# Convergent evolution of desiccation tolerance in grasses

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### 15 Abstract

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Desiccation tolerance has evolved repeatedly in plants as an adaptation to survive extreme environments. Plants use similar biophysical and cellular mechanisms to survive life without water, but convergence at the molecular, gene, and regulatory levels remains to be tested. Here, we explore the evolutionary mechanisms underlying the recurrent evolution of desiccation tolerance across grasses. We observed substantial convergence in gene duplication and expression patterns associated with desiccation. Syntenic genes of shared origin are activated across species, indicative of parallel evolution. In other cases, similar metabolic pathways are induced, but using different gene sets, pointing towards phenotypic convergence. Species-specific mechanisms supplement these shared core mechanisms, underlining the complexity and diversity of evolutionary adaptations to drought. Our findings provide insight into the evolutionary processes driving desiccation tolerance and highlight the roles of parallel and convergent evolution in response to environmental challenges. 

#### 38 Introduction

39 Anhydrobiosis, or life without water, is rare but widely distributed across life, spanning 40 microbial, animal, and plant lineages. Plants that can tolerate desiccation in their vegetative 41 tissues are known as resurrection plants due to their dramatic ability to revive from an extremely 42 dry state (water potential < -100 MPa or relative water content < 10%)<sup>1</sup>. Desiccation tolerance likely arose in plants during the Ordivocian period and is thought to have played a critical role in 43 44 facilitating the transition from aquatic to terrestrial environments by early land plants<sup>2</sup>. These 45 ancestral mechanisms of anhydrobiosis were retained in many non-seed plants (e.g., mosses, liverworts, ferns, and fern allies) and there is a high frequency of vegetative desiccation 46 tolerance among extant bryophytes and pteridophytes<sup>3</sup>. In contrast, vegetative desiccation 47 tolerance was lost, or suppressed, in the common ancestor of seed plants, presumably in a 48 49 tradeoff for other systems of drought avoidance and escape, such as annual life histories, water 50 transport, and retention mechanisms including stomata, vasculature, and roots<sup>4</sup>. Desiccation tolerance then re-evolved convergently in a subset of vascular plants, likely through the rewiring 51 of ancestral anhydrobiosis pathways maintained in seeds, spores, and pollen <sup>5-7</sup>. The retention 52 53 and re-evolution of desiccation tolerance seems to have been driven by a combination of 54 selective pressures in habitats with extreme water limitation, seasonal drought, and sporadic water availability <sup>8</sup> Consequently desiccation tolerance is more common in some lineages than 55 56 others, but diverse species of resurrection plants can often be found co-occurring in tightly 57 intertwined communities on rocky outcroppings in arid tropical and subtropical regions across the world <sup>3,9</sup>. 58

59 Despite more than 500 million years of evolution and divergence across extant resurrection plants, multiple biochemical and physiological mechanisms of desiccation tolerance 60 61 are shared across distantly related species. For example, all surveyed resurrection plants 62 accumulate small non-reducing sugars and other osmoprotectants to vitrify the cytoplasm and safeguard macromolecules during drying <sup>10</sup>. Dramatic shifts in carbohydrate and lipid 63 metabolism as well as the protection (or in some cases degradation) of photosynthetic apparati 64 are also observed in all resurrection plants during drying <sup>11–14</sup>. All surveyed desiccation tolerant 65 plants leverage robust anti-oxidant scavenging systems, mobilize numerous intrinsically 66 67 disordered and protective proteins, and have specialized cell wall properties that maximize flexibility and mitigate the mechanical strain of shrinkage <sup>3,14,15</sup>. These broad features of 68 69 anhydrobiosis are largely shared across organisms and tissues, but the specific metabolic 70 pathways, regulatory networks, and activated genes are notably complex and variable among species <sup>3,10,16</sup> and tissues <sup>17</sup>. 71

72 The recurrent evolution of desiccation tolerance offers an exciting opportunity to 73 understand how complex traits evolve independently across both broad and narrow phylogenetic distances. The evolution of complex traits can occur via multiple pathways <sup>18,19</sup>, 74 and it is often assumed that when closely related taxa evolve the same trait independently, they 75 do so by leveraging the same genetic pathways (parallelism) due to internal constraints within 76 77 that lineage <sup>20</sup>. In contrast, when distantly related taxa evolve the same trait independently they are expected to leverage divergent pathways and genes (convergence), due to contrasting 78 genetic starting points <sup>19,21</sup>. However, these patterns are not always observed in nature, and 79 contradictory examples exist, where distantly related taxa exhibit independent but identical 80 mutations and closely related taxa do not <sup>21</sup>. The recurrent evolution of desiccation tolerance at 81

multiple phylogenetic scales provides an ideal system to untangle the mechanisms of
convergent and parallel evolution. An important first step towards decoding the evolutionary
pathways to desiccation tolerance is characterizing the extent of shared genetic adaptations,

85 overlapping pathways, and lineage specific processes across resurrection plants .

86 Desiccation tolerance has received growing research attention in recent years and several resurrection plants have emerged as models for understanding this remarkable trait<sup>22</sup>. 87 88 Desiccation tolerance is found in at least ten angiosperm families, and is most common in 89 Poaceae, where it evolved independently at least six times across three subfamilies and is found in dozens of grass species<sup>3</sup>. Thus, the grasses are an excellent system to test if the same 90 91 pathways, regulatory modules, and mechanisms were recruited during the recurrent evolution of 92 desiccation tolerance. Most genomic studies of resurrection plants have investigated only a single species in isolation <sup>17,23-27</sup> or tolerant and sensitive taxon comparison <sup>7,28,29</sup>, but none 93 94 have identified core responses shared among independent lineages of resurrection plants. 95 Here, we quantify the extent of shared mechanisms of anhydrobiosis across resurrection grasses and investigate the roles of parallel mutation and convergent pathway adaptation in the 96 97 evolution of desiccation tolerance. We present highly contiguous genome assemblies of three 98 resurrection grasses native to Sub-Saharan Africa coupled with comprehensive gene 99 expression datasets and supporting physiological data. We leveraged comparative genomic and 100 transcriptomic approaches to investigate the evolution of desiccation tolerance in these three 101 species. We also extend these analyses to other desiccation tolerant and sensitive grasses to 102 describe a core signature that defines desiccation tolerance. 103

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### 105 Results

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### 107 **Comparative genomics of desiccation tolerant grasses**

108 We searched for signatures of convergent evolution across three grasses in two 109 Chloridoideae subtribes representing at least two independent origins of desiccation tolerance: 110 Microchloa caffra Nees. in subtribe Eleusininae and Oropetium capense Stapf. and Tripogon 111 minimus Steud. in the Tripogoninae subtribe. These three species have overlapping 112 distributions and tend to co-occur in shallow soils on rocky outcroppings, locally known as 113 ruwari, across Sub-Saharan Africa (Figure 1a). Microchloa caffra, commonly known as 114 pincushion grass, is distributed from Uganda to South Africa and is the largest of the three 115 species. Oropetium capense is smaller and grows as densely packed tufts on exposed rock 116 surfaces. Tripogon minimus. is a small but loosely tufted grass that occurs in shallow soils in 117 both western and southern Africa (Figure 1a). *Microchloa caffra* plants were collected from 118 Buffelskloof Private Nature Reserve in Mpumalanga and O. capense and T. minimus were 119 collected from Swebe Swebe Private Wildlife Reserve in Limpopo, South Africa. 120 We generated reference genome assemblies for each of the three grasses using PacBio 121 HiFi data. O. capense and T. minimus are diploid with haploid genome sizes of ~195 Mb based 122 on flow cytometry, and *M. caffra* is hexaploid with a 1.25 Gb haploid genome. Sequencing reads were assembled using Hifiasm <sup>30</sup>, producing near complete reference assemblies for O. 123 capense and T. minimus and a highly contiguous draft assembly of M. caffra (Table 1). Six and 124 125 nine of the ten chromosomes were assembled telomere-to-telomere for T. minimus and O.

126 capense respectively, and the remaining chromosomes were split into two contigs. The M. 127 caffra genome assembly was more fragmented, with 118 contigs spanning 968 Mb and a contig 128 N50 of 16 Mb. The monoploid genome size of *M. caffra* is 322 Mb, which is roughly 30% larger 129 than O. capense and T. minimus (237 and 223 Mb, respectively), and this expansion was driven 130 largely by DNA transposons. All three species have a similar proportion of long terminal repeat 131 retrotransposons (22-27%), but 27% of the *M. caffra* genome is composed of DNA transposons 132 compared to 12% in O. capense and 16% in T. minimus (Table 1). Despite this expansion of 133 transposons in *M. caffra*, the three Chloridoid grasses have very compact genomes compared to most grasses <sup>31</sup>. We used the MAKER-P pipeline to annotate these three genome 134 135 assemblies, with RNAseq data and protein homology as evidence. The O. capense and T. 136 *minimus* genome assemblies have 28,826 and 26,527 gene models respectively, which is 137 comparable to the well-annotated model resurrection plant Oropetium thomaeum (L.f.) Trin (28,835)<sup>24,32</sup>. The *M. caffra* genome assembly has 85,245 gene models, which matches the 138 expectations for a hexaploid genome (Table 1). We assessed annotation guality using the land 139 140 plant (Embryophyta) dataset of Benchmarking Universal Single-Copy Orthologs (BUSCO) and 141 found between 95.3-97.1% complete proteins across the three grasses, suggesting the genome 142 assemblies are largely complete and well-annotated (Table 1). 143 We leveraged comparative genomic approaches to identify evolutionary signatures

144 associated with desiccation tolerance and enable cross-species comparisons of gene 145 expression data. The three grass genomes are largely collinear with O. thomaeum, and have 146 considerable conserved gene content despite some notable structural rearrangements. Seven 147 pairs of O. thomeaum and O. capense chromosomes have near perfect synteny, with 148 chromosomes 8 and 9 showing a few large-scale inversions, and a telomeric translocation on 149 chromosome 2 (Supplemental Figure 1). Tripogon minimus has similar macrosynteny with O. 150 thomaeum, but has no rearrangements in chromosome 8. Synteny between M. caffra and O. 151 thomaeum is more fragmented because of phylogenetic divergence and each O. thomaeum 152 region has between 2-4 homeologous regions in *M. caffa* (Supplemental Figure 2). We 153 calculated the synonymous substitution rates (Ks) between homeologous gene pairs within M. 154 caffra to date the polyploid event(s). We observed a single Ks peak of 0.13 across all 155 homeologous gene pair combinations, suggesting the autohexaploidy event occurred ~4 million 156 years ago from rapidly successive polyploidy events (Supplemental Figure 2d). Using MCScan 157 with O. thomaeum as an anchor, we identified 18,428 syntenic orthologs (syntelogs) shared 158 among the three grasses, as well as previously published tolerant grasses Eragrostis nindensis Ficalho & Hierr<sup>28 33</sup>. These syntelogs were used to identify patterns of gene duplication 159 160 associated with desiccation tolerance across grasses and as anchor points to compare 161 expression of conserved genes across species. 162 To test for convergent evolution we characterized patterns of expansion and duplication in gene families with important roles in desiccation tolerance. The genomes of all sequenced 163 resurrection plants have large tandem arrays of early light induced proteins (ELIPs)<sup>34</sup>, and we 164

165 observed this same pattern across the desiccation tolerant grasses investigated here.

166 Oropetium capense, T. minimus, and M. caffra all have massive tandem arrays of 39, 31, and

167 58 ELIPs respectively, compared to an average of 4 in the genomes of desiccation sensitive

168 grasses <sup>34</sup>. This expansion of ELIPs is similar to other chlorophyll retaining (homiochlorophlylus)

169 resurrection plants and is generally higher than chlorophyll degrading (poikiochlorophyllus)

species. ELIPs are universally highly expressed in the diploid resurrection grasses *O. capense* 

- and *T. minimus* during drying, desiccation, and early rehydration, but only a subset of the ELIPs
- in the *M. caffra* tandem arrays have desiccation induced expression (Supplemental Figure 3).
- 173 We used CAFE <sup>35</sup> to test for changes in the dynamics of ELIP copy number evolution across
- 174 land plants. We found significant increases in the rate of ELIP expansion in all desiccation
- tolerant lineages of plants (Figure 2c). Within the grass family, ELIP expansion occurred
- 176 independently in subtribes Eleusininae, Sporobolinae, Eragrostidinae, and Tripogonae, but
- 177 *Oropetium* and *Tripogon* share a single origin of desiccation tolerance (Figure 2c). Other gene
- families with well-characterized roles in desiccation tolerance such as late embryogenesis
   abundant (LEAs) and heat shock proteins (HSPs) show no expansion in resurrection plants
- 180 based on OrthoFinder and or CAFE (Supplemental Figures 4-7).

181 We identified the origin of duplicated ELIPs to test if the same or different ancestral 182 copies were duplicated in each lineage using a synteny based approach. Tandem duplication of 183 ELIPs within the Tripogoninae occurred on Chromosome 8, and the Eleusininae and 184 Eragrostidinae subtribe species have no syntenic ELIPs in this region, despite otherwise high 185 collinearity (Figure 2a). Most ELIPs in Eleusininae and Eragrostidinae species are found in large 186 tandem arrays on Chromosome 7, compared to 4-5 ELIPS within Tripogoninae (Figure 2b). 187 Together, phylogenetic and comparative genomics analyses suggest these grass lineages 188 duplicated ELIPs independently, supporting the convergent evolution of desiccation tolerance 189 within Chloridoideae.

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# 2 Searching for overlapping signatures of desiccation tolerance

193 We collected dehydration and rehydration timecourses of O. capense, T. minimus, and 194 M. caffra plants under similar conditions in a climate controlled growth chamber. Plants reached 195 desiccation after ~17-20 days of natural drying, with a relative water content (RWC) < 10% and 196 photosystem II efficiency ( $F_v/F_m$ ) approaching 0.0 (Figure 1b-d). RWC and  $F_v/F_m$  recovered 197 within 12 hours of rehydration in O. capense and T. minimus, but  $F_v/F_m$  took longer to recover in 198 M. caffra (Figure 1b). We collected gene expression data (RNAseg) at six comparable 199 timepoints of drying and recovery for each of the three species. We quantified RNA abundance 200 and gene expression patterns across the dehydration-rehydration timecourse in each species 201 individually. RNAseg reads were pseudo-aligned to the respective genomes using Salmon (v 1.9.0) <sup>36</sup> and normalized counts were used for all downstream analyses. In general, gene 202 203 expression profiles were tightly associated with the hydration status of the plants. Correlation 204 matrices and principal component analysis (PCA) show tight clustering of samples by hydration 205 status, with hydrated, desiccated, and rehydrated samples forming distinct clusters for each 206 species (Supplemental Figure 8).

Using RWC as a covariate, we identified genes that were significantly up- and downregulated during dehydration and rehydration processes. Both dehydration and rehydration induced substantial changes in gene expression in all three desiccation tolerant grasses, with 35-52% of genes showing differential abundance during dehydration and 23-47% during rehydration (Figure 3a and Supplemental Figure 9). *Microchola caffra* had significantly more differentially expressed (DE) genes (Supplemental Figure 9) given its hexaploidy, but a lower proportion of DE genes compared to the other two grasses (Figure 3a). Broadly, desiccation and rehydration had inverse expression profiles, and most genes that increased in abundance
 during dehydration, dissipated during rehydration and vice versa (Supplemental Figure 9).

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217 To enable comparisons across species, we leveraged the 18,428 conserved syntelogs 218 and searched for overlapping patterns in the expression of these shared genes. There was 219 considerable overlap in gene expression across the three focal resurrection grasses, with ~18-220 24% of all DE syntelogs showing similar expression across species (Figure 3b and 221 Supplemental Figure 10a). The proportions of DEGs shared across the three resurrection 222 grasses for both up- and down-regulated genes was considerably more than observed in 223 previous studies or expected due to chance. In order to differentiate between desiccation 224 tolerance mechanisms and more general drought responses, we identified the extent of shared syntelog expression between these resurrection grasses and the desiccation sensitive species 225 E. tef, which was sampled along a similar dehydration timecourse in a previous study<sup>28</sup>. There 226 227 was considerable overlap in syntelog expression between the resurrection grasses and E. tef 228 (Supplemental Figure 11), reflecting deeply conserved mechanisms of drought tolerance in 229 grasses. We also detected a large set of genes that were expressed exclusively in the 230 resurrection grasses, which likely play desiccation specific roles to survive anhydrobiosis. 231 Species-specific expression patterns are also evident, particularly for E. tef.

232 Dimensionality reduction and co-expression analyses also point towards parallel 233 mechanisms of desiccation tolerance in resurrection grasess. Samples clustered primarily by 234 hydration status and secondarily by species in PCA (Figure 3c,d and Supplemental Figure 12). 235 We defined co-expression modules for each species and screened for shared network level 236 responses within co-expressed genes. High confidence modules were defined for each species, 237 and we grouped these into three broad classes based on the expression pattern of each 238 module: 1) elevated expression in hydrated conditions, 2) elevated expression during 239 dehydration, and 3) elevated expression during rehydration (Figure 4d). We identified 240 substantial overlap in gene module conservation with elevated expression during dehydration, 241 but less overlap in modules with high expression during rehydration (Figure 4a). We then 242 identified enriched gene ontology (GO) terms for each co-expression module and performed 243 hierarchical clustering on the enrichment p-values of GO terms. Modules clustered by their 244 expression profile rather than species identity, suggesting that hydration status is more 245 predictive of gene expression than species identity (Figure 4c) and pointing towards a shared 246 signature of desiccation tolerance in resurrection grasses.

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# 249 Functional characterization of the shared signatures of desiccation tolerance

250 Our analyses of syntelog expression tested for ancestral conservation and parallelism, 251 but it is also possible that different lineages of resurrection plants may utilize similar metabolic 252 strategies for achieving desiccation tolerance but through divergent genes and pathways. To 253 investigate this possibility, we used the Kyoto Encyclopedia of Genes and Genomes (KEGG) to 254 assign each gene to a predicted enzymatic function and metabolic pathways and compared the 255 overlap in these functional predictions across species. We detected substantially higher overlap 256 in KEGG terms across species (~30-40%) compared to DE syntelogs (only 18-24%) (Figure 5b 257 and Supplemental Figure 10c). The increased similarity at a metabolic level suggests that while

these species do not always leverage parallel gene copies, they induce similar metabolic
 mechanisms to survive anhydrobiosis, providing evidence of convergence across species.

260 We further investigated the functional roles of shared gene expression via GO 261 enrichment and KEGG analyses. We found that many hallmarks of desiccation tolerance were 262 shared across the three resurrection grasses, including the controlled downregulation of 263 photosynthesis and rapid induction of protective mechanisms. Enriched GO terms during 264 dehydration were related primarily to signaling and stress responses (e.g., stress perception 265 and ROS scavenging activities), developmental regulation (e.g., photoperiodism and germination processes), cellular reorganization (e.g., lipid droplet formation, vesicle fusion, 266 267 endocytosis), and modifications to transcription and translation (e.g., RNA modifications, 268 splicing, and protein degradation). In contrast, enriched GO terms during rehydration are related 269 to photosynthesis and metabolism (e.g., fructose biosynthesis, cellulose biosynthesis, and light 270 harvesting), pigment metabolism (e.g., chlorophyll biosynthesis and anthocyanin metabolism), 271 protein modification (e.g., protein phosphorylation and proteolysis), and some residual stress 272 response (e.g., response to cold and non photochemical guenching) (Figure 5a). Hierarchical 273 clustering of enriched GO terms also highlighted the inverse relationship between dehydration 274 and rehydration process (Figure 5a and Supplemental Figure 10d).

275 To differentiate between desiccation tolerance mechanisms and more typical drought 276 tolerance responses, we compared the enriched GO terms for DE syntelogs uniquely induced in 277 the resurrection grasses vs. those shared with desiccation sensitive E. tef (Supplemental Figure 278 11). Many of the classic stress response terms were shared across all species, reflecting deeply 279 conserved responses to water deprivation. For example, all species showed metabolic arrest 280 during drying with a particular emphasis on photosynthetic shut down. All species exhibited an 281 increase in classic stress response terms such as response to heat, response to water 282 deprivation, response to hydrogen peroxide, and sucrose metabolic process. These processes 283 represent core mechanisms of water deficit tolerance that likely form the foundation of 284 desiccation tolerance. Building on this foundation, resurrection grasses appear to activate 285 additional processes that enable more extreme resilience. For example, the resurrection grasses showed unique activation of nucleic acid processes including mRNA export, regulation 286 287 of chromosome condensation, and mRNA transcription by RNA polymerase II suggesting greater overall regulation of transcription and translation. Several terms associated with the 288 289 circadian rhythm and hormonal signaling were also uniquely upregulated in the resurrection 290 grasses, indicating a central role of circadian clock processes in preparing for desiccation. The 291 resurrection grasses exhibited a unique downregulation of tissue and cellular developmental 292 processes, implying a tightly regulated cessation of metabolism at later stages of drying. Taken 293 together, this suggests that resurrection grasses build on a shared foundation of drought 294 tolerance to achieve desiccation tolerance via a highly organized shift in cellular processes.

295 KEGG annotations revealed characteristic desiccation tolerance mechanisms shared 296 across resurrection grasses. Metabolic pathways associated with photosynthetic energy 297 metabolism were significantly down regulated in all three grasses. Interestingly, we observed an 298 increase of malate to pyruvate catalysis with concomitant regeneration of NADPH, which could 299 be related to NADPH's REDOX potential for antioxidant enzymes such as glutathione 300 reductase. We also detected noticeable changes to carbohydrate and energy metabolism, 301 including a shift towards the production of raffinose and stachyose under dehydrating conditions as seen in other resurrection plants (reviewed in <sup>10</sup>). Central carbohydrate metabolism appeared 302

operational suggesting that at low water contents, other solvents, such as natural deep eutectic 303 304 solvents within the mitochondria may facilitate glycolysis, the TCA, and electron transport <sup>37</sup>. 305 Amino acid metabolism favored degradative pathways with an increase in endoplasmic 306 reticulum-mediated ubiquitination and proteolysis, which could be serving a glucogenic role by 307 converting amino acids to pyruvate, or by generating an available amino acid pool for the rapid 308 assembly of thermo- and osmoprotective proteins. While amino acid metabolic pathways were 309 generally down-regulated, a few important pathways including glutathione metabolism were upregulated. Reduced glutathione (GSH) exerts numerous effects in the cell <sup>38</sup> from interaction 310 with hormones to acting as direct ROS guencher, and maintaining a steady supply of GSH is a 311 312 feature all three resurrection grasses share. Lipid metabolism showed a shift towards the 313 production of glycerolipids and glycerophospholipids which likely supports triacylglycerol 314 phosphatidylcholine production. The accumulation of phosphatidylcholine may further lead to phosphatidic acid synthesis, which has been implicated in numerous plant processes from 315 signaling to storage <sup>39–41</sup>. Pathways involved in the transcription and translation of genetic 316 317 information also showed an up-regulation of transcription factors, RNA polymerase, and 318 spliceosome activity, suggesting that active transcription and RNA processing are still occurring. 319 However, we observed substantial downregulation of ribosome activity, suggesting that RNA is 320 either differentially translated or delayed. Upon rehydration, up-regulated processes involved in 321 overall resumption of normal metabolic activity such as several photosystem I and II proteins, 322 light harvesting complexes, starch synthesis, and cell wall remodeling such as xyloglucan O-323 acetyltransferase, expansin, and pectinesterase were observed.

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#### Desiccation tolerance mechanisms are broadly conserved across grasses

326 Desiccation tolerance evolved independently in at least four subtribes of Chloridiodeae 327 (Eleusininae, Eragrostidinae, Sporobolinae, and Tripogoninae; Figure 2c), and we integrated 328 comparable desiccation and rehydration expression datasets from additional species to test for 329 patterns of convergence across grasses more broadly. Building on our detailed comparisons 330 across the three study species, we expanded our analysis to include publicly available RNAseq samples from desiccation tolerant *O. thomaeum*<sup>42</sup> and *E. nindensis*<sup>28</sup>, leveraging syntelogs for 331 332 cross-species comparisons. Similar to the three species comparisons described above, 333 dimensionality reduction across the five species generally separated samples by hydration 334 status along PC1 and PC2 (Supplemental Figure 12). While PCA provided some degree of 335 separation; residual heterogeneity, experimental differences, noise, or species level differences 336 in the datasets might have obscured underlying conserved biology. To account for this, we 337 employed a topological data analysis (TDA) approach to discern the underlying structure of the 338 expression datasets. We utilized the Mapper algorithm, which condenses the dataset into a 339 scalable, navigable representation. The Mapper algorithm is particularly well-suited for genome 340 scale analyses, as the underlying datasets are often characterized by high dimensionality and sparsity <sup>43</sup>. For our gene expression data, we constructed Mapper graphs using a "stress lens" 341 342 with the well-watered condition as a reference point. This model represents the baseline for 343 gene expression and we quantified the residuals or deviation of each sample from the baseline, 344 which represent the degree of water stress or recovery.

345 The resultant Mapper graph illustrates a clear topological shape that delineates 346 desiccation processes across grasses (Figure 6). Each node on the graph represents a cluster 347 of similar RNAseq samples, and the node color depicts the identity of samples within that 348 cluster. Connections between nodes signify shared samples among the intersecting clusters. 349 These graphs reveal a compelling topological depiction of the gene expression variations 350 induced by water stress across different species. Similar topology was observed for both 351 targeted comparison of the three focal species (Figure 6a and b) and for the larger dataset 352 including E. nindensis and O. thomaeum (Figure 6c and d). In both instances, clear delineation 353 between samples of different hydration statuses are evident, while the species are intermixed. 354 We then added the desiccation sensitive sister species *E. tef* in a final analysis (Supplemental 355 Figure 13), which revealed a similar topology across all species with notable gaps in E. tef. 356 Broadly, this supports our finding that similar ancestral mechanisms are being recruited for 357 foundational drought tolerance mechanisms, which are enhanced in resurrection plants via the 358 independent recruitment of specific desiccation tolerance pathways. 359

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# Species specific mechanisms underlying desiccation tolerance

362 Despite the considerable overlap in gene expression across all three focal species, 363 species-specific processes were also evident. In M. caffra, unique antioxidant responses were 364 induced including glutathione biosynthetic processes, glutamate decarboxylation, and L-365 ascorbic acid biosynthesis. Other processes enriched uniquely in M. caffra included seed-366 related terms such as seed oil body biogenesis and seed maturation. Several GO terms 367 associated with phytohormones were also uniquely induced in *M. caffra*, including overall 368 ethylene responses such as S-adenosylmethionine metabolic process, ethylene-activated 369 signaling pathway, and response to 1-aminocyclopropane-1-carboxylic acid, suggesting that 370 hormonal regulation might be exerting an effect on the partial breakdown of thylakoids and photosynthetic machinery as seen in classical senescence <sup>44</sup>. *Microchloa caffra* also exhibited 371 372 unique lipid, sphingolipid, riboflavin, and selenocompound metabolism, as well as 373 sesquiterpenoid and tripenoid biosynthesis. Microchloa caffra was the only species to have 374 multiple pathways involved in signal transduction up-regulated including phospholipase D and 375 calcium signaling. Uniquely down-regulated processes in *M. caffra* appear to center around 376 arresting growth and development, such as phototropism, gravitropism, leaf and root 377 morphogenesis, cell wall biogenesis, and regulation of auxin polar transport. There was also a 378 down-regulation of general amino acid-tRNA aminoacylation and nitrogen fixation and 379 assimilation.

380 Oropetium capense had fewer uniquely enriched processes compared to M. caffra, but 381 some notable patterns were detected. Uniquely upregulated processes in O. capense centered 382 around histone H3 and H4 acetylation, histone H3-K9 demethylation, and histone H2B 383 ubiquitination. The relative degree of acetylation of histones is directly related to the openness of chromatin which impacts transcription in specific drought-responsive genes <sup>45</sup>. Histone 384 demethylation <sup>46</sup> and H2B ubiquitination also regulate drought responsive genes <sup>47</sup>. 385 386 Interestingly, terms associated with chloroplast mRNA processing, poly(A)+ mRNA export from 387 the nucleus, ribosome assembly, and regulation of translation were also upregulated in O. 388 capense, suggesting continued translation and active processing of mRNA from both the 389 chloroplast and nucleus, presumably through increased transcriptional regulation due to histone 390 modifications. Oropetium capense exhibited unique up-regulation of C5-branched dibasic

metabolism and down-regulation within galactose metabolism. Uniquely down-regulated
processes in *O. capense* were minimal, but included regulation of salicylic acid metabolic
process and auxin polar transport. Monoterpenoid biosynthesis was up-regulated for *O. capense* and fatty acid degradation and steroid hormone biosynthesis was down-regulated.
Ferroptosis, an iron-dependent form of programmed cell death, was exclusively down-regulated
in *O. capense*.

397 Tripogon minimus also had fewer species-specific processes compared to M. caffra. 398 Uniquely up-regulated processes were centered around response to oxidative stress, 399 peroxisome organization, and removal of superoxide radicals. Tripogon minimus was the only 400 species to show upregulation of anthocyanin-containing compound biosynthetic process which is a typical response seen in the homoiochlorophyllous resurrection plants <sup>48</sup>. Similar to the 401 402 other two species, regulation of auxin-mediated signaling pathways were down-regulated as 403 was cellular response to salicylic acid stimulus. Other processes centered around mismatch 404 repair, chloroplast RNA processing, ribosome biogenesis, and plastid transcription. 405 Phosphonate and phosphinate, taurine and hypotaurine, and D-amino acid metabolism were 406 exclusively down-regulated in T. minimus whereas retinol metabolism was up-regulated.

407 Despite the unique pathways identified in each of the focal species, all three species 408 appear to respond to drought and desiccation stress by leveraging similar mechanisms. The 409 processes uniquely activated in each species are consistently centered around defense 410 mechanisms, the induction of guiescence, and reduction of normal growth and metabolism 411 under desiccated conditions. While nuanced variation in metabolism and defense responses are 412 evident, all species exhibit well known mechanisms of desiccation tolerance. Taken together, 413 the three species appear to share a core set of conserved mechanisms which are then 414 supplemented with convergent species-specific modules.

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### 417 Discussion

418 Our data suggest that the repeated evolution of desiccation tolerance within grasses 419 occurred via both parallel adaptations in the same ancestral genes and complementary 420 modifications to analogous pathways. We find evidence that core mechanisms of desiccation 421 tolerance are shared across resurrection grasses, and are supplemented with species-specific 422 adaptations. Many of these mechanisms overlap with typical drought responses and it is likely 423 that the evolution of anhydrobiosis builds on deeply conserved responses to water deficit shared 424 across all plants. Phenotypic and metabolic similarities in anhydrobiosis mechanisms have been 425 observed for decades, but the evolutionary pathways of convergence and parallelism have been 426 obscured by a lack of systems-level data and inconsistencies in experimental procedures <sup>49</sup>. 427 Here, we leveraged large scale genomic and transcriptomic datasets in a replicated and 428 standardized framework to characterize signatures underlying the recurrent evolution of 429 desiccation tolerance within chloridoid grasses.

The adaptations required for desiccation tolerance appear to be sufficiently narrow, such
that not any organism can, or will, evolve desiccation tolerance <sup>4</sup>. The physiological changes
that occur during the final stages of desiccation are dramatic and specialized biochemistry and
molecular mechanisms are required to protect the cellular macromolecules for life without water.
Achieving anhydrobiosis requires tight coordination and orchestration of multiple physiological

processes, and there may be only a few trajectories to evolve this trait. However, desiccation
 tolerance mechanisms overlap considerably with typical drought responses and many plants

437 possess the basic cellular machinery required to achieve desiccation tolerance <sup>28</sup>. Desiccation

438 tolerance is likely an ancestral adaptation in plants that evolved during terrestrialization.

439 subsequently formed the basis of seed pathways, and was later rewired again in vegetative

tissues <sup>5,25,42,50</sup>. While previous studies have found surprisingly little overlap in gene expression
across desiccation tolerant plants <sup>51,52</sup>, our data suggest that the repeated evolution of specific
genetic, biochemical, and physiological traits required for anhydrobiosis, are highly convergent
and build on more broadly conserved water deficit responses.

444 Convergence is thought to be driven primarily by exposure to external selective 445 pressures that lead to the same emergent phenotype, while parallelism is thought to be impacted more by internal constraints of the system <sup>18</sup> through independent mutations in the 446 same ancestral gene <sup>19,21</sup>. Because anhydrobiosis has evolved independently in both distantly 447 448 and closely related taxa, it is an ideal system in which to explore the roles of convergent and 449 parallel evolution. Numerous other independently evolved traits such as C4 and CAM 450 photosynthesis are highly complex, making their repeated evolution surprising <sup>53</sup> and difficult to characterize. In the case of C4 photosynthesis, both mutations in the same genes and 451 452 recruitment of unique pathways occurred in distantly related lineages to enable the emergent C4 phenotype <sup>54</sup>. Desiccation tolerance is similarly complex, involving the synchronized 453 454 orchestration of numerous pathways and genes, and it is likely that both external pressures (e.g. 455 selection in extremely xeric habitats) and internal constraints (lineage specific predispositions) 456 play a role in the recurrent evolution of desiccation tolerance. Here, we detected signatures of both processes and identified far more overlap in gene expression across resurrection grasses 457 than expected by chance or detected in previous studies <sup>29,51</sup>. The observed expansion of ELIP 458 459 tandem arrays coupled with activation of similar metabolic pathways driven by different gene 460 sets, suggests that both parallel and convergent processes contribute to the recurrent evolution 461 of desiccation tolerance in grasses.

462 Our systems-level analyses add to the growing literature on the mechanisms of 463 desiccation tolerance, and many of the patterns observed here corroborate previous findings <sup>17,25,29,34,51</sup>. We show that desiccation induces a major and reversible shift in gene expression 464 where normal growth and development are halted and numerous protective mechanisms are 465 induced <sup>13,14,55–57</sup>. Gene expression coalesced around a signature desiccation response during 466 drying with all three species initiating parallel processes <sup>58</sup>. The resumption of species specific 467 processes related to growth and development was evident upon rehydration. The shared 468 469 pathways of anhydrobiosis observed in these grasses pull on the deeply conserved architecture 470 of drought tolerance coupled with convergent and parallel mutations that provide the necessary 471 protection to survive extreme desiccation. This reflects the relatively narrow set of regulatory 472 networks and pathways in plants that can enable the evolution of desiccation tolerance, but also 473 hints as multiple evolutionary paths to anhydrobiosis 474

- 474
- 476 Methods
- 477
- 478 Field collections, plant growth, and maintenance

479 Plants for the current study were collected from two research sites in South Africa: 480 Buffelskloof Nature Reserve in Mpumalanga (-25.30229 S, 030.50631 E) (Microchloa caffra) 481 and Swebe Swebe Private Wildlife Reserve in Limpopo (-23.7949 S, 028.0705 E) (Oropetium 482 capense and Tripogon minimus). Voucher specimens of each species were collected, pressed, 483 and deposited at the National Herbarium of South Africa in Pretoria (specimen numbers: 484 PRE1004810-0, PRE1004793-0, and PRE1004794-0). Seeds of each species were also 485 collected and transported to Michigan State University under United States Department of 486 Agriculture (USDA) permit #537-22-37-10071 and according to the specifications in a Material 487 Transfer Agreement established between Drs. Jill M. Farrant, Robert VanBuren, and Rose A. 488 Marks. Seeds were cold stratified at 4°C for two weeks and then germinated on our standard 489 propagation mix (50:50 sure-mix:redi-earth) and grown in a climate controlled growth chamber 490 with a 16 hour photoperiod and internal temperature of 28/18°C. Six weeks after germination, 491 individual seedlings were transplanted into separate pots and grown to maturity and a single 492 plant (genetic line) of each species was selected for downstream experimentation. Seeds of 493 each genetic line were collected, cold stratified, and ~70 seedlings from each species were 494 germinated. One seedling from each species was used for genome sequencing and was 495 transplanted into a larger pot. The remaining seedlings were used for the desiccation and 496 rehydration timecourses experiments and three seedlings were transplanted into 4" pots. The three plants in each pot were pooled during sampling and treated as a single biological 497 498 replicate. These plants were grown for another two weeks prior to experimental treatments, 499 during which time they were maintained in constantly hydrated conditions in a growth chamber 500 set to the conditions described above.

501

### 502 Dehydration treatment and sample collection

503 After ~8 weeks of growth, plants were subjected to dehydration treatment. Prior to 504 treatment, any emerging reproductive tissues (e.g., panicles) were removed from plants. To 505 initiate dehydration treatment, plants were watered to full soil saturation and each pot was 506 weighed to ensure consistency across replicates. Water was then withheld until plants became 507 completely desiccated (between 2 and 3 weeks depending on the species). Plants were 508 sampled at targeted hydration states during the process of dehydration, including well watered, 509 partially dehydrated, fully desiccated, and rehydrated. We used visual cues to direct our 510 sampling and sampled plants at the first signs of visible leaf curling, partial pigmentation, deep 511 pigmentation, and full desiccation and validated the hydration status of tissues by measuring 512 relative water content (RWC). Plants were then rehydrated through a combination of watering 513 from the base and misting the aerial portions to simulate natural rainfall and sampled 24 and 48 514 hours post rehydration. We aimed to sample plants at biologically relevant water contents and 515 therefore directed our sampling by use of visual cues rather than a set number of hours. This 516 allowed us to compensate for different drying rates across species and plants due to subtle 517 variation in size, water use efficiency, and relative humidity in the growth chamber.

518 At each timepoint, we measured the photosynthetic efficiency ( $F_v/F_m$ ) and RWC, and 519 harvested tissue for RNAseq. Briefly,  $F_v/F_m$  was measured on dark adapted leaves using a Opti-520 Sciences OS30p+ chlorophyll fluorometer with the default test parameters. Relative water 521 content was measured using a set of 10-15 representative leaves from each pot / biological 522 replicate. Leaf mass was weighed immediately after collection (fresh weight), again after 48 hours submerged in dH20 in darkness at 4°C (turgid weight), and finally after 48 hours in a 70°C
drying oven (dry weight). RWC was calculated as (fresh weight - dry weight)/(turgid weight - dry
weight). Tissue for RNAseq was collected by harvesting all the vegetative tissue from each pot
and flash freezing in liquid nitrogen. Tissue samples were stored in a -80°C freezer prior to
downstream processing.

528

# 529 RNA extraction and sequencing

530 Frozen leaf tissue was ground to a powder by hand in a mortar and pestle with liquid 531 nitrogen. RNA was extracted from each sample using Spectrum Plant Total RNA kit according 532 to the manufacturer instructions. Total RNA was then cleaned to remove impurities and 533 contaminants using Zymo Clean & Concentrator kit. DNAse treatment was carried out during 534 clean and concentration steps according to manufacturer instructions. Sample concentration 535 was assessed on a qubit using the RNA broad range reagent set, purity was assessed with a 536 nanodrop, and RNA integrity was visualized on an agarose gel. RNAseq libraries were 537 constructed by Novogene following a standard polyA+ enrichment strategy including 538 fragmentation and cDNA synthesis. The resulting libraries were sequenced on an Illumina 539 HiSeq 4000 under 150 bp paired end mode

540

# 541 High molecular weight DNA extraction, and sequencing

542 Tissue for whole genome sequencing was collected from a single mature plant of each 543 species. Healthy green tissue was harvested and flash frozen in liquid nitrogen. Tissue was 544 ground by hand in a mortar and pestle for >20 minutes to liberate nuclei. Pure, high molecular 545 weight genomic DNA was extracted by first isolating nuclei with the Circulomics Nuclei Isolation 546 kit and then extracting DNA with the Circulomics Nanobind Plant Nuclei Big DNA kit. HiFi 547 libraries were constructed from the Genomic DNA and sequenced at the University of Georgia 548 Sequencing Core on a PacBio Sequel II machine.

549

# 550 Genome assembly

551 We used flow cytometry to estimate genome sizes (2C DNA values) for the three 552 grasses. Healthy leaf tissue was harvested from each genotype. Nuclei were isolated and 553 stained according to standard protocols. The stained nuclei were then run on a BD Accuri<sup>™</sup> C6 554 Plus Flow Cytometer at Plantploidy.com. Hosta plantaginea was used as an internal reference. 555 We built reference genomes for each species using high fidelity (HiFi) PacBio long read data. In 556 total, 70.1 Gb of HiFi reads were generated for *M. caffra*, 15.9 Gb for *O. capense*, and 20.2 Gb 557 for T. minimus, representing 56, 82, and 103 x genome coverage for each species respectively. K-mer analysis revealed that O. capense and T. minimus have low within genome 558 heterozyogisity and *M. caffra* is a highly heterozygous autopolyploid <sup>59</sup>. PacBio reads were 559 assembled using hifiasm (v 0.18) <sup>30,60</sup> with default settings for *O. capense* and *T. minimus* and 560 the number of haplotypes was set to 6 for *M. caffra* (flag: --n-hap 6). The resulting assemblies 561 562 were highly contiguous with six and nine of the ten chromosomes assembled telomere to 563 telomere for T. minimus and O. capense respectively, and 118 contigs across 968 Mb with an N50 of 16 Mb for *M. caffra* (Table 1). Raw assemblies were filtered for non-plant contigs using a 564 representative microbial database with BLAST <sup>61</sup>. Full length chloroplast and mitochondrial 565

566 genomes were identified and retained, and any additional partial or rearranged organelle 567 genomes were removed.

568

#### 569 Genome annotation

570 A library of repetitive elements was constructed for each of the three grass genomes using the EDTA package (v2.0.0)<sup>62</sup>. EDTA comprehensively identifies DNA-based transposable 571 elements using HelitronScanner<sup>63</sup>, and LTR retrotransposons using LTR FINDER<sup>64</sup> and 572 LTRharvest <sup>65</sup>. Protein coding genes were annotated using the MAKER-P pipeline (v2.31.10) <sup>66</sup> 573 574 with the following sets of input data for training. Transcript evidence was generated using the dehydration-rehydration timecourse RNAseq data from leaf tissue of each species described 575 below. Raw RNAseg reads were guality trimmed using fastp (v 0.23)<sup>67</sup> and aligned to the 576 unmasked genomes using the splice aware alignment program STAR (v2.6)<sup>68</sup>. A set of non-577 overlapping transcripts was identified from the aligned data using StringTie (v1.3.4)<sup>69</sup> with 578 579 default parameters. The resulting gff files were used as transcript evidence for MAKER. The 580 same protein evidence was used as training for each of the three grasses and this includes the full annotations of Oryza sativa <sup>70</sup>, Arabidopsis thaliana <sup>71</sup>, Oropetium thomaeum <sup>24,32</sup>, and 581 Eragrostis tef<sup>33</sup>. These datasets were used as input for MAKER and we utilized SNAP<sup>72</sup> and 582 Augustus (version 3.0.2)<sup>73</sup> for *ab initio* gene prediction, performing two rounds of iterative 583 584 training to refine our models. To filter out repetitive element-derived proteins, we used BLAST 585 using a non-redundant transposase library against the raw gene models produced by MAKER. 586 We assessed the completeness of our assembly using the plant-specific embryophyte set of Benchmarking Universal Single-Copy Orthologs (BUSCO v.2)<sup>74</sup>. These high-confidence gene 587 588 models were used for all downstream analyses.

589

### 590 Comparative genomics

591 The three desiccation tolerant grass genomes were compared to each other and other Chloridoid grasses using the MCScan toolkit (v1.1)<sup>75</sup> implemented in python 592 593 [https://github.com/tanghaibao/jcvi/wiki/MCscan-(Python-version)]. Syntenic orthologs were 594 identified across the three focal species, E. nindensis, E. tef, and Oropetium thomeaum using 595 the chromosome-scale O. thomeaum genome as an anchor. Syntenic blocks were identified 596 using gene models aligned using LAST with a minimum of five overlapping syntenic genes. The 597 macrosyntenic dot plots, histograms of depth, and microsynteny plots were generated using the 598 python version of MCScan. A set of 18,428 conserved syntenic orthologs across all six 599 desiccation tolerant grasses was created and used for downstream comparative genomic and 600 cross-species transcriptomic analyses. We identified orthologous genes across a subset of 33 601 land plant species to search for patterns of gene family expansion in desiccation tolerant 602 lineages as well as for downstream comparative genomic analyses. We included the following 603 species with desiccation tolerant species highlighted: Ananas comosus, Arabidopsis thaliana, 604 Brachypodium distachyon, Eleusine coracana, Eragrostis curvula, Eragrostis nindensis (DT), 605 Eragrostis pilosa, Eragrostis tef, Hordeum vulgare, Lindernia brevidens (DT), Lindernia 606 subracemosa, Microchloa caffra (DT), Marchantia polymorpha (DT), Medicago truncatula, 607 Oropetium capense (DT), Oryza sativa, Oropetium thomaeum (DT), Physcomitrium patens 608 (DT), Sorghum bicolor, Setaria italica, Selaginella lepidophylla (DT), Solanum lycopersicum, 609 Selaginella moellendorffii, Sporobolus pyramidalis, Sporobolus stapfianus (DT), Setaria viridis,

610 Triticum aestivum, Tripogon minimus (DT), Vitis vinifera, Xerophyta viscosa (DT), Zostera

611 *japonica, Zostera marina, and Zea mays.* Proteins were clustered into orthologous groups using

612 Orthofinder (v2.2.6)<sup>76</sup> with default parameters. For the orthogroup enrichment analysis, we

calculated a Z-score for each species within each orthogroup, compared it to a normal

distribution to obtain a p-value, and then adjust these p-values using the Benjamini and

Hochberg procedure to get q-values. We then searched for statistically enriched orthogroups

across all of the sequenced desiccation tolerant grasses. Using this approach, we identified

617 between 486 and 8,863 enriched orthogroups in the 33 species we included in our analysis, and 618 found none that are conserved across all desiccation tolerant grasses outside of ELIPs.

619

# 620 ELIP gene family evolution

To test the hypothesis that the ELIP gene family expansions are associated with the 621 evolution of desiccation tolerance, we used CAFÉ (v 5.1)<sup>35</sup>, which analyzes changes in gene 622 family size in a phylogenetic framework. The input tree was created from the amino acid 623 624 sequences from 36 land plant species, with a focus on Chloridoid grasses (ELIPs count 625 phylogeny figure; Supplemental Table 1). Sequences were first clustered using Orthofinder (v 2.4.1)<sup>76</sup>, filtered to remove any orthogroups that did not contain all taxa, and aligned using 626 MAFFT (v 7.305b) <sup>77</sup>. No single copy orthologs were found containing all taxa for species tree 627 628 construction. Instead, we pruned gene trees and alignments to the largest subtree containing 629 unique taxa using PhyloPyPruner (v 1.2.4) (https://gitlab.com/fethalen/phylopypruner); where 630 paralogs were monophyletic within a species, we randomly pruned all but one sequence prior to 631 extracting the largest subtree. The resulting pruned gene trees and alignments were further 632 filtered to remove any trees no longer containing at least 19 taxa. This final set of 195 alignments were concatenated and used to construct a phylogeny using IQ-TREE (v 2.3.0) 78 633 634 and time calibrated fast least-squares dating <sup>79</sup>.

ELIP gene family counts per haploid genome for non-focal taxa were done using BLASTP with the *Arabidopsis thaliana* (L.) Heynh. ELIP1 amino acid sequence as query for the remaining proteomes. We further investigated two other gene families with known roles in desiccation tolerance – heat shock proteins (HSPs) and late embryogenesis abundant proteins (LEAs) – along with 20 random selected orthogroups, to contextualize the tempo of ELIP evolution. These count data and the time calibrated phylogeny were used as input for CAFÉ under a single lambda model.

642

# 643 Transcriptomic analyses

644 RNA sequencing reads were processed following a pipeline developed by the VanBuren 645 Lab (https://github.com/pardojer23/RNAseqV2). Briefly, sequence read quality was assessed with fastQC (v 0.23) and reads were trimmed with trimmomatic (v 0.38)<sup>80</sup> to remove adapters 646 647 and low quality bases. Trimmed reads were sudo-aligned to reference genomes using Salmon (v 1.9.0)<sup>36</sup>, and the resulting quantification files were processed with tximport (v 3.18)<sup>81</sup> and to 648 649 generate normalized expression matrices of transcripts per million (TPM). A Principal 650 Component Analysis (PCA) was used to visualize replicate and sample relationships within 651 each species using the respective TPM expression values. A cross-species PCA was performed 652 using the TPM matrix of conserved syntenic orthologs across all species. To effectively quantify 653 gene expression while acknowledging the complexities introduced by polyploidy, we summed

654 the expression levels of all homeologs in *E. nindensis*, *E. tef*, and *M. caffra* to obtain a single 655 gene expression value to enable interspecies comparisons. This approach is grounded in the 656 logic that a unified expression value not only simplifies the analysis but also encapsulates 657 potential functional diversifications among homeologs. This methodology has been applied and 658 validated in our previous research <sup>17,29,82,83</sup>. We then computed the Z-score of the TPM for each 659 syntenic ortholog in each sample and ran PCA on the matrix of Z-scores of each syntenic 660 ortholog across all timepoints and species.

661

# 662 Differentially expressed genes

Differentially expressed genes (DEGs) were identified independently for each species 663 with DEseg2 R package (v 1.42.0)<sup>84</sup>. Briefly, transcript abundance estimates from Salmon were 664 665 imported into DEseq2 using tximport to generate counts matrices. Count matrices were 666 normalized, and hierarchical clustering was conducted for basic quality control and visualization 667 of relationships across experimental timepoints and biological replicates. We tested multiple 668 models for differential expression in DEseq2, including models that identified DEGs by pairwise 669 comparisons of each timepoint against well-watered, and models that used the continuous 670 variables of RWC or  $F_{v}/F_{m}$  as covariates. DEGs identified by pairwise comparisons were 671 summarized into a nonredundant list of up and down regulated genes during dehydration and 672 rehydration. DEGs identified using the continuous variables are based on a significant linear 673 association (positive or negative) with RWC or  $F_{v}/F_{m}$ . When identifying DEGs, we included the 674 term "process" in our model to differentiate between dehydration and rehydration processes. To 675 select the best performing model, we quantified similarities and differences in the number and 676 identity of DEGs defined by each model. There was a high degree of overlap in genes identified 677 by all three models. Ultimately, we selected the model based on RWC because it performed 678 well and is easily comparable across experiments regardless of sampling time, consistency 679 across replicates, or differences in experimental design. These analyses produced species-680 specific lists of DEGs during dehydration and rehydration with significant (FDR adjusted P-value 681 <0.05) associations with RWC. Log2foldchange values are calculated for one unit change in 682 RWC.

683 To gain insight into possible similarities and differences among the study species, we 684 looked at the overlap in DE syntenic orthologs. To do so, we used venn diagrams to identify the 685 shared syntelogs in up- and down-regulated genes during both dehydration and rehydration 686 across the three study species. We then compared the observed proportion of overlapping 687 DEGs in each category to the proportion of genes expected to overlap by chance (assuming 688 independent draws), and tested if these were significantly different using Fisher's exact test. 689 This analysis was then extended to include DEGs identified in the desiccation sensitive sister 690 species *E. tef* to distinguish between typical drought vs. pure desiccation responses. We then 691 conducted targeted analyses to look at the functional roles of DE syntelogs that were uniquely 692 shared across the three resurrection species vs. those that were common with E. tef. We also 693 investigated the functional signatures of differentially abundant transcripts that were unique to 694 each species. To do so we pulled the lists of DE syntelogs that were only found in one of the 695 focal species.

696

697 Functional annotation of DEGs

We annotated DE syntelogs with KEGG and GO terms to describe generalized
 metabolic and cellular processes responses shared across in the three study species. KEGG
 annotations were generated for each species using BLASTKoala

- 701 (https://www.kegg.jp/blastkoala/) on the complete set of annotated peptide sequences. These
- 702 KEGG terms were then assigned to syntenic orthologs and DE KEGG terms that were shared
- across all three species during dehydration and rehydration were identified and plotted using
- venn diagrams. These shared DE KEGG terms were used to generate metabolic pathway maps
   using the reconstruct function of the KEGGmapper tool
- 706 (https://www.genome.jp/kegg/mapper/color.html) for up- and down-regulated terms in
- 707 dehydration and rehydration. This returned a list of syntelogs per metabolic pathway and Brite
- descriptions. The difference between the number of syntelogs assigned to each pathway for up
- and down regulated genes was computed. The list was then sorted to determine pathways that
- 710 were primarily up or down regulated. Next, the assigned KO numbers were paired with
- 711 quantitative data on gene expression to identify which pathways were active at various
- timepoints. To summarize patterns, genetic information processes (transcription, translation,
- folding, sorting, and degradation, replication and repair, and processing in viruses) were
- 714 grouped together. Similarly Environmental information processing (membrane transport and
- signal transduction) and cellular processes (transport and catabolism, cell growth and death,
- cellular community- prokaryote and eukaryotes, and cell motility) were grouped together.
- 717 Pathways assigned to Organismal systems and Human disease were ignored. KEGG
- annotation is not without its limits as single KEGG identifiers can be present in multiplepathways.

720 GO terms were assigned through homology with the well annotated genome of sister species O. thomaeum<sup>26</sup>. This was done through a BLASTP (v 2.14.0)<sup>85</sup> search of all O. 721 722 thomaeum protein sequences against the protein sequences of each study species. Parameters 723 were set to return the single best match for each peptide and an e-value cutoff of 1e-10. We 724 assigned the GO terms from O. thomaeum to the homologous genes in our target species. We 725 then used TopGO R package (v 2.54.0) to identify significantly enriched GO terms (P-726 value<0.05) within sets of DEGs for up- and down-regulated genes during dehydration and 727 rehydration in each target species and for the different sets of overlapping and unique syntelogs 728 identified via cross-species comparisons.

729

### 730 Co-expression analyses

To complement the above analyses, we generated co-expression networks using Weighted Gene Co-expression Network Analysis (WGCNA) R package (v1.7) <sup>86</sup>. While DE analyses can be informative to identify and describe overarching patterns and large shifts in the data, more nuanced patterns of gene expression can be obscured. To investigate the more subtle temporal changes in gene expression, we used co-expression analyses to identify modules of co-expressed genes for each species.

For each species, we created a signed co-expression network using WGCNA. Each dataset was filtered to remove genes with no expression. To construct a weighted coexpression network, we determined a soft thresholding power for each dataset. This power was chosen to satisfy WGCNA's assumption that a weighted co-expression network is scale-free. An adjacency matrix, representing the strength of connections between genes in the network, was constructed for each network using the soft thresholding power. For module detection, this
matrix was then converted to a topological overlap matrix (TOM) and hierarchal clustering was
used on the TOM to group genes into modules based on similar expression patterns.

Additionally, we calculated connectivity of each gene within its network and its assigned moduleusing WGCNA's network analysis functions.

We identified shared and species-specific co-expressed genes using UpSet plots <sup>87</sup>. For all co-expressed genes, we identified the syntenic orthologs and computed the overlap across species. We combined all modules for a species that showed increased expression during dehydration, during rehydration, and under non-stressed conditions. We identified the sets of shared syntelogs as well as those that were only found in a single species. We then ran GO enrichment analysis of these sets of shared and unique co-expressed genes.

753

# 754 Topological data analysis

755 We employed a topological data analysis (TDA) approach following the pipeline 756 described at https://github.com/PlantsAndPython/plant-evo-mapper to discern the underlying 757 structure of the expression datasets. We utilized the Mapper algorithm, which condenses the 758 dataset into a scalable, navigable representation. The Mapper algorithm is particularly well-759 suited for genome scale analyses, as the underlying datasets are often characterized by high 760 dimensionality and sparsity. For our gene expression data, we constructed Mapper graphs 761 using a "stress lens" formulated by applying a linear model using the well-watered condition as 762 a reference point. This model represents the baseline for leaf expression and we quantified the 763 residuals or deviation of each sample from the baseline, which represents the degree of water 764 stress or recovery. We generated three different mapper graphs, one was constructed using the 765 syntelog expression matrix from just the three focal resurrection grasses; M. caffra, O. capense, 766 and T. minimus. The second mapper graph was constructed using the syntelog expression 767 matrix that included two additional resurrection grasses (E. nindensis and O. thomaeum) and 768 the third graph included the desiccation sensitive species E. tef. For the mapper graph, we 769 specified different intervals and overlap for the 3 species comparisons and the 5 species 770 comparisons. For the three species, we specified 110 intervals with a 90% overlap and for the 5 771 species comparison we specified 120 intervals with 95% overlap.

772

773 Acknowledgements: This work was funded by NSF IOS-PRFB-1906094 to RAM, DBI-774 2213983 to RAM and RV, and MCB-1817347 to RV. We thank landowners and stewards Pieter 775 and Jennie Pretorius, Ken Maude, Pieter and Nadine Vervoort, and Wayne and Messiah 776 Mudenda for assistance with field logistics and permission to collect plants. We thank the 777 University of Cape Town for access to facilities, Prof. Jill Farrant for her guidance, and Keren 778 Cooper for logistical support. We also thank the South African National Herbarium, Pretoria for 779 assistance identifying and vouchering of specimens and the United States Department of 780 Agriculture for providing import permits.

781

Author contributions: RAM and RV conceived of the study. RAM collected and curated data.
 RAM, LVP, JS, ISG, and RV conducted data analyses and contributed to data interpretation and
 conceptual framing of the manuscript. RAM and RV drew the figures. RAM, LVP, and RV wrote

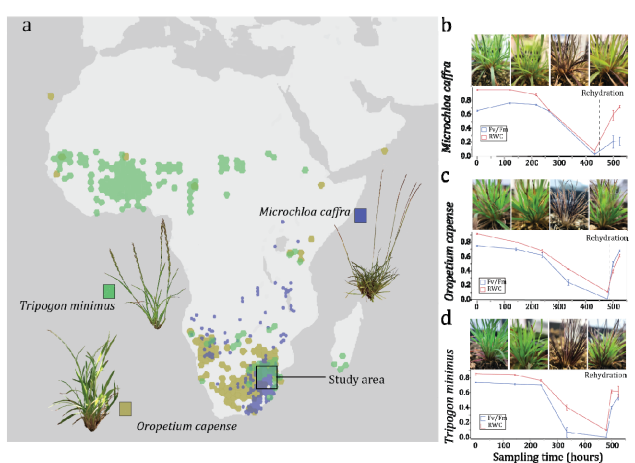
the manuscript. All authors edited and reviewed the manuscript.

786

- 787 Data availability: Sequence data associated with this study are deposited at NCBI under
- 788 BioProject PRJNA1044305 and BioSamples SAMN38380430-92. Genome assemblies are
- hosted on CoGe (<u>https://genomevolution.org/</u>) under the following IDs: 65089 (*T. minimus*),
- 790 65046 (O. capense), and 64494 (M. caffra). Metadata and other data summaries associated
- with this study are provided at Dryad https://doi.org/10.5061/dryad.kh18932c4.







#### 795 796 Figure 1. Overview of species distribution and experimental design to test for convergent evolution in

797 grasses. (a) Estimated distribution of the three desiccation tolerant grasses Microchloa caffra, Oropetium capense, 798 and Tripogon minimus. Distribution data were taken from GBIF.org (21 November 2023) GBIF Occurrence Download 799 https://doi.org/10.15468/dl.5jf47y. Collections for the current study were made in Mpumalanga and Limpopo 800 provinces of South Africa. Relative water content and F<sub>v</sub>/F<sub>m</sub> of plants during dehydration and rehydration timecourses 801 for (b) M. caffra, (c) O. capense, and (d) T. minimus. Three biological replicates were sampled at each timepoint for 802 each species. Error bars represent standard error of the mean.

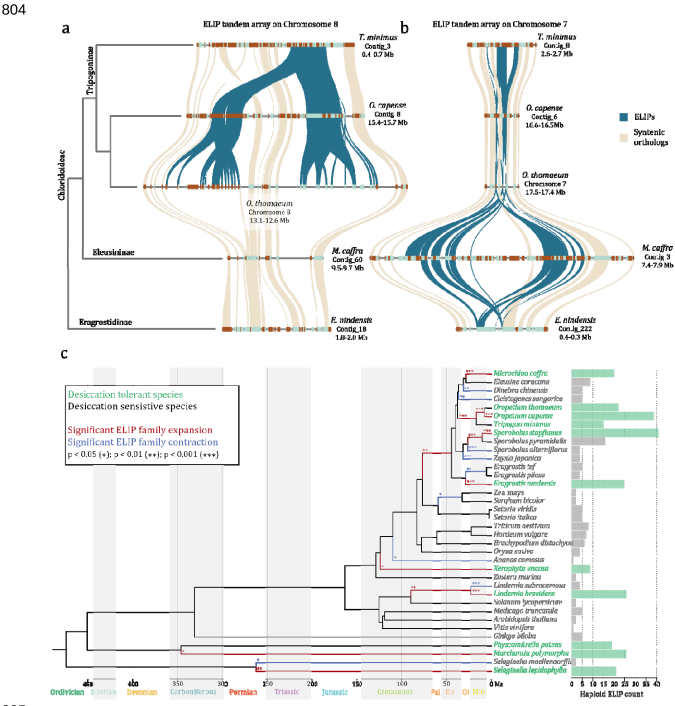




Figure 2. Independent tandem gene duplication of ELIPs in different resurrection grass lineages.

807 Microsyntenic regions of the Chromosome 8 (a) and Chromosome 7 (b) ELIP tandem arrays is shown for resurrection

grasses in the Tropogoninae (*T. minimus*, *O. thomaeum*, and *O. capense*), Eleusininae (*M. caffra*), and

809 Eragrostidinae (*E. nindensis*) subtribes of Chloridoideae. Syntenic orthologs between the species are shown in beige

and the ELIPs are highlighted in blue. Only a single syntenic region for autopolyploids *M. caffra* (hexaploid) and *E.* 

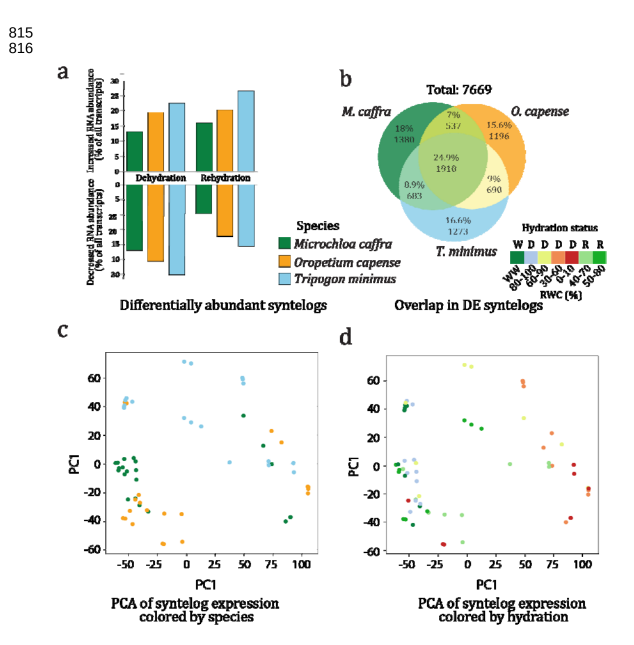
811 *nindensis* (tetraploid) is shown for simplicity, but each of the other haplotypes contain the same gene content in these

812 regions. (c) Evolutionary dynamics showing significant changes in the rates of gene family expansion (red) and

813 contraction (blue) of ELIPs inferred by CAFE. The haploid normalized number of ELIPs are plotted for desiccation

tolerant (green) and sensitive (gray) species.

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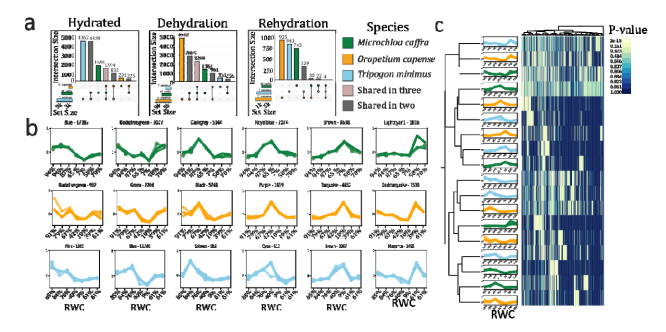


817

**Figure 3. Overlapping expression dynamics of conserved genes across species.** (a) Barplot showing the percentage of DEGs in each species for up- and down-regulated genes in dehydration and rehydration conditions. (b) Venn diagram showing the number of syntenic orthologs that increased in abundance during dehydration and overlap across species. The percentage and number of genes in each set are shown. (c,d) Principal component analysis of zscore transformed expression values for conserved syntelogs across all three species. Samples are colored by hydration status in (c) and by species identity in (d).

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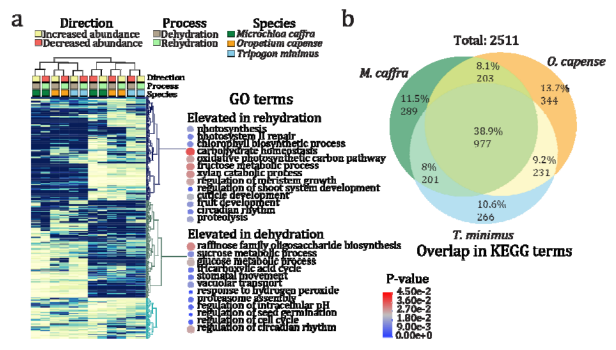
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826 827

Figure 4. Comparative co-expression network dynamics across resurrection grasses. (a) UpSet plots showing the number of shared and unique syntenic orthologs among co-expression modules characterized by elevated expression in hydrated, dehydrated, or rehydrated conditions. (b) Co-expression modules for each species. The Xaxis shows the approximate relative water content (RWC) of samples at each timepoint. The module name and total number of genes in the module are listed above. (c) Hierarchical clustering of enriched GO terms for each coexpression module. Secondary clustering performed on modules shows that modules are organized by expression profile rather than species.





836 837

Figure 5. Overlapping gene functions during desiccation and rehydration in resurrection grasses. (a)

838 Hierarchical clustering of p-values for enriched GO terms for each species and condition. Selected GO terms are

839 highlighted for genes that increased in abundance during dehydration (decreased in abundance during rehydration)

and genes that increased in abundance during rehydration (decreased in abundance during dehydration). Points are

colored by the average enrichment p-value across all species and sized by the number of genes assigned to that GO
 term. (b) Venn diagram showing the number of overlapping KEGG terms that increased in abundance during

843 dehydration. The percentage and number of KEGG terms in each set are shown.

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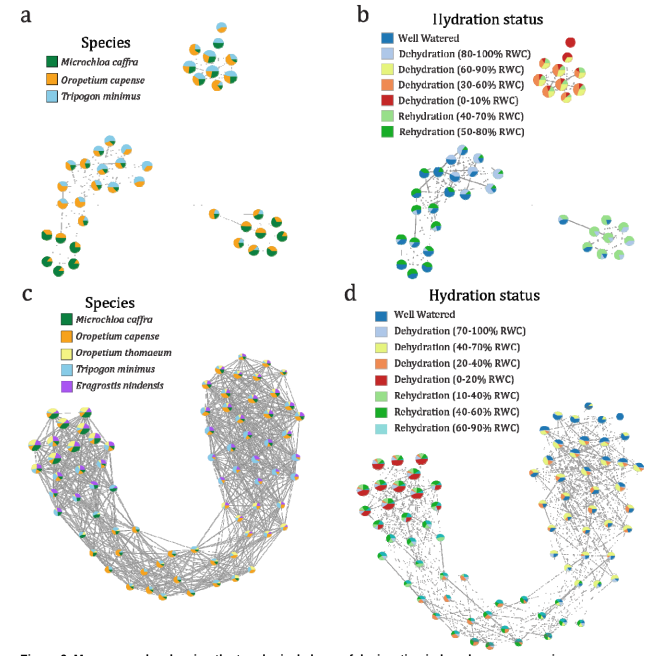


Figure 6. Mapper graphs showing the topological shape of desiccation induced gene expression across
species. Nodes within the graph represent clusters of RNAseq samples that are akin to one another, with the node
color indicating the identity of the samples contained within. Edges, or the connections between nodes, delineate
shared samples across intersecting clusters. Mapper graphs for the three species comparisons are shown in (a) and
(b), and Mapper graphs for the five species comparisons are shown in (c) and (d). Nodes within the graph are colored

by species (a and c) or hydration status (b and d).

Assembly stats	O. capense	T. minimus	M. caffra	
Ploidy	diploid	diploid	hexaploid	
Total assembly size (Mb)	237	223	968	
Number of contigs	14	57	118	
Contig N50	27,924,228	19,548,099	16,141,787	
Contig L50	4	5	22	
Number of genes	28,826	26,527	85,24	
Complete BUSCO (%)	97.1	95.3	96.2	
LTR elements (% of genome):	27.6	22.4	27.0	
Ty1/Copia (%)	4.0	5.2	5.	
Gypsy/DIRS1 (%)	21.4	13.2	15.	
DNA transposons (%)	12.2	15.9	27.0	
Total repeats (%)	41.7	39.4	56.	

# 852 Table 1. Assembly stats of the three resurrection grasses

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