# Light and life form interact to shape stomatal ratio among British angiosperms<sup>1</sup>

### Christopher D. Muir<sup>2</sup>

```
<sup>1</sup> Manuscript received _; revision accepted _.
```

Author for correspondence: Christopher D. Muir Tel: +17782284851 Email: chrisdmuir@gmail.com University of British Columbia 6270 University Blvd. Vancouver, BC, Canada V6T 1Z4

Short title: Shedding light on stomatal evolution

Word count:
Summary:
Introduction:
Methods and Results:
Discussion:
# Figures and # Tables, # references

[1] TRUE

 $<sup>^2</sup>$  Biodiversity Research Centre and Botany Department, University of British Columbia, Vancouver, British Columbia V6T 1Z4, Canada

### Acknowledgements

I thank Sally Otto, Matt Pennell, and Rob Salguero-Gómez for feedback on this manuscript.

#### Abstract

- In most plants, stomata are located only on the abaxial leaf surface (hypostomy), but many plants have stomata on both surfaces (amphistomy). Variation in stomatal ratio (the ratio of ab- and adaxial stomatal densities) is probably adaptive, but the ecological conditions that favor amphistomy are not well understood. In particular, high light and herbaceous growth form have been hypothesized to favor amphistomy, but these hypotheses have not been rigourously tested together.
- I leveraged a large dataset including stomatal ratio, Ellenberg light indicator value, Raunkiær lifeform, and phylogenetic relationships for 372 species of

  British angiosperms. I used phylogenetic comparative methods to test how light and/or growth form influence stomatal ratio.
  - (return to this) key results: L-value, growth form, and interaction are important
- I show for the first time that light and growth form interact to shape variation

in stomatal ratio; amphistomy is advantageous in high light, but mostly for herbs. These results improve our understanding of the adaptive significance of stomatal ratio, use stomatal ratio as proxy for paleo vegetation, and as a target for crop improvement.

### 19 Keywords

Adaptation, amphistomy, Ellenberg light indicator value, growth form, phylogenetic

#### 22 INTRODUCTION

comparative methods, stomata, stomatal ratio

Natural selection shapes leaf anatomy in order to optimize its photosynthetic func-

tion in a given environment (Haberlandt, 1914; Givnish, 1987; Smith et al., 1997).

 $_{25}$  By understanding the adaptive significance of leaf anatomical variation we can learn

26 about natural history, find targets for crop improvement, and identify anatomical

27 proxies for paleoclimates preserved in the fossil record [CITE]. The size, density, and

distribution of stomata on a leaf vary widely and impact functions like the maximum

<sup>29</sup> photosynthetic rate, water-use efficiency, photosynthetic nitrogen-use efficiency, and

susceptibility to foliar pathogens that infect through stomata [CITATIONS]. Hence, stomata have been especially useful in understanding plastic and evolutionary response to climate change and domestication (Royer, Ward, Woodward, Beerling, Milla et al...). While the density and size of stomata have been researched extensively [CITA-TIONS, the adaptive significance of stomatal distribution is less well understood. Stomata are most often found only on the lower leaf surface (hypostomy) but occur on both surfaces (amphistomy) in many species (Metcalfe and Chalk, 1950; Parkhurst, 1978; Mott et al., 1984). Theory and experiments demonstrate that amphistomy increases photosynthetic rates under many conditions. By creating a second parallel pathway for CO<sub>2</sub> diffusion within the mesophyll, amphistomy optimally supplies CO<sub>2</sub> (Parkhurst, 1978; Gutschick, 1984; Jones, 1985). Amphistomy is correlated with greater CO<sub>2</sub> diffusion (Beerling and Kelly, 1996) and higher photosynthetic rates (McKown et al., 2014). These observations are corroborated by experiments demonstrating that amphistomy increases maximum photosynthetic rates by up to 20% (Parkhurst and Mott, 1990). On the other hand, amphistomy can increase transpiration (Jones, 1985; Foster and Smith, 1986; Buckley et al., 2015). While transition to amphistomy is thus thought to increase transpiration, empirical studies

suggest greater water-use efficiency in amphistomatous species (Bucher et al., 2017).

Hence, amphistomy appears to benefit a plant's carbon use relative to water loss and should be favored when CO<sub>2</sub> limits photosynthetic rate. The open questions are under what ecological conditions does CO<sub>2</sub> supply most strongly limit photosynthetic rate (Peat and Fitter, 1994) and when is photosynthetic rate most important to fitness?

The leading, nonmutually exclusive hypotheses are that 1) open habitats favour amphistomy because CO<sub>2</sub> diffusion most strongly limits photosynthetic rate under high light and 2) herbaceous growth form favours amphistomy because traits that maximize photosynthetic rate are often under stronger selection in herbs. Salisbury (1927) first noted that amphistomy is most common in herbaceous plants from open habitats (i.e., with high light) of the British flora. These observations have been replicated in other studies (Mott et al., 1984; Peat and Fitter, 1994; Jordan et al., 2014; Muir, 2015) and may support physiological and ecological hypotheses that CO<sub>2</sub> most strongly limits photosynthesis in high light and/or photosynthesis contributes most to fitness in herbaceous plants. Under high light, CO<sub>2</sub> can strongly limit maximum photosynthetic rates, especially in thick leaves (Jones, 1985). Hence, having stomata on both surfaces relieves this limitation by adding a second parallel pathway for CO<sub>2</sub> diffusion. Parkhurst 1978 argued that greater leaf thickness per se selected for amphistomy, but there is little evidence for correlations between leaf thickness

- and stomatal ratio independent of light (Mott et al., 1984; Gibson, 1996; Muir, 2015).
- 69 Amphistomy is correlated with open habitat in warm desert plants of western North
- America (Mott et al., 1984; Gibson, 1996), among the Proteaceae (Jordan et al.,

Stomatal ratio is also associated with growth form. In the British flora, Salisbury

- 2014), and in continental European herbs (Bucher et al., 2017).
- 73 (1927) found that trees and shurbs are nearly always hypostomatous, whereas herbs
  74 from open habitats are amphistomatous. This pattern holds when data are averaged
  75 by family to coarsely control for phylogenetic nonindependence (Peat and Fitter,
  76 1994) or when using alternative classification schemes, such as Raunkiær life form
  77 (Peat and Fitter, 1994). Across plants from 90 familes worldwide, growth form is
  78 the strongest predictor of stomatal ratio when multiple factors are estimated simulta79 neously and controlling for phylogenetic nonindependence (Muir, 2015). These pat-

terns are consistent with other data indicating that many herbaceous plants are un-

der strong selection for high maximum photosynthetic rates. (Bazzaz, 1979; Körner

Although previous comparative studies have tested whether open habitat and growth form influence stomatal ratio, we do not know if these effects are independent of one another. Open habitat and growth form may not be independent because open habitats generally consist of more short-statured, herbaceous plants. Some authors have

et al., 1989). NEED MORE RECENT CITATIONS ON THIS.

- attempted to disentangle light and growth form by contrasting herbs from open and understory habiats (Salisbury, 1927). However, this is problematic if phylogenetic relationships are not controlled for, because shade species may share traits simply because they are more closely related to each other than they are to high light species. Finally, open habitat and groth form may also interact with one another. For example, amphistomy may only be favored when CO<sub>2</sub> strongly limits photosynthetic rate (e.g. in high light) and photosynthetic rate strongly limits fitness (e.g. in herbs).
- To better understand the adapative significance of stomatal ratio, I asked three main questions:
  - 1. Are light habitat and growth form correlated?

97

- 2. Do light habitat and growth form influence stomatal ratio additively, or do their effects interact?
- 3. Is evolution of stomatal ratio mediated by changes in stomatal density on the adaxial (upper) surface, abaxial (lower) surface, or both?
- The final question is important for telling whether amphistomy is part of a coordinated syndrome of traits that promote higher photosynthetic rate, as both the light and growth form hypotheses assume. If evolved increases in stomatal ratio are medi-

ated by shifting abaxial stomata to the adaxial surface, holding total stomatal density
constant, then the overall increase in CO<sub>2</sub> diffusion would be limited. In contrast,
if amphistomy evolves by increasing adaxial stomatal density while holding abaxial
density constant, then total stomatal density must increase as well. Evolutionary
coordination of amphistomy and high stomatal density would reinforce one another,
increasing CO<sub>2</sub> supply to chloroplasts more than changes in either trait would in
isolation.

To address these questions, I reanalyzed existing data on stomatal ratio, light habitat, and growth form in British angiosperms (Salisbury, 1927; Fitter and Peat, 1994; BEF) using phylogenetic comparative methods. The British angiosperm flora is well suited for these questions because this flora has been comprehensively surveyed for 115 many ecologically important traits, meaning it is probably the least biased survey 116 of stomatal trait variation. Salisbury's observations on stomata and ecology in the 117 British flora have heavily influenced plant ecophysiology, but many of his and subse-118 quent authors' analyses have significant limitations because of inadequate statistical 119 methods. For example, few analyses until recently account for phylogenetic nonin-120 dependence (Felsenstein, 1985), which can strongly influence inferences on stomatal 121 traits and growth form (Kelly and Beerling, 1995, this study did not consider light). 122 A species-level phylogeny of the entire British flora (Lim et al., 2014) now allows for 123

the first time rigorous analysis of evolutionary relationships among stomatal ratio, light, and growth form.

#### 6 METHODS

Data and annotated source code to generate this manuscript are available on GitHub (https://github.com/cdmuir/britstom) and Dryad (Muir, 2017).

## Data on stomatal ratio, light habitat, growth form, and phy-

#### 130 logenetic relationships

I obtained data on ab- and adaxial stomatal density on 395 species from British
Ecological Flora (Salisbury, 1927; Fitter and Peat, 1994; BEF). Following recent
comparative analyses (e.g. Bartelheimer and Poschlod, 2016; Salguero-Gómez et al.,
2016), I used Ellenberg light indicator values (Ellenberg, 1974) and Raunkiær life
form (Raunkiær, 1934) as measures of light habitat and growth form, respectively.
Hence, I am assuming that the species' light habitat is closely related to the type of
habitat (open versus closed) where that species is found. Both attributes have been
recently updated by taxonomic experts of the British flora (PLANTATT, Hill et al.

(2004)). Ellenberg light indicator values are hereafter abbreviated L-value. I used a dated molecular phylogeny of the British flora (Lim et al., 2014) available from 140 TreeBASE (http://treebase.org/; accession number 15105). 14 species (3.5%) in the 141 dataset were not present in the phylogeny. For 8 of these species, I used the position a 142 congeneric species as a proxy for the focal species. When multiple congeneric species 143 were present, I consulted the phylogenetic literature to identify the most closely 144 related proxy species (Scheen et al., 2004; Salmaki et al., 2013). For the remaining 6 missing species, I positioned them in the tree based on phylogenetic relationships 146 to other genera or families present in the tree (Fior et al., 2006). Because many 147 phylogenetic comparative methods do not allow polytomies, zero-length branches, 148 and non-ultrametric trees, I made several small adjustments to the tree. I resolved polytomies randomly using the 'multi2di' function in **phytools** version 0.5-64 (Revell, 150 2012). I added 0.02 my to all zero-length branches, as this was approximately the 151 length of the shortest nonzero branch length in the tree. After these changes, I 152 slightly altered terminal branch lengths to make the tree precisely ultrametric. I excluded data on hyrdrophytes (14 species) because many of these species are hyperstomatous (Fig. S1) due to the fact that leaves may rest on the water's surface, 155 selecting for stomata to be present on the upper surface only. I also excluded C<sub>4</sub> 156

(3 species) and CAM (2 species) plants. I limited this investigation to angiosperms

because only 4 non-angiosperms had stomata data. The final dataset contained 372 species. The R code accompanying this paper documents these decisions with citations to the relevant literature.

Following Muir (2015), I calculated stomatal ratio in two different ways depending on what was most appropriate for the question:

$$SR_{propAd} = \frac{SD_{ad}}{SD_{total}}$$
 (1)

$$SR_{even} = \frac{\min\{SD_{ab}, SD_{ad}\}}{\max\{SD_{ab}, SD_{ad}\}}$$
(2)

 $SD_{ab}$  and  $SD_{ad}$  are the stomatal densities on abaxial or adaxial surface, respectively.  $SD_{total} = SD_{ab} + SD_{ad}$ .  $SR_{propAd}$  is the proportion of stomata density on the adaxial  $SD_{total} = SD_{ab} + SD_{ad}$ .  $SR_{propAd}$  is the proportion of stomata density on the adaxial  $SD_{total} = SD_{ab} + SD_{ad}$ .  $SR_{propAd} = 0$ ,  $SR_{propAd} = 0$ ,  $SR_{propAd} = 0$ ,  $SR_{even}$  indicates how evenly stomatal densities are distributed across both leaf surfaces. This expression is useful because several hypotheses are based on the fact that  $SR_{even} = 0$ , and  $SR_{even} = 0$ , and  $SR_{even} = 0$ , are distributed across both leaf surfaces. This expression is useful because several hypotheses are based on the fact that

# Testing for an association between open habitat and growth form

I tested whether Raunkiær life form was associated L-value among British angiosperms using ANOVA with Type-2 sum of squares. I did not use phylogenetic ANOVA for this test because there was no phylogenetic signal in the regression fit using **phylolm** version 2.5 (Ho and Ané, 2014). See the R code accompanying this paper for further detail. I predicted that species with faster life histories, especially therophytes (annuals), would have greater L-values than species with slower life histories, especially phanerophytes, which are mostly long-lived trees.

#### Open habitat, growth form, and stomatal ratio

I compared phylogenetic linear models to test whether Raunkiær life form, L-value, or interactions between them predicted SR<sub>even</sub>. I used SR<sub>even</sub> rather than SR<sub>propAd</sub> as the response variable because the hypothesis is that faster life history and/or high light favor more even stomatal densities on each surface. I fit models using **phylolm** and extracted Akaike Information Criteria (AIC). For these and subsequent analyses, I assumed an Ornstein-Uhlenbeck process model for the residuals with the root character state integrated over the stationary distribution. I used a 10,000 para-

metric bootstrap samples of the full model (including main effects and interactions) to calculate parameter confidence intervals (Boettiger et al., 2012). Likewise, to 188 determine whether the interaction between Raunkiær life form and L-value was sta-189 tistically significant, I used a parametric bootstrap to generate the null distribution 190 of  $\triangle$ AIC values ( $\triangle$ AIC is the difference in AIC between competing models). Specif-191 ically, I sampled 1000 random datasets from the estimated model with main effects 192 of Raunkiær life form and L-value but no interaction. I fit these simulated datasets to models with and without interactions and calculated  $\Delta$ AIC. The statistical signif-194 icance of the observed  $\triangle AIC$  is the proportion of simulated  $\triangle AIC$  greater than the 195 observed.

# Does ab- or adaxial stomatal density contribute more to stomatal ratio evolution?

I used two complementary phylogenetic methods to assess the relative contribution of
ab- versus adaxial stomatal density to light-mediated stomatal ratio evolution. The
contribution of each can be formalized using standard variance decomposition methods as derived below. Because stomatal density is highly skewed, I log-transformed
values for normality:

$$SR_{even} = \frac{SD_{ad}}{SD_{ab}} \tag{3}$$

$$\log(SR_{even}) = \log(SD_{ad}) - \log(SD_{ad})$$
(4)

$$sr_{even} = sd_{ad} - sd_{ad}$$
 (5)

Lowercase variables (sr, sd) indicate log-transformed values. Because some species had zero adaxial stomata, I added one to all values prior to log-transformation. 205 To make the variance decomposition calculations tractable, I have defined SR<sub>even</sub> 206 here as the ratio of ad- to abaxial stomatal density because in most cases adaxial 207 stomatal density is lower than abaxial (see Eq. 2). This was not done in previous 208 analyses because I wanted to test what factors influenced the evenness of stomatal 209 densities, regardless of which surface had higher density. With this modified form, 210 the variance in  $sr_{even}$  can be decomposed into contributions of  $sd_{ad}$ ,  $sd_{ab}$ , and their 211 covariance: 212

$$Var(sr_{even}) = Var(sd_{ad}) + Var(sd_{ad}) - 2Cov(sd_{ad}, sd_{ab})$$
(6)

I estimated the phylogenetic covariance matrix between L-value, sd<sub>ab</sub>, and sd<sub>ad</sub> using a multivariate Ornstein-Uhlenbeck model fit in **Rphylopars** version 0.2.9 (Goolsby et al., 2016, 2017). From the covariance matrix, I estimated the contribution of abaxial density, adaxial density, and their covariance as:

$$\frac{Var(sd_{ad})}{Var(sr_{even})}, \frac{Var(sd_{ab})}{Var(sr_{even})}, \text{ and } \frac{Cov(sd_{ad}, sd_{ab})}{Var(sr_{even})},$$
 (7)

respectively. Note that when ab- and adaxial densities positively covary, the contribution will be negative because this reduces the variance in stomatal ratio.

I was interested in whether light-mediated evolution of stomatal ratio acted mostly 219 by increasing adaxial stomatal density while maintaining abaxial density, or keeping 220 total stomatal density the same, but shifting a greater proportion to the adaxial sur-221 face. The first scenario predicts that the phylogenetic regression of L-value on  $\mathrm{sd}_{\mathrm{ad}}$  is 222 stronger than that for sd<sub>ab</sub>. The second scenario predicts that L-value acts similarly 223 on both and that there is a negative covariance  $Cov(sd_{ad}, sd_{ab}) < 0$ . I tested these 224 competing predictions by fitting a simple phylogenetic structural equation model (SEM). The model uses the phylogenetic covariance matrix to simultaneously estimate regressions of L-value on sd<sub>ad</sub> and sd<sub>ab</sub> while allowing covariance between them (i.e. estimating  $Cov(sd_{ad}, sd_{ab})$ ). To fit the SEM, I used the R package **lavaan** version  $_{229}$  0.5-23.1097 (Rosseel, 2012). I tested whether parameter estimates were significantly different from zero using z-scores.

#### 231 RESULTS

#### Light tolerance varies among Raunkiær life forms

Ellenberg light indicator values (L-value) differed significantly among life forms (Fig. 1;ANOVA -  $F_{4,367} = 18.3$ ,  $P = 1.1 \times 10^{-13}$ ). Therophytes (annuals), hemicryptophytes (perennial herbs with buds near the soil surface), and chamaephytes (subshrubs) had greater L-values than phanerophytes (large woody plants) and geophytes (perennial herbs with storage organs) (Fig. 1).

# Interactions between light and Raunkiær life form determine

#### 239 stomatal ratio

Overall,  $SR_{even}$  increased with L-value, but there was a significant interaction ( $\Delta AIC > 241$  2, Table 1) between Raunkiær life form and L-value (Fig. 2). Both life form and L-value significantly increased model fit, though L-value had a markedly larger effect on model AIC (Table 1). The significant interaction is caused by different slopes

between life forms. Among life forms with the overall greatest L-value (therophytes, hemicryptophytes, and chamaephytes, see Fig. 1), there was a strong positive rela-245 tionship between L-value and SR<sub>even</sub>. Parametrically bootstrapped 95% confidence 246 intervals did not overlap zero (Fig. 2). The slope was weakly positive or not sig-247 nificantly different from zero in the most shade-adapted life forms (geophytes and phanerophytes), albeit the patterns were distinct in these groups. There were both 249 hypostomatous ( $SR_{even} \approx 0$ ) and amphistomatous ( $SR_{even} \approx 1$ ) geophytes, but these 250 were distibuted across L-values. In contrast, phanerophytes were nearly always hy-251 postomatous regardless of L-value. Allowing slopes to vary across life form signicantly 252 increased model fit (lower AIC, Table 1).

# Adaxial stomatal density contributes most of the variation in stomatal ratio

Adaxial ('upper') stomatal density contributed most to the evolutionary variation in stomatal ratio. The contributions of adaxial density, abaxial density, and their covariance are 1.14, 0.38, and -0.53, respectively. Recall that values can be greater than one for adaxial stomatal density and negative for the covariance when the latter value is positive. This implies that evolutionary variation in adaxial stomatal density is greater than that for stomatal ratio due to positive covariance between ab- and adaxial stomatal density.

Similarly, the phylogenetic SEM showed that changes in stomatal ratio associated 263 with L-value can be attributed mostly to evolution of adaxial stomatal density 264 (Fig. 3). Both sd<sub>ad</sub> and sd<sub>ab</sub> increased with L-value ( $P = 6.1 \times 10^{-7}$  and  $2.9 \times 10^{-5}$ , 265 respectively). However, the regression of L-value on sd<sub>ad</sub> was 2.1× that of L-value on 266  $\mathrm{sd}_{\mathrm{ab}}$  (0.21 versus 0.1). Because stomatal densities were natural log-transformed, this 267 implies an increase in L-value by one leads to a 1.23-fold change in adaxial stomatal 268 density versus a 1.1-fold change in abaxial stomatal density. The SEM also showed a significant positive covariance between stomatal densities on each surface (P = 1.7 $\times 10^{-11}$ ). These results together imply that total stomatal density increases with 271 L-value, but the response is mediated mostly by adaxial stomatal density.

#### 273 DISCUSSION

The ratio of stomatal densities on the abaxial ('lower') to that of the adaxial ('upper')
surface varies greatly across plant species, but the adaptive significance is not clear.
Comparative studies correlating stomatal ratio to ecological factors can distinguish
among competing hypotheses and reveal critical experiments for future work. Previ-

ous comparative studies suggested that high light and herbaceous growth form favor amphistomy (Mott et al., 1984; Jordan et al., 2014; Muir, 2015; Bucher et al., 2017), 279 particularly in the British flora (Salisbury, 1927; Peat and Fitter, 1994). However, 280 none of these studies have accounted for the fact thats light and growth form are often 281 confounded – open, high light habitats are necessarily dominated by herbs – or the 282 fact that species are not independent because of shared evolutionaey hisory. Here, I 283 reanalyzed data on stomata, light tolerance, and growth form in British angiosperms 284 using phylogenetic comparative methods. As expected, species' light tolerance (El-285 lenberg light indicator or L-value) is confounded with growth form (Raunkiær life 286 form; Fig. 1). Nevertheless, both L-value and Raunkiær life form affect stomatal 287 ratio, but these factors also interact; the influence of L-value on stomatal ratio varies 288 across forms. These novel findings provide further evidence that variation in stomatal 289 ratio is adaptive and have important implications for interpreting changes in stomatal ratio through the paleo record (Jordan et al., 2014) and during domestication 291 (Milla et al., 2013). 292

#### 293 Adaptive significance of amphistomy

Previously, associations between light, growth form, and stomatal ratio have been interpreted in isolation as indicating that either high light and/or herbaceous growth

form favors amphistomy. In British angiosperms, both factors are important, though statistical analyses suggest that light may be a stronger determinant than growth 297 form (Table 1). Unlike previous studies, I found a significant interaction between 298 light and growth form among British angiosperms, which suggests that amphistomy 290 may only be strongly favored when CO<sub>2</sub> strongly limits photosynthesies and pho-300 tosynthesis strongly limits fitness. The ideal way to test this would be to measure 301 selection on stomatal ratio in a species that varied quantitatively in both stomatal 302 ratio and life history (e.g. containing both annual and perennial forms). I predict 303 that amphistomy will be favored much more strongly in the annual form grown under 304 high light compared to an annual under low light or a perennial in high light. Similar 305 experiments could also be performed to test if and when light-mediated plasticity in 306 stomatal ratio is adaptive (Gay and Hurd, 1975; Mott and Michaelson, 1991). 307 The prevalence of amphistomatous species in high light habitats supports the hy-308 pothesis that amphistomy is an adaptation to maximize photosynthetic rates by 309 increasing CO<sub>2</sub> diffusion (Jones, 1985). It is also evidence against the hypothesis 310 that the principle fitness cost of amphistomy is water loss (Darwin, 1886; Foster 311 and Smith, 1986) or dehydration of pallisade mesophyll (Buckley et al., 2015). Since 312 evaporative demand increases under high insolation, under these hypotheses we would 313 expect plants in high light to be hypostomatous. Because stomatal conductances on

each surface can be regulated independently in response to the environment (Darwin, 315 1898; Pospíŝilová and Solárová, 1984; Smith, 1981; Reich, 1984; Mott and O'Leary, 316 1984), amphistomatous leaves likely cope with these stresses by rapidly closing adax-317 ial stomata when water supply cannot match evaporative demands. Instead, pat-318 terns in the British flora are at least consistent with the idea that adaxial stomata 319 increase susceptibility to foliar pathogens (Gutschick, 1984; McKown et al., 2014). 320 The cost of adaxial stomata may be greater in the shade because greater leaf wet-321 ness and lower ultraviolet light provide a more suitable microclimate for many foliar 322 pathogens. 323

#### Amphistomy as a proxy for open habitat

These observations from the British flora strongly support the hypothesis that amphistomy can be used a proxy for open habitat in paleoenvironment reconstruction (Carpenter, 1994; Jordan et al., 2014; Carpenter et al., 2015), but also point out previously unknown subtleties. These previous studies based their conclusions on data from Proteaceae, in which there is little quantitative variation in stomatal ratio; species are either completely hypostomatous ( $SR_{propAd} \approx 0$ ) or completely amphistomatous ( $SR_{propAd} \approx 0$ ). Stomatal ratio in British angiosperms is also bimodal (Peat and Fitter, 1994), but across many families there is also quantitative variation.

Importantly, this means that quantitative variation in stomatal ratio may provide a more precise, quantitative indicator of vegetation type, rather than simply 'open' or 'closed'. A quantitative relationship between L-value and stomatal ratio has already been shown for herbaceous perennials (Bucher et al., 2017), but we now know that it holds among annuals (therophytes), subshrubs (chamaephytes), and, to a lesser extent, geophytes as well (Fig. 2).

The nonsignificant relationship between L-value and stomatal ratio in geophytes and 339 phanerophytes suggests that in some cases amphistomy may not reliably indicate open habitat without further information. For example, perhaps amphistomatous geophytes from partially shaded habitats are spring ephemerals, so they experience high light during their growth phase, but this has not been tested. Likewise, phanerophytes (most tall trees) are almost always hypostomatous (see also Muir (2015)). Most British phanerophytes are tall, hypostomatous trees, but the exceptions are 345 For example, the most amphistomatous phanerophyte in this dataset is 346 Brassica oleracea, a short-statured biennial that has more in common physiologically with hemicryptophytes than other phanerophytes. The other amphistomatous 348 phanerophytes in this data set (*Populus nigra* and *Lavatera arborea*) are fast-growing 349 pioneer species.

351 Finally, phylogenetic information should improve inferences about paleoclimates be-

cause there is appreciable phylogenetic signal in stomatal ratio. The phylogenetic half-life of stomatal ratio evolution, after accounting for L-value and Raunkiær life form, is  $\log(2)/\alpha = 1.5$  my (see Table 1 for maximum likelihood estimates of  $\alpha$ ). This lag time may indicate that evolving to the 'optimum' is constrained by the shape of the fitness landscape (Muir, 2015) or that other unmeasured factors which affect stomatal ratio have some phylogenetic signal. Regardless of the mechanism, this fact means that researchers may be able to use data from closely related species to improve paleoenvironment reconstruction.

#### Why does adaxial stomatal density control stomatal ratio?

Variation in stomatal ratio is determined primarily by evolution of adaxial stomatal
density and is coordinated with increases in total leaf stomatal density summed across
both surfaces. Phylogenetic analyses show that changes in stomatal ratio and total
stomatal density, especially in response to L-value, are predominantly mediated by
changes in adaxial stomatal density. This highly nonrandom pattern among British
angiosperms mirrors evolutionary changes wrought by domesication (Milla et al.,
2013); crops species tend to have higher adaxial stomatal density than their wild
relatives. Note here that I am referring only to evolutionary variation in stomatal
ratio among species; different processes may mediate within species variation or

plastic responses.

There are at least two hypotheses that could explain why adaxial stomatal density 371 is the most responsive. The first I refer to as the 'real estate' hypothesis. In hy-372 postomatous plants, the lower surface is already crowded with stomata, and hence 373 plants must increase the real estate available for stomata by develoing them on the 374 upper surface whenever there is selection for greater stomatal density. When stomata 375 are packed too densely on one surface, stomatal interference limits their function-376 ing and hence may create a strong selective pressure for amphistomy (Parlange and 377 Waggoner, 1970; Dow et al., 2014). I refer to the second hypothesis as the 'coordination' hypothesis. In this scenario, 379 ecological conditions such as high light select for both increased total stomatal density 380 and for amphistomy because these traits work well in coordination with one another. 381 For example, if stomatal density were very high on a hypostomatous plant, then CO<sub>2</sub> 382 would be more strongly limited by the mesophyll. Adding a second parallel pathway 383 for diffusion by developing stomata on both surfaces would restore a more optimal 384 balance between stomatal and mesophyll limitations. Conversely, there would be 385 little benefit to amphistomy when total stomatal density is low because CO<sub>2</sub> diffusion 386 is strongly limited by stomatal resistance, and therefore photosynthetic rate is not 387 sensitive to changes in mesophyll diffusion mediated by stomatal ratio.

 $_{389}$  Conclusions - finish when analysis is complete

### References

- 391 ???? URL http://www.ecoflora.co.uk.
- Bartelheimer, M. and P. Poschlod, 2016. Functional characterizations of Ellenberg
- indicator values—a review on ecophysiological determinants. Functional Ecology
- 30:506-516.
- Bazzaz, F., 1979. The physiological ecology of plant succession. Annual Review of
- Ecology and Systematics 10:351–71.
- Beerling, D. J. and C. K. Kelly, 1996. Evolutionary comparative analyses of the
- relationship between leaf structure and function. New Phytologist 134:35–51.
- Boettiger, C., G. Coop, and P. Ralph, 2012. Is your phylogeny informative? mea-
- suring the power of comparative methods. Evolution 66:2240–2251.
- Bucher, S. F., K. Auerswald, C. Grün-Wenzel, S. I. Higgins, J. G. Jorge, and
- 402 C. Römermann, 2017. Stomatal traits relate to habitat preferences of herbaceous
- species in a temperate climate. Flora.
- Buckley, T. N., G. P. John, C. Scoffoni, and L. Sack, 2015. How does leaf anatomy
- influence water transport outside the xylem? Plant Physiology 168:1616–1635.
- 406 Carpenter, R. J., 1994. Cuticular morphology and aspects of the ecology and

- fossil history of North Queensland rainforest Proteaceae. Botanical Journal of
- the Linnean Society 116:249. URL + http://dx.doi.org/10.1111/j.1095-
- 8339.1994.tb00434.x.
- <sup>410</sup> Carpenter, R. J., M. K. Macphail, G. J. Jordan, and R. S. Hill, 2015. Fossil evidence
- for open, Proteaceae-dominated heathlands and fire in the Late Cretaceous of
- Australia. American Journal of Botany 102:2092–2107.
- Darwin, F., 1886. On the relation between the "bloom" on leaves and the distribution
- of the stomata. Botanical Journal of the Linnean Society 22:99–116.
- 415 ———, 1898. Observations on stomata. Philosophical Transactions of the Royal
- Society B: Biological Sciences 190:531–621.
- Dow, G. J., J. A. Berry, and D. C. Bergmann, 2014. The physiological importance
- of developmental mechanisms that enforce proper stomatal spacing in *Arabidopsis*
- thaliana. New Phytologist 201:1205–1217.
- Ellenberg, H., 1974. Indicator values of vascular plants in central Europe, Scripta
- 421 Geobotanica, vol. 9. Springer-Verlag, Göttingen, Germany.
- Felsenstein, J., 1985. Phylogenies and the comparative method. The American
- 423 Naturalist 1:1–15.
- Fior, S., P. O. Karis, G. Casazza, L. Minuto, and F. Sala, 2006. Molecular phylogeny

- of the Caryophyllaceae (Caryophyllales) inferred from chloroplast matk and nuclear
- rDNA ITS sequences. American Journal of Botany 93:399–411.
- Fitter, A. and H. Peat, 1994. The ecological flora database. Journal of Ecology 82:415–425.
- Foster, J. and W. Smith, 1986. Influence of stomatal distribution on transpiration
- in low-wind environments. Plant, Cell & Environment 9:751–759.
- Gay, A. and R. Hurd, 1975. The influence of light on stomatal density in the tomato.
- New Phytologist 75:37–46.
- 433 Gibson, A. C., 1996. Structure-Function Relations of Warm Desert Plants. Springer-
- Verlag, Berlin.
- Givnish, T. J., 1987. Comparative studies of leaf form: assessing the relative roles
- of selective pressures and phylogenetic constraints. New Phytologist 106:131–160.
- 437 Goolsby, E. W., J. Bruggeman, and C. Ané, 2016. Rphylopars: Phyloge-
- netic Comparative Tools for Missing Data and Within-Species Variation. URL
- https://CRAN.R-project.org/package=Rphylopars. R package version 0.2.9.
- 440 ——, 2017. Rphylopars: fast multivariate phylogenetic comparative methods for
- missing data and within-species variation. Methods in Ecology and Evolution
- 8:22-27.

- Gutschick, V. P., 1984. Photosynthesis model for C<sub>3</sub> leaves incorporating CO<sub>2</sub> trans-
- port, propagation of radiation, and biochemistry 2. ecological and agricultural
- utility. Photosynthetica 18:569–595.
- Haberlandt, G., 1914. Physiological Plant Anatomy. Macmillan and Co., London.
- 447 Hill, M., C. Preston, and D. Roy, 2004. PLANTATT Attributes of British and Irish
- Plants: Status, Size, Life History, Geography and Habitats. Centre for Ecology &
- 449 Hydrology, Huntingdon, Cambridgeshire.
- 450 Ho, L. S. T. and C. Ané, 2014. Intrinsic inference difficulties for trait evolution with
- Ornstein-Uhlenbeck models. Methods in Ecology and Evolution 5:1133–1146.
- Jones, H. G., 1985. Adaptive significance of leaf development and structural responses
- to environment. Pp. 155–173, in N. R. Baker, W. Davies, and C. K. Ong, eds.
- 454 Control of Leaf Growth, Society for Experimental Biology Seminar Series, vol. 27.
- 455 Cambridge University Press, Cambridge.
- 456 Jordan, G. J., R. J. Carpenter, and T. J. Brodribb, 2014. Using fossil leaves as
- evidence for open vegetation. Palaeogeography, Palaeoclimatology, Palaeoecology
- 458 395:168–175.
- 459 Kelly, C. and D. Beerling, 1995. Plant life form, stomatal density and taxonomic
- relatedness: a reanalysis of Salisbury (1927). Functional Ecology 9:422–431.

- Körner, C., M. Neumayer, S. P. Menendez-Riedl, and A. Smeets-Scheel, 1989. Functional morphology of mountain plants. Flora 182:353–383.
- Lim, J., M. J. Crawley, N. De Vere, T. Rich, and V. Savolainen, 2014. A phylogenetic
- analysis of the British flora sheds light on the evolutionary and ecological factors
- driving plant invasions. Ecology and Evolution 4:4258–4269.
- McKown, A. D., R. D. Guy, L. Quamme, J. Klápště, J. La Mantia, C. Constabel,
- 467 Y. A. El-Kassaby, R. C. Hamelin, M. Zifkin, and M. Azam, 2014. Association
- genetics, geography and ecophysiology link stomatal patterning in *Populus tri*-
- chocarpa with carbon gain and disease resistance trade-offs. Molecular Ecology
- 470 23:5771-5790.
- Metcalfe, C. R. and L. Chalk, 1950. Anatomy of the dicotyledons, Vols. 1 & 2. First
- ed. Oxford University Press, Oxford.
- Milla, R., N. de Diego-Vico, and N. Martín-Robles, 2013. Shifts in stomatal traits
- following the domestication of plant species. Journal of Experimental Botany
- 64:3137–3146.
- 476 Mott, K. A., A. C. Gibson, and J. W. O'Leary, 1984. The adaptive significance of
- amphistomatic leaves. Plant, Cell & Environment 5:455–460.
- 478 Mott, K. A. and O. Michaelson, 1991. Amphistomy as an adaptation to high light

- intensity in Ambrosia cordifolia (Compositae). American Journal of Botany 78:76– 479 79.
- Mott, K. A. and J. W. O'Leary, 1984. Stomatal behavior and CO<sub>2</sub> exchange char-481

acteristics in amphistomatous leaves. Plant physiology 74:47–51.

- Muir, C. D., 2015. Making pore choices: repeated regime shifts in stomatal ratio.
- Proc. R. Soc. B 282:20151498. 484

480

482

- 2017. from: Hight light interacts Data with 485
- form favor amphistomy in British angiosperms. URL to 486
- http://dx.doi.org/10.5061/dryad.????? 487
- Parkhurst, D. F., 1978. The adaptive significance of stomatal occurrence on one or
- both surfaces of leaves. The Journal of Ecology 66:367–383. 489
- Parkhurst, D. F. and K. A. Mott, 1990. Intercellular diffusion limits to CO<sub>2</sub> uptake
- in leaves studied in air and helox. Plant Physiology 94:1024–1032. 491
- Parlange, J.-Y. and P. E. Waggoner, 1970. Stomatal dimensions and resistance to 492
- diffusion. Plant Physiology 46:337–342. 493
- Peat, H. and A. Fitter, 1994. A comparative study of the distribution and density of
- stomata in the British flora. Biological Journal of the Linnean Society 52:377–393. 495

- Pospíŝilová, J. and J. Solárová, 1984. Environmental and biological control of diffu-
- sive conductances of adaxial and abaxial leaf epidermes. Photosynthetica 18:445–
- 498 453.
- Raunkiær, C. C., 1934. The Life Forms of Plants and Statistical Plant Geography.
- 500 Clarendon Press, Oxford.
- Reich, P., 1984. Relationships between leaf age, irradiance, leaf conductance, CO<sub>2</sub>
- exchange, and water-use efficiency in hybrid poplar. Photosynthetica 18:445–453.
- Revell, L. J., 2012. phytools: An R package for phylogenetic comparative biology
- (and other things). Methods in Ecology and Evolution 3:217–223.
- Rosseel, Y., 2012. lavaan: An R package for structural equation modeling. Journal
- of Statistical Software 48:1–36.
- Salguero-Gómez, R., O. R. Jones, E. Jongejans, S. P. Blomberg, D. J. Hodgson,
- 508 C. Mbeau-Ache, P. A. Zuidema, H. de Kroon, and Y. M. Buckley, 2016. Fast—
- slow continuum and reproductive strategies structure plant life-history variation
- worldwide. Proceedings of the National Academy of Sciences of the United States
- of America 113:230–235.
- Salisbury, E., 1927. On the causes and ecological significance of stomatal frequency,

- with special reference to the woodland flora. Philosophical Transactions of the Royal Society of London. Series B 216:1–65.
- Salmaki, Y., S. Zarre, O. Ryding, C. Lindqvist, C. Bräuchler, G. Heubl, J. Barber,
- and M. Bendiksby, 2013. Molecular phylogeny of tribe Stachydeae (Lamiaceae
- subfamily Lamioideae). Molecular Phylogenetics and Evolution 69:535–551.
- Scheen, A.-C., C. Brochmann, A. K. Brysting, R. Elven, A. Morris, D. E. Soltis, P. S.
- Soltis, and V. A. Albert, 2004. Northern hemisphere biogeography of Cerastium
- (Caryophyllaceae): insights from phylogenetic analysis of noncoding plastid nu-
- cleotide sequences. American Journal of Botany 91:943–952.
- Smith, W., 1981. Temperature and water relation patterns in subalpine understory plants. Oecologia 48:353–359.
- Smith, W. K., T. C. Vogelmann, E. H. DeLucia, D. T. Bell, and K. A. Shepherd,
  1997. Leaf form and photosynthesis. BioScience 11:785–793.

Table 1: Interaction between species' Ellenberg light indicator value (L-value) and Raunkiær lifeform shape stomatal ratio (SR<sub>even</sub>). I compared phylogenetic linear models using the Akaike Information Criterion (AIC), where AIC =  $2k - 2\log(\mathcal{L})$ . k is the number of model parameters and  $\mathcal{L}$  is the model likelihood. Given a set of candidate models, the difference in AIC between a model and the lowest AIC ( $\Delta$ AIC) indicates the relative fit of competing models. The correlation coefficient  $r^2$  is another indicator of model fit.  $\alpha$  and  $\sigma^2$  are the return rate and diffusion parameters of the Ornstein-Uhlenbeck model of trait evolution.

Model: $SR_{even} \sim$	$\alpha$	$\sigma^2$	$r^2$	k	$\log(\mathcal{L})$	AIC	$\Delta { m AIC}$
L-value $\times$ lifeform	0.46	0.068	0.34	12	-33.3	90.6	0
L-value + lifeform	0.47	0.072	0.32	8	-40.3	96.5	6
L-value	0.64	0.108	0.26	4	-59.3	126.6	36.1
lifeform	0.34	0.067	0.15	7	-79.2	172.4	81.8
1	0.29	0.067	0	3	-107.5	221	130.5

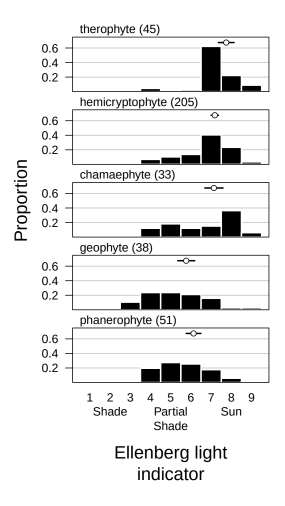


Figure 1: Lifeforms have different tolerances for sun and shade among British angiosperms. Each panel is the distribution of Ellenberg light indicator values on an integer scale of 1-9 for different Raunkiær life forms. Height of the bars indicate the raw proportion of species in each bin for that lifeform. The sample size for each lifeform is listed next in parentheses. The mean (open circle) and 95% confidence intervals (black line) around the mean Ellenberg light indicator value for each lifeform based on phylogenetic regression are above the histogram.

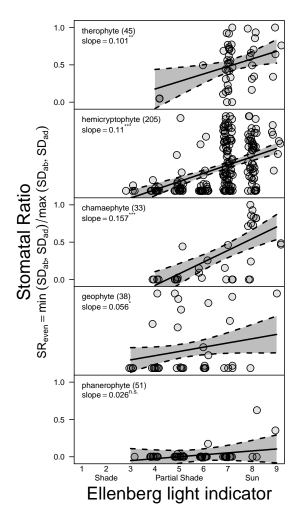


Figure 2: The effect of light on stomatal ratio depends on Raunkiær life form. Greater Ellenberg light indicator values (L-value) are associated with greater stomatal ratio (SR<sub>even</sub>) in therophytes, hemicryptophytes, and chamaephytes but not geophytes and phanerophytes. The maximum likelihood slope from phylogenetic regression is given with statistical significance based on 1000 parametric bootstrap samples. Numbers in parentheses next to Raunkiær life form are the sample sizes in the final dataset. Estimated slopes (solid line) and 95% bootstrapped confidence intervals (gray polygon between dashed lines) are plotted against raw data. Points have been jittered for visial clarity.

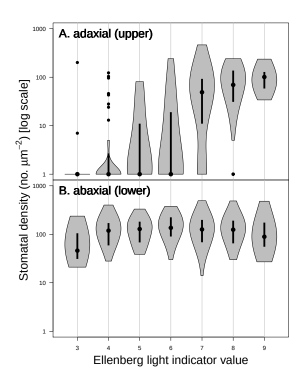


Figure 3: Light-mediated evolution of stomatal ratio is mostly driven by increased adaxial ('upper') stomatal density (Panel A), whereas abaxial ('lower') stomatal density (Panel B) is similar across Ellenberg light indicator values (L-value x-axis). The violin plot shows stomatal density (y-axis, log-scale) as a function of L-value. The width of the grey polygons indicates the density of data. Length of grey polygon indicate the range of the data; the point indicates the median; the thick lines indicate the 0.25 and 0.75 quantiles. Points outside the polygons are statistical outliers.

# 526 Supporting Information

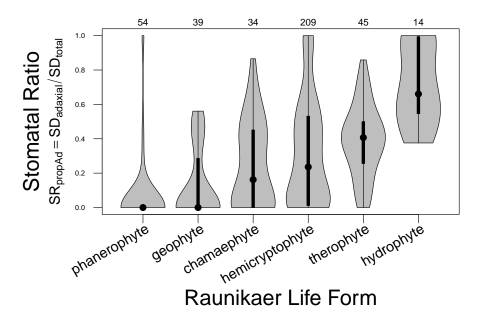


Figure S1: Most hydrophytes are hyperstomatous, having most stomata on the adaxial ('upper') surface (high  $SD_{propAd}$ ). The violin plot shows stomatal ratio as a function of Raunkiær lifeform. The width of the grey polygons indicates the density of data. Length of grey polygon indicate the range of the data; the point indicates the median; the thick lines indicate the 0.25 and 0.75 quantiles. Sample sizes per lifeform in the dataset are given above the upper plot margin.  $SD_{ad}$  and  $SD_{total}$  stand for the stomatal density on the adaxial surface and the total leaf surface (adaxial plus abaxial), respectively.