

From the Arctic to the tropics: multibiome prediction of leaf mass per area using leaf reflectance

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Summary

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- Leaf mass per area (LMA) is a key plant trait, reflecting tradeoffs between leaf photosynthetic function, longevity, and structural investment. Capturing spatial and temporal variability in LMA has been a long-standing goal of ecological research and is an essential component for advancing Earth system models. Despite the substantial variation in LMA within and across Earth's biomes, an efficient, globally generalizable approach to predict LMA is still lacking.
- We explored the capacity to predict LMA from leaf spectra across much of the global LMA trait space, with values ranging from 17 to 393 g m⁻². Our dataset contained leaves from a wide range of biomes from the high Arctic to the tropics, included broad- and needleleaf species, and upper- and lower-canopy (i.e. sun and shade) growth environments.
- Here we demonstrate the capacity to rapidly estimate LMA using only spectral measurements across a wide range of species, leaf age and canopy position from diverse biomes. Our model captures LMA variability with high accuracy and low error ($R^2 = 0.89$; root mean square error (RMSE) = 15.45 g m⁻²).
- Our finding highlights the fact that the leaf economics spectrum is mirrored by the leaf optical spectrum, paving the way for this technology to predict the diversity of LMA in ecosystems across global biomes.

Introduction

A key objective of plant ecology is to characterize the functional diversity of plants that have evolved different strategies for growth, reproduction and for coping with biotic and abiotic stressors (Wright *et al.*, 2004). Accurate characterization of this functional diversity in Earth system models (ESMs) will improve our ability to model the Earth system and understand the effect of global change on the cycling and storage of carbon (C), water and energy (Pavlick *et al.*, 2013; Fisher *et al.*, 2018). Therefore, an increasing number of ESMs are moving towards incorporating approaches that require a broader and more comprehensive representation of plant trait variation within and across biomes (Xu *et al.*, 2012; van Bodegom *et al.*, 2014; Wullschlegel *et al.*, 2014; Fisher *et al.*, 2015, 2018). As a result, considerable effort has been invested in the measurement and monitoring of plant traits across a range of biomes, and the storage and synthesis of that information in global databases (Wright *et al.*, 2004; Kattge *et al.*, 2011; Lebauer *et al.*, 2013; Butler *et al.*, 2017). Yet, the high degree of plant functional diversity and plasticity makes this apparently simple goal extremely challenging (Reich *et al.*, 1997, 1999;

Serbin *et al.*, 2014; Wu *et al.*, 2017; Osnas *et al.*, 2018). Consequently, the extent of global trait coverage is still woefully inadequate (Schimel *et al.*, 2015).

In recent decades, remote sensing has shown increasing promise as a means to capture plant traits across scales using spectroscopic approaches. For example, several recent studies have highlighted the capacity to connect remotely sensed spectra to characterize variation in a number of key functional traits across individual leaves, canopies and landscapes (Dahlin *et al.*, 2013; Asner *et al.*, 2015; Singh *et al.*, 2015; Shiklomanov *et al.*, 2016; Yang *et al.*, 2016; Wu *et al.*, 2017). Approaches have included empirical spectra–trait modeling, such as partial least-squares regression (PLSR; e.g. Serbin *et al.*, 2014; Singh *et al.*, 2015; Yendrek *et al.*, 2017) and spectral vegetation indices (SVIs; e.g. Feret *et al.*, 2011), as well as semimechanistic approaches, including the use of the PROSPECT leaf-level radiative transfer model (RTM; e.g. Shiklomanov *et al.*, 2016; Féret *et al.*, 2017). In general, all of these approaches rely on the fundamental biophysical connection between leaf chemistry and structure and the resultant optical properties of plants (Curran, 1989; Ustin *et al.*, 2004; Kokaly *et al.*, 2009; Ollinger, 2011), and, as a result, could be

used to fill critical gaps in our understanding of the variation in plant traits over landscapes and biomes (Schimel *et al.*, 2015; Jetz *et al.*, 2016).

However, most studies that have illustrated the strong promise of remote sensing to estimate foliar traits across scales have focused on a relatively narrow subsample of geographic regions (e.g. Townsend *et al.*, 2003; Feret *et al.*, 2011; Dahlin *et al.*, 2013; Asner *et al.*, 2015; Yang *et al.*, 2016), capturing only a small fraction of total trait and spectral space. In addition, semimechanistic methods such as PROSPECT may have other assumptions or limitations that could inhibit their broad application, for example, a limited number of supported traits (Féret *et al.*, 2017) or the generally poorer performance of needleleaf species (Malenovský *et al.*, 2006; Shiklomanov *et al.*, 2016). As a result, the general applicability of existing spectra–trait models and approaches across biomes and in the wider trait space is not known. In many cases, those models trained across a limited trait space have been shown to break down when applied more broadly (e.g. Sims & Gamon, 2002; Gitelson *et al.*, 2003; le Maire *et al.*, 2008).

Among plant traits, foliar morphology is commonly characterized using leaf mass per area (LMA) – the ratio of a leaf's dry mass to its surface area (g dry mass m^{-2} leaf area) – or its reciprocal, specific leaf area. LMA captures the tradeoff of a plant's investment in leaf structure and robustness vs leaf surface area and light harvesting for photosynthesis (Wright *et al.*, 2004; Shipley *et al.*, 2006; Poorter *et al.*, 2009). Given its strong linkage with overall plant functioning (Reich *et al.*, 1997; Wright *et al.*, 2004; Serbin *et al.*, 2012; Osnas *et al.*, 2018), LMA is a critical parameter in plant ecology. Illustrations of the importance of LMA include its use as a basis for monitoring biodiversity (Skidmore *et al.*, 2015), its role in modeling canopy radiation transfer (Jacquemoud *et al.*, 2009; Ollinger, 2011), and its widespread use as an input in ecosystem process models (Fisher *et al.*, 2014; Xu *et al.*, 2016; Ricciuto *et al.*, 2018).

Importantly, the substantial global variation in LMA, which ranges from 14 to 1500 g m^{-2} globally (Wright *et al.*, 2004), exists within and across species (Castro-Díez *et al.*, 2000; Wright *et al.*, 2004; Paula & Pausas, 2006; Poorter *et al.*, 2009) and is affected by local gradients in light, water and nutrient availability (Niinemets, 2007; de la Riva *et al.*, 2016; Liu *et al.*, 2017), leaf age (Wu *et al.*, 2016), as well as acclimation and adaptation to short- and long-term climate dynamics (Volin *et al.*, 2002; Paula & Pausas, 2006; Poorter *et al.*, 2009). LMA variation is attributed primarily to differences in leaf density and volume-to-area ratio (Poorter *et al.*, 2009; John *et al.*, 2017). The fundamental information on these attributes is found in the reflectance spectrum of a leaf, which captures its physical properties (e.g. thickness, density, the depth of palisade layers, albedo and elemental composition) (Ollinger, 2011). Given the coordination between leaf traits and optical properties, we expect that, using a spectroscopy approach, we can collapse the vast structural and functional diversity of leaves from different plant species, leaf types (e.g. grasses, forbs, broadleaf and needleleaf evergreen trees), across a wide range of environments, that is expressed as variation of LMA into a single generalizable model.

Here we present a broad, multibiome analysis linking fundamental covariation in LMA and spectroscopic (also known as hyperspectral) reflectance. We use a large dataset to develop a robust statistical model to infer LMA from corresponding leaf optical properties, and then validate this cross-biome model using independent datasets of additional spectra and LMA observations from a similar range of plant material as well as from external validation sources. The core training and validation datasets include leaves from the high Arctic to the tropics, and contain measurements from grasses, forbs, deciduous and evergreen shrubs, deciduous and evergreen broadleaf trees, needleleaf trees and crop species. They span a highly diverse range of leaf morphologies, including glabrous, highly reflective and waxy leaf types; microenvironments, including measurements from upper-canopy, sunlit leaves, and lower-canopy, shaded leaves; developmental stages, including recently emerged, mature and old leaves; and elevations, including measurements from sea level to $> 2000 \text{ m}$ above sea level. Thus, our analysis is based on data representing a large fraction of the global trait space for LMA.

Materials and Methods

Plant material

We collected and assembled a large dataset ($n = 2478$ leaves from > 176 species) of combined LMA and leaf reflectance spectra for model development from the high Arctic in northern Alaska to the tropics in central America and Brazil (Fig. 1). Our sites are distributed across a large proportion of the Earth's habitable climate space, encompassing a *c.* 40°C range of mean annual temperatures and a *c.* $200\text{--}2400 \text{ mm yr}^{-1}$ range in mean annual precipitation. Our sites in Alaska include coastal tundra vegetation within the Barrow Environmental Observatory, near Barrow (now Utqiagvik), Alaska (Brown *et al.*, 1980; Rogers *et al.*, 2017b) and dwarf and tall shrub vegetation on the Seward Peninsula (Rogers *et al.*, 2016; Serbin & Rogers, 2019). Our sites located in the Upper Midwest and northeastern US are dominated by northern temperate forest species, including deciduous broadleaf hardwoods and evergreen, needleleaf conifers (Serbin *et al.*, 2014). Measurements on Mediterranean and agricultural plants were conducted in the Coachella and Central valleys of California (Serbin *et al.*, 2015), and across an elevation gradient of sites ranging from low elevation woodlands to alpine forests in the Sierra Mountains (Goulden *et al.*, 2012; DuBois *et al.*, 2018). The data for tropical species were collected in several separate locations; a seasonal, wet evergreen Amazonian forest near Santarem, Brazil (Wu *et al.*, 2017); a seasonal, wet evergreen forest in the San Lorenzo Protected Area; a seasonal, dry forest in the Parque Natural Metropolitano near Panama City in the Republic of Panama (Wright *et al.*, 2003); trees within the Dona Inés Park urban forest site on the grounds of the Luis Muñoz Marín Foundation as well as other locations on the island of Puerto Rico (specific sites can be found in associated dataset metadata, Table S1), and a collection of plants grown in an artificial tropical forest within the Biosphere 2 facility (Walter & Carmen Lambrecht, 2004). All the data used in this study are

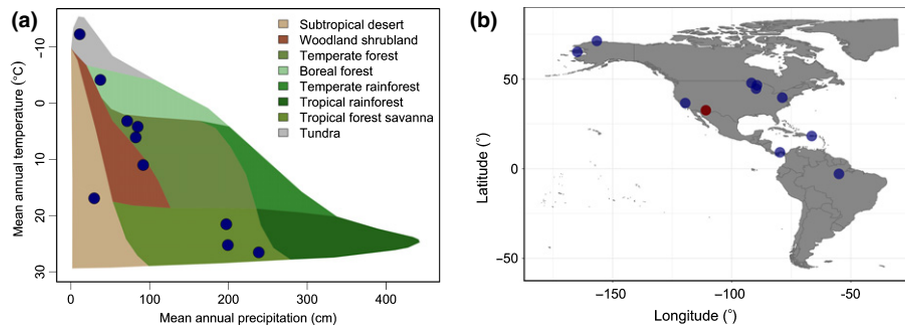


Fig. 1 General location of the 11 sites where leaf reflectance and leaf mass per area were measured in our model development dataset. Blue circles show field sites, and the red circle shows the location of Biosphere 2 from which we sampled tropical species within the glasshouse environment. (a) Sites plotted in climate space, binned by major climatic biomes (Whittaker, 1975); (b) general location in geographic space.

publicly available, and the published datasets include detailed descriptions of the sampling locations (Table S1).

Spectroscopic measurements

The approach, instrumentation and personnel used to measure leaf spectral reflectance varied across the different sites and projects included in this study. The leaf reflectance of Arctic plants was measured using a FieldSpec 3 (Analytical Spectral Devices (ASD) Inc., Longmont, CO, USA) with the ASD leaf clip assembly, or an HR-1024i full-spectrum spectroradiometer (Spectra Vista Corporation (SVC), Poughkeepsie, NY, USA) together with the standard leaf clip assembly as well as the more recent Leaf-Clip Reflectance-Probe Pro (LC-RP-Pro; SVC). Leaf spectra collected in the Upper Midwest and northeastern USA were measured with a FieldSpec pro or FieldSpec 3 (both ASD Inc.) with the ASD leaf clip assembly. The data collected in California were measured using either a FieldSpec 3 (ASD Inc.) or a PSR-3500 full-range spectrometer (Spectral Evolution, Lawrence, MA, USA) with an attached Spectral Evolution leaf clip assembly. Data from Biosphere2 and Brazil were measured with a FieldSpec Pro (ASD Inc.) and ASD leaf clip, the data from Puerto Rico with a PSR+ (Spectral Evolution) with an attached custom fiber optic connected to a LC-RP-Pro foreoptic (SVC), and data from the Republic of Panama were measured with an HR-1024i with the LC-RP-Pro attachment (SVC). All instruments had leaf clip assemblies that contained an internal calibrated light source, and all reflectance measurements were referenced against a 99% Spectralon reflectance standard. For needleleaf or very small leaf species, we used needle or leaf mats (Serbin, 2012). Needles of a similar age (i.e. current year, previous year, and older) were laid out edge to edge, creating a single layer and taped at the ends to hold the needles tightly together before inserting the mat into the leaf clip. No tape was visible to the fiberoptic inside the leaf clip. All spectral measurements were processed using the R-FIELDSPECTRA package (<https://github.com/serbinsh/R-FieldSpectra>). For SVC data, however, we first corrected the discontinuities in the spectra in the detector overlap areas using the vendor-provided software. Measurements at the edges of the spectral range of these spectrometers (350–500 nm, > 2400 nm) suffer from low signal-to-noise and were thus excluded from the analysis.

Measurements of LMA

Measurements of LMA also varied among the different studies and biomes. Various methods were used to determine leaf area, including measurement of length and width with a ruler and a hand lens (graminoid species), leaf disk punches (broadleaf species) and optical approaches. For small or compound leaves where it was not possible to take disks, area was measured with a leaf area meter (LI-3100C Area Meter; Li-Cor, Lincoln, NE, USA) calibrated for use with high edge-to-area ratio leaves and operated at high (0.1 mm²) resolution, or a flatbed scanner (evergreen needleleaf trees) followed by area estimation using IMAGEJ (Schneider *et al.*, 2012; as reported in Serbin *et al.*, 2014). Leaves and leaf sections of known area were then dried to constant mass in a ventilated oven (60–70°C) and leaf dry mass was measured on a top-pan balance. All leaf area data were provided on a projected leaf area basis.

LMA partial least-squares regression modeling

To relate the variability in LMA across sites, species, and environments we utilized a PLSR modeling approach (Geladi & Kowalski, 1986; Wold *et al.*, 2001) using the *pls* package (Mevik & Wehrens, 2007) in the R open source statistical environment (R Core Team, 2017). PLSR is widely utilized in spectroscopy and chemometric analyses given its ability to handle high predictor collinearity as well as a large number of predictor variables that may exceed the number of observations. PLSR minimizes the implications of these circumstances by reducing the number of predictor variables down to a relatively few, orthogonal latent components (Geladi & Kowalski, 1986; Wold *et al.*, 2001). Moreover, PLSR does not assume the measurement of predictor variables (reflectance values at given wavelengths in this case) was made without error.

Our PLSR model development has been described previously (e.g. Serbin *et al.*, 2014; Ely *et al.*, 2019) and is briefly summarized here. We first applied a square root transformation to the LMA data to reduce the right skewness distribution of the original data (Fig. S1) and satisfy the required normal distribution for the PLSR analysis. We then randomly split the full dataset into calibration (80%, $n = 1978$) and independent validation (20%,

$n = 500$) subsets (Fig. S2), ensuring that both subsets spanned the full range of observations and included measurements from each study and sample location. Using just the calibration dataset, we developed the multibiome leaf spectra–LMA model and then tested this final model using the validation data (500 observations) derived from our original dataset (2478 leaves). To avoid the potential to overfit the spectra–LMA calibration model, we optimized the number of PLSR components in the model by minimizing the prediction residual sum of squares (PRESS) statistic (Chen *et al.*, 2004). We calculated the PRESS statistic of successive model components through a cross-validation analysis where we minimized the PRESS statistic until successive PLSR components did not reduce model predictive error as assessed using a *t*-test (Serbin *et al.*, 2014). Lastly, we calculated the variable influence on projection (VIP) metric (Wold *et al.*, 2001) of the final model to identify the regions of the spectrum that contributed significantly to the prediction of LMA (Serbin *et al.*, 2014).

In addition to the general model development, we also conducted a PLSR model uncertainty analysis to characterize the predictive uncertainty, given the variability and error in measured LMA, spectra, and the relationship between the two. This was done by splitting the original calibration dataset into a balanced 70–30% via 1000 permutations and generating the same number of new model coefficient vectors, following Serbin *et al.* (2014). The result of this uncertainty analysis is an ensemble of PLSR models that can be used to predict new values of LMA based on spectral measurements plus the predictive uncertainty for each new value.

Finally, we quantified the performance of the multibiome LMA PLSR model using the independent validation dataset (500 observations). In this step, we validated the model and examined the residuals for model bias and predictive performance. We used four main evaluation metrics: the coefficient of determination (R^2), RMSE, the residual, and regression bias. All model and error results presented in the following are shown in original LMA units, not the square-root-transformed units that are the initial output of the PLSR model.

Examples of applying the PLSR model to new observations

We provide a simple R script (Methods S1) to illustrate the utility of our model and how it can be used to estimate LMA values from leaf spectral reflectance observations not used for model development. This script can be run to automatically download the foundational Leaf Optical Properties Experiment (LOPEX) and ANGERS spectral datasets (Hosgood *et al.*, 1994; Jacquemoud *et al.*, 2003), apply the PLSR model, and provide the results. The LMA model coefficients are provided through GitHub (https://github.com/serbinsh/SSerbin_etal_2019_NewPhytologist) and the leaf spectral data are provided from the Ecological Spectral Information System (EcoSIS) database (<https://ecosis.org>).

Furthermore, we also provide an extensive, external validation of our multibiome PLSR model with data collected in the Upper Midwest, US, and nine National Ecological Observatory Network (NEON) locations spanning seven NEON domains. The

first dataset consisted of fully expanded, peak-growing-season samples collected in and around the Madison, Wisconsin, area and the University of Notre Dame Environmental Research Center (UNDERC) between June and September 2017 (A. Chlus *et al.*, unpublished). A mix of sunlit and shaded foliar samples were collected from broadleaf trees ($n = 7446$), graminoids ($n = 74$), forbs ($n = 2017$) and vine ($n = 218$) species across the growing season. All samples were immediately scanned using a FieldSpec 3 (ASD Inc.), HR-1024i (SVC) and/or PSR+ (Spectral Evolution) with their respective leaf contact probes and external light sources. A Spectralon white reference was scanned before each sample to calculate the relative reflectance. After spectral measurement, the leaf area of each sample was immediately recorded using the LI-3100 benchtop leaf scanner (Li-Cor). Samples were subsequently flash-frozen in liquid nitrogen, freeze-dried in a VirTis lyophilizer (SP Scientific, Gardiner, NY, USA), and weighed with a precision balance.

The second dataset consisted of peak-growing-season foliar samples collected during the summer of 2017 at NEON sites in Wisconsin, Alabama, Georgia, Florida, Virginia, Maryland, Tennessee, Kansas and North Dakota, and also included trees ($n = 2584$), graminoids ($n = 381$) and forb ($n = 195$) species (Z. Wang *et al.*, unpublished). Spectra were collected immediately using the FieldSpec and/or PSR+ with their leaf contact probes. Leaves were scanned on a 600 dpi flatbed scanner (Epson, Nagano, Japan), oven-dried at 65°C for 48 h to a constant mass and weighed on a precision balance. The only processing applied to the spectra was the removal of spectral discontinuities (following Serbin *et al.*, 2012) at *c.* 1000 and 1900 nm where ‘jumps’ sometimes occur at overlapping wavelengths between detectors within the instruments. The final multibiome PLSR model was applied to all spectra and compared with laboratory measurements of LMA.

Data availability

All data used in this manuscript are publicly available through online data portals, including the U.S. Department of Energy NGEE-Arctic and NGEE-Tropics data portals as well as the EcoSIS spectral database (Table S1).

Results and Discussion

We employed an extensive dataset of leaf spectral reflectance and LMA across multiple biomes and spanning large environmental gradients (Fig. 1) to develop a generalized approach to model the variation in LMA (Fig. 2) using only leaf optical properties (Fig. 3). We found a very strong capacity for the empirical PLSR spectra–trait modeling approach to accurately model multibiome variation in LMA using spectral reflectance (Fig. 4). Our results show that spectra alone can explain 89% of the variation in LMA with a low model bias (0.96 g m^{-2}) and RMSE (15.45 g m^{-2}) when compared with our core validation dataset. We provide more details on our input datasets, PLSR modeling, and validation in the following, as well as a discussion of our work in the larger functional trait and remote-sensing research communities.

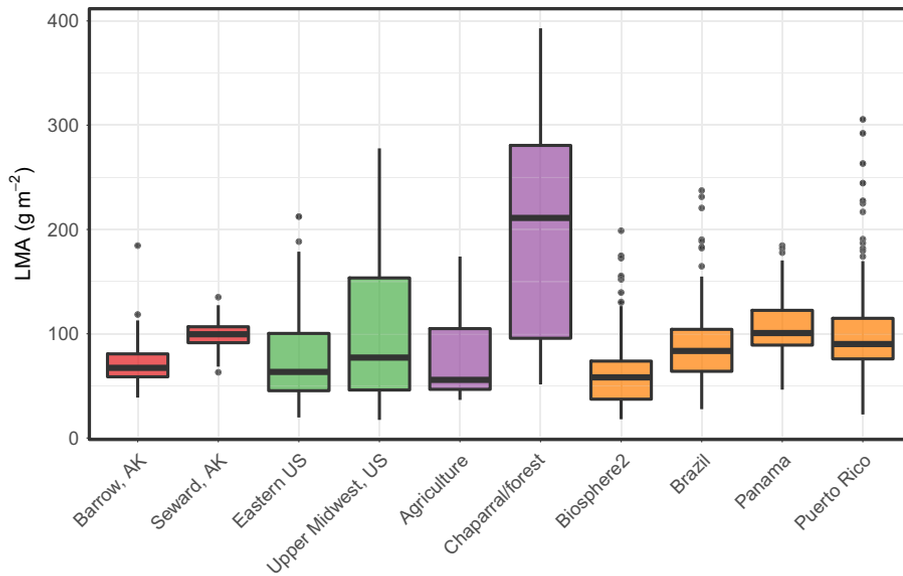


Fig. 2 Tukey box plots showing the range of leaf mass per area (LMA, g m^{-2}) in our model development dataset calculated from measurements of leaf area and dry mass for our 11 regions, including the Biosphere2 glasshouse location. Sites are color-coded by broad biome class (red, Arctic; green, boreal/temperate; purple, Mediterranean; orange, tropical). These are binned into four main biomes for clarity but correspond to the six Whittaker classes shown in Fig. 1. Box plots show the interquartile range (box), and median (solid horizontal line). The whiskers show lowest and highest datum still within $1.5\times$ interquartile range of the lower and upper quartiles. Outliers are shown as black dots. Sample sizes by region: 609 Arctic leaves, 935 boreal/temperate leaves, 102 Mediterranean leaves (including 33 agricultural samples), and 832 tropical leaves (including 72 Biosphere2 samples)

Biotic and abiotic variation in LMA

Across our sample sites (Fig. 1), we observed a broad range of LMA in our model development dataset, with values ranging from 17 to 393 g m^{-2} (Figs 2, S1). As expected, mean (\pm SD) top-of-canopy LMA of the needleleaf evergreen conifer species ($165 \pm 62 \text{ g m}^{-2}$) was significantly larger than those of broadleaf deciduous species, as well as grasses and forbs ($67 \pm 24 \text{ g m}^{-2}$). The LMAs of evergreen broadleaf species from arid environments ($169 \pm 72 \text{ g m}^{-2}$) were similar to those of needleleaf evergreen species, consistent with past observations (Paula & Pausas, 2006; de la Riva *et al.*, 2016), and owing to the well-documented differences in leaf life spans that typically require higher resource investment in leaf construction (Wright *et al.*, 2004; Poorter *et al.*, 2009). For the tropical species, LMA averaged $106 \pm 37 \text{ g m}^{-2}$ and ranged from 22 to 306 g m^{-2} .

Within a biome, LMA variation (Fig. S3) was large and related to biotic differences across species as well as factors such as leaf and plant age, and abiotic factors such as the leaf growth environment, including position in the canopy (Serbin *et al.*, 2014; Wu *et al.*, 2017), consistent with other studies (Niinemets *et al.*, 2015; John *et al.*, 2017; Osnas *et al.*, 2018). Notably, variation within species was often as large as that across species (data not shown), as has been noted in previous work (Butler *et al.*, 2017). On average, LMA in upper-canopy, fully sunlit leaves were $36 \pm 20\%$ higher than in fully shaded leaves of the same species, but this difference varied with growth form and leaf habit, probably owing to known differences in growth strategies, seasonality and the difference in the light penetration through a broadleaf or needleleaf canopy (Wright *et al.*, 2004; Ollinger, 2011; Butler *et al.*, 2017). Overall, the range in LMA within and across biomes showed similarities and overlap from the Arctic to the tropics, despite the vastly different leaf morphologies resulting from environmental and biological drivers, and these similarities were probably related to the underlying variation in leaf density and volume:area ratios, which can lead to similar values of LMA

with significantly different leaf geometries (Poorter *et al.*, 2009; John *et al.*, 2017).

Variation in leaf reflectance across biomes

As with LMA, we observed significant variation in the measured leaf-level spectral reflectance in our model development dataset within and across biomes (Figs 3a,b, S4). The biome-level spectral reflectance displayed a similar shape and comparable magnitude across the spectral range examined here (500–2400 nm; Fig. S4); however, the within-biome variation (presented as the coefficient of variation in reflectance by wavelength) showed some key differences (Fig. 3b) in the visible (i.e. 500–700 nm) and shortwave infrared (SWIR) regions (e.g. 1900–2500 nm). The biome-level mean reflectances appeared similar owing to the large variation in reflectance across wavelengths within each biome (Fig. S4), which mirrored the considerable within-biome variation in LMA (Figs 2, S3). This pattern has been observed in past research for similar vegetation types (Asner & Martin, 2008; Feret *et al.*, 2011; Yang *et al.*, 2016). Across the entire shortwave spectral region (i.e. 500–2400 nm), we also found that reflectance displayed the highest variation (Fig. 3b) in the visible region between about 500 and 700 nm (23–32% across biomes), in the SWIR region between 1300 and 1700 nm (15–17% across biomes), and in the far-SWIR region between 1900 and 2500 nm (24–37% across biomes). The near-infrared (NIR) region displayed only minor variation of *c.* 10% for all biome-level leaf spectra (Fig. 3b).

Exploring the connection between LMA and leaf spectral reflectance in the visible, NIR, and SWIR regions highlighted the strength and direction of the relationships between reflectance and LMA in different parts of the spectrum (Fig. S5). In the visible portion of the spectrum (Fig. S5a–c), the relationship between LMA and spectra is generally positive, as with the NIR (Fig. S5d) despite the much lower coefficient of variation in that region (Fig. 3b). There was a strong negative relationship

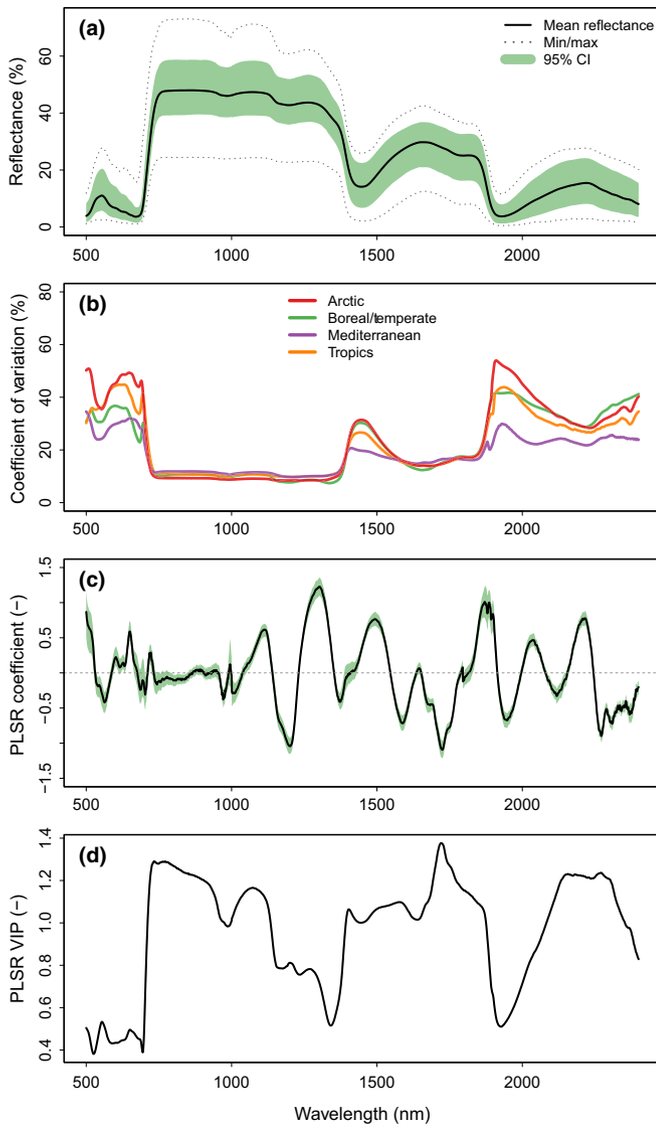


Fig. 3 Leaf reflectance and associated statistics. (a) Mean leaf reflectance, 95% confidence interval (green shading) and minimum and maximum reflectances (dotted lines) in our model development dataset. (b) Coefficient of variation for spectra from the four biomes represented in this study (red, Arctic; green, boreal/temperate; purple, Mediterranean; orange, tropical). (c) Plot of the partial least-squares regression (PLSR) model coefficients. (d) PLSR variable influence on projection (VIP).

between LMA and reflectance in the two SWIR (Fig. S5e,f) wavelengths (1800 and 2200 nm). Overall, these patterns are consistent with previous studies (le Maire *et al.*, 2008; Asner *et al.*, 2011; Feret *et al.*, 2011). For example, Feret *et al.* (2008) showed that spectral absorption of leaf dry matter content (the Cm or LMA parameter in the PROSPECT model) is generally near zero in the visible and NIR, but increases substantially when moving further out into the SWIR wavelengths. On the other hand, Slaton *et al.* (2001) illustrated the relationships of leaf morphology, thickness and other structural characteristics to changes in NIR reflectance across a range of leaf types and also found that there was a generally positive relationship between NIR reflectance (at 800 nm) and leaf structure parameters, including cuticle thickness and mesophyll properties.

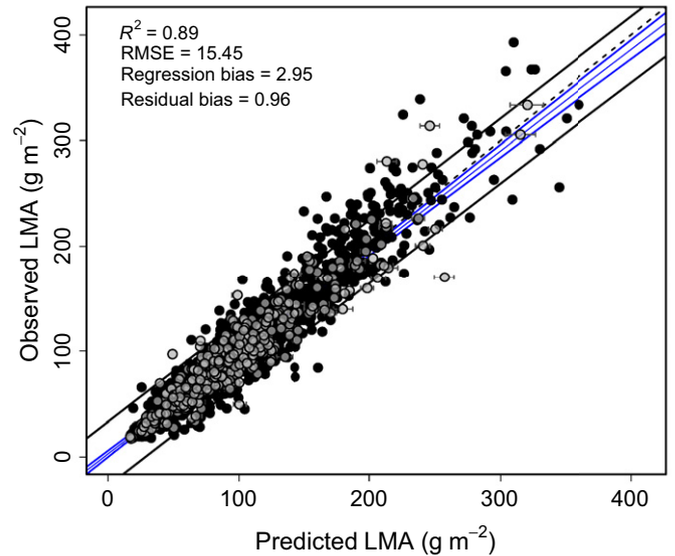


Fig. 4 Observed leaf mass per area (LMA) calculated from measured leaf area and dry mass vs LMA predicted using our spectral model. Our model development dataset ($n = 2478$) was split into two groups that were used to calibrate (black circles, $n = 1978$) and validate (gray circles, $n = 500$) the model. The validation points are shown with $\pm 95\%$ confidence interval (CI) error bars. For clarity, validation points are shown layered on top of calibration points. The 1 : 1 line is shown as a broken black line and the predictive interval of the model is shown as solid black lines. The regression between observed and predicted LMA is shown in blue (regression, thin blue line; 95% CI, thick blue line). The R^2 , root mean square error (RMSE) and regression bias (y-axis intercept) for the validation dataset are shown inside the panel. Data and statistics are presented in back-transformed standard LMA units.

Modeling the variation in LMA across biomes using reflectance spectroscopy

Using the observed variation in measured LMA (Figs 2, S1), leaf reflectance data (Figs 3a, S4), and the covariation between the two (Fig. S5) within our model development dataset, we then evaluated the capacity to build a widely applicable, multibiome model to estimate LMA from leaf spectral reflectance across the broad range of plant species, growth environments, leaf ages, measurement approaches, and spectrometer instrumentation reflected in our dataset. Our PLSR results (Figs 3c,d, 4) showed that a model based on leaf reflectance data was able to explain 89% of the variation in LMA in the independent validation data (and 91% in the calibration data using a cross-validation analysis; Fig. S6a). Our multibiome model also displayed a low overall independent validation error (RMSE = 16 g m⁻²) and bias (residual bias = 0.96 g m⁻², regression bias = 2.95 g m⁻²), with the model calibration and validation residuals both centered around zero (Fig. S6e,f). Within each biome in the core calibration and validation datasets, the model fit also showed low error, ranging from a minimum RMSE of 12 g m⁻² in the Arctic plants to 30 g m⁻² in the Mediterranean plants (Fig. S7). We further explored whether there were significant model biases attributable to variation with canopy position (Fig. S8) and across leaf ages (Fig. S9). Our results demonstrated that the model performance was very consistent across these different axes of variation,

although the leaf age evaluation displayed some bias in the young (expanding leaves or current year foliage) and mature (1 yr old) and mature (> 1 yr old) leaf age classes. Combined, the predictive error (dark black lines, 95% CI error bars on the points in Fig. 4) for the estimated LMA is small, particularly for the most commonly observed values of LMA (e.g. $LMA \leq 150 \text{ g m}^{-2}$; Fig. S1).

Our multibiome LMA PLSR model has a performance similar to or better than previous studies using leaves from a much smaller range of species, locations and growth environments (e.g. Asner *et al.*, 2011; Serbin *et al.*, 2014; Yang *et al.*, 2016). In addition, the model is able to cover a larger range of the global LMA trait-space (Figs 2, S1), habitable climates (Fig. 1) and variation in leaf optical properties (Fig. S4) than earlier work. This includes previous studies utilizing semimechanistic leaf RTMs, including the PROSPECT model (e.g. Feret *et al.*, 2008, 2011; Shiklomanov *et al.*, 2016). For example, Féret *et al.* (2018) found that performance of PROSPECT-D LMA inversion varied based on the input dataset (and species within), inversion approach and spectral domain, but that the overall PROSPECT-D inversion results across a smaller set of tropical to boreal samples were comparable to those shown here. Similarly, Shiklomanov *et al.* (2016) also found comparable results using a Bayesian inversion of PROSPECT-5b. However, in both cases, and with previous research (Malenovsky *et al.*, 2006), thicker leaves and needles have hindered the inversion accuracy of semimechanistic models like PROSPECT (e.g. Shiklomanov *et al.*, 2016) and, as a result, needleleaf species are typically removed before analysis (e.g. Féret *et al.*, 2018). Although approaches like PROSPECT are attractive to the remote-sensing and plant trait ecology communities, issues surrounding the handling of more complex leaf morphologies and absorption properties need to be addressed in order to facilitate the confident use of these models' global applications. On the other hand, our empirical approach was able to account for a broad range of morphologies and other drivers of leaf optical variation to produce a widely applicable multibiome model (Fig. 4).

The evaluation of the global model PLSR coefficient and VIP plots (Fig. 3c,d) highlights the regions of the spectrum that provide the strongest predictive power, many of which correspond to the areas showing the highest CV values, but note that the NIR region was also important. The SWIR spectral regions are known to contain absorption features related to structure, dry matter content, C compounds and internal leaf water content (Curran, 1989) which also covary with LMA (Elvidge, 1990; Poorter *et al.*, 2009; de la Riva *et al.*, 2016). Our global model showed relatively high VIPs in the NIR region (Fig. 3c,d), consistent with previous studies that also highlighted the importance of this region (Asner *et al.*, 2011; Yendrek *et al.*, 2017), particularly in the transition between visible and NIR reflectance (c. 750–800 nm) and the region between 950 and 1200 nm. Corresponding PLSR coefficient values in the NIR were highest in the c. 1000–1200 nm region. In general, the NIR has been shown to contain information connected to leaf internal scatter related to mesophyll layer thickness and water content, and varies with water, structural C, leaf thickness and variation in the epidermis layer (Ollinger, 2011), which also varies strongly with LMA

(Castro-Díez *et al.*, 2000; Jacquemoud *et al.*, 2009; Poorter *et al.*, 2009; de la Riva *et al.*, 2016).

Our results show the powerful capability of the spectral approach to estimate a key leaf trait (LMA) across a high diversity of plant species, growth environments, leaf ages, leaf morphology and biomes. Our synthesis represents the first time multiple datasets collected from different locations, by different groups, with different instrumentation across such large climatic and geographical ranges and from such a wide diversity of leaf types, including needleleaf species (Fig. 1), have been combined to test the ability to generalize the spectral PLSR modeling approach. Importantly, there appears to be a general pattern in the PLSR models shown for this and previous studies where similar portions of the visible, NIR and SWIR regions display high importance in the estimation of LMA with spectra (Asner *et al.*, 2011; Serbin *et al.*, 2014; Yang *et al.*, 2016). This strongly suggests that the coordination between leaf optical properties and traits can be used to develop generalized global models for leaf traits using the spectroscopic approach.

External validation of the multibiome LMA model

We further tested the predictive capability of our global leaf spectra–LMA model using three additional, completely independent datasets of measured leaf reflectance and LMA from trees, forbs, shrubs and grasses. These datasets include the foundational LOPEX and ANGERS spectral datasets (Hosgood *et al.*, 1994; Jacquemoud *et al.*, 2003), as well as data collected across NEON locations in the continental US. The first two datasets have been heavily used in remote-sensing literature to develop and test the PROSPECT model (Feret *et al.*, 2008). We applied our model coefficients (as shown in Fig. 3c) to the first two leaf reflectance datasets (i.e. LOPEX and ANGERS) and our results showed strong model performance ($R^2 = 0.66$) and low predictive error (RMSE = 21 g m^{-2} ; Fig. S10). Similarly, using the datasets collected in the Upper Midwest and at select NEON sites, we applied our model to the measured reflectance data to estimate LMA and found similar results (RMSE = 16.1 and 12.5 g m^{-2} for the Midwest and NEON locations, respectively; Fig. S11), suggesting the high model performance and generality of the spectra–trait approach for estimating LMA across a range of species and environments. As such, the development of globally applicable models like that presented here will reduce time and money spent developing site- or project-specific models to focus more on the rapid collection of much larger datasets by utilizing widely applicable models, like that presented here, to estimate traits such as LMA using only spectral measurements. The spectral approach has the added benefit of providing the capacity to repeat measurements of the same leaves during development or over a season, and during a manipulation or stress event, because it does not require destructive harvesting.

Main applications of our results

Our study of multibiome convergence in the leaf-level spectra–LMA relationships has four major implications. First, our

framework and approach can largely be extended to other leaf traits. In addition to LMA, some other plant functional traits (e.g. pigments (Chl a,b and carotenoid), chemical concentration (water%, N%, C%, isotopic N and C), and carboxylation capacity (V_{cmax}) are also very important inputs for ESMs, and critical measures of plant form and function (Xu *et al.*, 2012; van Bodegom *et al.*, 2014; Rogers, 2014; Butler *et al.*, 2017; Rogers *et al.*, 2017a; Ricciuto *et al.*, 2018). As these plant functional traits have been shown to be connected to leaf spectral reflectance (Asner & Martin, 2008; Feret *et al.*, 2008; Kokaly *et al.*, 2009; Serbin *et al.*, 2012, 2014, 2015; Wu *et al.*, 2019), we thus expect that similar globally convergent relationships with leaf spectra could be developed for these and other traits. To explore these additional generalities, future work can leverage emerging leaf spectra and trait databases, such as EcoSIS, which will facilitate much faster and easier development, testing and refinement of spectra–trait models.

Second, our work highlights the feasibility of using vegetation spectroscopy to advance large-scale monitoring of plant functional traits. This study evaluated the power of the spectroscopy approach at the leaf level, but an important next step is to explore the ability to use a range of imaging spectroscopy platforms to map traits, such as LMA, at the canopy and landscape scales using airborne and spaceborne instruments across larger regions than currently explored, and ultimately globally (Schimel *et al.*, 2015; Stavros *et al.*, 2017). Here we show the capacity to generalize the spectral approach at the leaf scale and there is strong evidence that similar approaches can be used at the canopy scale with imaging systems (e.g. Asner *et al.*, 2015; Singh *et al.*, 2015). This could augment or supplant the need for direct sampling and measurement of leaf traits at the global scale, especially given important upcoming satellite missions (e.g. Surface Biology and Geology Mission (National Academies of Sciences, Engineering, and Medicine, 2018), previously named *HyspIRI* (Lee *et al.*, 2015), and EnMAP (Guanter *et al.*, 2015)). Furthermore, functional trait mapping from imaging spectroscopy could supplement methods using remote sensing combined with climatology (Butler *et al.*, 2017; Moreno-Martínez *et al.*, 2018). These capabilities would significantly enhance the use of trait observations to inform ESMs and would address critical needs in biodiversity monitoring (Skidmore *et al.*, 2015; Jetz *et al.*, 2016). However, lack of global coverage, inconsistent processing workflows and other challenges with the use of imaging spectroscopy have limited the ability to derive consistent global trait maps. Furthermore, approaches are needed to separate trait retrieval from climatology to allow for the characterization of biotic and abiotic drivers, and for future mapping under novel climates (Fisher *et al.*, 2015). This can only be realized with a spaceborne mapping imaging spectroscopy mission (National Academies of Sciences, 2018).

Third, we hypothesized that a single, multibiome leaf reflectance model of LMA could be developed using datasets across diverse growth environments. We supported this hypothesis by showing a global convergence in the spectra–LMA relationship across samples representing a large portion of the global

trait-space for LMA (Figs 4, S1), which, when applied to external datasets from new, independent locations (Figs S10, S11) showed similar model performance for estimating LMA. Importantly, as leaf traits, particularly LMA, display adaptations and acclimation to their growth environment (Poorter *et al.*, 2009; Osnas *et al.*, 2018), the success of a generalized spectra–LMA model shown in this study further suggests that the spectroscopic approach could be an important, nondestructive means to help quantify and understand how plant traits acclimate to climatic variability and global change through rapid collection of traits with spectra (Shiklomanov *et al.*, 2019). Moreover, while the one-time costs of the spectrometer instrumentation utilized in this work can be high, this cost is low compared with the overall costs associated with traditional measurements of LMA (e.g. labor and supply costs over the lifetime of a project) and provides a much more rapid means for the retrieval of LMA.

Finally, the resulting multi-biome LMA PLSR model (see Methods S1 and associated links within) developed here can be downloaded and used by the scientific community. As measurement of leaf spectral reflectance is the only input required, our model enables the rapid, accurate and nondestructive estimation of LMA and can therefore be used for a broad range of additional applications, such as monitoring plant response to an emerging stress or evaluating physiological traits of interest to breeders in high-throughput phenotyping experiments (Silva-Perez *et al.*, 2017; Yendrek *et al.*, 2017; Ely *et al.*, 2019). As a result, databases such as EcoSIS can now be mined for increased coverage of important plant traits by applying this and other spectra–trait models, potentially increasing the amount of available data for modeling and ecological research.

Conclusions

In this work, we present a detailed synthesis of an extensive dataset containing coincident measurements of full-spectrum (i.e. 0.3–2.5 μm) leaf reflectance and LMA across multiple biomes and spanning large environmental gradients (Fig. 1). The variation in our dataset covered a large portion of the global variation observed for LMA (Fig. S1) and similarly contained a high degree of variation in leaf reflectance spectra (e.g. Figs 3, S4). Despite this high degree of variation in LMA and leaf reflectance observed across biomes, species, environment, leaf age and canopy position, our spectra–trait approach was able to collapse this variation into a single, widely applicable model (Fig. 4) that can be used to estimate LMA rapidly and accurately using only reflectance data (Figs S10, S11). These results are important because they highlight that the spectral approach presented here can be used to develop algorithms using datasets from a range of instruments, groups and locations to develop generalized models. Furthermore, this work illustrates the utility of the spectral approach in providing rapid, relatively low-cost and nondestructive (when the leaves are accessible) measurements of key plant traits across diverse plant species. The use of models such as that presented here will help to rapidly expand trait databases in order to address the known bias in observational datasets (Schimel *et al.*, 2015).

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Author contributions

SPS, JW and AR conceived the study, and defined the scope and focus of the manuscript. SPS developed the spectroscopic model of LMA. KSE, JW, RM, BTW, AR, and SPS participated in the collection of data in the Arctic and tropical field sites (including the Biosphere 2 experiment). SPS, PAT, ELK, AC and ZW collected the datasets in California, the Upper Midwest, and at the NEON locations. SPS, JW and AR wrote the first draft of the manuscript. All authors contributed to the final version.

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Supporting Information

Additional Supporting Information may be found online in the Supporting Information section at the end of the article.

Fig. S1 Comparison of the leaf mass per area (LMA) distribution in this article vs that found within the TRY global trait database.

Fig. S2 Distributions of the untransformed leaf mass per area (LMA) calibration and validation datasets used for model development and testing.

Fig. S3 The coefficient of variation of the measured leaf mass per area (LMA) across the four main biomes included in this research.

Fig. S4 The patterns of leaf reflectance across the four main biomes.

Fig. S5 The relationship between leaf mass per area (LMA) and spectral reflectance within the model development dataset.

Fig. S6 Model calibration and validation results for the leaf mass per area (LMA) partial least-squares regression (PLSR) model.

Fig. S7 Partial least-squares regression (PLSR) model fit by biome.

Fig. S8 Combined calibration and validation partial least-squares regression (PLSR) model residuals by canopy position.

Fig. S9 Combined calibration and validation partial least-squares regression (PLSR) model residuals by leaf age.

Fig. S10 Evaluation of the leaf mass per area (LMA) partial least-squares regression (PLSR) model using the LOPEX and ANGERS foundational datasets.

Fig. S11 External validation of the leaf mass per area (LMA) partial least-squares regression (PLSR) model using data collected from NEON sites.

Methods S1 An example R script illustrating the use of the multibiome PLSR model to estimate leaf mass per area for

LOPEX and Angers datasets, which are provided through the EcoSIS spectral library.

Table S1 Leaf spectra and LMA data sources.

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