Seasonal variability of multiple leaf traits captured by leaf spectroscopy at two temperate deciduous forests

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ABSTRACT

Understanding the temporal patterns of leaf traits is critical in determining the seasonality and magnitude of terrestrial carbon, water, and energy fluxes. However, we lack robust and efficient ways to monitor the temporal dynamics of leaf traits. Here we assessed the potential of leaf spectroscopy to predict and monitor leaf traits across their entire life cycle at different forest sites and light environments (sunlit vs. shaded) using a weekly sampled dataset across the entire growing season at two temperate deciduous forests. The dataset includes field measured leaf-level directional-hemispherical reflectance/transmittance together with seven important leaf traits [total chlorophyll (chlorophyll a and b), carotenoids, mass-based nitrogen concentration (Nmass), mass-based carbon concentration (Cmass), and leaf mass per area (LMA)]. All leaf traits varied significantly throughout the growing season, and displayed trait-specific temporal patterns. We used a Partial Least Square Regression (PLSR) modeling approach to estimate leaf traits from spectra, and found that PLSR was able to capture the variability across time, sites, and light environments of all leaf traits investigated (R² = 0.6–0.8 for temporal variability; R² = 0.3–0.7 for cross-site variability; R² = 0.4–0.8 for variability from light environments). We also tested alternative field sampling designs and found that for most leaf traits, biweekly leaf sampling throughout the growing season enabled accurate characterization of the seasonal patterns. Compared with the estimation of foliar pigments, the performance of Nmass, Cmass, and LMA PLSR models improved more significantly with sampling frequency. Our results demonstrate that leaf spectra-trait relationships vary with time, and thus tracking the seasonality of leaf traits requires statistical models calibrated with data sampled throughout the growing season. Our results have broad implications for future research that use vegetation spectra to infer leaf traits at different growing stages.

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1. Introduction

Leaf traits are important indicators of plant physiology and critical components in numerous ecological processes (Kattge et al., 2011; Wright et al., 2004). For example, leaf chlorophyll concentration represents the light harvesting potential and is an indicator of photosynthetic activity (Ninemets, 2007; Laisk et al. 2009), while accessory pigments such as carotenoids protect leaves from damage when exposed to excessive sunlight (Demmig-Adams & Adams, 2000). Leaf mass per area (LMA) describes plants’ investment to leaves in terms of carbon and nutrients to optimize sunlight interception (Poorter, Niinemets, Poorter, Wright, & Villar, 2009). Carbon is one of the major elements in cellulose and lignin, which are used to build the cell walls of various leaf tissues (Koike, Asner, Ollinger, Martin, & Wessman, 2009). Nitrogen is the key element in both carbon fixation enzyme RuBisCO and chlorophyll (Evans, 1989a, 1989b), and thus plays an important role in modeling leaf and canopy photosynthesis (Bonan, Oleson, Fisher, Lasslop, & Reichstein, 2012). The aforementioned leaf traits, as well as the corresponding spectral properties, strongly depend on leaf developmental stages and light environments (Yang, Tang, & Mustard, 2014; Lewandowska & Jarvis, 1977; Poorter et al., 2009; Wilson, Baldocchi, & Hanson, 2000; Wu et al., 2016a). Thus, capturing the spatial and temporal variations of these leaf traits is important for understanding terrestrial ecosystem functioning (Schimel et al., 2015).

Despite the importance and increasing interests in the temporal and spatial variability of these (and many other) leaf traits, the capacity to monitor these traits over seasons has not progressed accordingly. Wet
chemical analysis of these leaf traits is considered to be the standard method, yet the destructive and time-consuming protocols do not allow for rapid and repeated sampling of some traits. On the other hand, field spectroscopy can augment traditional approaches, and allows for repeated sampling of the same leaves and thus tracking time-sensitive changes such as frost damage (Asner & Martin, 2008; Couture, Serbin, & Townsend, 2013; Serbin, Singh, McNeil, Kingdon, & Townsend, 2014). Although spectroscopic approaches are promising, many previous efforts have only focused on mature sunlit leaves (e.g., Asner & Vitousek, 2005; Ustin, Roberts, Gamon, Asner, & Green, 2004; Wicklein et al., 2012; but see Sims and Gamon (2002)) and have not explored the ability of leaf spectral properties to track the continuous and developmental changes of leaf traits throughout the growing season. The temporal dimension of the spectra-traits relationship has mostly focused on leaf chlorophyll concentration (Belanger, Miller, & Boyer, 1995; Dillen, de Beeck, Hufkens, Buonanduci, & Phillips, 2012; Shen, Chen, Zhu, & Tang, 2009; Zhang et al., 2007), while it is largely unknown for other important leaf traits like nitrogen, carbon, and LMA. Moreover, the availability of high temporal resolution (weekly) datasets on important leaf traits and spectra is limited. These data would be very useful for assessing the utility of leaf spectral properties (i.e. reflectance) for estimating the temporal variability of leaf traits, as well as scaling to broader regions and informing process modeling activities.

Leaf traits not only change with time, but also with the light environments, such as moving from sun-lit to shaded light conditions and the concomitant changes in micrometeorology which also affect leaf traits (Ellsworth & Reich, 1993; Niinemets, 2007; Wu et al., 2016b), as a consequence of underlying fundamental evolutionary and eco-physiological constraints (Terashima, Miyazawa, & Hanba, 2001). For example, shaded leaves display lower chlorophyll a to b ratio and higher LMA compared with sunlit leaves (Niinemets et al., 2007). This variation in the vertical domain can be as much as the trait variation across space (Serbin et al., 2014). As such, it is important to not only explore trait variation across sites but also as in the vertical canopy light gradients to better capture ecosystem responses to global change.

Three categories of methods to estimate leaf traits from leaf spectral properties (i.e., reflectance and transmittance) are spectral vegetation indices (SVIs), statistical inversion methods exploiting the full wavelength (400–2500 nm), and leaf radiative transfer models like PROSPECT (Jacquemoud & Baret, 1990), which are limited to only a few leaf traits (do not include carbon and nitrogen) and thus are not the focus of this study. SVIs are typically calculated using the reflectance from two or three wavelengths (Huete et al., 2002; Richardson, Duigan, & Berlyn, 2002; Sims & Gamon, 2002). With proper calibration across a diverse range of vegetation types, SVIs can yield relatively robust models (Féret et al., 2011). Statistical methods such as Partial Least Square Regression (PLSR) modeling have become more popular in recent decades with the availability of high spectral resolution observations and increasing computational power (Asner & Martin, 2008; Couture et al., 2013; Wold, Sjöström, & Eriksson, 2001). Although both being widely used, these methods have not been thoroughly assessed, especially with respect to the robustness of PLSR models across time and different light environments (but see Serbin et al., 2014).

Here our primary goal was to assess the ability of leaf optical properties to track temporal variability of a suite of leaf traits across sites and different light environments. To explore this we collected a dataset of weekly-sampled leaf traits (including total chlorophyll (and chlorophyll a and b), carotenoids, mass-based nitrogen concentration (Nmass), mass-based carbon concentration (Cmass), and LMA) along with in situ directional-hemispherical reflectance/transmittance during the growing season at two temperate deciduous forests. We first present the temporal variations of leaf traits and spectra, and then highlight the ability of leaf spectra to track temporal variability of leaf traits. We investigate the robustness of the PLSR across season, sites, and light environments. We further explore the optimal field sampling strategy. We conclude by discussing the broad implications of our study.

2. Study area and methods

2.1. Study sites

Our field sampling was conducted in two temperate deciduous forests located in the northeastern United States. The first site, on the island of Martha’s Vineyard (MV, 41.362N, 70.578W), is a white oak (Quercus alba) dominated forest with a stand age of 80–115 years after natural recovery from abandoned cropland and pasture (Foster, Hall, Barry, Clayden, & Parshall, 2002). Mean annual temperature is 10 °C, and annual precipitation is about 1200 mm from 1981 to 2010 (Yang et al., 2014). The second site, in Harvard Forest (HF, 42.538N, 72.171W), has two dominating deciduous tree species: red oak (Quercus rubra) and red maple (Acer rubrum), with a few scattered yellow birch (Betula alleghenensis). The forest age is 70–100 years. The annual mean temperature is about 7.5 °C (Wofsy et al., 1993), and the annual precipitation is 1200 mm. Remote sensing studies suggested that the start of season in Martha’s Vineyard was about 10–20 days later than that of HF (Fisher & Mustard, 2007; Yang, Mustard, Tang, & Xu, 2012).

2.2. Measurements of leaf spectral properties and traits

We conducted two field campaigns to collect leaf traits at the sites in Martha’s Vineyard and Harvard Forest, respectively. In 2011, weekly (biweekly in August) sampling of leaves throughout the growing season (June–November) was conducted at the Martha’s Vineyard site on three white oak trees. For each sampling period, we cut two fully sunlit branches (each having –6 leaves) and one shaded branch using a tree pruner. The spectral properties of the leaves were immediately measured (see below). Then the leaves were placed in a plastic bag containing a moist paper towel, and all the samples were kept in a cooler filled with ice until being transferred back to the lab for further measurements. In 2012, the same weekly (biweekly from mid-July to late August) measurements in Harvard Forest were made on five individuals (two red oaks, two red maples and one yellow birch) from May to October. For each tree, two sunlit and one shaded branch were collected each time.

Directional-hemispherical leaf reflectance and transmittance were measured immediately after the sampling using a spectroradiometer (ASD FS-3, ASD Inc. Boulder, CO, USA; spectral range: 300–2500 nm, spectral resolution: 3 nm@700 nm, 10 nm@1400/2100 nm) and an integrating sphere (ASD Inc.). The intensity of light source in the integrating sphere decreases sharply beyond 2200 nm, with the signal in 2200–2500 nm being noisy (ASD Inc., personal communications), and thus is excluded from the spectral-leaf traits analysis below.

The measured leaf traits include total chlorophyll concentration (including chlorophyll a and chlorophyll b, μg/cm²), carotenoids (μg/cm²), leaf mass per area (LMA, g/m²), nitrogen concentration by mass (Nmass %), and carbon concentration by mass (Cmass %). Each branch was divided into two subsets. One subset was used to measure pigment concentrations. To measure the chlorophyll and carotenoids concentration, three leaf discs (~0.28 cm² each) were taken from each leaf using a hole puncher, and then ground in a mortar with 100% acetone solution and MgO (Asner, Martin, Ford, Metcalfe, & Liddell, 2009). After an 8-minute centrifugation, the absorbance of the supernatant was measured (see below). Then the leaves were placed in a plastic bag containing a moist paper towel, and all the samples were kept in a cooler filled with ice until being transferred back to the lab for further measurements. In 2012, the same weekly (biweekly from mid-July to late August) measurements in Harvard Forest were made on five individuals (two red oaks, two red maples and one yellow birch) from May to October. For each tree, two sunlit and one shaded branch were collected each time.

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\[ LMA = \frac{W_{dry}}{A_{leaf}} \]
Table 1
Simple vegetation indices (SVI) used in this study. These indices were calibrated using extensive datasets (Féret et al., 2011). Leaf traits were calculated based on a polynomial relationship: leaf trait = a × index^2 + b × index + c.

<table>
<thead>
<tr>
<th>Leaf trait</th>
<th>Index</th>
<th>Coefficients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chl (µg/cm²)</td>
<td>(R900 - R1212)/(R900 + R1212)</td>
<td>40.65 121.88 -0.77</td>
</tr>
<tr>
<td>Car (µg/cm²)</td>
<td>(R900 - R350)/(R900 + R350)</td>
<td>8.09 11.18 -0.38</td>
</tr>
<tr>
<td>LMA (g/cm²)</td>
<td>(R1358 - R1722)/(R1358 + R1722)</td>
<td>-0.1004 0.1286 -0.0044</td>
</tr>
</tbody>
</table>

where \( W_{dry} \) is leaf dry mass weight, \( A_{leaf} \) is the leaf area calculated from the scanned leaf using ImageJ (Schneider, Rasband, & Eliceiri, 2012). Dried leaves were then ground and analyzed for \( N_{mass} \) and \( C_{mass} \) with a CHNS/O analyzer (FLASH 2000, Thermo Scientific, Waltham, MA, USA).

2.3. Methods to estimate leaf traits using leaf spectral properties

We used two categories of methods to estimate leaf traits based on leaf spectral properties: vegetation indices that utilize the reflectance from two wavelengths, and statistical methods that exploit the information from the full leaf spectrum.

Based on extensive datasets from various types of biomes and plants, Féret et al. (2011) established polynomial relationships between SVIs and total chlorophyll concentration, carotenoids, and LMA (Table 1). We also obtained the best estimate of \( a, b, \) and \( c \) using our own dataset (see below for the division between training and validation dataset).

The second category of methods essentially is to build multivariate linear regression models between leaf spectra and leaf traits (Zhao, Valle, Popescu, Zhang, & Mallick, 2013):

\[
y = X \beta + \epsilon
\]

where \( y \) is an \( n \)-by-1 matrix of leaf traits (\( n \) equals to the number of leaf samples), \( X \) is an \( n \)-by-\( m \) matrix (\( m \) equals the number of bands from each spectrum, and thus in this study \( m = 1801 \): from 400 to 2200 nm), \( \epsilon \) is the \( n \)-by-1 estimation error that is to be minimized. PLSR modeling can be used to develop the best model for the given dataset while avoid over-fitting (Asner & Martin, 2008; Serbin et al., 2014). The numbers of independent factors used in the regression were determined by minimizing the Prediction Residual Error Sum of Squares (PRESS).

The above leaf traits and spectra from two sites were combined as one single dataset. To test the effectiveness of PLSR on this dataset, the whole dataset is divided into two parts (70%–30%), for the training and validation of PLSR, respectively. We used the Kennard-Stone algorithm to select the training subset that provides a uniform coverage of the whole dataset (Kennard & Stone, 1969). The training dataset was used to optimize the regression model parameters (\( \beta \)), and then the validation dataset was used to test and evaluate the PLSR models. Evaluation statistics include the \( R^2 \), Root Mean Square Error (RMSE) and normalized RMSE (NRMSE), which is the RMSE divided by the range of the estimated leaf traits.

The relative importance of reflectance or transmittance at each wavelength is determined by calculating the values of variable importance on projection (VIP) (Wold et al., 2001). VIP is an indicator of the importance of each wavelength for the modeling of both leaf traits (\( y \)) and spectra (\( X \)). Higher absolute values indicate greater importance of the corresponding wavelength. Generally, wavelengths with VIP value larger than 1 are considered being important (Mehmood, Liland, Snipen, & Sæbø, 2012).

2.4. Robustness of PLSR models and scenarios for field sampling design

To examine the robustness of PLSR models across time, light environment, and sites, we designed the following scenarios. In all scenarios, we used leaf traits and spectra of a subset of the whole dataset (e.g., leaf samples that are collected during only a certain period of time, or a certain level of light environment) to train PLSR models, and tested the performance of the models against the whole dataset.

For this we created five scenarios to examine how the timing of leaf sampling affects predictability of seasonality of leaf traits. Leaf traits and spectra in the first three scenarios were sampled only for the spring, summer, and fall, respectively. We defined these three seasons based on variations in total chlorophyll concentration: days before total chlorophyll reached a plateau in the mid-season were defined as spring; days when total chlorophyll started to decrease were defined as fall; and between spring and fall, days were defined as summer. The last two scenarios were that leaf traits and spectra were sampled monthly or biweekly (instead of weekly as in the full dataset). We then use the PLSR trained with leaf samples in the above scenarios to predict the leaf traits of the entire dataset. There are two reasons to choose the whole dataset for validation: 1) the whole dataset captures the temporal variability of leaf traits, which is the goal of this test and 2) it is necessary to have the same validation dataset to compare the performance of these five scenarios. Performance of these sampling strategies was measured by calculating the RMSE and \( R^2 \).

Finally, we also explored our capacity to develop a generalized approach for capturing seasonality in leaf traits with spectral observations. Two tests were conducted to examine the robustness of PLSR models at different light environment and sites. Test 1 used sunlit leaf traits and spectra to train a PLSR model, which was then used to predict shaded leaf traits with corresponding spectra. We then switched the training and validation datasets so that shaded leaves were used to train PLSR model while sunlit leaves were used to validate. Test 2 divided the entire dataset into two subsets by geographic location. For example, we used Martha’s Vineyard dataset to calibrate the model, and Harvard Forest dataset to validate, and vice versa.

3. Results

3.1. Temporal and spatial variability of leaf traits

All leaf traits displayed significant temporal variations throughout the growing seasons (Figs.1a–2). Overall, pigments from both sites displayed similar bell-shaped trajectories, despite being sampled from different species and locations within the canopy. Chlorophyll concentration rapidly increased from ~10 µg/cm² at the beginning of the season, and then stabilized around ~50 µg/cm² and ~40 µg/cm² in Martha’s Vineyard and Harvard Forest, respectively, during the summer followed by a decline in the fall to values similar to the beginning of the season prior to leaf abscission. The Harvard Forest samples were collected from three different species, and showed much larger variability compared with Martha’s Vineyard, especially for the shaded leaves (Fig.1e–h). The carotenoids concentration was ~3 µg/cm² at the beginning/end of the season and ~10 µg/cm² at the peak season. The total chlorophyll concentration relative to the carotenoids concentration (Chl/Car) increased during the early seasons. In the fall, though both chlorophyll and carotenoids decreased, Chl/Car decreased steadily, as a result of faster decline of chlorophyll relative to the carotenoids (Fig.5a).

The remaining three leaf traits (LMA, \( N_{mass} \), and \( C_{mass} \)) displayed different seasonal patterns compared with leaf pigments (Fig. 2). For example, LMA rapidly increased in the spring, but showed only a minor decline by the end of the measurement period. \( N_{mass} \) was higher (~4–5%) at the start of the season, and remained stable around 2% during the summer, followed by ~1% decrease in the fall, presumably caused by nitrogen resorption (Eckstein, Karlsson, & Weih, 1999). Similar to
LMA, $C_{mass}$ accumulated 2–4% in the spring and stabilized for the rest of the growing seasons around 50%. The rapid increase of LMA in the spring was accompanied by a similar increase of $C_{mass}$ and decrease of $N_{mass}$, which all ended at the same time (DOY ~194 in Martha’s Vineyard, and DOY ~170 in Harvard Forest).

Mean annual values of leaf traits from Martha’s Vineyard were significantly different from those at Harvard Forest (Table 2). For example, leaf chlorophyll in Martha’s Vineyard is 5.64 μg/cm² (17.5%) higher than that from Harvard Forest ($p < 0.0001$). LMA in Martha’s Vineyard showed much larger variation than that from Harvard Forest, and the

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**Fig. 1.** Seasonal patterns of pigments of sunlit (filled) and shaded (open) leaves from two deciduous forests. Martha’s Vineyard, year 2011: (a) Total chlorophyll; (b) chlorophyll $a$; (c) chlorophyll $b$; (d) carotenoids. Harvard Forest year 2012: (e) Total chlorophyll; (f) chlorophyll $a$; (g) chlorophyll $b$; (h) carotenoids. Each dot is the mean value of all the samples collected that day. Error bars are standard deviations.
mean LMA was 39.85 g/m² (37.5%) higher than that from Harvard Forest. Similar situation applies to all other leaf traits except for Cmass, for which the value at Harvard Forest were higher than that at Martha’s Vineyard.

Sunlit leaves contained more total chlorophyll and carotenoids (Fig. S2) and the carotenoids to the total chlorophyll ratio was significantly higher for sun-lit leaves comparing with shaded leaves (MV, \(p < 0.0001\); HF, \(p = 0.0182\)). Chlorophyll a/b was also significantly larger for sunlit leaves in both sites (MV, \(p < 0.0001\); HF, \(p < 0.0001\)). Similarly, LMA and Cmass values were significantly higher in the sun-lit leaves versus shaded foliage, with the only exception of Nmass, in which both sun-lit and shaded leaves were indistinguishable throughout the two seasons (Fig. 2b).

A linear regression analysis highlighted various levels of correlation among leaf traits (Fig. 3). Close correlation was found among leaf pigments: total chlorophyll concentration was highly correlated with carotenoids concentration (\(R^2 = 0.85\)), suggesting a tight coupling among those pigments throughout the growing season despite the faster decrease of chlorophyll concentration during the senescence (Fig. S1). For the entire dataset (across all sunlit and shaded leaves from different species), Nmass was weakly correlated with pigments. LMA showed positive correlation with all pigments while a negative correlation was observed with Nmass and Cmass.

### 3.2. Seasonal variability of leaf spectral properties

The full leaf reflectance and transmittance spectrum showed significant variability in both amplitude and shape (Fig. 4). The visible (VIS, 400–700 nm) and near infrared (NIR, 700–1000 nm) changed dramatically throughout the season, while shortwave infrared (SWIR 1000–2500 nm) was relatively stable. Data from Martha’s Vineyard showed larger variability in NIR compared to Harvard Forest.

The R, G, and B reflectance at both sites showed a U-shape pattern (Fig. S3a, S3c): all of them decreased in the beginning of the season; and increased in the end of the season after a stable summer. The NIR from Martha’s Vineyard showed a consistent decline in the mid-
summer and then increased in the fall, while the NIR from Harvard Forest was relatively stable throughout the season. Leaf transmittance at each band had similar patterns as the reflectance (Fig. S3b, S3d).

3.3. Comparisons of methods of leaf traits estimation

We compared two categories of methods to estimate leaf traits from leaf spectra. Overall, PLSR consistently outperformed the SVIs in estimating leaf traits, showing an improved performance when the SVIs were trained by the original datasets or our own dataset (Table 3). The PLSR models using leaf reflectance (PLSR_ref hereafter) had slightly better performance compared with those using leaf transmittance (PLSR_tra hereafter) when assessed with the independent dataset. For different leaf traits, the performance of these methods varied, as described in details below.

Leaf chlorophyll from the validation dataset was well estimated by PLSR_ref (Fig. 5. $R^2 \geq 0.70$ and NRMSE $< 10\%$). The SVI for chlorophyll showed slightly larger prediction error (~0.5 μg/cm² larger) compared

![Correlation Matrix](image1)

**Fig. 3.** Correlation matrix of all the leaf traits. Histograms of each leaf trait are on the diagonal positions. Number on each subplot indicates $R^2$ (Red means $p < 0.05$). See Table 2 for units.

![Examples of leaf directional-hemispherical reflectance and transmittance](image2)

**Fig. 4.** Examples of leaf directional-hemispherical reflectance and transmittance measured on (a, b) Martha’s Vineyard and in (c, d) Harvard Forest.
with PLSR\textsubscript{ref} and PLSR\textsubscript{tra} (Table 3). The two components of chlorophyll (Chl \(a\) and \(b\)) were also well captured by the PLSR\textsubscript{ref} approach with \(\text{NRMSE} < 10\%\) and \(\text{R}^2 > 0.63\) and \(0.66\) respectively. Similarly, carotenoids were estimated relatively well by PLSR\textsubscript{ref} and PLSR\textsubscript{tra} (\(R^2 > 0.65\)) but the SVI for carotenoids had \(30\%\) higher RMSE compared with PLSR\textsubscript{ref}.

\(N_{\text{mass}}\) was well captured by leaf spectra especially with the reflectance dataset (Fig. 5). \(R^2 > 0.6\) and \(\text{NRMSE} < 5\%\). Similarly, both PLSR\textsubscript{ref} and PLSR\textsubscript{tra} explained \(60\%\) of the variance in \(C_{\text{mass}}\) (\(R^2 > 0.6\) and \(\text{NRMSE} < 7\%\)). PLSR also displayed a strong capacity to predict LMA (\(R^2 = 0.80\) and \(\text{NRMSE} < 9\%\)), where the SVI for LMA could not capture \(>20\%\) of the variation in LMA and had more than double the RMSE of PLSR\textsubscript{ref} mainly due to a saturation effect (data not shown).

The VIP values of PLSR show the relative importance of each wavelength in predicting leaf traits (Fig. 6). Visible and near-infrared wavelengths were important to the prediction of leaf pigments; there are three peaks (400, 550 and 730 nm) that are related to the chlorophyll absorption in the red (620–750 nm) and blue (400–450 nm), and reflection in the green (495–570 nm). The two components of chlorophyll (\(a\) and \(b\)) were mainly contributing to the red/NIR region (600–750 nm), and the main contributing bands for chl \(b\) shifted towards green comparing to those for chl \(a\) (Fig. 6b and 6c) (Ustin et al., 2009). Carotenoids have a similar VIP curve comparing with the chlorophyll, with one distinction: the VIP values for carotenoids between 650 nm and 700 nm are relatively higher to those of chlorophyll.

Compared with the pigments, \(N_{\text{mass}}, C_{\text{mass}}\) and LMA have relatively smooth VIP curves. For \(N_{\text{mass}},\) wavelengths around 700 nm and beyond 1900 nm are important to the prediction of \(N_{\text{mass}}\) presumably because the visible region is controlled by pigments and nitrogen is an important component in leaf pigments, and the SWIR region near 2000 nm is controlled by protein absorption features (Kokaly et al., 2009). Both \(C_{\text{mass}}\) and LMA were related to the leaf structure and were largely contributing to the reflectance at NIR and SWIR.

Table 3

<table>
<thead>
<tr>
<th>Leaf traits</th>
<th>RMSE (NRMSE)</th>
<th>(R^2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(N_{\text{mass}}) ((\mu g/cm^2))</td>
<td>3.93</td>
<td>0.71</td>
</tr>
<tr>
<td>Chl (a) ((\mu g/cm^2))</td>
<td>3.99</td>
<td>0.71</td>
</tr>
<tr>
<td>Chl (b) ((\mu g/cm^2))</td>
<td>1.62</td>
<td>0.73</td>
</tr>
<tr>
<td>Car ((\mu g/cm^2))</td>
<td>1.07</td>
<td>0.73</td>
</tr>
<tr>
<td>(N_{\text{mass}}) (%)</td>
<td>0.22</td>
<td>0.73</td>
</tr>
<tr>
<td>(C_{\text{mass}}) (%)</td>
<td>0.24</td>
<td>0.64</td>
</tr>
<tr>
<td>LMA ((g/cm^2))</td>
<td>18.11</td>
<td>0.64</td>
</tr>
</tbody>
</table>

3.4. Robustness of the PLSR approach across time, sites and light environment

We examined the performance of the PLSR\textsubscript{ref} models under five scenarios where different sampling strategies were applied. The performance of the PLSR models generally improved in the order of spring, fall, summer, monthly and biweekly (Table 4). As expected, more sampling throughout the season (and the increasing size and representativeness of the calibration dataset) increased \(R^2\) and reduced RMSE. When comparing the three seasons, summer-only sampling yielded higher model performance relative to the other two scenarios, yet the improvements from scenarios 2 (summer-only) to scenario 4 (monthly) were not as obvious for pigments as \(N_{\text{mass}}, C_{\text{mass}}\) and LMA. Sampling scenario 5 (biweekly) largely improved the performance of PLSR, especially for \(N_{\text{mass}}\) and \(C_{\text{mass}}\) (\(R^2\) increased from \(<0.4\) to \(>0.6\)).

Examining the seasonal patterns of predicted and observed leaf traits revealed time-dependent performance of each scenario. In spring-only scenario where leaf samples only from the spring were used for PLSR calibration, all leaf traits during the first four weeks of the growing seasons were well estimated. However, fall season leaf traits were overestimated except for LMA in Martha’s Vineyard (Fig. 5am). By contrast, in the fall-only scenario, spring and summer leaf traits were underestimated except for \(C_{\text{mass}}\) (Fig. 5ak). Summeronly scenario showed a better ability to capture the seasonal patterns of leaf traits, only underestimated the \(N_{\text{mass}}\) peak in the early spring at Harvard Forest (Fig. 5ji). The monthly sampling scenario improved estimation of all leaf traits, in which the improvement on estimating LMA was the most obvious (\(R^2\) from 0.26 in the summer case to 0.76 in the monthly sampling case, Fig. 5mj, 5nj). Biweekly sampling scenario appeared to produce a satisfactory result for all the leaf traits studied here (Fig. 5j).

PLSR\textsubscript{ref} models trained using sunlit leaves explain 35–70% of the variability in shaded leaves with highest \(R^2\) for pigments while lowest \(R^2\) for \(C_{\text{mass}}\) (Fig. 5s, Table 1). However, PLSR\textsubscript{ref} was less accurate for leaf traits like LMA in terms of RMSE (Fig. 5l m), for which the difference between sun-lit and shaded leaves was significant (Fig. 2). Similarly, PLSR\textsubscript{ref} models trained with shaded leaves were able to predict the sunlit leaf traits, but with lower model performance compared to when trained with sunlit foliage. Depending on the leaf traits, the variability explained by PLSR ranges from 35% to 70% (Fig. 5lm).

PLSR\textsubscript{ref} models trained using data from Harvard Forest (Test 1) were able to capture 60–70% of variability of the pigments from Martha’s Vineyard, except for \(N_{\text{mass}}\) and \(C_{\text{mass}}\) (Table 5). Similar results were obtained from PLSR\textsubscript{ref} trained using Martha’s Vineyard data (Test 2) and validated with Harvard Forest data. VIP values for pigments in Test 1 were similar to those from Test 2. This is in stark contrast with VIP values for Nmass, Cmass, and LMA from both experiments. The locations of important wavelengths were quite different between two tests (Fig. 5l).

4. Discussion

4.1. Can we track the seasonality of leaf traits using leaf spectroscopy?

Here we show that the seasonal variability of leaf traits can be well captured with leaf spectroscopy approaches (Fig. 5, Table 3). All leaf properties (seven leaf traits and leaf spectra) display seasonal dynamics that are also related to the location and microclimate (i.e., sunlit vs. shaded, and the accompanying changes in humidity and temperature). The PLSR models explain 60–80% of variability of these leaf traits in our study, supporting the hypothesis that leaf spectra can capture the seasonal variability of leaf traits. Indeed, each leaf trait has its own spectral fingerprint (Curran, 1989; Kokaly et al., 2009), as we have seen from the VIP values of PLSR models (Fig. 6). Patterns of VIP values were similar to previous studies (Asner et al., 2009; Serbin et al., 2014) and consistent with our understandings of leaf physiology (Ustin et al., 2009). This is an important result as collecting leaf spectra is much more time-efficient than traditional approaches and allows for repeated sampling of the same leaves throughout the season. SVIs can be an alternative
Fig. 5. Comparisons between the observed leaf traits and predicted traits from PLSRref. For detailed statistics, refer to Tables 2 and 3. Observations are from the independent validation dataset selected using the Kennard-Stone method. The red dashed lines are 1:1 line.
for the estimation of total chlorophyll concentration when there are limits on available instruments or, for example, using two-band LED sensors (e.g., Garrity, Vierling, & Bickford, 2010; Ryu et al., 2010). The result also has implications for the current and future use of field spectrometers that measure leaf or canopy reflectance at high temporal frequency (e.g., Hilker et al., 2009). Our well-calibrated PLSR models can be used on leaf reflectance to track the seasonality of multiple leaf traits in temperate deciduous forests.

The tests on the robustness of leaf spectra-trait relationships suggested that the overlap between the training dataset and an independent validation dataset is important for a good prediction. Traditionally, the development of the leaf traits-spectra relationship has been focused on a single time point, typically mid-season and mature leaves. Summer mature leaves displayed higher pigments concentration and LMA, while lower Nmass compared with young leaves (Fig. 1, Fig. 2). In addition, the corresponding leaf spectra were significantly different (Fig. 4). We have shown here that if we apply an empirical relationship between spectra and traits derived from one period (for example, summer) to another (spring or fall), leaf traits will likely be over or under-estimated (Fig. S4–S6). These findings mirror that observed by McKown, Guy, Azam, Drewes, and Quamme, 2013 which used traditional trait measurements across key phenophases of a temperate forest species, *Populus trichocarpa*, to show that the direction and magnitude of many trait-trait relationships is strongly tied to phenological state and can change over a season. However, we have also illustrated that with proper calibration, we can adequately characterize the seasonality of a range of leaf traits, despite the impacts of phenology, which is critical for monitoring ecosystems and informing large-scale modeling activities (Table 5).

VIP values can help to explain the prediction power of PLSR models. For example, in the case of PLSR models trained with data from one site to predict another (Tests 1 & 2), VIP values of leaf pigments overlap different (Fig. 4).

### Table 4

Performance of all scenarios (spring, summer, fall, monthly, and biweekly) in terms of the goodness-of-fit (RMSE, $R^2$).

<table>
<thead>
<tr>
<th>Leaf traits</th>
<th>RMSE</th>
<th>$R^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Chl ($\mu g/cm^2$)</td>
<td>Spring</td>
<td>Summer</td>
</tr>
<tr>
<td></td>
<td>8.64</td>
<td>7.23</td>
</tr>
<tr>
<td>Chl a ($\mu g/cm^2$)</td>
<td>5.97</td>
<td>5.25</td>
</tr>
<tr>
<td>Chl b ($\mu g/cm^2$)</td>
<td>2.73</td>
<td>2.06</td>
</tr>
<tr>
<td>Car ($\mu g/cm^2$)</td>
<td>1.71</td>
<td>1.29</td>
</tr>
<tr>
<td>Nmass (%)</td>
<td>1.62</td>
<td>0.51</td>
</tr>
<tr>
<td>Cmass (%)</td>
<td>1.59</td>
<td>1.26</td>
</tr>
<tr>
<td>LMA (g/cm$^2$)</td>
<td>61.13</td>
<td>24.86</td>
</tr>
</tbody>
</table>
well, indicating both sites share similar wavelength regions (Fig. S11). As a result, cross-site prediction of leaf pigments showed reasonable accuracy (Table 5). It also has important implications for the design of multi-band sensors and imagers as it can select the wavelengths that are most useful for the leaf traits of interest (Nijland et al., 2014; Ryu et al., 2010).

The variability of our seven leaf traits was not equally captured (Table 3). The absorption features of pigments are well understood and clearly represented in the VIP value plots (Fig. 6). While for \( C_{\text{mass}} \) and \( N_{\text{mass}} \), although there have been studies on the possible linkage between certain components in the leaves (e.g., protein, cellulose) and leaves’ optical properties, the impact on leaf spectra is less pronounced in any single spectral range as compared with pigments and instead is more generally spread across the visible and shortwave infrared spectral regions (Curran, 1989; Kokaly et al., 2009). This may partly explain the less accurate PLSR models for the \( C_{\text{mass}} \) and \( N_{\text{mass}} \). Moreover, in fresh leaves the spectral absorption by proteins can be partially obscured by water absorption, which can impact the performance of statistical (e.g., PLSR, SVIs) and other spectral inversion approaches, such as radiative transfer models (RTMs).

As expected, the PLSR approach, which can exploit the full spectrum information to estimate leaf traits performed better than traditional SVIs (Table 3). While SVIs that were calibrated with extensive datasets displayed a similar performance to PLSR in estimating total chlorophyll concentration, we observed better performance of PLSR for the \( \mu \text{g/cm}^2 \) and \( \mu \text{g/cm}^2 \). Performance of PLSR re

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### Table 5

<table>
<thead>
<tr>
<th>Leaf traits</th>
<th>RMSE</th>
<th>( R^2 )</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( \text{MV} \rightarrow \text{HF} )</td>
<td>( \text{HF} \rightarrow \text{MV} )</td>
</tr>
<tr>
<td>Total Chl (( \mu \text{g/cm}^2 ))</td>
<td>6.17</td>
<td>7.44</td>
</tr>
<tr>
<td>Chl ( a ) (( \mu \text{g/cm}^2 ))</td>
<td>4.39</td>
<td>5.29</td>
</tr>
<tr>
<td>Chl ( b ) (( \mu \text{g/cm}^2 ))</td>
<td>1.85</td>
<td>1.99</td>
</tr>
<tr>
<td>Car (( \mu \text{g/cm}^2 ))</td>
<td>1.19</td>
<td>1.54</td>
</tr>
<tr>
<td>( N_{\text{mass}} ) (%)</td>
<td>0.56</td>
<td>0.72</td>
</tr>
<tr>
<td>( C_{\text{mass}} ) (%)</td>
<td>2.89</td>
<td>2.90</td>
</tr>
<tr>
<td>LMA (( \mu \text{g/cm}^2 ))</td>
<td>35.62</td>
<td>59.45</td>
</tr>
</tbody>
</table>

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4.3. Broad implications of using leaf spectroscopy for ecological studies

Understanding the seasonality of leaf traits has recently gained attention as an effort to improve our modeling of terrestrial carbon and water cycles (Bauera et al., 2012; Grassi, Winnelli, Ponti, Cantoni, & Magnani, 2005; Medvigy, Jeong, Clark, Skowrons, & Schäfer, 2013). For example, in the Community Land Model, \( N_{\text{mass}} \) and LMA control the maximum rate of carboxylation, \( V_{\text{max}} \), which is highly variable temporally and across different species and light environments (Oleson, Lawrence, Bonan, et al., 2013). Our time-series of \( N_{\text{mass}} \) capture two important features: (1) the seasonal peak at the beginning of the spring, suggesting that nitrogen was allocated to the leaves early in the season. As leaves matured, other types of elements such as carbon accumulated at a faster rate, resulting in an increase in \( C_{\text{mass}} \) relative to \( N_{\text{mass}} \) ratio. (2) A decline of \( N_{\text{mass}} \) by the end of the season. \( N_{\text{mass}} \) and LMA was relatively stable at both sites during the summer (Fig. 2a and b), thus leaf age does not appear to be affecting the nitrogen concentration during the peak season (Field & Mooney, 1983). This finding as that of others (McKown et al., 2013; Wilson et al., 2000) highlights the importance of tracking the seasonality of leaf traits, and our work demonstrates that leaf spectroscopy can provide a rapid means to routinely measure leaf traits. Importantly, these results highlight that spectroscopy observations can provide key information on the individual differences in multiple leaf traits that can feed into ecosystem models (Medvigy, Wofsy, Munger, Hollinger, & Moorcroft, 2009) or be used to test key ecological hypotheses (Rowland et al., 2015). Our results suggest the important capability of monitoring ecosystem dynamics across a range of spatial and temporal scales with hyperspectral observations from leaves, towers, as well as with new instruments mounted on unmanned and piloted aircraft and satellite platforms (Asner & Martin, 2008; Hilker et al., 2010; Yang et al., 2014; Yang et al., 2015).
spectra to track seasonally varying leaf traits, and thus supports the use of automated field spectrometers, airborne and satellite hyperspectral sensors to track leaf traits repeatedly throughout the season and across broad regions (Roberts, Quattrochi, Hulley, Hook, & Green, 2012; Singh, Serbin, McNeil, Kingdon, & Townsend, 2015; Yang et al., 2015).

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Appendix A. Supplementary data

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.rse.2016.03.026.

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