

Intraspecific variability and reaction norms of forest understorey plant species traits

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Summary

1. Trait-based models of ecological communities typically assume intraspecific variation in functional traits is not important, although such variation can change species trait rankings along gradients in resources and environmental conditions, and thus influence community structure and function.
2. We examined the degree of intraspecific relative to interspecific variation, and reaction norms of 11 functional traits for 57 forest understorey plant species, including: intrinsic water-use efficiency (iWUE), $\Delta^{15}\text{N}$, five leaf traits, two stem traits and two root traits along gradients in light, nitrogen, moisture and understorey cover.
3. Our results indicate that interspecific trait variation exceeded intraspecific variation by at least 50% for most, but not all traits. Intraspecific variation in $\Delta^{15}\text{N}$, iWUE, leaf nitrogen content (LNC) and root traits was high (47–70%) compared with most leaf traits and stem traits (13–38%).
4. $\Delta^{15}\text{N}$ varied primarily along gradients in abiotic conditions, while light and understorey cover were relatively less important. Intrinsic water-use efficiency was related primarily to light transmission, reflecting increases in photosynthesis relative to stomatal conductance. Leaf traits varied mainly as a function of light availability, with some reaction norms depending on understorey cover. Plant height increased with understorey cover, while stem-specific density was related primarily to light. Resources, environmental conditions and understorey cover did not contribute strongly to the observed variation in root traits.
5. Gradients in resources, environmental conditions and competition all appear to control intraspecific variability in most traits to some extent. However, our results suggest that species cross-over (i.e. trait rank reversals) along the gradients measured here are generally not a concern.
6. Intraspecific variability in understorey plant species traits can be considerable. However, trait data collected under a narrow range of environmental conditions appears sufficient to establish species rankings and scale between community and ecosystem levels using trait-based models. Investigators may therefore focus on obtaining a sufficient sample size within a single set of conditions rather than characterizing trait variation across entire gradients to optimize sampling efforts.

Key-words: functional traits, herbaceous layer, Pacific Northwest, stable isotopes, water-use efficiency

Introduction

Trait-based models of ecological communities are used increasingly in community ecology because they promise

greater generality, predictive power, and ability to scale between community and ecosystem levels of organization (McGill *et al.* 2006; Suding *et al.* 2008). Large collaborative databases composed of average species trait values now facilitate the large-scale adoption of trait-based approaches (e.g. Wright *et al.* 2004; Kattge *et al.* 2011).

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However, the use of species averages may discount the importance of intraspecific variation in community assembly processes, species coexistence and associated ecosystem functions (Bolnick *et al.* 2011; Laughlin *et al.* 2012; Violle *et al.* 2012; Hart, Schreiber & Levine 2016). Empirical studies are therefore necessary to evaluate the degree of intraspecific variation in traits, and to determine whether accounting for intraspecific variation may improve or modify trait-based models of species assemblages (Siefert *et al.* 2015; Shipley *et al.* 2016).

The mass ratio theory posits that the relative contribution of species in a community is proportional to its contribution to primary production, and that ecosystem processes are determined by the traits of dominant plant species (Grime 1998). Mass ratio theory provides the basis for using community-aggregated traits to model responses of plant communities to variation in the environment and effects of community composition on ecosystem processes and/or services. Species average trait values are commonly weighted by their relative abundances and summed to calculate community-aggregated traits (Pérez-Harguindeguy *et al.* 2013). This is considered sound when interspecific variability exceeds intraspecific variability, or when species rankings are maintained across gradients in resources and environmental conditions (Fig. S1a in Supporting Information; Garnier *et al.* 2001; Kazakou *et al.* 2014). Conversely, trait reaction norms (i.e. responses of traits to gradients in resources and environmental conditions) may result in species cross-over, here defined as shifts in rankings of species traits along gradients in resources or competition (*sensu* Givnish, Montgomery & Goldstein 2004; Fig. S1b). Species cross-over could result in different community-aggregated trait values along those gradients for the same community and/or differences in species composition for a given community-aggregated trait value – requiring researchers to account for intraspecific variation when calculating community-aggregated traits. There is evidence for cross-over in the physiological performance among coexisting and/or closely related species across environmental gradients (Chazdon 1992; Kaelke, Kruger & Reich 2001; Givnish, Montgomery & Goldstein 2004). Traits are considered proxies for physiological performance. However, despite the increasing use of trait-based approaches, the assumption that species trait rankings are constant across gradients in resources, environmental conditions and competition has been evaluated sparingly (Garnier *et al.* 2001; Albert *et al.* 2010; Lepš *et al.* 2011; Auger & Shipley 2013; Kazakou *et al.* 2014). More intensive sampling within species to account for intraspecific trait variability and cross-over may improve trait-based models of plant communities, but likely comes at a cost to the number of species that can be sampled when resources are limited (Paine, Baraloto & Díaz 2015).

Trait-based approaches are rapidly being adopted to study effects of disturbance, forest management, climate change and interactions thereof on understorey plant communities and associated ecosystem services (Neill &

Puettmann 2013; Kern *et al.* 2014; Sabatini *et al.* 2014; Sonnier *et al.* 2014). Forest understorey plant communities in the temperate zone typically contain 2–20+ times the number of species as the overstorey (Gilliam 2007). Understorey plant species are sensitive indicators of resources and environmental conditions (Daubenmire 1976), and may be partitioned along gradients in soil properties including moisture and nutrients, light transmission as it relates to overstorey tree structure, and climate (Ares, Berryman & Puettmann 2009; Burton *et al.* 2011, 2014). Similarly, understorey species physiological performance, reflected in morphological and physiological traits, is likely to vary along these gradients (McGill *et al.* 2006). Recent meta-analyses show that intraspecific variation in whole-plant traits is greater than biochemical traits, which exceeds intraspecific variation in morphological traits (Siefert *et al.* 2015). To date, much research has focused on adaptation and acclimation of leaf traits and associated physiological processes to shade (e.g. Givnish 1988; Chazdon 1992; Ellsworth & Reich 1992, 1993; Kaelke, Kruger & Reich 2001; Givnish, Montgomery & Goldstein 2004), yet little empirical work exists for temperate understorey species. Intraspecific variation in leaf traits along gradients in soil properties, and climatic conditions, and effects of interactions among these gradients are even less well understood (e.g. Roche, Díaz-Burlinson & Gachet 2004; Nicotra *et al.* 2010; Funk *et al.* 2016). Moreover, little is known about how whole-plant (e.g. water-use efficiency, nitrogen use strategy), stem [e.g. plant height, stem-specific density (SSD)], and root traits [e.g. specific root length (SRL), rooting depth] vary along these gradients within species.

Our goal was to investigate the assumptions underlying the common practice of using species means in trait-based modelling of plant communities. We examined intraspecific relative to interspecific trait variability. Additionally, we assessed alternative models of trait reaction norms and cross-over along specific gradients in light, soil nitrogen, understorey cover and climatic conditions using hierarchical mixed models. All models include random effects accounting for the nested structure of the sampling design. We do not control for the effects of local adaptation or genetic variation, which may vary along environmental and resource gradients with traits (e.g. Ravenscroft, Fridley & Grime 2014). We evaluated the hypotheses that trait variation among species exceeds variation within species, and species maintain rankings along environmental and resource gradients. Finally, we interpreted trait reaction norms considering expected physiological responses (Table 1). We assessed a suite of leaf, stem and root traits for plant species found in the understorey of Douglas-fir forests in western Oregon. We also examined whole-plant traits including intrinsic water-use efficiency (iWUE) based on stable carbon ($\delta^{13}\text{C}$) isotopes (Brooks *et al.* 1997; Foster & Brooks 2005), and nitrogen (N) stable isotope discrimination relative to soil ($\Delta^{15}\text{N}$) – a metric of niche partitioning in nitrogen use strategies among plants (Nadelhoffer *et al.* 1996; Gubsch *et al.* 2011).

Table 1. Leaf, stem and root traits examined and their hypothesized reaction norms (positive or negative) along gradient in resources and environmental conditions relevant to understorey plant species

Trait	Gradients in resources and environmental conditions			
	GLI	Understorey cover	Soil nitrogen	CMD
Whole-plant traits				
$\Delta^{15}\text{N}$	+	–	+	–
iWUE	+	–	+	–
Leaf traits				
SLA	–	+	+	–
LNC	–	+	+	–
N_{area}	+	–	+	–
Leaf size	–	+	+	–
LDMC	+	–	–	+
Stem traits				
Plant height	+	+	+	–
SSD	–	+	–	+
Root traits				
SRL	+	–	+	–
Rooting depth	–	+	–	+

GLI, gap light index; CMD, climatic moisture deficit; $\Delta^{15}\text{N}$, nitrogen stable isotope discrimination; iWUE, intrinsic water-use efficiency; SLA, specific leaf area; LNC, leaf nitrogen content; N_{area} , leaf nitrogen per unit area; LDMC, leaf dry matter content; SSD, stem-specific density; SRL, specific root length.

Materials and methods

STUDY AREA

We collected functional trait data at seven sites located in western Oregon Coast Range and western Oregon Cascades, USA. These sites are the locations of a replicated manipulative experiment known as the Density Management Study. For more detailed information about experiment, history, soils and climate of the study sites, see Cissel *et al.* (2006). The sites are distributed across the western hemlock zone (Franklin & Dyrness 1988) covering a broad geographic (sites range between 10 and 245 km apart) and climatic gradient (across sites average 2001–2010 mean annual temperatures range from 8.6 to 11.7 °C, mean annual precipitation ranges from 1274 to 2080 mm; Wang *et al.* 2012). The climate is Mediterranean with mild, wet winters and warm, dry summers (Cissel *et al.* 2006). Soils are well- to poorly drained (highly weathered) Ultisols and (younger, less structured) Inceptisols, and vary widely among sites in nitrogen (N) availability (Thiel & Perakis 2009). Forests were thinned ~60- to 80-year-old Douglas-fir (*Pseudotsuga menziesii*) stands with varying abundances of western hemlock on some sites. Other conifer species, such as western redcedar (*Thuja plicata*), and hardwood species including bigleaf maple (*Acer macrophyllum*), red alder (*Alnus rubra*), Pacific dogwood (*Cornus nuttallii*), Pacific madrone (*Arbutus menziesii*) and golden chinquapin (*Chrysolepis chrysophylla*) were minor components of the overstorey.

The Density Management Study uses a randomized complete block design with one replicate of four density treatments at each of seven 94–131 ha sites (Cissel *et al.* 2006). This experimental structure ensured a broad gradient of overstorey structures and associated resources and environmental conditions for the understorey plants (Appendix S1). We used overstorey and understorey data collected from permanent plots to select dominant

understorey species and characterize local overstorey conditions (Appendix S1).

TRAIT DATA

We focused on a suite of eleven whole-plant, leaf, stem and root traits commonly used to infer ecological strategies of plants (Table 1). Foliar stable isotopes (δ) for C and N provide information about ecological strategies at the whole-plant level. Higher values of foliar C isotope ($\delta^{13}\text{C}$) generally indicate higher intrinsic water-use efficiency of plants (iWUE, A/g_s), which may be sensitive to variation in light, soil moisture and microclimatic conditions in the forest understorey (Farquhar, Ehleringer & Hubick 1989b; Farquhar *et al.* 1989a; Ehleringer 1991). Foliar N isotopes ($\delta^{15}\text{N}$) are influenced by many aspects of the environmental physiology of plant N uptake including N form(s) used (i.e. inorganic; NO_3^- vs. NH_4^+ vs. organic N), timing and depth of N uptake as well as mycorrhizal influence and within-plant N partitioning. These many factors, their potential interactions and their environmental dependence complicate attempts to resolve specific cause(s) of high vs. low plant $\delta^{15}\text{N}$ values in natural settings (Evans 2001). However, in general wider variation in $\delta^{15}\text{N}$ values broadly reflects greater diversity in plant N use strategies (Nadelhoffer *et al.* 1996; Gubsch *et al.* 2011). Leaf traits, including specific leaf area (SLA, $\text{mm}^2 \text{mg}^{-1}$), LNC (mg g^{-1}), leaf nitrogen per area (N_{area} , g m^{-2}), leaf size (cm^2) and leaf dry matter content (LDMC, mg g^{-1}), indicate major leaf economic trade-offs between high rates of resource acquisition and resource conservation (Diaz *et al.* 2004; Wright *et al.* 2004; Pierce *et al.* 2017). We also measured stem traits, including vegetative height, as an indicator of a potential trade-off between height growth and photosynthetic and conductive tissues maintaining water transport (Givnish 1982, 1995), and SSD (mg mm^{-3}), as an indicator of a potential trade-off between growth, and strength and decay resistance (Chave *et al.* 2009). Rooting depth (cm) can affect resource acquisition and persistence (e.g. Antos & Halpern 1997), while SRL (m g^{-1}), defined as the ratio of a standard unit of resource acquisition (root length) to the resource investment (mass), is positively related to rates of nutrient and water uptake and relative growth rate, but negatively related to root life span (Eissenstat 1991). Plants are therefore expected to exhibit variation in these traits in response to variation in resources and environmental conditions (Table 1).

Field data collection

We measured traits of all understorey plant species comprising $\leq 80\%$ of the cumulative importance (the average of the relative frequency and relative abundance, measured here as per cent cover) at each site following standard protocols, with modifications for SRL (Pérez-Harguindeguy *et al.* 2013; see Appendix S1 for details). Plant samples were collected in 2015, 3–5 years following a second experimental treatment. At each permanent vegetation survey plot, individual plants located closest to plot centre were sampled provided they were not severely suppressed by other understorey vegetation and did not exhibit signs of damage due to, e.g. herbivory or diseases. Plant height and rooting depth were measured *in situ* (Appendix S1). Once all *in situ* morphological measurements were recorded, we collected each specimen for processing in the laboratory (Appendix S1). Plant samples were stored in a 3 °C dark cold storage room until processed in the laboratory.

Calculating plant intrinsic water-use efficiency and leaf $\Delta^{15}\text{N}$

Soil $\delta^{15}\text{N}$ values in forests vary widely in response to local topography, N fixation and disturbance history, which can contribute to

variation in plant tissue $\delta^{15}\text{N}$ among sites (Perakis, Sinkhorn & Compton 2011; Perakis, Tepley & Compton 2015). To control for background soil variation and enable comparisons of $\delta^{15}\text{N}$ as a trait-based measure of plant N use, we calculated $\Delta^{15}\text{N}$ as $\delta^{15}\text{N}_{\text{soil}}$ (measured in each plot) subtracted from leaf $\delta^{15}\text{N}$ prior to statistical analysis (e.g. Gubsch *et al.* 2011).

$$\Delta^{15}\text{N} = \delta^{15}\text{N}_{\text{leaf}} - \delta^{15}\text{N}_{\text{soil}} \quad \text{eqn 1}$$

From foliar $\delta^{13}\text{C}$ values, we calculated iWUE (Farquhar, Ehleringer & Hubick 1989b; Farquhar *et al.* 1989a; Ehleringer 1991). The $\delta^{13}\text{C}$ values in plant leaves are influenced by the variation in the atmospheric CO_2 isotopic composition ($\delta^{13}\text{C}_{\text{air}}$), and by biochemical and biochemical processes in the plant. To calculate iWUE, we first calculated carbon isotope discrimination ($\Delta^{13}\text{C}$) to account for variation due to $\delta^{13}\text{C}_{\text{air}}$ (Farquhar, O'Leary & Berry 1982):

$$\Delta^{13}\text{C} = \frac{\delta^{13}\text{C}_{\text{air}} - \delta^{13}\text{C}_{\text{leaf}}}{1 + \delta^{13}\text{C}_{\text{leaf}}/1000} \quad \text{eqn 2}$$

We estimated $\delta^{13}\text{C}_{\text{air}}$ for each individual sample using the model of Buchmann, Brooks & Ehleringer (2002) because $\delta^{13}\text{C}_{\text{air}}$ varies vertically within forest understories as a result of respired CO_2 and low wind speeds:

$$\delta^{13}\text{C}_{\text{air}} = \delta^{13}\text{C}_{\text{trop}} - \frac{0.023 \times L}{h} \quad \text{eqn 3}$$

where L = leaf area index (LAI), h = height and $\delta^{13}\text{C}_{\text{trop}}$ is $\delta^{13}\text{C}$ of the troposphere, well above the influence of the canopy. We estimated LAI from hemispherical photos taken at plot centres (described below). Field measurements of plant height were used for h . We estimated $\delta^{13}\text{C}_{\text{trop}}$ during the 2015 growing season to be -8.55‰ using a simple linear regression of growing season $\delta^{13}\text{C}_{\text{trop}}$ on year using Mauna Loa data from 1990 to 2014 ($r^2 = 0.98$).

We then used $\Delta^{13}\text{C}$ values to estimate iWUE [i.e. photosynthesis (A), divided by stomatal conductance (g_s)] using the relationship between $\Delta^{13}\text{C}$ and the ratio of internal CO_2 to atmospheric CO_2 (c_i/c_a) described by Farquhar *et al.* (1989a):

$$\Delta^{13}\text{C} = a + (b - a) \left(\frac{c_i}{c_a} \right) \quad \text{eqn 4}$$

where a = diffusion (4.4‰), b = RuBisCO $\sim 29\text{‰}$, c_i and c_a are internal and ambient CO_2 , respectively. Intrinsic water-use efficiency can then be estimated from c_i and c_a as follows:

$$\text{iWUE} = \frac{A}{g} = \frac{c_a - c_i}{1.6} \quad \text{eqn 5}$$

where A is the rate of photosynthesis ($\mu\text{mol m}^{-2} \text{s}^{-1}$), g is stomatal conductance ($\text{mmol m}^{-2} \text{s}^{-1}$), 1.6 is the ratio of diffusivities of water and CO_2 in air and c_a is predicted to be 400 p.p.m. using a simple linear regression of growing season CO_2 on year using Mauna Loa data from 1969 to 2014 ($r^2 = 0.998$).

RESOURCES AND ENVIRONMENTAL CONDITIONS

To characterize local light transmission and associated environmental conditions (impacts on humidity and temperature) in the understory, we took hemispherical photographs using a Nikon Coolpix 5000 digital camera (Melville, NY, USA) and FC-E8 fish-eye lens adapter. Photos were taken at plot centres after mounting and levelling the camera on a tripod at 1 m height under variable weather conditions (i.e. sunny as well as cloudy skies). Hemispherical photos were analysed for LAI and gap light index (GLI) using Hemisphere v. 2.16, © Patrick Schleppei, WSL. In addition, we estimated total vascular plant cover (per cent) within a 2.5 m

radius of the specimen as a proxy measure for understory competition intensity (Wagner & Radosevich 1998). We collected one mineral soil sample in the centre of each plot using a 5.8 cm diameter corer to a depth of 13 cm, sieved samples through a 2 mm sieve, and ground and analysed samples for total C and N, $\delta^{13}\text{C}$, and $\delta^{15}\text{N}$ as described above. To understand the integrated effects of temperature and precipitation on plant traits, we used annual climatic moisture deficit (CMD) calculated as the sum of the monthly differences between potential evapotranspiration and precipitation (Wang *et al.* 2012). Annual CMD data (2001–2010) were obtained for all plots based on geographic coordinates and elevation from downscaled spatial interpolations of monthly data, accounting for effects of local topography, coastal influences and temperature inversions using ClimateWNA (Wang *et al.* 2012).

DATA ANALYSIS

We analysed the trait data using linear mixed effects models to account for the hierarchical sampling design. First, we analysed intercept-only models of traits with random effects to quantify the (i) variation associated with differences among species ($\sigma_{\text{species}}^2$) and (ii) variability within species across sites (sites nested in species, σ_{sites}^2). Residuals therefore represent intraspecific variability within sites (σ_{resid}^2). To assess whether intraspecific variability was on average lower than interspecific variability, we examined the ratio of the variance associated with interspecific variation ($\sigma_{\text{species}}^2$) to the total variance associated with intraspecific variation ($\sigma_{\text{sites}}^2 + \sigma_{\text{resid}}^2$).

Then we analysed a series of alternative models composed of fixed and random effects. Fixed effects accounted for the effects of resources and environmental conditions that vary over progressively broader spatial scales on observed trait values, including understory cover, GLI, soil nitrogen (N) and moisture deficits. We treated species as a fixed effect to be able to sort out the roles of resources and environmental conditions after accounting for differences among species, and to examine the importance of interactions between species and resource/environmental variables potentially resulting in cross-over. In this context, the random effects structure accounts for the hierarchical experimental design (i.e. site, treatment nested in site and plot nested within treatment and site were modeled as random intercepts).

For each trait, we used a multi-step modelling process to compare a sequence of alternative models comprised of progressively more variables (Table S1). In step one, we fit null, intercept-only models consisting of random effects only. In step two, we added the effect of species. In step three we added variables describing effects of fine-scale variation in overstorey and understory vegetation structure on resources (e.g. light, soil moisture, available nitrogen) indexed by GLI and understory cover, and plausible interactions. In step four, we considered intermediate-scale variation in soil nitrogen (total) and plausible interactions with variables selected in steps two and three (i.e. species, GLI and understory cover) were assessed. In step five, we assessed alternative models integrating the best model selected in previous steps with additional effects of broad-scale variation in CMD and plausible interactions. We tested for species cross-over along all environmental/resource gradients by comparing alternative models with and without interactions between species and the gradient variable of interest. For each step, we evaluated alternative models consisting of various plausible combinations of variables and appropriate two-way interactions using AIC_c, a bias-corrected version of the Akaike Information Criterion (AIC) for small sample sizes (Burnham & Anderson 2002). Three-way interactions were considered when the categorical effect for species was one of the three terms. The best model was selected and carried forward to the next step; if top-ranking models did not differ substantially ($\Delta\text{AIC}_c < 2$) we used the model with the fewest parameters. We

parameterized the best model from all steps, and plotted marginal predictions of species traits along selected gradients to interpret reaction norms in light of hypotheses in Table 1.

For each alternative model, we estimated the variance explained by the marginal fixed effects alone (R_m^2), and by the fixed effects conditioned on random effects (R_c^2) (Nakagawa & Schielzeth 2013). Changes in R_c^2 and R_m^2 following the addition of fixed effects reflected the importance of that effect to explaining variation in traits. In addition, to compare the relative importance of variables, we calculated semi-partial R_c^2 following Edwards *et al.* (2008) for all fixed effects in the selected models. Semi-partial R^2 values (R_β^2) measured the marginal contributions of predictor variables conditioned on other predictor variables in the models. All analysis was done with SAS version 9.4, using the mixed procedure (SAS Institute, © 2002-2012, Cary, NC, USA).

Results

INTER- AND INTRASPECIFIC TRAIT VARIABILITY

All whole-plant, leaf, stem and root traits exhibited considerable levels of inter- and intraspecific variability (Fig. 1). Intraspecific variation was highest for whole-plant measures (the ratio of interspecific-to-intraspecific variation for $\Delta^{15}\text{N} = 0.38$ and $\text{iWUE} = 0.47$), followed by root traits (interspecific : intraspecific for specific root length, $\text{SRL} = 0.89$, rooting depth = 0.50). With the exception of mass-based leaf nitrogen content, LNC, (interspecific : intraspecific = 0.79), intraspecific variation was lowest for leaf traits (interspecific : intraspecific for SLA = 2.67, nitrogen per unit area, $\text{N}_{\text{area}} = 1.58$, LDMC = 3.06 and leaf size = 6.53) and stem traits (interspecific : intraspecific for height = 3.06, $\text{SSD} = 1.94$). These results are largely consistent with those of Siefert *et al.* (2015).

TRAIT REACTION NORMS

Whole-plant traits

The selected model indicated that $\Delta^{15}\text{N}$ was related to differences among species (semi-partial $R_\beta^2 = 0.37$), understorey cover ($R_\beta^2 = 0.02$), soil N ($R_\beta^2 = 0.15$) and CMD ($R_\beta^2 = 0.30$) but not GLI. Specifically, plant $\Delta^{15}\text{N}$ (i.e. $\delta^{15}\text{N}_{\text{plant}}$ normalized to $\delta^{15}\text{N}_{\text{soil}}$) decreased with

understorey cover and increased with soil N and CMD (Fig. 2; Table 2). These fixed effects explained 35% of the variation (R_m^2), with an additional 12% explained by random effects ($R_c^2 = 0.47$).

In contrast, the best supported model for iWUE showed that in addition to being related to differences among species ($R_\beta^2 = 0.54$), iWUE increased with GLI ($R_\beta^2 = 0.36$), and decreased with understorey cover ($R_\beta^2 = 0.02$) and soil N ($R_\beta^2 = 0.05$; Fig. 3). Fixed effects accounted for the majority of variation explained for iWUE ($R_m^2 = 0.49$, $R_c^2 = 0.54$; Table 2).

Leaf traits

Variation in leaf traits was related primarily to differences among species (R_β^2 ranges 0.56–0.88), GLI and understorey cover (Fig. 4, Table 2). Specific leaf area ($R_\beta^2 = 0.11$), LNC ($R_\beta^2 = 0.06$) and leaf size ($R_\beta^2 = 0.27$) decreased, whereas LDMC ($R_\beta^2 = 0.12$) and N_{area} ($R_\beta^2 = 0.29$) increased with GLI. Relationships of these traits to understorey cover were relatively weak and in the opposite direction, as for GLI (Fig. S2). The model for SLA ($\Delta\text{AIC}_c = 0.5$) and LDMC also included an interaction between understorey cover and GLI ($R_\beta^2 = 0.01$ and 0.04, respectively). In addition, N_{area} was negatively related to soil N ($R_\beta^2 = 0.06$; Fig. S3). Fixed effects explained between 53 and 87% of the variation in leaf traits (R_m^2), with random effects explaining a relatively small amount (1–6%, R_c^2 , Table 2).

Stem traits

Variation in stem traits was related to variation among species, and in understorey cover and GLI. Plant height varied among species ($R_\beta^2 = 0.77$) and increased with understorey cover ($R_\beta^2 = 0.19$). Stem-specific density varied among species ($R_\beta^2 = 0.69$), decreased with understorey cover ($R_\beta^2 = 0.01$) and increased with GLI ($R_\beta^2 = 0.11$; Fig. 5).

Root traits

Root traits (i.e. SRL and root depth) varied only among species. These models explained 49 and 43% of the

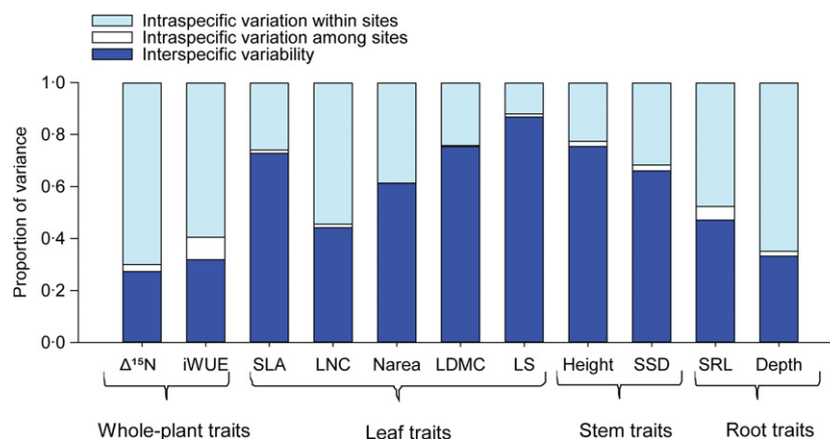


Fig. 1. Sources of variation in whole-plant, leaf, stem and root traits. Relative variance decomposition at the species and within-species levels. Intraspecific variation was partitioned among and within sites. $\Delta^{15}\text{N}$, nitrogen stable isotope discrimination; iWUE , intrinsic water-use efficiency; SLA, specific leaf area; LNC, leaf nitrogen content; N_{area} , leaf nitrogen per unit area; LDMC, leaf dry matter content; LS, leaf size; SSD, stem-specific density; SRL, specific root length.

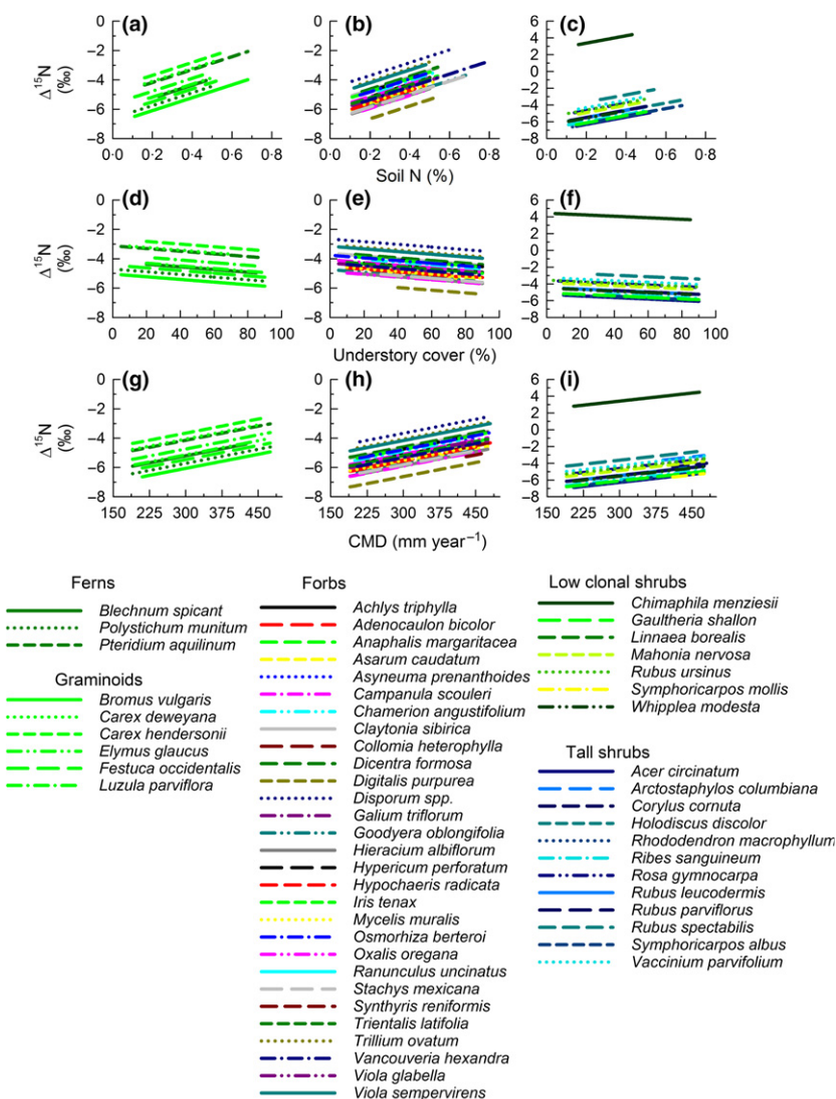


Fig. 2. Relationships of foliar stable isotope discrimination ($\Delta^{15}\text{N}$) to soil N (a–c), understory cover (d–f) and climatic moisture deficit (g–i). Ferns and graminoids are shown in the first column (a, d, g), forbs in the second (b, e, h) and shrubs in the third (c, f, i). Lines show model estimates from selected model (Table 2) for individual species plotted across the range of conditions in which they were sampled. Note differences in scale range for shrubs (c, f, i). $\Delta^{15}\text{N}$, nitrogen stable isotope discrimination; CMD, climatic moisture deficit.

variation, with random effects accounting for an added 4 and 1%, respectively (Table 2).

Variation explained by fixed effects

Fixed effects explained between 35% ($\Delta^{15}\text{N}$) and 88% (leaf size) of the variability in all traits (R_m^2 in Table 2). Additional effects, including species - gradient interactions indicative of species cross-over, did not substantially increase the variation explained (Tables 2 and S3). Apart from the models for iWUE and $\Delta^{15}\text{N}$, in which additional terms led to increases of 10 and 9%, none of the variables added in steps three through five did much to increase the variance explained beyond that explained by species alone (Table S3). Moreover, small differences between R_m^2 and R^2 conditioned on random effects (R_c^2) suggest that only a small proportion of variation traits was due to unexplained variation among sites, experimental treatments and plots (Tables 2 and S3).

Finally, our analysis of semi-partial coefficients (R_β^2) showed that after accounting for other fixed effects in the model, the majority of variation (i.e. $R_\beta^2 = 0.37\text{--}0.88$) in all traits is related to differences among species (Fig. 6). Following variation associated with species, variation in leaf traits is most strongly associated with GLI (SLA = 0.11, LNC = 0.06, $N_{\text{area}} = 0.29$, leaf size = 0.27 and LDMC = 0.12). Understorey cover was the only additional fixed effect in the model for plant height (explaining 19% of the variation after accounting for species). The model for SSD also included GLI [$R_\beta^2 = 0.11$ accounting for species ($R_\beta^2 = 0.69$) and understory cover ($R_\beta^2 = 1\%$)]. Resource and environmental variables were relatively more important in explaining variation in isotope-derived whole-plant traits. For iWUE, the order of importance of resources and environmental variables was GLI ($R_\beta^2 = 0.36$) > soil N ($R_\beta^2 = 0.05$) > understory cover ($R_\beta^2 = 0.02$), whereas the order for $\Delta^{15}\text{N}$ was CMD ($R_\beta^2 = 0.23$) > soil N ($R_\beta^2 = 0.14$) > understory cover ($R_\beta^2 = 0.2$).

Table 2. Comparisons of ΔAIC_c , AIC weights (w) and variability explained by fixed effects (R_m^2) and fixed and random effects ($k =$ number of parameters) combined (R_c^2) among three top models selected using multistep modelling processes (Table S1)

Model by dependent variable	k	ΔAIC_c	w	R_m^2	R_c^2
Whole-plant traits					
$\Delta^{15}N$					
Species – Cover_U + N_{soil} + CMD	63	0.00	0.85	0.35	0.47
Species – Cover _U + N _{soil} + CMD – CMD × N _{soil} + CMD × Cover _U	65	3.74	0.13	0.35	0.47
Species + Cover _U + N _{soil} – N _{soil} × Cover _U	64	8.59	0.01	0.32	0.44
iWUE					
Species – Cover _U + GLI – N _{soil} + N _{soil} × Cover _U	64	0.00	0.25	0.49	0.54
Species – Cover _U + GLI – N _{soil}	63	0.01	0.24	0.49	0.54
Species – Cover_U + GLI	62	0.96	0.15	0.49	0.53
Leaf traits					
Log (specific leaf area)					
Species – Cover _U + GLI + Cover _U × GLI + N _{soil}	64	0.00	0.36	0.80	0.81
Species – Cover_U – GLI + Cover_U × GLI	63	1.14	0.20	0.80	0.80
Species + Cover _U – GLI + Cover _U × GLI + N _{soil} + N _{soil} × GLI – N _{soil} × Cover _U	66	1.72	0.15	0.80	0.80
Log (leaf nitrogen content)					
Species – Cover_U – GLI + Cover_U × GLI	64	0.00	0.37	0.49	0.55
Species – Cover _U – GLI + Cover _U × GLI – CMD	65	2.21	0.12	0.53	0.57
Species – Cover _U – GLI + Cover _U × GLI + N _{soil}	65	2.30	0.12	0.55	0.59
Log (leaf nitrogen per unit area)					
Species + GLI – N_{soil}	63	0.00	0.37	0.66	0.66
Species + GLI – N _{soil} – CMD	64	0.90	0.23	0.66	0.66
Species + GLI + N _{soil} – GLI × N _{soil}	64	2.10	0.13	0.66	0.66
Log (leaf size)					
Species + Cover_U – GLI	62	0.00	0.25	0.87	0.88
Species + Cover _U – GLI + Cover _U × GLI	63	0.00	0.25	0.88	0.88
Species + Cover _U – GLI + N _{soil}	63	0.30	0.22	0.88	0.88
Log (LMDC)					
Species + Cover_U + GLI – Cover_U × GLI	64	0.00	0.41	0.80	0.81
Species + Cover _U + GLI – Cover _U × GLI – CMD	65	0.32	0.35	0.80	0.81
Species + Cover _U + GLI – Cover _U × GLI – N _{soil}	65	1.57	0.19	0.80	0.81
Stem traits					
Log (height)					
Species + Cover _U + N _{soil}	62	0.00	0.21	0.78	0.78
Species + Cover _U – GLI	63	0.12	0.20	0.78	0.78
Species + Cover_U	62	0.18	0.19	0.78	0.78
Log (stem-specific density)					
Species + Cover_U + GLI – Cover_U × GLI	51	0.00	0.40	0.71	0.71
Species + Cover _U + GLI – Cover _U × GLI – CMD	52	2.24	0.13	0.71	0.71
Species – Cover _U + GLI	50	2.46	0.12	0.70	0.71
Root traits					
Log (specific root length)					
Species	61	0.00	0.36	0.49	0.53
Species + Cover _U	62	2.32	0.11	0.49	0.53
Species – GLI	62	2.33	0.11	0.49	0.53
Log (Root depth)					
Species	51	0.72	0.17	0.41	0.42
Species – Cover _U	52	0.53	0.19	0.41	0.42
Species – Cover _U + GLI	53	0.00	0.25	0.41	0.42

The selected model is shown in bold text. Results from all model comparisons are provided in Table S3. Cover_U, understorey cover; N_{soil}, soil nitrogen. See Table 1 for others.

Discussion

Trait-based models of ecological communities are appealing because they promise greater generality, predictive power and ability to scale between community and ecosystem levels of organization (McGill *et al.* 2006; Suding *et al.* 2008). These approaches commonly ignore trait variation within species, an assumption that has rarely been

tested (Shipley *et al.* 2016; Siefert *et al.* 2015). Our results highlight the importance of understanding the inference (sampling) scope when using single trait values to represent species in a plant community. Although interspecific variability exceeded intraspecific variability for most traits, levels of intraspecific variability were considerable. Within species, most traits varied significantly along multiple gradients (i.e. in light, understorey cover and/or soil N), but

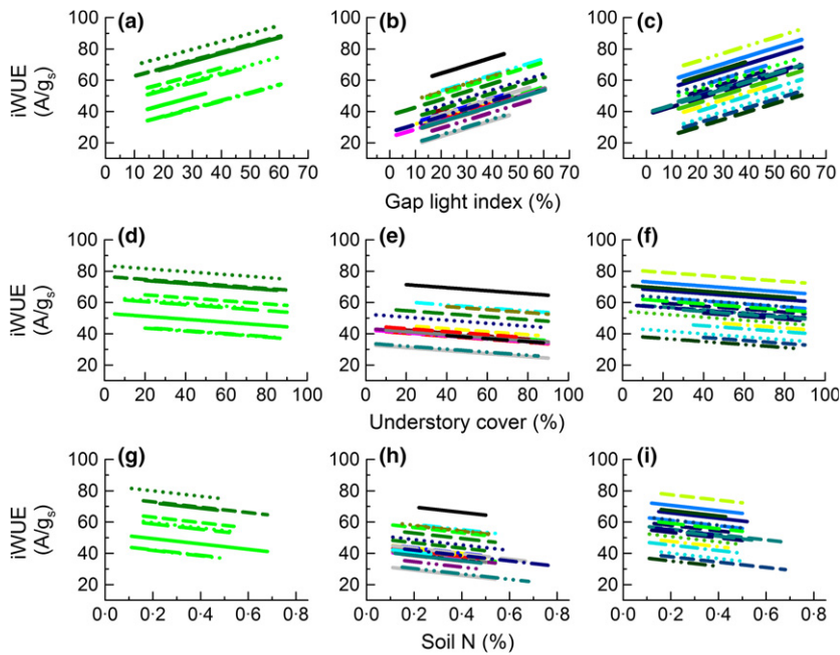


Fig. 3. Relationships of intrinsic water-use efficiency (iWUE) to gap light index (a–c), understorey cover (d–f) and soil N (g–i). Ferns and graminoids are shown in the first column (a, d, g), forbs in the second (b, e, h) and shrubs in the third (c, f, i). Lines show model estimates from selected models (Table 2) for individual species plotted across the range of conditions in which they were sampled (see Fig. 2 for legend).

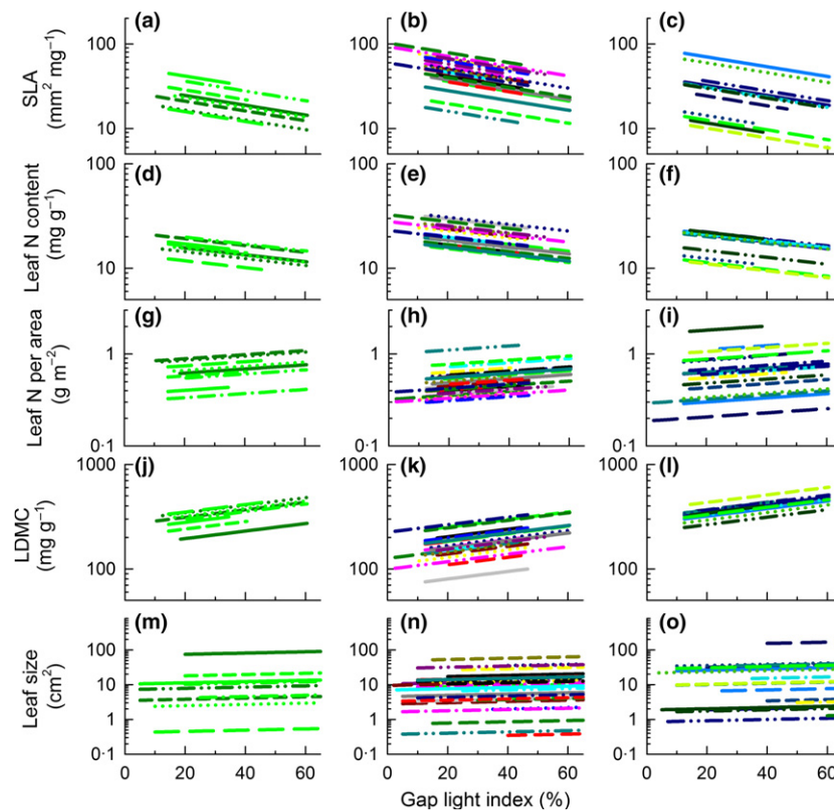


Fig. 4. Relationships between leaf traits (rows) and gap light index (GLI). Ferns and graminoids are shown in the first column (a, d, g, j, m), forbs in the second (b, e, h, k, n) and shrubs in the third (c, f, i, l, o). Lines show model estimates from selected models (Table 2) for individual species plotted across the range of conditions in which they were sampled (see Fig. 2 for legend). Understorey cover was held at ‘low’ levels (20%, rather than species means) for plotting trait – GLI relationships when final models included an interaction between GLI and understorey cover (SLA and LDMC) because relationships between traits and GLI were less pronounced at high understorey cover. Similarly, in these cases GLI was held constant at high levels (0–40) for plotting trait–understorey cover relationships (Fig. S3.1). Relationships are plotted on the log scale (y-axis) to ease interpretation. Note differences in scale range for height between (b), and (a) and (c). SLA, specific leaf area; LDMC, leaf dry matter content.

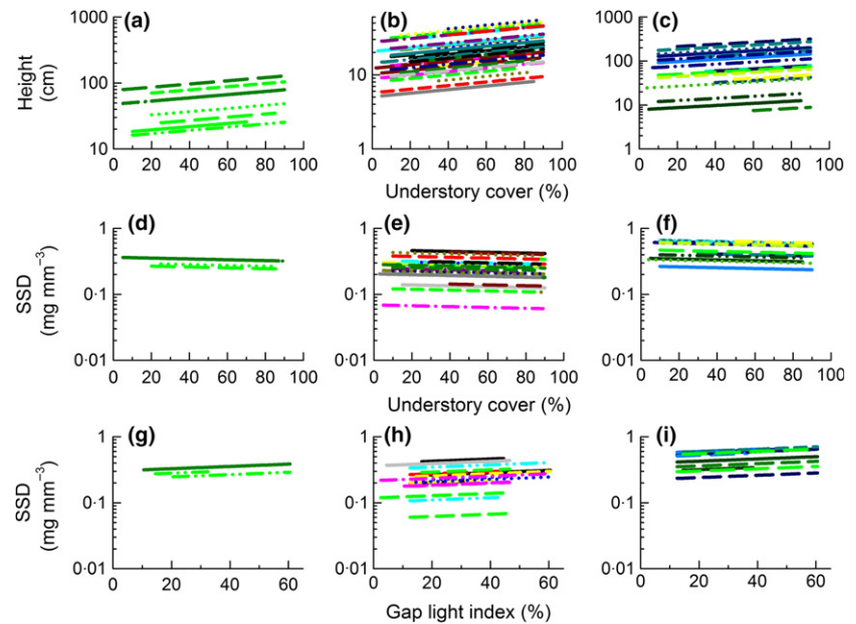


Fig. 5. Relationship between (a) plant height and understory cover and (b) stem-specific density (SSD) and gap light index at low understory cover. Lines show model estimates from selected models (Table 2) for individual species plotted across the range of conditions in which they were sampled (see Fig. 2 for legend). Relationships are plotted on the log scale (y-axis) to ease interpretation.

that variation did not lead to species cross-over. Thus, species rankings established in one set of environmental and resource conditions appear to hold under a broader range of conditions. In contrast, rankings may not be valid when individual species are sampled under different conditions (e.g. a shade-tolerant species sampled in the shade compared to an intolerant species sampled in a gap). Future studies examining the generality of our findings in other ecosystems, for other plant groups and over larger gradients in resources and environmental conditions would be worthwhile. High levels of intraspecific variability suggest that a larger sample size may be required when characterizing reaction norms across a wider range of environmental conditions.

STABLE ISOTOPE-DERIVED WHOLE-PLANT TRAITS

The range of variation in foliar N stable isotope discrimination relative to soil ($\Delta^{15}\text{N}$) provides a metric of niche partitioning in N use strategies among plants due to differences in the forms, timing and depth of N uptake from soil, as well as internal plant N distribution and plant-mycorrhizal associations (e.g. Nadelhoffer *et al.* 1996; Gubsch *et al.* 2011). Estimates of $\Delta^{15}\text{N}$ also control for foliar $\delta^{15}\text{N}$ tracking of soil $\delta^{15}\text{N}$ across sites, which can otherwise obscure or confound differences in plant uptake strategies (Houlton *et al.* 2007). We observed significant variation in $\Delta^{15}\text{N}$ that confirms high intraspecific variation in N use strategies for the understory species examined. Nearly all species displayed negative $\Delta^{15}\text{N}$ values, which is typical of plants that rely on soil inorganic N (Nadelhoffer *et al.* 1996). For these species, the decrease in $\Delta^{15}\text{N}$ (i.e. indicating a broader span between plant and soil $\delta^{15}\text{N}$) with increasing understory cover indicates intensified niche partitioning and diminished variation in niche breadth (e.g. timing, depth and form) of N uptake when

plant competition for N is high (Gubsch *et al.* 2011). However, the low semi-partial coefficient (R^2_{β}) for understory cover (0.02) suggests effects of competition on $\Delta^{15}\text{N}$ are relatively weak. Interspecific differences (i.e. phylogenetic N use strategies) and broader-scale variation in abiotic variables (soil N and CMD) were relatively more important. Low $\Delta^{15}\text{N}$ values (i.e. broader span between plant and soil $\delta^{15}\text{N}$) in most species were also associated with low soil N, consistent with patterns of overstorey $\delta^{15}\text{N}$ in these forests (Perakis, Sinkhorn & Compton 2011; Perakis, Tepley & Compton 2015). This can reflect greater N niche partitioning at low N sites, increased reliance on NH_4^+ uptake and/or greater $\delta^{15}\text{N}$ discrimination by mycorrhizae (Gubsch *et al.* 2011). A majority of species also displayed increasing $\Delta^{15}\text{N}$ values (i.e. narrower span between plant and soil $\delta^{15}\text{N}$) with higher moisture deficits, consistent with greater uptake of N from deeper soil horizons (Nadelhoffer *et al.* 1996) and/or uptake of soil nitrate that has been isotopically enriched by denitrification (Austin & Vitousek 1998). At the level of individual species, the unusually high $\delta^{15}\text{N}$ (i.e. the only species with positive values of $\Delta^{15}\text{N}$) and rather ordinary $\delta^{13}\text{C}$ values of *Chimaphila menziesii* is noteworthy. Species in this genus can be myco-heterotrophic, and our stable isotope patterns suggest it derived most N directly from fungal partners while deriving C from either photosynthesis or recently produced plant photosynthates (Zimmer *et al.* 2007).

Summer moisture deficits are generally believed to limit photosynthesis and stomatal conductance in the Pacific Northwest (e.g. Waring & Franklin 1979). Our results showed that iWUE of understory species increased, rather than decreased, with GLI. This suggests increases in photosynthetic carbon assimilation relative to stomatal conductance at low overstorey densities are due to resource levels and/or increasing stomatal conductance associated with higher relative humidity in closed canopies. A positive

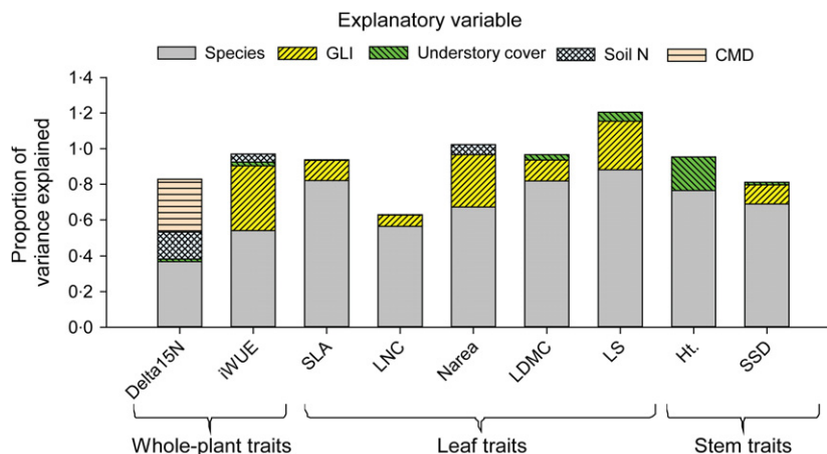


Fig. 6. Proportion of the total variance explained by fixed effects in the selected models for each whole-plant, leaf, stem and root trait. GLI, gap light index; CMD, climatic moisture deficit; $\Delta^{15}\text{N}$, nitrogen stable isotope discrimination; iWUE, intrinsic water-use efficiency; SLA, specific leaf area; LNC, leaf nitrogen content; N_{area} , leaf nitrogen per unit area; LDMC, leaf dry matter content; LS, leaf size; SSD, stem-specific density.

correlation between iWUE and leaf nitrogen per unit area (N_{area} ; $r = 0.38$, $P < 0.0001$), which also increases with GLI (Fig. 6), indicates that greater assimilation rates with lower overstorey densities was the more likely mechanism. The negative relationship between iWUE and understorey cover may reflect effects of understorey cover on light below the 1 m camera height that is not captured by GLI. However, in contrast to previous work, which focused on overstorey trees (e.g. Waring & Franklin 1979) our results suggest a dominant role of light in determining iWUE of understorey species and that the role of water limitations in forests in the Pacific Northwest may vary for different vegetation layers.

LEAF TRAITS

The relationships between leaf traits and resources, environmental conditions and understorey cover may have resulted from a trade-off between high structural investments per unit area in well-lit conditions vs. light absorption in shady conditions. This interpretation is supported by the negative relationship of SLA and leaf size to GLI, and the positive of LDMC to GLI (Fig. 6). This is consistent with the reversed response of SLA and LDMC (i.e. positive) in response to understorey cover (Fig. S2). Decreases in SLA with irradiance is associated with a lower density of thylakoids per stroma volume/grana and increases in palisade parenchyma cell thickness, chloroplasts and nitrogen per unit area (N_{area}), biochemical photosynthetic capacity and respiration (Givnish 1988; Lambers, Chapin & Pons 1998). In contrast, higher SLA in shade is associated with increased mass-based LNC and an increased proportion of spongy mesophyll leading to more efficient light capture per unit biomass and longer leaf life spans. Longer leaf life spans without additional structural investment are favoured in forest understoreys characterized by relatively low wind speeds and high

relative humidity (Westoby *et al.* 2002; Lusk *et al.* 2008). Lower mass-based, and higher area-based, LNC in sun relative to shade, respectively, have been documented for a wide variety of plants (Chazdon 1992; Ellsworth & Reich 1992, 1993; Givnish, Montgomery & Goldstein 2004). Deciduous and evergreen species show similar plastic responses of SLA (or LMA) to shade as found in our study, likely as a result of selection for low construction and maintenance costs (Lusk *et al.* 2008).

STEM TRAITS

The observed relationships of stem traits to light transmission and understorey cover may be related to trade-offs between competition in dense understoreys and mechanical safety where canopy conditions are more open (Givnish 1995). Stem-specific density increased with GLI and decreased with understorey cover, whereas height increased (Fig. 6). Shade and competition with neighbours can decrease mechanical stability as a result of lower stem diameter relative to height, reducing SSD and root : shoot ratios. However, plants can increase SSD, and root : shoot ratios after overstorey removal and decrease leaf area to control stem deflection in wind (Henry & Thomas 2002; Briggs *et al.* 2012). Allocation of a greater proportion of a plant's resources to stems in the form of height or SSD comes at a cost of allocation to photosynthetically active leaves. Our results empirically support the theoretical prediction that the optimal allocation to stems increases with competition (i.e. understorey cover) and productivity (Givnish 1982). Additional effects of soil N did not improve models of SSD and height, suggesting productivity effects can be predicted locally with understorey cover and light transmission. Alternatively, local variation in soil total N is not clearly reflective of plant-available N. High SSD can lead to lower hydraulic conductance (Meinzer *et al.* 2008; Chave *et al.* 2009). However, the benefits of

increasing SSD for mechanical safety at high GLI and low understorey cover appear to outweigh the potential costs to hydraulic conductance.

ROOT TRAITS

Variation in SRL and rooting depth was not strongly related to resource or environmental variables assessed here. Root development patterns may be similar to those for leaves, where longer root investments per unit mass can increase the capacity for uptake and proliferation at the expense of tolerance for xeric or infertile soils (Fitter 1985; Eissenstat 1991; Eissenstat & Yanai 1997). As such, positive relationships between SRL and resources, especially, soil N and moisture, and negative relationships with competition may be expected. On the other hand, negative relationships between SRL and soil resources, and positive relationships between SRL and competition support the hypothesis that plant allocation to roots is lower where resources are high, and necessarily higher where resources are low (Agren & Franklin 2003; Ostonen *et al.* 2007). The lack of clear support for either alternative in our data might suggest that SRL generally does not vary in response to resources levels, at least within the resource range measured in our study. It is possible, however, that relevant resources were not measured at an appropriately fine scale. For example, soil properties may vary significantly within the sampling radius of our plots as a result of variation in tree and understorey species composition, pit and mound topography and soil disturbance (e.g. Ettema & Wardle 2002). Moreover, understorey species vary with respect to root branching patterns and the presence of fine roots. We focused on the outermost live roots, including only fine roots when possible. For species lacking fine roots (e.g. *Goodyera oblongifolia*, *Disporum* spp., *Asarum caudatum*), we measured the finest of the live coarse roots. Controlling for variation among root branching orders would have allowed us to better detect such relationships, but was not possible in our study (Pregitzer *et al.* 2002; Ostonen *et al.* 2007). Similar interpretations may apply to the results for maximum rooting depth (Antos & Halpern 1997).

Authors' contributions

J.I.B., K.J.P. and S.S.P. conceived the ideas and designed the methodology; C.E.L., S.C.M. and J.I.B. collected the data; J.I.B. analysed the data; J.I.B. led the writing of the manuscript. All authors contributed critically to the drafts and gave final approval for publication.

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Data accessibility

Data will be deposited in the Dryad Digital Repository <http://dx.doi.org/10.5061/dryad.8125b> (Burton *et al.* 2017).

References

- Agren, G.I. & Franklin, O. (2003) Root: shoot ratios, optimization and nitrogen productivity. *Annals of Botany*, **92**, 795–800.
- Albert, C.H., Thuiller, W., Yoccoz, N.G., Soudant, A., Boucher, F., Saccone, P. & Lavorel, S. (2010) Intraspecific functional variability: extent, structure and sources of variation. *Journal of Ecology*, **98**, 604–613.
- Antos, J.A. & Halpern, C.B. (1997) Root system differences among species: implications for early successional changes in forest of western Oregon. *American Midland Naturalist*, **138**, 97–108.
- Ares, A., Berryman, S.D. & Puettmann, K.J. (2009) Understorey vegetation response to thinning disturbance of varying complexity in coniferous stands. *Applied Vegetation Science*, **12**, 472–487.
- Auger, S. & Shipley, B. (2013) Inter-specific and intra-specific trait variation along short environmental gradients in an old-growth temperate forest. *Journal of Vegetation Science*, **24**, 419–428.
- Austin, A.T. & Vitousek, P. (1998) Nutrient dynamics on a precipitation gradient in Hawai'i. *Oecologia*, **113**, 519–529.
- Bolnick, D.I., Amarasekare, P., Araujo, M.S. *et al.* (2011) Why intraspecific trait variation matters in community ecology. *Trends in Ecology & Evolution*, **26**, 183–192.
- Briggs, N.A., Kuehne, C., Kohnle, U. & Bauhus, J. (2012) Root system response of naturally regenerated Douglas-fir (*Pseudotsuga menziesii*) after complete overstorey removal. *Canadian Journal of Forest Research*, **42**, 1858–1864.
- Brooks, J.R., Flanagan, L.B., Buchmann, N. & Ehleringer, J.R. (1997) Carbon isotope composition of boreal plants: functional grouping of life forms. *Oecologia*, **110**, 301–311.
- Buchmann, N., Brooks, J.R. & Ehleringer, J.R. (2002) Predicting daytime carbon isotope ratios of atmospheric CO₂ within forest canopies. *Functional Ecology*, **16**, 49–57.
- Burnham, K.P. & Anderson, D.R. (2002) *Model Selection and Multimodel Inference: A Practical Information-Theoretic Approach*. Springer, New York, NY, USA.
- Burton, J.I., Mladenoff, D.J., Clayton, M.K. & Forrester, J.A. (2011) The roles of environmental filtering and colonization in the fine-scale spatial patterning of ground-layer plant communities in north temperate deciduous forests. *Journal of Ecology*, **99**, 764–776.
- Burton, J.I., Mladenoff, D.J., Forrester, J.A. & Clayton, M.K. (2014) Experimentally linking disturbance, resources and productivity to diversity in forest ground-layer plant communities. *Journal of Ecology*, **102**, 1634–1648.
- Burton, J.I., Perakis, S.S., McKenzie, S.C., Lawrence, C.E. & Puettmann, K.J. (2017) Data from: Intraspecific variability and reaction norms of forest understorey plant species traits. *Dryad Digital Repository*, <http://dx.doi.org/10.5061/dryad.8125b>.
- Chave, J., Coomes, D., Jansen, S., Lewis, S.L., Swenson, N.G. & Zanne, A.E. (2009) Towards a worldwide wood economics spectrum. *Ecology Letters*, **12**, 351–366.
- Chazdon, R.L. (1992) Photosynthetic plasticity of two rain forest shrubs across natural gap transects. *Oecologia*, **92**, 586–595.
- Cissel, J.H., Anderson, P.D., Olson, D., Puettmann, K.J., Berryman, S.D., Chan, S. & Thompson, C. (2006) BLM density management and riparian buffer study: Establishment report and study plan. US Geological Survey, Scientific Investigations Report, 2006–5087.
- Daubenmire, R. (1976) The use of vegetation in assessing the productivity of forest lands. *The Botanical Review*, **42**, 115–143.
- Diaz, S., Hodgson, J.G., Thompson, K. *et al.* (2004) The plant traits that drive ecosystems: evidence from three continents. *Journal of Vegetation Science*, **15**, 295–304.

- Edwards, L.J., Muller, K.E., Wolfinger, R.D., Qaqish, B.F. & Schabenberger, O. (2008) An R^2 statistic for fixed effects in the linear mixed model. *Statistics in Medicine*, **27**, 6137–6157.
- Ehleringer, J.R. (1991) $^{13}C/^{12}C$ fractionation and its utility in terrestrial plant studies. *Carbon Isotope Techniques*, **1**, 187.
- Eissenstat, D.M. (1991) On the relationship between specific root length and the rate of root proliferation: a field study using citrus rootstocks. *New Phytologist*, **118**, 63–68.
- Eissenstat, D. & Yanai, R. (1997) The ecology of root lifespan. *Advances in Ecological Research*, **27**, 60.
- Ellsworth, D.S. & Reich, P.B. (1992) Leaf mass per area, nitrogen content and photosynthetic carbon gain in *Acer saccharum* seedlings in contrasting forest light environments. *Functional Ecology*, **6**, 423–435.
- Ellsworth, D. & Reich, P. (1993) Canopy structure and vertical patterns of photosynthesis and related leaf traits in a deciduous forest. *Oecologia*, **96**, 169–178.
- Ettema, C.H. & Wardle, D.A. (2002) Spatial soil ecology. *Trends in Ecology & Evolution*, **17**, 177–183.
- Evans, R.D. (2001) Physiological mechanisms influencing plant nitrogen isotope composition. *Trends in Plant Science*, **6**, 121–126.
- Farquhar, G.D., Ehleringer, J.R. & Hubick, K.T. (1989b) Carbon isotope discrimination and photosynthesis. *Annual Review of Plant Biology*, **40**, 503–537.
- Farquhar, G., Hubick, K., Condon, A. & Richards, R. (1989a) Carbon isotope fractionation and plant water-use efficiency. *Stable Isotopes in Ecological Research*, **68**, 21–40.
- Farquhar, G.D., O'Leary, M.H. & Berry, J.A. (1982) On the relationship between carbon isotope discrimination and the intercellular carbon dioxide concentration in leaves. *Functional Plant Biology*, **9**, 121–137.
- Fitter, A.H. (1985) Functional significance of root morphology and root system architecture. *Ecological Interactions in Soil* (eds D. Atkinson, A.H. Fitter, D.J. Read & M.B. Usher), pp. 87–106. Blackwell Scientific, Oxford, UK.
- Foster, T.E. & Brooks, J.R. (2005) Functional groups based on leaf physiology: are they spatially and temporally robust? *Oecologia*, **144**, 337–352.
- Franklin, J.F. & Dyrness, C.T. (1988) *Natural Vegetation of Oregon and Washington*. Oregon State University Press, Corvallis, OR, USA.
- Funk, J.L., Larson, J.E., Ames, G.M. et al. (2016) Revisiting the Holy Grail: using plant functional traits to understand ecological processes. *Biological Reviews*, **92**, 1156–1173.
- Garnier, E., Laurent, G., Bellmann, A., Debain, S., Berthelot, P., Ducout, B., Roumet, C. & Navas, M.L. (2001) Consistency of species ranking based on functional leaf traits. *New Phytologist*, **152**, 69–83.
- Gilliam, F.S. (2007) The ecological significance of the herbaceous layer in temperate forest ecosystems. *BioScience*, **57**, 845–858.
- Givnish, T.J. (1982) On the adaptive significance of leaf height in forest herbs. *American Naturalist*, **120**, 353–381.
- Givnish, T.J. (1988) Adaptation to sun and shade – a whole-plant perspective. *Australian Journal of Plant Physiology*, **15**, 63–92.
- Givnish, T.J. (1995) Plant stems: biomechanical adaptation for energy capture and influence on species distributions. *Plant Stems: Physiology and Functional Morphology* (ed. B.L. Gartner), pp. 3–49. Chapman and Hall, New York, NY, USA.
- Givnish, T.J., Montgomery, R.A. & Goldstein, G. (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.
- Grime, J.P. (1998) Benefits of plant diversity to ecosystems: immediate, filter and founder effects. *Journal of Ecology*, **86**, 902–910.
- Gubsch, M., Roscher, C., Gleixner, G., Habekost, M., Lipowsky, A., Schmid, B., Schultze, E.D., Steinbeiss, S. & Buchmann, N. (2011) Foliar and soil $\delta^{15}N$ values reveal increased nitrogen partitioning among species in diverse grassland communities. *Plant, Cell & Environment*, **34**, 895–908.
- Hart, S.P., Schreiber, S.J. & Levine, J.M. (2016) How variation between individuals affects species coexistence. *Ecology Letters*, **19**, 825–838.
- Henry, H.A. & Thomas, S.C. (2002) Interactive effects of lateral shade and wind on stem allometry, biomass allocation, and mechanical stability in *Abutilon theophrasti* (Malvaceae). *American Journal of Botany*, **89**, 1609–1615.
- Houlton, B.Z., Sigman, D.M., Schuur, E.A. & Hedin, L.O. (2007) A climate-driven switch in plant nitrogen acquisition within tropical forest communities. *Proceedings of the National Academy of Sciences*, **104**, 8902–8906.
- Kaelke, C.M., Kruger, E.L. & Reich, P.B. (2001) Trade-offs in seedling survival, growth, and physiology among hardwood species of contrasting successional status along a light-availability gradient. *Canadian Journal of Forest Research*, **31**, 1602–1616.
- Kattge, J., Diaz, S., Lavorel, S. et al. (2011) TRY—a global database of plant traits. *Global Change Biology*, **17**, 2905–2935.
- Kazakou, E., Violle, C., Roumet, C., Navas, M.L., Vile, D., Kattge, J. & Garnier, E. (2014) Are trait-based species rankings consistent across data sets and spatial scales? *Journal of Vegetation Science*, **25**, 235–247.
- Kern, C.C., Montgomery, R.A., Reich, P.B. & Strong, T.F. (2014) Harvest-created canopy gaps increase species and functional trait diversity of the forest ground-layer community. *Forest Science*, **60**, 335–344.
- Lambers, H., Chapin, F.S. & Pons, T.L. (1998) *Plant Physiological Ecology*. Springer, New York, NY, USA.
- Laughlin, D.C., Joshi, C., Bodegom, P.M., Bastow, Z.A. & Fulé, P.Z. (2012) A predictive model of community assembly that incorporates intraspecific trait variation. *Ecology Letters*, **15**, 1291–1299.
- Lepš, J., de Bello, F., Šmilauer, P. & Doležal, J. (2011) Community trait response to environment: disentangling species turnover vs intraspecific trait variability effects. *Ecography*, **34**, 856–863.
- Lusk, C.H., Reich, P.B., Montgomery, R.A., Ackerly, D.D. & Cavender-Bares, J. (2008) Why are evergreen leaves so contrary about shade? *Trends in Ecology & Evolution*, **23**, 299–303.
- McGill, B.J., Enquist, B.J., Weiher, E. & Westoby, M. (2006) Rebuilding community ecology from functional traits. *Trends in Ecology & Evolution*, **21**, 178–185.
- Meinzer, F.C., Woodruff, D.R., Domec, J.-C., Goldstein, G., Campanello, P.I., Gatti, M.G. & Villalobos-Vega, R. (2008) Coordination of leaf and stem water transport properties in tropical forest trees. *Oecologia*, **156**, 31–41.
- Nadelhoffer, K., Shaver, G., Fry, B., Giblin, A., Johnson, L. & McKane, R. (1996) ^{15}N natural abundances and N use by tundra plants. *Oecologia*, **107**, 386–394.
- Nakagawa, S. & Schielzeth, H. (2013) A general and simple method for obtaining R^2 from generalized linear mixed-effects models. *Methods in Ecology and Evolution*, **4**, 133–142.
- Neill, A.R. & Puettmann, K.J. (2013) Managing for adaptive capacity: thinning improves food availability for wildlife and insect pollinators under climate change conditions. *Canadian Journal of Forest Research*, **43**, 428–440.
- Nicotra, A.B., Atkin, O.K., Bonser, S.P. et al. (2010) Plant phenotypic plasticity in a changing climate. *Trends in Plant Science*, **15**, 684–692.
- Ostonen, I., Püttsepp, Ü., Biel, C. et al. (2007) Specific root length as an indicator of environmental change. *Plant Biosystems*, **141**, 426–442.
- Paine, C., Baraloto, C. & Díaz, S. (2015) Optimal strategies for sampling functional traits in species-rich forests. *Functional Ecology*, **29**, 1325–1331.
- Perakis, S.S., Sinkhorn, E.R. & Compton, J.E. (2011) $\delta^{15}N$ constraints on long-term nitrogen balances in temperate forests. *Oecologia*, **167**, 793–807.
- Perakis, S.S., Tepley, A.J. & Compton, J.E. (2015) Disturbance and topography shape nitrogen availability and $\delta^{15}N$ over long-term forest succession. *Ecosystems*, **18**, 573–588.
- Pérez-Harguindeguy, N., Díaz, S., Garnier, E. et al. (2013) New handbook for standardised measurement of plant functional traits worldwide. *Australian Journal of Botany*, **61**, 167–234.
- Pierce, S., Negreiros, D., Cerabolini, B.E.L. et al. (2017) A global method for calculating plant CSR ecological strategies applied across biomes world-wide. *Functional Ecology*, **31**, 444–457.
- Pregitzer, K.S., DeForest, J.L., Burton, A.J., Allen, M.F., Ruess, R.W. & Hendrick, R.L. (2002) Fine root architecture of nine North American trees. *Ecological Monographs*, **72**, 293–309.
- Ravenscroft, C.H., Fridley, J.D. & Grime, J.P. (2014) Intraspecific functional differentiation suggests local adaptation to long-term climate change in a calcareous grassland. *Journal of Ecology*, **102**, 65–73.
- Roche, P., Díaz-Burlinson, N. & Gachet, S. (2004) Congruency analysis of species ranking based on leaf traits: which traits are the more reliable? *Plant Ecology*, **174**, 37–48.
- Sabatini, F.M., Burton, J.I., Scheller, R.M., Amatangelo, K.L. & Mladenoff, D.J. (2014) Functional diversity of ground-layer plant communities in old-growth and managed northern hardwood forests. *Applied Vegetation Science*, **17**, 398–407.
- Shipley, B., Bello, F., Cornelissen, J.H.C., Laliberté, E., Laughlin, D.C. & Reich, P.B. (2016) Reinforcing loose foundation stones in trait-based plant ecology. *Oecologia*, **180**, 923–931.
- Siefert, A., Violle, C., Chalmandrier, L. et al. (2015) A global meta-analysis of the relative extent of intraspecific trait variation in plant communities. *Ecology Letters*, **18**, 1406–1419.

- Sonnier, G., Johnson, S.E., Amatangelo, K.L., Rogers, D.A. & Waller, D.M. (2014) Is taxonomic homogenization linked to functional homogenization in temperate forests? *Global Ecology and Biogeography*, **23**, 894–902.
- Suding, K.N., Lavorel, S., Chapin, F.S. *et al.* (2008) Scaling environmental change through the community-level: a trait-based response-and-effect framework for plants. *Global Change Biology*, **14**, 1125–1140.
- Thiel, A.L. & Perakis, S.S. (2009) Nitrogen dynamics across silvicultural canopy gaps in young forests of western Oregon. *Forest Ecology and Management*, **258**, 273–287.
- Violle, C., Enquist, B.J., McGill, B.J., Jiang, L., Albert, C.H., Hulshof, C., Jung, V. & Messier, J. (2012) The return of the variance: intraspecific variability in community ecology. *Trends in Ecology & Evolution*, **27**, 244–252.
- Wagner, R.G. & Radosovich, S.R. (1998) Neighborhood approach for quantifying interspecific competition in coastal Oregon forests. *Ecological Applications*, **8**, 779–794.
- Wang, T., Hamann, A., Spittlehouse, D.L. & Muddock, T.Q. (2012) ClimateWNA—high-resolution spatial climate data for western North America. *Journal of Applied Meteorology and Climatology*, **51**, 16–29.
- Waring, R.H. & Franklin, J.F. (1979) Evergreen coniferous forests of the Pacific Northwest. *Science*, **204**, 1380–1386.
- Westoby, M., Falster, D.S., Moles, A.T., Vesk, P.A. & Wright, I.J. (2002) Plant ecological strategies: some leading dimensions of variation between species. *Annual Review of Ecology and Systematics*, **33**, 125–159.
- Wright, I.J., Reich, P.B., Westoby, M. *et al.* (2004) The worldwide leaf economics spectrum. *Nature*, **428**, 821–827.
- Zimmer, K., Hynson, N.A., Gebauer, G., Allen, E.B., Allen, M.F. & Read, D.J. (2007) Wide geographical and ecological distribution of nitrogen and carbon gains from fungi in pyrolloids and monotropoids (Ericaceae) and in orchids. *New Phytologist*, **175**, 166–175.

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Supporting Information

Details of electronic Supporting Information are provided below.

Appendix S1. Detailed information related to the study area, data collection and statistical analysis.

Fig. S1. Illustration of species cross-over.

Fig. S2. Relationship between leaf traits and understorey cover.

Fig. S3. Relationship of leaf N per area to soil N.

Table S1. Step-wise model selection process.

Table S2. Number of individuals sampled by species and trait.

Table S3. Alternative model comparisons.

Table S4. Correlations among fixed effects.