# Amphistomy increases leaf photosynthesis more in coastal than montane plants of Hawaiian 'ilima (Sida fallax)

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## 12 ABSTRACT —

Premise of the study The adaptive significance of stomata on both upper and lower leaf surfaces, called amphistomy, is unresolved. A widespread association between amphistomy and open, sunny habitats suggests the adaptive benefit of amphistomy may be greatest in these contexts, but this hypothesis has not been tested experimentally. Understanding amphistomy informs its potential as a target for crop improvement and paleoenvironment reconstruction.

Methods We developed a method to quantify "amphistomy advantage", AA, as the log-ratio of photosynthesis in an amphistomatous leaf to that of the same leaf but with gas exchange blocked through the upper surface (pseudohypostomy). Humidity modulated stomatal conductance and thus enabled comparing photosynthesis at the same total stomatal conductance. We estimated AA and leaf traits in six coastal (open, sunny) and six montane (closed, shaded) populations of the indigenous Hawaiian species 'ilima (Sida fallax).

Key results Coastal 'ilima leaves benefit 4.04 times more from amphistomy than montane leaves. Evidence was equivocal with respect to two hypotheses – that coastal leaves benefit more because 1) they are thicker and have lower CO<sub>2</sub> conductance through the internal airspace, and 2) that they benefit more because they have similar conductance on each surface, as opposed to most conductance being through the lower surface.

Conclusions This is the first direct experimental evidence that amphistomy increases photosynthesis, consistent with the hypothesis that parallel pathways through upper and lower mesophyll increase CO<sub>2</sub> supply to chloroplasts. The prevalence of amphistomatous leaves in open, sunny habitats can partially be explained the increased benefit of amphistomy in 'sun' leaves, but the mechanistic basis remains uncertain.

<sup>34</sup> **Keywords:** amphistomy, Hawai'i, leaf, light, Malvaceae, photosynthesis, *Sida fallax*, stomata

## 35 INTRODUCTION —

Amphistomy, the presence of stomata on both lower and upper surfaces of broad leaves, should in-36 crease carbon gain by reducing the average diffusion pathlength between stomata and chloroplasts, yet 37 paradoxically this seemingly simple adaptation is uncommon in nature and we don't know why. Un-38 derstanding variation in stomatal traits like amphistomy is imperative because these tiny pores play 39 an outsized ecological role in the global carbon and water cycles (Hetherington and Woodward, 2003; Berry et al., 2010). A widely applicable, accurate representation of how stomata mediate the relation-41 ship between CO<sub>2</sub> gained through photosynthesis and water lost through transpiration is essential to 42 predict future climate using Earth Systems Models (Jarvis, 1976; Ball et al., 1987; Collatz et al., 1991; 43 Leuning, 1995; Sellers et al., 1997). Optimality models accurately predict the major cause of water loss, 44 stomatal conductance  $(g_{sw})$ , by assuming plants maximize carbon gain minus a cost of water (Cowan 45 and Farquhar, 1977; Givnish, 1986; Medlyn et al., 2011; Lin et al., 2015; Wang et al., 2017; Franks et 46 al., 2018; Deans et al., 2020; Franklin et al., 2020; Wang et al., 2020; Harrison et al., 2021). Despite

the success of optimality modeling in predicting  $g_{\rm sw}$ , the same modeling approach has so far failed to explain the rarity of amphistomatous leaves (Muir, 2019). This gap between theory and observations strongly implies that we remain ignorant about some key benefits and costs associated with stomata.

Where are amphistomatous leaves found and why aren't they more common? Among terrestrial flower-52 ing plants, amphistomatous leaves are rarely found on woody plants and shade-tolerant herbs, but they 53 are common in annual and perennial herbs from sunny habitats (Salisbury, 1928; Parkhurst, 1978; Mott et al., 1982; Peat and Fitter, 1994; Gibson, 1996; Jordan et al., 2014; Muir, 2015, 2018; Bucher et al., 55 2017). Even in resupinate leaves where the abaxial surface faces up toward the sky, stomata develop on 56 the lower adaxial surface (Lyshede, 2002). Exceptions to this general pattern include some arid woody 57 plants which typically have vertically oriented, isobilateral leaves (Wood, 1934; Jordan et al., 2014; 58 Boer et al., 2016; Drake et al., 2019) and floating/amphibious leaves of aquatic plants (Kaul, 1976; 59 Doll et al., 2021). The dearth of amphistomatous leaves should be quite surprising and has been de-60 scribed as one of the most important unsolved problems in the study of leaf structure-function relations despite some recent progress (Grubb, 1977, 2020). 62

Amphistomatous leaves should be common because, all else being equal, a leaf with a given number 63 of stomata per area could increase its photosynthetic rate simply by apportioning approximately half its stomata to each surface (Parkhurst, 1978; Gutschick, 1984a, b). The key difference between a 65 hypo- and amphistomatous leaf, holding all other factors constant, is that an amphistomatous leaf has 66 two parallel diffusion paths through the internal airspace to any given chloroplast. Those airspaces 67 pose a resistance for CO<sub>2</sub> diffusion, so CO<sub>2</sub> concentration drops as it approaches chloroplasts. Shorter 68 pathways mean a smaller drop in CO<sub>2</sub> concentration. Thus, chloroplasts in amphistomatous leaves experience higher CO<sub>2</sub> concentrations than in hypostomatous leaves, thereby increasing photosynthesis. 70 The airspace resistance (or its inverse, the airspace conductance,  $g_{ias}$ ) is rarely measured directly and 71 there is disagreement between empirical (Parkhurst and Mott, 1990; Morison et al., 2005; Evans et al., 2009; Tomás et al., 2013; Earles et al., 2018; Šantrůček et al., 2019; Nobel, 2020; Harwood et al., 2021; 73 Márquez et al., 2023) and theoretical models (Tholen and Zhu, 2011; Ho et al., 2016; Théroux-Rancourt 74 et al., 2021). The  $g_{ias}$  in thin, porous leaves may be so large as to be inconsequential given much lower 75 conductances for other components of the diffusion pathway, whereas the  $g_{\rm ias}$  of thick leaves with 76 little airspace may greatly hinder CO<sub>2</sub> diffusion to chloroplasts. Amphistomy should confer the largest 77 photosynthetic benefit in leaves with intrinsically low  $g_{ias}$ . The airspace conductance is one component 78 of the overall mesophyll conductance,  $g_{\rm m}$ , which is often strongly influenced by the chloroplast surface 79 area exposed to airspace and mesophyll cell wall thickness (Evans et al., 2009; Gago et al., 2020; Flexas et al., 2021). Hence, thicker leaves may compensate for lower  $g_{\rm ias}$  through increased chloroplast surface 81 area exposed to airspace (Terashima et al., 2006), but will still benefit from amphistomy as long as  $g_{ias}$ 82 is finite. 83

Amphistomy should also enhance photosynthesis when leaf boundary layer resistance is high, because apportioning total flux between two boundary layers rather than one results in a smaller  ${\rm CO_2}$  concentration drop between the atmosphere and stomata. A similar effect has been validated with a computer model and measurements for transpiration: amphistomatous leaves lose somewhat more water for the same vapor pressure deficit and total  $g_{\rm sw}$  (Foster and Smith, 1986), but the additional carbon gain

should be enough to offset this cost under most realistic conditions (Muir, 2019). However, if minimal stomatal conductance is related to stomatal density (Drake et al., 2013; Márquez et al., 2022) and the upper boundary layer conductance is higher, then amphistomy could cause additional, unavoidable water loss.

The most promising adaptive hypothesis is that amphistomy is important for maximizing photosynthetic 93 rate under high light. Mott et al. (1982) proposed that "plants with a high photosynthetic capacity, 94 living in full-sun environments, and experiencing rapidly fluctuating or continuously available soil water" would benefit most, in terms of increased carbon gain, from having amphistomatous leaves. As described above, herbs from sunny habitats are often amphistomatous. Most variation in stomatal 97 density ratio (SR, the ratio of stomatal density between the upper and lower surfaces) among species 98 is assumed to be genetic, but there is also putatively adaptive plasticity in response to light. Leaves of 99 Ambrosia cordifolia, a desert perennial herb, are hypostomatous under low light (photosynthetic photon 100 flux density, PPFD = 110  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup>) but develop ~20% of their stomata on the upper surface 101 under high light (1700  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup>) (Mott and Michaelson, 1991). Similarly, Solanum lycopersicum 102 leaves are hypostomatous when grown in the shade but develop ~20% of their stomata on the upper 103 surface grown under high light-intensity (Gay and Hurd, 1975). Adult leaves of Eucalyptus globulus 104 are amphistomatous, but the proportion of adaxial stomata increases from ~10-20% under low light 105 to ~30-40% under high light (James and Bell, 2001). In summary, both genetic and plastic responses 106 evince a widespread association between light and SR. 107

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The association between high light and amphistomy suggests that 'sun' leaves have the most to gain in terms of increased photosynthesis from having stomata on both surfaces, as Mott et al. (1982) hypothesized. Parkhurst (1978) proposed quantifying this benefit as 'amphistomy advantage' (AA), which we adopt here with some modification (see Materials and Methods). This hypothesis has never been tested directly by comparing the photosynthetic rate of an amphistomatous leaf to that of an otherwise identical hypostomatous leaf with the same total stomatal conductance under the same conditions. We propose a straightforward method to do this by experimentally creating a pseudohypostomatous leaf with gas exchange blocked through the upper surface (see Materials and Methods). We use humidity to modulate stomatal conductance so that amphi- and pseudohypostomatous leaves can be compared at the same total stomatal conductance. One reason that sun leaves might have greater AA is that they are usually thicker or denser (Poorter et al., 2019), which will often result in lower  $g_{ias}$  either by increasing the diffusion path length (Parkhurst, 1978) or making the airspace less porous. A nonmutually exclusive hypothesis is that if sun leaves have a stomatal density ratio closer to 0.5 (same density on each leaf surface), this will confer a greater advantage than an amphistomatous leaf with most stomata on one surface. In other words, amphistomy doesn't make much difference if one leaf surface has few open stomata on it. We therefore predict that sun leaves will have greater AA possibly because they have thicker leaves and/or SR closer to 0.5. We actually report  $g_{\text{smax,ratio}}$ , which is similar to SR except that it accounts for differences in both stomatal density and size between surfaces.

The native flora of the Hawaiian archipelago is a excellent system to test the relationship between light habitat and AA. Many lineages have adapted to different light habitats after colonization and leaf anatomical traits such as SR and thickness vary within and among closely related species. It is hypothesized that the common ancestor in many Hawaiian clades was a weedy species with high dispersal

ability adapted to open habitats (Carlquist, 1966). Colonization was followed by adaptive radiation into higher elevation, montane, closed, forested habitats. Consequently, adaptation to sun and shade is a common axis of phenotypic variation among Hawaiian plants such as lobeliads (Givnish et al., 2004; Montgomery and Givnish, 2008; Givnish et al., 2009; Givnish and Montgomery, 2014; Scoffoni et al., 2015), *Bidens* (Carlquist, 1966; Knope et al., 2020), *Scaevola* (Robichaux and Pearcy, 1984; McKown et al., 2016), *Euphorbia* (Sporck, 2011), and *Plantago* (Dunbar-Co et al., 2009).

Here we focus on variation within an indigenous plant species Sida fallax Walp. (Malvaceae), known in the Hawaiian language as 'ilima. 'Ilima is found from sea level to elevations > 1000 mas on mul-137 tiple Hawaiian islands. Coastal populations are morphologically different from montane populations 138 (Fig. 1). Coastal regions of Hawai'i are characterized by high sun exposure, warmer temperatures, 139 high winds, salinity, and variation in water availability. Coastal populations of 'ilima tend to be short 140 and prostrate which likely helps them to withstand the windy environment (Fig. 1a). The leaves of 141 these populations are covered on both surfaces in dense, soft hairs that give the leaves a silvery green 142 appearance (Fig. 1b), which helps mitigate water loss by reflecting solar radiation, thereby lowering 143 leaf temperature (Ehleringer and Björkman, 1978). Montane regions, on the other hand, provide very 144 different challenges. Many other tall species grow on the slopes of these wet mountainous regions, 145 which makes light competition a factor that plants may need to adapt to. Possibly due to this, montane 146 populations are erect and shrub- or tree-like, capable of growing meters tall with strong, woody stems. 147 These individuals have smooth, green foliage with serrated edges. Montane populations exhibit traits 148 that may help them to compete for light availability. This montane morphology is not found in S. fallax 149 populations on other Pacific Islands (Pejhanmehr et al., 2023). 150

Because of their contrasting habitat and morphology, we treat leaves from coastal and montane plants as representatives of sun and shade leaves, respectively, for testing hypotheses about amphistomy advantage. Specifically, the objectives of our study are to test whether 1) sun leaves of coastal 'ilima plants will have greater AA than shade leaves of montane plants; and if so, is this because 2a) coastal plants have thicker leaves than montane plants and/or 2b) coastal plants have a  $g_{\text{smax,ratio}}$  closer to 0.5?

## 156 MATERIALS AND METHODS —

#### 157 Plant sampling and climate —

We identified 7 suitable natural populations of 'ilima on O'ahu and 5 on Hawai'i Island by consulting 158 Yorkston and Daehler (2006) and citizen scientist records on iNaturalist (Anon, 2022) (Fig. 1c; Table 1). 159 We avoided sites that appeared to be cultivated. We visited sites between August and November 2022. 160 For logistical reasons, the sites on Hawai'i were sampled during one three-day trip. We haphazardly 161 sampled eight plants distributed evenly between the highest and lowest elevation plants along a transect 162 at each site. For safety and conservation reasons, transects were along a trail or road. We did not 163 sample small individuals if there was risk removing leaves would cause mortality. From each plant, we 164 collected two fully expanded leaves for trait measurements. We sampled stomatal traits on all leaves; 165 leaf thickness on one leaf from three randomly selected plants per site; and, due to limited time, a 166

single leaf from a single plant at the middle of each transect for gas exchange measurements. We 167 downloaded climatic data on mean annual temperature, solar radiation, and vegetation height from 168 the Climate and Solar Radiation of Hawai'i databases (Giambelluca et al., 2014) using the latitude 169 and longitude at the middle of each transect. We also downloaded mean annual precipitation from 170 1978-2007 from the Rainfall Atlas of Hawai'i (Giambelluca et al., 2013). The spatial resolution is 171 approximately  $234 \times 250$ m. The temperature data are calibrated from networks of meteorological 172 stations operating in the late 20th century and 21st century; the solar radiation data are calibrated from 173 satellite measurements collected between 2002 and 2009 (Giambelluca et al., 2014). We tested whether 174 climatic variables differed among our coastal and montane populations using Welch's two-sample t-175 176

#### 177 Leaf traits —

#### 178 Stomata —

We estimated the stomatal density and size on ab- and adaxial leaf surfaces from all leaves. For 179 pubescent leaves (usually coastal), we dried and pressed leaves for  $\approx 1$  week (Hill et al., 2014), care-180 fully scraped trichomes off with a razor blade, and rehydrated the leaf. Rehydration restores leaf area to 181 its fresh value (Blonder et al., 2012). For glabrous leaves, we used fresh leaves. We applied clear nail 182 polish to both leaf surfaces of fresh or rehydrated leaves in the middle of the lamina away from major 183 veins. After nail polish dried, we mounted impressions on a microscope slide using transparent tape 184 (Mott and Michaelson, 1991). We digitized a portion of each leaf surface impression using a brightfield 185 microscope (Leica DM2000, Wetzlar, Germany). We counted all stomata and divided by the visible 186 leaf area (0.890 mm<sup>2</sup>) to estimate density and measured guard cell length from five randomly chosen 187 stomata per field using ImageJ (Schneider et al., 2012). 188

#### 189 Leaf thickness —

We cut thin sections using two razor blades taped together. We sectioned the leaf in a petri dish of water, wet-mounted sections onto a slide, and took digital micrographs using a brightfield microscope, as described above. Leaf thickness is measured as the length from upper cuticle to lower cuticle.

#### 193 Gas exchange measurements —

At each site, we selected one representative leaf from one plant near the middle of the transect for gas exchange measurements using a portable infrared gas analyzer (LI-6800PF, LI-COR Biosciences, Lincoln, Nebraska, USA). We estimated the photosynthetic rate (A) and stomatal conductance to water vapor ( $g_{\rm sw}$ ) at saturating light (photosynthetic photon flux density (PPFD) =  $2000~\mu{\rm mol~m^{-2}~s^{-1}}$ ), ambient CO<sub>2</sub> (415 ppm), and  $T_{\rm leaf} = 25.0$ –29.3°C. The midday irradiance in coastal 'ilima typically meets or even exceeds a PPFD of  $2000~\mu{\rm mol~m^{-2}~s^{-1}}$  and previous experiments with sun leaves revealed that

2000  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup> is always at or near saturating irradiance. Even though lower irradiance may be 200 saturating for montane leaves, we used this higher value for all leaves to standardize conditions. 201

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We also estimated 'amphistomy advantage' (AA) sensu Parkhurst (1978), but with modification. For each leaf, we measured the photosynthetic rate of an untreated amphistomatous leaf  $(A_{amphi})$  over a range of  $g_{\rm sw}$  values. We refer to this as an  $A-g_{\rm sw}$  curve, which is described in more detail below. We 204 compared the  $A-g_{sw}$  curve of the untreated leaf to the photosynthetic rate of pseudohypostomatous leaf  $(A_{\text{hypo}})$ , which is the same leaf but with gas exchange through the upper surface blocked by a neutral density plastic (propafilm). Hypostomy refers to leaves with stomata only present on the lower, typically abaxial, surface. We refer to the untreated and partially blocked leaves as "amphi" and "pseudohypo", respectively. AA is calculated as the log-response ratio of A compared at the same total  $g_{sw}$ :

$${\rm AA} = \log(A_{\rm amphi}/A_{\rm hypo})$$

The log-response ratio is commonly used social and biological sciences (e.g. Hedges et al. (1999)). It is straightforward to interpret because values above 0 indicate a photosynthetic advantage of amphistomy, whereas values less than 0 indicate a disadvantage. The log-response ratio is preferable to the absolute difference because it indicates a proportional change in A, which facilitates comparisons across leaves and environments with different baseline photosynthetic rates. The irradiance of the light source in the pseudohypo leaf was higher because the propafilm reduces transmission. To compensate for reduced transmission, we increased incident PPFD for pseudohypo leaves by a factor 1/0.91, the inverse of the measured transmissivity of the propafilm. We also set the stomatal conductance ratio, for purposes of calculating boundary layer conductance, to 0 for pseudohypo leaves following manufacturer directions.

Appendix S1 (see the Supplementary Data with this article), Fig. S1 illustrates our method for collecting  $A-g_{sw}$  curves. We collected two curves per leaf, an amphi (untreated) curve and a pseudohypo (treated) curve. To control for order effects, we alternated between starting with amphi or pseudohypo leaf measurements, though we did not detect an effect of treatment order on AA (results not shown). In the field, we acclimated the focal leaf to saturating light and high relative humidity (RH = 70%), as described above, until A and  $g_{\rm sw}$  reach their maximum. We used these data as our estimates of maximum A and  $g_{\rm sw}$ . After that, we decreased RH to  $\approx 10\%$  to induce rapid stomatal closure without biochemical downregulation. Hence,  $A_{\text{amphi}}$  and  $A_{\text{hypo}}$  were both measured at low chamber humidity after the leaf had acclimated to high humidity. All other environmental conditions in the leaf chamber remained the same. We logged data until  $g_{sw}$  reached its nadir. We then repeated the process of acclimating the leaf to 70% RH and inducing stomatal closure with low RH with the other treatment (amphi or pseudohypo).

#### 231 Data analysis —

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Objective 1: Do coastal leaves have greater amphistomy advantage than montane leaves? —

It is not feasible to record  $A_{\rm amphi}$  and  $A_{\rm hypo}$  at the exact same  $g_{\rm sw}$ . To overcome this, we fit  $A-g_{\rm sw}$  curves using a linear regression of  $\log(g_{\rm sw})$  on A to interpolate modeled A for amphi and pseudohypo leaves at the same  $g_{\rm sw}$ . Let  $\hat{A}_{\rm amphi}$  and  $\hat{A}_{\rm hypo}$  be the estimated A of the amphi and pseudohypo leaves, respectively. We estimated these quantities at the same  $g_{\rm sw}$  using fitted parameters ( $\hat{\beta}$ 's):

$$\begin{split} \hat{A}_{\text{amphi}} &= \hat{\beta}_{0,\text{amphi}} + \hat{\beta}_{1,\text{amphi}} \times \log(g_{\text{sw}}) \\ \hat{A}_{\text{hypo}} &= \hat{\beta}_{0,\text{hypo}} + \hat{\beta}_{1,\text{hypo}} \times \log(g_{\text{sw}}) \end{split}$$

In 10 of 12 leaves, the minimum  $g_{\rm sw}$  of the amphi curve was smaller than the maximum  $g_{\rm sw}$  of the pseudohypo curve (i.e. the curves overlapped for a range of  $g_{\rm sw}$  values). In those cases, we estimated  $\hat{A}_{\rm amphi}$  and  $\hat{A}_{\rm hypo}$  at the  $g_{\rm sw}$  value in the middle of the range of overlap between the curves. In 2 of 12 leaves, the  $A-g_{\rm sw}$  curves did not quite overlap because the minimum  $g_{\rm sw}$  of the amphi curve was slightly greater than the maximum  $g_{\rm sw}$  of the pseudohypo curve. In those cases, we estimated AA by extrapolating slightly,  $1.98\times 10^{-3}$  and  $3.29\times 10^{-3}$  mol m<sup>-2</sup> s<sup>-1</sup>, beyond the measured curves to the  $g_{\rm sw}$  value in between the curves. The vertical lines in Appendix S1, Fig. S2 show the  $g_{\rm sw}$  for each leaf. We estimated AA from  $\hat{A}_{\rm amphi}$  and  $\hat{A}_{\rm hypo}$  for each leaf using the log-response ratio shown above.

To estimate  $\hat{\beta}$ 's from the  $A-g_{\rm sw}$  curve for each leaf, we fit Bayesian regressions using the R package brms version 2.20.4 (Bürkner, 2017) with MCMC sampling in Stan (Stan Development Team, 2023). We used CmdStan version 2.33.1 and cmdstanr version 0.6.1 (Gabry et al., 2023) to interface with R version 4.3.1 (R Core Team, 2023). We sampled the posterior distribution from 4 chains with 1000 iterations each after 1000 warmup iterations per chain. We estimated parameters and confidence intervals as the median and 95% quantile intervals of the posterior, respectively. The key prediction is that  $AA_{\rm coastal} > AA_{\rm montane}$ , meaning the 95% confidence intervals of  $AA_{\rm coastal} = AA_{\rm montane}$  should be positive and not encompass 0.

254 Objective 2a: Are coastal leaves thicker than montane leaves? —

We tested whether leaf thickness (log-transformed) varied between coastal and montane populations and among individuals within populations using a Bayesian mixed-effects model with habitat as a fixed effect and individual plant and site as random effects. We used the *R* package **brms** version 2.20.4 (Bürkner, 2017) to fit the model in *Stan* (Stan Development Team, 2023) with CmdStan version 2.33.1 and **cmdstanr** version 0.6.1 (Gabry et al., 2023). We sampled the posterior distribution from 4 chains with 1000 iterations each after 1000 warmup iterations per chain. We estimated the relationship between population average leaf thickness and AA measured from a single individual per population. We used this approach because most of the variation in leaf thickness was among sites and the plant

selected for gas exchange measurements was not always among the plants randomly selected for leaf thickness, precluding individual level correlation. We propagated uncertainty about in AA and leaf thickness estimates by integrating over the entire posterior distribution sample for each variable. The key prediction is that the effect of leaf thickness on AA is positive, meaning the 95% confidence interval of the slope should be positive and not encompass 0.

Objective 2b: Is  $g_{\text{smax, ratio}}$  closer to 0.5 in coastal leaves than montane leaves? —

We tested whether  $g_{\text{smax,ratio}}$  varied between coastal and montane populations and among individuals 269 within populations using a Bayesian mutliresponse, mixed-effects model. The modeled response vari-270 ables are stomatal count and guard cell length on each surface. Counts were modeled as negative 271 binomially distributed variable from a latent stomatal density and a parameter  $\phi$  to estimate overdis-272 persion in counts relative to a Poisson model. For all traits, the explanatory variables were habitat as a fixed effect and leaf within individual plant, individual plant, and site as random effects. We used 274 the R package **brms** version 2.20.4 (Bürkner, 2017) to fit the model in *Stan* (Stan Development Team, 275 2023) with CmdStan version 2.33.1 and **cmdstanr** version 0.6.1 (Gabry et al., 2023). We interpolated 276 missing adaxial guard cell lengths from 6 out of 185 samples with zero adaxial stomata using the "mi" 277 function in **brms** package. We sampled the posterior distribution from 4 chains with 1000 iterations 278 each after 1000 warmup iterations per chain. From each posterior sample, we calculated  $g_{\text{smax,ratio}}$  as: 279

$$g_{\rm smax,ratio} = \frac{g_{\rm smax,upper}}{g_{\rm smax,lower} + g_{\rm smax,upper}}, \label{eq:gsmax}$$

where  $g_{\rm smax,lower}$  and  $g_{\rm smax,upper}$  are maximum stomatal conductance to water vapor at  $T_{\rm leaf}=25^{\circ}$  C on the lower and upper surface, respectively. The maximum stomatal conductance was calculated from stomatal density and length, assuming that stomata are fully open, following Sack and Buckley (2016):

$$g_{\rm smax} = bmds^{0.5}.$$

In this equation, b is a biophysical constant, m is a morphological constant, d is the stomatal density, and s is the stomatal complex area. We assume that b, which is determined by the molecular species, temperature, and air pressure, is the same for both surfaces; we assume that m, which is determined by guard cell allometry is also the same for both surfaces. Hence, the b and m constants cancel out of  $g_{\rm smax,ratio}$  and only density and length (l), which is proportional to the square root of area, affect the ratio:  $g_{\rm smax} \propto dl$ .

We estimated the relationship between leaf  $g_{\rm smax,ratio}$  and AA measured from a single leaf per population. We propagated uncertainty about AA and  $g_{\rm smax,ratio}$  by integrating over the entire posterior distribution sample for each variable. The key prediction is that the effect of  $g_{\rm smax,ratio}$  on AA is positive until  $g_{\rm smax,ratio} < 0.5$ , meaning the 95% confidence interval of the slope should be positive and not encompass 0 in the domain  $g_{\rm smax,ratio} < 0.5$ .

## 5 RESULTS —

Coastal 'ilima are surrounded by shorter vegetation than their montane counterparts (Fig. 1d; Welch Two Sample *t*-test,  $t_{6.67}=5.13$ , P=0.002). The montane site with the lowest vegetation height is a remnant dry forest (Koai'a tree sanctuary) in a matrix of cattle pasture, hence the satellite derived vegetation height may be lower than what existed prior to human disturbance. Coastal sites receive greater average solar radiation at the top of the canopy (Fig. 1d; Welch Two Sample *t*-test,  $t_{10.86}=-2.22$ , P=0.049); coastal sites are significantly warmer (Fig. 1d; Welch Two Sample *t*-test,  $t_{6.01}=-2.96$ , P=0.025); and coastal sites receive less precipitation (Fig. 1d; Welch Two Sample *t*-test,  $t_{7.45}=2.73$ ,  $t_{7.45}=2.73$ ,  $t_{7.45}=0.028$ ).

#### 304 Amphistomy advantage is greater in coastal leaves —

Amphistomy increases photosynthesis in leaves of coastal 'ilima plants more than that of montane plants. 305 AA was significantly greater than 0 (95% confidence intervals did not overlap 0) in 5 of 6 coastal leaves, 306 but only 1 of 6 montane leaves (Fig. 2; see Appendix S1, Fig. S2 for individual curves). Overall, the av-307 erage AA among coastal and montane leaves is 0.12 [0.077 - 0.15] and 0.027 [-0.0034 - 0.057], respec-308 tively; the difference in average AA between habitat types is  $AA_{coastal} - AA_{montane} = 0.09 [0.039 - 0.14]$ . 309 Posterior predictions closely match observed values of A (Appendix S1, Fig. S3), indicating an ade-310 quate model fit from which we can interpolate between measurements reliably. It also suggests that 311 slight extrapolation beyond the data should be reliable, but this is less certain. When we remove 312 two leaves where we extrapolated slightly beyond fitted  $A-g_{\rm sw}$  curves, we estimate that  $AA_{\rm coastal}$  is still positive, 0.081 [0.023-0.13], but the difference between coastal and montane leaves is smaller, 0.053 [-0.012-0.12], and confidence intervals slightly overlap 0. Maximum photosynthetic rate was 315 slightly, but not significantly higher in coastal leaves (Welch Two Sample t-test,  $t_{9.65} = 1.6$ , P =316 0.14); total stomatal conductance was similar (Welch Two Sample *t*-test,  $t_{9.71} = -0.09$ , P = 0.93) 317 in coastal and montane leaves (Appendix S1, Fig. S4). Water-use efficiency ( $A/g_{\rm sw}$ ) was significantly 318 higher in coastal leaves (Welch Two Sample *t*-test,  $t_{9.99} = 2.54$ , P = 0.03). 319

#### Leaf thickness is associated with amphistomy advantage between but not within habitats —

Coastal 'ilima leaves are  $91 [26-164] \mu m$  thicker than their montane counterparts. Although coastal leaves are thicker and have greater AA, there is little relationship between leaf thickness and AA within habitats (Fig. 3A; slope = -0.11 [-0.28-0.035]).

#### $g_{ m smax,ratio}$ is not associated with amphistomy advantage —

Coastal and montane leaves have similar average  $g_{\rm smax,ratio}$ , the ratio of adaxial (upper) to total  $g_{\rm smax}$ , the anatomical maximum stomatal conductance to water vapor (Appendix S1, Fig. S5); coastal leaves have 0.059~[-0.14-0.28] higher  $g_{\rm smax,ratio}$  than montane leaves, but the 95% confidence intervals overlap 0

difference. The  $g_{\rm smax,ratio}$  is somewhat bimodal among sites. Some sites in both habitats have leaves with  $g_{\rm smax,ratio} < 0.07$  and others with  $g_{\rm smax,ratio} > 0.2$  (Appendix S1, Fig. S5). This is particularly noticeable in montane sites where those on the Big Island of Hawai'i all have low  $g_{\rm smax,ratio}$  whereas those on O'ahu have relatively high  $g_{\rm smax,ratio}$ . There is no relationship between  $g_{\rm smax,ratio}$  and AA in either habitat (Fig. 3B; slope = 0.14 [-0.057–0.34]) in our sample.

# 3 DISCUSSION —

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Amphistomy is a seemingly simple way that leaves can increase carbon gain without significant additional water loss, yet it is rare in nature and we do not know why. The strong association between amphiostomy and sunny, open habitats suggests that amphistomy may benefit sun leaves more than shade leaves, but progress has been limited by the lack of evidence that amphiostomy per se improves photosynthesis in a given leaf. By experimentally blocking gas exchange through the upper surface in a controlled environment, we directly compared an amphistomatous leaf to an otherwise identical pseudohypostomatous leaf. This allows us to quantity the amphistomy advantage (AA) holding all else constant. Taking advantage of the steep climatic gradients in the Hawaiian archipelago, we applied this new method to show for the first time that sun leaves benefit 4.04 times more from amphistomy than shade leaves on 'ilima ( $Sida\ fallax$ ) plants ( $AA_{coastal}=0.12\ vs.\ AA_{montane}=0.027$ ). Coastal and montane 'ilima leaves are likely good representatives of classic sun and shade leaf syndromes because 1) they vary in traits like reflective pubescence (Ehleringer and Björkman, 1978) and leaf thickness (Terashima et al., 2001) that typically characterize sun-shade adaptation; and 2) since 'ilima shrubs are typically < 1m tall, they are shaded by trees in montane, but not coastal habitats (Fig. 1d). While this result has not yet been validated in other species, our results indicate that part of the reason amphistomatous leaves are found most commonly in high light habitats is that the adaptive benefit is greater in such environments.

If AA is typically greater in sun leaves than shade leaves, it could partially explain the distribution of amphi- and hypostoamtous leaves, but the precise mechanism(s) require further study. One hypothesis is that the internal airspace conductance,  $g_{ias}$ , from stomata to mesophyll cell walls is lower in thicker sun leaves (Parkhurst, 1978). All else being equal, a leaf with lower  $g_{ias}$  will benefit more from amphistomy. Our results partially support this hypothesis. Coastal 'ilima leaves with high AA (Fig. 2) are thicker than montane leaves, but the relationship between AA and leaf thickness within habitats is actually slightly negative (Fig. 3a), opposite our prediction. Since coastal and montane leaves differ in many respects besides thickness, we do not have enough data to conclude that leaf thickness explains the variation in AA between habitats. Alternatively, other biochemical or anatomical differences between coastal and montane leaves may explain why AA is greater in coastal leaves. The negative relationship, albeit nonsignificant in that 95% confidence intervals encompassed 0, between leaf thickness and AA could be explained if thicker leaves compensated by having a more porous mesophyll and/or less tortuous airspaces (Théroux-Rancourt et al., 2021).

A second natural hypothesis is that amphistomatous leaves with few adaxial (upper) stomata benefit less than those with similar densities on both surfaces. We predicted that leaves with  $g_{\text{smax.ratio}}$  closer to

0.5 would have higher AA based on biophysical models (Gutschick, 1984a). The logic is that a small number of stomata on the upper surface are insufficient to supply the entire upper mesophyll due to limited lateral diffusion (Morison et al., 2005). Our results do not support this hypothesis. Montane leaves from Big Island sites had low  $g_{\mathrm{smax,ratio}}$  and low AA whereas low montane leaves on Oʻahu had high  $g_{\text{smax,ratio}}$ , but similarly low AA (Fig. 3b). Among coastal sites, the site with the lowest  $g_{\text{smax,ratio}}$ had the lowest AA, but there was little variation in  $g_{\text{smax,ratio}}$  among coastal leaves in our sample. We therefore cannot rule out that a larger sample of coastal leaves with greater variance in  $g_{\text{smax,ratio}}$  might support this hypothesis. 

Two major implications from our study are that 1) photosynthesis in hypostomatous leaves is likely limited by CO<sub>2</sub> concentration drawdown within leaf airspaces; and 2) amphistomy *per se* contributes to, but is not wholly responsible for, higher photosynthetic rates among amphistomatous leaves. The amphistomy advantage we observe in 'ilima leaves implies decreased CO<sub>2</sub> supply in pseudohypostomatous leaves because of concentration drawdowns in the leaf airspace. Limited diffusion through the airspace has long been hypothesized to depress photosynthesis in hypostomatous leaves (Parkhurst, 1994), with empirical support from helox studies (Parkhurst and Mott, 1990). However, these studies relied on interspecific comparisons of amphi- and hypostomatous leaves that differ systematically in many traits that affect gas exchange and photosynthesis (Xiong and Flexas, 2020). Our experimental approach overcomes this limitation and implies that the drop in CO<sub>2</sub> concentration from substomatal cavities to the upper surface depresses photosynthesis.

Among land plants grown in a common garden, amphistomatous leaves have on average nearly  $2 \times 1000$  higher area-based photosynthetic rates (Xiong and Flexas, 2020), naively implying an AA  $\approx 1000$  log 2 = 0.69. This is much higher than our estimate of 0.12 among coastal 'ilima leaves. The most likely explanation is that amphistomy is not the only cause of high photosynthetic rate. Indeed, species adapted to open, high light habitats with amphistomatous leaves also have higher concentrations of Rubisco, overall stomatal conductance, and photosynthetic capacity (Smith et al., 1997; Xiong and Flexas, 2020). For a leaf with high photosynthetic capacity that is well illuminated and hydrated, the major limitation becomes  $CO_2$ . Under these conditions, amphistomy may substantially increase photosynthesis, as we observe in coastal 'ilima leaves. Selection on increased photosynthesis under similar conditions may explain why crop leaves tend to increase stomatal density ratio during domestication (Milla et al., 2013).

Three limitations of this study are the small sample size, experimental design that precludes distinguishing genetic from environmental differences in leaf traits, and potentially confounding effects of other environmental differences besides light environment. Understanding the mechanistic basis of higher AA in sun leaves would require much larger sample sizes. Sun leaves tend to be thicker, more densely packed with mesophyll cells, and have greater photosynthetic capacity and higher stomatal conductance, among other traits (Lambers et al., 2008). Each of these factors and others potentially modulate AA. Quantifying the contribution of all these factors requires larger samples and additional measurements that are beyond the scope of this study, but exciting avenues for future research on leaf structure-function relations. Although many morphological traits that distinguish coastal and montane 'ilima populations persist in a common environment (Yorkston and Daehler, 2006), we cannot distinguish between genetic effects and plastic responses to habitat as causes of difference in AA because

we measured naturally occurring plants in situ. While disentangling genetic and plastic contributions 407 is not necessarily important for understanding the distribution of amphistomatous leaves, it would be 408 insightful to know about genetic and environmental contributions to trait variation. A reciprocal trans-409 plant would be able to determine the genetic and environmental contributions, as well their interaction, 410 to trait variance in nature. However, reciprocal transplants cannot control for other differences between 411 coastal and montane habitats besides vegetation height, such as temperature and precipitation. Experi-412 mental studies in controlled environments will be necessary to isolate the effects of light quantity and 413 quality on AA. 414

#### 15 CONCLUSIONS —

This study reports the first direct experimental evidence that having stomata open on both leaf surfaces, 416 amphistomy, increases photosynthesis for a given total stomatal conductance, particularly in leaves 417 from the type of open, sunny habitats where this trait is most common. By developing a straightfor-418 ward experimental method to block gas exchange through the upper surface, we directly compared the 419 photosynthetic rate of a leaf with gas exchange through both surfaces or just one, holding all other fac-420 tors constant. In doing so, we found that coastal leaves of the indigenous Hawaiian 'ilima (Sida fallax) enjoyed a greater photosynthetic benefit from amphistomy than nearby montane leaves living in more 422 closed forest. This is not because amphistomatous leaves necessarily have greater leaf surface available 423 for stomata, although that likely influences realized photosynthetic rates in natural populations. Rather, 424 our experiments show that coastal amphistomatous leaves with the same total leaf stomatal conduc-425 tance photosynthesize more than identical hypostomatous leaves. We cannot yet ascribe the difference 426 in amphistomy advantage between coastal and montane leaves to particular physiological or anatomical 427 variation, but this is a promising area for future research. 428

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#### 435 Author Contributions —

GT and CDM contributed equally to all stages of this project; TNB contributed to development of the method and helped edit the manuscript.

# Data Availability Statement —

- Custom scripts are available on a GitHub repository (https://github.com/cdmuir/stomata-ilima) and archived on Zenodo (https://doi.org/10.5281/zenodo.10369114). Raw data are deposited on Dryad (https://doi.org/10.5061/dryad.rxwdbrvfw).
- <sup>442</sup> Supporting Information —

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- Additional supporting information may be found online in the Supporting Information section at the end of the article.
  - Appendix S1: Supplemental figures and table

#### 46 Literature Cited —

- 447 Anon. 2022. iNaturalist.
- Ball, J. T., I. E. Woodrow, and J. A. Berry. 1987. A Model Predicting Stomatal Conductance and its Contribution to the Control of Photosynthesis under Different Environmental Conditions. *In J.*Biggins [ed.], Progress in Photosynthesis Research, 221–224. Springer Netherlands, Dordrecht.
- Berry, J. A., D. J. Beerling, and P. J. Franks. 2010. Stomata: Key players in the earth system, past and present. *Current Opinion in Plant Biology* 13: 232–239.
- Blonder, B., V. Buzzard, I. Simova, L. Sloat, B. Boyle, R. Lipson, B. Aguilar-Beaucage, et al. 2012.

  The leaf-area shrinkage effect can bias paleoclimate and ecology research. *American Journal of Botany* 99: 1756–1763.
- Boer, H. J. de, P. L. Drake, E. Wendt, C. A. Price, E.-D. Schulze, N. C. Turner, D. Nicolle, and E. J.
   Veneklaas. 2016. Apparent overinvestment in leaf venation relaxes leaf morphological constraints on photosynthesis in arid habitats. *Plant Physiology* 172: 2286–2299.
- Bucher, S. F., K. Auerswald, C. Grün-Wenzel, S. I. Higgins, J. Garcia Jorge, and C. Römermann. 2017.
   Stomatal traits relate to habitat preferences of herbaceous species in a temperate climate. *Flora* 229: 107–115.
- Bürkner, P.-C. 2017. **Brms**: An *r* Package for Bayesian Multilevel Models Using *stan*. *Journal of Statistical Software* 80.
- Carlquist, S. 1966. The Biota of Long-Distance Dispersal. II. Loss of Dispersibility in Pacific Compositae. *Evolution* 20: 30.
- Collatz, G. J., J. T. Ball, C. Grivet, and J. A. Berry. 1991. Physiological and environmental regulation of
   stomatal conductance, photosynthesis and transpiration: A model that includes a laminar boundary
   layer. Agricultural and Forest Meteorology 54: 107–136.
- Cowan, I. R., and G. D. Farquhar. 1977. Stomatal function in relation to leaf metabolism and environment. *Symposia of the Society for Experimental Biology* 31: 471–505.
- Deans, R. M., T. J. Brodribb, F. A. Busch, and G. D. Farquhar. 2020. Optimization can provide the fundamental link between leaf photosynthesis, gas exchange and water relations. *Nature Plants* 6: 1116–1125.
- Doll, Y., H. Koga, and H. Tsukaya. 2021. *Callitriche* as a potential model system for evolutionary studies on the dorsiventral distribution of stomata. *Plant Signaling & Behavior* 16: 1978201.
- Drake, P. L., H. J. de Boer, S. J. Schymanski, and E. J. Veneklaas. 2019. Two sides to every leaf: Water and CO<sub>2</sub> transport in hypostomatous and amphistomatous leaves. *New Phytologist* 222: 1179–1187.
- Drake, P. L., R. H. Froend, and P. J. Franks. 2013. Smaller, faster stomata: Scaling of stomatal size, rate of response, and stomatal conductance. *Journal of Experimental Botany* 64: 495–505.
- Dunbar-Co, S., M. J. Sporck, and L. Sack. 2009. Leaf trait diversification and design in seven rare taxa of the Hawaiian *p*lantago radiation. *International Journal of Plant Sciences* 170: 61–75.
- Earles, J. M., G. Theroux-Rancourt, A. B. Roddy, M. E. Gilbert, A. J. McElrone, and C. R. Brodersen.

  2018. Beyond porosity: 3D leaf intercellular airspace traits that impact mesophyll conductance.

  Plant Physiology 178: 148–162.
- Ehleringer, J. R., and O. Björkman. 1978. Pubescence and leaf spectral characteristics in a desert shrub, Encelia farinosa. *Oecologia* 36: 151–162.

- Evans, J. R., R. Kaldenhoff, B. Genty, and I. Terashima. 2009. Resistances along the CO<sub>2</sub> diffusion pathway inside leaves. *Journal of Experimental Botany* 60: 2235–2248.
- Flexas, J., M. J. Clemente-Moreno, J. Bota, T. J. Brodribb, J. Gago, Y. Mizokami, M. Nadal, et al. 2021.
   Cell wall thickness and composition are involved in photosynthetic limitation P. Manavella [ed.],
   Journal of Experimental Botany 72: 3971–3986.
- Foster, J. R., and W. K. Smith. 1986. Influence of stomatal distribution on transpiration in low-wind environments. *Plant, Cell & Environment* 9: 751–759.
- Franklin, O., S. P. Harrison, R. Dewar, C. E. Farrior, Å. Brännström, U. Dieckmann, S. Pietsch, et al. 2020. Organizing principles for vegetation dynamics. *Nature Plants* 6: 444–453.
- Franks, P. J., G. B. Bonan, J. A. Berry, D. L. Lombardozzi, N. M. Holbrook, N. Herold, and K. W. Oleson. 2018. Comparing optimal and empirical stomatal conductance models for application in Earth system models. *Global Change Biology* 24: 5708–5723.
- Gabry, J., R. Češnovar, and A. Johnson. 2023. Cmdstanr: R Interface to 'CmdStan'.
- Gago, J., D. M. Daloso, M. Carriquí, M. Nadal, M. Morales, W. L. Araújo, A. Nunes-Nesi, et al. 2020.
   The photosynthesis game is in the "inter-play": Mechanisms underlying CO2 diffusion in leaves.
   Environmental and Experimental Botany 178: 104174.
- Gay, A. P., and R. G. Hurd. 1975. The influence of light on stomatal density in the tomato. *New Phytologist* 75: 37–46.
- Giambelluca, T. W., Q. Chen, A. G. Frazier, J. P. Price, Y.-L. Chen, P.-S. Chu, J. K. Eischeid, and D. M.
   Delparte. 2013. Online Rainfall Atlas of Hawai'i. Bulletin of the American Meteorological Society
   94: 313–316.
- Giambelluca, T. W., X. Shuai, M. L. Barnes, R. J. Alliss, R. J. Longman, T. Miura, Q. Chen, et al. 2014.
   Evapotranspiration of Hawai'i.
- Gibson, A. C. 1996. Structure-Function Relations of Warm Desert Plants. Springer Berlin / Heidelberg,
   Berlin, Heidelberg.
- Givnish, T. J. ed. 1986. On the economy of plant form and function. Cambridge University Press, Cambridge.
- Givnish, T. J., K. C. Millam, A. R. Mast, T. B. Paterson, T. J. Theim, A. L. Hipp, J. M. Henss, et al.
   2009. Origin, adaptive radiation and diversification of the Hawaiian lobeliads (Asterales: Campanulaceae). Proceedings of the Royal Society B: Biological Sciences 276: 407–416.
- Givnish, T. J., and R. A. Montgomery. 2014. Common-garden studies on adaptive radiation of photosynthetic physiology among Hawaiian lobeliads. *Proceedings of the Royal Society B: Biological Sciences* 281: 20132944–20132944.
- Givnish, T. J., R. A. Montgomery, and G. Goldstein. 2004. Adaptive radiation of photosynthetic physiology in the Hawaiian lobeliads: Light regimes, static light responses, and whole-plant compensation points. *American Journal of Botany* 91: 228–246.
- Grubb, P. J. 2020. Leaf structure and function. *In A. Dobson*, D. Tilman, and R. D. Holt [eds.], Unsolved
   Problems in Ecology, 124–144. Princeton University Press, Princeton.
- Grubb, P. J. 1977. Leaf structure and function. *In* R. Duncan, and M. Weston-Smith [eds.], The encyclopedia of ignorance, 317–330. Pergamon, Oxford.
- Gutschick, V. P. 1984a. Photosynthesis model for C<sub>3</sub> leaves incorporating CO<sub>2</sub> transport, propagation of radiation, and biochemistry 1. Kinetics and their parameterization. *Photosynthetica* 18: 549–568.

- Gutschick, V. P. 1984b. Photosynthesis model for C<sub>3</sub> leaves incorporating CO<sub>2</sub> transport, propagation of radiation, and biochemistry 2. Ecological and agricultural utility. *Photosynthetica* 18: 569–595.
- Harrison, S. P., W. Cramer, O. Franklin, I. C. Prentice, H. Wang, Å. Brännström, H. Boer, et al. 2021.
   Eco-evolutionary optimality as a means to improve vegetation and land-surface models. New Phytologist 231: 2125–2141.
- Harwood, R., G. Théroux-Rancourt, and M. M. Barbour. 2021. Understanding airspace in leaves: 3D anatomy and directional tortuosity. *Plant, Cell & Environment* 44: 2455–2465.
- Hedges, L. V., J. Gurevitch, and P. S. Curtis. 1999. The meta-analysis of response ratios in experimental ecology. *Ecology* 80: 1150–1156.
- Hetherington, A. M., and F. I. Woodward. 2003. The role of stomata in sensing and driving environmental change. *Nature* 424: 901–908.
- Hill, K. E., G. R. Guerin, R. S. Hill, and J. R. Watling. 2014. Temperature influences stomatal density
   and maximum potential water loss through stomata of Dodonaea viscosa subsp. Angustissima along
   a latitude gradient in southern Australia. Australian Journal of Botany 62: 657.
- Ho, Q. T., H. N. C. Berghuijs, R. Watté, P. Verboven, E. Herremans, X. Yin, M. A. Retta, et al. 2016.
   Three-dimensional microscale modelling of CO<sub>2</sub> transport and light propagation in tomato leaves enlightens photosynthesis: 3-D modelling of photosynthesis in leaves. *Plant, Cell & Environment* 39: 50–61.
- James, S. A., and D. T. Bell. 2001. Leaf morphological and anatomical characteristics of heteroblastic Eucalyptus globulus ssp. Globulus (Myrtaceae). *Australian Journal of Botany* 49: 259.
- Jarvis, P. G. 1976. The interpretation of the variations in leaf water potential and stomatal conductance found in canopies in the field. *Philosophical Transactions of the Royal Society of London. B,* Biological Sciences 273: 593–610.
- Jordan, G. J., R. J. Carpenter, and T. J. Brodribb. 2014. Using fossil leaves as evidence for open vegetation. *Palaeogeography, Palaeoclimatology, Palaeoecology* 395: 168–175.
- Kaul, R. B. 1976. Anatomical observations on floating leaves. *Aquatic Botany* 2: 215–234.
- Knope, M. L., V. A. Funk, M. A. Johnson, W. L. Wagner, E. M. Datlof, G. Johnson, D. J. Crawford, et al. 2020. Dispersal and adaptive radiation of *bidens* (Compositae) across the remote archipelagoes of Polynesia. *Journal of Systematics and Evolution* 58: 805–822.
- Lambers, H., F. S. Chapin, and T. L. Pons. 2008. Plant Physiological Ecology. Springer New York, New York, NY.
- Leuning, R. 1995. A critical appraisal of a combined stomatal-photosynthesis model for C3 plants.

  Plant, Cell and Environment 18: 339–355.
- Lin, Y.-S., B. E. Medlyn, R. A. Duursma, I. C. Prentice, H. Wang, S. Baig, D. Eamus, et al. 2015.

  Optimal stomatal behaviour around the world. *Nature Climate Change* 5: 459–464.
- Lyshede, O. B. 2002. Comparative and functional leaf anatomy of selected Alstroemeriaceae of mainly Chilean origin. *Botanical Journal of the Linnean Society* 140: 261–272.
- Márquez, D. A., H. Stuart-Williams, L. A. Cernusak, and G. D. Farquhar. 2023. Assessing the CO<sub>2</sub> concentration at the surface of photosynthetic mesophyll cells. *New Phytologist* 238: 1446–1460.
- Márquez, D. A., H. Stuart-Williams, G. D. Farquhar, and F. A. Busch. 2022. Cuticular conductance
   of adaxial and abaxial leaf surfaces and its relation to minimum leaf surface conductance. New
   Phytologist 233: 156–168.
- McKown, A. D., M. E. Akamine, and L. Sack. 2016. Trait convergence and diversification arising from

- a complex evolutionary history in Hawaiian species of Scaevola. *Oecologia* 181: 1083–1100.
- Medlyn, B. E., R. A. Duursma, D. Eamus, D. S. Ellsworth, I. C. Prentice, C. V. M. Barton, K. Y. Crous,
   et al. 2011. Reconciling the optimal and empirical approaches to modelling stomatal conductance.
   Global Change Biology 17: 2134–2144.
- 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.
- Montgomery, R. A., and T. J. Givnish. 2008. Adaptive radiation of photosynthetic physiology in the Hawaiian lobeliads: Dynamic photosynthetic responses. *Oecologia* 155: 455–467.
- Morison, J. I. L., E. Gallouët, T. Lawson, G. Cornic, R. Herbin, and N. R. Baker. 2005. Lateral diffusion of CO<sub>2</sub> in leaves is not sufficient to support photosynthesis. *Plant Physiology* 139: 254–266.
- Mott, K. A., A. C. Gibson, and J. W. O'Leary. 1982. The adaptive significance of amphistomatic leaves.

  \*\*Plant, Cell & Environment 5: 455–460.
- Mott, K. A., and O. Michaelson. 1991. Amphistomy as an adaptation to high light intensity in *Ambrosia* cordifolia (Compositae). *American Journal of Botany* 78: 76–79.
- Muir, C. D. 2019. Is amphistomy an adaptation to high light? Optimality models of stomatal traits along light gradients. *Integrative and Comparative Biology* 59: 571–584.
- Muir, C. D. 2018. Light and growth form interact to shape stomatal ratio among British angiosperms.

  New Phytologist 218: 242–252.
- Muir, C. D. 2015. Making pore choices: Repeated regime shifts in stomatal ratio. Proceedings of the
   Royal Society B: Biological Sciences 282: 20151498.
- Nobel, P. S. 2020. Physicochemical and environmental plant physiology. Fifth edition. Academic Press, an imprint of Elsevier, London, United Kingdom; San Diego, CA.
- Parkhurst, D. F. 1994. Diffusion of CO<sub>2</sub> and other gases inside leaves. *New Phytologist* 126: 449–479.
- Parkhurst, D. F. 1978. The adaptive significance of stomatal occurrence on one or both surfaces of leaves. *The Journal of Ecology* 66: 367.
- Parkhurst, D. F., and K. A. Mott. 1990. Intercellular diffusion limits to CO<sub>2</sub> uptake in leaves: Studies in air and helox. *Plant Physiology* 94: 1024–1032.
- Peat, H. J., and A. H. 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.
- Pejhanmehr, M., M. Yorkston, and C. W. Morden. 2023. Low Genetic Diversity in the Highly Morphologically Diverse Sida fallax Walp. (Malvaceae) Throughout the Pacific 1. *Pacific Science* 76.
- Poorter, H., Ü. Niinemets, N. Ntagkas, A. Siebenkäs, M. Mäenpää, S. Matsubara, and T. L. Pons. 2019.

  A meta□analysis of plant responses to light intensity for 70 traits ranging from molecules to whole plant performance. *New Phytologist*: nph.15754.
- R Core Team. 2023. R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing, Vienna, Austria.
- Robichaux, R. H., and R. W. Pearcy. 1984. Evolution of C<sub>3</sub> and C<sub>4</sub> plants along an environmental moisture gradient: Patterns of photosynthetic differentiation in Hawaiian <i>Scaevola<\i>and <i>Euphorbia<\i>species. American Journal of Botany 71: 121–129.
- Sack, L., and T. N. Buckley. 2016. The developmental basis of stomatal density and flux. *Plant Physiology* 171: 2358–2363.
- 616 Salisbury, E. J. 1928. I. 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, Containing Papers of a Biological Character 216: 1–65.
- Santrůček, J., L. Schreiber, J. Macková, M. Vráblová, J. Květoň, P. Macek, and J. Neuwirthová. 2019.
  Partitioning of mesophyll conductance for CO<sub>2</sub> into intercellular and cellular components using carbon isotope composition of cuticles from opposite leaf sides. *Photosynthesis Research*: 1–19.
- Schneider, C. A., W. S. Rasband, and K. W. Eliceiri. 2012. NIH Image to ImageJ: 25 years of image analysis. *Nature Methods* 9: 671–675.
- Scoffoni, C., J. Kunkle, J. Pasquet-Kok, C. Vuong, A. J. Patel, R. A. Montgomery, T. J. Givnish, and
   L. Sack. 2015. Light-induced plasticity in leaf hydraulics, venation, anatomy, and gas exchange in
   ecologically diverse Hawaiian lobeliads. New Phytologist 207: 43–58.
- Sellers, P. J., R. E. Dickinson, D. A. Randall, A. K. Betts, F. G. Hall, J. A. Berry, G. J. Collatz, et al.
   1997. Modeling the exchanges of energy, water, and carbon between continents and the atmosphere.
   Science 275: 502–509.
- 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.
- Sporck, M. J. 2011. The Hawaiian C<sub>4</sub> *euphorbia* adaptive radiation: An ecophysiological approach to understanding leaf trait variation. Ph. {D}. University of Hawaii.
- 634 Stan Development Team. 2023. Stan Modeling Language Users Guide and Reference Manual.
- Terashima, I., Y. T. Hanba, Y. Tazoe, P. Vyas, and S. Yano. 2006. Irradiance and phenotype: Comparative eco-development of sun and shade leaves in relation to photosynthetic CO<sub>2</sub> diffusion. *Journal of Experimental Botany* 57: 343–354.
- Terashima, I., S.-I. Miyazawa, and Y. T. Hanba. 2001. Why are sun leaves thicker than shade leaves?
   Consideration based on analyses of CO<sub>2</sub> diffusion in the leaf. *Journal of Plant Research* 114:
   93–105.
- Théroux-Rancourt, G., A. B. Roddy, J. M. Earles, M. E. Gilbert, M. A. Zwieniecki, C. K. Boyce, D.
  Tholen, et al. 2021. Maximum CO<sub>2</sub> diffusion inside leaves is limited by the scaling of cell size and
  genome size. *Proceedings of the Royal Society B: Biological Sciences* 288: 20203145.
- Tholen, D., and X.-G. Zhu. 2011. The mechanistic basis of internal conductance: A theoretical analysis of mesophyll cell photosynthesis and CO<sub>2</sub> diffusion. *Plant Physiology* 156: 90–105.
- Tomás, M., J. Flexas, L. Copolovici, J. Galmés, L. Hallik, H. Medrano, M. Ribas-Carbó, et al. 2013.
   Importance of leaf anatomy in determining mesophyll diffusion conductance to CO2 across species:
   Quantitative limitations and scaling up by models. *Journal of Experimental Botany* 64: 2269–2281.
- Wang, H., I. C. Prentice, T. F. Keenan, T. W. Davis, I. J. Wright, W. K. Cornwell, B. J. Evans, and C. Peng. 2017. Towards a universal model for carbon dioxide uptake by plants. *Nature Plants* 3: 734–741.
- Wang, Y., J. S. Sperry, W. R. L. Anderegg, M. D. Venturas, and A. T. Trugman. 2020. A theoretical and empirical assessment of stomatal optimization modeling. *New Phytologist* 227: 311–325.
- Wood, J. G. 1934. The physiology of xerophytism in Australian plants: The stomatal frequencies,
   transpiration and osmotic pressures of sclerophyll and tomentose-succulent leaved plants. *Journal* of Ecology 22: 69–87.
- Xiong, D., and J. Flexas. 2020. From one side to two sides: The effects of stomatal distribution on photosynthesis. *New Phytologist* 228: 1754–1766.

Yorkston, M., and C. C. Daehler. 2006. Interfertility between Hawaiian ecotypes of *Sida fallax* (Mal-vaceae) and evidence of a hybrid disadvantage. *International Journal of Plant Sciences* 167: 221–230.

Table 1: 'Ilima study site location information.

Site	Island	Habitat	Latitude	Longitude	Elevation (mas)
Kahuku Point	Oʻahu	coastal	21.710	-157.982	4
Kaloko beach	Oʻahu	coastal	21.293	-157.661	4
Kaloko-Honokōhau national historical park	Hawaiʻi	coastal	19.676	-156.024	6
Ka'ena Point	Oʻahu	coastal	21.574	-158.278	4
Makapu'u beach	Oʻahu	coastal	21.313	-157.661	3
Puakō petroglyph park	Hawaiʻi	coastal	19.957	-155.858	8
Hawai'i loa ridge	Oʻahu	montane	21.294	-157.727	352
Hāloa 'Āina	Hawaiʻi	montane	19.552	-155.793	1567
Ka'ohe game management area	Hawaiʻi	montane	19.817	-155.616	1946
Koai'a tree sanctuary	Hawaiʻi	montane	20.048	-155.737	970
Mau'umae Ridge	Oʻahu	montane	21.305	-157.779	248
Waʻahila ridge	Oʻahu	montane	21.314	-157.793	357

# 663 Figure Legends —

Figure 1: A. Typical growth form of montane (left) and coastal (right) 'ilima plants and B. leaves. C. Map of the sites that were sampled on the islands of O'ahu and Hawai'i (aka Big Island). D. Climatic, light, and vegetation height comparisons between montane (blue) and coastal (orange) habitats sampled in this study. Open circles are values for the midpoint of each site transect; closed circles and intervals are the mean  $\pm 1$  standard error. The habitats differ significantly in mean annual precipitation (top-left), solar radiation (bottom-left), temperature (top-right), and vegetation height (bottom-right). MAP = mean annual precipitation; MAT = mean annual temperature; ns = not significant; \* indicates  $0.01 \le P < 0.05$ ; \*\* indicates  $0.01 \le P < 0.01$ .

Figure 2: Coastal leaves benefit more from amphistomy than montane leaves. A positive amphistomy advantage (y-axis) means that the photosynthetic rate of an amphistomatous leaf is greater than that of an identical pseudohypostomatous leaf at the same overall  $g_{\rm sw}$ . Each point-interval is the median posterior estimate plus 95% confidence interval of amphistomy advantage for that leaf. Each leaf is from a different montane (blue) or coastal (orange) site, arranged by habitat and ascending amphistomy advantage within habitat. The longer horizonal bars are the average amphistomy advantage for montane and coastal leaves.  $g_{\rm sw}$ , stomatal conductance to water vapor.

Figure 3: Relationships between leaf amphistomy advantage, (A) leaf thickness and (B)  $g_{\rm smax,ratio}$  among 'ilima ( $Sida\,fallax$ ) plants from montane (blue) and coastal (orange) habitats in Hawai'i. A positive amphistomy advantage (y-axis) means that the photosynthetic rate of an amphistomatous leaf is greater than that of an identical pseudohypostomatous leaf at the same overall  $g_{\rm sw}$ . Each point-interval is the median posterior estimate plus 95% confidence interval of the trait value. Each leaf is from a different montane (blue) or coastal (orange) site. Lines are the estimated linear regression of (A) log(leaf thickness) and (B)  $g_{\rm smax,ratio}$  on amphistomy advantage; ribbons are the 95% confident bands of the regression. Symbols:  $g_{\rm smax,ratio}$ , anatomical maximum stomatal conductance ratio;  $g_{\rm sw}$ , stomatal conductance to water vapor.