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1	Drought frequency predicts life history strategies in <i>Heliophila</i>
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Abstract

Explaining variation in life history strategies is a long-standing goal of evolutionary biology. 12 For plants, annual and perennial life histories are thought to reflect adaptation to 13 environments that differ in the frequency of stress events such as drought. Here we test this 14 hypothesis in *Heliophila* (Brassicaceae), a diverse genus of flowering plants native to Africa, 15 by integrating 34 years of satellite-based drought measurements with 2192 herbaria 16 occurrence records. Consistent with predictions from classic life history theory, we find that 17 perennial *Heliophila* species occur in environments where droughts are significantly less 18 frequent compared to annuals. These associations are predictive while controlling for 19 phylogeny, lending support to the hypothesis that drought related natural selection has 20 influenced the distributions of these strategies. Additionally, the collection dates of annual 21 and perennial species indicate that annuals escape drought prone seasons during the seed 22 phase of their life cycle. Together, these findings provide empirical support for classic 23 hypotheses about the drivers of life history strategy in plants - that perennials out compete 24 annuals in environments with less frequent drought and that annuals are adapted to 25 environments with more frequent drought by escaping drought prone seasons as seeds. 26

Keywords: drought adaptation, life history evolution, remote sensing, phylogeography,
 herbaria records

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Drought frequency predicts life history strategies in Heliophila

Introduction

Understanding the causes and consequences of life history variation is a longstanding 31 goal of ecology and evolutionary biology (Cole, 1954). In plants, life histories are especially 32 diverse, with herbaceous species that complete their life cycle in a number of weeks to trees 33 that live for thousands of years (Brown, 1996). Along this continuum in angiosperms an 34 important division exists, distinguishing annuals which complete their seed to seed life cycle 35 within a single calendar year from perennials which can persist over multiple years. Annual 36 plants flower once, set seed, senesce, and then die, spending at least some portion of the year 37 as a seed, where they are relatively protected from environmental stress. In contrast, 38 perennial plants can continue vegetative growth after reproduction and must survive 39 conditions experienced during all seasons. These represent fundamentally different life 40 history strategies, but the ecological factors that explain their evolution and distributions 41 remain empirically uresolved (Friedman & Rubin, 2015). 42

Classical theory predicts shorter life spans in environments where adult mortality is 43 high (Charnov & Schaffer, 1973; Stearns, 1992; Franco & Silvertown, 1996). In plants, this 44 has been extended to the hypothesis that annuality is adaptive when it allows plants to 45 escape drought (Schaffer & Gadgil, 1975). Lack of water is perhaps the greatest threat to 46 survival during vegetative or reproductive growth and annuals can remain dormant (and 47 protected as a seed) during drought. Thus, environments with greater seasonal drought 48 frequency may select for annual life histories that complete reproduction prior to drought 49 prone seasons. Conversely, environments with less frequent drought may select for perennial 50 species, which benefit from multiple bouts of reproduction and competitive advantage by 51 preventing recruitment of annual species (Corbin & D'Antonio, 2004). These predictions 52 have been supported by the observation of annuals in arid environments in Oryza perennis 53 (Morishima et al., 1984) and Oenothera (Evans et al., 2005). Additionally, annual and 54

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⁵⁵ perennial species of *Nemesia* were qualitatively associated with winter rather and summer
⁵⁶ rainfall environments respectively (Datson *et al.*, 2008) and annual species of *Scorzoneroides*⁵⁷ were associated with environments classified as unpredictable (Cruz-Mazo *et al.*, 2009).
⁵⁸ However, whether the history frequency of drought events indeed predicts the distributions
⁵⁹ annual or perennial life history strategies has yet to be tested.

Here we combine a long-term global dataset of satellite detected drought events with 60 metadata from natural history collections to test these classic hypotheses within the African 61 endemic mustard genus, *Heliophila L.* (Brassicaceae). If annuality is an adaptive strategy 62 allowing plants to escape drought prone seasons, then drought frequency should predict the 63 distribution of life history strategies across landscapes, and annual species should be more 64 commonly associated with drought prone regions than perennial species. Furthermore, if 65 annual species have adapted to escape drought prone seasons, observations of growing annual 66 species (i.e. occurring in forms other than seed) should be rare during drought prone seasons. 67 Phylogenetic relatedness can influence tests of associations between species' traits and their 68 environments (Felsenstein, 1985; Barrett et al., 1996), and therefore we assessed the 69 relationship between life history distribution and drought frequency in a phylogenetic 70 context. 71

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Materials and Methods

73 Data

Availability. All analyses were performed using R. All data and the source code to
produce this manuscript are available at https://github.com/greymonroe/heliophila.
Software used is listed in the supplement.

Satellite-detected drought data. Remotely sensed data is a powerful tool for
characterizing seasonal patterns in drought because it is less limited in spatial and temporal

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⁷⁹ scope and resolution than weather stations or field observations (AghaKouchak *et al.*, 2015).

⁸⁰ To quantify the frequency of drought during different seasons across landscapes, we used the

⁸¹ remotely sensed Vegetative Health Index (VHI), which measures landscape scale reductions

⁸² in plant cover and temperature conditions characteristic of drought (Kogan, 2001).

⁸³ Generated from data collected by NOAA AVHRR satellites since 1981, the VHI combines

⁸⁴ Normalized Difference Vegetation Index (NDVI) derived measures of vegetative stress

⁸⁵ (Vegetative Condition Index - VCI) with temperature stress indicated by anomalies in

thermal spectra (Temperature Condition Index - TCI). The VHI of year y during week w of

 $_{87}$ [1,52] at pixel *i* is derived from the following equations, where *n* is the number of years

88 observed.

$$VCI_{y,w,i} = 100 \frac{NDVI_{y,w,i} - NDVI_{min}}{NDVI_{max} - NDVI_{min}}$$

$$TCI_{y,w,i} = 100 \frac{T_{y,w,i} - T_{min}}{T_{max} - T_{min}}$$

$$VHI_{y,w,i} = 0.5(VCI_{y,w,i}) + 0.5(TCI_{y,w,i})$$

⁸⁹ where $NDVI_{min} = min(NDVI_{1981,w,i}...NDVI_{1981+n,w,i})$ and

⁹⁰
$$NDVI_{max} = max(NDVI_{1981,w,i}...NDVI_{1981+n,w,i})$$
 and $T_{min} = min(T_{1981,w,i}...T_{1981+n,w,i})$
⁹¹ and $T_{max} = max(T_{1981,w,i}...T_{1981+n,w,i})$

Thus, VHI measurements are standardized according to conditions historically observed at each locations. These measurements have been validated and generally used for evaluating drought risk and predicting crop yields in agriculture (e.g., Rojas *et al.*, 2011; Kogan *et al.*, 2016). But they also present a new tool to study seasonal patterns in the frequency of drought across environments and to test hypotheses about the effect of drought on ecological and evolutionary processes (Kerr & Ostrovsky, 2003). As such, the VHI has

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⁹⁸ been applied recently to study drought related ecology of natural species and proven useful ⁹⁹ for predicting intraspecific variation in drought tolerance traits and genes (Mojica *et al.*, ¹⁰⁰ 2016; Dittberner *et al.*, 2018; Monroe *et al.*, 2018b). Here, we accessed VHI data at $16km^2$ ¹⁰¹ resolution from 1981 to 2015

(https://www.star.nesdis.noaa.gov/smcd/emb/vci/VH/vh_ftp.php) to characterize the
seasonal drought frequencies experienced by annual and perennial *Heliophila* species.

Life history data for *Heliophila*. *Heliophila* is a genus of flowering plants 104 endemic to the southern portion of Africa including the Cape Floristic and Succulent Karoo 105 Regions. These are among the most botanically diverse environments on Earth and the 106 Heliophila species occurring there are considered to make up the most diverse genus of the 107 family Brassicaceae (Mummenhoff et al., 2005; Mandáková et al., 2012). This genus includes 108 both perennial and annual species and this change in life history strategy has likely arisen 109 multiple independent times (Appel & Al-Shehbaz, 1997; Mummenhoff et al., 2005). 110 Furthermore, the fine scale climatic heterogeneity of Southern Africa is ideal for studying the 111 distribution of traits in relation to environmental parameters (Sayre *et al.*, 2013). We used 112 life histories reported by Mummenhoff et al. (2005), grouping species with annual or 113 perennial life histories. Perenniality was defined based any form of perennial life history (e.g., 114 herbs, shrubs, mixed, etc). Because the nature of species reported with mixed traits were 115 unknown (i.e. plasticity vs. genetic variation), we classified these species here as perennial 116 since they can maintain vegetative growth after reproduction at least to some capacity. 117

Heliophila occurrence records. Botanists have collected and maintained over 350 million botanical specimens worldwide over the past 300 years (Thiers, 2016). Herbarium specimens and their associated metadata have been used since the 1960s to study species' geographical distributions (reviewed by Willis *et al.* (2017) and Lang *et al.* (2018)). And as they become digitized (Soltis, 2017), these collections have been used to study relationships between trait distributions, geography, and climate (Davis *et al.*, 2015; Stropp *et al.*, 2016;

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Wolf et al., 2016; Václavı'k et al., 2017). To characterize the distributions of annual and
perennial *Heliophila* species, all records for the genus *Heliophila* were downloaded from the
Global Biodiversity Information Facility (gbif.org) on July 21, 2018 (GBIF, 2018).

Sequence data for phylogeny. An alignment of ITS I and II sequences for
 Heliophila species was obtained from the authors of Mandáková *et al.* (2012). Individual ITS
 I and II sequences for *Aethionema grandiflorum*, *Alliaria petiolata*, *Cardamine matthioli*,
 Chamira circaeoides, and *Rorippa amphibia* were downloaded from Genbank.

131 Analyses

Drought frequency calculations. To characterize drought regimens across the 132 distributions of annual and perennial species of *Heliophila*, we calculated drought during 133 different seasons at the location of observations for *Heliophila* records using the VHI. 134 Specifically, we created global maps of the frequencies of observing drought conditions 135 (VHI<40, NOAA) during the winter (quarter surrounding winter solstice), spring (quarter 136 surrounding spring equinox), summer (quarter surrounding summer solstice) and fall (quarter 137 surrounding fall equinox) from 1981 to 2015. From these maps, the drought frequency during 138 the winter, spring, summer, and fall were extracted for the locations of all GBIF records. 139

Filtering of occurrence records. To avoid instances with spurious location data,
we filtered raw GBIF by restricting our analyses to include only:

- records for species with reported life history
- records with geospatial data
- records without known geospatial coordinate issues (i.e., coordinates reported are those
 of herbarium)
- records from collection sites classified as land pixels in the VHI dataset
- records from Africa (to exclude locations of cultivation)

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• records without duplicates (i.e., identical species, location, collection date)

Phylogeny construction. Out group (*Aethionema grandiflorum*, *Alliaria petiolata*, 149 Cardamine matthioli, Chamira circaeoides, and Rorippa amphibia) and ingroup Heliophila 150 ITS I and II sequences were assigned using MAFFT (Katoh et al., 2002) with strategy 151 G-INS-I, offset value 0.1, and all other options set as default. The $GTR + \Gamma$ model of 152 nucleotide substitution was determined to best fit the data based on AIC using iModelTest2 153 (Guindon & Gascuel, 2003; Darriba et al., 2012). A maximum clade credibility tree with 154 branch lengths as relative time was estimated by summarizing data from six runs of 155 100.000.000 generations of Bayesian Markov chain Monte Carlo conducted in BEAST 2 156 (Bouckaert *et al.*, 2014). Model selection and phylogenetic analyses were conducted through 157 the CIPRES Science Gateway (Miller et al., 2010). 158

Comparison of drought frequency between annual and perennial species. 159 To evaluate the hypothesis that annual and perennial life history strategies reflect 160 adaptations to alternative drought regimes, we tested the corresponding prediction that the 161 observed distributions of annual and perennial *Heliophila* species would be significantly 162 associated with historic drought frequency. First, we compared the frequency of drought 163 during the winter, spring, summer, and fall between total occurrence records of annual and 164 perennial species by t-tests. To account for variation in the number of occurrence records per 165 species, we next calculated the mean drought frequency during the winter, spring, summer 166 and fall for each species. Because shared evolutionary history of closely related species can 167 lead to spurious associations between traits and environments (Felsenstein, 1985), we tested 168 for a relationship between life history strategy and drought frequency while controlling for 169 phylogeny using phylogenetic logistic regression (Ives & Garland, 2010). 170

Collection dates. To test the hypothesis that annual species have adapted to
 escape drought prone seasons as seeds, collection dates for herbarium specimens were
 compared between annual and perennial species. Comparisons of distributions were made by

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¹⁷⁴ Two-sample Kolmogorov-Smirnov test and Barlett variance test.

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Results

Out of 8670 *Heliophila* GBIF records, 6634 were for species with reported life history (Mummenhoff *et al.*, 2005), 2856 had geospatial data, 2833 did not have geospatial issues, 2684 were located on pixels classified as land having drought measurements, 2543 were located in Africa, 2192 were not duplicated. Thus, after all filtering steps, 2192 records for 42 species (Figure 1, Table S1) passed for further analyses. The number of samples varied between species, with a mean of 52.19 samples per species. *H. rigidiuscula* had the most records, 201, and *H. cornellsbergia* the fewest, 2 (Table S1).

There were clear visual differences between the distributions of the 960 annual and the 1232 perennial *Heliophila* observation records (see Figure S1 for maps of individual species). While annual species were generally found in the western regions of South Africa and Namibia, primarily in the Cape Floristic Region and Succulent Karoo (Figure 2a), the occurrence of perennials extended to the east coast of South Africa (Figure 2b).

The frequency of drought varied considerably across the ranges of *Heliophila* species 188 (Figure 2c-f). This heterogeneity is expected, given that this is one of the most climatically 189 diverse regions of the Earth (Savre et al., 2013). It is worth noting the east to west cline in 190 drought frequency observed during the summer, which distinguishes the high drought 191 frequency of the Kalahari Sands and Namid Desert phytogeographic regions from the low 192 drought frequency of the Drakensberg Mountains and Coastal Zambesian phytogeographic 193 regions. In the Cape phytogeographic region there was finer scale heterogeneity in drought 194 frequency during the summer. 195

Theory predicts that annuality should be adaptive in places where stresses such as drought are more common. Conversely, perenniality should be adaptive in places where such

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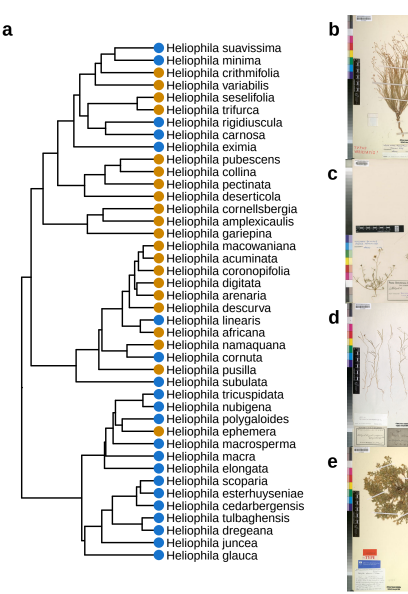


Figure 1. Species and examples of herbaria specimens of *Heliophila* (a) Phylogeny and life history strategies of species studied. Orange circles at branch tips mark annual species and blue circles mark perennial species. Example herbaria specimens accessed via GBIF of (a) *H. minima*, (b) *H. deserticola*, (c) *H. coronopifolia* and (d) *H. ephemera*. Images (a,c,d) courtesy of The Bavarian Natural History Collections (CC BY-SA 4.0) and (b) The London Natural History Museum (CC BY 4.0). Links to images are found in the supplement.

¹⁹⁸ stresses are less frequent. We found that the frequency of drought was significantly higher at ¹⁹⁹ the locations of occurrence records for annual species. When comparing across all occurrence

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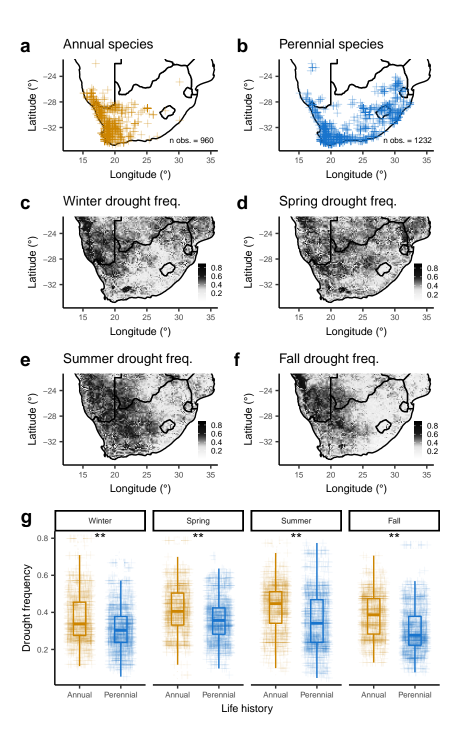


Figure 2. Locations of occurrence records of (a) annual and (b) perennial *Heliophila*. Drought frequency during the (c) winter, (d) spring, (e) summer and (f) fall measured using the VHI. (g) Drought frequencies during each season at the observation locations of annual and perennial *Heliophila* (t tests, $^{**} = p < 0.01$).

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records (all records rather than species means, Figure 2g), the frequency of drought was 200 significantly higher at the location of annuals during the winter (t = 10.65, p = 0.00), spring 201 (t = 10.73, p = 0.00), summer (t = 12.67, p = 0.00), and fall (t = 15.26, p = 0.00). Because 202 a comparison across all occurrence records does not account for variation in the number of 203 records per species (Table S1) or species relatedness (Figure 1a), we also tested whether 204 mean drought frequency values of each species were significantly different between annuals 205 and perennials using phylogenetic logistic regression. We found that the mean drought 206 frequencies were significantly higher ($\alpha = 0.05$) in annual species during the spring, summer, 207 and fall (Table 1, Figure 3a). These findings indicate that common acestry alone does not 208 explain differences the drought frequencies experienced between the environments of annual 209 and perennial *Heliophila*. 210

The preceding results indicate that annual species are found in environments where 211 droughts are significantly more frequent, especially in the summer and fall. Classic life 212 history theory hypothesizes that annuality reflects adaptation to such environments because 213 it allows species to escape stressful conditions. If this is the case, we would expect that 214 annuals spend the drought prone seasons of summer and fall as seeds. To test this 215 hypothesis, we compared the dates of occurrence records between annual and perennial 216 Heliophila species. The distributions reveal a considerable difference in the timing of 217 observation of these two life histories. In comparison to perennials, which appear to be 218 collected throughout the year, annuals are almost exclusively observed during the winter and 219 spring (Figure 3b). The differences between the distribution of collection dates were 220 significant by all tests (ks.test D = 0.25, p = 0; bartlett.test K2 = 503.18, p = 0.00) This is 221 consistent with a model of life history in which annual species flower in the spring, set seed, 222 senesce, and die before the summer. Thus, these annual species are likely to remain dormant 223 during the summer and fall, when drought is the strongest predictor of the distributions of 224 annual and perennial life histories (Figure 3a). 225

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Table 1

Phylogenetic logistic regressions between life history, and the mean drought frequency observed at specimen sites of Heliophila species the winter, spring, summer, and fall.

Predictor	Estimate	Р
Intercept	0.7231	0.6636
Winter drought freq.	-1.5452	0.7274
Intercept	5.0107	0.0534
Spring drought freq.	-12.9014	0.0464
Intercept	7.7093	0.0054
Summer drought freq.	-19.9056	0.0042
Intercept	7.0162	0.0082
Fall drought freq.	-20.8174	0.0067

Note. Annual species were scored as 0 and perennial species as 1.

Discussion

To test the hypothesis that annual and perennial plants reflect adaptation to alternative drought environments we examined the landscape distribution of life history strategies in the large and diverse mustard genus, *Heliophila*. Using metadata of 2192 occurrence records and a 34 year dataset of satellite-detected droughts, we tested the prediction that annual species are more often observed in drought-prone locations than perennial species, when controlling for phylogenetic relatedness. We found that drought frequency is significantly different between the distributions of annual and perennial species,

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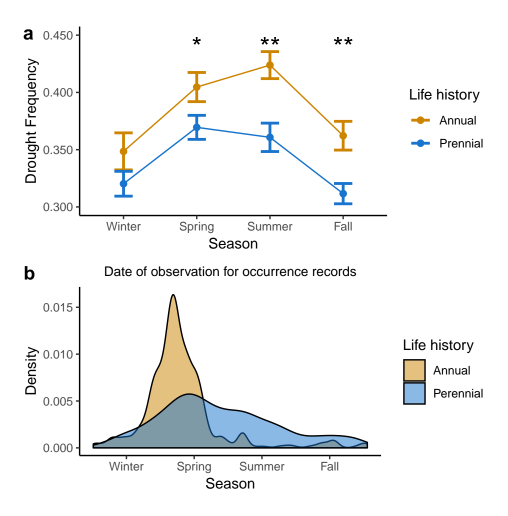


Figure 3. (a) Comparison (mean +- SE) of drought frequency across seasons measured at the GBIF records of annual and perennial species of *Heliophila*. (phylogenetic logistic regression, * = p < 0.05, ** = p < 0.01) (b) Collection dates of GBIF records of annual and perennial species of *Heliophila*.

with annuals being found in environments with more frequent drought, and that this signal is strongest during the seasons when annuals are likely escaping via seed dormancy. These results remain significant while controlling for the phylogenetic relationships of *Heliophila* species, yielding support for the role that natural selection has played in driving contemporary distributions of these alternatives strategies in relation to drought regimens.

We cannot eliminate the possibility that confounding traits or environmental variables

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are the causative factors explaining variation in the distributions of annual and perennial 240 species. Nevertheless, these results provide quantitative support for the classic prediction 241 that annual species are found in environments that experience more frequent drought than 242 perennial species. These findings complement previous reports of qualitative associations 243 between annuality with environments characterized as having increased aridity (Evans *et al.*, 244 2005), alternative precipitation defined habitats (Morishima et al., 1984; Datson et al., 2008), 245 or greater unpredictability (Cruz-Mazo et al., 2009). However, to our knowledge this is the 246 first study to demonstrate a significant association between life history and drought in a 247 phylogenetic context informed by large scale species distribution data and long term drought 248 measures. 240

Unfortunately, herbarium collections and their associated data do not represent 250 systematic or random sampling of a species distribution. Significant biases in collecting exist, 251 which we have not necessarily controlled for here, and may have some effect on our findings, 252 such as a bias toward collecting near roads or near the locations of natural history collections 253 (Daru et al., 2018). Future research will benefit from systematic sampling efforts to avoid 254 these noted biases. However, the ecosystems of southern Africa include several biodiversity 255 hotspots and are among the most botanically well sampled regions on Earth (Daru *et al.*, 256 2018), suggesting that this may currently be the optimal region for our analyses of life 257 history distribution. Indeed, we were able to use 2192 occurrence records to study 42 species, 258 which represents a significant advance over relying on personal observations to characterize 259 species distributions. 260

These findings support classical theoretical predictions about the adaptive value of annual and perennial life history strategies. Taken together, they suggest that in *Heliophila*, annual species are adapted to environments with increased summer droughts by avoiding these seasons in a dormant seed phase of their life cycle. They also suggest that perenniality is adaptive in environemnts where droughts are less frequent. While most previous work has

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focused on describing the evolutionary origins of annuality (Barrett et al., 1996; Conti et al., 266 1999; Andreasen & Baldwin, 2001; Verboom et al., 2004; Friedman & Rubin, 2015) there are 267 at least a few other cases where perenniality appears to have arisen from an annual ancestor 268 (Bena et al., 1998; Tank & Olmstead, 2008). And while early theory predicted selection for 269 annuality when adult morality is high (Stearns, 1992), we also find evidence that perenniality 270 could be explained by reduced frequency of drought. The phylogeny reveals several 271 transitions from annual to perennial life history (Figure 1a) and the distributions of 272 perennial *Heliophila* extend into regions where drought frequency is low (Figure 2b, Figure 273 S1). Perennials may be able to out complete annual relatives in environments where the 274 infrequency of drought favors strategies that allow plants to benefit from growth over many 275 seasons. This also suggests that annuals rely on drought as a source of disturbance for 276 seedling recruitment when competing with perennials (Corbin & D'Antonio, 2004). Indeed, 277 no annual species were observed in the low drought regions of eastern South Africa (Figure 2, 278 Figure S1). 279

These findings suggest that species with locally adaptive life history strategies could be threatened by rapidly changing drought regimens (Dai, 2011). This could have impacts on ecosystem functioning and processes such as carbon cycling if life history traits evolve or the composition of annual and perennial species changes in response (Garnier *et al.*, 1997; Roumet *et al.*, 2006; Monroe *et al.*, 2018a). Furthermore, the frequency of drought may be an important factor when considering the use of perennial cropping systems (Parry *et al.*, 2005; Lelièvre & Volaire, 2009).

In conclusion, we find strong support for classic life history theory which predicts that annuality is adaptive in environments where droughts occur more frequently. Additionally, we report evidence consistent with a life history model in annuals in which they escape drought prone seasons during the seed phase of their life cycle. Finally, we find evidence that the distributions of perennial lineages may indicate a competitive advantage in areas where

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droughts are infrequent. More broadly, this work highlights the irreplaceable value of natural history collections and demonstrates the power of combining such information with large scale remote sensing data to address outstanding classic hypotheses in ecology and evolution.

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Author contributions

JGM, BG, KGT and JKM contributed to the design of the research, interpretation, and writing the manuscript. JGM, BG, and KGT contributed to the performance of the research and data analysis.

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References

DROUGHT AND LIFE HISTORY

304	AghaKouchak A, Farahmand A, Melton F, Teixeira J, Anderson M, Wardlow BD, Hain C.
305	2015. Remote sensing of drought: Progress, challenges and opportunities. Reviews of
306	<i>Geophysics</i> 53 : 452–480.

³⁰⁷ Alfaro M, Santini F, Brock C, Alamillo H, Dornburg A, Rabosky D, Carnevale G, Harmon

- L. 2009. Nine exceptional radiations plus high turnover explain species diversity in
- jawed vertebrates. Proceedings of the National Academy of Sciences of the United

310 States of America **106**: 13410–13414.

- Andreasen K, Baldwin BG. 2001. Unequal evolutionary rates between annual and perennial
 lineages of checker mallows (sidalcea, malvaceae): Evidence from 18S–26S rDNA
 internal and external transcribed spacers. *Molecular Biology and Evolution* 18:
 936–944.
- Appel O, Al-Shehbaz IA. 1997. Generic limits and taxonomy of hornungia, pritzelago, and
 hymenolobus (brassicaceae). Novon: 338–340.
- Aust F, Barth M. 2018. papaja: Create APA manuscripts with R Markdown.
- Barrett SCH, Harder LD, Worley AC. 1996. The comparative biology of pollination and
 mating in flowering plants. *Phil. Trans. R. Soc. Lond. B* 351: 1271–1280.

³²⁰ Bates D, Maechler M. 2018. Matrix: Sparse and dense matrix classes and methods.

Bena G, Lejeune B, Prosperi J-M, Olivieri I. 1998. Molecular phylogenetic approach for
 studying life-history evolution: The ambiguous example of the genus medicago l.
 Proceedings of the Royal Society of London B: Biological Sciences 265: 1141–1151.

Bouckaert R, Heled J, Kühnert D, Vaughan T, Wu C-H, Xie D, Suchard M, Rambaut A,
 Drummond A. 2014. BEAST 2: A software platform for bayesian evolutionary
 analysis. *PLoS Computational Biology* 10: doi:10.1371/journal.pcbi.1003537.

DROUGHT AND LIFE HISTORY

- Brown PM. 1996. OLDLIST: A database of maximum tree ages. Tree rings, environment,
 and humanity. Radiocarbon 1996: 727–731.
- ³²⁹ Charnov EL, Schaffer WM. 1973. Life-history consequences of natural selection: Cole's
 result revisited. *The American Naturalist* 107: 791–793.
- Cole LC. **1954**. The population consequences of life history phenomena. *The Quarterly Review of Biology* **29**: 103–137.

Conti E, Soltis DE, Hardig TM, Schneider J. 1999. Phylogenetic relationships of the silver
 saxifrages (saxifraga, sect. Ligulatae haworth): Implications for the evolution of
 substrate specificity, life histories, and biogeography. *Molecular Phylogenetics and Evolution* 13: 536–555.

- ³³⁷ Corbin JD, D'Antonio CM. 2004. Competition between native perennial and exotic annual
 ³³⁸ grasses: Implications for an historical invasion. *Ecology* 85: 1273–1283.
- ³³⁹ Cruz-Mazo G, Buide M, Samuel R, Narbona E. 2009. Molecular phylogeny of
- scorzoneroides (asteraceae): Evolution of heterocarpy and annual habit in
- ³⁴¹ unpredictable environments. *Molecular phylogenetics and evolution* **53**: 835–847.
- ³⁴² Dai A. 2011. Drought under global warming: A review. Wiley Interdisciplinary Reviews:
 ³⁴³ Climate Change 2: 45–65.
- Darriba D, Taboada G, Doallo R, Posada D. 2012. JModelTest 2: More models, new
 heuristics and parallel computing. *Nature Methods* 9: 772.
- Daru BH, Park DS, Primack RB, Willis CG, Barrington DS, Whitfeld TJ, Seidler TG,
 Sweeney PW, Foster DR, Ellison AM *et al.* 2018. Widespread sampling biases in
 herbaria revealed from large-scale digitization. *New Phytologist* 217: 939–955.

DROUGHT AND LIFE HISTORY

- Datson P, Murray B, Steiner K. 2008. Climate and the evolution of annual/perennial
 life-histories in nemesia (scrophulariaceae). *Plant Systematics and Evolution* 270:
 39–57.
- ³⁵² Davis CC, Willis CG, Connolly B, Kelly C, Ellison AM. 2015. Herbarium records are
 ³⁵³ reliable sources of phenological change driven by climate and provide novel insights
 ³⁵⁴ into species' phenological cueing mechanisms. *American Journal of Botany* 102:
 ³⁵⁵ 1599–1609.
- Dittberner H, Korte A, Mettler-Altmann T, Weber A, Monroe G, Meaux J de. 2018.
 Natural variation in stomata size contributes to the local adaptation of water-use
 efficiency in arabidopsis thaliana. *bioRxiv*: 253021.
- Eastman J, Alfaro M, Joyce P, Hipp A, Harmon L. 2011. A novel comparative method for
 identifying shifts in the rate of character evolution on trees. *Evolution* 65: 3578–3589.
- Evans ME, Hearn DJ, Hahn WJ, Spangle JM, Venable DL. 2005. Climate and life-history
 evolution in evening primroses (oenothera, onagraceae): A phylogenetic comparative
 analysis. Evolution 59: 1914–1927.
- Felsenstein J. 1985. Phylogenies and the comparative method. American Naturalist 125:
 1-15.
- Franco M, Silvertown J. **1996**. Life history variation in plants: An exploration of the fast-slow continuum hypothesis. *Phil. Trans. R. Soc. Lond. B* **351**: 1341–1348.
- Friedman J, Rubin MJ. 2015. All in good time: Understanding annual and perennial
 strategies in plants. American journal of botany 102: 497–499.
- Garnier E, Cordonnier P, Guillerm J-L, Sonié L. 1997. Specific leaf area and leaf nitrogen
 concentration in annual and perennial grass species growing in mediterranean

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- old-fields. *Oecologia* **111**: 490–498.
- ³⁷³ GBIF. **2018**. GBIF occurrence download.
- Genz A, Bretz F. **2009**. Computation of multivariate normal and t probabilities. Heidelberg: Springer-Verlag.
- ³⁷⁶ Guindon S, Gascuel O. **2003**. A simple, fast and accurate method to estimate large ³⁷⁷ phylogenies by maximum-likelihood. *Systematic Biology* **52**: 696–704.
- ³⁷⁸ Hadfield JD. **2010**. MCMC methods for multi-response generalized linear mixed models:

The MCMCglmm R package. Journal of Statistical Software 33: 1–22.

- Harmon L, Weir J, Brock C, Glor R, Challenger W. 2008. GEIGER: Investigating
 evolutionary radiations. *Bioinformatics* 24: 129–131.
- Heinze G, Ploner M. 2018. Logistf: Firth's bias-reduced logistic regression.
- Henry L, Wickham H. 2018. Purr: Functional programming tools.
- ³⁸⁴ Hijmans RJ. 2018. Raster: Geographic data analysis and modeling.
- Ho LST, Ane C. 2014. A linear-time algorithm for gaussian and non-gaussian trait
 evolution models. Systematic Biology 63: 397–408.
- Ives A, Garland T. 2010. CPhylogenetic logistic regression for binary dependent variables.
 Systematic Biology 59: 9–26.
- Katoh, Misawa, Kuma, Miyata. 2002. MAFFT: A novel method for rapid multiple
 sequence alignment based on fast fourier transform. Nucleic Acids Research 30:
 3059–3066.

DROUGHT AND LIFE HISTORY

- Kerr JT, Ostrovsky M. 2003. From space to species: Ecological applications for remote
 sensing. Trends in ecology & evolution 18: 299–305.
- Kogan FN. 2001. Operational space technology for global vegetation assessment. Bulletin
 of the American Meteorological Society 82: 1949–1964.
- Kogan F, Guo W, Strashnaia A, Kleshenko A, Chub O, Virchenko O. 2016. Modelling and
 prediction of crop losses from noaa polar-orbiting operational satellites. *Geomatics*,
 Natural Hazards and Risk 7: 886–900.
- Lang PL, Willems FM, Scheepens J, Burbano HA, Bossdorf O. 2018. Using herbaria to
 study global environmental change. PeerJ Preprints.
- Lelièvre F, Volaire F. 2009. Current and potential development of perennial grasses in
 rainfed mediterranean farming systems. Crop Science 49: 2371–2378.
- Mandáková T, Mummenhoff K, Al-Shehbaz IA, Mucina L, Mühlhausen A, Lysak MA.
 2012. Whole-genome triplication and species radiation in the southern african tribe
 heliophileae (brassicaceae). Taxon 61: 989–1000.
- Miller M, Pfeiffer W, Schwartz T and. 2010. Creating the cipres science gateway for
 inference of large phylogenetic trees. *Proceedings of the Gateway Computing Environments Workshop*: 1–8.

Mojica JP, Mullen J, Lovell JT, Monroe JG, Paul JR, Oakley CG, McKay JK. 2016.
Genetics of water use physiology in locally adapted Arabidopsis thaliana. *Plant Science*.

Monroe J, Markman D, Beck W, Felton A, Vahsen M, Pressler Y. 2018a. Ecoevolutionary
dynamics of carbon cycling in the anthropocene. *Trends in ecology & evolution* 33:
213–225.

DROUGHT AND LIFE HISTORY

415	Monroe J, Powell T, Price N, Mullen J, Howard A, Evans K, Lovell J, McKay J. 2018b.
416	Drought adaptation in nature by extensive genetic loss-of-function. $eLife:$ DOI:
417	10.7554/eLife.41038.

Morishima H, Sano Y, Oka H. 1984. Differentiation of perennial and annual types due to
habitat conditions in the wild rice oryza perennis. *Plant Systematics and Evolution*144: 119–135.

- Mummenhoff K, Al-Shehbaz IA, Bakker FT, Linder HP, Mühlhausen A. 2005. Phylogeny,
 morphological evolution, and speciation of endemic brassicaceae genera in the cape
 flora of southern africa. Annals of the Missouri Botanical Garden: 400–424.
- ⁴²⁴ Müller K. 2018. Bindrcpp: An 'rcpp' interface to active bindings.
- ⁴²⁵ Müller K, Wickham H. 2018. *Tibble: Simple data frames.*
- ⁴²⁶ Orme D, Freckleton R, Thomas G, Petzoldt T, Fritz S, Isaac N, Pearse W. 2018. Caper:
 ⁴²⁷ Comparative analyses of phylogenetics and evolution in r.
- Paradis E, Schliep K. 2018. Ape 5.0: An environment for modern phylogenetics and
 evolutionary analyses in R. *Bioinformatics* xx: xxx-xxx.
- Parry M, Flexas J, Medrano H. 2005. Prospects for crop production under drought:
 Research priorities and future directions. Annals of Applied Biology 147: 211–226.
- ⁴³² Pebesma EJ, Bivand RS. 2005. Classes and methods for spatial data in R. R News 5: 9–13.
- Plummer M, Best N, Cowles K, Vines K. 2006. CODA: Convergence diagnosis and output
 analysis for mcmc. *R News* 6: 7–11.
- R by Thomas Lumley VJCP to, author. BRN that maintainers are not available to give
 advice on using a package they did not. 2015. Gee: Generalized estimation equation

DROUGHT AND LIFE HISTORY

solver.

438	R Core Team. 2018. R: A language and environment for statistical computing. Vienna,
439	Austria: R Foundation for Statistical Computing.
440	Revell LJ. 2012. Phytools: An r package for phylogenetic comparative biology (and other
441	things). Methods in Ecology and Evolution 3 : 217–223.
442	Richard A. Becker OS code by, Ray Brownrigg. Enhancements by Thomas P Minka ARWR
443	version by, Deckmyn. A. 2018. Maps: Draw geographical maps.
	Rojas O, Vrieling A, Rembold F. 2011 . Assessing drought probability for agricultural areas
444	
445	in africa with coarse resolution remote sensing imagery. <i>Remote sensing of</i>
446	<i>Environment</i> 115 : 343–352.
447	Roumet C, Urcelay C, Dı'az S. 2006. Suites of root traits differ between annual and
448	perennial species growing in the field. New phytologist 170: 357–368.
449	Sayre RG, Comer P, Hak J, Josse C, Bow J, Warner H, Larwanou M, Kelbessa E, Bekele
450	T, Kehl H et al. 2013. A new map of standardized terrestrial ecosystems of africa.
451	African Geographical Review.
452	Schaffer W, Gadgil M. 1975. Selection for optimal life histories in plants. Ecology and
453	evolution of communities.: 142–157.
454	Slater G, Harmon L, Wegmann D, Joyce P, Revell L, Alfaro M. 2012. Fitting models of
455	continuous trait evolution to incompletely sampled comparative data using
456	approximate bayesian computation. Evolution 66: 752–762.
	· · ·
457	Soetaert K. 2017. Diagram: Functions for visualising simple graphs (networks), plotting
458	flow diagrams.

DROUGHT AND LIFE HISTORY

- ⁴⁵⁹ Soetaert K. **2018**. *Shape: Functions for plotting graphical shapes, colors.*
- Soltis PS. 2017. Digitization of herbaria enables novel research. American journal of botany
 104: 1281–1284.
- ⁴⁶² Stearns SC. **1992**. The evolution of life histories.
- 463 Stropp J, Ladle RJ, M. Malhado AC, Hortal J, Gaffuri J, H. Temperley W, Olav Skøien J,
 464 Mayaux P. 2016. Mapping ignorance: 300 years of collecting flowering plants in
 465 africa. Global Ecology and Biogeography 25: 1085–1096.
- Tank DC, Olmstead RG. 2008. From annuals to perennials: Phylogeny of subtribe
 castillejinae (orobanchaceae). American Journal of Botany 95: 608–625.
- Thiers B. 2016. Index herbariorum: A global directory of public herbaria and associated
 staff. New york botanical garden's virtual herbarium. http://sweetgum. nybg. org/ih.
- ⁴⁷⁰ Václavı'k T, Beckmann M, Cord AF, Bindewald AM. 2017. Effects of uv-b radiation on
 ⁴⁷¹ leaf hair traits of invasive plants—combining historical herbarium records with novel
 ⁴⁷² remote sensing data. *PloS one* 12: e0175671.
- ⁴⁷³ Venables WN, Ripley BD. 2002. Modern applied statistics with s. New York: Springer.
- ⁴⁷⁴ Verboom GA, Linder HP, Stock WD. 2004. Testing the adaptive nature of radiation:
- Growth form and life history divergence in the african grass genus ehrharta (poaceae:
 Ehrhartoideae). American Journal of Botany 91: 1364–1370.
- 477 Wickham H. 2016. Ggplot2: Elegant graphics for data analysis. Springer-Verlag New York.
- 478 Wickham H. 2017. Tidyverse: Easily install and load the 'tidyverse'.
- 479 Wickham H. 2018a. Forcats: Tools for working with categorical variables (factors).

DROUGHT AND LIFE HISTORY

491

- Wickham H. 2018b. Stringr: Simple, consistent wrappers for common string operations. 480
- Wickham H, François R, Henry L, Müller K. 2018. Dplyr: A grammar of data 481 manipulation. 482
- Wickham H, Henry L. 2018. Tidyr: Easily tidy data with 'spread()' and 'qather()' functions. 483
- Wickham H, Hester J, Francois R. 2017. Readr: Read rectangular text data. 484
- Willis CG, Ellwood ER, Primack RB, Davis CC, Pearson KD, Gallinat AS, Yost JM, 485
- Nelson G, Mazer SJ, Rossington NL et al. 2017. Old plants, new tricks: 486
- Phenological research using herbarium specimens. Trends in ecology & evolution 32: 487 531 - 546.488
- Wolf A, Zimmerman NB, Anderegg WR, Busby PE, Christensen J. 2016. Altitudinal shifts 489 of the native and introduced flora of c alifornia in the context of 20th-century 490 warming. Global ecology and biogeography 25: 418–429.

DROUGHT AND LIFE HISTORY

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Supplement

⁴⁹³ Images used. https://www.gbif.org/occurrence/1099023487

https://www.gbif.org/occurrence/1057389408 https://www.gbif.org/occurrence/1099023562
https://www.gbif.org/occurrence/1099023490

496 Software used. We used R (Version 3.5.1; R Core Team, 2018) and the R-packages

ape (Version 5.2; Paradis & Schliep, 2018; Orme *et al.*, 2018; Soetaert, 2018), *bindrcpp*

⁴⁹⁸ (Version 0.2.2; Müller, 2018), *caper* (Version 1.0.1; Orme *et al.*, 2018), *coda* (Version 0.19.2;

⁴⁹⁹ Plummer *et al.*, 2006), *diagram* (Version 1.6.4; Soetaert, 2017), *dplyr* (Version 0.7.8;

⁵⁰⁰ Wickham et al., 2018), forcats (Version 0.3.0; Wickham, 2018a), gee (Version 4.13.19; R by

⁵⁰¹ Thomas Lumley & author., 2015), geiger (Version 2.0.6; Alfaro et al., 2009; Harmon et al.,

⁵⁰² 2008; Eastman et al., 2011; Slater et al., 2012), ggplot2 (Version 3.1.0; Wickham, 2016),

⁵⁰³ logistf (Version 1.23; Heinze & Ploner, 2018), maps (Version 3.3.0; Richard A. Becker et al.,

⁵⁰⁴ 2018), MASS (Version 7.3.51.1; Venables & Ripley, 2002), Matrix (Version 1.2.15; Bates &

⁵⁰⁵ Maechler, 2018), *MCMCglmm* (Version 2.26; Hadfield, 2010), *mvtnorm* (Version 1.0.8; Genz

⁵⁰⁶ & Bretz, 2009), *papaja* (Version 0.1.0.9842; Aust & Barth, 2018), *phylolm* (Version 2.6; Ho &

⁵⁰⁷ Ane, 2014), *phytools* (Version 0.6.60; Revell, 2012), *purrr* (Version 0.2.5; Henry & Wickham,

⁵⁰⁸ 2018), raster (Version 2.8.4; Hijmans, 2018), readr (Version 1.2.1; Wickham et al., 2017),

⁵⁰⁹ shape (Version 1.4.4; Soetaert, 2018), sp (Version 1.3.1; Pebesma & Bivand, 2005), stringr

⁵¹⁰ (Version 1.3.1; Wickham, 2018b), tibble (Version 1.4.2; Müller & Wickham, 2018), tidyr

⁵¹¹ (Version 0.8.2; Wickham & Henry, 2018), and *tidyverse* (Version 1.2.1; Wickham, 2017) for

⁵¹² all our analyses.

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Supplementary tables and figures.

Table S1

Heliophila species records and the mean drought frequencies during

different seasons at the location of records

Species	LH	n	Winter	Spring	Summer	Fall
Heliophila acuminata	a	28	0.32	0.38	0.41	0.36
Heliophila africana	a	91	0.33	0.35	0.34	0.34
Heliophila amplexicaulis	a	60	0.32	0.36	0.39	0.33
Heliophila arenaria	a	65	0.34	0.37	0.38	0.34
Heliophila carnosa	р	129	0.33	0.37	0.39	0.31
Heliophila cedarbergensis	р	3	0.40	0.43	0.32	0.27
Heliophila collina	a	16	0.35	0.47	0.48	0.45
Heliophila cornellsbergia	a	2	0.33	0.42	0.35	0.21
Heliophila cornuta	р	101	0.35	0.40	0.40	0.34
Heliophila coronopifolia	a	40	0.37	0.42	0.40	0.37
Heliophila crithmifolia	a	97	0.35	0.42	0.45	0.38
Heliophila descurva	a	12	0.36	0.38	0.38	0.29
Heliophila deserticola	a	133	0.48	0.48	0.46	0.45
Heliophila digitata	a	30	0.33	0.38	0.44	0.38
Heliophila dregeana	р	17	0.33	0.37	0.33	0.32
Heliophila elongata	р	82	0.26	0.32	0.30	0.25
Heliophila ephemera	a	3	0.14	0.27	0.31	0.26
Heliophila esterhuyseniae	р	3	0.21	0.30	0.37	0.27
Heliophila eximia	р	12	0.42	0.41	0.32	0.34
Heliophila gariepina	a	12	0.50	0.53	0.48	0.41
Heliophila glauca	р	35	0.29	0.35	0.34	0.33
Heliophila juncea	р	150	0.32	0.37	0.39	0.35
Heliophila linearis	р	94	0.32	0.33	0.28	0.30

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Heliophila macowaniana	a	31	0.33	0.38	0.44	0.39
Heliophila macra	р	22	0.30	0.30	0.32	0.29
Heliophila macrosperma	р	5	0.28	0.36	0.35	0.25
Heliophila minima	р	35	0.36	0.45	0.51	0.39
Heliophila namaquana	a	16	0.39	0.46	0.48	0.39
Heliophila nubigena	р	19	0.31	0.36	0.43	0.38
Heliophila pectinata	a	16	0.27	0.34	0.50	0.34
Heliophila polygaloides	р	12	0.40	0.48	0.42	0.34
Heliophila pubescens	a	9	0.31	0.40	0.48	0.39
Heliophila pusilla	a	45	0.32	0.38	0.38	0.34
Heliophila rigidiuscula	р	201	0.30	0.33	0.28	0.24
Heliophila scoparia	р	106	0.31	0.37	0.36	0.31
Heliophila seselifolia	a	80	0.36	0.42	0.45	0.40
Heliophila suavissima	р	92	0.30	0.39	0.42	0.31
Heliophila subulata	р	103	0.29	0.33	0.31	0.29
Heliophila tricuspidata	р	8	0.28	0.33	0.38	0.30
Heliophila trifurca	a	77	0.45	0.48	0.48	0.43
Heliophila tulbaghensis	р	3	0.36	0.41	0.36	0.35
Heliophila variabilis	a	97	0.35	0.41	0.40	0.37

Note. LH = Life history (a = annual, p = perennial). n=sample size of GBIF records. Seasons are mean drought frequencies observed at locations of records.

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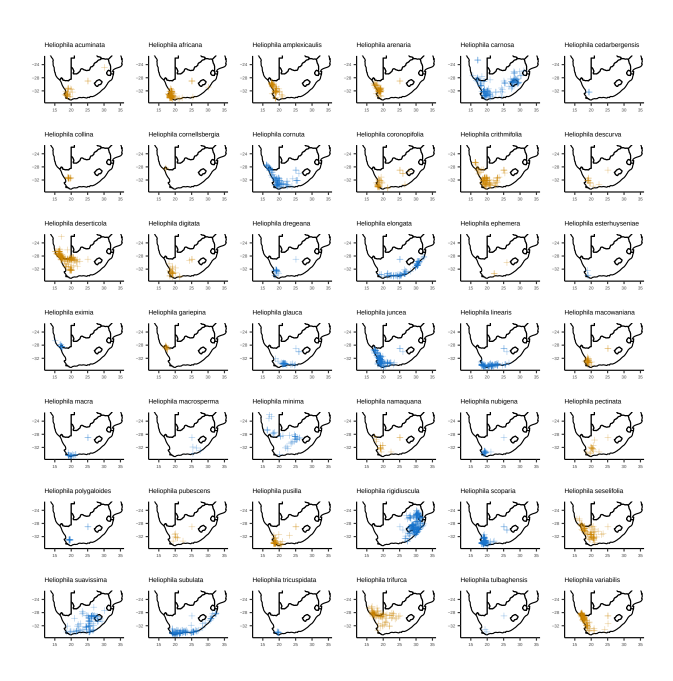


Figure S1. Maps of occurrence records for individual species. Orange points indicate annual species. Blue points indicate perennial species.