RESEARCH PAPER

Plot-level rapid screening for photosynthetic parameters using proximal hyperspectral imaging

Katherine Meacham-Hensold1,2,*, Peng Fu2,*, Jin Wu3,4, Shawn Serbin3, Christopher M. Montes1, Elizabeth Ainsworth1,2,5, Kaiyu Guan6,7, Evan Dracup5, Taylor Pederson2, Steven Driever2,8 and Carl Bernacchi1,2,5,†

1 Department of Plant Biology, University of Illinois at Urbana-Champaign, Champaign, IL, USA
2 Carl R. Woese Institute for Genomic Biology, University of Illinois at Urbana-Champaign, Champaign, IL, USA
3 Environmental & Climate Science Department, Brookhaven National Laboratory, Upton, New York, USA
4 School of Biological Sciences, University of Hong Kong, Pokfulam, Hong Kong
5 USDA ARS Global Change and Photosynthesis Research Unit, Urbana, IL, USA
6 Department of Natural Resources and Environmental Sciences, University of Illinois at Urbana-Champaign, Champaign, IL, USA
7 National Center of Supercomputing Applications, University of Illinois at Urbana-Champaign, Champaign, IL, USA
8 Center for Crop Systems Analysis, Wageningen University, The Netherlands.

* These authors contributed equally to this work.
† Correspondence: Carl.Bernacchi@USDA.gov

Received 10 February 2020 Accepted 10 February 2020

Abstract

Photosynthesis is currently measured using time-laborious and/or destructive methods which slows research and breeding efforts to identify crop germplasm with higher photosynthetic capacities. We present a plot-level screening tool for quantification of photosynthetic parameters and pigment contents that utilizes hyperspectral reflectance from sunlit leaf pixels collected from a plot (~2 m×2 m) in <1 min. Using field-grown Nicotiana tabacum with genetically altered photosynthetic pathways over two growing seasons (2017 and 2018), we built predictive models for eight photosynthetic parameters and pigment traits. Using partial least squares regression (PLSR) analysis of plot-level sunlit vegetative reflectance pixels from a single visible near infra-red (VNIR) (400–900 nm) hyperspectral camera, we predict maximum carboxylation rate of Rubisco (V_c,max, R^2=0.79) maximum electron transport rate in given conditions (J_{1800}, R^2=0.59), maximal light-saturated photosynthesis (P_max, R^2=0.54), chlorophyll content (R^2=0.87), the Chl a/b ratio (R^2=0.63), carbon content (R^2=0.47), and nitrogen content (R^2=0.49). Model predictions did not improve when using two cameras spanning 400–1800 nm, suggesting a robust, widely applicable and more ‘cost-effective’ pipeline requiring only a single VNIR camera. The analysis pipeline and methods can be used in any cropping system with modified species-specific PLSR analysis to offer a high-throughput field phenotyping screening for germplasm with improved photosynthetic performance in field trials.

Keywords: Field phenotyping, food security, hyperspectral imaging, photosynthesis, proximal sensing, spectral reflectance.
Introduction

Projected population increase and pressures on land and agricultural resource availability induced by a changing global climate is placing increased demand to secure global food supply in the coming decades (Tilman et al., 2009; Foley et al., 2011). Improving photosynthetic capacity has become a target to enable crop yield increases (Monteith and Moss, 1977; Long et al., 2006; Zhu et al., 2010; Evans, 2013). Inefficiencies in the photosynthetic pathway have inspired research efforts to exploit natural variation in photosynthetic capacity (Lawson et al., 2012), and to improve photosynthetic pathways transgenically (Ott et al., 2015). Thus, crop scientists and breeders face the challenge of characterizing genetic improvements in field trials in a high-throughput manner as a screening tool to identify ‘photosynthetically superior’ germplasm (Furbank and Tester, 2011). While photosynthetic capacity has been successfully estimated from hyperspectral imaging at the ecosystem scale (Serbin et al., 2018), it has only too coarse in spatial resolution to discriminate in mixed germplasm field trials. While hyperspectral analysis has predicted leaf-level photosynthetic capacities and pigment contents (Serbin et al., 2012; Ainsworth et al., 2014; Yendrek et al., 2017; Silva-Perez et al., 2018), it has limitations as leaf clip measurements only pinpoint a few individual leaves in a plot canopy. Currently there are limited tools to screen a whole plot, rather than individual leaves, for photosynthetic performance. Plot-level estimations with proximal sensing platforms are needed to allow rapid capture of reflectance from all sunlit vegetation in the sensor range, eliminating the need to make assumptions about plot performance based on leaf-level samples, and expanding the spatial and temporal capabilities of analysis to capture hundreds of plots in a single day.

The maximum carboxylation rate of Rubisco ($V_{\text{c,max}}$) and maximum electron transport rate in given conditions ($J_{\text{max}}$) are widely used as determinants of photosynthetic capacity for the carbon reduction cycle and the electron transport chain, respectively (von Caemmerer and Farquhar, 1981; von Caemmerer, 2000), and are traditionally derived at the leaf level with infrared gas exchange analysis. The response of leaf-level CO$_2$ assimilation to incrementing CO$_2$ is measured (Long and Bernacchi, 2003) and analyzed (Sharkey et al., 2007) according to the mechanistic model of photosynthesis (Farquhar et al., 1980). The quantum yield of CO$_2$ fixation ($\phi$CO$_2$) and maximum light-saturated photosynthetic rates ($P_{\text{max}}$) are also used as determinants of photosynthetic operating efficiency, as derived from leaf-level gas exchange measurements of the response of CO$_2$ assimilation to incrementing photosynthetically active radiation (PAR) (Ögren and Evans, 1993). Due to the wealth of physiological information provided, leaf-level gas exchange has dominated retrieval of these photosynthetic parameters for decades, but it is limited and time restrictive for the sampling required to measure large crop trials. Additionally, upscaling from leaf gas exchange to determine plot or canopy photosynthetic capacity from gas exchange often requires complex modeling with many assumptions (de Wit, 1965; Evans and Farquhar, 1991; De Pury and Farquhar, 1997; Yin and Struik, 2017; Wu et al., 2019).

Recently, advances have been made in quantifying photosynthesis from spectral analysis at the leaf to ecosystem scales. At the leaf level, with a hand-held spectral leaf gun, photosynthetic capacity ($V_{\text{c,max}}$ and $J_{\text{max}}$) and chlorophyll, carbon (C) and nitrogen (N) content have been predicted successfully from hand-held reflectance spectroscopy across the full electromagnetic spectrum (400–2500 nm) for tree species (Serbin et al., 2012, 2016), productive cropping systems (Ainsworth et al., 2014; Yendrek et al., 2017; Silva-Perez et al., 2018; Ely et al., 2019), and in field trials of Nicotiana tabacum with altered photosynthetic pathways (Fu et al., 2019; Meacham-Hensold et al., 2019). Partial least squares regression (PLSR) analysis of reflectance spectra has also been applied to predict photosynthetic capacity with airborne hyperspectral imaging at the agroecosystem canopy scale (Serbin et al., 2015); however, the most advanced satellite hyperspectral systems capture ~1 pixel per 10–30 m (Transon et al., 2018), which is too sparse in spatial resolution to identify genotypic variation within field trials of many small plots. Advanced UAV (unmanned aerial vehicle) systems are able to capture greater spatial resolution (~40 cm per pixel) (Zarco-Tejeda et al., 2013; Kuwaimana et al., 2018), but still fall short of the millimeter resolution required to build models to predict photosynthetic capacities at the scale of individual leaves in small plots. While multispectral cameras are widely available at higher resolution and used to derive plot-level spectral vegetation indices (SVIs) from discrete spectral wavelengths (Curran et al., 1990; Gamon et al., 1992; Thoenkabail et al., 2000; Zarco-Tejada et al., 2002; Haboudane et al., 2004), SVIs are not able to determine photosynthetic parameters beyond structural inference on physiological processes from discrete spectral bands. Satellite-mounted multispectral imaging systems have also been widely exploited to derive spectral indices such as the enhanced vegetation index (EVI) and normalized difference vegetation index (NDVI), and, more recently, solar-induced fluorescence (SIF) (Guanter et al., 2014; Porcar-Castell et al., 2014; Guan et al., 2016) and linked to ecosystem gross primary productivity (GPP) (Smith et al., 2002; Wylie et al., 2003; Rahman et al., 2005; Zhang et al., 2014, 2018; Barnes et al., 2017; Shi et al., 2017; He et al., 2019). Multispectral SVI and SIF estimates have been incorporated into terrestrial biosphere models to predict photosynthetic capacities at the ecosystem scale (Demarty et al., 2007; Kattge et al., 2009; Zhang et al., 2014), but have not been used to predict photosynthetic capacity in smaller scale plot trials.

Resolving hyperspectral analysis of photosynthetic parameters at the plot level holds many practical and technical challenges. First, hyperspectral cameras and sensors that capture reflectance at the spatial and spectral resolution required for plot-level analysis are often limiting in terms of availability, affordability, and suitability for field trial scanning. Secondly, field phenotyping proximal sensing platforms (Deery et al., 2014) to house such sensors are not currently commercially available and need to be fabricated for purpose. Thirdly, hyperspectral imaging systems generate memory-intensive three-dimensional data sets with two spatial dimensions ($S_x$ and $S_y$) and one spectral ($S_z$) dimension, forming ‘hypercubes’ (Bannon, 2009), necessitating advanced data storage systems and custom analysis pipelines. Fourthly, at the plot level, plant geometrical structure, leaf...
scattering properties, background soil, and dynamic environmental conditions (Verhoef, 1984; Vogelman et al., 1996; Gao et al., 2000; Jay et al., 2016) need to be resolved against leaf-level ‘ground truth’ measurements to accurately infer photosynthetic performance upscaled from leaf to plot level. Finally, ensuring use of this technology answers important physiological questions requires effective interdisciplinary collaboration between engineering, computational, and biological specialists.

In this study, we present a plot-level high-throughput phenotyping platform housing two hyperspectral cameras. One visible near infra-red (VNIR) camera captured reflectance from 400 nm to 900 nm (spectral resolution 2.1 nm) and the second near infra-red (NIR)/shortwave infra-red (SWIR) camera from 900 nm to 1800 nm (spectral resolution 4.9 nm). We created an automated hyperspectral imaging processing pipeline that extracts plot-level sunlit vegetation pixel reflectance spectrum to predict $V_{c,max}, J_{800}$ chlorophyll content, Chl $a\text{b}$, C content, N content, $P_{max}$, and $\phi$CO$_2$. From PLSR analysis of plot-level reflectance spectra from hyperspectral images, we predict these photosynthetic traits in field trials of wild-type and genetically modified lines of Nicotiana tabacum. We assess the contribution of spectral regions and the applicability of this technique to the field phenotyping community, and offer a tool for high-throughput phenotyping of large-scale crop trials to facilitate screening for increasing crop yields.

Materials and methods

Data from two growing seasons (2017 and 2018) were used in this study, presented in two performance tests. For performance test 1, three wild-type and seven transgenic N. tabacum lines were measured over the 2017 and 2018 growing seasons (Table 1). Measurements in 2017 were taken from 22 June to 1 August and in 2018 on 24 and 25 July. For performance test 2, two wild-type and eight transgenic N. tabacum plants were measured in 2018 on 26, 27, and 28 July. In performance test 1, predictive models were built from hyperspectral reflectance (both leaf and plot level) with ground truth data from gas exchange measurement of CO$_2$ response curves for $V_{c,max}$ and $J_{800}$ and leaf pigment extractions for chlorophyll content, Chl $a\text{b}$, C content, and N content. For performance test 2, predictive models for plot- and leaf-level $P_{max}$ and $\phi$CO$_2$ were trained with ground truth data from gas exchange measurement of light response curves.

Plant material

In 2017, three wild-type N. tabacum cultivars and six transgenically modified lines (described in detail in Table 1) were grown at the University of Illinois Energy Farm Facility in Urbana, Illinois (40°03'46.4''N, 88°12'25.4''W, 215 m above sea level). All experiments consisted of four replicated plots of each genotype arranged in a 6 x 6 grid and spaced 0.38 m apart with 36 plants per plot. Each plot measured ~2 x 2 m. All transgenic material is expressed in the Petite Havana background, with the exception of the Rubisco antisense lines in the W38 background. Seedlings were germinated in greenhouse conditions in flat trays using the exception of the Rubisco antisense line (4-KO). Single Rubisco antisense plants were planted to the field without screening. The 4-KO seedlings were screened 8 d post-emergence with chlorophyll fluorescence imaging to identify and select only plants with the PbS knockout phenotype for low non-photochemical quenching (NPQ).

In 2018, two wild-type, five previously grown transgenic lines, and three newly added transgenic lines (described in detail in Table 1) were grown according to the same protocol as in 2017. All transgenic plant material was homozygous, with the exception of the single Rubisco antisense and decreased PbS line (4-KO). Single Rubisco antisense plants were planted to the field without screening. The 4-KO seedlings were screened 8 d post-emergence with chlorophyll fluorescence imaging to identify and select only plants with the PbS knockout phenotype for low non-photochemical quenching (NPQ).

Hyperspectral image collection

A ground-based field phenotyping platform was built to house two hyperspectral push-broom cameras mounted on a horizontal beam (Fig. 1A). The first hyperspectral imaging camera (PIKA II; Resonon, Inc., Bozeman, MT, USA) captured spectral radiation from 400 nm to 900 nm in 2.1 nm contiguous bands (240 spectral bands in total) with 640 spatial channels. The second camera (PIKA NIR; Resonon, Inc.) recorded spectral radiation from 900 nm to 1800 nm in 4.9 nm contiguous bands (164 spectral bands) with 320 spatial channels. Both cameras were mounted at a height of 1.6 m from the soil and were triggered simultaneously above a plot to acquire two images during an ~30 s scan. Images were captured in high-irradiance conditions during a 3 h window around solar noon and stored using SpectronPro software (Resonon, Inc.). A 99% reflective white Teflon panel was mounted horizontally and level with the top of the plant canopy and captured in the field of view for each image (Fig. 1B). Images were captured and stored in raw data mode. The cameras were calibrated to remove electrical and dark current daily prior to data acquisition. Camera integration time was set at 20% below the saturation point according to the radiance signal from the Teflon panel before each scan to avoid saturation.

Hyperspectral image analysis pipeline

An automated image analysis pipeline was created using Python (Python Software Foundation, https://www.python.org/), to extract spectral reflectance from images acquired in raw data mode (Fig. 2A). Data from each camera went through the same three phases of processing: first, conversion of raw data in digital numbers to radiance using radiometric calibration; secondly, the classification of pixels (Fig. 2B); and, thirdly, conversion of radiance pixels to reflectance (Fig. 2C). For the first phase, raw data were converted to absolute radiance using radiometric calibration files from the camera manufacturer. In the second phase, the image was segmented to represent six matter classifications using K-means clustering (Spath, 1985) which separated pixels of interest (sunlit leaves and Teflon) from shaded leaves, soil, platform shadow, and non-biological matter. Thirdly, reflectance ($R$) was calculated using the radiance signature from the Teflon white reference captured in each image against a lab-calibrated Teflon standard using Equation 1:

$$R = \frac{S_{sunlit}}{S_{ref}}$$

where $S_{sunlit}$ is radiance from sunlit leaves, $S_{ref}$ is radiance from the Teflon panel, and $R_{ref}$ is the percentage reflectance from the lab-calibrated Teflon standard. Brightness in digital numbers was converted to percentage reflectance (Fig. 2D), before reflectance from all sunlit pixels in each image was averaged per plot (Fig. 2E). Spectral reflectance from both cameras in the same plot was joined to give reflectance for sunlit leaves per plot from 400 nm to 1800 nm. Spectral reflectance were filtered with a polynomial order of 2 using 11 spectral measurements (nm) as the window length (Savitzky and Golay, 1964). Prior to PLSR analysis, spectral bands below 450 nm and above 1700 nm were removed due to atmospheric water absorption (Hill and Jones, 2000; Serbin et al., 2015).

Leaf spectral measurements

Leaf-level spectral measurements were made using a spectroradiometer (FieldSpec4; Analytical Spectral Devices – ASD, Boulder, CO, USA), with a leaf clip attached to a fiber optic cable. Leaf spectral reflectance was measured in situ from 400 nm to 2500 nm with spectral resolution of 2.1 nm.
3 nm in the visible and NIR spectra (350–1000 nm) and 8 nm in SWIR (1000–2500 nm) spectra. The device houses a radiometrically calibrated light source which was standardized for relative reflectance prior to measurement using a Spectralon (Labsphere Inc., North Dutton, NH, USA) panel for white reference. In 2017, three leaves were sampled per plot and five per plot in 2018. Measurements were made on the last fully expanded leaf, maintaining natural leaf orientation avoiding the leaf midrib and edges. On a single leaf, six reflectance spectra were recorded using the leaf clip attachment in different regions of the same leaf. The six spectra for a single leaf were then averaged to give a mean spectrum per leaf. Each
A single measurement was the mean of 10 scans at a scan speed of 100 ms. A spectral splice correction was applied to each spectrum to remove heat drift effects that may shift the sensors and align the visible and SWIR sensors to the NIR sensors within the FieldSpec4, using the FieldSpectra package in R according to Serbin et al. (2015). For quality control, spectra with abnormally high light levels at 450 nm were excluded from analysis to ensure the leaf clip was properly fastened onto the leaf for each measurement. Spectral samples with a deviation from the mean reflectance >2% were eliminated from analysis along with leaves with fewer than four viable spectra.

Predictive PLSR models for all traits were built at both leaf and canopy levels for comparisons. For plot-level models, the averaged ground truth subsamples (three leaf measurements in 2017 and five leaf measurements in 2018) from each plot were used as input for model building and validation. For leaf-level models, each individual leaf subsample measurement was included as a training data point without averaging per plot.

**PLSR analysis**

Predictive models were built for eight traits, following PLS principles (Wold et al., 2001) according to the protocol of Serbin et al. (2015), modified for *N. tabacum*. Although in our previous work (Fu et al., 2019), multiple stacked machine learning algorithms showed increased predictability (+5% for $R^2$) of photosynthetic capacity ($V_{\text{c,max}}$), we used PLSR only in this work given the ability to derive scaling coefficients across the electromagnetic spectra from this technique, which allow inference of important contributing regions of the spectra for trait prediction. Unlike other predictive algorithms, PLSR coefficient loadings can be calculated to infer the physiological importance of specific spectral bands based on known vegetation spectral properties, and thus can be used to confirm the biological relevance of model builds between different devices and scales.

We used the open-source PLS package (Mevik and Wehrens, 2007) in R, (The R Foundation for Statistical Computing, Wien, Austria) to create a linear model of waveband coefficients that account for trait variation in reflectance spectra. The optimal number of components (latent variables: LVs) for each model build was determined from the minimum root mean square error (RMSE) of the predicted residual sum of squares (PRESS) statistic (Esbensen et al., 2002), using a leave-one-out cross-validation (CV) approach that then makes a prediction for the out-of-sample observation (Siegmund and Jarmer, 2015). This prevents overfitting. Models were trained with data pairs of leaf or plot reflectance and a ground truth.
measurement, and cross-validated with 1000 times random resampling to determine model stability. All model $R^2$ values presented herein are from this CV. Loading weights indicate known spectral peaks or profiles for each model and are translated to the variable importance in projection (VIP), calculated as the weighted sum of squares of PLS weights (Wold et al., 2001; Fearn et al., 2015).

Temperature corrections were not applied to bring photosynthetic parameters to a standard temperature prior to model fitting; absolute plot temperature was not measured at the time of image capture. As such, all leaf- and plot-level models include temperature variation. No outliers were removed from the predictive models presented.

**Infra-red gas exchange measurements**

**CO$_2$ response**

Photosynthetic ($A$) versus intercellular CO$_2$ ($C_i$) response curves were collected within 30 min of the leaf spectral measurements on the same last fully expanded leaves to determine $V_{c,max}$ and $J_{1800}$ for each leaf using a portable leaf gas exchange system with a leaf cuvette (LI-6800; LICOR Biosciences, Lincoln, NE, USA). Four machines were used by four operators to ensure unbiased sampling. Leaf temperature was determined as the mean of three measurements with a hand-held IR gun (FLIR TG54, FLIR® Systems, Inc., Wilsonville, OR, USA). Leaf temperature for gas exchange was set to match this mean leaf temperature prior to each CO$_2$ response curve, and relative humidity was set to 65%. PAR was set to 1800 µmol m$^{-2}$ s$^{-1}$ and CO$_2$ concentrations were adjusted stepwise over a range of 50–2000 µmol mol$^{-1}$ in set increments as follows: 400, 200, 50, 100, 300, 400, 600, 900, 1200, 1500, 1800, and 2000. Leaves were acclimated to chamber conditions for a minimum of 160 s prior to each $A$/$C_i$ curve with a minimum and maximum wait time of 160 s and 200 s, respectively, before each individual measurement of a response curve. $V_{c,max}$ and $J_{1800}$ were determined from these $A$/$C_i$ curves according to the mechanistic model of photosynthesis (Farquhar et al., 1980) and analyzed using a curve fitting utility developed by Sharkey et al. (2007). While light response curves were carried out prior to analysis to determine saturating light intensity as $=\sim 1800$ µmol m$^{-2}$ s$^{-1}$, we refer to maximum electron transport as $J_{1800}$ rather than $J_{c,max}$ to avoid potential false claims of true maximal capacity (Sharkey, 2016). Mesophyll conductance ($g_m$) was constrained according to values for tobacco at 25°C reported previously, with temperature dependency incorporated from the linear relationship of $g_m$ with temperature where $\psi = 0.44 + 0.058x$ (Evans and von Caemmerer, 2013).

**Light response**

In performance test 2, to train the $P_{max}$ and $\phi$CO$_2$ PLSR models, photosynthetic ($A$) versus irradiance ($Q$) response curves were collected within 30 min of leaf spectral measurements, on the same leaves, with a portable leaf gas exchange system (LI-6800; LICOR Biosciences). All environmental settings matched those for $A$/$C_i$ response curves (temperature to match ambient, relative humidity 65%), but with CO$_2$ set to 400 µmol mol$^{-1}$. Irradiance concentrations were adjusted stepwise over a range of 2000–0 µmol m$^{-2}$ s$^{-1}$ in set increments as follows: 2000, 1800, 1400, 1000, 600, 400, 200, 150, 100, 75, 50, and 0. Leaf absorption for each genotype was determined using an integrating sphere (LI-1800; LICOR Biosciences) connected to a spectrometer (USB-2000; Ocean Optics Inc., Dunedin, FL, USA) as the mean absorbance of six last fully expanded leaves (Supplementary Fig. S1G), and $\phi$CO$_2$ was calculated as the slope of the relationship between $A$ and absorbed irradiance below 150 µmol m$^{-2}$ s$^{-1}$. $P_{max}$ was calculated by a non-rectangular curve fit according to Thoron and Johnson (1990) as:

$$P_{max} = \frac{I_s + P_{max} - \sqrt{(I_s + P_{max})^2 - 4I_s}}{2} \theta P_{max} - R_d$$

where $P_{max}$ is maximum light-saturated photosynthesis, $\phi$ is quantum yield, $I_s$ is absorbed irradiance, $\theta$ is the curvature factor, and $R_d$ is the dark respiration rate.

**Chlorophyll, carbon, and nitrogen content**

In performance test 1, immediately following each leaf spectral measurement, a 2.01 cm$^2$ leaf disc was destructively harvested from each leaf using a cork borer, placed in 2 ml tubes and flash-frozen in liquid nitrogen. To determine leaf chlorophyll (mg m$^{-2}$), one leaf disc from each leaf was incubated in 96% (v/v) ethanol for 24 h at 4°C. The bleached material and ethanol were mixed (100 µl of solution for each sample) and analyzed with a Synergy 2 spectrophotometer (BioTek Instruments, Inc, Winooski, VT, USA) at 470, 649, and 665 nm (Lichtenthaler and Wellburn, 1983). To determine leaf carbon and nitrogen content (%), three more 2.01 cm$^2$ leaf discs were destructively harvested, and dried until constant mass, and a subset of ground tissue of known mass (3±0.5 mg) was combusted with oxygen in an elemental analyzer (Costech 4010; Costech Analytical Technologies) and calibrated to %N and %C against an acetylaniide standard curve.

**Results**

**Physiological and spectral characteristics**

Our models captured a wide range of natural and genetically altered trait variation over consecutive growing seasons (2017 and 2018). For performance test 1, averaged plot-level measurements of observed $V_{c,max}$, $J_{1800}$ chlorophyll content, Chl a/b, N content, and C content (Supplementary Fig. S1A–F) include variation of environmental and meteorological conditions (between three and five subsamples per plot), with $V_{c,max}$ from 13.4 µmol m$^{-2}$ s$^{-1}$ to 359.3 µmol m$^{-2}$ s$^{-1}$ (Supplementary Fig. S1A), $J_{1800}$ from 54.9 µmol m$^{-2}$ s$^{-1}$ to 362.1 µmol m$^{-2}$ s$^{-1}$ (Supplementary Fig. S1B), chlorophyll content from 0.1 mg m$^{-2}$ to 0.3 mg m$^{-2}$ (Supplementary Fig. S1C), Chl a/b from 1.7 to 3.7 (Fig. S1D), N content from 2.5% to 8.4% (Supplementary Fig. S1E), and C content from 36.2% to 47.4% (Supplementary Fig. S1F). In performance test 2, from light response curves measured between 26 and 29 July in 2018, $P_{max}$ ranged between 4.1 µmol m$^{-2}$ s$^{-1}$ and 77.7 µmol m$^{-2}$ s$^{-1}$ (Supplementary Fig. S1G, and $\phi$CO$_2$ ranged between 0.024 µmol m$^{-2}$ s$^{-1}$ and 0.064 µmol m$^{-2}$ s$^{-1}$ (Supplementary Fig. S1H). Hyperspectral reflectance from all sunlit pixels per plot used to build PLSR models for all traits exhibit a peak centering at ~550 nm and high reflectivity in the NIR from 800 nm to 1300 nm, and a smaller peak developing from 1440 nm to 1800 nm, following the expected spectral profile pattern. However, reflectance values are slightly lower than expected between 900 nm and 1250 nm (Fig. 3).

**Plot-level PLSR predictions**

The corresponding reflectance spectrum from all sunlit pixels per plot (Fig. 3) paired with the observed, measured traits (Supplementary Fig. S1) produced robust predictive plot-level models for all traits other than $\phi$CO$_2$. Mean spectra used for each model build in performance test 1 varied slightly, as spectra without a paired ‘ground truth’ sample for each trait were eliminated from model build data sets (Fig. 3). For example, in the SsU genotype, $J_{1800}$ could not be determined from gas exchange as the low Rubisco content meant this genotype was never electron transport limited but instead always Rubisco limited. Given that $J_{1800}$ could not be calculated, the $J_{1800}$ spectral sample size is reduced compared with the $V_{c,max}$ model build (Fig. 3A, B).
Similarly, a small number of leaf disc samples for leaf chlorophyll, carbon, and nitrogen content were lost in transportation, storage, or during analysis, creating slight variation in spectral sample used for chlorophyll (Fig. 3C), and C and N (Fig. 3D) model builds. Using reflectance spectra from 450–900 nm only, collected with a single VNIR hyperspectral camera, $V_c$,max ($R^2$ 0.79, RMSE% 11.9), $J_{1800}$ ($R^2$ 0.59, RMSE% 11.5), chlorophyll content ($R^2$ 0.87, RMSE% 10), Chl $a:b$ ($R^2$ 0.63, RMSE% 18.5), and $P_{max}$ ($R^2$ 0.54, RMSE% 10.6) were highly predictable from PLS hyperspectral regression models (Fig. 4; Table 2). PLSR predictions performed moderately well for C content ($R^2$ 0.47, RMSE% 18.7, Fig. 4E) and N content ($R^2$ 0.49, RMSE% 15.9%, Fig. 4F), but offered no predictability for $\phi CO_2$ ($R^2$ 0.02, RMSE%, Fig. 4H; Table 2). When compared with the single camera models, PLSR models using both hyperspectral cameras (Fig. 5) had weakened predictive power (lower $R^2$ and increased RMSE%) for all traits, except Chl $a:b$ (Table 2). Using both cameras, $V_c$,max ($R^2$ 0.74, RMSE% 13.1, Fig. 5B), $R^2$ decreased by 5% and RMSE% increased by 1.9%. However, for Chl $a:b$, predictability increases when both cameras are used ($R^2$ 0.77, RMSE% 14, Fig. 5D), where $R^2$ increases by 14%, and RMSE% decreases by 4.5% (Table 2).

Model loading weights indicate the importance of regions of the reflectance spectra for trait variation. For plot-level PLSR predictions with a single VNIR camera (450–900 nm), the region around 700 nm is important for all traits (Fig. 3). When translated to a VIP score for easier interpretation (Fig. 6), 700 nm is shown to be the most important region for $V_c$,max, $J_{1800}$, and chlorophyll content predictions. While ~700 nm is important for all other traits, for C and N content regions from 500 nm to 650 nm and from ~820 nm and ~870 nm in the NIR also hold importance (Fig. 6C, D). For Chl $a:b$ and $P_{max}$, the entire NIR from 700 nm to 900 nm holds weight. When plot-level model loadings (Fig. 6) and VIP scores (Fig. 7) are compared with those from leaf-level PLSR models built using the same leaves that ground truth the plot-level models, they generally follow the same response pattern for all traits, with the exception of VIP scores for Chl $a:b$ (Fig. 7D) and $\phi CO_2$ (Fig. 7H).

Leaf-level PLSR models

When leaf-level PLSR models were built to include different spectral ranges (500–900, 500–1700, and 500–2400 nm), only $V_c$,max, N content, and $P_{max}$ predictability showed minor improvement...
with greater spectral range (Table 3). The CV $R^2$ for $V_{c,max}$ remained the same when the model used reflectance from 500 nm to 1700 nm, rather than from 500 nm to 900 nm, but there was a 2% increase when the full spectrum was used (500–2400 nm). For $P_{max}$ CV, $R^2$ increased by 7% when the spectral bandwidth matched that of both hyperspectral cameras (500–1700 nm).
rather than with the single VNIR only (500–900 nm), but no benefit was seen with the addition of the SWIR (1700–2400 nm). Leaf N content is the only trait for which improved predictability correlated with increased spectral range, with a 3% increase in CV \( R^2 \) using 500–1700 nm, and a further 7% increase using 500–2400 nm (Table 2). Unlike the plot-level \( \phi CO_2 \) model, \( \phi CO_2 \) was highly predictable from PLSR analysis of leaf-level spectral reflectance (CV \( R^2 \)) between 0.61 and 0.63, Table 3.

For PLSR models built at the leaf level for three different spectral ranges (500–900, 500–1700, and 500–2400 nm, Fig. 8A–H), VIP scores in the VNR from 400 nm to 800 nm were larger than those in the NIR and SWIR from 800 nm to 2400 nm, for all traits other than leaf C and N contents. For C (Fig. 8E) and N (Fig. 8F) contents, VIP peaks at ~1400 nm and 1900 nm suggest these regions also hold high importance for predictability. Comparing VIP scores for models built with a single VNIR camera (Pika II, 450–900 nm) showed greater variability than those for models built with two cameras (Pika II and Pika NIR, Fig. 8I–J). While all models had high VIP scores between 450 nm and 900 nm, and C and N contents followed a similar pattern seen at the leaf level (Fig. 8M, N), \( V_{\text{c,max}} \) and \( J_{\text{1800}} \) also had VIP peaks at ~1100 nm and 1700 nm (Fig. 8I, J). VIP scores for plot-level \( \phi CO_2 \) models were not shown due to the lack of predictability of this parameter with hyperspectral imaging in this study.

### Discussion

Results show that photosynthetic capacity (\( V_{\text{c,max}} \) and \( J_{\text{1800}} \)), maximum light-saturated photosynthesis (\( P_{\text{max}} \)), and associated photosynthetic pigment contents (C, N, chlorophyll, and Chl \( a:b \)) can be predicted using high-throughput proximal plot-level hyperspectral imaging. PLSR analysis of reflectance spectra is now well established as a robust tool for estimating photosynthetic performance at the leaf level (Serbin et al., 2012; Ainsworth et al., 2014; Yendrek et al., 2017; Silva-Perez et al., 2018), and the technique holds integrity when used on plants with altered photosynthetic pathways (Meacham-Hensold et al., 2019). At a broader spatio–temporal scale, data collected with the Airbourne Visible Infrared Imaging spectrometer (AVIRIS) has been used with PLSR analysis of reflectance spectra to successfully predict photosynthetic capacity \( (V_{\text{c,max}}) \) at the agroecosystem canopy level, providing lessons for ecosystem and earth system models (Serbin et al., 2015). The results here offer a tool to measure between these contrasting scales to derive photosynthetic capacity as a crop breeding selection tool. The predictive models presented in this study show the utility of hyperspectral imaging as a tool for plot-level phenotyping for superior photosynthetic performance in large-scale field trials, offering potential to screen hundreds of accessions in a single day.

### Spectral compositional features

Electromagnetic energy in the visible range provides the energy for photosynthesis, and absorption in the visible region specifically between 660 nm and 700 nm is of high importance for photosynthetic predictions from reflectance spectra (Serbin et al., 2012; Silva-Perez et al., 2018; Fu et al., 2019; Meacham-Hensold et al., 2019). Similarly, the region of transition from low reflectivity in the visible range to higher reflectivity in the NIR (~750 nm), termed ‘red-edge’, has been utilized to predict \( V_{\text{c,max}} \) (Dillen et al., 2012) and is also heavily weighted in previous PLSR predictive model loadings (Yendrek et al., 2017; Silva-Perez et al., 2018; Meacham-Hensold et al., 2019). These relationships are underpinned by the importance of chlorophyll, nitrogen, and Rubisco in photosynthetic processes (Evans, 1989) and the dominating influence of these pigments on reflectance spectra from 500 nm to 800 nm (Curran, 1989; Elvidge, 1990; Ustin et al., 2009). VIP scores quantify the contribution of each variable (spectral bands) to overall variance and, in this study, when models were built using data from a single VNIR camera (450–900 nm), the greatest peaks in VIP scores are also in the chlorophyll absorption bands and the red-edge regions for \( V_{\text{c,max}} \) and \( J_{\text{1800}} \). Pmax, chlorophyll content, and N content (Fig. 7), fitting with previous spectral reflectance compositional observations (Farré et al., 2015).

Previous leaf-level studies show that some regions of the lower energy NIR, particularly ~1400 nm, are also important
for photosynthetic PLSR predictions (Yendrek et al., 2017). However, in this study, plot-level models built using reflectance in the VNIR range only (450–900 nm) give greater predictability than those using reflectance from a greater spectral range (450–1700 nm) (Figs 4, 5; Table 3). This was unexpected and may be the result of compounding factors. In our
plot-level analysis using both cameras (450–1700 nm), we removed reflectance between 1313 nm and 1440 nm given convolution of reflectance spectra in that region from atmospheric water absorption properties (Hill and Jones, 2000; Serbin et al., 2015), where removal of these bands is unnecessary when using a leaf clip with an artificial light source. Thus, it follows, with the absence of reflectance at ~1400 nm, that the spectral region detected by the single VNIR camera (400–900 nm)
captures the most important regions for photosynthetic predictions. This offers one possible explanation for the strength of PLSR predictions for all parameters in this study from the single VNIR camera (Fig. 4). In addition, when reflectance spectra from both cameras (450–1700 nm) were used to build predictive models, VIP scores for chlorophyll content (Fig. 8K), Chl a:b (Fig. 8L), and \( P_{\text{max}} \) (Fig. 8P) show that reflectance from the NIR above 900 nm holds little or no importance (Fig. 8). This is not surprising given that the absorption of chlorophyll occurs in the visible range (Ustin et al., 2009) and that \( P_{\text{max}} \) should be highly related to pigment and pigment pool distributions. However, for \( V_{\text{c,max}} \) (Fig. 8I), \( J_{1800} \) (Fig. 8J), C content (Fig. 8M), and N content (Fig. 8N), while VIP peaks between 400 nm and 900 nm dominate, peaks at ~1150 nm and 1750 nm are present, suggesting secondary importance of these regions. Despite the known spectral properties for N and C contents in these regions (Curran, 1989; Asner and Martin, 2008), and similarly high VIP scores around ~1100 nm in predictions of \( V_{\text{c,max}} \) from airborne spectroscopy (Serbin et al., 2015), models for these three parameters built with reflectance from both cameras (450–1700 nm) rather than just the VNIR (450–900 nm) are weaker (Figs 4, 5; Table 2).

Chl a:b is the only trait for which predictions improve when two cameras (450–1700 nm) are used for the model build rather than the single VNIR (450–900 nm) camera (Figs 4D, 5D). With known chlorophyll absorption dominant only in the visible range, supported by the low loading values for the leaf level Chl a:b models above 900 nm (Fig. 8D), this raises questions as to the cause of improved predictability when adding reflectance spectra above 900 nm. This is probably due to the dilution effect for spectral regions of physiological importance when a ratio of two physiological traits is presented. While the Chl a:b model is unlikely to be overfit given the reliance on the PRESS statistic in latent variable number selection, physiological importance is reduced, allowing ‘statistical’ number training rather than physiologically based ‘trait’ training. Thus care should be taken to eliminate spectral regions shown to hold little weight for the original trait pair when using this PLSR technique to predict ratio values.

Leaf-level comparisons

In attempts to understand the relationship between spectral range and predictability power of PLSR models, we built leaf-level models for all of the plot-level ground truth material measured in this study at three different spectral ranges (Table 3). For each trait, we built models first using reflectance spectra measured with the FieldSpec4 from 500 nm to 900 nm, secondly from 500 nm to 1700 nm, and thirdly from 500 nm to 2400 nm. At the leaf level, with a single device measuring from 400 nm to 2500 nm and an artificial light source, the only trait prediction that improved with greater spectral range inclusion was leaf N (500–900 nm \( R^2 = 0.66 \), 500–1700 nm \( R^2 = 0.69 \), 500–2400 nm \( R^2 = 0.76 \), Table 3). The predictability of all other parameters was not increased with increased spectral range. This may be due to the almost equal importance of VIP peaks around 1400 nm and 1900 nm when compared with the chlorophyll and red-edge regions from 500 nm to 800 nm for N content (Fig. 8F). In contrast, at the leaf level for all other predicted traits in this study, the highest VIP scores occur between 500 nm and 800 nm, with only small peaks in the NIR and SWIR (Fig. 8A–H), which may explain the

Fig. 7. Comparison of variable importance projection (VIP) scores from leaf-level and plot-level PLSR models from 450–900 nm for all traits: \( V_{\text{c,max}} \) (A), \( J_{1800} \) (B), chlorophyll content (C), Chl a:b (D), C content (E), and N (F) in performance test 1, and \( P_{\text{max}} \) (G) and \( \phi_{\text{CO}_2} \) (H) in performance test 2.
lack of correlation between PLSR prediction power and spectral range included in the leaf-level model builds for \( V_{\text{c,max}} \), \( J_{1800} \), chlorophyll content, Chl \( a:b \), C content, \( P_{\text{max}} \), and \( \phi_{\text{CO}_2} \). While \( \phi_{\text{CO}_2} \) is not predictable with PLSR analysis at the plot level (Figs 4H, 5H), it is highly predictable at the leaf level (Table 3; Supplementary Fig. S2), highlighting the need for high variation in observed trait values, to cover greater ‘trait space’ (Ely et al., 2019) for building robust models (Meacham-Hensold et al., 2019). Where observed leaf traits are averaged (between three and five subsamples) at the plot level for \( \phi_{\text{CO}_2} \), observed measurement repetitions are thus reduced, shrinking the trait space and consequently the model prediction strength.

Vegetative structural reflective properties and the comparative loading and VIP scores for leaf and plot-level models from 450 nm to 900 nm (Figs 6, 7) support the strength of plot-level models built with a single VNIR camera (Fig. 4). Loadings and VIP scores may support a lack of improved predictability when models for the same traits are built with reflectance from two cameras that span a greater spectral range (400–1700 nm), but they do not explain the apparent reduction in predictive power (Table 3). N and C content predictions, in particular, should perhaps be improved when lower energy regions of the NIR are included in analysis with both cameras, due to the known absorption features properties of C and N in the NIR (Curran, 1989), and the strong VIP peaks at ~1100 nm and 1700 nm (Fig. 8M, N). This unexpected reduced model strength with increased spectral range is likely to be due to instrumentation limitations. Hyperspectral imaging equipment for phenotyping in field trials is limited. We used two hyperspectral cameras, with different spectral resolution (Pika II, 2.1 nm; Pika NIR, 4.9 nm), different spatial resolution (Pika II, 7.4 µm pixel size; Pika NIR, 30 µm pixel size), and different signal to noise ratios (Pika II,198; Pika NIR, 1885), given the lack of affordability and availability of a single sensor to cover the full electromagnetic spectra. The NIR camera has greater intrinsic error.

### Improving plot-level hyperspectral predictions

The quality of the signal from the Pika NIR (900–1800 nm) camera presents a key challenge throughout this work. Model predictions using two cameras are probably weakened due to technical limitations rather than lack of importance of particular NIR spectral regions for physiological trait prediction. The reflectance profile from the Pika NIR imaging system, ~900–1250 nm, is lower than expected when compared with reflectance measured with a leaf clip. Working with spectral reflectance measured by imaging systems using sunlight rather than a leaf clip with an artificial light source presents

---

### Table 3. PLSR models built at leaf level for all traits using three different spectral ranges (500–900, 500–1700, and 500–2400 nm)

| Spectral range (nm) | \( R^2 \) Train | \( R^2 \) CV | RMSECV (trait unit) | RMSE (%) | Model bias (trait unit) | Latent variable (LV) no. |
|---------------------|----------------|-------------|---------------------|---------|-------------------------|------------------------|
| 500–900             | 0.71           | 0.67        | 48.33               |         |                         | 7                      |
| 500–1700            | 0.75           | 0.69        | 45.21               |         |                         | 10                     |
| 500–2400            | 0.79           | 0.69        | 41.67               |         |                         | 11                     |
| 500–900             | 0.59           | 0.40        | 38.58               |         |                         | 11                     |
| 500–1700            | 0.58           | 0.39        | 39.15               |         |                         | 11                     |
| 500–2400            | 0.53           | 0.40        | 41.38               |         |                         | 8                      |
| 500–900             | 0.82           | 0.78        | 0.02                |         |                         | 10                     |
| 500–1700            | 0.78           | 0.74        | 0.03                |         |                         | 6                      |
| 500–2400            | 0.80           | 0.77        | 0.03                |         |                         | 6                      |
| Chlorophyll content |                 |             |                     |         |                         |                        |
| 500–900             | 8.82           |             | 0.00007             |         |                         | 10                     |
| 500–1700            | 9.76           |             | 0.00003             |         |                         | 6                      |
| 500–2400            | 9.32           |             | 0.00001             |         |                         | 6                      |
| Chl \( a:b \)       |                 |             |                     |         |                         |                        |
| 500–900             | 8.56           |             | -0.003              |         |                         | 14                     |
| 500–1700            | 8.84           |             | 0.00001             |         |                         | 15                     |
| 500–2400            | 7.50           |             | 0.005               |         |                         | 13                     |
| C content (%)       |                 |             |                     |         |                         |                        |
| 500–900             | 7.85           |             | -0.011              |         |                         | 15                     |
| 500–1700            | 8.30           |             | 0.007               |         |                         | 15                     |
| 500–2400            | 7.84           |             | 0.016               |         |                         | 15                     |
| N content (%)       |                 |             |                     |         |                         |                        |
| 500–900             | 8.50           |             | 0.011               |         |                         | 15                     |
| 500–1700            | 8.65           |             | 0.007               |         |                         | 15                     |
| 500–2400            | 7.50           |             | 0.005               |         |                         | 15                     |
| \( P_{\text{max}} \) (µmol m\(^{-2}\) s\(^{-1}\)) |                 |             |                     |         |                         |                        |
| 500–900             | 10.92          |             | 0.12                |         |                         | 9                      |
| 500–1700            | 9.71           |             | -0.04               |         |                         | 13                     |
| 500–2400            | 9.55           |             | 0.04                |         |                         | 13                     |
| \( \phi_{\text{CO}_2} \) (quanta/A µmol m\(^{-2}\) s\(^{-1}\)) |                 |             |                     |         |                         |                        |
| 500–900             | 8.82           |             | 0.000004            |         |                         | 11                     |
| 500–1700            | 8.63           |             | -0.000001           |         |                         | 12                     |
| 500–2400            | 9.24           |             | 0.000044            |         |                         | 9                      |
challenges, with light having been influenced by the atmosphere before reaching the leaf and again after reflection before detection by a sensor. This results in a more complex signal compared with reflectance from integrated full-spectrum leaf-level devices. For example, quantification of leaf angles, removal of background noise from scattered reflectance at lower
canopy levels, removal of background noise from soil (Verhoef, 1984; Gao et al., 2000), and correction for plot temperature at the time of image capture (Serbin et al., 2015) could all improve plot-level model strength. Our plot-level reflectance spectra are also lower between 900 nm and 1250 nm than those from aircraft and other proximal hyperspectral imagers. Proximal hyperspectral imagery usually presents data captured from nadir sensors rather than push-broom scanning platforms. At the time of our data collection, for mounting ~1 m above the target vegetation on a proximal sensing push-cart, push-broom line sensors offered the greatest spatial resolution and affordability. However, the camera angle rotation increases directional anisotropy and, coupled with light scattering from background vegetation, increases our signal to noise ratio. While our automated analysis pipeline (Fig. 3) very accurately accounts for radiance at the time of image capture using a Teflon reference panel for accurate conversion to reflectance (Fig. 2B), the signal could probably be improved with an updated nadir scanner and future incorporation of more complex radiative transfer modelling to account for background scattering. Leaf-level VIP scores show less variation than plot-level scores (Fig. 8), particularly in the NIR. While VIP scores are higher at the plot level, peaks do follow the same trends, thus the variation is likely to be indicative of scattering detected by the NIR hyperspectral camera and sensor noise rather than a need to question the true importance of these regions for prediction of a given trait.

The variation in plot-level ground truthing also presents a known challenge as plot-level estimations are trained with leaf-level measurements. While currently this is the only realistic ground truth method for canopy photosynthetic measurements, it is not ideal given the known limitations of applying leaf-level measurements to canopy estimations (Amthor, 1994; Baldocchi and Harley, 1995; De Pury and Farquhar, 1997; Wu et al., 2016) and the known variation in photosynthetic rates and capacities within crop canopies of the same germplasm and even within plant crowns at the highest levels of a canopy due to variation in light environment (Niinemets, 2007). More robust plot-level models could be trained with a greater number of ground truth samples, but the time taken to obtain gas exchange measurements of photosynthetic capacity is a limitation. These challenges persist for the high-throughput phenotyping and the remote-sensing community and, as equipment develops and sensor integration capabilities advance, predictive models of the nature presented in this study will probably improve. Despite these challenges, this study proposes robust plot-level predictions of key photosynthetic parameters and structural traits that are the focus of current research efforts to increase crop yields for global food security (Evans, 2013; Ort et al., 2015).

The challenges facing agricultural production in the face of resource limitation and changing climate necessitates methods for rapid screening of large field trials for productivity and performance. The results from the automated hyperspectral image analysis pipeline we present synthesize high-resolution plot-level information to a single sunlit plot leaf reflectance spectrum for use in a variety of applications. Photosynthetic predictions from PLSR analysis of this output offers a tool for rapid field phenotyping for photosynthetic performance. Such synthesis of large spatial and temporal data sets with user-friendly analysis pipelines that derive biologically relevant outcomes will be increasingly important in the fight for increased global food production. The success of predictive models with a single VNIR hyperspectral camera widens the relevance and potential application of this technique for greater utility, as reduced spectral bandwidth equates to reduced cost of acquisition and operation of hyperspectral imaging systems.

Supplementary data

Supplementary data are available at JXB online.

Table S1. Leaf absorption values used to correct $P_{\text{max}}$ and $\phi CO_2$ for genotypes in performance test 2.

Fig. S2. ‘Ground truth’ $V_{\text{c,max}}$, chlorophyll content, Chl $a:b$, N content, C content, $P_{\text{max}}$ and $\phi CO_2$ values used to train predictive models.

Fig. S3. Comparison between observed photosynthetic parameters and those predicted from PLS regression of leaf-level reflectance using ASD FieldSpec4 with leaf clip attachment for $\phi CO_2$ from reflectance from 500 nm to 900 nm and from 500 nm to 1700 nm.

Dataset 1. Spectrum collected with hyperspectral imaging cameras used for model builds for each trait as presented in Fig. 3.

Dataset 2. PLSR model predictions from a single VNIR hyperspectral camera (450–900 nm), as shown in Fig. 4.

Dataset 3. PLSR model predictions from two hyperspectral cameras (450–1800 nm), as shown in Fig. 5.

Acknowledgements

This research was supported by a Bill and Melinda Gates Foundation grant OPP1060461, ‘RIPE—Realizing increased photosynthetic efficiency for sustainable increases in crop yield’ and the USDA-ARS through Project Number 5012-21000-030-00D to the Global Change and Photosynthesis Research Unit. We thank David Drag, Ben Harbaugh, Ben Thompson, and Ron Edquilang for greenhouse and field plant management. Professor Susanne von Caemmerer (ARC Centre of Excellence for Translational Photosynthesis Research, Australian National University) kindly provided the Rubisco antisense Nicotiana tabacum. Johannes Kromdijk, Katarzyna Glowka, Steven Driever, and Stephen P. Long provided transgenic N. tabacum lines VPZ-23, PSBS-43, pbs-4, LMD, and LCD, and Paul South and Donald R. Ort provided lines Bypass AP3 and Bypass AP3/RNAi. Co-authors SPS and JW were supported by the United States Department of Energy contract no. DE-SC0012704 to Brookhaven National Laboratory. We also thank Caitlin Moore, Amanda Cavanagh, Marshall Mitchell, Emily Timms, Justine Brumm, Kyle Coffland, Morgan Prinn, Alyssa Dwyer, Alex Riley, Isaac Howenstein, Jennifer Ward, Sam Jameson, and Elena Pelech for assistance with the field work. Any opinions, findings, and conclusions or recommendations expressed in this publication are those of the author(s) and do not necessarily reflect the views of the U.S. Department of Agriculture (USDA). Mention of trade names or commercial products in this publication is solely for the purpose of providing specific information and does not imply recommendation or endorsement by the USDA. USDA is an equal opportunity provider and employer.
References

Ainsworth EA, Serbin SP, Skonczeka JA, Townsend PA. 2014. Using leaf optical properties to detect ozone effects on foliar biochemistry. Photosynthesis Research 119, 65–76.

Amthor J. 1994. Scaling CO2–photosynthesis relationships from the leaf to the canopy. Photosynthesis Research 39, 321–350.

Asner GP, Martin RE. 2008. Spectral and chemical analysis of tropical forests: scaling from leaf to canopy levels. Remote Sensing of Environment 112, 3958–3970.

Baldocchi D, Harley P. 1995. Scaling carbon dioxide and water vapour exchange from leaf to canopy in a deciduous forest. II. Model testing and application. Plant, Cell & Environment 18, 1157–1173.

Bannon D. 2009. Hyperspectral imaging: cubes and slices. Nature Photonics 3, 627.

Barnes ML, Breshears DD, Law DJ, van Leeuwen WJD, Monson RK, Fojtik AC, Barron-Gafford GA, Moore DJP. 2017. Beyond greenness: detecting temporal changes in photosynthetic capacity with hyperspectral reflectance data. PLoS One 12, e0189539.

Curran PJ. 1989. Remote sensing of foliar chemistry. Remote Sensing of Environment 30, 271–278.

Curran PJ, Dungan JL, Gholz HL. 1990. Exploring the relationship between reflectance red edge and chlorophyll content in slash pine. Tree Physiology 7, 33–48.

Deery D, Jimenez-Berni J, Jones H, Sirault X, Furbank R. 2014. Proximal remote sensing buggies and potential applications for field-based phenotyping. Agronomy 4, 349–379.

Demarty J, Chevallier F, Friend A, Viovy N, Piao S, Ciais P. 2007. Assimilation of global MODIS leaf area index retrievals within a terrestrial biosphere model. Geophysical Research Letters 34, doi: 10.1029/2007GL030014.

De Purry DGG, Farquhar GD. 1997. Simple scaling of photosynthesis from leaves to canopies without the errors of big-leaf models. Plant, Cell & Environment 20, 537–557.

de Wit CT. 1965. Photosynthesis of leaf canopies. Wageningen: Pudoc.

Dillen SY, de Beeck MO, Hufkens K, Buonanduci M, Phillips NG. 2016. Improving yield by exploiting close-range imaging spectroscopy. Remote Sensing of Environment 197, 220–236.

Ely KS, Burnett AC, Lieberman-Cribbin W, Serbin SP, Rogers A. 2019. Spectroscopy can predict key leaf traits associated with source–sink balance and carbon–nitrogen status. Journal of Experimental Botany 70, 1789–1799.

Esbensen KH, Guyt D, Westad F, Hounoller LP. 2002. Multivariate data analysis: in practice. An introduction to multivariate data analysis and experimental design. Chapman & Hall.

Evans J, Farquhar G. 1991. Modelling canopy photosynthesis from the biochemical of the C3 chloroplast. In: Boote KJ, I R, eds. Modelling crop photosynthesis: from biochemistry to canopy. Madison, WI: Crop Science Society of America, 1–15.

Evans JR. 1989. Photosynthesis and nitrogen relationships in leaves of C3 plants. Oecologia 78, 9–19.

Evans JR. 2013. Improving photosynthesis. Plant Physiology 162, 1780–1793.

Evans JR, von Caemmerer S. 2013. Temperature response of carbon isotope discrimination and mesophyll conductance in tobacco. Plant, Cell & Environment 36, 745–756.

Farquhar GD, von Caemmerer S, Berry JA. 1980. A biochemical model of photosynthetic CO2 assimilation in leaves of C3 species. Planta 149, 78–90.

Farrés M, Platikanov S, Tsakovski S, Tauler R. 2015. Comparison of the variable importance in projection (VIP) and of the selectivity ratio (SR) methods for variable selection and interpretation. Journal of Chemometrics 29, 528–536.

Foley JA, Ramankutty N, Brauman KA, et al. 2011. Solutions for a cultivated planet. Nature 478, 337–342.

Fu P, Meacham-Hensold K, Guan K, Bernacchi CJ. 2019. Hyperspectral leaf reflectance as proxy for photosynthetic capacities: an ensemble approach based on multiple machine learning algorithms. Frontiers in Plant Science 10, 730.

Furbank RT, Tester M. 2011. Phenomics—technologies to relieve the phenotyping bottleneck. Trends in Plant Science 16, 635–644.

Gamon J, Penuelas J, Field C. 1992. A narrow-wavelength spectral index that tracks diurnal changes in photosynthetic efficiency. Remote Sensing of Environment 41, 35–44.

Gao X, Huete AR, Ni W, Miura T. 2000. Optical–biophysical relationships of vegetation spectra without background contamination. Remote Sensing of Environment 74, 609–620.

Glwacka K, Kromdijk J, Kucera K, Xie J, Cavanagh AP, Leonelli L, Leakey ADB, Ort DR, Niyogi KK, Long SP. 2018. Photosystem II Subunit S overexpression increases the efficiency of water use in a field-grown crop. Nature Communications 9, 868.

Glwacka K, Kromdijk J, Leonelli L, Niyogi KK, Clemente TE, Long SP. 2016. An evaluation of new and established methods to determine T-DNA copy number and homozygosity in transgenic plants. Plant, Cell & Environment 39, 908–917.

Guan K, Berry JA, Zhang Y, Joiner J, Guanter L, Badgley G, Lobell DB. 2016. Improving the monitoring of crop productivity using spaceborne solar-induced fluorescence. Global Change Biology 22, 716–726.

Guanter L, Zhang Y, Jung M, et al. 2014. Global and time-resolved monitoring of crop photosynthesis with chlorophyll fluorescence. Proceedings of the National Academy of Sciences, USA 111, E1327–E1333.

Haboudane D, Miller JR, Pattey E, Zarco-Tejada PJ, Strachan IB. 2004. Hyperspectral vegetation indices and novel algorithms for predicting green LAI of crop canopies; modeling and validation in the context of precision agriculture. Remote Sensing of Environment 90, 337–352.

He L, Chen JM, Liu J, et al. 2019. Diverse photosynthetic capacity of global ecosystems mapped by satellite chlorophyll fluorescence measurements. Remote Sensing of Environment 232, 111344.

Hill C, Jones RL. 2000. Absorption of solar radiation by water vapor in clear and cloudy skies: implications for anomalous absorption. Journal of Geophysical Research: Atmospheres 105, 9421–9428.

Hudson GS, Evans JR, van Caemmerer S, Arvidsson YB, Andrews TJ. 1992. Reduction of ribulose-1,5-bisphosphate carboxylase/oxygenase content by antisense RNA reduces photosynthesis in transgenic tobacco plants. Plant Physiology 98, 294–302.

Jay S, Bendoula R, Hadoux X, Férét J-B, Gorretta N. 2016. A physically-based model for retrieving foliar biochemistry and leaf orientation using close-range imaging spectroscopy. Remote Sensing of Environment 177, 220–236.

Kattge J, Knorr W, Raddatz T, Wirth C. 2009. Quantifying photosynthetic capacity and its relationship to leaf nitrogen content for global-scale terrestrial biosphere models. Global Change Biology 15, 976–991.

Kromdijk J, Glowacka K, Leonelli L, Gablyl ST, Iwai M, Niyogi KK, Long SP. 2016. Improving photosynthesis and crop productivity by accelerating recovery from photoprotection. Science 354, 857–861.

Lawson T, Kramer DM, Raines CA. 2012. Improving yield by exploiting mechanisms underlying natural variation of photosynthesis. Current Opinion in Biotechnology 23, 215–220.

Lichtenthaler HK, Wellburn AR. 1983. Determinations of total carotenoids and chlorophylls of leaf extracts in different solvents. Biochemical Society Transactions 11, 591–592.

Long SP, Bernacchi CJ. 2003. Gas exchange measurements, what can they tell us about the underlying limitations to photosynthesis? Procedures and sources of error. Journal of Experimental Botany 54, 2393–2401.

Long SP, Zhu XG, Naidu SL, Ort DR. 2006. Can improvement in photosynthesis increase crop yields? Plant, Cell & Environment 29, 315–330.

Meacham-Hensold K, Montes CM, Wu J, et al. 2019. High-throughput field phenotyping using hyperspectral reflectance and partial least squares regression (PLSR) reveals genetic modifications to photosynthetic capacity. Remote Sensing of Environment 231, 111176.

Mevik BH, Wehrens R. 2007. The pls package: principal component and partial least squares regression. Journal of Statistical Software 2, 1–24.

Monelith JL, Moss CJ. 1977. Climate and the efficiency of crop production in Britain [and Discussion]. Philosophical Transactions of the Royal Society B: Biological Sciences 281, 277–294.

Niinemets U. 2007. Photosynthesis and resource distribution through plant canopies. Plant, Cell & Environment 30, 1052–1071.
Ögren E, Evans JR. 1993. Photosynthetic light–response curves. Planta 189, 182–190.

Ort DR, Merchant SS, Alric J, et al. 2015. Redesigning photosynthesis to sustainably meet global food and bioenergy demand. Proceedings of the National Academy of Sciences, USA 112, 8529–8536.

Porcar-Castell A, Tystsjärvi E, Atherton J, van der Tol C, Flexas J, Pfündel EE, Moreno J, Frankenberg C, Berry JA. 2014. Linking chlorophyll a fluorescence to photosynthesis for remote sensing applications: mechanisms and challenges. Journal of Experimental Botany 65, 4065–4095.

Rahman A, Sims D, Cordova V, El-Masri B. 2005. Potential of MODIS EVI and surface temperature for directly estimating per-pixel ecosystem C fluxes. Geophysical Research Letters 32, doi: 10.1029/2005GL024127.

Ruwayimana M, Satyanarayana B, Otero V, M Muslim A, Syafiq A M, Ibrahim S, Raymaekers D, Koedam N, Dahdouh-Guebas F. 2018. The advantages of using drones over space-borne imagery in the mapping of mangrove forests. PLoS One 13, e0200289.

Savitzky A, Golay MJ. 1964. Smoothing and differentiation of data by simplified least squares procedures. Analytical Chemistry 36, 1627–1639.

Serbin SP, Dillaway DN, Kruger EL, Townsend PA. 2012. Leaf optical properties reflect variation in photosynthetic metabolism and its sensitivity to temperature. Journal of Experimental Botany 63, 489–502.

Serbin SP, Singh A, Desai AR, Dubois SG, Jablonski AD, Kingdon CC, Kruger EL, Townsend PA. 2015. Remotely estimating photosynthetic capacity, and its response to temperature, in vegetation canopies using imaging spectroscopy, Remote Sensing of Environment 167, 78–87.

Serbin SP, Singh A, McNeil BE, Kingdon CC, Townsend PA. 2016. Spectroscopic determination of leaf morphological and biochemical traits for northern temperate and boreal tree species. Ecological Applications 24, 1651–1669.

Sharkey TD. 2018. What gas exchange data can tell us about photosynthesis. Plant, Cell & Environment 39, 1161–1163.

Sharkey TD, Bernacchi CJ, Farquhar GD, Singsaas EL. 2007. Fitting photosynthetic carbon dioxide response curves for C3 leaves. Plant, Cell & Environment 30, 1035–1040.

Shi H, Li L, Eamus D, Huete A, Cleverly J, Tian X, Yu Q, Wang S, Montagnani L, Magliulo V. 2017. Assessing the ability of MODIS EVI to estimate terrestrial ecosystem gross primary production of multiple land cover types. Ecological Indicators 72, 153–164.

Siegmund B, Jarmer T. 2015. Comparison of different regression models and validation techniques for the assessment of wheat leaf area index from hyperspectral data. International Journal of Remote Sensing 36, 4519–4534.

Silva-Perez V, Molero G, Serbin SP, Condon AG, Reynolds MP, Furban RT, Evans JR. 2018. Hyperspectral reflectance as a tool to measure biochemical and physiological traits in wheat. Journal of Experimental Botany 69, 483–496.

Smith M-L, Ollinger SV, Martin ME, Aber JD, Hallett RA, Goodale CL. 2002. Direct estimation of aboveground forest productivity through hyperspectral remote sensing of canopy nitrogen. Ecological Applications 12, 1286–1302.

South PF, Cavanagh AP, Liu HW, Ort DR. 2019. Synthetic glycolate metabolism pathways stimulate crop growth and productivity in the field. Science 363, eaat9077.

Spath H. 1985. The cluster dissection and analysis theory fortran programs examples. Prentice-Hall, Inc.

Thenkabail PS, Smith RB, De Pauw E. 2000. Hyperspectral vegetation indices and their relationships with agricultural crop characteristics. Remote Sensing of Environment 71, 158–182.

Thornley JH, Johnson IR. 1990. Plant and crop modelling. Oxford: Clarendon Press.