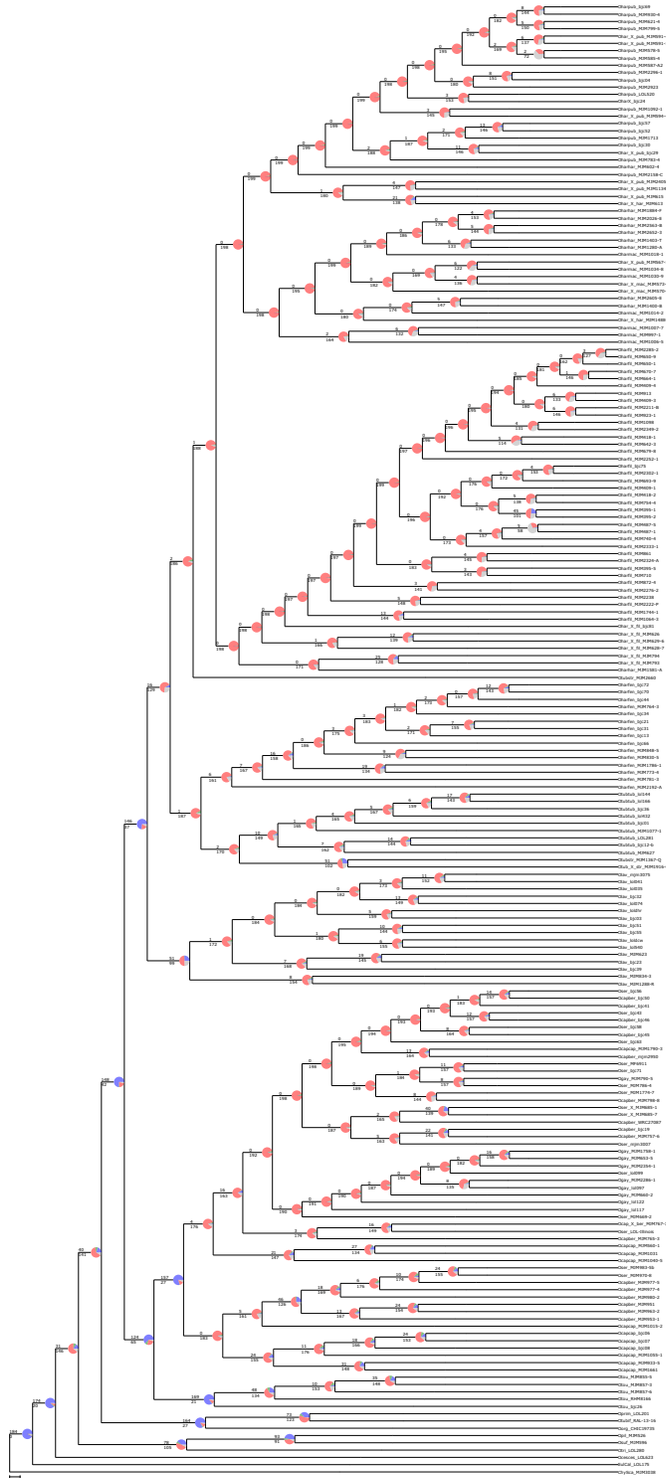
1113 S4.



1114
1115 Concatenation tree constructed using the exon-only dataset and 100 bootstraps. Bootstrap values
1116 indicated at relevant nodes.

1129

1130 S8.



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1134

Phyparts piecharts ASTRAL-II tree constructed using the supercontig dataset. Piechart colors correspond to: blue = concordant, green = top alternative bipartition, red = all other alternative bipartitions, black = uninformative for that node.

1135 S9.

1136 Summary of support values for current and proposed taxonomic treatments, by analysis. ‘e’
 1137 signifies trees based on exon-only data, “e+i” signifies trees based on supercontigs. ‘p’ indicates
 1138 paraphyletic, unsupported taxon treatment according to tree topology.

Taxon	Concat. e	Concat. e+i	ASTRAL-II e	ASTRAL-II e+i	SVDQ e+i	Phyparts	gCF iQtree e+i	sCF iQtree e+i
a. <i>Oenothera hartwegii</i> (Towner 1977)	p	p	p	p	p	p	p	p
b. <i>Oenothera hartwegii</i> subsp. <i>fendleri</i> (Towner 1977)	<50	p	81	99	p	1/ 188	1	38
c. <i>Oenothera hartwegii</i> subsp. <i>filifolia</i> (Towner 1977)	p	p	<50	<50	<50	1/ 190	0	36
d. <i>Oenothera hartwegii</i> subsp. <i>hartwegii</i> (Towner 1977)	p	p	p	p	p	p	p	p
e. <i>Oenothera hartwegii</i> subsp. <i>macartii</i> (Towner 1977)	p	p	p	p	p	p	p	
f. <i>Oenothera hartwegii</i> subsp. <i>hartwegii</i> + subsp. <i>macartii</i>	p	p	<50	<50	<50	0/ 200	0	33
g. <i>Oenothera hartwegii</i> subsp. <i>pubescens</i> (Towner 1977)	p	p	<50	<50	<50	0/ 200	0	36
h. <i>Oenothera lavandulifolia</i> (Towner 1977)	100	p	100	100	100	147/37	74	83
i. <i>Oenothera tubicula</i> (Towner 1977)	p	p	p	p	p	1/ 188	p	p
j. <i>Oenothera tubicula</i> subsp. <i>tubicula</i> (Towner 1977)	p	p	81	99	<50	2/ 171	1	38
k. <i>Oenothera tubicula</i> subsp. <i>strigulosa</i> (Towner 1977)	p	p	81	100	<50	p	0	37
l. <i>Oenothera fendleri</i> + <i>Oenothera tubicula</i> subsp. <i>tubicula</i>	p	p	100	100	<50	16/129	10	40
m. <i>Oenothera toumeyi</i> (Towner 1977)	100	100	100	100	100	125/65	62	56
n. <i>Oenothera toumeyi</i> + subsect. <i>Calylophus</i>	100	100	100	100	100	149/42	76	65
o. <i>Oenothera capillifolia</i> (Towner 1977)	p	p	p	p	p	p	p	p
p. <i>Oenothera capillifolia</i> subsp. <i>capillifolia</i> (Towner 1977)	p	p	<50	p	<77	p	p	p
q. <i>Oenothera capillifolia</i> subsp. <i>berlandieri</i> (Towner 1977)	p	p	p	p	p	p	p	p
r. <i>Oenothera capillifolia</i> subsp. <i>berlandieri</i> (New Taxon; South Texas coastal populations)	p	p	<50	100	100	157/27	79	86
s. <i>Oenothera serrulata</i> (Towner 1977)	p	p	p	p	p	p	p	p
t. <i>Oenothera ‘australis’</i> (New Taxon; South Texas <i>O. serrulata</i>)	p	100	p	p	p	10/175	p	p
u. <i>Oenothera gayleana</i> (Turner & Moore 2014)	p	p	p	p	p	p	p	p
v. <i>Oenothera gayleana</i> (Revised Taxon)	p	p	p	<50	<50	0/ 193	0	35

1139

1140 S10.

1141 List of admixture hypotheses tested, Zscore, P-value and Gamma results from HyDe analysis.

1142 Each row represents a triplet set that was tested consisting of a putative hybrid individual and

1143 two parent groups; Parent 1 and Parent 2 (see “HyDe Group” Appendix I for group

1144 membership).

1145

Putative hybrid individuals	Parent 1 Group	Parent 2 Group	Zscore	P	Gamma
Ohar_X_fil_MJM629.6	CoreOharfil	CoreOharpub	-99999.9	1	-0.049
Ohar_X_fil_MJM628	CoreOharfil	CoreOharpub	-99999.9	1	-0.057
Ohar_X_fil_MJM626	CoreOharfil	CoreOharpub	-99999.9	1	-0.001
Ohar_X_fil_MJM793	CoreOharfil	CoreOharpub	-99999.9	1	-0.057
Ohar_X_fil_MJM794	CoreOharfil	CoreOharpub	-99999.9	1	-0.034
Ohar_X_fil_BJC81	CoreOharfil	CoreOharpub	-99999.9	1	-0.042
Oharhar_MJM1581	CoreOharpub	Oharhar/harmac	-99999.9	1	1.706
Ohar_X_pub_BJC29	CoreOharpub	Oharhar/harmac	2.378	0.009**	0.338
Oharpub_BJC30	CoreOharpub	Oharhar/harmac	-99999.9	1	-0.215
Oharpub_MJM2158.C	CoreOharpub	Oharhar/harmac	1.113	0.133	0.229
Ohar_X_pub_MJM1134.2	CoreOharpub	Oharhar/harmac	0.043	0.483	0.015
Ohar_X_pub_MJM2405.A	CoreOharpub	Oharhar/harmac	0.882	0.189	0.158
Ohar_X_pub_MJM594.4	CoreOharpub	Oharhar/harmac	2.094	0.018*	0.332
Ohar_X_pub_MJM615	CoreOharpub	Oharhar/harmac	0.413	0.340	0.096
Oharmac_MJM997.1	CoreOharpub	Oharhar/harmac	1.563	0.059	0.224
Ohar_X_har_MJM613	CoreOharpub	Oharhar/harmac	-99999.9	1	-0.495
Otub_X_str_MJM1916.E	Otubtub	Oharhar/harmac	5.585	0.000***	0.947
Oharfen_MJM2192.A	CoreOharfen	CoreOharfil	0.305	0.380	0.001
Oharfen_MJM781.3	CoreOharfen	CoreOharpub	0.025	0.490	9.69E-05
Ocapcap_MJM1015.2	CoreOcapcap	S. TX Ocapber	0.467	0.320	0.003
Ocap_X_ber_MJM767.3	central OK Ocapber	Pecos River/Southern Plains	-99999.9	1	-0.003

• Indicates value meets significance threshold of P < 0.05
 ** Indicates value meets significance threshold of P < 0.001
 *** Indicates value meets significance threshold of P < 0.0001

1146

1147 S11.

1148 See excel table “S11 Morphometric Data”.

1149 S12.

1150 See excel table “S12 Pollen Counts”.

1151

1152 S13.

1153 ***Oenothera HybSeq CTAB Silica DNA extractions***
1154 ***b. cooper 11/10/2015***

1155 Adapted from JISA Protocol, J. Fant's Protocol and Sharma and Puohit (2012) for Silica-dried Leaves of *Oenothera*
1156 *sect. Calylophus*

1157
1158 **SAMPLE PREPARATION & GRINDING**

- 1159 1. Warm CTAB buffer to 65°C in water bath in the hood and pour liquid nitrogen into dipping canister (You
1160 will only need a few inches of liquid in the Dewar, make sure to pour it back into the large holding canister
1161 when you are done to prevent evaporation).
- 1162 2. Put a small volume of 1:1 sterilized sand and 3 metal/ceramic beads into centrifuge tubes, then put tubes
1163 in freezer (-20) until envelopes are ready.
- 1164 3. Keeping envelopes/leaf tissue on ice to prevent thawing; take envelopes and centrifuge tubes out of
1165 freezer and weigh/estimate small amount of dried tissue (e.g., 10-50 mg of leaves), put in labeled
1166 microfuge tubes. After each tube is filled, place tube in freezer.
 - 1167 a. Make sure leaf tissue is broken up into small pieces when adding to ensure full grinding.
 - 1168 b. Keeping tissue frozen prevents phenolic compounds from oxidizing, and enzymes from activating and
1169 breaking down DNA
 - 1170 c. Follow DNA free lab protocols using bleach to prevent contamination/cross contamination of samples.
- 1171 4. Mix 45 (= .3%) μ l 2-mercaptoethanol to 15 ml CTAB buffer in small beaker (enough for ~15
1172 samples...double for 30 samples, etc.)
 - 1173 a. Mercaptoethanol must be measured and added in the fume hood.
 - 1174 b. CTAB is a detergent that lyses (breaks down) the cell membranes. The PVP in the buffer helps bind the
1175 polysaccharides and might co-ppt
- 1176 5. Dip tubes in Liquid Nitrogen (or store in -80 for ~30 minutes) and **immediately** Grind tissue **thoroughly** in
1177 Fast Prep machine (thorough grinding may require 2-4 cycles, re-dip tubes in liquid nitrogen after 2 cycles
1178 to ensure no thawing occurs).
- 1179 6. **Immediately** after grinding add 600 μ l of CTAB/mercaptoethanol buffer to each tube
 - 1180 a. **BE CAREFUL.** It is important to prevent cross-contamination when opening tubes. **Powdered tissue can**
1181 **spread easily through the air or on your fingers.** Make sure open tubes are spaced apart from one another
1182 on the tray; open each tube away from other open tubes/caps; and place caps a safe distance from one
1183 another. Make sure leaf powder is not stuck to your gloves before touching consecutive tubes/caps.
 - 1184 b. Grind on FastPrep for one cycle

1185 **EXTRACTION AND ISOLATION OF DNA (once leaf tissue is well-ground)**

- 1186 1. Incubate tubes for 60 minutes at 60°C on Thermoblock or water bath
 - 1187 a. Shake tubes several times ~ every 10 minutes during incubation (for compacted samples, vortex on high
1188 setting to ensure mixing)
 - 1189 b. When incubation is complete, turn Thermoblock down to 37°C
- 1190 2. Add 600 μ L chloroform-isoamyl (24:1) to tube and vortex on High setting to ensure thorough mixing.
 - 1191 a. **BE CAREFUL.** It is important to prevent cross-contamination at this step when opening/handling tubes.
1192 Follow steps outlined previously.
 - 1193 b. Chlorophyll and other pigments get transferred into the Cl layer because pigments are non-polar and
1194 dissolve into the highly non-polar chloroform. The DNA remains in the aqueous layer.
- 1195 3. Spin for 10 min at 9,000 RPM.
 - 1196 a. After spinning, the upper layer of each tube should have a clear liquid: this is where your DNA is.
- 1197 4. Transfer top (clear) liquid to a new (labeled) 1.5 ml eppendorf tube. **Do not be Greedy**, leaving some liquid
1198 in tube is fine – the important thing is not to pipette **ANY** interphase material.
 - 1199 a. A yield of ~400-500 μ L of supernatant is common.
 - 1200 b. If you accidentally disturb the interphase, or believe that you have pipetted interphase, return supernatant to
1201 microfuge tube and re-centrifuge for 10 min at 9000 rpm. Repeat step 4.

1202 **DNA PRECIPITATION**

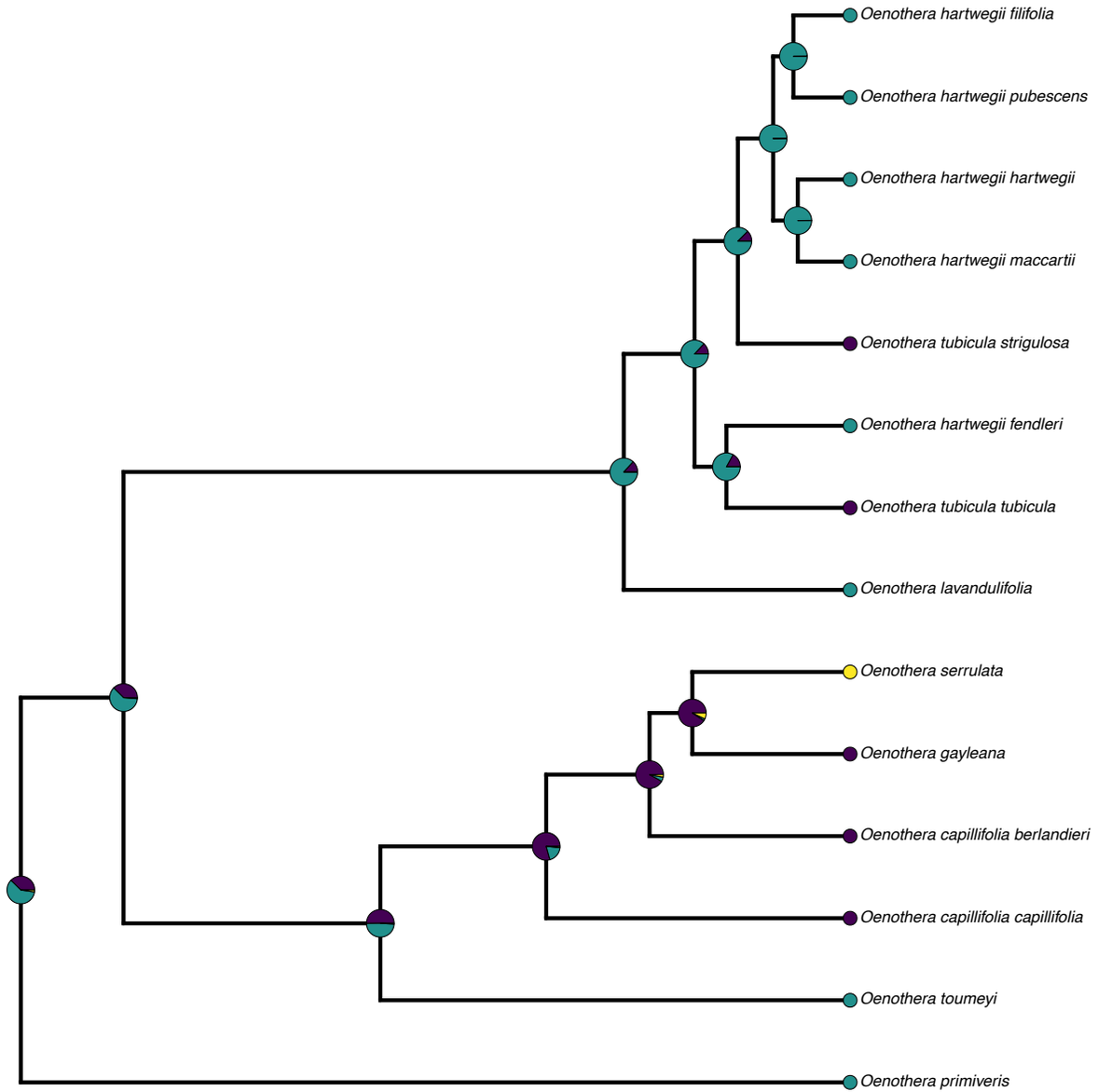
- 1203 1. Add 1/10 volume 3M sodium acetate and ½ volume 5M NaCl (i.e. 500ul of solution add 250 ul of NaCl, 50
1204 ul of Na-Acetate).
- 1205 2. Add 2/3rd volume of cold isopropanol (to previous volumes = 500 ul isop)
 - 1206 a. Isopropanol is heavy alcohol so allows DNA ppt with lower volume

- 1207 3. Mix & store in -20°C (freezer) for 1-2 hours
1208 **(NOTE: POTENTIAL STOPPING POINT FOR A DAY OR MORE...SAMPLES CAN STAY AT -20°C FOR DAYS)**
1209 4. Spin for 3 min at **10K-12K RPM.**
1210 5. Decant the supernatant & drain; make sure pellet stays at bottom of tube.
1211 a. Pellets are usually white but sometimes they can even be brown, this is not necessarily a problem.
1212 6. Wash pellet with 500ul of 70% EtOH & flick tubes to dislodge pellet from bottom, then vortex on high to
1213 clean.
1214 7. Spin for 3 min at 10K-12K RPM.
1215 8. Decant the supernatant making sure pellet stays at bottom of tube
1216 9. Leave tubes open and place tubes on Speedvac (vacuum on -2.5, heat = low, spin) for ~20 minutes or until
1217 all EtOH has evaporated.
1218 a. If pellets are not dry after 20 minutes, check frequently to prevent over drying which makes the pellet
1219 difficult to re-suspend.
1220 b. Alternatively, can air dry on thermoblock
1221 **(NOTE: POTENTIAL STOPPING POINT ...SAMPLES CAN STAY AT 20°C FOR DAYS – place in cupboard but cover with kimwipe)**
1222 **FINAL CLEANING & RESUSPENSION OF DNA**
1223 1. Add 500 ul of 0.5 mL High Salt TE Buffer and close lids
1224 2. Incubate tubes on Thermoblock at 37-50°C (45) until pellet dissolves (15-30 minutes);
1225 a. Vortex frequently if this is taking a long time
1226 **(NOTE: POTENTIAL STOPPING POINT FOR A DAY OR MORE...SAMPLES CAN STAY AT -20°C FOR DAYS)**
1227 3. Add 3 volumes of binding buffer (3 volumes for every 1 volume DNA pellet = ~100-150 ul BB), **let stand for**
1228 **20 minutes**
1229 4. Centrifuge at 550 g for 10 min
1230 5. transfer supernatant to clean 2ml eppi tube leaving any colored or gelatinous precipitate behind.
1231 6. Add 300 µl of silica suspension and mix for 30 minutes by regular and frequent gentle inversion
1232 7. Centrifuge at 550 g for 10 minutes, discard supernatant
1233 8. Re-suspend silica pellet in 1.5 mL of wash buffer 1
1234 9. Centrifuge at 3000 g for 15 seconds, decant supernatant
1235 10. Re-suspend silica pellet in 1.5 mL of wash buffer 2
1236 11. Centrifuge at 3000 g for 15 seconds, decant supernatant, dry pellet completely on speed vac
1237 12. suspend silica pellet in 300 ul TE buffer, incubate at 50° mixing regularly by vortex
1238 13. centrifuge at 11600 g for 1 min. then transfer supernatant into clean 1.5 mL eppi tube.
1239 14. Precipitate DNA by adding 50 ul of 3M sodium Acetate and 500 uL 100% Etoh
1240 15. Store at -20° C for at least 2 hours
1241 **(NOTE: POTENTIAL STOPPING POINT FOR A DAY OR MORE...SAMPLES CAN STAY AT -20°C FOR DAYS)**
1242 16. Centrifuge at 11600 g for 5 min, discard supernatant, dry pellet on speed vac
1243 17. Dissolve in 50 uL 1X TE buffer (it may not look like there is anything in the tube at this point, but there is
1244 lots of pure DNA in there! Add the TE buffer)
1245 18. Store in -20°C (freezer)
1246 19. In rare circumstances that an extraction still has coloration at this stage (yellow or brown usually) you can
1247 now put it through a Qiagen Qiaquick spin column or other proprietary cleanup column to further purify
1248 the DNA.
1249
1250 **REAGENTS (recipes below) and SUPPLIES**
1251 • CTAB buffer (in glass container in refrigerator)
1252 • Liquid Nitrogen
1253 • 2-mercaptoethanol (in fume hood with gloves)
1254 • 24:1Chloroform:isoamyl alcohol - glass container in fumehood use gloves and keep in hood -(Or 25:24:1
1255 phenol-chloroform:isoamyl)
1256 • 5M Nacl
1257 • 3M M sodium acetate
1258 • Isopropanol and 70% and 100% EtOH
1259 • High Salt TE buffer
1260 • Binding Buffer

- 1261 • Silica Matrix
1262 • Wash Buffer 1
1263 • Wash Buffer 2
1264 • 1X TE buffer
1265 • 1.5 ml eppi tubes
1266 • 2 ml eppi tubes
1267
- 1268 **CTAB BUFFER** [100 mM Tris-HCl (pH=8.0), 1.4 M NaCl, 20 mM EDTA, 2% (or 4%) CTAB
1269 (hexadecyltrimethylammonium bromide), 2% PVP-40 (polyvinylpyrrolidone, m.w. 40000), + 0.3% β-
1270 mercaptoethanol (add in fume hood in a beaker to a premeasured volume of CTAB buffer required for the number
1271 of extractions planned plus one).]
1272
- 1273 For 200 mL CTAB buffer 2% (4%):
1274 20 ml 1M Tris-Cl
1275 1.4 M NaCl (16.36 g NaCl)
1276 • The addition of NaCl at concentrations higher than 0.5 M, along with CTAB, is known to remove
1277 polysaccharides during DNA extraction
1278 8 ml 0.5 M EDTA
1279 4 g CTAB (8 g for 4%)
1280 4 g PVP-40
1281 ~120 ml dH₂O, then fill to 200 mL
1282
- 1283 **TE buffer:** [10 mM Tris-HCl (pH=8.0), 1 mM EDTA.]
- 1284 10 mL 1 M Tris-HCl
1285 2 ml 0.5 M EDTA
1286 982 ml dH₂O
1287
- 1288 **High Salt TE Buffer:** [10 mM Tris-HCl (pH=8.0), 1 mM EDTA, 1M NaCl.]
1289 10 ml 1 M Tris-HCl
1290 2 ml 0.5 M EDTA
1291 200 ml 5M NaCl
1292 788 ml dH₂O
1293
- 1294 **Binding Buffer:** [50mM Tris (pH7.5), 6M NaClO₄, 1mM EDTA]
1295 For 200 ml:
1296 10mL Tris
1297 168.552 g NaClO₄
1298 400 ul EDTA
1299 top off to 200 mL with dH₂O
1300
- 1301 **Wash Buffer 1:** [3 volumes binding buffer, 1 volume water]
1302 For 60 samples:
1303 90 ml Binding Buffer
1304 30 ml dH₂O
1305
- 1306 **Wash Buffer 2:** [1 volume 40 mM Tris (pH 8.0), 4 mM EDTA, 0.8 M NaCl, 1 Volume ethanol]
1307 For 60 samples:
1308 1.8 ml 1M tris
1309 360 ul .5M EDTA
1310 7.2 ml 5M NaCl

- 1311 35.64 ml dh20
1312 45 ml 100% etoh
1313
1314 **24:1 Chloroform: Isoamyl**
1315 240ml Chloroform
1316 10ml Isoamyl alcohol
1317
1318 25:24:1 Chloroform: Isoamyl Saturated with 10 mM Tris, pH 8.0, 1 mM EDTA. – Purchase to ensure right pH
1319 250ml Phenol
1320 240ml Chloroform
1321 10ml Isoamyl alcohol
1322
1323 **Silica suspension (use lab grade silicon dioxide 99%)**
1324 1. Suspend silicon dioxide powder in ~20 volumes of water (50 ml : 1L)
1325 a. use magnetic stirring rod to bring silica into uniform suspension
1326 2. Decant suspension into clean beaker leaving behind heavy sediment at bottom
1327 (discard this)
1328 3. Allow suspension to settle by gravity for 2-3 hours
1329 4. Decant and discard supernatant with fine material keeping silicon dioxide that
1330 has settled
1331 5. Re-suspend silicon dioxide in dH2O, repeating step until fine particles removed
1332 (2-3X total is good)
1333 a. use magnetic stirring rod to bring silica into uniform suspension
1334 6. Re-suspend silicon dioxide in dH2O and transfer to 15 ml falcon tubes
1335 7. Collect silicon dioxide by low speed centrifugation (up to 3200 rpm for 3 min)
1336 8. Add ~15 ml of 10% bleach solution to each aliquot and mix frequently by
1337 inversion/vortexing for 15-30 minutes
1338 9. Re-suspend diatomite in dH2O by vortexing, then centrifuge at low speed for 3
1339 minutes to pellet
1340 10. Discard supernatant
1341 11. Repeat step 9 and 10 two more times
1342 12. Re-suspend in dH2O and autoclave
1343 13. Re-suspend silica dioxide pellet in 1 volume PCR grade DNA/RNA free H2O
1344 14. Aliquot into 1.5 ml eppi tubes and store at 4° C
1345 15. Vortex thoroughly to bring silica into uniform suspension before using
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1361 S14.



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1363 Ancestral State Reconstruction of reproductive system in sect. *Calylophus* using supercontigs
1364 and accessions grouped into taxa. Pie-charts on nodes represent likelihood of ancestral
1365 reproductive system at each node (teal = hawkmoth pollination, purple = bee pollination, yellow
1366 = PTH).
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