



## Abstract

The vegetation indices normalized difference vegetation index (NDVI) and photochemical reflectance index (PRI) provide indicators of pigmentation and photosynthetic activity that can be used to model photosynthesis from remote sensing with the light-use efficiency model. To help develop and validate this approach, reliable proximal NDVI and PRI sensors have been needed. We tested new NDVI and PRI sensors, “SRS” sensors recently developed by Decagon Devices, during spring activation of photosynthetic activity in evergreen and deciduous stands. We also evaluated two methods of sensor cross-calibration, one that considered sky conditions (cloud cover) at midday only, and the other that also considered diurnal sun angle effects. Cross-calibration clearly affected sensor agreement with independent measurements, with the best method dependent upon the study aim and time frame (seasonal vs. diurnal). The seasonal patterns of NDVI and PRI differed for evergreen and deciduous species, demonstrating the complementary nature of these two indices. Over the spring season, PRI was most strongly influenced by changing chlorophyll:carotenoid pool sizes, while over the diurnal time scale PRI was most affected by the xanthophyll cycle epoxidation state. This finding demonstrates that the SRS PRI sensors can resolve different processes affecting PRI over different time scales. The advent of small, inexpensive, automated PRI and NDVI sensors offers new ways to explore environmental and physiological constraints on photosynthesis, and may be particularly well-suited for use at flux tower sites. Wider application of automated sensors could lead to improved integration of flux and remote sensing approaches to studying photosynthetic carbon uptake, and could help define the concept of contrasting vegetation optical types.

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# 1 Introduction

The photochemical reflectance index (PRI) was originally derived as a measure of xanthophyll cycle activity determined using proximal remote sensing of leaves and canopies on a diurnal time scale (Gamon et al., 1992, 1997). In this context, the xanthophyll cycle is a facultative response that changes readily as a means of dissipating extra light energy non-destructively (Demmig-Adams and Adams, 1992). Because this xanthophyll response reflects changing light energy distribution within the photosynthetic system, it can provide a useful measure of short-term changes in photosynthetic light-use efficiency, typically expressed as the photosynthetic rate normalized by the incident or absorbed photosynthetically active radiation (Gamon et al., 1992, 1997; Peñuelas et al., 1995). However, studies at larger scales spanning species or seasons often reveal a different story; when sampled at these larger spatial or temporal scales, PRI is strongly influenced by evolving carotenoid:chlorophyll ratios (Sims and Gamon, 2002; Stylinski et al., 2002; Filella et al., 2009; Garrity et al., 2011; Gamon and Berry, 2012; Porcar-Castell et al., 2012; Wong and Gamon, 2014). Unlike the diurnal xanthophyll cycle activity, these pigment pool size adjustments comprise a constitutive response to *chronic* stress, ontogeny, or phenology, determined by more slowly changing physiological states in response to time-integrated environmental conditions. For example, during seasonal transitions from a dormant to an active growth phase, evergreen plants adjust their chlorophyll : carotenoid ratios over many weeks in response to changing temperatures (Adams et al., 2002) and this adjustment can be readily detected by PRI (Stylinski et al., 2002; Filella et al., 2009; Porcar-Castell et al., 2012; Wong and Gamon, 2014). Both the facultative and constitutive PRI responses are strongly correlated with photosynthetic activity, but over different time scales and using different mechanisms, both of which involve photoprotective carotenoid pigments.

The normalized difference vegetation index (NDVI) was developed as a measure of vegetation greenness. Typically, it is used to evaluate seasonal phenology or productivity of vegetation as it changes gradually with the growth and senescence

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of vegetation (Gamon et al., 1995). NDVI is also a common product of many satellite sensors and is widely used for tracking vegetation phenology, and mapping potential photosynthetic activity or productivity (Defries and Townshend, 1994; Running et al., 2004). However, NDVI-based approaches often miss more subtle, short-term responses to stress that can determine how much of the photosynthetic potential is actually realized, particularly for species showing little structural responses to stress. For example, in annual or deciduous canopies, NDVI is highly correlated with morphological changes (green biomass or leaf area index) that affect seasonally changing photosynthetic capacity. In evergreens where canopy structure is relatively stable over the year, NDVI changes little with season, and fails to detect the onset and cessation of photosynthesis early and late in the growing season (Gamon et al., 1995). So, while NDVI is well-suited for detecting photosynthetic potential defined by light absorption and canopy structure, it misses many of the more subtle photosynthetic dynamics arising from alterations in physiological activity (e.g. photosynthetic downregulation during short-term stress). For this, PRI is often a useful counterpart.

When combined, NDVI and PRI can provide complementary information regarding photosynthetic activity. Together, NDVI and PRI can be used to estimate photosynthetic rate, typically using a light-use efficiency model, with NDVI providing a means to estimate light absorption by green vegetation and PRI providing a measure of the efficiency with which that absorbed light is converted to fixed carbon (Gamon and Qiu, 1999; Gamon et al., 2001). When integrated over time (typically a growing season) the photosynthetic rate estimated from vegetation indices can provide a good measure of net primary production, NPP (Goward et al., 1985). Singly or in combination, these two indices provide a means to estimate photosynthetic phenology and activity, with the exact use depending upon the particular application and time frame.

Originally, most field studies employing these vegetation indices were conducted with portable spectrometers capable of measuring reflectance in many wavebands simultaneously. While these instruments represent the “gold standard” for field





1.5 m × 1.5 m arrays comprising two synthetic plant stands, providing closed canopy monocultures for viewing by the sensors.

## 2.2 Optical measurements

The automated sensors consisted of five PRI sensors and five NDVI Spectral Reflectance Sensors (SRS, Decagon Inc, Pullman WA, USA). These were early prototypes of the SRS sensors currently available from Decagon Devices, Inc. The detectors of the prototype PRI sensors were photodiodes paired with interference filters centered at the 532 and 570 nm PRI wavelengths, similar to those used by Garrity et al. (2010). The interference filters have a bandpass of 10 nm at full width half maximum (FWHM). Following Ryu et al. (2010), prototype NDVI sensors used light emitting diodes (LEDs). LEDs had peak sensitivity at 630 and 800 nm with bandpass widths of 50 and 40 nm, respectively. Note that since our study was completed, the manufacturer has changed the NDVI sensor to be based on a photodiode design.

Of the five PRI (“P”) sensors, three were downward-looking sensors (“Pr1”, “Pr2”, and “Pr3”, where “r” indicates radiance), with a field-of-view of approximately 20° full angle, and two were upward-looking hemispherical sensors (“Pi4” and “Pi5”, where “i” indicates irradiance), with a field of view of approximately 180° (full angle). Of the five NDVI (“N”) sensors, three were downward-looking sensors (“Nr1”, “Nr2”, and “Nr3”, where “r” indicates radiance) and two of which were upward-looking sensors (“Ni4” and “Ni5”, where “i” indicates irradiance). The upward-looking hemispherical sensors provided reference values of sky irradiance against which we normalized the downward looking sensor values of canopy radiance using a cross-calibration procedure described in Sect. 2.3.

The ten SRS sensors were positioned above the two plant stands at a height of approximately 2 m above the ground. Due to differences in canopy height, the exact distance between the sensors and the tops of the canopies varied as follows: 45 cm for lodgepole pine, and 50 cm for aspen. In all cases, downward-looking canopy radiance sensors were positioned over the center of the plant stands to avoid possible edge

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effects. The upward-looking sensors were mounted above the middle of the canopies to monitor sky irradiance.

Each sensor was logged every 5 s and expressed as 1 min averages by a datalogger (CR1000, Campbell Scientific, Logan UT, USA). To calculate reflectance, data from each of the three downward-looking PRI and NDVI (radiance) sensors were compared to the average of the coincident measurements made by the two upward-looking PRI and NDVI (irradiance) sensors, respectively. For each waveband, uncorrected reflectance was first calculated by dividing the radiance by the irradiance values:

$$\frac{Pr_{532\text{ nm}}}{Pi_{532\text{ nm}}} \quad (1a)$$

$$\frac{Pr_{570\text{ nm}}}{Pi_{570\text{ nm}}} \quad (1b)$$

$$\frac{Nr_{630\text{ nm}}}{Ni_{630\text{ nm}}} \quad (1c)$$

$$\frac{Nr_{800\text{ nm}}}{Ni_{800\text{ nm}}} \quad (1d)$$

These uncorrected reflectance values ( $Pr/Pi$  and  $Nr/Ni$ ) calculated for each waveband were then used to calculate uncorrected PRI (using Eq. 2) and NDVI (using Eq. 3), respectively.

$$PRI = \frac{Pr/Pi_{532\text{ nm}} - Pr/Pi_{570\text{ nm}}}{Pr/Pi_{532\text{ nm}} + Pr/Pi_{570\text{ nm}}} \quad (2)$$

$$NDVI = \frac{Nr/Ni_{800\text{ nm}} - Nr/Ni_{630\text{ nm}}}{Nr/Ni_{800\text{ nm}} + Nr/Ni_{630\text{ nm}}} \quad (3)$$

Each uncorrected reflectance measurement was further modified by a sensor cross-calibration coefficient (see below), yielding corrected reflectance, and allowing us to evaluate the effect of this coefficient on the PRI or NDVI signals.

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## 2.3 Sensor cross-calibration

Previous studies (Gamon et al., 2006) have illustrated the need for cross-calibration to properly match radiance and irradiance sensor outputs when calculating reflectance from dual-detector (radiance and irradiance) optical sensors. This need arises from the different sensor responses and foreoptics, which must be normalized to yield correct reflectance and index values. In this study, a similar cross-calibration was performed by correcting each radiance sensor against the matching pair of irradiance sensors used for raw reflectance calculation. The cross-calibration procedure involved the insertion of a 25 cm × 25 cm, 99 % reflective white panel (Spectralon, Labsphere Inc., North Sutton, NH, USA) covering the field-of-view of each downward-looking sensor while the upward-looking sensors sampled sky irradiance. For each cross-calibration, the panel was held under the downward-looking sensors at an approximate distance of 20 cm for 5 consecutive minutes and the measurements made during this period were averaged to obtain a single cross-calibration for that sensor at that time for those particular sky conditions (cloud cover and sun angle). This procedure yielded a cross-calibration ratio (“cross calibration”), expressed as  $Pr_{\text{panel}}/Pi_{\text{sky}}$  and  $Nr_{\text{panel}}/Ni_{\text{sky}}$  for each band and sensor pair.

To explore the effect of cloud cover on seasonally changing indices, midday measurements from 11:00 to 15:00 LT (where solar noon was approximately 13:30 LT) were used. These midday cross-calibration ratios were plotted as a function of cloud cover (expressed as the ratio of sun visibility), to evaluate the relationship between the cross-calibration ratios and cloud cover. The sun visibility ratio was calculated by comparing the actual PPFd (PAR irradiance) measured on several dates (9, 15, 28, and 30 May and 4 and 5 June 2013) to a modeled PPFd assuming perfectly sunny conditions. Actual PPFd was sampled with a quantum sensor (model S-LIA-M003, Onset Computer Corporation, Bourne, MA, USA) and datalogger (HOBO U30, Onset Computer Corporation, Bourne, MA, USA). Modeled PPFd was calculated with the Solar Radiation Calculator (SolRad), using the Ryan–Stolzenbach modelled

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18:30 LT. Consequently, corrected reflectance and indices for diurnal experiments used an empirical cross-calibration derived for each sensor using the hourly cross-calibration ratios collected closest in time to that sample, incorporating the combined effects of both sun angle and sky conditions on the cross-calibration. These empirical, whole-day corrections were compared to corrections using midday cross-calibrations (Eqs. 6 and 7) and to “raw” sensor index values uncorrected by cross-calibration (Eqs. 2 and 3). For plotting, the corrected one-minute PRI and NDVI samples were averaged over 15 min (for diurnal studies) or 1 h (for seasonal studies), creating a single value for each time period with the error estimate expressed as the standard error of the mean during that time period.

## 2.4 Spectrometer readings (stand reflectance)

To provide independent measures of PRI and NDVI, stand spectral reflectance was calculated from measurements made by a dual detector spectrometer (UniSpec DC, PP Systems, Amesbury MA, USA). For the downward-looking (radiance) detector, the spectrometer foreoptics consisted of a fiber optic (Uni-684, PP Systems, Amesbury, MA, USA) with a FOV restrictor (Hypo-Tube, PP Systems, Amesbury, MA, USA). This yielded a nominal 20° FOV, but our measurements of the actual FOV yielded estimates closer to 15°, providing a smaller view of the canopy than the SRS sensors having a 20° FOV. For the upward-looking (irradiance) detector, the spectrometer foreoptics consisted of a similar fiber optic (Uni-686, PP Systems, Amesbury MA, USA) with a cosine head (Uni-435, PP Systems, Amesbury MA, USA). For seasonal studies, 12 canopy reflectance spectra were sampled over each plant stand near solar noon at the same height as the SRS sensors, covering the canopy regions sampled by the SRS sensors, and these 12 scans were expressed as averages ( $\pm$  SEM).

Reflectance was calculated by referencing the downward-looking (radiance) detector to the upward-looking (irradiance) detector (calculating uncorrected reflectance), and then correcting this ratio by a cross-calibration procedure using panel measurements similar to that described above (see Gamon et al., 2006 for further details). A standard

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reference panel (Spectralon, LabSphere, North Sutton, NH, USA) was used as the reference for all reflectance and cross-calibration calculations. To facilitate comparison with the index values of the SRS sensors, identical wavelengths were selected (532 and 570 nm for PRI, and 630 and 800 nm for NDVI).

## 2.5 Leaf reflectance

The PRI and NDVI were also measured at the leaf level using a spectrometer (UniSpec SC, PP Systems, Amesbury MA, USA) configured for leaf reflectance measurements. In this configuration, the spectrometer foreoptics consisted of a bifurcated fiber optic (UNI-410, PP Systems, Amesbury MA, USA) equipped with a needle leaf clip (UNI-501, PP Systems, Amesbury MA, USA) to hold the fiber tip at a fixed angle and position relative to the leaf surface. Six plants were randomly selected for leaf reflectance. Five random leaves per plant from the illuminated upper-canopy regions were measured weekly near 13:00 LT (solar noon). Dark measurements and white reference scans (Spectralon, LabSphere, North Sutton, NH, USA) were taken before each sample.

## 2.6 Pigment assays

For pigment assays, leaf tissue samples were collected periodically over the course of the study, during midday (for seasonal experiments), and over the course of a single day (for diurnal experiments). For seasonal studies, leaf samples were collected from the same six plants as the leaf reflectance and 1 cm long segments from each of the six plants were pooled together for each date and analyzed as a single average. For diurnal studies, four plants from the corners of the plot were selected and two leaves from each plant were obtained in 3 cm long segments and analyzed as a single average ( $\pm$ SEM).

For seasonal studies, needle segments were excised within 30 min of leaf reflectance (Sect. 2.5), measured with calipers for diameter, and stored in liquid nitrogen until transferred to a  $-80^{\circ}\text{C}$  freezer for long-term storage. For diurnal studies, needles were

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frozen within a minute of leaf reflectance. To estimate sample area, each segment length was multiplied by the diameter and analyzed with high-performance liquid chromatography (HPLC, 1260 Infinity, Agilent Technologies, Santa Clara, CA, USA). To quantify the concentrations of various carotenoid and chlorophyll pigments, we used the method of Thayer and Björkman (1990). The chlorophyll:carotenoid ratio was calculated as the sum of chlorophyll *a* and *b* divided by the sum of all carotenoids including neoxanthin, violaxanthin (*V*), antheraxanthin (*A*), lutein, zeaxanthin (*Z*) and  $\beta$ -carotene. The epoxidation state (EPS), a measure of xanthophyll cycle activity, was calculated as:

$$\text{EPS} = \frac{V + 0.5A}{V + A + Z} \quad (8)$$

where *V*, *A*, and *Z* represent the area-based concentrations of violaxanthin, antheraxanthin, and zeaxanthin, respectively.

## 3 Results

### 3.1 Cross-calibrations

For each sensor, cross-calibration ratios varied with sun visibility, which ranged from near-zero under heavy cloud conditions to approximately one under sunny conditions (Fig. 1). In cases of sunny conditions with cumulus clouds, sun visibility values typically exceeded one due to the additional skylight reflected from clouds. On average, cross-calibration ratios approximated the theoretical expectation for radiance/irradiance values of  $1/\pi$ , or 0.318 (Monteith, 1973), but clearly varied with sky conditions. Typically, the resulting cross-calibration ratios were higher during sunny than during cloudy conditions and exhibited strong linearly relationships with sun visibility (Fig. 1). From these responses, we derived an empirical equation for each sensor band, enabling automatic correction of the mid-day PRI and NDVI values. These equations were subsequently applied to all midday index calculations involving seasonal trends.

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### 3.2 Seasonal trends: NDVI and PRI during spring recovery

The NDVI and PRI were monitored during spring photosynthetic recovery for both species, illustrating the complementary nature of these two indices (Fig. 2). During this time, air temperature increased from a daily average of approximately 0 °C in late April to approximately 15 °C by early June (not shown). In *P. contorta* (lodgepole pine), PRI showed an initial increase coincident with a period of increasing chlorophyll : carotenoid ratios and photosynthetic activity (Wong and Gamon, 2014; Wong and Gamon, 2015). On the other hand, in the *P. tremuloides* (aspen) canopy, PRI was relatively flat, with the exception of a slight rise in canopy-level PRI in May during leaf expansion, followed by a slight decline toward mid-June as leaves matured. This pattern of PRI rise and fall was more apparent in the spectrometer PRI than in the SRS sensor PRI (Fig. 2c), in part due to the greater short-term dynamics in the SRS sensor values. SRS sensor cross-calibration improved the agreement with spectrometer PRI for *P. contorta*, but slightly decreased the agreement for *P. tremuloides* (different sensor pairs were used for each species).

For the pine species, the NDVI trend was nearly flat, but for the aspen stand it showed a marked increase during initial bud burst and leaf expansion. For the pine, these patterns were consistent across instruments and sampling scale (leaf vs. stand). For the aspen, leaf NDVI showed a relatively flat response during the sudden increase in stand NDVI during leaf expansion in early May. Earlier leaf-level sampling was not possible in the aspen because leaves had not yet emerged from the buds. Unlike the effect on PRI, cross-calibration of SRS sensors yielded little change in the SRS NDVI values.

The strong rise in midday PRI for *P. contorta* during spring photosynthetic activation is shown in more detail, along with midday PPFD and pigment trends (Fig. 3). Because the SRS sensors were not available for the early part of this period, PRI values from a spectrometer were added to the plot to show the full period of spring transition associated with photosynthetic activation (for detailed information on these spring

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physiological changes, see Wong and Gamon, 2014; Wong and Gamon, 2015). In this case, cross-calibration of the SRS PRI improved the agreement with spectrometer PRI (Fig. 3a). The spring rise in PRI was coincident with a rise in chlorophyll:carotenoid pigment ratios but not with the xanthophyll cycle epoxidation state (EPS), which increased about three weeks earlier than either the PRI or pigment ratios (Fig. 3b). The considerable short-term variability in the SRS PRI signal (particularly visible in late-May to early June) was largely attributable to the day-to-day variation in midday PPFD, with PRI declining during sunny days, and rising during cloudy days (Fig. 3a).

### 3.3 Diurnal experiments

Next, we explored the ability of the SRS PRI sensors to resolve diurnal patterns related to xanthophyll cycle activity, as affected by diurnal irradiance. Both xanthophyll cycle EPS and PRI declined towards midday as PPFD increased and recovered in the afternoon as PPFD declined (Fig. 4). At very low sun angles, when PPFD values were low and the sun was sometimes partly obscured by objects near the horizon (before 8 a.m. and after 7 p.m.), PRI values were extremely noisy (indicated by the erratic pattern and large error bars). To test the effect of cross-calibrations on diurnal PRI responses, we first applied midday cross-calibration equations (Fig. 1) using the sun visibility values prevailing during each sample. We also applied empirical cross-calibrations closest in time to each sample, considering both sky conditions and sun angle. The exact PRI pattern was strongly influenced by which of the two cross-calibration methods were applied. The most noticeable effect of the midday cross-calibration was a downward shift in the absolute PRI values, similar to the effect seen in the seasonal PRI patterns for *P. contorta* (Fig. 4c, solid black line). Application of empirical cross-calibrations (using the values closest in time to each sample) further changed the *shape* of the diurnal PRI pattern, leading to a more pronounced dip and recovery in PRI (Fig. 4c, solid red line) that more closely matched the diurnal pattern of the xanthophyll cycle pigment epoxidation state (EPS) (Fig. 4b).

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### 3.4 Comparing PRI to pigments over diurnal and seasonal time scales

To evaluate the cause of PRI variation over diurnal and seasonal time periods, PRI values recorded by the SRS sensors were compared to pigment data (chlorophyll:carotenoid ratios and xanthophyll cycle epoxidation state). Seasonal measurements spanned the period of spring recovery of photosynthesis in *P. contorta* (3 May 2013–4 June 2013) (Fig. 3) and diurnal measurements were collected from a single experiment on 25 July 2013 (Fig. 4).

These comparisons illustrated that over the seasonal time scale, PRI was correlated with the chlorophyll:carotenoid pigment ratios, but not with the xanthophyll cycle EPS (Fig. 5a and b). Time trends showed that spring recovery of EPS occurs 3–4 weeks before the increase of chlorophyll:carotenoid pigment ratios (Fig. 3), and it is these pigment ratios (not EPS) that best corresponded to the spring increase in PRI (Fig. 5). However, the reverse was true over the diurnal time scale, when PRI was clearly correlated with the xanthophyll cycle EPS (Figs. 4 and 5d), and not the chlorophyll:carotenoid pigment ratios (Figs. 4 and 5c). This result was consistent with the similar diurnal patterns of both PRI and EPS, combined with the relatively flat diurnal pattern of the pigment pool ratios (Fig. 4b).

## 4 Discussion

As expected, NDVI and PRI showed complementary behaviour in evergreen and deciduous canopies in early spring. In this sense, evergreens and deciduous species represent distinct optical types, as revealed by their contrasting NDVI and PRI behaviour. Stand-level NDVI increased in deciduous canopies during leaf emergence and expansion in early spring, but not in evergreen canopies that did not add new leaves during this period. By contrast, PRI detected the changing pigment pool sizes during spring in the evergreen stands, and showed relatively little change in the deciduous stands during this period. These complementary patterns emerged

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both with the SRS sensors and with the field spectrometer, and demonstrate that automated NDVI and PRI sensors can provide useful information on the contrasting photosynthetic phenology of evergreen and deciduous species. Particularly intriguing is the ability of PRI to detect changing chlorophyll : carotenoid ratios in the evergreen pine stands. Recent work (Wong and Gamon, 2014, 2015) has shown that these changing pigments (and hence PRI) can provide an indicator of evergreen spring physiological activation, a process that is hard to detect with the eye or with conventional optical remote sensing. On the other hand, NDVI readily captures the changing photosynthetic capacity associated with bud burst and leaf development, but not the less visible evergreen pigment changes during spring. Based on these findings, we would expect that ecosystems from different biomes having contrasting evergreen and deciduous stand composition would show contrasting behaviour of NDVI and PRI (Garbulsky et al., 2011). Due to the lack of suitable sensors, this hypothesis has been hard to test extensively, and the advent of automated NDVI and PRI sensors could now enable such comparative tests across contrasting ecosystems. Since these two indices relate to the two terms of the light-use efficiency model (Gamon and Qiu, 1999), better understanding of their complementary behaviour could help improve the application of the light-use efficiency model for prediction of ecosystem photosynthesis.

In the evergreen lodgepole pine stand, the SRS PRI was clearly affected by two different processes operating over different time scales. Over the diurnal time scale, the SRS PRI followed the changing xanthophyll cycle epoxidation state (EPS), as has been reported before (Gamon et al., 1992). Over a longer time period of several weeks, the midday PRI values followed the changing chlorophyll : carotenoid pool sizes associated with spring photosynthetic activation. Thus, PRI provided a sensitive indicator of pigment changes associated with photosynthetic activity, but in different ways and with different mechanisms. These contrasting mechanisms have been termed the “facultative” (diurnal xanthophyll activity) and “constitutive” (longer-term pigment pool size shifts) responses (Gamon and Berry, 2012; Wong and Gamon, 2014). Recently, Wong and Gamon (2014) reported similar findings for evergreens; during seasonal







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sensor distance and sampling angle accordingly. While not a central part of our study due to the limited number of sensors, further studies should investigate the role of sensor position and sampling angle, as well as the required replication needed to obtain representative samples of stand optical properties. This becomes particularly critical if the goal is to relate proximal optical sampling to larger footprints, as is often the case when validating satellite measurements or comparing to flux tower measurements.

Missing from our short-term study was a full consideration of long-term sensor stability. Temperature stability and ability to withstand moisture are key considerations, particularly if sensors are to be useful over one or more annual cycles, and these factors were not fully considered in our study. Since completion of our experiment, the manufacturer (Decagon Devices) has changed the NDVI sensor from the LED version used in this study to a photodiode design, to attain greater temperature stability. We recommend that additional studies be conducted over a range of environments to more fully test the behaviour, utility, stability and longevity of the SRS sensors. For such studies, the cross-calibration methods described here could be essential, not just for obtaining accurate index values, but also to check and correct for sensor drift and enable proper comparison of values across sites. Ideally, such tests would include ecosystems and biomes with contrasting optical behaviour and environmental constraints on photosynthesis, allowing us to more fully develop the concept of optical types. Our hope is that the initial findings reported here can provide a first step in developing protocols for such a study.

## 5 Conclusions

PRI and NDVI detected complementary processes during spring transition in evergreen and deciduous canopies. As expected, NDVI was primarily sensitive to leaf emergence in deciduous aspen stands, and PRI was sensitive to changing pigment ratios in evergreen pine stands. PRI was also able to detect diurnal changes in xanthophyll



cycle epoxidation state, although the primary cause of PRI increase during spring was the increasing chlorophyll:carotenoid ratio, and not the xanthophyll cycle.

The diurnal and seasonal patterns were clearly sensitive to the method of cross calibration. For each sensor, sun visibility (cloud cover) had a predictable effect on the cross calibration, allowing us to model this correction for each sensor. Determining this response for each sensor should facilitate automated application of optical sensors where regular calibration would not be feasible. On the other hand, due to the combined effects of sun angle and sky conditions, obtaining accurate diurnal responses may require frequent manual calibration that may present challenges for sensor automation.

Automated, low cost NDVI and PRI sensors offer new opportunities for monitoring photosynthetic phenology. We recommend further tests be applied over longer time periods at flux tower sites across a range of ecosystems, with a particular focus on the optical responses of contrasting vegetation types. Such studies would help improve our understanding of the component terms of the light-use efficiency model and could help reveal contrasting controls on carbon flux for different ecosystems.

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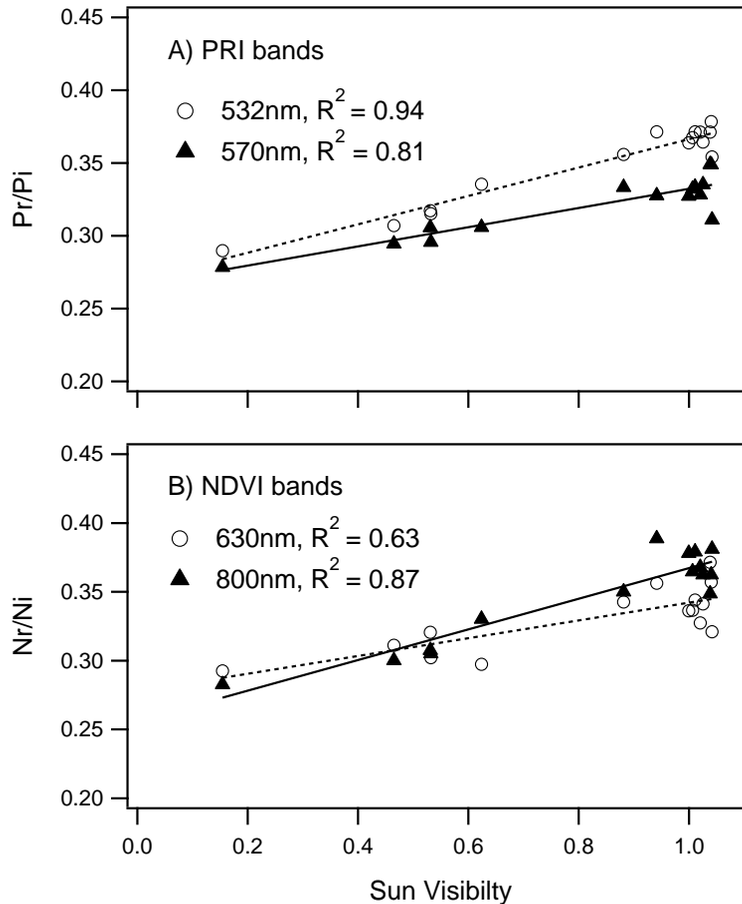
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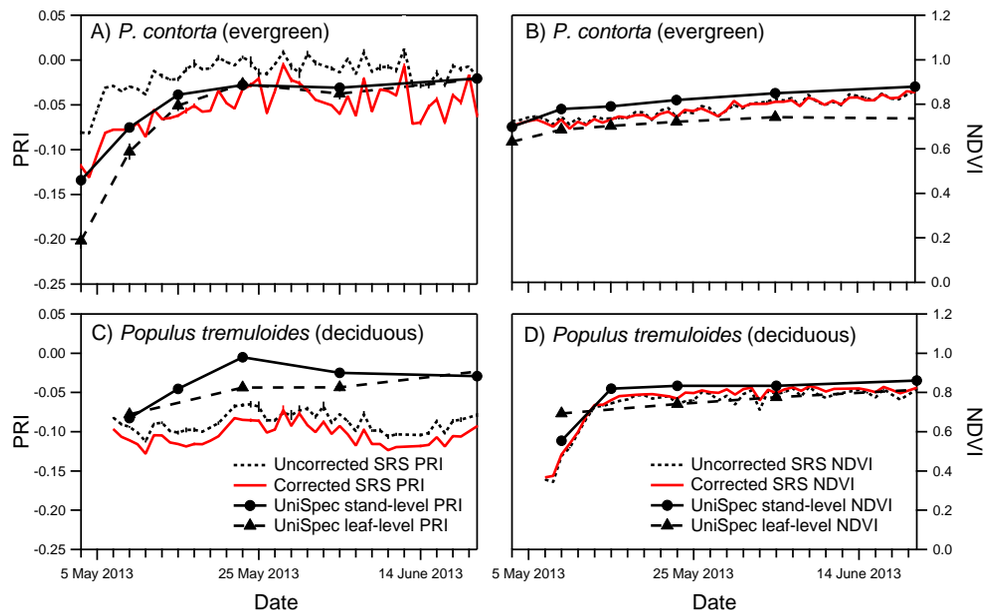




**Figure 1.** Representative cross-calibration ratios for a single sensor set as a function of sun visibility ratios for PRI bands ( $P_R/P_I$ , **a**) and NDVI bands ( $N_R/N_I$ , **b**). Sun visibility: 0 = darkness, 1 = clear, sunny skies, with intermediate values indicating varying degrees of cloudiness.

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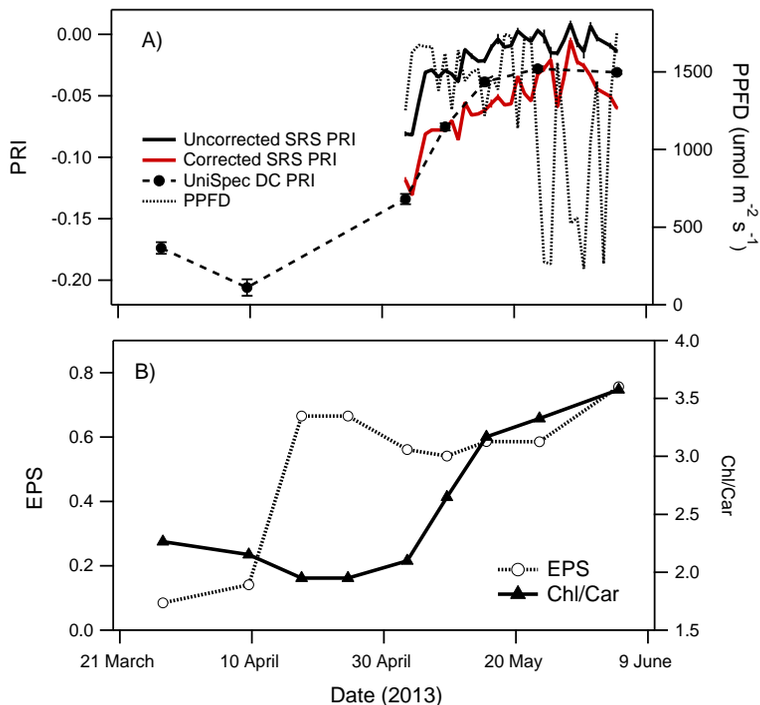
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**Figure 2.** Midday PRI (a and c) and NDVI (b and d) time trends (3 May 2013–21 June 2013) for *Pinus contorta* (lodgepole pine) and *Populus tremuloides*, (trembling aspen) sampled using both SRS sensors and spectrometers at both leaf and stand scales. Aspen bud burst began on 5 May, and full expansion was reached on 16 May.

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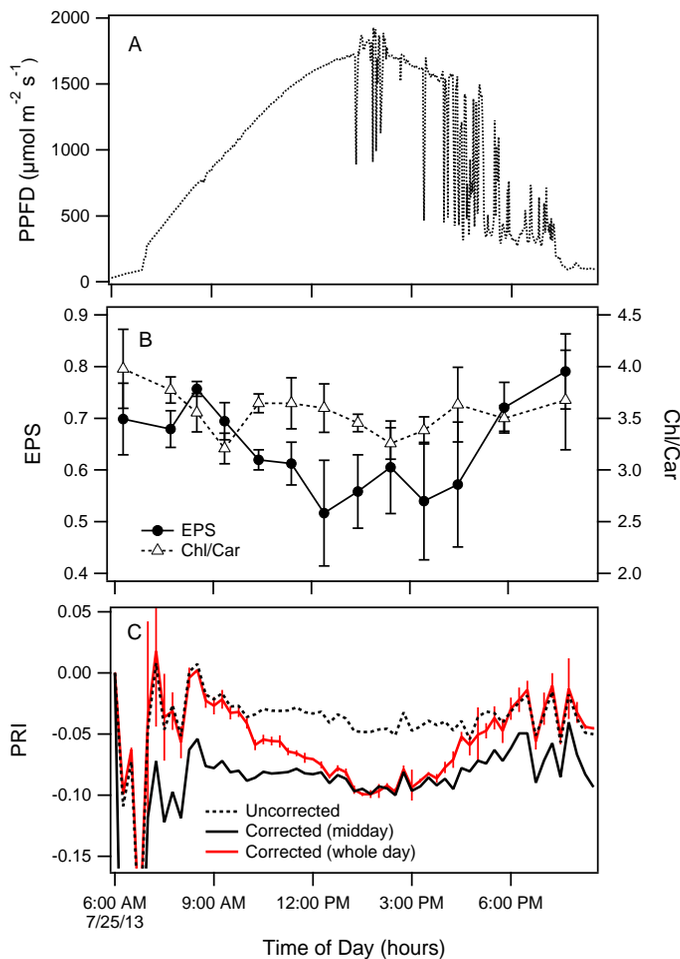
**Figure 3.** Midday PRI and PPFD trends (a) and pigment trends (b) of *P. contorta* during spring photosynthetic activation. The PRI values were measured by the Decagon SRS sensors and a dual-channel spectrometer (UniSpec DC, PP Systems, Amesbury, MA). Corrected PRI (mid-day correction) were produced by applying the empirical mid-day cross calibration equations, derived from linear trends (Fig. 1). PRI error bars are standard error of the mean. Chl / Car ratios and xanthophyll cycle epoxidation state (EPS) were single values with no error bars (see methods).

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**Figure 4. (a)** PAR irradiance (PPFD) over the course of the day (25 July 2013). **(b)** Pigment values (epoxidation state and chlorophyll:carotenoid pigment ratios) over the course of the day. **(c)** Uncorrected and corrected PRI values plotted at 15 min intervals over the course of the day. Uncorrected PRI values were calculated using Eq. (2). Corrected PRI data (mid-day correction) were calculated by applying the empirical mid-day cross calibrations (Fig. 1), where the sun visibility (cloud cover) is taken into consideration, but not the sun angle. Corrected PRI data (whole day correction) were calculated using the hourly white panel ( $P_r/P_i$ ) ratios obtained throughout the day, using the ratio nearest in time. Error bars are  $\pm 1$  SEM for EPS, Chl / Car, and PRI. For clarity, only the error bars for the corrected (whole day) PRI are shown.

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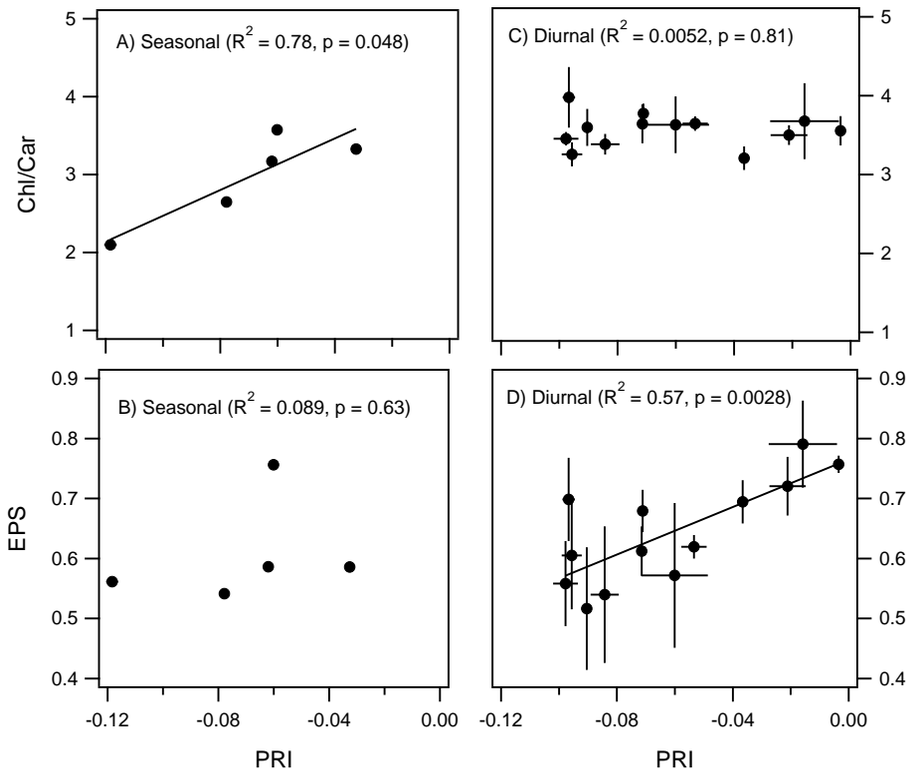
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**Figure 5.** Corrected PRI vs. pigment measures (xanthophyll cycle epoxidation state, EPS, or chlorophyll : carotenoid ratio, Chl / Car). Seasonal data (**a**, **b**) spans the spring recovery period (3 May 2013–4 June 2013; see Fig. 3). Diurnal data (**c**, **d**) are from 25 July 2013 (see Fig. 4). Error bars indicate  $\pm$ SEM for PRI and diurnal pigment data. Linear regressions are shown for significant ( $p < 0.05$ ) fits only.