Supplementary File 1

S1.1 Obligatory criteria

2

- 3 Our differential expression analyses required non-normalised RNA-seq data for the hybrid
- 4 and both parental species in each allopolyploid or homoploid hybrid complex. Only datasets
- 5 that included at least two biological replicates of each species were considered. A genome or
- 6 gene sequences for at least one of the parental species was needed for the utilisation of the
- 7 HyLiTE pipeline (Duchemin et al., 2015) in the differential expression analyses. The
- 8 implementation of GO for functional analyses required genome annotations for at least one
- 9 parental species, however, both a genome and genome annotation were not required,
- therefore if a dataset only had gene annotations available it was still considered. The RNA-
- seq data for the parents and hybrid had to be extracted from the same tissue, or from cells
- grown in the same medium, where applicable, and extracted under the same conditions.

13 S1.2 Preferential criteria

- We gave preference to datasets where the respective hybrid species from which RNA-seq
- data was obtained were naturally occurring (it is common in the literature to see synthetic
- 16 hybrids; commonly formed via artificial crosses for homoploid hybrids or colchicine
- treatment to obtain allopolyploids), to mitigate the influence of synthetic formation on any
- observed gene expression patterns. We also gave preference to allopolyploid and homoploid
- 19 hybrid complexes that were as taxonomically close as possible, to minimise taxon-specific
- 20 differences in observed gene expression patterns. Moreover, we preferred that both parental
- 21 species were extant, where possible, as opposed to extant relatives of the extinct parents.
- 22 A broad search of the literature and online databases was performed with the above criteria
- considered. These are the resulting options and their justifications for use, or omission, in
- 24 differential gene expression analysis.

S1.3 Animals

25

- 26 Achondrostoma oligolepis x Pseudochondrostoma polylepis
- Naturally-occurring homoploid hybrid of two Iberian Leuciscinae fish (Pereira et al., 2014).
- No comparative transcriptomic studies are available of this parent-hybrid complex.
- 29 Ambystoma

- 1 McElroy et al. (2017) studied an allotriploid Ambystoma (mole salamander) with three
- 2 parental species. This would have complicated our analyses as the classes used in the
- 3 categorisation of differential gene expression are based on a two parent system (Cox et al.,
- 4 2014).
- 5 In other publications, it was unclear as to exactly how many species were parents (parental
- 6 species could potentially range from 2-4) of a given hybrid, or if the hybrid had a duplicated
- 7 genome (Bogart et al., 2007; Schmid et al., 2015).
- 8 We additionally searched NCBI BioProjects for Ambystoma. There were some hybrids
- 9 amongst the results, but they did not specify polyploidy/homoploidy, and as no papers have
- 10 given specific hybrid/parent complexes other than McElroy et al, Ambystoma were not used
- in the analyses.
- 12 Bacillus
- 13 Stick insect genus purportedly containing allopolyploid species (Mallet, 2007). Subsequent
- searches of Google to identify specific species were unsuccessful and no data could be found
- on NCBI SRA.
- 16 Bufo viridis
- 17 The B. viridis subgroup contains a complex of diploid, triploid and tetraploid green toad
- species from Central Asia. The parental species of the triploid and tetraploid species are
- unclear (Stöck et al., 2005; Betto-Colliard et al., 2015).
- 20 Bulinus truncates
- 21 Allopolyploid freshwater snail briefly mentioned in (Mallet, 2007). Searches of NCBI SRA
- found no data.
- 23 Carassius auratus x Cyprinus carpio
- 24 Both parental species of this allopolyploid fish (carp) had genome sequences and annotations;
- 25 the issue was a lack of homoploid hybrid complex from within the carp or carp-like fishes.
- Due to the lack of an accompanying publication, it was also unclear if this allopolyploid
- 27 (NCBI BioProject PRJNA86641) was naturally formed.
- 28 Carassius auratus gibelio

- 1 Naturally-occurring allopolyploid fish (carp) formed from Carassius auratus x Cyprinus
- 2 carpio pairing (Li et al., 2014). Only an mtDNA genome was available and limited gene
- 3 models (13), however, there were extensive genome annotations available for the parental
- 4 species. The issue was a lack of homoploid hybrid complex from within the carp fishes.
- 5 Carassius auratus red var. × Cyprinus carpio
- 6 Allopolyploid fish (carp) formed via artificial cross (Wang et al., 2015).
- 7 Carassius auratus red var. × Megalobrama amblycephala
- 8 Allopolyploid fish (carp) with genome sequence, however, only the transcriptome sequence
- 9 for one biological replicate (Qin et al., 2016).
- 10 Cichlids
- Within the literature, there exist a number of studies on cichlids (fish, family: Cichlidae), e.g.
- 12 (Stelkens et al., 2009; Genner & Turner, 2011; Selz et al., 2014; Santos et al., 2016),
- 13 however, none produced RNA-seq data from a comparative analysis between hybrid and
- 14 parents.
- 15 Cobitis taenia x Colbitis elongatoides
- These two diploid fish hybridised to form both diploid and polyploid lineages. RNA-seq data
- was only available for one parental species (*C. taenia*) (Juchno et al., 2007; Juchno & Boroń,
- 18 2018).
- 19 Cottus gobio
- 20 Homoploid hybrid fish derived from C. perifretum x C. rhenanum (Mallet, 2007; Stemshorn
- et al., 2011). No RNA-seq data was available for the hybrid.
- 22 Ctenopharyngodon idellus x Megalobrama amblycephala
- 23 This interspecific diploid fish pairing can form both diploid and triploid hybrids (He et al.,
- 24 2013), however, there is no transcriptomic study that encompasses both parental species and
- 25 the hybrids in the literature.
- 26 Two transcriptome accessions for triploid Ctenopharyngodon idellus x Megalobrama
- 27 amblycephala hybrids can be found under NCBI BioProject PRJNA240314.

- 1 Danio rerio x Danio nigrofasciatus
- 2 A homoploid hybrid zebrafish complex (Spiewak et al., 2018). Although extensive genomic
- 3 information is available for *D. rerio*, there are no comparative transcriptomic data available
- 4 for this complex.
- 5 Artificial triploid *D. rerio* can be generated, however these are not allopolyploid (Mizgireuv
- 6 et al., 2004).
- 7 Daphnia
- 8 The water flea genus Daphnia does contain hybrid species, such as D. galeata \times D. hyalina
- 9 (Wolinska et al., 2006) and some allopolyploids within the D. pulex complex (Dufresne,
- 10 2011). Comparative RNA-seq data could not be found for either complex.
- 11 Drosophila
- 12 Interspecific homoploid hybrids are common in the fruit fly genus *Drosophila* (Goulielmos &
- Alahiotis, 1989; Landry et al., 2005; Kelleher et al., 2012). Some Drosophila species do
- exhibit mosaic polyploidy (Fox et al., 2010), but there are no entirely allopolyploid species.
- 15 Epinephelus coioides x Epinephelus lanceolatus
- 16 Under NCBI BioProject PRJNA431058 there are transcriptome accessions for diploid and
- 17 triploid hybrids from the Epinephelus coioides x Epinephelus lanceolatus (grouper fish)
- 18 pairing. However, there are no accompanying parental transcriptomes, and only one
- 19 biological replicate at each ploidy level with two different tissues sampled.
- 20 BioProject PRJNA497240 has transcriptomic sequences for *E. coioides*, however, there is
- 21 only one replicate subjected to different treatment condition. Likewise, BioProject
- 22 PRJNA413272 has transcriptomic sequences for a single replicate of *E. lanceolatus*, sampled
- over a time series, post-hatch.
- 24 Gasterosteus nipponicus x Gasterosteus aculeatus
- 25 Parent-hybrid fish complex with 6 or 7 RNA-seq runs available for each species, found under
- 26 NCBI BioProject PRJDB7242. Genome with gene models also available for G. aculeatus
- 27 (three-spined stickleback fish). However, as data was unpublished, it was unclear if the
- 28 hybrid was homoploid and if it was naturally or synthetically formed.

- 1 Gila seminuda
- 2 Homoploid hybrid fish formed from the hybridisation of *Gila robusta* x *Gila elegans* (Mallet,
- 3 2007; Mavárez & Linares, 2008). Only an organellar genome is available for G. robusta on
- 4 NCBI, in addition to 13 gene models. Further, no comparative RNA-seq data is available for
- 5 this hybrid complex.
- 6 Megalobrama amblycephala x Cyprinus carpio
- 7 RNA-seq data for the hybrid was unavailable (Wang et al., 2017). The research group was
- 8 contacted but no response was obtained.
- 9 Morone chrysops x Morone saxatilis
- 10 Six transcriptome accessions of homoploid hybrid striped bass available under NCBI
- BioProject PRJNA382266. There were no accompanying parental transcriptome accessions.
- 12 A single transcriptomic replicate for M. chrysops and M. saxatilis were found under
- BioSample SRR1187194 and SRR1187193, respectively.
- 14 Oncorhynchus
- 15 This genus in the Salmonidae family contains a number of hybrid species, including
- Oncorhynchus keta x Oncorhynchus gorbuscha (Zhivotovsky et al., 2016), and diploid and
- triploid *Oncorhynchus kisutch* (Withler et al., 1995). No comparative transcriptomic analyses
- were found for either an allopolyploid or homoploid *Oncorhynchus* complex.
- 19 Pelophylax esculentus
- 20 The hybrid water frog of Pelophylax lessonae x Pelopyhlax ridibundus (Doležálková-
- 21 Kaštánková et al., 2018). Searches of NCBI database and https://amphibiaweb.org did not
- 22 find any genome sequences or comparative RNA-seq analyses.
- 23 Poecilia formosa
- Naturally-occurring allopolyploid fish (molly) formed from *Poecilia Mexicana* x *Poecilia*
- 25 latipinna (Schedina et al., 2014). Genome sequences and annotations were available for both
- parental species and the allopolyploid. The issue was a lack of homoploid hybrid complex
- within the mollies.
- 28 Salmo salar

- 1 Alantic salmon can occur as natural diploid and artificial triploid hybrids (Refstie & Gjedrem,
- 2 1975; Vera et al., 2017; Murray et al., 2018). However, there are no comparative
- 3 transcriptomic analyses of parental species and hybrids in the literature.
- 4 Squalius
- 5 A genus of Cyprinidae fish with allopolyploid and homoploid hybrids produced from the
- 6 same parental pairing. S. alburnoides exists as natural populations of hybrid allotriploid PAA,
- 7 homodiploid hybrid PA, and the parental-like diploid AA genomotypes. The parental species
- 8 of S. alburnoides are S. pyrenaicus (PP) and an extinct species related to Anaecypris
- 9 hispanica (Pala et al., 2010; Matos et al., 2015).
- 10 There are genome sequences and annotations for S. pyrenaicus available at
- 11 ftp://ftp.ncbi.nlm.nih.gov/genomes/all/GCA/001/403/095/GCA_001403095.1_pp_cds_nonRe
- dudant, there is also a GenBank record (accession CVRK00000000) for the S. pyrenaicus
- whole genome shotgun sequencing project, but it contains no sequence data.
- Pala et al. (2008) performed cDNA sequencing of allotriploid S. alburnoides, but only for 6
- 15 genes.
- 16 There is comparative small RNA data available for this complex under GEO Series
- 17 GSE38691 (Inácio et al., 2012).
- 18 Comparative RNA-seq data for allopolyploid and homoploid Squalius complexes can be
- 19 found under ArrayExpress E-MTAB-3174 (Matos et al., 2015, 2019). This genus is a strong
- 20 candidate for our homoploid and allopolyploid animal representatives.
- 21 Takifugu rubripes x Takifugu flavidus
- 22 An artificially formed homoploid hybrid fish (pufferfish), with transcriptome accessions only
- 23 available for one replicate each of the parental species and hybrid (Gao et al., 2013).
- 24 Tigriopus californicus
- 25 Copepod species whose hybridisations are limited to being inter-populational only (Pritchard
- 26 et al., 2013; Barreto et al., 2014).
- 27 Xenopus laevis
- 28 Allopolyploid African clawed frog with unknown parental species (Session et al., 2016).

1 *S1.4 Fungi*

- 2 Aspergillus flavus x Aspergillus parasiticus
- 3 These Ascomycete molds form interspecific hybrid strains through artificial crosses, some
- 4 with evidence of allopolyploidisation (Olarte et al., 2015). Genome sequences and
- 5 annotations for both parental species are available through the NCBI database. No
- 6 comparative transcriptomic studies available.
- 7 Blumeria graminis f. sp. triticale
- 8 A homoploid species of powdery mildew formed from the hybridisation of B. g. f. sp. tritici
- 9 and B. g. f. sp. secalis. There is an annotated genome available for B. g. tritici and RNA-seq
- data for all members of this complex available. However, these transcriptomic data did not
- originate from a single study, so the conditions around RNA extraction could not be
- 12 guaranteed as controlled across all samples, therefore we would not use this as a first option
- dataset (Menardo et al., 2016; Praz et al., 2018).
- Additional comparative transcriptomic analyses relating to this complex involve only B. g.
- 15 *tritici* (Hu et al., 2018).
- 16 Epichloë canadensis
- 17 Allopolyploid endophyte hybrid of *E. elymi* x *E. amarillans* with RNA-seq data with
- 18 replicates for all members of the complex, in addition to genomes and gene sequences for
- both the parental species (data unpublished). A strong candidate for our allopolyploid fungal
- 20 complex.
- 21 Neotyphodium lolii x Epichloë typhina Lp1
- 22 Allopolyploid endophyte. RNA-seq data with replicates available for all members of the
- complex, in addition to gene sequences for both the parental species (Cox et al., 2014;
- 24 Campbell et al., 2017).
- 25 Ophiostoma ulmi x Ophiostoma novo-ulmi
- 26 Transient hybrids of *Ophiostoma ulmi* x *Ophiostoma novo-ulmi* do occur in nature (Brasier,
- 27 2001). No comparative transcriptomic analyses of this homoploid hybrid complex are

- available; only a single study in which the O. novo-ulmi transcriptome was compared with
- 2 that of Candida albicans and Histoplasma capsulatum (Nigg et al., 2015).
- 3 Saccharomyces cerevisiae x Saccharomyces bayanus
- 4 Homoploid hybrid yeast. Genome annotations for both parental species are available at
- 5 http://www.saccharomycessensustricto.org/cgi-
- 6 bin/s3.cgi?data=Annotations&version=current, in addition to RNA-seq reads from a
- 7 comparative transcriptomic analysis of this complex. The hybrids were generated
- 8 synthetically through mating a single haploid individual of each parental species, growing a
- 9 clonal population of the diploid, and checking it by PCR (Schraiber et al., 2013).
- 10 Saccharomyces cerevisiae x Saccharomyces kudriavzevii
- 11 There is a comparative transcriptomic study available that used two biological replicates of
- this homoploid hybrid yeast, however, it used a microarray and only compared the hybrids at
- varying temperatures (Tronchoni et al., 2017).
- 14 Saccharomyces cerevisiae x Saccharomyces paradoxus
- 15 Homoploid hybrid yeast. RNA-seq-based transcriptomic analyses of multiple replicates are
- 16 available. The hybrids were generated synthetically through mating a single haploid
- individual of each parental species, growing a clonal population of the diploid, and checking
- it by PCR (Schraiber et al., 2013). Genome annotations for both parental species available at
- 19 http://www.saccharomycessensustricto.org/cgi-
- bin/s3.cgi?data=Annotations&version=current. A strong candidate for our homoploid fungal
- 21 complex.
- Swain Lenz et al. (2014) also did a comparative transcriptomic analysis of this complex,
- 23 however, due to the genetic manipulations performed during their synthetic crossing of the
- parental species, we have opted to use the data from Schraiber et al.
- 25 Saccharomyces cerevisiae x Saccharomyces uvarum
- 26 Synthetically formed homoploid hybrids (Pfliegler et al., 2012). Although genome
- 27 annotations are available for both parental species on
- 28 http://www.saccharomycessensustricto.org/cgi-
- 29 bin/s3.cgi?data=Annotations&version=current, the only comparative transcriptomic studies

- 1 available relating to this complex are of a single hybrid replicate grown at varying
- 2 temperatures, available under NCBI BioProject PRJNA505697.
- 3 Saccharomyces pastorianus (syn. Saccharomyces carlsbergensis)
- 4 This allopolyploid yeast is formed from the mating of S. cerevisiae and S. eubayanus. The
- 5 allopolyploid does not have a genome sequence or annotations available, but both parental
- 6 species do through the NCBI database. A comparative transcriptomic analysis of this
- 7 complex was performed by (Gibson et al., 2010), however, they used a microarray. It was
- 8 unclear whether the S. pastorianus they used was a naturally occurring allopolyploid, so the
- 9 authors were emailed to clarify. Their response was that their experimental procedure
- involved an industrial strain that had been in use for decades, thus its ancestry is not entirely
- 11 certain. However, the author did not think it had originated from a synthetic protocol. The
- 12 literature also suggests that S. pastorianus arose through instantaneous speciation during an
- interspecific hybridisation event (Dunn & Sherlock, 2008).
- 14 Zygosaccharomyces parabailii
- A homoploid hybrid species of *Z. bailii sensu stricto* and an unidentified *Zygosaccharomyces*
- species. There is an RNA-seq study of the response to lactic acid stress in Z. parabailii, but it
- does not involve comparisons to its parental species (Ortiz-Merino et al., 2018).
- 18 Zygosaccharomyces rouxii
- 19 Yeast with allopolyploid strains. Zygosaccharomyces rouxii NBRC 1876 draft genome
- sequence is available (Sato et al., 2017). RNA-seq data for this species can be found under
- 21 NCBI BioProject PRJNA437612. However, no comparative transcriptomic studies were
- 22 found through NCBI SRA.
- 23 Zymoseptoria pseudotritici
- 24 Zymoseptoria pseudotritici is a fungal homoploid hybrid grass pathogen with unknown
- parental species (Stukenbrock et al., 2012).
- 26 *S1.5 Plants*
- 27 Arabidopsis suecica

- 1 Natural allopolyploid of A. thaliana x A. arenosa (Jeffrey Chen et al., 2004). Comparative
- 2 RNA-seq data of this complex is available under NCBI BioProject PRJNA393427 with two
- 3 biological replicates for each member, in addition to genome annotations available through
- 4 NCBI SRA. The main issue was no suitable *Arabidopsis* homoploid hybrid complex.
- 5 Arabidopsis thaliana x Arabidopsis lyrata
- 6 Artificial homoploid hybrid with two RNA-seq replicates available for the hybrid and A.
- 7 lyrata (Zhu et al., 2017). The issue was that this study did not perform RNA-seq for A.
- 8 *thaliana*, so there was no guarantee that the conditions under which extraction was performed
- 9 were held constant across all species.
- 10 Arachis hypogaea
- Peanut is an allopolyploid of A. ipaensis and A. duranensis. RNA-seq data, genomes and
- genome annotations for the parental species and the allopolyploid are available (Bertioli et
- 13 al., 2019).
- 14 Further RNA-seq data for allopolyploid A. hypogaea is also available, without biological
- replicates (Han et al., 2017).
- 16 The major issue was a lack on a homoploid hybrid complex from within this genus.
- 17 Brassica juncea
- Allotetraploid hybrid of *B. rapa* and *B. nigra*. (Yang et al., 2016) did RNA-seq analysis of *B.*
- 19 *juncea* only, in addition to a genome annotation of *B. juncea*. The genome annotation of *B.*
- *rapa* is available through NCBI SRA.
- 21 Brassica napus
- 22 An allotetraploid hybrid of B. rapa x B. oleracea. RNA-seq data with three biological
- 23 replicates for each member of the complex is available under NCBI BioProject
- 24 PRJNA449400 (Wu et al., 2018). All members have genes and genomes available through
- NCBI SRA. This is a possible option for an allopolyploid complex.
- 26 Brassica napus x Brassica carinata
- 27 Homoploid hybrid artificially produced via hand pollination and embryo rescue. Comparative
- 28 RNA-seq analysis of this homoploid hybrid complex with three biological replicates for each

- 1 member, and functional gene annotations is available (Chu et al., 2014). Extensive genome
- 2 annotations are available for *B. napus* through NCBI SRA. Limited gene models are available
- 3 for B. carinata through NCBI SRA and http://brassicadb.org. This is a possible option for a
- 4 homoploid hybrid plant complex.
- 5 Brassica napus x Brassica rapa
- 6 Comparative RNA-seq analysis of this homoploid hybrid complex (Zhang et al., 2015), with
- 7 no biological replicates.
- 8 Camellia
- 9 Camellia reticulata can exist as an allopolyploid (Gu & Xiao, 2003), but there are no
- 10 comparative transcriptomic studies on the allopolyploid complex. The only related RNA-seq
- data on NCBI SRA is of five different *C. reticulata* tissues (Yao et al., 2016).
- A homoploid hybrid complex exists in Camellia azalea x Camellia amplexicaulis, with
- 13 RNA-seq data on each member available. There are no biological replicates; the RNA
- libraries were formed from pooled RNA of three individuals (Zhang et al., 2018).
- 15 Capsella bursa-pastoris
- Natural allopolyploid of *C. orientalis* and *C.grandiflora*. Multiple replicates of RNA-seq data
- are available for the allopolyploid and parental species (Kryvokhyzha et al., 2019).
- 18 Capsella rubella x Capsella grandiflora
- 19 Reciprocal homoploid hybrids with RNA-seq data, however, no accompanying RNA-seq data
- 20 is available for the parental species (Rebernig et al., 2015).
- 21 *Chrysanthemum nankingense* x *Tanacetum vulgare*
- Homoploid hybrid with comparative RNA-seq data available for hybrid and parental species
- 23 (Wang et al., 2013; Qi et al., 2018). Qi et al. also generated allopolyploids through colchicine
- 24 treatment of homoploid hybrids in addition to performing functional annotations of genes,
- 25 making this a possible option for our analyses.
- 26 Coffea arabica

- 1 Natural allopolyploid of C. eugenioides and C. canephora. Data from a microarray-based
- 2 comparative analysis of this complex are available (Bardil et al., 2011), as well as an RNA-
- 3 seq-based comparative analysis of *C. arabica* grown under different temperatures (Combes et
- 4 al., 2013).
- 5 Glycine
- 6 No records of an RNA-seq-based study of Glycine homoploid hybrids. However, an
- 7 intraspecific comparative transcriptome analysis of homoploid G. max hybrids has been done
- 8 (Zhang et al., 2017).
- 9 *Glycine dolichocarpa*
- 10 A naturally formed recent allopolyploid of G. syndetika and G. tomentella. RNA-seq data for
- only the allopolyploid is available (Coate et al., 2014).
- A comparative transcriptomic study of the allopolyploid and its parents is found here, with 3
- biological replicates per member (Ilut et al., 2012). https://soybase.org only has G. max and
- 14 G. soja genome and gene sequences. NCBI SRA has 127 gene models available for each of
- 15 *G. syndetika, G. tomentella and G. dolichocarpa.*
- 16 Transcriptomic analyses of this complex can also be found here (Coate et al., 2012).
- 17 Gossypium
- 18 Microarray-based analysis of natural allopolyploid cotton (G. hirsutum) and synthetic
- 19 homoploid hybrid progeny of *G. arboreum* and *G. raimondii* (Flagel et al., 2008).
- 20 RNA-seq based analysis of natural allopolyploid (G. hirsutum) and synthetic homoploid
- 21 hybrid progeny of G. arboreum and G. raimondii (Yoo et al., 2013). There are 3 biological
- 22 replicates of RNA-seq data available for each member under NCBI BioProject
- 23 PRJNA171342. The synthetic hybrid was formed by emasculation of the A2 parent in the
- evening and subsequent cross-pollination the following morning. They are a strong option for
- our homoploid and allopolyploid plant representatives.
- 26 Helianthus

- 1 The sunflower genus *Helianthus* contains several homoploid hybrid species (Ungerer et al.,
- 2 2006). RNA-seq data is available (Renaut et al., 2014), in addition to a reference genome for
- 3 the parental species *H. annuus* (Badouin et al., 2017).
- 4 There are some purportedly allopolyploid *Helianthus* species (Espinasse et al., 1995), but
- 5 data on these are very limited, especially RNA-seq.
- 6 Mimulus
- 7 Mimulus peregrinus is a naturally formed recent allopolyploid of M. luteus x M. guttatus.
- 8 Genomes and genome annotations for both parents found at
- 9 https://datadryad.org/resource/doi:10.5061/dryad.d4vr0. Edger et al. (2017) performed a
- 10 comparative RNA-seq analysis of this complex, in addition to an artificially formed M. x
- 11 robertsii of the same parent pairing. Mimulus x robertsii was formed by hand pollination via
- diallel cross. They also did GO and KEGG. The issue was there were no biological replicates
- for any of the RNA-seq data.
- 14 Oryza
- 15 Comparative RNA-seq analysis of intraspecific rice hybrids and their derived allopolyploids
- 16 (Xu et al., 2014).
- 17 Comparative RNA-seq analysis of a synthetic interspecific rice hybrid and its two parental
- species (Wu et al., 2016). However, the hybrid is a triploid produced from a diploid and
- tetraploid, so its genome size relative to one parent has increased and to the other, decreased,
- and thus it would confound the interpretation of our analyses.
- 21 Saccharum
- 22 Modern allopolyploid and homoploid sugarcane cultivars are commonly formed from
- pairings between domesticated S. officinarum and wild S. spontaneum (Song et al., 2016).
- 24 RNA-seq data for S. officinarum, S. spontaneum, and their F₁ hybrid are available under
- 25 NCBI BioProject PRJNA335885. Saccharum intermedium, an allopolyploid of S.
- 26 angustifolium and S. villosum, has recently been described (Welker et al., 2017), but the
- 27 complex lacks accompanying RNA-seq data.
- 28 Senecio

- 1 Comparative transcriptomic analyses of both an allopolyploid (S. cambrensis = S. vulgaris x
- 2 S. squalidus) and homoploid hybrid ($S \times baxteri = S$. vulgaris $\times S$. squalidus) of this ragwort
- 3 genus are available (Hegarty et al., 2006), however, they are microarray-based. Further
- 4 microarray-based transcriptomic analyses of this genus are also available (Hegarty et al.,
- 5 2005; Hegarty et al., 2008, 2009).
- 6 Spartina
- 7 The cordgrass genus contains a recently formed natural allopolyploid, S. anglica (Ainouche
- 8 et al., 2004), two interspecific hybrids S. x neyrautii and S. x townsendii, and the parental
- 9 species S. maritima and S. alterniflora. Non-normalised Illumina data is available for these
- 10 complexes as assembled reference transcriptomes (Boutte et al., 2016), but comparative
- 11 RNA-seq data is as yet unavailable.
- Microarray data for allopolyploid, homoploid hybrid and parental *Spartina* are also available
- 13 (Chelaifa et al., 2010).
- 14 Tragopogon
- There is RNA-seq data with biological replicates for the allopolyploid species *T. mirus* and *T.*
- 16 *miscellus* and their diploid parents available under NCBI BioProject PRJNA210897.
- 17 A study of *T. miscellus*, its parental species, and their synthetically produced homoploid
- 18 hybrid has been done, but it was not RNA-seq based, rather surveying expression of 144
- duplicated gene pairs derived from the parental species (Buggs et al., 2011).
- 20 Triticeae
- 21 Synthetic allopolyploid of Aegilops sharonensis and Triticum monococcum ssp. aegilopoides
- obtained through colchicine treatment of F₁ hybrids. Gene expression was analysed through
- cDNA-AFLP (Kashkush et al., 2002).
- 24 Synthetic allopolyploid of Aegilops longissimi and Triticum urartu. Gene expression of the
- 25 allopolyploid and its diploid parental species was analysed through RNA-seq (Wang et al.,
- 26 2016). The main issue was that the allopolyploid was synthetic, as well as the lack of RNA-
- seq data for a homoploid hybrid complex within these genera.
- 28 Vigna reflexo-pilosa var. glabra

- 1 Allopolyploid mungbean species with hypothetical parental species, one of which is thought
- 2 to be extinct. RNA-seq assemblies are available for the allopolyploid (Kang et al., 2014)
- 3 Zea mays
- 4 Maize is purportedly of allotetraploid origin that has subsequently undergone rediploidisation
- 5 (Gaut et al., 2000). Thus, it would not be suitable for investigating the impact of genome
- 6 doubling on gene expression.
- 7 Zea mays x Tripsacum dactyloides
- 8 Interspecific hybrid of maize and gamagrass (Shavrukov & Sokolov, 2015) with
- 9 transcriptomic studies of each parent available (Wu et al., 2017; Gault et al., 2018), albeit
- 10 none for the entire complex.

- 1 S1.6 References
- 2 Ainouche, M. L., Baumel, A., & Salmon, A. (2004). Spartina anglica C. E. Hubbard: a
- 3 natural model system for analysing early evolutionary changes that affect allopolyploid
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