- 1 sppIDer: a species identification tool to investigate hybrid genomes with high-throughput
- 2 sequencing
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- 11 Abstract:

12 The genomics era has expanded our knowledge about the diversity of the living world, 13 yet harnessing high-throughput sequencing data to investigate alternative evolutionary 14 trajectories, such as hybridization, is still challenging. Here we present sppIDer, a pipeline for 15 the characterization of interspecies hybrids and pure species, that illuminates the complete 16 composition of genomes. sppIDer maps short-read sequencing data to a combination genome 17 built from reference genomes of several species of interest and assesses the genomic contribution 18 and relative ploidy of each parental species, producing a series of colorful graphical outputs 19 ready for publication. As a proof-of-concept, we use the genus Saccharomyces to detect and 20 visualize both interspecies hybrids and pure strains, even with missing parental reference 21 genomes. Through simulation, we show that sppIDer is robust to variable reference genome 22 qualities and performs well with low-coverage data. We further demonstrate the power of this 23 approach in plants, animals, and other fungi. sppIDer is robust to many different inputs and provides visually intuitive insight into genome composition that enables the rapid identification 24 25 of species and their interspecies hybrids. sppIDer exists as a Docker image, which is a reusable, 26 reproducible, transparent, and simple-to-run package that automates the pipeline and installation 27 of the required dependencies (https://github.com/GLBRC/sppIDer).

28

29 Introduction:

Interspecies hybrids play a large role in both natural and in industrial settings (Dunn and
Sherlock 2008; Soltis et al. 2015; Payseur and Rieseberg 2016; Peris et al. 2017c). However,

32 identification and characterization of the genomic contributions of hybrids can be difficult. High-33 throughput sequencing can be used to address many of the barriers to identifying and 34 characterizing hybrids. With the influx of sequencing data, the quality and number of reference genomes available is increasing at a rapid pace. Population genomic, ecological diversity, and 35 36 gene expression projects are underway in many fields. These studies are yielding a high volume 37 of short-read data, but determining the best way to leverage these data can be challenging. A key goal of the modern genomic era is to be able to integrate and synthesize these data to further our 38 39 understanding of natural diversity (Richards 2017), including addressing key questions about the 40 frequency and genomic identities of hybrid and admixed lineages in the wild.

The number of reference genomes available has rapidly increased, but it is not complete 41 42 in most clades. To avoid the drawbacks of limited reference genomes, several new phylogenetic 43 approaches have been developed that do not require sequence alignments or whole-genome 44 assemblies, such as phylogeny-building approaches using kmers (Fan et al. 2015), de novo 45 identification of phylogenetically informative regions (Schwartz et al. 2015), and local 46 assemblies of target genes (Allen et al. 2015; Johnson et al. 2016). These methods can accurately 47 reconstruct known and simulated phylogenies of pure lineages. However, these methods have not been tested on hybrid or admixed lineages. As hybrids are the result of an outcrossing event 48 49 between two independently evolving lineages, their origin is inherently not tree-like. Therefore, 50 placing hybrids on a bifurcating tree will not reflect the topology observed with pure lineages. Placing hybrids on a phylogenetic network is more apt, but it is still untested with alignment-free 51 52 phylogenetic approaches. Other species identification methods based on local assembly of target 53 genes could lead to erroneous identification, depending on which parent the gene of interest is 54 retained from in the hybrid, or could lead to the assembly of a chimeric gene if the hybrid has 55 retained copies from multiple parents. Therefore, in organisms with alternative evolutionary 56 trajectories, such as hybrids with complex genomes, applying alignment-free phylogenetic 57 methods is difficult and could potentially result in imprecise conclusions.

58 Other methods to detect interspecies hybrids have been adapted from methods developed 59 for intraspecies diversity, such as F_{ST}, STRUCTURE analysis, phylogenetic discordance, linkage 60 disequilibrium, and PCA approaches (Payseur and Rieseberg 2016). There are numerous 61 drawbacks to using these methods to detect interspecies hybrids. For example, most definitions 62 of speciation require the cessation of gene flow and the accumulation of sequence divergence well beyond the levels observed between populations, which are therefore beyond the
expectations of most of these approaches. Many of these methods were also developed for
diploid obligately outcrossing species, which makes problematic their application to
allopolyploids or species that primarily undergo selfing or other forms of inbreeding. Indeed, the
basic assumptions of these methods, including gene flow, demographic history, and natural
selection, are violated by most interspecies hybrids.

69 Here we present sppIDer as a novel, assumption-free method that rapidly provides visual 70 and intuitive insight into ancestry genome-wide, which will aid in the discovery and 71 characterization of interspecies hybrids. This method maps short-read data to combination 72 genome, built from available reference genomes chosen by the user. sppIDer allows for the 73 analysis and visualization of the genomic makeup of a single organism of interest, facilitating the 74 rapid discovery of hybrids and individuals with other unique genomic features, such as aneuploidies and introgressions. Therefore, sppIDer is an unbiased method that provides unique 75 76 and intuitive insights into complex genomic ancestry and regions of differing evolutionary 77 history, which can complement existing methods in the characterization of hybrids.

78

79 New Approaches

80 Here we describe and make available a user-friendly short-read data analysis pipeline that 81 utilizes existing bioinformatic tools and custom scripts to determine species identity, hybrid 82 status, and chromosomal copy-number variants (CCNVs). Short-reads are mapped to a 83 combination reference genome of multiple species of interest, and the output is parsed for where, how well, and how deeply the reads map across this combination genome. A colorful automated 84 85 output allows end-users to rapidly and intuitively assess the genomic contribution, either from a 86 single species or multiple species, and relative ploidy of an organism. Figure 1 illustrates the 87 basic workflow in a flow chart of each step. An upstream step creates a combination reference 88 genome, which is a concatenation of reference genomes of interest, before the main pipeline is 89 run. The main pipeline starts with mapping short-read data to this combination reference 90 genome. Then, this output is parsed for percentage and quality of reads that map to each 91 individual reference genome within the combination reference and percentage of unmapped 92 reads; this summary is then plotted so these metrics can be visualized. In parallel, the mapping 93 output is analyzed for depth of coverage. Reads with a mapping quality (MQ) greater than three

are retained and sorted into the combination reference genome order; then, coverage across the
combination reference genome is computed. A custom script then calculates the mean coverage
for each species, and the combination reference genome broken into windows. The output of
these analyses is then plotted so that coverage across the combination reference genome can be
visualized.

99 We have given this computational pipeline and wrapper a portmanteau of the pluralized 100 abbreviation of species (spp.) and identifier (IDer), to reflect its ability to identify hybrids of multiple species. sppIDer also detects CCNVs, such as those caused by aneuploidy and other 101 102 genomic changes that do not meet the textbook definition of an euploidy, including interspecies 103 loss-of-heterozygosity events, interspecies unbalanced translocations, and other differences in 104 relative ploidy. sppIDer is provided as an open source Docker (http://www.docker.com), which 105 organizes the pipeline and all the dependencies into a reusable, reproducible, transparent, and 106 simple-to-run package (https://github.com/GLBRC/sppIDer).

107 Here we present several applications of sppIDer in yeast, plant, and animal genomes. 108 Through simulations, we show that sppIDer can detect hybrids of closely or distantly related 109 species, and of recent or ancient origin. We use the genus Saccharomyces to 1) detect both interspecies hybrids and pure strains; 2) detect hybrids, even with missing reference genomes; 110 111 and 3) determine how divergent lineages and poor-quality data and reference genomes affect 112 sppIDer's performance. Next, we test sppIDer's utility in non-Saccharomyces systems: another 113 yeast genus, Lachancea; an animal genus, Drosophila; and a plant genus, Arabidopsis. Finally, 114 we test an extension for non-nuclear DNA using mitochondrial genome data. Overall, sppIDer is 115 robust to many different inputs and can be used across organisms to provide rapid insight into the 116 species identity, hybrid status, and CCNVs of an organism.

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118 **Results and Discussion:**

119 Species and interspecies hybrid identifications:

To test sppIDer, we first used the well-studied genus *Saccharomyces* (Hittinger 2013).
Seven of the eight species have reference genomes scaffolded at a near-chromosomal level, and
there are many interspecies hybrids (Goffeau et al. 1996; Fischer et al. 2000; Dunn and Sherlock
2008; Liti and Carter et al. 2009; Scannell and Zill et al. 2011; Liti et al. 2013; Baker et al. 2015;
Naseeb et al. 2017; Peris et al. 2017c). To test sppIDer's species-level classification ability for a

125 natural isolate, we used the short-read data available for a Saccharomyces eubayanus strain 126 isolated in New Zealand (P1C1) (Gayevskiy and Goddard 2016). The reads from this wild S. 127 *eubayanus* strain mapped preferentially to the S. *eubayanus* reference genome (Figure 2a), as seen by normalized coverage only being above zero for the S. eubayanus genome. This strain 128 129 belongs to the same diverse lineage as the reference strain (Peris and Langdon et al. 2016), but as 130 the first isolate from Oceania, these results show that sppIDer can easily classify, to the species 131 level, a divergent wild strain isolated from a novel environment. To test sppIDer's utility for 132 industrial strains, we used short reads from an ale strain, Fosters O (Gonçalves et al. 2016). This 133 test shows that this brewing strain is a pure species; the S. cerevisiae genome is the only genome 134 that had normalized coverage above zero. However, normalized coverage differed within the S. 135 *cerevisiae* genome (Figure 2b & Figure S1a), implying an euploidies. Coverage was lower for 136 chromosomes VII and XIV and increased for chromosome XIII, in comparison with the genome-137 wide average coverage, indicating that there are more copies of chromosome XIII and fewer 138 copies of chromosome VIII and XIV. Additionally, we could detect regions of CCNV within a 139 chromosome, such as the small region within chromosome VII where the normalized coverage 140 returned to the genome average.

141 To test sppIDer's ability to delineate hybrids, we used short-read data from two S. 142 cerevisiae X S. eubayanus lager yeast lineages, Saaz (strain CBS1503) and Frohberg (strain 143 W34/70). These results recapitulated the known relative ploidy and rearrangements, where 144 ploidy differs both within and between genomes. Specifically, the Frohberg lineage contains 145 approximately two copies of each chromosome from both S. cerevisiae and S. eubayanus. Thus, 146 what was observed matched this expectation, where the average normalized coverage across both 147 the S. cerevisiae and S. eubayanus genomes were approximately at the same level, but there were 148 clear fluctuations, indicating ploidy changes (Figure 2c & Figure S1b). In our test with a 149 representative of the Saaz lineage, we observed that the S. cerevisiae genome had an average 150 normalized coverage of ~0.5, that fluctuated from none to two, and the S. eubayanus genome had 151 an average normalized coverage of 1.5, that fluctuated from zero to three (Figure 2d & Figure 152 S1c). These results match with previous observations that the Saaz lineage is approximately 153 haploid for the S. cerevisiae genome and diploid for the S. eubayanus genome. Additionally, 154 from the sppIDer plots, we also easily inferred the previously described aneuploidies and 155 translocations (Figure 2c-d) (Dunn and Sherlock 2008; Okuno et al. 2016).

156 As an additional hybrid test, we used short-read data from the wine strain Vin7, a S. 157 cerevisiae X Saccharomyces kudriavzevii hybrid. From the normalized coverage plot (Figure 158 2e), we could determine that Vin7 has retained complete copies of both parental genomes, but at 159 different ploidy levels. Specifically, the normalized coverage for S. cerevisiae was around two 160 across the genome, while the normalized coverage for S. kudriavzevii was consistently around 161 one across the genome. Here we could infer that this strain has double the number of copies of S. 162 cerevisiae chromosomes as it does of S. kudriavzevii chromosomes. Although exact ploidy 163 cannot be measured without direct measures of DNA content, the inferred ploidy is consistent 164 with previous studies (Borneman et al. 2012; Peris et al. 2012; Borneman et al. 2016). 165 As a final test of interspecies hybrids, we used data from the cider strain CBS2834 (Almeida et al. 2014). Here sppIDer detected large genetic contributions from S. cerevisiae, S. 166 167 kudriavzevii, and Saccharomyces uvarum, as well as introgressed contributions from S. 168 eubayanus (Figure 2f & Figure S1d). Although the S. eubayanus genetic contribution is quite

small, seen on chromosomes XII and XIV, it was still easily detected by sppIDer. These
examples show that sppIDer can easily detect higher-order interspecies hybrids, even those with
minor contributions from several species.

172

173 <u>Testing the limits of sppIDer with a simulated phylogeny:</u>

174 To test sppIDer's performance with hybrids of varying levels of parental divergence, we 175 used a simulated phylogeny. To build this phylogeny we started with the S. cerevisiae reference 176 genome and produced a phylogeny of 10 species through several rounds of simulating short-read 177 sequencing data, applying a set mutation rate, and assembling those reads. For these simulated 178 genomes, sister species were $\sim 4\%$ diverged, and the most distantly related species were $\sim 20\%$ 179 diverged (Figure 3a). This simulated phylogeny allowed us to test pseudo-hybrids from closely 180 and distantly related lineages. Further, the iterative process of phylogeny building allowed us to 181 create ancient pseudo-hybrids that simulated the result from hybridization of a common ancestor 182 predating a lineage split. sppIDer accurately mapped pure lineages to their corresponding 183 reference genome (Figure 3b). For all 10 species, >90% of the reads mapped to their 184 corresponding reference genome. The read simulation and assembly process resulted in varying 185 quality final references, but despite differences in genome quality, all reads still mapped 186 accurately and were not biased to the best reference genome.

187 To determine sppIDer's applicability to hybrids of both closely and distantly related 188 parents and of recent and ancient origin, we tested sppIDer with pseudo-hybrids of different 189 combinations of simulated species. sppIDer accurately detected all true hybrid parents. When 190 pseudo-hybrids were between sister species, <0.01% of the reads mapped promiscuously to other 191 species (Figure 3c). When we used more divergent pseudo-hybrids, sppIDer still detected the 192 true parents, with <5% of the reads mapped promiscuously to the sister species (Figure 3d). 193 Additionally, we simulated ancient pseudo-hybrids, between common ancestors before lineage 194 splits, and found that sppIDer mapped the reads of these hybrids to the references of the lineages 195 that descended from the ancestors that hybridized (Figure 3e). With complete knowledge of this 196 simulated phylogeny, we were able to test many different potential hybrid arrangements and 197 found that sppIDer detected the true parents of all hybrids.

198 Finally, we tested a scenario, which is common in biology, of incomplete knowledge of 199 the clade of interest. This dearth could due to many variables, such as a described species lacking 200 a reference genome or a species being unknown to science altogether. To test the effect of 201 missing a species, we removed one species' reference genome from the combination reference 202 genome, then mapped pure lineage and pseudo-hybrid reads to this permuted genome. With 203 reads of a simulated pseudo-hybrid of sister species, G and H, we observed that, when one parent 204 genome was missing, the reads mapped primarily to the reference genome of the remaining 205 parent, reference H, with slightly increased promiscuous mapping of reads to the next-closest 206 clade, references I and J (Figure 3f). Therefore, with incomplete reference genome knowledge, 207 detecting hybrids of closely related species is limited. However, we could still detect hybrids of 208 more distantly related species, such as a pseudo-hybrid of E and G and a pseudo-hybrid of the 209 common ancestor of A and the common ancestor of G and H (Figure S2), though our inference of parentage was biased by the availability of reference genomes. Therefore, with incomplete 210 211 knowledge of reference genomes, hybrid detection is limited, and the inference of true parentage 212 can suffer in specific cases, but generally, distant and ancient hybrids can be detected.

213

214 <u>Hybrid detection with missing reference genomes:</u>

To empirically address how sppIDer would be affected by missing reference genomes, such as for hybrids whose parents are themselves unknown (Hoot et al. 2004; Pryszcz et al. 2014), we focused again on the genus *Saccharomyces*. Specifically, we used the *S. cerevisiae* X S. kudriavzevii hybrid (Vin7) and the S. cerevisiae X S. eubayanus Frohberg lager yeast
(W34/70) as examples. We tested the performance of sppIDer on short-read data from both
hybrids by removing the S. cerevisiae reference genome and, in a separate test, removing the
reference genome of the other parent. Our expectation was that reads would map to the genome
of the sister species, if it were available, or that they would fail to map or be distributed across
other genomes, if there were no close relatives.

224 When we removed the S. eubayanus reference genome for the lager example, the 225 proportion of reads that failed to map increased, as did those reads that mapped to S. uvarum, its 226 sister species (~93% identical in DNA sequence, Libkind and Hittinger et al. 2011), albeit with a 227 decreased mapping quality (MQ) (Figure 4c). We then tested sppIDer on Vin7 and W34/70 when 228 the S. cerevisiae reference genome was removed (Figure 4a & d). In both examples, the 229 proportion of reads that mapped to Saccharomyces paradoxus, S. cerevisiae's sister species 230 (~87% identical in DNA sequence), increased (Figure 4a & d). Thus, the absence of a reference 231 genome for one of the parents of a hybrid led to increased mapping to its sister species, instead. 232 We also tested removing the S. kudriavzevii reference genome for Vin7. Since there is not a 233 sister species closely related to S. kudriavzevii, the number of unmapped reads increased, and the 234 remaining reads mapped to the reference genomes other species of the genus in approximately 235 equal proportions (Figure 4f).

236 From these tests, we would have easily inferred that W34/70 was a hybrid, regardless of 237 whether either parent genome was withheld (the actual state of affairs for *S. eubayanus* before 238 Libkind and Hittinger et al. 2011). Using the coverage plots, we were still even able to infer the 239 same CCNVs for W34/70 that we observed with the full suite of reference genomes. With Vin7, 240 we still easily inferred its hybrid status without including the S. cerevisiae genome. Without the S. kudriavzevii reference genome, Vin7 produced an unusually high number of unmapped reads 241 242 without a decrease in mapping quality to S. cerevisiae, a result that should spur the investigator 243 to perform more detailed analyses to search for evidence of contributions by an unknown 244 species, such as de novo genome assembly and phylogenetics. Therefore, even without a full 245 complement of reference genomes, sppIDer can still be useful for rapid inference of interspecies 246 hybrids.

247

248 Hybrid detection with simulated low-quality reference genomes:

249 To test a scenario where not all of the reference genomes are ideal, we used iWGS (Zhou 250 et al. 2016) to independently simulate reads and then assemble de novo genomes for S. 251 cerevisiae, S. kudriavzevii, S. uvarum, and S. eubayanus. These simulations resulted in reference 252 genomes with many more scaffolds and with a lower N50 than the published genomes (Table 253 S1). These low-quality genomes were independently swapped for the high-quality references in 254 the combination reference genome and tested with short-read data. We started by testing 255 simulated pseudo-lager short reads where we expected reads to map both to the S. cerevisiae and 256 S. eubayanus reference genomes. Whether we swapped in the low-quality S. cerevisiae reference 257 (Figure S3a) or the low-quality *S. eubayanus* reference (Figure S3b), the reads still mapped 258 equally to the references that were used to simulate the reads with minimal promiscuously 259 mapped reads to their sister species reference genomes.

260 We next tested the limits of sppIDer with the empirical data for CBS2834 because it has the most complex arrangement of contributions from four species. Tests with each simulated 261 262 low-quality reference genome independently showed that we could indeed recapitulate the same 263 inference of ancestry and that roughly the same proportion of reads mapped to each reference 264 genome (Figure S4) as with high-quality reference genomes (Figure S1d). Here, the inference of 265 approximate ploidy became more difficult, and visually interpreting translocations between 266 species was impossible. When both high-quality S. cerevisiae and S. kudriavzevii reference 267 genomes were used, we could infer translocations between these two genomes on chromosomes IV, X, and XV due to mid-chromosome ploidy changes that are compensated for in the other 268 269 genome. There were more promiscuously mapped reads to the high-quality reference genomes of the sister species, but not at the same level as mapped to the true parent reference genomes. 270 271 These tests with simulated low-quality de novo genomes showed that, both with simulated and 272 empirical data, proper hybrid genome contributions can still be identified, and ploidy shifts still 273 detected, despite the poor-quality reference genomes, but the inference of translocations and 274 ploidies of specific chromosomes becomes difficult.

275

276 <u>Hybrid detection with low-coverage and long-read data</u>:

To further explore the power of sppIDer, we wanted to test how little coverage was
needed to still detect the proper ancestry (Figure S5). Using data simulated at varying coverages,
we found that only 0.5X coverage was needed to recover the true ancestry for a single species

(Figure S5a-b), single species with aneuploidies (Figure S5c-d), and interspecies hybrids (Figure S5e-f). We also tested empirical data by down-sampling the FASTQ files of CBS2834 and found that we could still detect contributions from the four species at as low as ~0.05X coverage
(Figure S5g), but we lost the ability to infer ploidy at around ~0.5X coverage (Figure S5h).
These low coverage tests show how powerful sppIDer is, even with scant data, which could be a boon in many systems with large genomes or when sequencing resources are limited.

We also tested sppIDer with simulated PacBio long-read data from the *S. cerevisiae* genome and a hybrid pseudo-lager genome with equal contributions from the *S. cerevisiae* and *S. eubayanus* reference genomes. We found that we could still easily determine the species contribution for each (Figure S6), suggesting sppIDer's utility will continue if long-read technologies eventually supplant short-read sequencing technologies.

291

292 <u>Divergent lineages and poor-quality data:</u>

293 Since sppIDer relies on reference genomes, we recognized that it might be biased in its 294 ability to work with lineages that were highly divergent from the reference genome, as might be 295 the case in many systems. We tested this scenario with an example from S. paradoxus, one of the 296 most diverse *Saccharomyces* species (Liti and Carter et al. 2009; Leducq et al. 2016). Compared 297 to a representative of the reference genome's lineage, fewer reads from the divergent lineage 298 (~96% identical) mapped and with poorer quality (Figure S7a-b). We also tested this effect in S. 299 kudriavzevii using poor-quality data (36-bp reads from a first-generation Illumina Genome 300 Analyzer run by Hittinger et al. 2010) and found qualitatively similar results, but many more unmapped reads. Thus, while divergence from the reference genome affected map-ability, 301 302 sppIDer still worked generally as expected. However, when mapping percentage and quality 303 decline substantially, such as seen in these test cases, sppIDer can provide an early indication 304 that the organism may be highly divergent from the reference genome, which may merit further 305 investigation.

306

307 <u>Comparison to alignment-free phylogenetic methods:</u>

Alignment and assembly (AA)-free phylogeny-building methods are gaining popularity, but they have not previously been applied to hybrid data. Therefore, we also tested how AA-free phylogenetic methods, such as AAF (Fan et al. 2015) or SISRS (Schwartz et al. 2015),

311 performed in detecting and visualizing hybrids compared to sppIDer. We found that these 312 methods performed well when given only pure lineages, but when hybrids were included, they 313 either failed completely or produced incorrect phylogenies. We tested both our simulated 314 phylogeny and empirical Saccharomyces data. For the simulated data, both AAF and SISRS 315 produced the correct phylogeny when given the 10 simulated species. However, when given any 316 hybrid data, AAF failed to produce the correct phylogeny and instead clustered the hybrid with 317 its parents, while SISRS failed to complete at all. With the empirical data, we saw similar results; 318 with AAF, we could recapitulate the phylogeny of the genus *Saccharomyces* when using only 319 pure samples, but when we included any hybrid, an incorrect phylogeny was produced (Figure 320 S8a and c). SISRS had similar issues with producing the correct phylogeny with hybrids, but its 321 output allowed for more nuanced network visualizations. For CBS2834, the SISRS output 322 allowed us to infer the shared background with S. cerevisiae, S. kudriavzevii, S. uvarum, and S. 323 eubayanus (Figure S8d), but the proportion of contribution from each species was difficult to estimate compared to the sppIDer output. Overall, we found that these methods have serious 324 325 limitations when used with hybrids, but they could be used as a complement to sppIDer to make 326 inferences about pure parental lineages.

327 Methods that assemble targeted genes from short read-data, such as aTRAM and 328 HybPiper, can be used with poor-quality references and/or references that may be missing genes 329 of interest. We tested these tools with a panel of loci that can be used to delineate the S. 330 eubayanus populations (Peris and Langdon et al. 2016). HybPiper and aTRAM were able to 331 match short-reads to a locus of interest 59% or 34% of the time, respectively, but they could only assemble these reads 23% or 28% of the time, respectively. Neither method could assemble one 332 333 locus for all 15 strains tested, including both hybrids and non-hybrids (Table S1). While these 334 methods can be powerful when applied in a targeted manner to pure strains, they fail when 335 applied to hybrid data.

336

337 <u>Non-Saccharomyces examples</u>

338 *Lachancea*: refining the interpretation of voucher specimens

With the publication of 10 high-quality *Lachancea* genome sequences (Vakirlis et al.

340 2016) and another two recently-described and fully-sequenced species (González et al. 2013;

Freel et al. 2015; Sarilar et al. 2015; Freel et al. 2016), this genus is becoming a powerful yeast

342 model. As molecular techniques improve, initial identifications in culture and museum 343 collections can yield new interpretations. For example, the strain CBS6924 was initially 344 identified as Lachancea thermotolerans, but recent evidence suggested it as a candidate for a 345 novel species (Lachancea fantastica nom. nud. Vakirlis et al. 2016). Its closest relative, 346 Lachancea lanzarotensis, was also recently described (González et al. 2013). To test sppIDer's 347 utility for determining whether a strain or voucher specimen is or is not properly classified, we 348 tested mapping reads from CBS6924 to a combination genome with all Lachancea reference 349 genomes (Figure S9a), then removing the 'L. fantastica' reference genome (Figure S9b), and then 350 removing both the 'L. fantastica' and L. lanzarotensis reference genomes (Figure S9c). When 351 both reference genomes were removed, the reads were spread across many genomes, and the 352 initial classification of the strain as L. thermotolerans would have been easily falsified. By 353 including the L. lanzarotensis reference genome, most reads mapped to that reference, but still 354 poorly enough to warrant additional investigation. When the 'L. fantastica' reference genome was 355 included, CBS6924 reads mapped unambiguously to this reference. These results demonstrate 356 sppIDer's utility outside of the genus *Saccharomyces* to aid in reclassifying provisional species 357 identifications of voucher specimens from culture and museum collections.

358

359 Drosophila

360 To test sppIDer with larger genomes, we examined the animal genus *Drosophila*, which is a large genus with many available reference genomes (Adams et al. 2000; Drosophila 12 361 362 Genomes Consortium 2007; Alekseyenko et al. 2013; Sanchez-Flores et al. 2016), as well as 363 several species still lacking reference genomes. The difference in reference genome qualities led 364 us to remove contigs <10Kb from our combination reference genome; this paring down sped up 365 computation time, reduced memory usage, and improved the visualization, but otherwise did not 366 affect the results (Figure S10a). To test the ability of sppIDer to distinguish closely related species, we started with the Drosophila yakuba species complex (Turissini et al. 2015), where D. 367 368 yakuba has a sequenced reference genome available, but its close relative Drosophila santomae 369 and more distant relative Drosophila teissieri do not. Here we observed that short reads from a 370 D. yakuba (Comeault et al. 2016) representative mapped well to the D. yakuba reference 371 genome. As we moved from the close relative *D*. santomae (Figure 5b) to a more distant one, *D*. 372 *teissieri* (Figure 5c), the mapping percentage and quality decreased with increased promiscuous

mapping to other relatives (Figure 5a-c). Thus, as in yeasts, sppIDer can classify pure speciesand their close relatives well and provide insight to guide downstream analyses.

We also used *Drosophila* short-read data to test sppIDer's ability to detect hybrids in non-fungal systems. In this case, we used genomic data from a pure parent and RNA-seq data from a F_1 interspecies hybrid (Coolon et al. 2014). We found that sppIDer could easily detect hybrids in an animal model, but as expected, detection of CCNVs using RNA-seq was not possible (Figure S11).

380

381 Arabidopsis

382 The study of hybrid speciation and allopolyploidy in plants has a long history (Rieseberg 383 1997; Soltis et al. 2015), and we choose Arabidopsis as our plant test case because it has 384 reference genomes available for Arabidopsis halleri, Arabidopsis thaliana, and Arabidopsis 385 lyrata (Swarbreck et al. 2008). There are drastic differences in the quality of reference genomes 386 available: the A. thaliana reference has seven scaffolds with an N50 of 23,459,830, whereas the 387 A. halleri reference has 282,453 scaffolds with an N50 of 17,686. To control for this limitation, 388 we again removed contigs <10KB from our combination reference genome, which helped with 389 run time and memory usage but did not affect the conclusions (Figure S10b). These tests in 390 Arabidopsis provide an empirical illustration of sppIDer's performance with differing quality 391 reference genomes. Arabidopsis also provides a useful test of detecting hybrids in a plant system, 392 as there are two well-supported allotetraploid species in the genus, Arabidopsis suecica and 393 Arabidopsis kamchatica (Shimizu-Inatsugi et al. 2009; Schmickl et al. 2010). First, we tested 394 short-read data from a divergent lineage of A. thaliana (Durvasula et al. 2017) and found that the 395 reads mapped well to the A. thaliana reference genome (Figure 5d). As expected, reads from the 396 interspecies hybrid A. kamchatica (Novikova et al. 2016) mapped both to A. lyrata and to A. 397 halleri (Figure 5e), approximately equally, confirming that A. kamchatica indeed has genomic 398 contributions from these two species and that sppIDer can detect hybrids, even when the 399 combination reference genome contains reference genomes of substantially varying quality. 400 Thus, sppIDer can accurately detect interspecies hybrid in a plant model and will likely become 401 more generally useful in other plant systems, where allopolyploidy is frequent (Soltis et al. 402 2015), as more reference genomes become available.

403

404 <u>mitoSppIDer</u>

405 Applications of sppIDer with non-nuclear sequencing data are also of considerable 406 interest. Organelle genomes (e.g. mitochondria, chloroplast) have a different mode of 407 inheritance, and increasing data suggest widespread reticulation and cases where their ancestries 408 differ from the nuclear genomes (Peris et al. 2014; Wu et al. 2015; Leducq et al. 2017; Peris et 409 al. 2017a; Peris et al. 2017c; Sulo et al. 2017). We developed mitoSppIDer as an extension to 410 explore these non-nuclear inherited elements. Since mitochondrial genomes are generally small, 411 the coding regions can be easily visualized, which allows precise mapping of introgressions in 412 both coding and non-coding regions. However, more cautious interpretation is warranted, 413 because mitochondrial reads are often at low and variable abundance, and quality can differ 414 between DNA isolations and sequencing runs. Again, we tested using the genus Saccharomyces 415 because of the availability of mitochondrial reference genomes (Foury et al. 1998; Procházka et 416 al. 2012; Baker et al. 2015). We first tested mitoSppIDer with a strain of S. uvarum (ZP1021) 417 (Almeida et al. 2014) and found that, of the reads that mapped to any mitochondrial genome, 418 >99% mapped to the S. uvarum mitochondrial genome (Figure S12a). Next, we examined Vin7, 419 a hybrid strain of S. cerevisiae X S. kudriavzevii, and mitoSppIDer revealed that this strain 420 inherited the mitochondrial genome of S. kudriavzevii with intergenic introgressions from 421 multiple non-S. kudriavzevii mitochondrial genomes (Figure S12b) (Peris et al. 2017c). As with 422 conventional sppIDer, mitoSppIDer rapidly highlights interesting regions for further analysis, 423 such as detailed phylogenetic analyses of introgression candidates.

424

425 <u>Summary</u>

426 Altogether, these tests show the versatility of sppIDer across clades: in fungi, plants, and 427 animals. sppIDer allows for the rapid exploration and visualization of short-read sequencing data 428 to answer a variety of questions, including species identification; determination of the genome 429 composition of natural, synthetic, and experimentally evolved interspecies hybrids; and inference 430 of CCNVs (Brickwedde et al. 2017; Gorter de Vries et al. 2017; Peris et al. 2017b). With 431 examples from the genus Saccharomyces, sppIDer could detect contributions from up to four 432 species and recapitulated the known relative ploidy and aneuploidies of brewing strains. From a 433 simulated phylogeny, we found that sppIDer accurately detected hybrids from a range of 434 divergences in the parents and even detected ancient hybrids. In systems with low-quality or

435 varying quality references genomes, sppIDer performs well without much promiscuous mapping 436 between varying reference qualities, but its ability to infer translocations and CCNVs is limited. 437 Even in systems missing reference genomes, sppIDer still enables rapid inferences by using the 438 reference genomes of closely related species, with the caveat that mapping quality declines with 439 sequence divergence. Additionally, sppIDer works on long-read data and with coverage as low 440 as 0.5X. Finally, sppIDer can be extended to non-nuclear data, allowing for the exploration of 441 alternative evolutionary trajectories of mitochondria or chloroplasts. As more high-quality 442 reference genomes become available across the tree of life, we expect sppIDer will become an 443 increasingly useful and versatile tool to quickly provide a first-pass summary and intuitive 444 visualization of the genomic makeup in diverse organisms and interspecies hybrids.

445

446 Methods:

447 The sppIDer workflow to identify pure species, interspecies hybrids, and CCNVs consists 448 of one main pipeline that utilizes common bioinformatics programs, as well as several custom 449 summary and visualization scripts (Figure 1). An upstream step is required to prepare the 450 combination reference genome to test the desired comparison species. The inputs for the main 451 sppIDer pipeline are this combination reference genome and short-read FASTQ file(s) from the 452 organism to test. The output consists of several plots showing to which reference genomes the 453 short-reads mapped, how this mapping varies across the combination reference genome, and 454 several text files of summary information. Additionally, the pipeline retains all the intermediate 455 files used to make the plots and summary files; these contain much more detailed information 456 and may be useful as inputs to various other potential downstream analyses. We are releasing 457 sppIDer as a Docker, which runs as an isolated, self-contained package, without the need to 458 download dependencies and change environmental settings. Packaging complex bioinformatics 459 pipelines as Docker containers increases their reusability and reproducibility, while simplifying 460 their ease of use (Boettiger 2015; Di Tommaso et al. 2015). sppIDer can be found here 461 (https://github.com/GLBRC/sppIDer), where a transparent Dockerfile lays out the technical 462 prerequisites, platform, how they work in combination, and is a repository for all the custom 463 scripts. A manual for sppIDer can be found both at the GitHub page and at 464 http://sppider.readthedocs.io.

465

466 <u>The pipeline:</u>

Before running the main sppIDer script, a combination reference genome must first be 467 468 created and properly formatted (top of Figure 1). This is a separate script, 469 combineRefGenomes.py, that takes multiple FASTA-formatted reference genomes and a key 470 listing the reference genomes to use and a unique ID for each. The script concatenates the 471 reference genomes together in the order given in the input key, outputting a combination 472 reference FASTA where the chromosomes/scaffolds are renamed to reflect their reference 473 unique ID and their numerical position within the reference-specific portion of the combination 474 output. For reference genomes that contain many short and uninformative scaffolds, there is an 475 option to remove scaffolds below a desired base-pair length. This option improves speed, memory usage, and visual analysis for large genomes with many scaffolds and low N50 values. 476 477 Setting a threshold usually does not affect the conclusions (Figure S10), but we recommend 478 trying different thresholds to determine how much information is lost. The choice of reference 479 genomes to concatenate is completely at the discretion of the user and their knowledge of the 480 system to which they are applying sppIDer. We recommend choosing multiple phylogenetically 481 distinct lineages or species, where gene flow and incomplete lineage sorting are limited, from a 482 single genus. We caution that, for ease of analysis and interpretation, less than 30 reference 483 genomes should be used at once. To illustrate the power of sppIDer, for our examples, we used 484 all available species-level reference genomes for the genera tested, but we excluded lineages and 485 strains within species. However, sppIDer could be applied iteratively with different combinations 486 of reference genomes that are more targeted for a particular lineage or question. For example, 487 with an experimentally evolved hybrid, just the parental genomes could be included to detect 488 CCNVs that occurred during the evolution, but with a suspected hybrid isolated from the wild or 489 industry, all potential parent species reference genomes should be included.

The main body of sppIDer (Figure 1b) uses a custom python 2 (Python Software Foundation) script to run published tools and custom scripts to map short-reads to a combination reference genome and parse the output. The first step uses the mem algorithm in BWA (Li and Durbin 2009) to map the reads to the combined concatenated reference genome. Two custom scripts use this output to count and collect the distribution of mapping qualities (MQ) for the reads that map to each reference genome and produce plots of percentage and MQ of reads that map to each reference genome. The BWA output is also used by samtools view and sort 497 (Li et al. 2009) to keep only reads that map with a MQ > 3, a filter that removes reads that map 498 ambiguously. From here, the number of reads that map to each base pair can be analyzed using 499 bedtools genomeCoverageBed (Quinlan and Hall 2010), for smaller genomes using the 500 per-basepair option (-d) and, for large genomes, the -bga option. The depth of coverage output 501 is used by an R (Wickham 2009; R Core Team 2013) script that determines the mean coverage of 502 the combined reference genome that is subdivided into 10,000 windows of equal size. Finally, a 503 plot for the average coverage for each component reference genome and a second plot of average 504 coverage for the windows are produced.

505

506 <u>The metrics:</u>

507 Several different metrics are used to summarize the data. Depth of coverage is a count of 508 how many reads cover each base pair or region of the genome. Coverage can vary greatly from 509 sequencing run to sequencing run; hence, a log, conversion is used to normalize to the mean 510 coverage. As discussed in the Results, depth of coverage plots can be used to infer the species, 511 the parents of hybrids, and ploidy changes either between or within a genome. sppIDer also 512 reports the percentage of reads that map to each reference genome. Finally, sppIDer uses the 513 established MAPPing Quality (MQ) score introduced in Li et al. (2008) to bin reads by their 514 map-ability on a 0-60 scale. A score of zero is used for reads where it is unlikely that their placement is correct, so sppIDer reports these as "unmapped", along with reads that cannot be 515 516 mapped and therefore do not receive a MQ score. The mapping quality scale can therefore 517 provide a rough assessment of data quality, as well as divergence to the provided reference 518 genomes.

519

520 <u>Tested reference genomes and data:</u>

For the *Saccharomyces* tests, we used reference genomes that are scaffolded to a chromosomal level. In some cases, there is only one reference genome available per species, and for the others, we used the first available near-complete reference; see Table S1 for those used. For systems with multiple reference genomes available, the choice could be more targeted, such as utilizing lineage specific references or references that contains unplaced scaffolds with genes of interest. Alternatively, for systems where few genomes are available, we have shown here that a close relative works as a proxy. For the *Saccharomyces* references, each ordered "ultra-

528 scaffolds" genome was downloaded from http://www.saccharomycessensustricto.org/ or for S. 529 arboricola and S. eubayanus from NCBI. The published S. uvarum genome (Scannell and Zill et 530 al 2011) had chromosome X swapped with chromosome XII, which was fixed manually. These 531 genomes were concatenated together using the python script combineRefGenomes.py, creating a 532 combination reference FASTA with all Saccharomyces species. This combination reference 533 genome can then be used repeatedly to test any dataset of interest. For the Saccharomyces tests, 534 we used publicly available FASTQ data from a number of publications, all available on NCBI (Table S1 contains all accession numbers). Using the data for each strain separately and the 535 536 combination reference genome created above, we then called sppIDer.py with, --out uniqueID, --537 ref SaccharomcyesCombo.fasta, --r1 read1.fastq, and optionally --r2 read2.fastq. sppIDer is 538 written to test one sample's FASTQ file(s) against one combination reference genome at a time, 539 but this could be easily parallelized.

540 For the tests to determine if hybrids could be detected with missing reference genomes, 541 new combination reference genomes without one species' genomes were created by removing 542 the desired species' reference name from the reference genome key before running 543 combineRefGenomes.py. Since both Vin7 and W34/70 contain contributions from S. cerevisiae, the combination reference genome lacking the S. cerevisiae reference was tested for each set for 544 545 FASTQ files for Vin7 and W34/70. The same process was followed to remove the S. 546 kudriavzevii reference genome from the combination reference to test Vin7, as well as to remove 547 the S. eubavanus reference genome from the combination reference to test W34/70.

For the *Lachancea* test, all of the genomes were available and downloaded from http://gryc.inra.fr/. The FASTQ data for CBS6924 was downloaded from NCBI. A combination reference genome with all available genomes was created and used. Then, sequentially, the *"Lachancea fantastica"* and *Lachancea lanzarotensis* genomes were removed by modifying the input key and rerunning combineRefGenomes.py. The FASTQ data for CBS6924 was tested against all three of these combination reference genomes. See Table S1 for the full accessions.

For the non-*Saccharomyces* tests, we used the most complete reference genome available for each species in the genus (accessions Table S1). Therefore, there is quite a bit of variation between different references. For the *Drosophila* and *Arabidopsis* genomes, we tested removing contigs, using the --trim option, with combineRefGenomes.py, as well as not removing contigs, and found the cleanest results when we removed contigs less than 10 Kb. The combined 559 reference genomes of both *Drosophila* and *Arabidopsis* were both larger than four gigabases; 560 therefore, the --byGroup option was used with sppIDer.py to speed up processing and reduce 561 memory usage. The data we tested came from a variety of publications, but we targeted data of divergent or hybrid lineages. See Table S1 for complete information. 562 563 For the mitoSppIDer test, we used the complete species-level Saccharomyces 564 mitochondrial reference genomes available on NCBI, which do not necessarily correspond to the 565 same strain that was used to build nuclear genomic reference (Table S1). Again, 566 combineRefGenomes.py was used to concatenate these references. An additional script, 567 combineGFF.py, was used to create a combination GFF file that was used to denote the coding 568 regions on the output plots. mitoSppIDer.py has an additional flag for the GFF file, but it 569 otherwise runs in a similar manner to sppIDer.py; the same input FASTQ file(s) can even be 570 used. Whole genome sequencing data contains varying amounts of mitochondrial sequences; 571 therefore, using the raw FASTQ data works sufficiently, even when many of the genomic reads 572 will be classified as "unmapped". 573 574 Simulations:

To create the simulated low-quality de novo genomes, we used the software *iWGS* (Zhou et al. 2016) to simulate 100bp paired-end reads with an average inter-read insert size of 350bp (sd 10) at 2X coverage from the reference genomes of *S. cerevisiae*, *S. kudriavzevii*, *S. uvarum*, and *S. eubayanus*. For the simulated de novo *Saccharomyces* genomes the N50 scores ranged from 1254-1274 and the number of scaffolds ranged from 10023-10426 (Table S1).

580 To simulate short-read data, we used DWGSIM (https://github.com/nh13/DWGSIM), 581 which allowed us to vary the coverage, error rate, and mutation rate as needed. The S. cerevisiae 582 reference genome was used to simulate single species reads and a concatenation of the S. 583 cerevisiae and S. eubayanus reference genomes was used for hybrid pseudo-lager reads. As a test 584 of an aneuploid genome, we also manually manipulated the S. cerevisiae reference genome so 585 that it contained zero copies of chromosomes I and III and duplicate copies of chromosome XII,. 586 All simulated reads were 100bp paired-end reads with an average insert size of 500bp. For the 587 coverage tests, we varied the coverage from 0.01-10X. For the short reads used against the low-588 quality de novo genomes, we used 10X coverage and a 3% mutation rate. To simulate PacBio589 style long reads, we used iWGS on the hybrid pseudo-lager concatenated genome with the 590 default settings of 30X coverage, average read accuracy of 0.9, and SD of read accuracy 0.1.

591 To make our simulated phylogeny, we used the S. cerevisiae reference genome as a base 592 and simulated reads with DWGSIM at a 2% mutation rate as 100bp paired-end reads with an 593 average insert size of 500bp at 10X coverage. iWGS was used to assemble these reads. The resulting assembly was again simulated with a 2% mutation rate, and those reads were 594 595 assembled. This procedure was followed for 6 rounds with one lineage being independently 596 simulated twice each round to produce a speciation event. This simulation resulted in 10 species 597 in the phylogenetic arrangement shown in Figure 3a. Summaries of the final assemblies can be 598 found in Table S1, but the median of the final assemblies was 5100 scaffolds, N50 of 1335, and 599 total length of 6.4MB. Each simulated species was $\sim 12\%$ diverged from S. cerevisiae, the most 600 closely related species were $\sim 4\%$ diverged, and the most distantly related species were $\sim 20\%$ 601 diverged. The reads used to produce the final assemblies were used to test whether sppIDer 602 mapped each set of reads to their corresponding reference genomes. The reads of different 603 references were concatenated to simulate pseudo-hybrids of different divergences. To simulate 604 ancient hybrids, the reads from earlier rounds of simulation, before speciation events, were 605 concatenated and tested against the final assemblies with sppIDer. As with the empirical data, to 606 simulate a missing reference genome, that reference was removed from the input key prior to 607 running combineRefGenomes.py.

608

609 <u>Alignment-free phylogenetic methods:</u>

610 We tested four alignment-free phylogenetic methods: two that build phylogenies using 611 short-read data, SISRS (Schwartz et al. 2015) and AAF (Fan et al. 2015), and two that assemble 612 targeted loci from short-read data, aTRAM (Allen et al. 2015) and HybPiper (Johnson et al. 613 2016). We simulated 10X coverage paired-end, 100bp data for each Saccharomyces reference 614 genome at a mutation rate of 0 with DWGSIM to use as input for these methods. For SISRS, we 615 used the default settings with a genome size of 12Mb, first using only the reference 616 Saccharomyces data, then including empirical data for hybrids. SISRS failed at the missing data 617 filtering step when data from the lager strain W34/70 was used, even when we allowed for all but 618 one sample to have missing data. SISRS nexus outputs were visualized with SplitsTree 619 (Huson and Bryant 2006). For AAF, we found that a k of 17 accurately recapitulated the

620 *Saccharomyces* phylogeny, even with the inclusion of empirical data from other pure lineages.

- 621 Once we determined the optimal k, we tested including empirical hybrid data. We also used AAF
- 622 with our simulated phylogeny, which constructed the tree that matched the simulations with the
- default k of 25. The output of AAF was visualized with iTol (Letunic and Bork 2016).
- For the targeted loci methods, we used 13 loci that can delineate *S. eubayanus*
- 625 populations (Peris and Langdon et al. 2016), as well as the ITS sequences for *S. cerevisiae*
- 626 (AY046146.1) (Kurtzman and Robnett 2003) and S. eubayanus (JF786673.1) (Libkind and
- 627 Hittinger et al. 2011) as bait, all obtained from NCBI. We tested the simulated *Saccharomyces*
- reads, as well as the empirical data for P1C1, Fosters O, CBS1503, CBS2834, Vin7, and
- 629 W34/70. For aTRAM, we used the default settings and the option for the Velvet assembler. For
- 630 HybPiper, we used the default settings and the SPADES assembler.
- 631

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- 645

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837 Figure legends

838

839 Figure 1. Workflow of sppIDer. (a) An upstream step concatenates all the desired reference 840 genomes (represented by colored bars). Generally, references should be distinct species (see 841 methods for advice about choosing references). This combination reference genome can be used 842 for many analyses. (b) The main sppIDer pipeline. First, reads (short lines) are mapped. This 843 output is used to parse for quality and percentage (left) or for coverage (right). On the left, 844 quality (high MQ black lines versus low MQ light lines) is parsed, and the percentage of reads 845 that map to each genome or do not map (grey bar) is calculated. To determine coverage, only 846 MQ>3 reads (black lines) are kept and sorted into the combination reference genome order. 847 These reads are then counted, either for each base pair or, for large genomes (combination length 848 >4Gb), in groups. Then, the combination reference genome is broken into equally sized pieces, 849 and the average coverage is calculated. (c) Several plots are produced. Shown here are examples 850 of Percentage Mapped and Mapping Quality plots, a plot showing average coverage by species, 851 and two ways to show coverage by windows with species side-by-side or stacked. Scer = S. 852 cerevisiae, Spar = S. paradoxus, Smik = Saccharomyces mikatae, Skud = S. kudriavzevii, Sarb =853 Saccharomyces arboricola, Suva = S. uvarum, Seub = S. eubayanus. 854 Figure 2. Normalized coverage plots of Saccharomyces test cases. (a) Reads from a New 855 Zealand isolate of S. eubayanus, P1C1, mapped to the S. eubayanus reference genome 856 (magenta). (b) Reads from an ale strain, FostersO, mapped to the S. cerevisiae reference genome 857 (red), with visually detectable aneuploidies. (c) Reads from a hybrid Frohberg lager strain, 858 W34/70, mapped to both the S. cerevisiae and S. eubayanus reference genomes in an average 859 approximately 1:1 ratio with visually detectable translocations and aneuploidies. (d) Reads from 860 a hybrid Saaz lager strain, CBS1503, mapped to both S. cerevisiae and S. eubayanus reference 861 genomes in an average approximately 1:2 (respectively) ratio with visually detectable 862 translocations and aneuploidies. (e) Reads from a wine hybrid strain, Vin7, mapped to S. 863 cerevisiae and S. kudriavzevii (green) reference genomes in an average approximately 2:1 864 (respectively) ratio. (f) Reads from a hybrid cider-producing strain, CBS2834, mapped to four 865 reference genomes: S. cerevisiae, S. kudriavzevii, S. uvarum (purple), and S. eubayanus. 866 Figure 3. Simulated phylogeny of 10 species and sppIDer's detection of hybrids from this 867 phylogeny. (a) Phylogeny built with AAF. (b) Reads from G mapped to the G reference genome.

(c) Reads from a pseudo-hybrid of the closely related species G and H mapped to the G and H
references. (d) Reads from more distant pseudo-hybrid of E and G mapped to references E and
G. (e) Reads of ancient pseudo-hybrid of A and a common ancestor of G and H mapped to the
references of A, G, and H, which are the lineages that descended from the hybrid's parents. (f)
Without the G reference genome, reads from a pseudo-hybrid of the closely related species G
and H mapped to the H reference genome, with some mapped promiscuously to references I and
J.

875 Figure 4. Comparison of the percentage of reads that mapped when different reference genomes 876 were excluded, compared to when all possible reference genomes for Saccharomyces were 877 available (middle panels). (a) When the S. cerevisiae reference genome was not provided and 878 reads from a Frohberg lager strain, W34/70, were mapped, more reads failed to map (grey) or 879 mapped to the S. paradoxus reference genome (yellow). (b) When the full array of 880 Saccharomyces genomes was provided, reads for the lager strain mapped to both S. cerevisiae 881 and S. eubayanus. (c) When the S. eubayanus reference genome was removed, more reads from 882 the lager strain failed to map or mapped to the S. uvarum reference genome (purple). (d) With 883 the removal of the S. cerevisiae reference genome, reads from the S. cerevisiae X S. kudriavzevii 884 hybrid strain Vin7, which would normally map to S. cerevisiae, instead failed to map or mapped 885 to S. paradoxus. (e) When all genomes were used, reads mapped to both S. cerevisiae and S. 886 kudriavzevii. (f) With the removal of the S. kudriavzevii reference genome, reads that would 887 normally map to S. kudriavzevii instead failed to map or were distributed across all other 888 genomes.

Figure 5. Examples using animal and plant genomes. (a) Reads from a *D. yakuba* individual

890 mapped primarily (>99%) to the *D. yakuba* reference genome. (b) Reads from the sister species

891 *D. santomae* mapped best to the *D. yakuba* reference genome with some mapped promiscuously

to other reference genomes. (c) Reads from the more distantly related species D. teissieri mapped

893 mostly to the *D*. yakuba reference genome, but with more reads not mapped and mapped

- 894 promiscuously to other related reference genomes. (d) Reads from an Arabidopsis thaliana
- accession from Tanzania mapped back to the European reference genome for *A. thaliana*. The

repetitive nature of centromeres causes the coverage to fluctuate around those regions. (e) Reads

from the hybrid species *A. kamchatica* mapped to the two parental reference genomes: *A. halleri*

898 and *A. lyrata*.

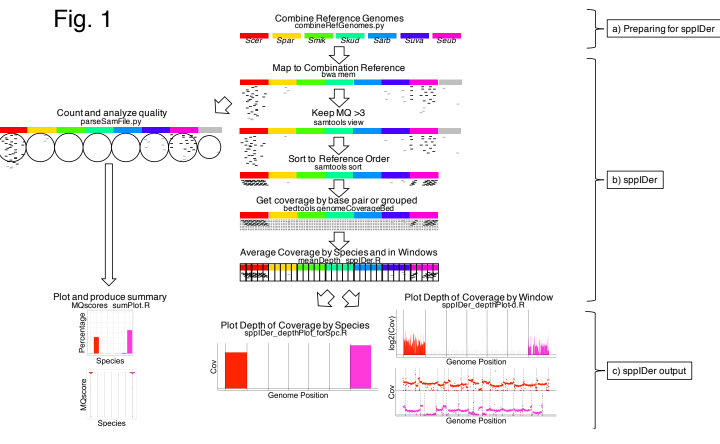
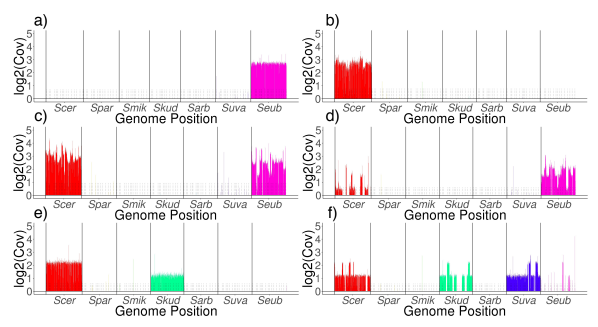


Fig. 2



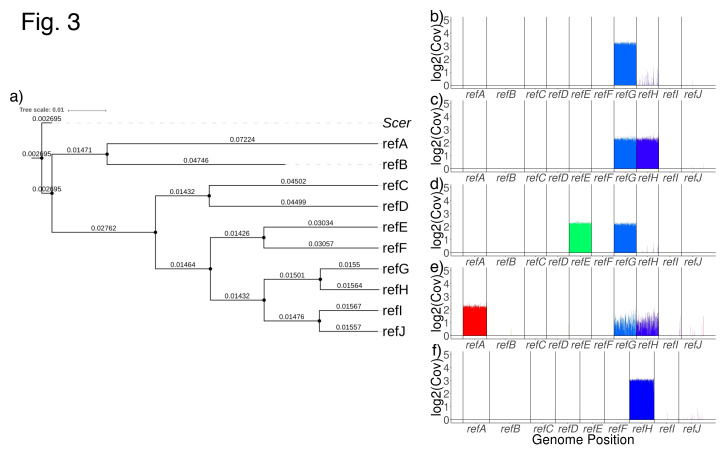
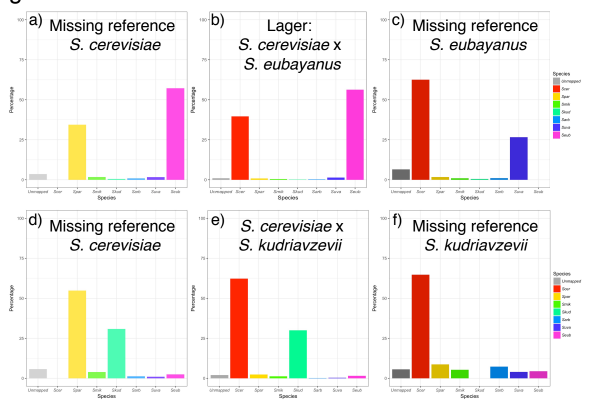


Fig. 4



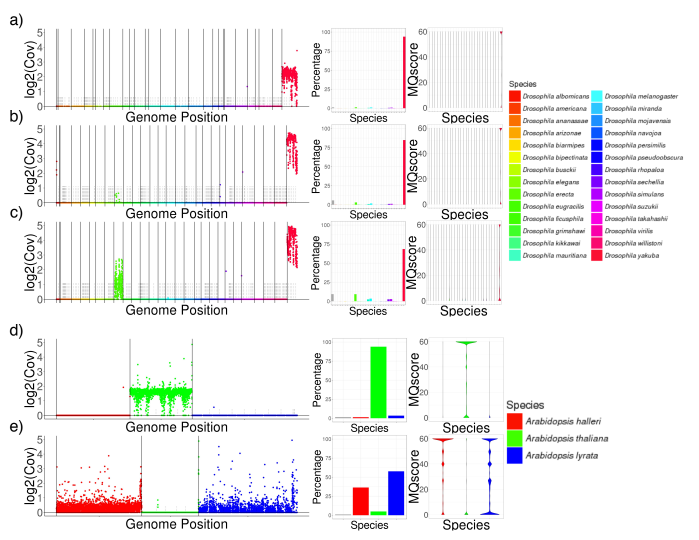


Fig. 5