Re-evaluating homoploid reticulate evolution in the annual sunflowers.

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15 Abstract:

16 Sunflowers of the genus *Helianthus* are models for hybridization research and contain three

- 17 of the best studied examples of homoploid hybrid speciation. To understand the broader picture
- 18 of hybridization within the annual sunflowers, we used whole genome resequencing to conduct
- 19 a phylogenomic analysis and test for gene flow between lineages. We find that all annual
- 20 sunflower species tested have evidence of admixture, suggesting hybridization was common

21 during the radiation of the genus. Support for the major species tree decreases with 22 recombination rate, consistent with hybridization and introgression contributing to discordant 23 topologies. Admixture graphs found hybridization to be associated with the origins of the three 24 putative hybrid species (H. anomalus, H. deserticola, and H. paradoxus). However, the 25 hybridization events are more ancient than suggested by previous work. Furthermore, H. 26 anomalus and H. deserticola appear to have arisen from a single hybridization event involving 27 an unexpected donor, rather than through multiple independent events as previously proposed. 28 Using a broader data set that covers the whole *Helianthus* genus, including perennial species, 29 we find that signals of introgression span the genus and beyond, suggesting highly divergent 30 introgression and/or the sorting of ancient haplotypes. Thus, Helianthus can be viewed as a 31 syngameon in which largely reproductively isolated species are linked together by occasional or 32 frequent gene flow.

33 Introduction

34 The evolutionary importance of hybridization and introgression has been explored since 35 Darwin's time (Darwin 1859). While initially zoologists thought that hybridization was relatively 36 rare and unimportant (Mayr 1963), botanists have long emphasized the role of hybridization and 37 introgression in both adaptation and speciation (Anderson 1949; Heiser 1949; Anderson and 38 Stebbins 1954). Genomics eventually proved botanists correct in this regard; evidence of past 39 hybridization is common in both animals and plants (Figueiró et al. 2017; Fontaine et al. 2015; 40 Mallet et al. 2016; Calfee et al. 2021; Dagilis et al. 2021). In several cases, alleles responsible 41 for ecologically important traits have been found to be acquired through introgression (Suarez-42 Gonzalez et al. 2018; Jones et al. 2018; Oziolor et al. 2019), supporting the notion that 43 introgression not only occurs, but can be evolutionarily important.

44 Hybridization can also result in the formation of new species through homoploid hybrid
45 speciation, in which an admixed lineage acquires reproductive isolation from its progenitors

46 through a combination of alleles or chromosomal rearrangements from both parents (Grant 47 1958; Schumer et al. 2014; Stebbins 1957). One way this can occur is if two parental species 48 differ at both habitat and mate choice. A hybrid population between these two may have 49 alternate parental alleles for these two traits, and therefore be separated from one parental 50 species by habitat and the other by mate choice (Wang et al., 2021). Hybrid speciation can also 51 occur when hybridization results in the ability to colonize a new ecological niche, reject parental 52 species as potential mates or have unique combinations of chromosomal rearrangements 53 (Gross and Rieseberg 2005: Melo et al. 2009: Rieseberg et al. 1995). Schumer et al. (2014) 54 argued that three criteria must be satisfied to demonstrate homoploid hybrid speciation: (1) 55 reproductive isolation from parental species, (2) prior admixture, and (3) evidence that 56 reproductive barriers arose via hybridization. Evaluation of the natural hybridization literature 57 indicates that while admixed lineages are relatively common (criterion 2), few case studies 58 satisfy criteria 1 and 3 (Schumer et al. 2014). Three examples of hybrid species thought to 59 address all three criteria are within the sunflower genus, Helianthus.

60 The widespread species Helianthus annuus and H. petiolaris are thought to be the parents 61 of three independent homoploid hybrid species, H. anomalus, H. deserticola and H. paradoxus. 62 These were initially identified as hybrid species through early genetic studies that found they 63 had a mixture of chloroplast DNA, allozyme and ribosomal DNA markers from both parental 64 species (Rieseberg et al. 1990a: Rieseberg 1991). The hybrid species have strong ecological 65 separation based on habitat choice; Helianthus anomalus occurs on sand dunes, H. deserticola 66 on sandy soil of the desert floor and *H. paradoxus* in brackish salt marshes (Heiser et al. 1969). 67 Comparative QTL mapping of hybrids and parental species suggested that this ecological 68 separation was achieved by combinations of parental alleles that allowed for extreme 69 (transgressive) phenotypes (Rieseberg et al. 2003). Subsequent experiments found that such 70 transgressive phenotypes were favored when segregating hybrids of *H. annuus-H. petiolaris*

were transplanted into the natural habitats of the three ancient hybrid species, with some
synthetic hybrids rivaling the ancient hybrids in fitness (Lexer et al. 2003; Gross et al. 2004;
Ludwig et al. 2004).

74 In addition to ecological isolation, the three hybrid species are partially reproductively 75 isolated by hybrid pollen sterility caused mainly by chromosomal rearrangements (Lai et al. 76 2005). This was hypothesized to be due to the hybrid species having a combination of parental 77 chromosomal rearrangements as well as unique changes (Rieseberg et al. 1995; Lai et al., 78 2005). Synthetic hybrids of *H. annuus-H. petiolaris* not only recovered fertility in a small number 79 of generations, but also were more compatible with the putative hybrid species than were the 80 parental species (Rieseberg et al. 1996; Rieseberg 2000, 2006). Microsatellite, chloroplast, and 81 karyotypic analyses suggested all hybrid species originated between 63,000 and 208,000 82 generations before present and a single (H. paradoxus) or multiple (H. anomalus and H. 83 deserticola) origins depending on the species (Schwarzbach and Rieseberg 2002; Welch and 84 Rieseberg 2002: Gross et al. 2003). The strong link between admixture and reproductive 85 isolation, in terms of ecology and karyotype, comparisons with experimental hybrid lineages, 86 and partial replication across three species have made *Helianthus* a model for understanding 87 homoploid hybrid speciation.

88 Although the *Helianthus* hybrid species were probed with cutting-edge molecular markers 89 when initially identified, they have been relatively unexamined in the genomics age. Studies 90 using Genotyping-By-Sequencing grouped H. anomalus and H. deserticola together in a 91 phylogenetic tree but did not probe further (Baute et al. 2016). Moody and Rieseberg (2012) 92 sequenced ten nuclear genes for multiple individuals of most annual sunflowers, including the 93 hybrid species. They found extensive discordance between individual gene trees and notably 94 that H. anomalus and H. deserticola grouped together and that H. paradoxus grouped with H. 95 annuus, although the overall species tree did not agree with previous expectations. The most

96 recent phylogeny of *Helianthus* did not include any hybrid species (Stephens et al., 2015). In 97 each of these cases, modern approaches for identifying gene flow and hybrid ancestry were not 98 performed. This is important because sunflower species are both young — Helianthus crown 99 age is 3.6 mya — and generally have high effective population size (Strasburg and Rieseberg 100 2008). This means that incomplete lineage sorting (ILS) is likely high and discordant topologies 101 that could be produced by hybridization may also be present without any interspecies gene flow. 102 In addition, signals of hybridization can sometimes derive from unexpected sources (Owens et 103 al. 2021).

104 Beyond the three well-characterized *Helianthus* hybrid species, numerous other cases of 105 hybridization and introgression have been reported in the genus (Heiser et al. 1969; Rieseberg 106 et al. 1991). leading Rieseberg (1991) to conclude that evolution in the group "must be 107 considered reticulate, rather than exclusively dichotomous and branching." While many of these 108 early cases - which were based on relatively limited evidence - have been subsequently 109 confirmed with genomic data (Kane et al. 2009; Owens et al. 2016; Mondon et al. 2018; Lee-110 Yaw et al. 2019; Zhang et al. 2019), others have not (Owens et al. 2021), and several remain to 111 be tested. This highlights that the combination of modern genomic datasets and modern 112 phylogenetic techniques allow for hypotheses about hybridization to be rigorously tested 113 (Hibbins and Hahn 2022).

Here we use modern phylogenomic techniques to assess the extent of admixture during the evolution of the genus. To do this, we first use whole genome sequencing data for most annual sunflowers to infer the genome-wide phylogeny of this group. We then apply tests of introgression and gene flow across the annual clade, including the three hybrid species, *H. anomalus*, *H. deserticola* and *H. paradoxus*. This allows us to revisit criterion 2 of homoploid hybrid speciation for these three species, and to gain greater insight regarding their ancestry and their relationship to the broader patterns of admixture within the annual sunflower clade.

Lastly, using a previously published phylogenomic data (Stephens et al., 2015), we extend ouranalyses across the entire genus.

123 **Results:**

124 Data filtering

We generated whole genome sequence data for two *Helianthus paradoxus* samples. We called variants against the *H. annuus* HA412-HOv2 reference genome for these samples as well as previously sequenced data for two samples of eight other annual species, or subspecies, and four outgroups (supplementary table 1; Todesco et al. 2020). We retained 5,877,513 variable and 28,274,146 invariant sites after filtering for quality, mappability and call rate (supplementary figure 1). We divided the genome into a total of 11,797 10 kbp non-overlapping windows and retained 3,267 that had \geq 2,000 called bases.

132 Species and gene trees

The species topology for non-hybrid taxa from ASTRAL (fig. 1A,B) was mostly consistent
with previous phylogenies that used fewer markers (Stephens et al. 2015; Baute et al. 2016;
Zhang et al. 2019). Quadripartition support at nodes ranged from 38 to 84%, suggesting
substantial variation in topology at gene trees. The maximum likelihood concatenated tree also
found the same topology as ASTRAL and monophyly for all samples of the same species (fig.
18).

As outlined in the introduction, the three putative homoploid hybrid taxa, *H. paradoxus*, *H. deserticola* and *H. anomalus* are thought to have originated from hybridization between *H. annuus* and *H. petiolaris* (Rieseberg 1991). Since ASTRAL does not account for hybridization, we would expect the potential hybrid species to group near to the dominant parental species.
We found that *H. paradoxus* grouped with *H. annuus* and its sister species *H. argophyllus*, whereas *H. anomalus* and *H. deserticola* grouped with *H. niveus canescens* (fig. 1B).

145 Quadripartition support for these groupings is low (each 39%), as expected for hybrid lineages,

146 although this was not unique to branches involving the previously identified hybrid species.

We further examined variation in gene tree topology by calculating the gene concordance factor for each branch on our concatenated tree using gene trees. We found relatively low support at branches separating species compared to branches separating samples within species (fig. 1C). When including all samples, each tree was unique. When we subsampled tips down to a single individual per species we found higher agreement; the most common tree was found 6 times out of 3,242 trees.

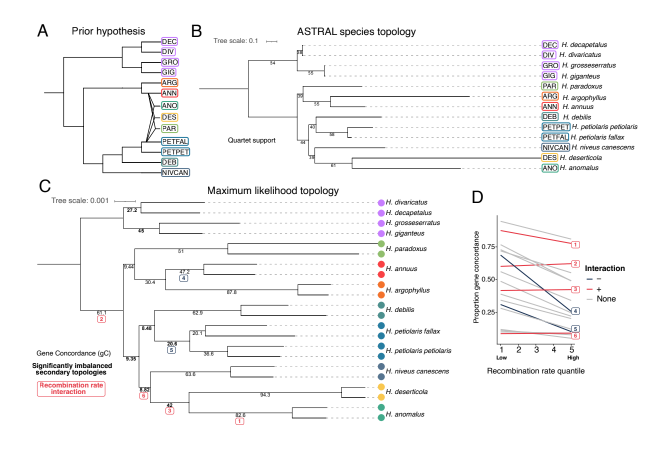
153 Gene tree discordance

To explore why gene concordance values were low, we compared the proportion of trees matching the two alternate topologies for every quadripartition using IQ-TREE2's gene concordance measure (Minh et al. 2020). Under purely incomplete lineage sorting, we expect that both alternate topologies should be found at equal proportions, while introgression will lead to imbalances. We found that at five nodes within the annual species, there was significant (Xsquared, p<0.05) imbalance in alternate topologies (fig. 1C).

160 Selection against introgressed ancestry can lead to reduced introgression in regions of low 161 recombination (Schumer et al. 2018; Martin et al. 2019). We explored this by using a binomial 162 regression to quantify the relationship between support for the species topology and the 163 recombination rate (as identified from the H. annuus genetic map) at each node with > 10% 164 support for alternate topologies. In other words, are genomic regions with lower recombination 165 more likely to show the species topology? We found a significant negative effect of recombination rate ($\beta_1 = -0.30$, p = e⁻¹⁴) on support for the species topology (fig. 1C,D). 166 167 Interestingly, we find a significant interaction between node and recombination rate at six nodes. 168 At four nodes, most of which include the *H. anomalus/H. deserticola* clade, the relationship is

less negative, or is positive (fig. 1D). At the node separating the *H. annuus* samples, therelationship with recombination is most strongly negative.

Helianthus niveus canescens was previously considered a variety of *H. petiolaris*, with which it intergrades for part of its range (Heiser et al. 1969). Our species topology places *H. niveus canescens* with *H. anomalus* and *H. deserticola* instead of *H. petiolaris*, but with relatively low support and overall less support at low recombination regions (fig. 1D). We counted the number of trees where *H. niveus canescens* grouped with *H. petiolaris petiolaris* and *H. petiolaris fallax* to the exclusion of *H. anomalus* and *H. deserticola*, as well as the reverse (supplementary figure 2). We found more support for *H. niveus canescens* together with *H. petiolaris*.



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Figure 1: Phylogenetics of annual sunflowers. A) The prior hypothesis of annual species
relationships (Rieseberg 1991; Stephens et al. 2015). B) Phylogenetic tree from combined 10

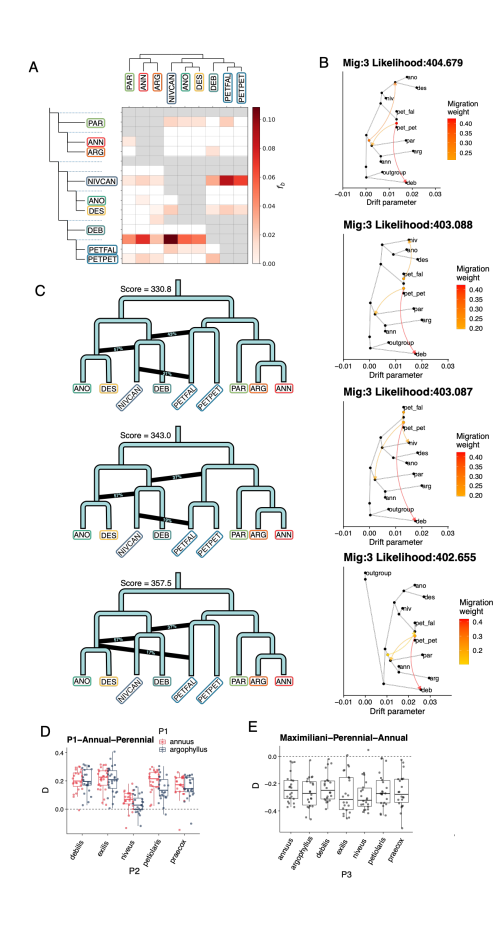
kbp gene trees used as input for ASTRAL species topology. Quartet support plotted on
branches. C) Concatenated maximum likelihood tree using IQ-TREE2. Gene concordance
plotted on nodes. Bolded values have significantly imbalanced secondary topologies. D) The
relationship between gene concordance proportion and recombination quantile for each node
with <90% overall gene concordance. Significant interactions with node ID are highlighted here
and on panel C.

187 Admixture in the annual sunflowers

188 To assess the extent of hybridization during the evolution of the annual sunflowers, we used 189 comprehensive approaches to test for admixture across all species. The f_{branch} analysis, which 190 uses the f4-ratio test to identify introgression within a phylogenetic framework, found many 191 significant signals (fig. 2A, supplementary table 2). The two strongest signals show that H. 192 petiolaris is closer to all other species, when compared to H. debilis, and H. niveus canescens is 193 closer to H. petiolaris and H. debilis, than H. anomalus and H. deserticola are. The fbranch 194 patterns are dependent on the underlying phylogeny, so to explore this effect we repeated fbranch 195 using the top nine most common species tree topologies (supplementary fig. 3). While no 196 topology completely removes significant f_{branch} signal, this shows that introgression signals from 197 fbranch can come from either close phylogenetic position or introgression. For example, in three of 198 the top nine trees, *H. paradoxus* is placed within the *H. petiolaris* clade, but its f_{branch} signal 199 shows it is much more similar to H. annuus and H. argophyllus than others in the H. petiolaris 200 clade (supplementary fig. 3). The three hybrid species do not stand out in terms of the extent of 201 admixture.

A different picture comes from the TreeMix analysis, which uses the covariance of population allele frequency to estimate a phylogeny and migration edges. We found that three was the optimal number of migration edges and explained 99.9% of variance. Above four edges, there was diminishing returns in likelihood gain and variance explained (supplementary 206 figure 4). Admixture was found to be associated with the origins of the three putative hybrid 207 species in the most likely TreeMix graphs at both three and four edges (supplementary figure 5). 208 The origin of *H. paradoxus* appears to involve ancient hybridization between the ancestor of the 209 H. annuus clade and H. petiolaris, although this admixture is inherited by the other sampled 210 members of the *H. annuus* clade. Likewise, there is an admixture event between a basal lineage 211 of the *H. annuus* clade and the ancestor of *H. deserticola* and *H. anomalus*. The third signal in 212 common between the top three admixture graphs was between H. debilis and H. petiolaris. 213 However, this likely is an artifact of misplacing *H*, debilis as basal in the tree and may instead 214 suggest that it has ancestry from a lineage at the base of the annual sunflower clade. When we 215 look other graphs that are close in likelihood to the top graphs, we frequently find admixture 216 between H. niveus canescens and H. petiolaris fallax, although this results in the loss of the 217 admixture event associated with *H. anomalus* and *H. deserticola*.

218 We also used ADMIXTOOLS2, which builds an admixture graph that explains f-statistics 219 between populations. Admixture scenarios seen using ADMIXTOOLS2 have both similarities 220 and differences with the TreeMix graphs. Out-of-sample scores decreased with increasing 221 admixture edges from one to two, and from two to three, but not from three to four, suggesting 222 that three admixture edges explain much of the f_2 incongruence (supplementary figure 6). While 223 bootstrapping was unable to distinguish between the top three graphs with three admixture 224 nodes (p > 0.1), they each present a similar pattern that distinguishes them from the species 225 tree (fig. 2C). The strongest signal is associated with origin of two of the putative hybrid species 226 (H. anomalus and H. deserticola), but from admixture with the ancestor of H. niveus canescens 227 and an unsampled or extinct basal lineage. Helianthus petiolaris is placed at the base of the 228 annuus clade in the ADMIXTOOLS2 graph, implying that it shares ancestry with members of the 229 annuus clade, including H. paradoxus. In addition, recent admixture is seen between H. niveus 230 canescens and H. petiolaris fallax similar to that observed in the TreeMix analysis.



232 Figure 2: Introgression across sunflowers. A) F_{branch} statistic for annual species using the 233 ASTRAL species topology. B) The four best supported TreeMix graph with three admixture 234 edges. Higher likelihood score is better. C) The best supported admixture graph from 235 ADMIXTOOLS2 with three admixture edges. Branch lengths are not to scale. Lower score is 236 better. D) ABBA-BABA with the groupings [H. annuus OR H. argophyllus], other annual species, 237 perennial species, *Phoebanthus*. Positive values indicate more ABBA counts. E) ABBA-BABA 238 with the groupings *H. maximiliani*, other perennials, annual species, *Phoebanthus*.

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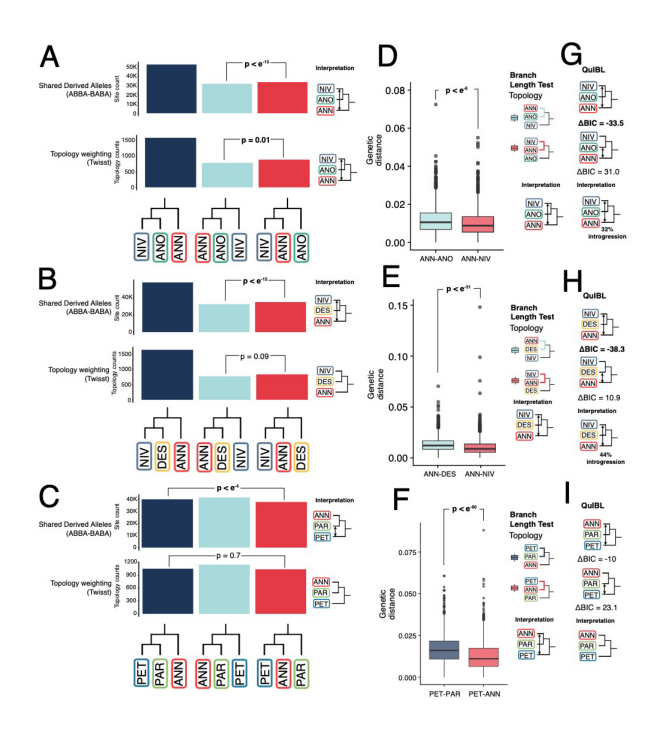
Triplet tests of hybrid ancestry

240 While the admixture graphs (above) detected signals of hybridization associated with the 241 origins of the three hybrid species, there were differences relative to previous hypotheses 242 (Rieseberg 1991). Most notably, the origin of *H. anomalus* and *H. deserticola* appears to involve 243 hybridization with the ancestor of *H. niveus canescens* rather *H. petiolaris fallax*. Also, there was 244 uncertainty as to whether the signal of admixture associated with the origin of *H. paradoxus* was 245 independent from other members of the annuus clade. Therefore, we explicitly tested 246 hypotheses of hybrid ancestry for H. anomalus, H. deserticola and H. paradoxus against their 247 putative parents. In the case of *H. paradoxus*, the testing is relative to *H. annuus* and *H.* 248 petiolaris fallax, whereas for H, anomalus and H, deserticola, the putative parents are H, annuus 249 and *H. niveus canescens*. For each test, we expect to see that the hybrid species is most 250 closely related to the major parental species, and that tests of introgression show that the 251 dominant signal of introgression involves the hybrid species and its minor parental species, 252 rather than the parental species. We used four different approaches. Patterson's D compares 253 the amounts of shared derived alleles in a trio (Green et al. 2010; Malinsky et al. 2021). In this 254 case, we are using Twisst to compare the counts of different gene tree topologies (Martin and 255 Van Belleghem 2017). The branch length test compares the branch lengths in alternate 256 topology gene trees. Lastly, QuIBL models ILS and introgression based on branch lengths. We

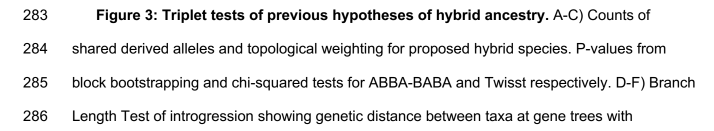
note that the results of triplet tests should be interpreted with caution because of the long history
of introgression between the putative parental clades (Yatabe et al. 2007; Strasburg and
Rieseberg 2008; Kane et al. 2009). For the topology-based tests (Patterson's D and Twisst),
this means that the extent of introgression from the minor parent must be significantly greater
than that between the parental lineages. For the distance-based tests (branch length and
QuIBL), both the extent and the timing of introgression becomes critical, as the distance-based
tests have little power for detecting ancient admixture.

264 For *H. paradoxus*, we consistently find *H. annuus* is the closest species, but inconsistent 265 evidence of admixture with H. petiolaris fallax (fig. 3). Patterson's D finds significant H. 266 paradoxus – H. petiolaris fallax admixture. The results from Twisst are in the same direction, but 267 are not significant. In contrast, the branch length test supports greater H. annuus – H. petiolaris 268 fallax introgression and QuIBL supports neither (fig. 3). We find qualitatively similar results when 269 using *H. petiolaris petiolaris* as a potential parental species. The branch length test is more 270 sensitive to recent introgression, such as that ongoing between H. annuus and H. petiolaris 271 (Yatabe et al. 2007), whereas D has greater power to detect ancient hybridization, perhaps 272 accounting for the apparent discrepancy between the different tests. At the genomic window 273 level, there is considerable variation in D with values ranging from -1 to 1 (supplementary figure 274 7).

For *H. anomalus* and *H. deserticola*, we see a consistent pattern for all four tests; they are more closely related to *H. niveus canescens* and there is more admixture between the potential parental species, than between the hybrid species and *H. annuus* (fig. 3). This could indicate that the hybrid species are ancient, and that the minor parent is at the base of the *H. annuus* clade (as suggested by TreeMix), and therefore has a relatively small contribution from contemporary *H. annuus*, or that the other parent of *H. anomalus* and *H. deserticola* is a now extinct basal lineage, as suggested in the ADMIXTOOLS2 graph.



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secondary topologies. G-I) QuIBL delta BIC values and interpretation. Lower values indicatemore support for introgression.

289 Admixture across the sunflower genus

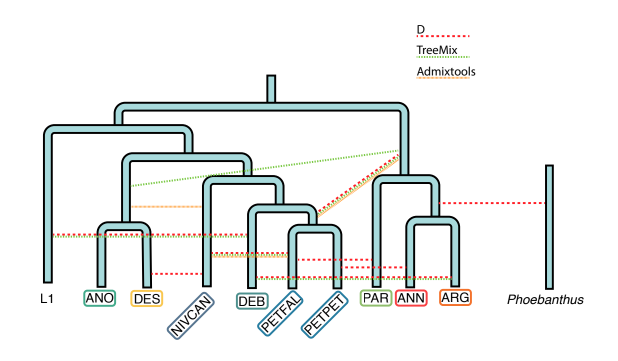
290 To assess introgression across the rest of the genus, as well as to confirm our findings within the annual clade, we expanded our analysis of gene flow using a previously published 291 292 sequence capture dataset spanning the genus. With this data, we used Dsuite to calculate fbranch 293 and D for all trios (supplementary figure 8, supplementary table 3). We found evidence of 294 introgression between perennial species, including H. verticilliatus – H. giganteus, H. divaricatus 295 - H. arizonensis and H. atrorubens - H. mollis. This dataset also affirms the signal of 296 introgression between *H. petiolaris* and both *H. niveus* and *H. annuus*. However, no 297 introgression was found involving *H. debilis*. Surprisingly, we also find introgression-like signals 298 between annual and perennial species. Helianthus maximilliani shares more derived alleles with 299 annual species than other perennials (fig. 2E). Strangely, both H. annuus and H. argophyllus 300 share fewer derived alleles with perennials than other annual species, which is consistent with 301 widespread annual-perennial introgression excluding H. annuus/H. argophyllus, or introgression 302 between the *H. annuus/H. argophyllus* clade and the outgroup *Phoebanthus* (fig. 2D). 303

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Figure 4: The proposed annual sunflower phylogeny with introgression events
 identified in the present study. L1 represents an unknown lineage seen through admixture
 analyses. Dotted lines indicate signatures of admixture seen in different analyses.

312 **Discussion**:

313 For three-fourths of a century, Helianthus sunflowers have served as a model for the study 314 of hybridization and its evolutionary role (Heiser 1947; Rieseberg 1991; Todesco et al. 2020). 315 Early studies documented hybrid swarms between different sunflower species (Heiser 1947; 316 Heiser 1949; Heiser 1951), demonstrated that chromosomal rearrangements contributed to 317 hybrid sterility (Chandler et al. 1986; Lai et al. 2005), and speculated on possible ecotypes and 318 species that might be the products of hybridization (Heiser 1949; Heiser 1958; Rieseberg 1991). 319 Later molecular phylogenetic work and early low-resolution genomic studies supported some of 320 these hypotheses (Rieseberg et al. 1990a; Yatabe et al. 2005), but not others (Rieseberg et al. 321 1988), and also suggested new hypotheses, including the putative origins of three homoploid 322 hybrid species (Rieseberg 1991). Here, we have taken advantage of new high resolution

323 genomic data and computational methods to further explore the role of hybridization in 324 sunflower evolution. We pay particular attention to the three homoploid hybrid species, but we 325 also examine other well-studied cases of hybridization in the *Helianthus* (and potential new 326 cases) with the goal of reconciling the new data and analyses with previous work. Lastly, we 327 explore the effects of variation in recombination rate on tree topology and introgression.

328 Hybridization in sunflowers

329 The whole genome phylogeny for sunflower is consistent with that found by previous multi-330 locus studies (Stephens et al. 2015; Baute et al. 2016; Zhang et al. 2019). Leveraging this 331 phylogeny, we employed several different methods to search for possible cases of admixture 332 during the evolution of the genus. Remarkably, all species included in our study appear to be 333 admixed, although the level of admixture depends in part on the particular analysis performed 334 (fig. 2,4). Overall, the strongest signal of hybridization occurs between two subspecies of H. 335 petiolaris and H. annuus. Helianthus petiolaris is sympatric with H. annuus and contemporary 336 hybridization is well-documented (Heiser 1947; Rieseberg et al. 1998). Previous studies have 337 documented "rampant" introgression between the species (Yatabe et al. 2005) and have 338 suggested that interspecific gene flow was common in the past as well (Strasburg and 339 Rieseberg 2008).

340 Another strong signal of admixture is found between *H. petiolaris* and *H. niveus canescens*, 341 confirming a previous report (Zhang et al. 2019). Helianthus petiolaris intergrades with H. niveus 342 canescens phenotypically, and Heiser et al. (1969) suggested that it might be due to 343 hybridization. Although both types of phylogenetic analysis suggested H. niveus canescens 344 grouped with H. anomalus and H. deserticola rather than H. petiolaris, there were more gene 345 trees that placed *H. niveus canescens* with *H. petiolaris*. Considering the weight of evidence, we 346 support that H. niveus canescens was involved with admixture of the H. anomalus/deserticola 347 ancestor but was originally a member of the *H. petiolaris* clade. The beach sunflower, *H. debilis*,

is sister to *H. petiolaris* in our tree, but f_{branch} shows that it is less similar to all other annual
sunflower species (fig. 2A). This implies that it has ancestry from a lineage at the base of the
annual sunflowers, as also suggested by TreeMix. Lastly, both the TreeMix and ADMIXTOOLS2
graphs suggest a shared hybrid origin for *H. anomalus* and *H. deserticola*, which we will discuss
in more length below.

353 We used an alternate dataset generated by Stephens et al. (2015) to assess introgression 354 across the entire Helianthus genus. We found evidence of hybridization between H. verticilliatus 355 and *H. giganteus*. Helianthus verticilliatus was once thought to be a hybrid species but was later 356 disproven by molecular data (Heiser et al. 1969; Ellis et al. 2006). Interestingly, H. verticilliatus 357 was previously thought to be a subspecies of *H. giganteus*, and the two species overlap in 358 range (Farwall 1916). Similarly, introgression between H. mollis and H. atrorubens is highly 359 plausible considering they are known to hybridize (Beatley 1969). In contrast, the proposed 360 introgression between H. divaricatus and H. arizonensis is difficult to explain because the 361 species do not share any current range and there are no plausible proxy ancestry donors.

362 Surprisingly, there was identified introgression between the annual and perennial clades, 363 despite strong post-pollination reproductive barriers between them. Most crosses between 364 annual and perennial sunflower species fail to set seed, and the few successful hybrids that 365 have been reported typically have low pollen viability (Heiser et al. 1969). Crosses involving the 366 hexaploid perennials, represent an exception to this rule (Atlagić and Terzić 2006), but we found 367 the strongest signal of introgression with a diploid perennial H. maximiliani, which is incompatible with members of the annual clade (Henn et al. 1998). Thus, we likely are seeing 368 369 the outcome of ancient hybridization, which occurred before reproductive isolation was 370 complete. However, the most unexpected pattern is a signal of ancestral allele sharing between 371 H. annuus/argophyllus and the outgroup, Pheobanthus (fig. 2E). These two lineages diverged 372 2.5-5.4 mya, and are therefore expected to be completely reproductively isolated (Owens and

373 Rieseberg 2014; Mason 2018), again suggesting that the signal of admixture is from ancient 374 hybridization with a now extinct species. Nonetheless, we recommend experimental crosses be 375 undertaken between *Pheobanthus* and members of the annual clade to see if hybrids can be 376 made. It is also possible that this signal represents a change in evolutionary rate in *H. annuus* 377 and *H. argophyllus*. This has been proposed to explain similar patterns of introgression signals 378 in *Papilio* butterflies (Xiong et al. 2022). More broadly, our results suggest that admixture has 379 been common in Helianthus throughout its evolutionary history, and that it has had a profound 380 effect on phylogenomic relationships in the genus (fig. 4). Thus, Helianthus can be viewed as a 381 syngameon, in which gene flow connects otherwise distinct species (Grant 1981). Our findings 382 also address earlier concerns that hybridization in *Helianthus* might be a recent consequence of 383 anthropogenic disturbance and therefore of little significance to its long-term evolution 384 (Schemske 2000). Whether these ancient hybridization events triggered diversification, as 385 suggested in other groups (e.g., Meier et al. 2017), remains unclear.

386 Ancestry of putative homoploid hybrid species

387 The TreeMix and ADMIXTOOLS2 graphs indicate that the three hybrid species are admixed, 388 thereby fulfilling the second criterion for homoploid hybrid speciation. However, there are 389 differences relative to the original scenario put forward by Rieseberg (1991). In particular, the 390 hybridization events associated with the origins of the hybrid species appear to have occurred 391 further back in time than previously believed. For *H. paradoxus*, a potential hybrid speciation 392 event is older than previously anticipated (Welch and Rieseberg 2002) since individual gene 393 trees place *H. paradoxus* most often at the base of the annuus clade, therefore at least older 394 than 1 mya. For *H. anomalus* and *H. deserticola*, our phylogenomic results suggest they are 395 sister species, and share an admixture close in time to their origin. This would also place a 396 potential hybrid speciation event further in the past than previous work suggested (Schwarzbach and Rieseberg 2002; Gross et al. 2003; see supplementary discussion for reconciliation with
previous microsatellite dating of the hybrid species origin).

399 Another important difference is that *H. anomalus* and *H. deserticola* are sister species and 400 their major parent appears to be H. niveus canescens rather than H. petiolaris fallax as 401 originally proposed (Rieseberg 1991). While this was unexpected, H. niveus canescens 402 intergrades phenotypically with H. petiolaris fallax and the two taxa share chloroplast and 403 nuclear ribosomal DNA haplotypes, which were the markers employed to hypothesize a hybrid 404 origin in the first place (Heiser et al. 1969; Beckstrom-Sternberg et al. 1991). As a consequence, 405 some taxonomic treatments have considered H. niveus canescens to be a variety of H. 406 petiolaris (Blake 1942; Schilling 2020). The present study indicates that H. niveus canescens is 407 genetically distinct from H. petiolaris fallax, although there is considerable admixture between 408 them, which might account for their phenotypic resemblance (Zhang et al. 2019). Fortunately, 409 recognition that the ancestor of H. niveus canescens was a parent of H. anomalus and H. 410 deserticola has little impact on the interpretation of previous work. The population of H. 411 petiolaris fallax that was employed for most of the experimental studies of hybrid speciation 412 came from the zone of intergradation with H. niveus canescens (Rieseberg et al. 2003; Lexer et 413 al. 2003; Gross et al. 2004; Ludwig et al. 2004). The taxa are indistinguishable morphologically 414 and ecologically in this area, and may recapitulate the ancestor of *H. niveus canescens*.

Despite the power of the phylogenomic analyses, some uncertainties remain (see supplementary discussion about the challenges for interpreting admixture signals in sunflower). For *H. paradoxus*, it is unclear whether the signal of admixture associated with its origin is independent from other members of the *annuus clade*. The triplet tests of introgression provide indirect support for independence. For D, there is a significant enrichment for *H. petiolaris – H. paradoxus* sharing as predicted under the hybrid origin scenario; gene tree topology counts show the same pattern but are not significant. However, the branch length test, which has 422 greater power to detect recent introgression, indicates more introgression between H. annuus – 423 H. petiolaris. These results may be reconciled by the long history of introgression between the 424 annuus and petiolaris clades (Strasburg and Rieseberg 2008), with the branch length test 425 successfully detecting recent introgression between H. annuus and H. petiolaris, and D offering 426 evidence of more ancient hybridization associated with the evolution of H. paradox. Likewise. H. 427 paradoxus shares more derived alleles with H. anomalus and H. deserticola than do other 428 members of the annuus clade, offering further indirect support that H. paradoxus is the product 429 of an independent hybridization event (supplementary table 2).

430 There is also uncertainty regarding the parentage of *H. anomalus* and *H. deserticola*. While 431 the role of *H. niveus canescens* as one parent is strongly supported, the other hybrid parent is 432 less clear. Is it the *H. annuus* lineage? The most likely TreeMix graph suggests the ancestor of 433 the *H. annuus* clade is the other parent, but this pattern is not strongly supported by admixtools, 434 the triplet testing or D. The hybrid species share fewer derived alleles with H. annuus and H. 435 argophyllus when compared to H. debilis, H. niveus canescens or H. petiolaris (supplementary 436 table 2), but they share more derived alleles with *H. paradoxus* when compared to *H. debilis*. 437 This may be indicative of more recent introgression between the *H. annuus* lineage and 438 members of the *petiolaris* clade sampled in this study, all of which overlap in geographic 439 distribution and hybridize with H. annuus. Alternatively, H. anomalus and H. deserticola are a 440 combination of *H. niveus canescens* and an unsampled or extinct basal lineage, as is suggested 441 in the admixtools tree. Future comparisons with an allopatric control from the H. petiolaris clade 442 (e.g., *H. niveus niveus*) will be needed to distinguish between these hypotheses.

One perhaps surprising aspect of our results is that the amount of admixture in "hybrid species" is not exceptional compared to non-hybrid species. This highlights that the amount of hybrid ancestry is not critical for hybrid speciation, only that admixture was involved in reproductive isolation (Rieseberg 1997; Schumer et al. 2014), which we do not directly address 447 in this study. Nonetheless, our results have implications for the interpretation of previous 448 experimental studies that have attempted to make this link (e.g., Rieseberg et al. 1995, 1996, 449 2003; Lexer et al. 2003; Gross et al. 2004; Ludwig et al. 2004). For example, evidence that the 450 homoploid hybrid species are more ancient than previously believed makes it more difficult to 451 determine which ecological, phenotypic and genomic changes are associated with hybridization 452 and which changes arose as a consequence of divergence after hybrid speciation (Ungerer et 453 al. 2006). We suspect that the clusters of parental markers found to be linked to QTLs in 454 segregating hybrids, and which were also found in the hybrid species genomes (Rieseberg et al. 455 2003), likely correspond to inversions. Such inversions are frequent in Helianthus (Ostevik et al. 456 2020) and often appear to be ancient, pre-dating the species they are found in (Todesco et al. 457 2020). Future studies attempting to connect reproductive barriers with admixture should be 458 conducted within a phylogenetic framework to better assess when key differences arose.

Previous work attempted to dissect parental ancestry across the putative hybrid genomes, finding strings of linked parental markers suggestive of rapid establishment of the hybrid genomes (Ungerer et al. 1998; Buerkle and Rieseberg 2008). However, incomplete lineage sorting, multiple ancient admixture events, structural variation, and the auto-correlational effects of recombination rate variation may all affect ancestry assignments across the genome. Future attempts at identifying and interpreting ancestry tracks will have to contend with this complexity.

Are there other examples of homoploid hybrid speciation in *Helianthus*? Probably – although the relevant data come from earlier studies rather than the present analysis (see supplementary discussion about outcomes of hybridization in the annual sunflower clade). Independently derived dune ecotypes of the prairie sunflower are isolated from non-dune populations by multiple reproductive barriers (Heiser 1958; Ostevik et al. 2016). Recent population genomic studies indicate that most of the traits underlying reproductive isolation map to ancient chromosomal inversions that appear to have originated via introgression from a basal lineage in 472 the annual clade (or even earlier) (Huang et al. 2020; Todesco et al. 2020). In an example 473 involving more recent hybridization, a 77-day difference in flowering between coastal and barrier 474 island ecotypes of *H. argophyllus* was found to result from a 30 Mb introgression from *H.* 475 annuus containing a functional copy of HaFT1 (Todesco et al. 2020). The dune ecotypes 476 arguably represent new species, and one of them is described as H. neglectus in most 477 taxonomic treatments (Heiser et al. 1969; Schilling 2020). Ecotypic differentiation in H. 478 argophyllus represents an earlier stage of the speciation process, but it illustrates how 479 introgression of a single gene can cause significant reproductive isolation. It is noteworthy that 480 all three potential new cases of homoploid hybrid speciation involve the colonization of 481 ecologically divergent habitats that provide ecogeographic isolation similar to that seen in the 482 ancient hybrids.

483 **Recombination rate and introgression**

484 Local recombination rate is a critical population genetics parameter that can affect ILS, gene 485 flow and genetic diversity (Ortíz-Barrientos et al. 2002; Nachman and Payseur 2012; Haenel et 486 al 2018). For example, introgressed loci are more likely to persist in regions of high 487 recombination because they more quickly unlink from deleterious neighbouring loci, such as 488 genetic incompatibilities (Brandvain et al. 2012: Schumer et al. 2018). In our dataset, higher 489 recombination regions are less likely to support the inferred species topology. This is likely a 490 combination of reduced introgression in low recombination regions, as well as lower Ne, which 491 reduces ILS (Martin et al. 2019; Li et al. 2019; Pease and Hahn 2013). The effect is strongest 492 for the node separating *H. annuus* samples, where the recombination rate estimates are most 493 accurate. Sunflowers have highly labile chromosome structure and there are numerous large-494 scale rearrangements between species in this tree, but this pattern is retained even between 495 our outgroup perennial sunflower species, suggesting that recombination rate is relatively 496 conserved even if genome structure changes (Ostevik et al. 2020). At several nodes we find

that the relationship is reversed, which may be because at those nodes we have the
introgressed topology, rather than the true species topology. Exploring the relationship between
recombination rate and tree topology is useful for identifying an accurate species topology.

500 Conclusions

501 Despite vast improvements in the quantity of genetic data available to current researchers, 502 disentangling introgression patterns is often challenging (Hibbins and Hahn 2022). When 503 multiple introgression events have occurred, this can lead to false positives or false negatives, 504 which is especially true when the true donors are not sampled or, in some cases, no longer 505 extant. Broad sampling of species can help reduce these issues and broad sampling of 506 populations within species can identify when introgression is geographically localized. Thus, 507 when evaluating the plausibility of introgression signals detected using genomic data, it is 508 important to consider factors such as reproductive compatibility and the likelihood of geographic 509 overlap over the course of evolutionary divergence. That being said, the reconstruction of 510 historical ranges is an inexact science, and information on extinct congeners is completely 511 unknown for most wild genera.

512 Phylogenomic analysis below the genus level often finds evidence of introgression, 513 especially in plants (Dagilis et al., 2021). This suggests that a significant proportion of plant 514 species have some admixed ancestry. Despite this, homoploid hybrid species are rare because 515 most studies fail to link admixture with the evolution of reproductive barriers (Schumer et al., 516 2014). Although attempts have been made to identify admixture-derived reproductive barriers 517 through purely bioinformatic analyses (e.g. Sun et al. 2020; Wang et al. 2022), links between 518 phenotypes, reproductive isolation and introgressed ancestry are required to confirm the 519 homoploid hybrid speciation. A notable exception to this is work by Wang et al. (2021), which 520 identified an admixed species in Ostryopsis, quantified reproductive barriers and transgenically 521 tested candidate genes acquired through admixture. Widespread introgression across sunflower species raises the possibility that admixture played a role in other speciation events. In
particular, chromosome rearrangements are both common and are known to cause reproductive
isolation in sunflowers (Ostevik et al. 2020; Lai et al. 2005), so future work should leverage
chromosome resolved genomes and QTL mapping to identify how whether the introgressed
rearrangements cause speciation.

527 Methods:

528 Data generation

529 We generated whole genome resequencing data for two *H. paradoxus* samples. DNA was 530 extracted from individual seedlings or dried leaf tissue using a modified CTAB extraction 531 protocol (Murray et al. 1980; Zeng et al. 2002), and an indexed Illumina sequencing library was 532 created, which included a Duplex Specific Nuclease (DSN) repeat depletion step (Todesco et al. 533 2020). This data was added to previously published whole genome resequencing data to create 534 a dataset of nine species, or subspecies, of annual sunflowers (Hubner et al. 2019; Todesco et 535 al. 2020; Owens et al. 2021). For each, we randomly selected two samples to be used in the 536 analysis. We only included two per species because we were interested in understanding 537 between species relationships, rather than within species diversity. Additionally, using fewer 538 samples substantially reduced the computational bottleneck in both variant calling and 539 subsequent phylogenetic analyses. To act as outgroups, we also included one sample each for 540 four diploid perennial sunflower species (See supplementary table 1).

541 For each sample, Illumina sequence data was aligned to the *Helianthus annuus* HA412-542 HOv2 genome using NextGenMap (v0.5.3) (Sedlazeck et al. 2013), PCR duplicates were 543 marked with samtools v0.1.19 (Danecek et al. 2021), and variants were called using GATK 544 (v4.1.4.1) (Mckenna et al. 2010). We specifically included invariant sites in GATK 545 genotypeGVCFs to facilitate downstream analyses. The sunflower genome contains a large proportion of transposable element repeats that hamper read alignment. To avoid those regions,
we used GenMap (-K 50 -E 2) (v1.3) to calculate k-mer uniqueness or mappability for all
positions in the genome (Pockrandt et al. 2020). We then removed all contiguous regions with
mappability < 1 that are greater than 100 bp. This removed 2.43 GBp out of 3.23 Gbp, but
should highly enrich retained regions for single copy sequences.

- 551 For our dataset, we filtered variant and invariant sites separately. For variant sites, we
- removed all indels and visualized the distribution of quality metrics of SNPs using bcftools
- 553 (v1.14) to extract values and ggplot (v3.3.5) in R (v4.1.2) to plot (Whickham 2011; R core team
- 554 2021). We visually selected thresholds to remove outliers based on the distribution (bcftools
- 555 view -e 'INFO/DP > 1000' -e 'INFO/FS > 50' -e 'INFO/QD < 2' -e 'INFO/SOR > 4' -e 'INFO/MQ <
- 556 30') (supplementary figure 1). Variant and invariant sites were combined, and then filtered to
- 557 require \geq 4 reads per genotype and \geq 80% of samples genotyped using VCFtools (v0.1.16)
- 558 (Danecek et al. 2011).
- 559 Code used to conduct analyses and make figures is deposited at
- 560 https://github.com/owensgl/helianthus_hybrid_species_2021.

561 Gene and species tree creation

We took two approaches to explore genome-wide gene trees. In the first, we equalized information across windows by requiring each genomic window have 10,000 called bp, leading to variable physical size. In the second approach, we equalized physical size but not information by dividing the entire genome into non-overlapping 10,000 bp windows and retained windows with \ge 2,000 called bases. In both cases, heterozygous sites were retained and coded using IUPAC coding.

568 For each of our genomic windows, we calculated a maximum likelihood phylogeny using IQ-569 Tree (v2.0.6) (Minh et al. 2020a) with *H. giganteus*, a perennial species, as the outgroup. Additionally, a single concatenated species tree using all retained bases was created using
maximum likelihood and model selection in IQ-TREE. We used a coalescent approach in
ASTRAL (v5.7.3) to estimate the species tree using both sets of gene trees separately (Zhang
et al. 2018).

We explored the diversity of gene tree topologies by using the tool findCommonTrees.py (Edelman et al. 2020) to identify and count the occurrence of each topology. We found that every tree was unique, so we subsampled trees down to a single sample per species and a single perennial outgroup species to reduce the possible tree space. We repeated the counts of trees and visualized the most common gene trees.

579 Using the concatenated phylogeny as a backbone, we measured gene concordance using 580 IQ-Tree2 (Minh et al. 2020b). This measures the percent of gene trees that support the topology 581 and the two possible alternate topologies for each branch quartet. We matched the *H. annuus* 582 recombination rate for each genomic window and calculated the percent support for each 583 topology for each recombination quintile. To determine if recombination rate affected support for 584 the species topology, we used a binomial regression with the formula:

585

gC ~ node_ID * cm_rate

586

Broad tests of introgression

We used several methods that integrate signals of introgression between all samples. Using Dsuite, we calculated the f_{branch} statistic for the species phylogeny (Malinsky et al. 2021). This statistic incorporates all valid f_4 ratios (similar to Patterson's D) as well as the phylogeny to identify branches that share allelic imbalance. This can help identify introgression events that predate speciation times and therefore have a signal in multiple species.

592 We also used TreeMix to identify introgression events (Pickrell and Pritchard 2012). This 593 program builds a graph model of population splits based on the covariance of allele frequencies. 594 It allows for migration edges to be added to the graph that explain remaining covariance not 595 accounted for by the initial graph topology. Since TreeMix relies on covariance, we pruned our dataset for SNPs in LD by only including sites with > 0.8 r^2 in 100,000 bp using `bcftools prune` 596 597 (Danececk et al. 2021), and did not include sample size correction. We ran TreeMix with 0 to 10 598 migration edges, with 99 replicates using different starting seeds, and selected the optimal 599 number of migration edges using the Evanno method in OptM (Fitak 2021). We visualized the 600 replicates with the likelihood values within 10 of the highest likelihood for each number of 601 migration edges (i.e. the models that were not the best, but were close).

602 Lastly, we built an admixture graph for the phylogeny using ADMIXTOOLS 2 (Patterson et 603 al. 2012). We first converted our VCF file to eigenvector format using a custom Perl script, and 604 then calculated all f2 statistics using the extract f2 and f2 from precomp functions. We used a 605 5 Mbp window for block bootstrapping. Admixture graphs were found using the find graphs 606 command with stop gen=200, stop gen2=20 and perennials set as the consistent outgroup. 607 This method finds graphs that fit the observed f-statistics in the data. The graph search can get 608 stuck at local optima, so we repeated each search 100 times for each number of admixture 609 events, from one to four, and retained the best fitting graph in each iteration. From this, we 610 picked the top three scoring graph for each number of admixture events. We then used 611 bootstrap-resampling of the out-of-sample scores for each graph using the function 612 gpgraph resample multi to ask whether adding additional nodes was significantly better 613 supported, by comparing the best graph with n admixture events with the best graph with n+1 614 admixture events, as well as comparing graphs.

615 Trio based tests of introgression

Given the high levels of discordant gene trees, and conflicting signals from broad tests of introgression, we explored trio-based tests of introgression using Dsuite to calculate Patterson's D (Green et al. 2010; Malinsky et al. 2021). This test looks at quartets with the relationship (((A,B),C),O) and asks if there is greater derived allele sharing between A and C or B and C. Under ILS, both should share equal counts of derived alleles, but introgression will lead to imbalance. Significance was tested using a block bootstrap algorithm. We used the perennial species as an outgroup and tested all trio sets consistent with the species phylogeny.

623 For the putative hybrid species and their parents, we used three additional approaches, the 624 branch length test (BLT), QuIBL and Twisst. For these tests, we used genomic windows with 625 equal physical size, as described above. The BLT test compares the tip-to-tip distance between 626 inferred sister clades in the minor topologies using a Mann-Whitney test (Suvorov et al. 2022). 627 Under ILS, both minor topologies should have equal distance, but if introgression is occurring 628 then the topology it produces should have reduced distance due to its more recent coalescence. 629 These tests were done in R using TreeTools (Smith 2019), using the subsampled trees with 630 only a single sample per species. QuIBL examines trios in gene trees and asks whether the 631 branch length distribution fits better to a model that includes both ILS and later introgression, or 632 purely ILS (Edelman et al. 2020). As before, we used our perennial samples as outgroups. We 633 considered models with Bayesian information criterion score difference of \geq 30 to be evidence of 634 introgression. We used Twisst to calculate the topological weighting for each possible tree using 635 the equal size gene trees (Martin and Van Belleghem 2017). This takes into account 636 phylogenetic position variation amongst individuals within a species and calculates a weight for 637 how common a particular topology is at each gene tree. We normalized weights such that each 638 genomic window had a total weight of one and counted the total weight for each topology. 639 Similar to the Discordant Count Test from Suvorov et al. (2020), we compared the counts of the 640 two minor topologies using a chi-squared test, under a null hypothesis that ILS should produce

equal counts of both minor topologies. We also calculated D across the genome using Dsuite,
with a window size of 50 SNPs and a step size of 25 for each possible candidate trio.

643 **Quantifying introgression across the genus**

644 Given the amount of gene flow identified within annual sunflowers, we expanded our 645 analysis of gene flow by using a previously published sequence capture set encompassing a 646 majority of diploid species in the genus (Stephens et al. 2015). This dataset included 170 647 aligned sequences totalling 106,862 bp and 11,407 parsimony informative sites. Since these 648 markers were anonymous and genome positions are required for block bootstrapping, we 649 aligned them to the H. annuus HA412-HOv2 reference genome using BLAST and selected the 650 location hit with the highest e-value. Sequence capture gene alignments include numerous 651 indels, so we were not able to combine this dataset with our previous WGS dataset. As before, 652 we used Dsuite to calculate D, f₄ and f_{branch}. For the species topology, we used the MP-EST tree 653 from Stephens et al. (2015), with polytomies randomly resolved.

654 Acknowledgements:

655 We would like to thank Sarah Yakimowski and Robert Sivinski for providing tissue used for 656 DNA extractions. This work was funded by a Banting Fellowship to GLO, as well as a Discovery 657 Grant from the Natural Sciences and Engineering Research Council of Canada to LHR.

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