# 1 Running Head: Phased Target Enrichment for Polyploids

2	Phasing Alleles Improves Network Inference with Allopolyploids
3	George P. Tiley <sup>1,†,*</sup> , Andrew A. Crowl <sup>1,†</sup> , Paul S. Manos <sup>1</sup> , Emily B. Sessa <sup>2</sup> , Claudia Solís-
4	Lemus <sup>3</sup> , Anne D. Yoder <sup>1</sup> , J. Gordon Burleigh <sup>2</sup>
5	<sup>1</sup> Department of Biology, Duke University, Durham NC, 27708, USA
6	<sup>2</sup> Department of Biology, University of Florida, Gainesville FL, 32611, USA
7	<sup>3</sup> Wisconsin Institute for Discovery and Department of Plant Pathology, University of Wisconsin –
8	Madison, Madison WI, 53706, USA
9	<sup>†</sup> These authors contributed equally
10	*Author for correspondence: george.tiley@duke.edu
11	
12	

#### 13 Abstract

Accurately reconstructing the reticulate histories of polyploids remains a central 14 15 challenge for understanding plant evolution. Although phylogenetic networks can provide 16 insights into relationships among polyploid lineages, inferring networks may be hampered by the 17 complexities of homology determination in polyploid taxa. We use simulations to show that 18 phasing alleles from allopolyploid individuals can improve inference of phylogenetic networks 19 under the multispecies coalescent. Phased allelic data can also improve divergence time 20 estimates for networks, which is helpful for evaluating allopolyploid speciation hypotheses and 21 proposing mechanisms of speciation. To achieve these outcomes, we present a novel pipeline 22 that leverages a recently developed phasing algorithm to reliably phase alleles from polyploids. 23 This pipeline is especially appropriate for target enrichment data, where depth of coverage is 24 typically high enough to phase entire loci. We provide an empirical example in the North 25 American Dryopteris fern complex that demonstrates how phasing can help reveal the mode of 26 polyploidization and improve network inference. We establish that our pipeline (PATÉ: Phased 27 Alleles from Target Enrichment data) is capable of recovering a high proportion of phased loci 28 from both diploids and polyploids, and that these data improve network estimates compared to 29 using haplotype consensus assemblies. This approach is shown to be especially effective in 30 reticulate complexes where there are multiple hybridization events. The pipeline is available at: 31 https://github.com/gtiley/Phasing.

32

33 Key words: Introgression; Hybridization; Reticulate Evolution; Multispecies Coalescent;

34 Divergence Time Estimation; Polyploidy; Target Enrichment; Dryopteris

35

36

### 37 INTRODUCTION

38 The phenomenon of polyploidy, or whole-genome duplication, occurs throughout the tree 39 of life. Nowhere, perhaps, is its evolutionary significance more evident than in plants, with recent 40 estimates suggesting up to 35% of vascular plant species are of recent polyploid origin (Wood 41 et al. 2009; Barker et al. 2016). Despite advances in genomic data generation and a long-term 42 interest in understanding the role of whole-genome duplication in driving plant speciation and 43 local adaptation (reviewed in Soltis et al. 2014), polyploids remain a central challenge for the 44 field of phylogenetics. One persistent problem when analyzing sequence data from polyploid 45 taxa, and especially allopolyploids, is identifying the alleles and divergent homeolog copies from 46 parental lineages. Most bioinformatic tools for processing next generation sequence data were 47 developed with diploids, or specifically humans, in mind. These approaches often collapse 48 variable homeolog sequences into a single consensus sequence for de novo assemblies or 49 assume the organism is diploid when performing genotyping and phasing for reference-based 50 assembly. For polyploids, this creates chimeric sequences that may interfere with phylogenetic 51 reconstruction and obscure signals of polyploidy and polyploid mode-of-origin. Using allelic data 52 that more accurately capture the complex genomic histories of polyploids should enable the 53 incorporation of divergent signals from polyploid loci into phylogenomic inference, distinguish 54 allopolyploidy from autopolyploidy, and identify parental taxa. However, few studies have 55 examined the potential benefits of using phased versus unphased data to reconstruct polyploid 56 histories, and there are few formal methods and little guidance for phasing alleles from polyploid 57 taxa. Here we explore the value of using phased data to reconstruct polyploid networks 58 leverage recent algorithmic advances in polyploid phasing (Xie et al. 2016) to develop a 59 bioinformatic pipeline that can phase alleles from polyploids using target enrichment sequence 60 data.

61 Previous studies of reticulate complexes have suggested phasing alleles is crucial for 62 accurate evolutionary reconstruction, at least when sampling relatively few loci (e.g., 4 to 10;

63 Rothfels et al. 2017; Eriksson et al. 2018). The applications of phased sequencing data for 64 phylogenomic studies of polyploid complexes are thus enticing; however, it remains challenging 65 to genotype and phase next generation sequencing data from polyploids. Methods exist to 66 genotype consensus loci from target enrichment data, but these have been either limited to 67 diploids (Kates et al. 2018; Andermann et al. 2019) or manual curation of variants with polyploids where both parental populations are available (Eriksson et al. 2018). Otherwise, 68 69 obtaining phased sequence data for polyploids has largely depended on costly long-read 70 sequencing to recover complete haplotype sequences (e.g., Rothfels et al. 2017), or cloning 71 PCR products (e.g., Sessa et al. 2012; Oberprieler et al. 2017). 72 Target enrichment (or HybSeg), where specific regions of the genome are isolated and 73 sequenced (Faircloth et al. 2012; Lemmon et al. 2012), is an increasingly common method for 74 collecting large-scale phylogenomic datasets, and these data can provide insights into the

evolutionary history of reticulate complexes (e.g., Crowl et al. 2017; Karimi et al. 2020) and

sources of gene tree discordance (e.g., Morales-Briones et al. 2018; Stull et al. 2020). Probe

kits for target enrichment have been developed in many land plant lineages (Wolf et al. 2018;

Johnson et al. 2019; Liu et al. 2019; Breinholt et al. 2021), and there are bioinformatic pipelines

available for custom probe design (e.g., Jantzen et al. 2020). The most common approach to

80 assemble phylogenetic datasets from target enrichment data has been to use *de novo* assembly

pipelines (Faircloth 2016; Johnson et al. 2016; Andermann et al. 2018; Breinholt et al. 2018).

82 The assembly algorithms within these pipelines typically treat variable base calls as sequencing

83 errors and consider only the most frequent nucleotide sequence while discarding the

84 alternatives (Bankevich et al. 2012; Iqbal et al. 2012; Luo et al. 2012). This results in loci in

85 which variable positions are collapsed to a single base call (haplotype consensus loci), losing

86 information related to heterozygosity. While this may be appropriate, or at least benign, for

87 phylogenetic analyses of diploid taxa (Kates et al. 2018), it may pose substantial problems when

88 attempting to investigate the evolutionary history of polyploid taxa or reticulate lineages.

89 While phylogenetic studies in plants often infer strictly bifurcating trees, the complexities 90 of allopolyploid evolution can be represented more accurately using networks. Simulations and 91 empirical analyses have suggested that phylogenetic networks can recover the reticulate 92 histories of polyploid lineages with few loci, at least when gene tree discordance due to 93 incomplete lineage sorting (ILS; Hudson et al. 1983; Pamilo and Nei 1988) is low (Oberprieler et 94 al. 2017) using parsimony methods (Huber et al. 2006; Lott et al. 2009), or even with moderate 95 ILS when explicitly modeled (Jones et al. 2013). Contemporary phylogenetic network models 96 and software packages jointly consider gene tree variation due to allele sampling error as 97 described by the multispecies coalescent (MSC; Rannala and Yang 2003) and gene flow 98 modeled as episodic introgression events (Solis-Lemus and Ané 2016; Wen et al. 2016; Zhang, 99 Ogilvie et al. 2018, Flouri et al. 2020). Depending on the complexity and goals of the research 100 question, these methods can search for networks with a constrained number of reticulation 101 events using guartet-based maximum pseudolikelihood (Solis-Lemus and Ané 2016; Wen et al. 102 2018) or a full-likelihood Bayesian model where the number of reticulations is a parameter (Wen 103 et al. 2018; Zhang et al. 2018). Also, it is possible to estimate model parameters (divergence 104 times, population sizes, and the fraction of introgressed genes) on a fixed species network using 105 a full-likelihood Bayesian model that allows efficient computation with large numbers of loci 106 (Flouri et al. 2020). We refer to these network models from here on as the multispecies 107 coalescent with introgression (MSci), consistent with Flouri et al. (2020); although, other names 108 have been used, such as the network multispecies coalescent (NMSC; e.g Zhu and Degnan 109 2017) and multispecies network coalescent (MSNC; e.g., Wen et al. 2016). We emphasize that 110 network approaches to investigate polyploid complexes are not novel (e.g., Huber et al. 2006; 111 Lott et al. 2009; Jones et al. 2013), but the difficulty of collecting appropriate genomic data from 112 polyploids for such analyses has limited their use.

113 To address the issues outlined above, we have developed a pipeline, PATÉ (Phased 114 Alleles from Target Enrichment data), that can phase genotyping data for individuals of a known

ploidy without the need for sampling their parental lineages. PATÉ was designed with scalability 115 116 and population-level sampling in mind for target enrichment projects where deep coverage from 117 paired-end Illumina data allow calling of high-quality variants. In this study, we first use 118 simulations to explore the ability of network approaches to reconstruct the history of 119 allopolyploidy in the presence of ILS, and whether phasing the data affects the accuracy of the 120 reconstruction. We show that using phased allelic data can improve network estimation and 121 divergence time estimation compared to using haplotype consensus sequences, but also 122 highlight scenarios where phasing may not be necessary or beneficial. Next, we describe the 123 individual steps used by PATÉ to phase target enrichment data. For an empirical example of the 124 benefits of PATÉ, we compare phased and unphased (haplotype consensus) data to infer the 125 evolutionary history of the North American Dryopteris complex, a model system for reticulate 126 polyploid evolution (Sessa et al. 2012a; Sessa et al. 2012b), using new targeted enrichment 127 data. The system includes four diploid species, as well as one extinct diploid, that have formed 128 five allopolyploids in which there is high confidence in the parent-progeny relationships (Fig. 1), 129 although numerous sterile allopolyploid species have also been reported within the complex 130 (Montgomery and Paulton 1981). The allopolyploid species have relatively ancient origins, with 131 the best estimates placing hybridization events between six and 13 Ma (Sessa et al. 2012b). 132 PATÉ is largely successful in recovering phased haploid sequences from polyploid individuals, 133 and networks inferred from phased data more accurately represent the North American 134 Dryopteris complex than those inferred from unphased data.

135

#### 136 MATERIALS & METHODS

### 137 Testing the Effects of Phasing on Network Inference through Simulation

Simulating phased and unphased sequence data for an allopolyploid — We simulated
 gene trees and their nucleotide sequence data using the MSC model with BPP v1.4.1 (Flouri et
 al. 2018) under a five-species network (Fig. 2). All simulations used the Jukes-Cantor (Jukes

141 and Cantor 1969) model of sequence evolution with no rate heterogeneity. The allopolyploid species *E* was treated as two lineages (*E* sister to *B* and *F* sister to *C*) whose alleles were 142 pooled to form the hybrid species E at time  $\tau_h$ . This makes species E a tetraploid hybrid with the 143 144 parents B and C.  $\theta$  was constant among lineages and set at 0.01. Assuming a per-generation mutation rate ( $\mu$ ) of  $1 \times 10^{-8}$  and one year per generation yields an effective population size 145 146  $(N_e)$  of 250,000 and root age of 10 Ma for the simulation network (Fig. 2). We also simulated 147 data under a shallow divergence scenario, in which all node ages were divided by 10, and a 148 deep divergence scenario, in which all node ages were multiplied by 10. This changes the root 149 age  $(\tau_r)$  to 0.01 (1 Ma) and 1.0 (100 Ma), respectively.

150 Because the distance between speciation nodes is 0.025 and  $\theta = 0.01$  for the simulation 151 network (Fig. 2), there are five coalescent units between nodes,  $T = \frac{\tau}{\left(\frac{\theta}{2}\right)}$  (Yang 2006), which

152 implies a near-zero probability of gene tree discordance due to ILS (Hudson 1983). We

incorporated ILS into the simulation by reducing the node heights  $\tau_u$  and  $\tau_s$  to 0.0375 and 0.05

154 (for a low level of ILS) and 0.03 and 0.035 (for a moderate level of ILS). This increases the

probability of ILS at node *u* and *s* to 0.05 and 0.25, respectively. Thus, we used a total of nine

simulation conditions that combined three levels of evolutionary distance and three levels of ILS.

157 While our simulations do not explore extreme levels of gene tree discordance, they allow us to

158 learn about some general features of increasing ILS on network inference with different data

types. We simulated 1000 gene trees and their sequences, 500bp in length, under the MSC for

160 each of the nine conditions. We also explored the effects of sampling fewer genes on

161 downstream analyses. For each replicate of 1000 gene trees and their sequences, we randomly

sampled 400, 40, and four genes without replacement.

All simulations sampled haploid data. Two haploid sequences were sampled for each diploid species and four haploid sequences were sampled for the allopolyploid species *E*, where two sequences came from each parental lineage. We then investigated unphased data in three

166 ways. First, to generate unphased genotype sequences, the simulated haploid sequences for 167 each species were collapsed into a single sequence in which heterozygous sites were 168 represented by IUPAC ambiguity codes (genotype). The allopolyploid species was not restricted 169 to only biallelic sites. Second, we generated haploid consensus sequences, where for each 170 variable site, only one base was randomly retained (consensus). This could represent a case in 171 which read coverage across a locus is highly uneven such that a haploid sequence is actually a 172 chimera of two or more alleles. Finally, we simply picked one phased haploid sequence, which 173 is possible when one parental haplotype has a majority of reads for a locus (pick one). This 174 scenario where only one parent's sequence would be recovered in the offspring could be 175 anticipated in real data due to subgenome dominance (e.g., Buggs et al. 2014; Bird et al. 2018). 176 In practice, we expect most *de novo* assemblers to generate output in between the haploid 177 consensus data and pick one data.

178

179 Inferring species networks with phased and unphased simulated data — We estimated 180 species networks with PhyloNetworks v0.12.0 (Solis-Lemus et al. 2017) using Julia v.1.4.1 181 (Bezanson et al. 2015) from either the true gene trees used to simulate the data, or gene trees 182 estimated from the phased or unphased sequence data. For estimated trees, we used IQTREE 183 v1.6.10 (Nguyen et al. 2015) with the same model used for simulation. Each PhyloNetworks 184 analysis used the species tree (A,((B,E),(C,D))) as the starting tree and allowed zero, one, or 185 two reticulation events. Each analysis included ten independent optimizations of the 186 pseudolikelihood score. We considered larger numbers of reticulations an improvement if they 187 were two or more pseudolikelihood units lower than the best model. We compared the 188 estimated networks with one reticulation to the true network with the hardwiredClusterDistance 189 function (Huson et al. 2010) in PhyloNetworks. This allowed us to score the number of 190 replicates that 1) recovered the correct number of reticulations and 2) matched the true network 191 when the number of reticulations was set to one. We estimated networks for samples of four,

40, 400, and 1000 gene trees for each of the 30 replicates for each of the nine simulationdivergence and ILS conditions.

194

195 Effect of phasing on divergence time estimation — We also used our simulated phased 196 and unphased data to estimate divergence times under the MSci model (Flouri et al. 2020) 197 using BPP v4.2.9. Here, we estimate divergence times ( $\tau s$ ), population sizes ( $\theta s$ ), and the 198 proportion of introgressed loci ( $\varphi$ ) on the correct fixed species network. MSCi analyses used 199 diffuse priors on  $\tau_r$  and  $\theta$  with a mean on their simulated values with  $\varphi \sim \beta(1,1)$ . Phased, 200 consensus, and pick one sequences were treated as haploids while genotype sequences were 201 treated as unphased diploids and used the analytical phasing (Gronau et al. 2011) implemented 202 in BPP. Although this is not correct for the tetraploid, it is arguably more appropriate than 203 treating all of the genotype sequences as haploid. Each Markov chain Monte Carlo (MCMC) analysis collected 10,000 posterior samples, saving every 100 generations, while discarding the 204 205 first 100,000 generations (i.e., 10% of the total run) as burnin. All scripts for simulation and 206 subsequent analyses of simulated data are available in Dryad (X).

207

### 208 A Phasing Pipeline for Polyploids

209 Target enrichment data — We were motivated by the general premise of using phased 210 data to infer reticulate evolutionary histories of polyploids based on the success of empirical 211 studies where phasing was informative about hybridization or introgression events (e.g., 212 Oberprieler et al. 2017; Eriksson et al. 2018). We were aware of few instances of phasing 213 genomic or phylogenomic data in polyploids, except in cases where chromosome-level whole-214 genome assemblies have characterized subgenomes in allopolyploid crops (Yang et al. 2017; 215 Colle et al. 2019) or emerging results that are dependent on the sampling of parental lineages 216 (Freyman et al. 2020; Nauheimer et al. 2020). We designed PATÉ for target enrichment data

because of the availability of such data for many of taxa, but it is applicable to other types ofdata with paired-end Illumina reads.

219 The end product of a *de novo* target enrichment assembly pipeline (such as HybPiper: 220 Johnson et al. 2016) generally is a single consensus sequence for each locus for each 221 individual. Allelic variation may be represented by ambiguous nucleotide codes within the single 222 consensus sequence or lost when the pipeline outputs the haplotype consensus sequence 223 where the majority vote from a collection of reads is used. We use these existing de novo 224 assembly pipelines as a starting point to provide the reference sequence for each locus for each 225 individual and leverage a recent phasing algorithm with high-quality variant calls to recover 226 phased haplotype sequences for taxa with known ploidy levels. Ploidy levels were well-227 characterized for individuals in our Dryopteris analyses, but for unknown systems there are 228 existing methods to estimate ploidy directly from target enrichment data (Weiss et al. 2018; 229 Viruel et al. 2019) in the absence of other sources, such as flow cytometry (Farhat et al. 2019).

230

*Phasing alleles within loci* — PATÉ (Fig. 3) starts with assembled target enrichment loci,
such as the supercontig files output from HybPiper (Johnson et al. 2016) that contain a single
haplotype consensus sequence from each individual per locus. Reads for each individual are
then realigned to their consensus locus using BWA v0.7.17 (Li and Durbin 2009). PCR

235 duplicates are flagged with MarkDuplicates in Picard v2.9.2

236 (http://broadinstitute.github.io/picard), and variant calls are computed with HaplotypeCaller in 237 GATK v.4.1.4 (McKenna et al. 2010). We applied the following hard filters with VariantFiltration 238 in GATK: (1) QD < 2.0, (2) FS > 60.0, (3) MQ < 40.0, (4) ReadPosRankSum < -8.0, (5) AF < 239 0.05 || AF > 0.95. These loosely follow community recommendations on filters for germline 240 variant discovery (DePristo et al. 2011). Notably, we do not perform quality score recalibration 241 or filter on the mapping quality rank sum, as we anticipate allopolyploids could have a lower 242 mapping quality associated with an alternate allele due to sequence divergence or structural 243 variation among homeologous chromosomes. We also consider a very narrow window for 244 filtering on allele frequency. Because increasing ploidy levels will generate smaller anticipated 245 ratios of alternate to reference alleles, coupled with sequencing error and read stochasticity, we 246 only aim to remove the most extreme cases. For example, if almost all reads support the 247 alternate allele at a site, it is difficult to diagnose if the error lies in the consensus assembly or 248 the read alignment. In these cases, only the reference site is retained, and the variant does not 249 pass the allele frequency filter. However, the allele frequency filter could be removed if 250 investigators are focused on organisms with extremely high ploidy levels.

251 Biallelic SNPs that pass filters are then phased with H-PoPG v.0.2.0 (Xie et al. 2016). H-252 PopG solves a heuristic phasing problem efficiently using dynamic programming. Although not 253 guaranteed to be an optimal solution, H-PoPG has been shown to have high accuracy while 254 also being fast (Xie et al. 2016; He et al. 2018; Moeinzadeh et al. 2020). Phasing variants in polyploids is difficult because for *n* variants and *k* ploidy, there are  $2^{n-1}(k-1)^n$  possible ways 255 256 to link the sites together. H-PoPG evaluates possible solutions efficiently by grouping reads into k groups in such a way that differences within the groups are minimized. Focusing on target 257 258 enrichment data also constrains the complexity of the phasing problem compared to whole-259 genome alignments (i.e., haplotype blocks are constrained to about 1000 bp). We then used the 260 phased variants to create individual allele sequences, where invariable sites are filled in based 261 on the reference sequence. In cases where there is no linkage information to phase across the 262 entire locus, we retain the phasing only for the longest block. The variants for the shorter 263 haplotype blocks can be collapsed into IUPAC ambiguity codes or treated as missing data based on the investigator's preferences. PATÉ outputs analysis-ready fasta files with multiple 264 265 alleles per species. Variants are only phased within loci; we do not attempt to assign loci to 266 parental subgenomes. While this may complicate analyses of concatenated multi-locus 267 datasets, it is ideal for the MSC that assumes free recombination between loci and can leverage 268 multiple alleles per species for estimating  $\theta s$ . Those interested in concatenated analyses can

269 use other recent approaches that assign gene copies to parental subgenomes (Freyman et al.

270 2020; Nauheimer et al. 2020).

271

# 272 Analyses of a Species Complex with Allopolyploidy

The North American wood fern complex (Dryopteris) — We tested PATÉ using new 273 274 target enrichment data from nine North American Dryopteris species, including both 275 allotetraploid and allohexaploid taxa, with well-studied reticulate relationships (Sessa et al. 276 2012a, b) as well as two outgroup taxa from the sister genus *Polystichum*. All putative parental 277 lineages are represented in our dataset, with the exception of a hypothesized extinct lineage (D. 278 semicristata: Sessa et al. 2012b). We sampled two or three individuals for each Dryopteris 279 taxon (Table S1). The target enrichment data were generated from the GoFlag 408 flagellate 280 land plant probe set (Breinholt et al. 2021) at RAPID Genomics (Gainesville, FL). The target 281 regions for this probe set are 408 exons found in 229 single or low-copy nuclear genes. We 282 generated haplotype consensus assemblies for each with HybPiper (Johnson et al. 2016). The 283 resulting supercontig sequences became our reference sequences for genotyping and phasing 284 with PATÉ. We aligned both phased and unphased (i.e., the reference supercontig sequences) 285 with MUSCLE with default settings (Edgar 2004).

286

287 *Three species tests* — We first explored the value of phasing data when estimating 288 reticulate relationships among three species, including two diploid parental lineages (D. 289 expansa and D. intermedia) and their putative allotetraploid descendent (D. campyloptera). The 290 two diploid parents last shared a common ancestor during the Late Eocene and Early Miocene, 291 approximately 23 Ma (Sessa et al. 2012b). We used both a full-likelihood Bayesian approach 292 and a topology-based pseudolikelihood approach to estimate the correct species relationships 293 from phased and unphased data. First, using BPP v.4.1.4 (Flouri et al. 2020), we estimated log-294 marginal likelihoods (In mL) with stepping-stone sampling (Xie et al. 2011) for the three possible

rooted three taxon trees and twelve possible network models that imply differences for the
timing and direction of allopolyploidy and the presence of unsampled ancestral lineages
(Supplementary Fig. S1). Each ln *mL* estimate used 24 steps, and each step had a posterior
sample of 10,000, saving every 100 generations after a 100,000 generation burnin (10% of the
total run). The ln *mL* values were then used to calculate the fifteen model probabilities following
equation 1 (e.g., Beerli et al. 2019).

Р

$$(model) = \frac{exp(ln \ mL_{model} \ - \ ln \ mL_{max} \ )}{\Sigma_i [exp(ln \ mL_i \ - \ ln \ mL_{max} \ )]} \qquad Equation \ 1$$

302 We repeated analyses for 30 random subsets of 40 and then four loci to explore the 303 effects of the number of loci on inferring allopolyploidy. Next, we used PhyloNetworks to test the 304 presence and placement of gene flow between the three species. For that analysis, we also 305 included the sequences from the two *Polystichum* outgroups. IQ-TREE v1.6.10 (Nguyen et al. 306 2015) and model selection by ModelFinder (Kalyaanamoorthy et al. 2017) was used to estimate 307 gene trees for the phased and unphased data. The starting tree was obtained with ASTRAL III 308 v5.6.3 (Zhang, Rabiee et al. 2018). We tested the presence of zero, one, or two reticulations 309 with slope heuristics (Solis-Lemus and Ané 2016). Each analysis used ten independent 310 optimizations. In addition to the dataset of all loci, we analyzed the same 30 random subsets of 311 40 and four loci from the marginal likelihood analyses.

312

Nine species tests — We also investigated the differences in results from phased versus consensus sequences when estimating a network for the nine-species complex, which involves multiple reticulation events on an edge and thus should be difficult for network estimation (Solis-Lemus and Ané 2016). Because the increased number of species and complexity of reticulation in *Dryopteris*, evaluation with marginal likelihoods was not computationally feasible. Instead, we performed analyses of the nine-species complex and two outgroups with phased and unphased haplotype consensus data with PhyloNetworks as described above, but we allowed up to six

reticulation events. We used ASTRAL III v5.6.3 (Zhang, Rabiee et al. 2018) to generate the
starting species tree for network estimation using gene trees inferred from IQ-TREE v1.6.10
(Nguyen et al. 2015) with the best model selected by ModelFinder (Kalyaanamoorthy et al.

323 2017).

324

325 **Results** 

#### 326 Simulation Shows Benefits of Phased Data

327 *Network inference* — In our simulation results, both phased (i.e., the haplotypic allele 328 sequences) and unphased data (i.e., genotype, consensus, and pick one) performed well when 329 the goal is only to detect the correct number of reticulations (Supplementary Fig. S2). The only 330 case where analyses did not converge to the true number of reticulations was in the presence of 331 moderate ILS and high nucleotide divergence; however, this appears largely due to gene tree 332 estimation error, as analyses using the true simulated gene trees greatly outperformed those 333 using gene trees estimated from the simulated sequence data. However, using phased data 334 provides more accurate estimates of the placement of the reticulation edge in comparison to 335 using genotype data, and to a lesser extent, consensus sequences (Fig. 4). When the true gene 336 trees were used, which have information about the allopolyploid's hybridization event (i.e., the 337 allele sequences are sister to their respective parents in every tree), the correct network can be 338 inferred with 40 or fewer loci when nucleotide divergence is moderate. The gene trees 339 estimated from phased data perform equally well, although they require a few more gene trees 340 when nucleotide divergence is low and ILS moderate. Analyses using the genotype data almost 341 never recover the true network for these medium and low divergence scenarios, regardless of 342 the amount of ILS. Analyses based on pick one data perform similarly well to the phased and 343 true data when sampling 400 loci, but they are less accurate for four or 40 loci at low and 344 medium divergences. Analyses using the consensus data perform poorly for low numbers of loci 345 under a low divergence and no ILS scenario, but they are capable of recovering the true

reticulation when sampling 400 or more loci for the other low and medium divergence cases. In
the high divergence simulations, all data types could infer the true network if enough genes
were sampled, but analyses with the phased data required fewer genes than others.

349

350 *Divergence times* — Using phased data also improves divergence time estimates when 351 nucleotide divergence is low, but not when divergence is moderate or high (Fig. 5). When 352 divergence was low, the timing of reticulate events was accurately estimated when using 353 phased haplotypic data, but was overestimated when using genotype and especially consensus 354 data. For analyses with genotype and consensus data, as the number of loci increased and 355 uncertainty in the posterior was reduced, the posterior mean did not converge to the true 356 estimate and the simulated value was not within the highest posterior density (HPD) interval. 357 Additionally, all other nodes in the species network were overestimated with genotype or 358 consensus data, while the phased data were capable of recovering the simulated divergence 359 times (Supplementary Figs. S3-S5). Aside from some cases with the consensus sequences, all 360 four data types performed similarly with four loci; however, this is likely due to the posterior 361 being dominated by the prior in the absence of enough data. There was little improvement in 362 divergence time estimates when going from 40 to 400 loci, aside from further reduction in the 363 HPD intervals.

364 For medium sequence divergence, there was little difference between the phased and 365 unphased data. Phased sequences slightly underestimated the timing of hybridization while 366 unphased data slightly overestimated the timing of these events. However, phased data 367 accurately estimated the age of older speciation nodes that were again systematically 368 overestimated with unphased data (Supplementary Figs. S6-S8). At a high level of nucleotide 369 divergence, genotype data were capable of accurately estimating all divergence time 370 parameters while the phased data underestimated the timing of hybrid events (Fig. 5; 371 Supplementary Figs. S9-S11). Notably, the pick one data performed very well for all divergence

time estimation scenarios. Divergence times were not strongly affected by increasing levels of
ILS, likely because estimates were performed with the MSci model and we did not explore very
high ILS scenarios, but age estimates improved slightly for the genotypes, consensus, and pick
one data with increasing ILS.

376

### 377 Analyses of target enrichment data from Dryopteris

378 Recovery of Phased Loci — On average, 62% of loci sequenced for an individual were 379 phased with eight variants passing filters (Table S2). The ploidy level appears to be strongly 380 associated with the number of phased loci. Among diploids, only 30% of loci were phased; the 381 other 70% of diploid loci were either homozygous or had too few linked variants for phasing. For 382 tetraploids and hexaploids, 87% and 94% of loci, respectively, were phased such that two or 383 more phased haplotype sequences could be recovered. Among loci where phasing was 384 possible, variants were almost always resolved as a single haplotype block, as opposed to 385 being split into two or more blocks because not enough reads were available to physically link 386 variants. For polyploids, the number of phased haplotype sequences most frequently matched 387 the ploidy level except in the case of a single *D. campyloptera* individual (B087-D08), which also 388 had few recovered loci. Phasing data only extended sequence alignment length by about four 389 base pairs, but it more than doubled the number of parsimony informative sites (Table S3).

390

391 Placing a single reticulation event — For our three-species full-likelihood analyses with 392 the MSCi, both phased and unphased data recovered the anticipated reticulation hypothesis, 393 identifying *D. campyloptera* as an allotetraploid with the two diploid parental lineages *D.* 394 expansa and *D. intermedia*, when using all loci (Table 1). Model probabilities show decisive 395 support for a scenario where there are two unsampled ancestral populations that were the 396 progenitors of *D. campyloptera*, as opposed to *D. campyloptera* being a hybrid species with 397 extant *D. expansa* and *D. intermedia* as parents. All model parameters converged for both

398 phased (Supplementary Figs. S12 and S13) and unphased data (Supplementary Figs. S14 and 399 S15). Divergence time estimates were older in the analysis of phased data, although the relative 400 order of divergence events was the same using phased and unphased data (Fig. 6). There was 401 more uncertainty in the  $\theta$  estimates, especially for the allophyploid species *D. campyloptera* and 402 the two ancestral populations of the parental lineages that formed the polyploid ancestor 403 (Supplementary Figure S16). Repeating the analyses with fewer loci did not always produce the 404 same result, but there was either decisive support for a model or multiple plausible models that 405 all had the correct species relationships and direction of introgression for both phased and 406 unphased data when using 40 loci (Fig. 7). The only difference between models was the 407 presence or absence of ancestral  $\theta s$  for unsampled lineages. Analyses with four loci produced 408 less reliable results for both phased and unphased data. The four-locus analyses of phased and 409 unphased data found some non-negligible model probability for trees without hybridization or 410 networks where hybridization was incorrect in nine and twelve out of 30 replicates, respectively 411 (Fig. 7).

When performing a network search based on gene tree distributions, both the phased 412 413 and unphased data were able to recover the hypothesized allopolyploidy event (Fig. 8; 414 Supplementary Table S5). Both analyses inferred the major branch to be D. intermedia with the 415 minor branch from *D. expansa*. Phased data estimated a slightly smaller inheritance probability 416 compared to the unphased data. These findings from PhyloNetworks are consistent with 417 parameter estimation under the MSci model, where phased data estimated a slightly smaller 418 mean  $\varphi_h$  compared to unphased data (Table S4). When sampling 100 replicates of 40 loci, 98% 419 of replicates for phased data and 100% for unphased data were capable of detecting a single 420 hybrid event in the data. When sampling four loci, this drops to 66% and 85%, respectively (Fig. 421 8). Phased data more frequently recovered the correct network (58%) compared to unphased 422 data (38%) with 40 loci; however, the converse is true for four loci, with 34% correct for phased 423 and 41% correct for unphased (Fig. 8). Unphased data also got the network wrong more

frequently than phased data, such that phased data had the direction of introgression incorrect
in 36% and 12% of replicates while unphased data were incorrect in 60% and 38% replicates for
40 and four loci, respectively (Fig. 8).

427

428 Inferring relationships among a complex with multiple reticulation events — The network 429 recovered for phased data identified three out of five hypothesized reticulation events among 430 the nine Dryopteris species (Fig. 9; Supplementary Table S5). The allotetraploid D. celsa was 431 correctly identified with diploid D. ludoviciana and D. goldiana as parents. Analysis of the 432 phased data detected a low level of gene flow from the common ancestor of this clade into 433 tetraploid *D. cristata*, which has *D. ludoviciana* as one hypothesized parent while the other 434 parent lineage (D. semicristata) is assumed to be extinct (Sessa et al. 2012b). Dryopteris 435 carthusiana is another tetraploid that is assumed to share the extinct common ancestor with D. 436 cristata, but has experienced introgression from *D. intermedia*, with a high inheritance probability of 0.43 (Fig. 9). However, the phased data missed the putative allotetraploid case of 437 438 D. campyloptera, despite the strong evidence for this reticulation event in our earlier three-taxon 439 analyses. Our network with phased data also failed to identify the putative reticulate 440 evolutionary history of *D. clintoniana*, an allohexaploid where allotetraploid *D. cristata* and 441 diploid *D. goldiana* are assumed to be the parents (Sessa et al. 2012b). The spectra of quartet 442 concordance factors were overall similar between the phased and unphased data 443 (Supplementary Fig. S17), but the phased data were arguably more accurate. 444 Although the phased data were not successful in recovering all hypothesized reticulate 445 relationships, they performed better than the unphased data. Analyses of unphased data were 446 capable of finding the allotetraploid history of *D. celsa* with an inheritance probability similar to

the phased data (Fig. 9). Hybridization between *D. intermedia* and *D. carthusiana* was also

detected; however, the directionality was reversed, with gene flow going from the allotetraploid

into the diploid. A third reticulation edge was found in the unphased analysis, from the common

450 ancestor of *Dryopteris* into the common ancestor of *D. clintoniana* and its sister clade. This 451 hybrid edge is difficult to reconcile because of the hypothesized extinct common ancestor that 452 contributed to both *D. cristata* and *D. carthusiana*. *Dryopteris clintoniana* is correctly placed in 453 the major species tree topology, as a grade between *D. cristata* and *D. goldiana*. This 454 reticulation edge from the *Dryopteris* common ancestor may reflect the extinct lineage and a 455 high degree of gene tree variation.

456

#### 457 DISCUSSION

458 New phylogenetic network methods offer the promise of elucidating the often complex 459 reticulate histories of polyploid lineages, even in the presence of ILS. Our results demonstrate 460 that phasing polyploid target enrichment data can improve the accuracy of such network 461 inferences as well as divergence time estimates for the networks, and we describe a novel 462 pipeline (PATÉ) to address the difficult problem of phasing polyploid data. Although PATÉ could 463 handle different types of genomic data, such as transcriptomes and whole genomes, target 464 enrichment data are ideal for investigation because they often yield high and even coverage 465 across loci. Because MSC methods assume treat loci independently and assume free 466 recombination between loci, it is not necessary to assign individual loci to parental subgenomes. 467 However, the allele sequences output by PATÉ can also be used as input for the recently 468 developed Homologizer (Freyman et al. 2020) or HybPhaser (Nauheimer et al. 2020), which 469 attempt to phase across loci and recover parental subgenomes. Our methods also enable 470 population genomic studies of polyploids where accurate estimation of site frequency spectra 471 can be used for demographic analyses otherwise complicated by polyploidy (e.g., Excoffier et 472 al. 2013; Liu and Fu 2020) or SNP-based network inference in the absence of variation suitable 473 for gene tree estimation (Blischak et al 2018; Olave and Meyer 2020).

474

### 475 Promises and Pitfalls of Phasing

476 The prospect of using alleles from phased genomic data presents an exciting step 477 towards revealing the evolutionary history of polyploids, which remains a critical impediment 478 within the plant evolution community (McKain et al. 2018). Strategies for explicitly addressing 479 this challenge are only now emerging (Freyman et al. 2020; Nauheimer et al. 2020), and PATÉ 480 can be a useful tool by phasing variants for many individuals while leveraging genotyping 481 information. Our simulations demonstrate that phasing can improve estimates of reticulate 482 evolutionary relationships using network methods. Phased data can more accurately recover 483 the placement and directionality of hybrid edges than various types of unphased data in 484 simulations (Fig. 4) and empirical analyses with limited numbers of loci (Fig. 7). The advantages 485 of using phased versus unphased data for network estimation decrease when a large number of 486 loci are sampled (Table 1; Fig. 8).

487 Perhaps an underappreciated aspect of phased data is their ability to improve 488 divergence times estimates (Fig. 5; Supplementary Figs. S3-S11; Anderman et al. 2019). Our empirical analyses also demonstrated how the timing of introgression  $\tau_h$  can be greatly affected 489 490 by whether phased or consensus data are used. In our Dryopteris analyses, the estimate from 491 phased data was nearly four times older than the estimate from unphased data (Supplementary 492 Table S4). The direction of this difference was unanticipated, because our simulations 493 suggested the consensus data should overestimate age compared to phased data. This 494 highlights the difficulties of simulating data that capture real complexities and makes deciding which estimate is more reliable somewhat difficult; however, the uncertainty of  $\tau_h$  for haplotype 495 496 consensus data reflected in its posteriors (Supplementary Fig. S14) gives us more confidence in 497 the phased estimates (Supplementary Fig. S12). Because we used the MSci model for 498 divergence time estimation, we did not observe any effect of ILS on age estimates in our 499 simulations, but if we were using concatenation methods to date the divergence times for the

allopolyploids two subgenomes, nodes affected by ILS should be overestimated (Tiley et al.2020).

502 In most cases phasing appears to be beneficial, but it may be problematic when the 503 parental lineages are deeply diverged. Although the phased data were able to accurately 504 estimate the age of older speciation nodes (Supplementary Figs. S3-S11), as phylogenetic 505 information is lost from multiple hits, the influence of the prior becomes more substantial. These 506 deep divergence simulation scenarios may border on being biologically unrealistic, as identifying 507 an allopolyploid and its two parental lineages becomes more difficult over time due to extinction 508 and population genetic processes, but it provides some expectations for the performance of 509 phased and unphased data in the presence of high nucleotide divergence. Our simulations 510 showed that when alleles are phased but only one is sampled, as in our pick one simulations, 511 network and divergence time estimation is similar to having all phased alleles present. We also 512 showed where consensus data can perform poorly through simulation (Figs. 3 and 4) as well as 513 empirical analyses where the direction of introgression was more frequently reversed in a 514 simple example of two parental lineages and an allopolyploid (Fig. 8). When enough reads are 515 available to call high-confidence variants, we suggest that phasing with PATÉ can improve 516 network and divergence time estimation for species complexes with low to moderate sequence 517 divergence. However, when investigating very ancient hybrid events, unphased genotype data 518 may be preferential, and using analytical approaches that integrate over phases (Gronau et al. 519 2011; Flouri et al. 2020) may outperform analyses with phased sequences because allele-520 specific information no longer captures shared ancestry with parental lineages and haplotype 521 blocks become smaller due to recombination.

522

#### 523 Challenges of Network Estimation

524 Our analyses highlight the difficulty of estimating the evolutionary histories of reticulate 525 complexes using phylogenetic methods, regardless of data type. The full-likelihood

526 implementation of the MSci model appears to be useful for limited cases, but these methods are computationally demanding. Thus, they may not be practical for generating hypotheses and 527 528 exploring unknown relationships for large numbers of taxa (e.g., Zhang et al. 2018). Quartet-529 based methods are fast and accurate when there are limited numbers of reticulations (Solis-530 Lemus and Ané 2016) and show similar accuracy to full-likelihood methods for estimating 531 introgression probabilities (Flouri et al. 2020). However, there are scenarios where true network 532 topologies become non-identifiable (Solis-Lemus and Ané 2016). For example, when multiple 533 introgression events affect the same lineage, the expected guartet distribution under the MSci 534 model becomes a poor fit for the empirical data (Cai and Ané 2020) and the network estimated 535 may be incorrect. These effects were evident in our empirical analyses where the relationship 536 between *D. campyloptera*, *D. expansa*, and *D. intermedia* was missing in the nine-species 537 analysis (Fig. 9), which we expect is due to a reticulation edge present between D. intermedia 538 and *D. carthusiana*. Phasing sequence data adds information that can improve estimates (Supplementary Table S3), but unsampled or extinct lineages, such as the hypothesized D. 539 540 semicristata, can create significant barriers to recovering the true evolutionary history of 541 reticulate complexes, regardless of how many loci or individuals are available.

542

### 543 Insights into Dryopteris Evolution

544 The North American Dryopteris complex has been well-characterized through the study 545 of multi-locus nuclear and chloroplast phylogenies, morphology, cytological observations, 546 chromatography, and isozyme analyses (reviewed in Sessa et al. 2012b). This makes it a useful 547 system for testing our phasing pipeline, and our analyses add nuance to our understanding of 548 some of the relationships among Dryopteris species. For example, our results indicate D. 549 campyloptera has received slightly more loci from D. intermedia than D. expansa (Fig. 8), and 550 D. intermedia is thought to be the likely maternal progenitor (Sessa et al. 2012b). Although our 551 analysis of marginal likelihoods for all target enrichment loci suggested the presence of

552 unsampled ancestral populations (Table 1), the age of introgression and divergence of the D. 553 campyloptera ancestral population from the D. intermedia and D. expanse parental lineages is 554 consistent with hybrid speciation rather than a lineage that was isolated from D. intermedia and 555 received more recent gene flow from D. expansa (Fig. 6). Following the allopolyploidy event, the 556 D. intermedia genome was likely dominant, providing some selective advantage for D. 557 campyloptera in its distribution at the time (Bird et al. 2018). Similar insights can be gained from 558 the nine-taxon analyses, which suggests *D. ludoviciana* is the dominant genome in *D. celsa*. 559 Dryopteris ludoviciana is the maternal parent of D. celsa, again suggesting some bias in 560 retaining homeologous alleles from the maternal lineage, which provides the chloroplast 561 genome in ferns (Sessa et al. 2012b). 562 The nine-taxon analyses also support the hypothesis of the unsampled diploid lineage D. 563 semicristata, based on the placement of D. carthusiana as sister to the rest of Dryopteris in the 564 phased nine-taxon analyses, but with D. carthusiana having received over 40% of its genes from *D. intermedia* more recently. Our analyses suggest that *D. cristata* did not have *D.* 565 566 ludoviciana as a progenitor, but rather an unsampled common ancestor of D. goldiana and D. 567 ludoviciana. In the case of D. cristata, both parental diploid lineages, including D. semicristata, 568 may have gone extinct. 569

570 Conclusions

571 Combining phased data with recent network methods holds much promise for 572 confronting a major challenge of plant phylogenetics: resolving the complex histories of 573 polyploids. The PATÉ pipeline can enhance systematic, speciation genomic, and population 574 genomic analyses of groups containing polyploids. While haplotype consensus sequences may 575 be adequate for resolving single reticulation events where both parents are sampled, using 576 phased sequences can improve inferences of more complicated allopolyploid events, 577 demonstrating how allelic variation can be leveraged for MSC methods that account for

578	reticulation. Still, some reticulate complexes can be difficult to disentangle with any data when					
579	there are multiple hybrid events involving the same branch. PATÉ is available through GitHub					
580	(https://github.com/gtiley/Phasing) and can be run on any UNIX environment after installing					
581	basic genotyping software and H-PoPG.					
582						
583	DATA AVAILABILITY					
584	PATÉ is freely available through GitHub at https://github.com/gtiley/Phasing. Simulated					
585	and empirical data supporting findings and files for replicating analyses are available from the					
586	Dryad Digital Repository: (X). Raw Fastq reads for Dryopteris individuals are available through					
587	the NCBI SRA database and are associated with BioProject PRJNA725004. Individual SRA					
588	Identifiers are available in Supplementary Table S1.					
589						
590	SUPPLEMENTARY MATERIAL					
591	Data available from the Dryad Digital Repository: (X).					
592						
593	ACKNOWLEDGEMENTS					
594	The authors thank A.M. Duffy, K. Imwattana, M. Nieto-Lugilde, B.T. Piatkowski, and A.J.					
595	Shaw for helpful discussions and providing feedback on the manuscript. We also thank M.G					
596	Johnson, J. Mendez Reneau, and L. Nauheimer for discussions and sharing their strategies for					
597	phasing sequence data. We are grateful to the people who made data collection possible; we					
598	used Dryopteris samples collected by C.J. Rothfels, M.A. Sundue, and W. Testo, and DNA					
599	extractions were performed by S.B. Carey and E. Lockwood.					
600						
601	FUNDING					
602	Funding was provided by National Science Foundation awards DEB-2038213 to AAC					
603	and PSM, and from DEB-1541506 to JGB and EBS. This work was also partially supported by					

- 604 the Department of Energy award DE-SC0021016 to CSL. ADY and GPT gratefully acknowledge
- 605 support from Duke University.
- 606
- 607

### 608 LITERATURE CITED

Andermann T., Cano A., Zizka A., Bacon C., Antonelli A. 2018. SECAPR-a bioinformatics

- pipeline for the rapid and user-friendly processing of targeted enriched Illumina sequences, fromraw reads to alignments. *PeerJ* 6:e5175.
- 612
- Andermann T., Fernandes A.M., Olsson U., Topel M., Pfeil B., Oxelman B., Aleixo A., Faircloth
- B.C., Antonelli A. 2019. Allele Phasing Greatly Improves the Phylogenetic Utility of
- 615 Ultraconserved Elements. *Syst. Biol.* 68:32-46. 616
- 617 Baaijens J.A., Schonhuth A. 2019. Overlap graph-based generation of haplotigs for diploids and 618 polyploids. *Bioinformatics* 35:4281-4289.
- 619
- Bankevich A., Nurk S., Antipov D., Gurevich A.A., Dvorkin M., Kulikov A.S., Lesin V.M.,
- Nikolenko S.I., Pham S., Prjibelski A.D., Pyshkin A.V., Sirotkin A.V., Vyahhi N., Tesler G.,
   Alekseyev M.A., Pevzner P.A. 2012. SPAdes: a new genome assembly algorithm and its
- 623 applications to single-cell sequencing. *J. Comput. Biol.* 19:455-477.
- 624 625 Barker M.S., Arrigo N., Baniaga A.E., Li Z., Levin D.A. 2016. On the relative abundance of
- autopolyploids and allopolyploids. *New Phytol.* 210:391-398.
- Beerli P., Mashayekhi S., Sadeghi M., Khodaei M., Shaw K. 2019. Population Genetic Inference
  With MIGRATE. *Curr. Protoc. Bioinformatics* 68:e87.
- Berger E., Yorukoglu D., Peng J., Berger B. 2014. HapTree: a novel Bayesian framework for
  single individual polyplotyping using NGS data. *PLoS Comput. Biol.* 10:e1003502.
- 634 Bezanson J., Edelman A., Karpinski S., Shah V.B. 2015. Julia: A Fresh Approach to Numerical 635 Computing. *arXiv*1411.1607v4.
- 637 Bird K.A., VanBuren R., Puzey J.R., Edger P.P. 2018. The causes and consequences of 638 subgenome dominance in hybrids and recent polyploids. *New Phytol.* 220:87-93.
- Blischak P.D., Chifman J., Wolfe A.D., Kubatko, L.S. 2018. HyDe: a Python package for
  genome-scale hybridization detection. *Syst. Biol.*, 67:821-829.
- 642

- Breinholt J.W., Carey S.B., Tiley G.P., Davis E.C., Endara L., McDaniel S.F., Neves L.G., Sessa
  E.B., von Konrat M., Chantanaorrapint S., Fawcett S., Ickert-Bond S.M., Labiak P.H., Larraín J.,
  Lehnert M., Lewis L.R., Nagalingum N.S., Patel N., Rensing S.A., Testo W., Vasco A., Villarreal
  J.C., Williams E.W., Burleigh J.G. 2021. A target enrichment probe set for resolving the
  flagellate plant tree of life. *Appl. Plant Sci.* 9:e11406.
- 648
- 649 Breinholt J.W., Earl C., Lemmon A.R., Lemmon E.M., Xiao L., Kawahara A.Y. 2018. Resolving 650 relationships among the megadiverse butterflies and moths with a novel pipeline for anchored 651 phylogenomics. *Syst. Biol.* 67:78-93.
- 652
- Buggs J.A., Wendel J.F., Doyle J.J., Soltis D.E., Soltis P.S., Coate J.E. 2014. The legacy of diploid progenitors in allopolyploid gene expression patterns. *Philos. Trans. R. Soc. Lond. B*
- 655 *Biol. Sci.* 368:20130354.
- 656

657 Cai R., Ané C. 2020. Assessing the fit of the multi-species network coalescent to multi-locus 658 data. Bioinformatics btaa863. doi: https://doi.org/10.1093/bioinformatics/btaa863 659 660 Colle M., Leisner C.P., Wai C.M., Ou S., Bird K.A., Wang J., Wisecaver J.H., Yocca A.E., Alger E.I., Tang H., Xiong Z., Callow P., Ben-Zvi G., Brodt A., Baruch K., Swale T., Shiue L., Song G.-661 Q., Childs K.L., Schilmiller A., Vorsa N., Buell C.R., VanBuren R., Jiang N., Edger P.P. 2019. 662 663 Haplotype-phased genome and evolution of phytonutrient pathways of tetraploid blueberry. 664 GigaScience 8:giz012. 665 666 Crowl A.A., Myers C., Cellinese N. 2017. Embracing discordance: Phylogenomic analyses 667 provide evidence for allopolyploidy leading to cryptic diversity in a Mediterranean Campanula 668 (Campanulaceae) clade. Evolution 71:913-922. 669 DePristo M., Banks E., Poplin R., Garimella K., Maguire J., Hartl C., Philippakis A., del Angel G., 670 Rivas M.A., Hanna M., McKenna A., Fennell T., Kernytsky A., Sivachenko A., Cibulskis K., 671 672 Gabriel S., Altschuler S., Daly M. 2011. A framework for variation discovery and genotyping 673 using next-generation DNA sequencing data. Nat. Genet. 43:491-498. 674 Edgar R.C. 2004. MUSCLE: multiple sequence alignment with high accuracy and high 675 676 throughput. Nucleic Acids Res. 32:1792-1797. 677 678 Eriksson J.S., de Sousa F., Bertrand Y.J.K., Antonelli A., Oxelman B., Pfeil B.E. 2018. Allele 679 phasing is critical to revealing a shared allopolyploid origin of Medicago arborea and M. 680 strasseri (Fabaceae). BMC Evol. Biol. 18:9. 681 682 Excoffier L., Dupanloup I., Huerta-Sanchez E., Sousa V.C., Foll M. 2013. Robust demographic 683 inference from genomic and SNP data. PLoS Genet. 9:e1003905. 684 685 Faircloth B.C. 2016. PHYLUCE is a software package for the analysis of conserved genomic 686 loci. Bioinformatics 32:786-788. 687 688 Faircloth B.C., McCormack J.E., Crawford N.G., Harvey M.G., Brumfield R.T., Glenn T.C. 2012. 689 Ultraconserved elements anchor thousands of genetic markers spanning multiple evolutionary 690 timescales. Syst. Biol. 61:717-726. 691 692 Farhat P., Hidalgo O., Robert T., Siljak-Yakovlev S., Leitch I.J., Adams R.P., Bou Dagher-693 Kharrat M. 2019. Polyploidy in the Conifer Genus Juniperus: An Unexpectedly High Rate. Front. 694 Plant Sci. 10:676. 695 696 Flouri T., Jiao X., Rannala B., Yang Z. 2020. A Bayesian Implementation of the Multispecies 697 Coalescent Model with Introgression for Phylogenomic Analysis. Mol. Biol. Evol. 37:1211-1223. 698 699 Flouri T., Jiao X., Rannala B., Yang Z. 2018. Species Tree Inference with BPP Using Genomic 700 Sequences and the Multispecies Coalescent. Mol. Biol. Evol. 35:2585-2593. 701 702 Freyman W.A,. Johnson M.G., Rothfels C.J. 2020. Homologizer: Phylogenetic phasing of gene 703 copies into polyploid subgenomes. bioRxiv doi: https://doi.org/10.1101/2020.10.22.351486. 704 705 Gronau I., Hubisz M.J., Gulko B., Danko C.G., Siepel A. 2011. Bayesian inference of ancient 706 human demography from individual genome sequences. Nat. Genet. 43:1031-1034. 707

708 He D., Saha S., Finkers R., Parida L. 2018. Efficient algorithms for polyploid haplotype phasing. 709 BMC Genomics 19:110. 710 711 Huang J., Flouri T., Yang Z. 2020. A simulation study to examine the information content in 712 phylogenomic datasets under the multispecies coalescent model. Mol. Biol. Evol. 37:3211-3224. 713 714 Huber K.T., Oxelman B., Lott M., Moulton V. 2006. Reconstructing the evolutionary history of 715 polyploids from multi-labelled trees. Mol. Biol. Evol. 23:1784-1791. 716 717 Hudson R.R. 1983. Testing the Constant-Rate Neutral Allele Model with Protein Sequence 718 Data. Evolution 37:203-217. 719 720 Huson D.H., Rupp R., Scornavacca C. 2010. Phylogenetic networks: concepts, algorithms and 721 applications. Cambridge University Press. 722 723 Igbal Z., Caccamo M., Turner I., Flicek P., McVean G. 2012. De novo assembly and genotyping 724 of variants using colored de Bruijn graphs. Nat. Genet. 44:226-232. 725 726 Jantzen J.R., Amarasinghe P., Folk R.A., Reginato M., Michelangeli F.A., Soltis D.E., Cellinese 727 N., Soltis P.S. 2020. A two-tier bioinformatic pipeline to develop probes for target capture of 728 nuclear loci with applications in Melastomataceae. Appl. Plant Sci. 8:e11345. 729 730 Johnson M.G., Gardner E.M., Liu Y., Medina R., Goffinet B., Shaw A.J., Zerega N.J., Wickett 731 N.J. 2016. HybPiper: Extracting coding sequence and introns for phylogenetics from high-732 throughput sequencing reads using target enrichment. Appl. Plant Sci. 4:1600016. 733 734 Johnson M.G., Pokorny L., Dodsworth S., Botigue L.R., Cowan R.S., Devault A., Eiserhardt 735 W.L., Epitawalage N., Forest F., Kim J.T., Leebens-Mack J.H., Leitch I.J., Maurin O., Soltis 736 D.E., Soltis P.E., Wong G.K.-S., Baker W.J., Wickett N.J. 2019. A Universal Probe Set for 737 Targeted Sequencing of 353 Nuclear Genes from Any Flowering Plant Designed Using k-738 Medoids Clustering. Syst. Biol. 68:594-606. 739 740 Jones G., Sagitov S., Oxelman B. 2013. Statistical inference of allopolyploid species networks in 741 the presence of incomplete lineage sorting. Syst. Biol. 62:467-478. 742 743 Jukes T.H., Cantor C.R. 1969. Evolution of protein molecules. In: Munro H.N., editor. 744 Mammalian Protein Metabolism. New York, NY: Acedemic Press. p. 21-132. 745 746 Kalyaanamoorthy S., Minh B.Q., Wong T.K.F., von Haeseler A., Jermiin L.S. 2017. 747 ModelFinder: fast model selection for accurate phylogenetic estimates. Nat. Methods 14:587-748 589. 749 750 Karimi N., Grover C.E., Gallagher J.P., Wendel J.F., Ané C., Baum D.A. 2020. Reticulate 751 Evolution Helps Explain Apparent Homoplasy in Floral Biology and Pollination in Baobabs 752 (Adansonia; Bombacoideae; Malvaceae). Syst. Biol. 69:462-478. 753 754 Kates H.R., Johnson M.G., Gardner E.M., Zerega N.J.C., Wickett N.J. 2018. Allele phasing has 755 minimal impact on phylogenetic reconstruction from targeted nuclear gene sequences in a case 756 study of Artocarpus. Am. J. Bot. 105:404-416. 757

758 Lemmon A.R., Emme S.A., Lemmon E.M. 2012. Anchored hybrid enrichment for massively 759 high-throughput phylogenomics. Syst. Biol. 61:727-744. 760 761 Li H., Durbin R. 2009. Fast and accurate short read alignment with Burrows-Wheeler transform. 762 *Bioinformatics* 25:1754-1760. 763 764 Liu X., Fu Y.X. 2020. Stairway Plot 2: demographic history inference with folded SNP frequency 765 spectra. Genome Biol. 21:280. 766 767 Liu Y., Johnson M.G., Cox C.J., Medina R., Devos N., Vanderpoorten A., Hedenas L., Bell N.E., 768 Shevock J.R., Aguero B., Quandt D., Wickett N.J., Shaw A.J., Goffinet B. 2019. Resolution of 769 the ordinal phylogeny of mosses using targeted exons from organellar and nuclear genomes. 770 Nat. Commun. 10:1485. 771 Lott M., Spillner A., Huber K.T., Moulton V. 2009. PADRE: a package for analyzing and 772 773 displaying reticulate evolution. *Bioinformatics* 25:1199-1200. 774 775 Luo R., Liu B., Xie Y., Li Z., Huang W., Yuan J., He G., Chen Y., Pan Q., Liu Y., Tang J., Wu G., 776 Zhang H., Shi Y., Liu Y., Yu C., Wang B., Lu Y., Han C., Cheung D.W., Yiu S.-M., Peng S., 777 Xiaogian Z., Liu G., Liao X., Li Y., Yang H., Wang J., Lam T-W., Wang J. 2012. SOAPdenovo2: 778 an empirically improved memory-efficient short-read de novo assembler. GigaScience 1:18. 779 780 McKain M.R., Johnson M.G., Uribe-Convers S., Eaton D., Yang Y. 2018. Practical 781 considerations for plant phylogenomics. Appl. Plant Sci. 6:e1038. 782 783 McKenna A., Hanna M., Banks E., Sivachenko A., Cibulskis K., Kernytsky A., Garimella K., 784 Altshuler D., Gabriel S., Daly M., DePristo M.A. 2010. The Genome Analysis Toolkit: a 785 MapReduce framework for analyzing next-generation DNA sequencing data. Genome Res. 786 20:1297-1303. 787 788 Moeinzadeh M.H., Yang J., Muzychenko E., Gallone G., Heller D., Reinert K., Haas S., Vingron 789 M. 2020. Ranbow: A fast and accurate method for polyploid haplotype reconstruction. PLoS Comput. Biol. 16:e1007843. 790 791 792 Montgomery J.D., Paulton E.M. 1981. Dryopteris in North America. Fiddlehead Forum 8:25-31. 793 794 Morales-Briones D.F., Liston A., Tank D.C. 2018. Phylogenomic analyses reveal a deep history 795 of hybridization and polyploidy in the Neotropical genus Lachemilla (Rosaceae). New Phytol. 796 218:1668-1684. 797 798 Nauheimer L., Weigner N., Joyce E., Crayn D., Clarke C., Nargar K. 2020. HybPhaser: a 799 workflow for the detection and phasing of hybrids in target capture datasets. bioRxiv. doi: 800 https://doi.org/10.1101/2020.10.27.354589. 801 802 Nguyen L.-T., Schmidt H.A., von Haeseler A., Minh B.Q. 2015. IQ-TREE: a fast and effective 803 stochastic algorithm for estimating maximum-likelihood phylogenies. Mol. Biol. Evol. 32:268-804 274. 805 806 Oberprieler C., Wagner F., Tomasello S., Konowalik K. 2017. A permutation approach for 807 inferring species networks from gene trees in polyploid complexes by minimising deep 808 coalescences. Methods in Ecology and Evolution 8:835-849.

809 810 Olave M., Meyer A. 2020. Implementing Large Genomic Single Nucleotide Polymorphism Data 811 Sets in Phylogenetic Network Reconstructions: A Case Study of Particularly Rapid Radiations of 812 Cichlid Fish. Syst. Biol. 69:848-862. 813 814 Pamilo P., Nei M. 1988. Relationships between gene trees and species trees. Mol. Biol. Evol. 815 5:568-583. 816 817 Rothfels C.J., Pryer K.M., Li F.-W. 2017. Next-generation polyploid phylogenetics: rapid 818 resolution of hybrid polyploid complexes using PacBio single-molecule sequencing. New Phytol. 819 213:413-429. 820 821 Saada O.A., Tsouris A., Freidrich A., Schachrer J. 2020. nPhase: An accurate and contiguous 822 phasing method for polyploids. *bioRxiv* doi:https://doi.org/10.1101/2020.07.24.219105. 823 824 Sessa E.B., Zimmer E.A., Givnish T.J. 2012a. Reticulate evolution on a global scale: a nuclear 825 phylogeny for New World Dryopteris (Dryopteridaceae). Mol. Phylogenet. Evol. 64:563-581. 826 827 Sessa E.B., Zimmer E.A., Givnish T.J. 2012b. Unraveling reticulate evolution in North American 828 Dryopteris (Dryopteridaceae). BMC. Evol. Biol. 12:104. 829 830 Solis-Lemus C., Ané C. 2016. Inferring Phylogenetic Networks with Maximum Pseudolikelihood 831 under Incomplete Lineage Sorting. PLoS Genet. 12:e1005896. 832 833 Solis-Lemus C., Bastide P., Ané C. 2017. PhyloNetworks: A Package for Phylogenetic 834 Networks. Mol. Biol. Evol. 34:3292-3298. 835 836 Soltis, D.E., Visger C.J., Soltis P.S. 2014. The polyploidy revolution then...and now: Stebbins 837 revisited. American Journal of Botany 101: 1057–1078. 838 839 Stull G.W., Soltis P.S., Soltis D.E., Gitzendanner M.A., Smith S.A. 2020. Nuclear phylogenomic 840 analyses of asterids conflict with plastome trees and support novel relationships among major 841 lineages. Am. J. Bot. 107:790-805. 842 843 Tiley G.P., Poelstra J.W., dos Reis M., Yang Z., Yoder A.D. 2020. Molecular Clocks without 844 Rocks: New Solutions for Old Problems. *Trends Genet.* 36:845-856. 845 846 Viruel J., Conejero M., Hidalgo O., Pokorny L., Powell R.F., Forest F., Kantar M.B., Soto Gomez 847 M., Graham S.W., Gravendeel B., Wilkin P., Leitch I.J. 2019. A Target Capture-Based Method 848 to Estimate Ploidy from Herbarium Specimens. Front. Plant Sci. 10:937. 849 850 Weiss C.L., Pais M., Cano L.M., Kamoun S., Burbano H.A. 2018. nQuire: a statistical framework 851 for ploidy estimation using next generation sequencing. BMC Bioinformatics 19:122. 852 853 Wen D., Yu Y., Nakhleh L. 2016. Bayesian Inference of Reticulate Phylogenies under the 854 Multispecies Network Coalescent. PLoS Genet. 12:e1006006. 855 856 Wen D., Yu Y., Zhu J., Nakhleh L. 2018. Inferring Phylogenetic Networks Using PhyloNet. Syst. 857 Biol. 67:735-740. 858

- Wolf P.G., Robison T.A., Johnson M.G., Sundue M.A., Testo W.L., Rothfels C.J. 2018. Target
  sequence capture of nuclear-encoded genes for phylogenetic analysis in ferns. *Appl. Plant Sci.*6:e01148.
- 862
- Wood T.E., Takebayashi N., Barker M.S., Mayrose I., Greenspoon P.B., Rieseberg L.H. 2009.
  The frequency of polyplid speciation in vascular plants. *Proc. Natl. Acad. Sci. U. S. A.*106:13875-13879.
- 865 866
- Xie M., Wu Q., Wang J., Jiang T. 2016. H-PoP and H-PoPG: heuristic partitioning algorithms for single individual haplotyping of polyploids. *Bioinformatics* 32:3735-3744.
- 869
- Xie W., Lewis P.O., Fan Y., Kuo L., Chen M.H. 2011. Improving marginal likelihood estimation
   for Bayesian phylogenetic model selection. *Syst. Biol.* 60:150-160.
- Yang J., Moeinzadeh M-H., Kuhl H., Helmuth J., Xiao P., Haas S., Liu G., Zheng J., Sun Z., Fan
  W., Deng G., Wang H., Hu F., Zhao S., Fernie A.R., Boerno S., Timmermann B., Zhang P.,
  Vingron M. 2017. Haplotype-resolved sweet potato genome traces back its hexaploidization
  history. *Nat Plants* 3:696-703.
- 878 Yang Z. 2006. Computational molecular evolution: Oxford University Press.
- Zhang C., Ogilvie H.A., Drummond A.J., Stadler T. 2018. Bayesian Inference of Species
   Networks from Multilocus Sequence Data. *Mol. Biol. Evol.* 35:504-517.
- 882

877

- Zhang C., Rabiee M., Sayyari E., Mirarab S. 2018. ASTRAL-III: polynomial time species tree
   reconstruction from partially resolved gene trees. *BMC Bioinformatics* 19:153.
- 885886 Zhu S., Degnan J.H. 2017. Displayed trees do not determine distinguishability under the
- 887 network multispecies coalescent. *Syst. Biol.* 66:283-298.
- 888

# 889 TABLES

Topology	Phased Marginal InL	Unphased Marginal InL	Phased Model Probability	Unphased Model Probability	
1	-613405.3154	-552720.6776	1.006E-278	3.2574E-214	
2	-613223.0755	-552534.6540	1.4073E-199	2.0039E-133	
3	-613366.4128	-552644.7603	7.9026E-262	3.0431E-181	
4	-612866.6368	-552245.2218	8.86633E-45	1.00164E-07	
5	-612828.7348	-552243.7091	2.56082E-28	4.54611E-07	
6	-612822.4509	-552238.6462	1.37227E-25	7.18535E-05	
7†	-612765.2027	-552229.1054	1	0.999927592	
8	-613211.8583	-552484.8320	1.047E-194	8.6956E-112	
9	-613168.7562	-552479.3732	5.4825E-176	2.0419E-109	
10	-613102.1509	-552454.2002	4.6266E-147	1.74796E-98	
11	-613079.9316	-552426.8867	2.0654E-137	1.27244E-86	
12	-613379.6350	-552538.6937	1.4303E-267	3.5276E-135	
13	-613390.9585	-552533.6798	1.7287E-272	5.3085E-133	
14	-613272.7561	-552561.3165	3.7359E-221	5.2787E-145	
15	-613247.7503	-552566.6741	2.7054E-210	2.4873E-147	

890

891 **Table 1 — Marginal Likelihoods for Possible Topological Hypotheses.** Topologies of the 15

892 models are displayed in Figure S1.

893 <sup>†</sup>*a priori* allopolyploid hypothesis

### 895 FIGURES





897 Figure 1 — Hypothesized Relationships among North American Dryopteris. Synthesis of 898 results from Sessa et al. 2012a and Sessa et al. 2012b. A) Links between shapes show the 899 putative parents and their allopolyploid derivatives. Black circles are diploids, squares are 900 tetraploids, and the hexagon is the one hexaploid species in the group. Dryopteris semicristata 901 is presumed extinct. B) Placement of allopolyploids in the context of the backbone relationships 902 among diploids. Tetraploids are indicated with solid orange lines and the hexaploid with dotted 903 lines. The grey line denoting a sister relationship between D. semicristata and D. expansa reflects one possible placement for the extinct taxon based on previous analyses (Sessa et al., 904 905 2012b).



**Figure 2** — **Species network used for simulation.** The divergence times in expected substitutions per site are given for each node, and *h* is the hybrid node where two alleles enter from both *t* and *v*. E is an allotetraploid while other species are diploid. Nucleotide divergence was reduced by dividing all  $\tau$  by 10 or increased by multiplying all  $\tau$  by 10. ILS was increased by halving the distance between  $\tau_h$  and  $\tau_u$  and between  $\tau_u$  and  $\tau_s$  either once (for the medium ILS condition) or twice (for the high ILS condition).

Input data	Flow cytometry Genomic data	HybPiper	Assemble reads into consensus loci	bwa	Realign reads to consensus loci	gatk	Ploidy-aware variant calling	H-PoPG + custom	Phase alleles	Downstream analyses
	[Ploidy = 4x] TGGCG TGGGGCA GCGCA GGGCA TGGCGCA		>L7141 TGGGGCA		>L7141 TGGGGCA TGGCG TGGGGCA GCGCA GGGGCA TGGGGCA		genotype files .bam .vcf	scripts	<pre>&gt;L7141_1 TGGGGCAGMCTY &gt;L7141_2 TGGGGCAGMCTY &gt;L7141_3 TGGCGCAGMCTY &gt;L7141_4 TGGCGCAGMCTY</pre>	

916 **Figure 3 — PATÉ Phasing Pipeline.** Overview of data input, output, and steps taken to phase

917 alleles. Input data are assumed to be paired-end Illumina reads and reference sequences for

918 each individual are required (consensus loci from HybPiper can be used). The ploidy of each

919 individual must be specified. Only biallelic sites are used for phasing.

920



921

Figure 4 — Proportion of simulations that correctly identify the allopolyploid lineage. The
x-axes are the number of loci sampled for each simulation. Th y-axes are the proportions of
correct networks. Results are based on networks estimated with a single reticulation, even if
that network was considered less optimal than networks with zero or two reticulations. We
saved the true gene trees from the simulations, while we estimated gene trees with the phased
and unphased (Genotype, Consensus, and Pick One) data.



929

Figure 5 — Estimating the Timing of Introgression. Divergence times are for node *h* in Figure 2. The low divergence simulation corresponds to the y-axis units of  $\tau_h \times 10^{-2}$  while the high divergence case is represented by  $\tau_h$ . Divergence times are measured as the expected number of substitutions per site. The dashed line represents the true simulated values. Points are posterior means, and error bars are 95% HPD intervals, averaged across 30 replicates.



936

Figure 6— *Dryopteris* Divergence Times under the MSci Model. Divergence times are
measured as the expected number of substitutions per site. The x-axis shows estimates from
phased data and the y-axis shows estimates from unphased (haplotype consensus) data for the
inset network (equivalent to model 7 from Supplementary Fig. S1). The dashed one-to-one line
shows where older age estimates are consistently obtained from phased data. Error bars on
points show the 95% HPD intervals for phased and unphased data.



945 Figure 7— Probabilities for three-taxon Dryopteris MSci Models with fewer Loci. Marginal 946 likelihoods were estimated for the 15 MSci models (Supplementary Fig. S1). Model weights 947 were used to obtain probabilities. We consider model probability greater than 0.95 as decisive 948 evidence in favor of a model, and a probability less than 0.05 is evidence against a model. We 949 considered probabilities between 0.05 and 0.95 to be ambiguous. Models four through seven (in 950 bold) all have the correct reticulate relationships between the parental diploid lineages and the 951 allopolyploid. Models one through three do not have introgression, and models eight through 15 952 have incorrect introgression events. Each row represents one of 30 sampling replicates.



954 Figure 8 — Network search results for three-taxon Dryopteris example. Both phased and 955 unphased haplotype consensus data recovered the same network topology with introgression 956 occurring in the expected direction. The major topology is indicated by the blue edge and the 957 minor edge (direction of introgression) is shown in tan. The inheritance probability  $\gamma$  was slightly 958 higher in the haplotype consensus data ( $\gamma_{unphased} = 0.455$ ). Bar plots show the proportion of 959 100 replicates when sampling four or 40 loci that correctly detect one reticulation based on the 960 pseudolikelihood scores (top), correctly estimate the network with introgression going from one 961 of the diploids into D. campyloptera (middle), and estimate a network where the direction of 962 introgression is from *D. campyloptera* into one of the diploids (bottom).

963

953



964

965 Figure 9 — Networks for Nine-Taxon Dryopteris example. Both data types recovered 966 optimal networks with three reticulation events. The major topology edge is blue and the minor 967 (reticulation) edges shown in tan, with the direction of introgression flowing into the major edge. 968 The position of *D. carthusiana* changes in the major topology between phased and unphased 969 data but the relationships are otherwise the same. All three reticulation events in the phased 970 data are plausible, but in the unphased data, the direction of introgression from D. carthusiana 971 into D. intermedia is incorrect and the reticulate edge from the common ancestor of Dryopteris is 972 difficult to reconcile. Inheritance probabilities for each introgression event are shown next to 973 reticulation edges.