THE PHOTOCHEMICAL REFLECTANCE INDEX FROM DIRECTIONAL CORNFIELD REFLECTANCES: OBSERVATIONS AND SIMULATIONS

[Research Paper]

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1 ABSTRACT

2 The two-layer Markov chain Analytical Canopy Reflectance Model (ACRM) was linked with *in situ* hyperspectral leaf optical properties to simulate the Photochemical Reflectance Index 3 4 (PRI) for a corn crop canopy at three different growth stages. This is an extended study after a successful demonstration of PRI simulations for a cornfield previously conducted at an early 5 vegetative growth stage. Consistent with previous in situ studies, sunlit leaves exhibited lower 6 PRI values than shaded leaves. Since sunlit (shaded) foliage dominates the canopy in the 7 reflectance hotspot (coldspot), the canopy PRI derived from field hyperspectral observations 8 9 displayed sensitivity to both view zenith angle and relative azimuth angle at all growth stages. Consequently, sunlit and shaded canopy sectors were most differentiated when viewed along the 10 azimuth matching the solar principal plane. These directional PRI responses associated with 11 12 sunlit/shaded foliage were successfully reproduced by the ACRM. As before, the simulated PRI values from the current study were closer to *in situ* values when both sunlit and shaded leaves 13 were utilized as model input data in a two-layer mode, instead of a one-layer mode with sunlit 14 leaves only. Model performance as judged by correlation between *in situ* and simulated values 15 was strongest for the mature corn crop (r = 0.87, RMSE = 0.0048), followed by the early 16 vegetative stage (r = 0.78; RMSE = 0.0051) and the early senescent stage (r = 0.65; RMSE = 17 0.0104). Since the benefit of including shaded leaves in the scheme varied across different 18 growth stages, a further analysis was conducted to investigate how variable fractions of 19 sunlit/shaded leaves affect the canopy PRI values expected for a cornfield, with implications for 20 remote sensing monitoring options. Simulations of the sunlit to shaded canopy ratio near 50/50 21 \pm 10 (e.g., 60/40) matching field observations at all growth stages were examined. Our results 22

23	suggest in the importance of the sunlit/shaded fraction and canopy structure in understanding and					
24	interpreting PRI.					
25						
26	Highlights:					
27	> Demonstrating PRI responses to illumination conditions and viewing geometry at least					
28	and canopy level.					
29	Validating the capability of the two-layer Analytical Canopy Reflectance Model for PRI					
30	simulations in a cornfield at different growth stages.					
31	> Investigating how canopy structure associated with variable fraction of sunlit/shaded					
32	leaves affect the PRI values.					
33						
34	Keywords: hyperspectral, two-layer Analytical Canopy Reflectance Model (ACRM),					
35	photochemical reflectance index (PRI), cornfield					
36						

37 1. INTRODUCTION

38 Remotely sensed spectral bio-indicators have the potential to play a critical role in monitoring and modeling processes in time and space for our Earth's ecosystems, including the 39 exchange of carbon between the biosphere and the atmosphere. This is because uncertainties 40 exist in how ecosystems will function and what feedbacks to expect, especially under 41 disturbances induced by the changing climate (Garbulsky et al., 2011; Middleton et al., 2011). 42 One of the widely used concepts to model carbon assimilation by plants is the light use 43 efficiency (LUE) model (Monteith, 1972; Monteith, 1977). This approach describes carbon 44 assimilation, in the form of gross or net primary productivity (GPP, NPP), as the product of the 45 absorbed photosynthetically active radiation (APAR) and LUE. Previous studies have shown 46 47 that LUE can vary based on vegetation type, environmental conditions, and temporal resolution of the observations (Anderson et al., 2000; Garbulsky et al., 2011; Gower et al., 1999; King et 48 49 al., 2011; Middleton et al., 2011; Peñuelas et al., 2011).

The importance of accurate LUE estimation has been emphasized in recent studies (e.g., 50 (Lin et al., 2011; Peñuelas et al., 2011), reporting that errors in LUE are a major contributor to 51 52 biases in annual carbon assimilation estimates. Current tools and methods developed for LUE estimation usually utilize a look-up table of maximum possible LUE, which is then downscaled 53 by an adjustment coefficient determined using meteorological data (e.g., air temperature and 54 VPD) to account for non-optimal environmental effects (Law and Waring, 1994; Mahadevan et 55 al., 2008; Prince and Goward, 1995; Xiao et al., 2004). This approach is used for the satellite 56 57 data product available from the Terra and Aqua Moderate Resolution Imaging Spectroradiometers (MODIS), the MOD17 GPP product (Heinsch et al., 2003; Heinsch et al., 58 2006). However, with this approach, errors are usually introduced into LUE estimates due to 59

uncertainties about the fixed values within the look-up table and the meteorological data used for
scaling factors. Moreover, these meteorological data usually have a much larger footprint than
the area of interest, and hence, are not always representative for local LUE (Middleton et al.,
2011). On the other hand, a spectral bio-indicator directly derived from vegetation optical
properties has been shown capable of providing useful estimates of LUE without needing
ancillary information or relying on meteorological data (Garbulsky et al., 2011; Hall et al., 2011;
Huemmrich et al., 2009; Middleton et al., 2009; Middleton et al., 2011; Peñuelas et al., 2011).

The LUE of plants is closely linked to the reversible photoprotective responses of the 67 68 foliar xanthophyll pigment cycle to illumination conditions, especially as induced by saturating mid-day irradiances. These responses are expressed by a spectral bio-indicator, the 69 Photochemical Reflectance Index (PRI; (Gamon et al., 1990; Gamon et al., 1992; Gamon et al., 70 71 1997; Peñuelas et al., 1995). This PRI information can be used to model the down-regulation of photosynthesis (Demmig-Adams and Adams III, 1996). The PRI utilizes 72 a narrow physiologically active green band centered at 531 nm and a reference band most typically 73 centered at 570 nm, in the form of a normalized difference index (i.e., [p531-74 ρ 570]/[ρ 531+ ρ 570]). The PRI has been increasingly used and examined for its correlation with 75 LUE across various vegetation types and scales (Cheng et al., 2009; Coops et al., 2010; Filella et 76 al., 1996; Gamon et al., 1993; Gamon et al., 1992; Gamon et al., 1997; Garbulsky et al., 2011; 77 Garbulsky et al., 2008; Hall et al., 2011, 2012; Hilker et al., 2011; Hilker et al., 2012; Inoue et 78 al., 2008; Middleton et al., 2009; Middleton et al., 2011; Nichol et al., 2002; Peñuelas et al., 79 1995; Peñuelas and Inoue, 2000; Peñuelas et al., 1997). 80

However, studies have also shown that various factors affect the remote sensing-based
PRI:LUE relationship at canopy or ecosystem scales, including viewing geometry, canopy

structure, leaf area index (LAI), soil background, pigment content and shadow fraction (Barton 83 and North, 2001; Cheng et al., 2009; Drolet et al., 2005; Gamon et al., 2001; Hall et al., 2008; 84 Hernández-Clemente et al., 2011; Hilker et al., 2008a; Hilker et al., 2008b; Middleton et al., 85 2009; Nichol and Grace, 2010; Sims and Gamon, 2002; Sims et al., 2006; Stylinski et al., 2002). 86 Furthermore, previous studies have also shown the importance of taking both sunlit and shaded 87 foliage into account to explain PRI behaviors at the canopy level, since sunlit foliage is more 88 likely to experience high light-induced environmental stress, and to have lower LUE, and hence, 89 lower PRI values (Cheng et al., 2009; Cheng et al., 2010; Hall et al., 2008; Hilker et al., 2008b; 90 91 Middleton et al., 2009; Peñuelas et al., 1995). It follows that we must have more understanding about the relative roles of sunlit and shaded foliage in canopies, and associated canopy structure, 92 to improve our knowledge regarding PRI:LUE relationships. Radiative Transfer (RT) models 93 provide a powerful tool to study this topic since they are designed to quantitatively examine 94 changes in vegetation optical properties with leaf biochemical and canopy biophysical properties 95 (Cheng et al., 2006; Jacquemoud et al., 1996; Verhoef, 1984; Zarco-Tejada et al., 2003; Zhang et 96 al., 2011). 97

Canopy PRI was studied for water stress detection using the PROSPECT leaf model 98 linked with the SAILh and FLIGHT canopy RT models to produce a non-stressed version of the 99 PRI in two tree-structured orchards and a maize field (Suárez et al., 2009). In a more recent 100 study, the leaf model LIBERTY was coupled with the canopy model INFORM to study PRI as a 101 physiological stress indicator in conifer forests (Hernández-Clemente et al., 2011). Both studies 102 focused on PRI acquired at near nadir angles. Cheng et al. (2010) utilized in situ leaf optical 103 properties coupled with a Markov chain Analytical two-layer Canopy Reflectance Model 104 105 (ACRM; (Kuusk, 1995a, b, 2001) to simulate nadir and directional PRI at the canopy level in a

106 cornfield, which was compared and validated with *in situ* canopy PRI observations. That study 107 showed that ACRM successfully simulated PRI under various viewing geometries for a corn crop in the early vegetative stage without noticeable environmental stressors present, and 108 109 explored how several canopy structure parameters affected PRI values. The ACRM-simulated 110 PRI showed the best agreement with *in situ* values when the model was run in a two layer 111 simulation mode, using leaf optical properties from sunlit leaves as the upper layer and shaded leaves as the lower canopy (Cheng et al., 2010). In the current study, we took a step further to 112 examine the robustness of the same algorithm to simulate PRI through three different growth 113 114 stages for a corn crop, examining early vegetative, fully mature, and senescent canopies. Our objective was to determine whether the directional PRI responses previously observed for a 115 young, vigorous canopy also continue to be present throughout the growing season, and to 116 117 characterize and evaluate them. We also investigated how the vertical distribution of sunlit and shaded leaves affect an important structure-related variable, the canopy sunlit/shaded foliage 118 119 ratio, and associated canopy PRI values.

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121 **2.** Methods

122 2.1. Study Site and Field Data Collection

During the summer of 2010, field campaigns were conducted on a corn crop (*Zea mays* L.) in an experimental cornfield at the Optimizing Production Inputs for Economic and Environmental Enhancement (OPE3) site (39.0304°N, 76.8458°W) maintained by the USDA Beltsville Agricultural Research Center (BARC) in Beltsville, Maryland, U.S.A. Measurements were acquired on three dates representing three different growth stages: an early vegetated canopy when plants had nine fully expanded leaves (V9) and were ~1 m tall on 07/01; a fully 129 mature canopy having 13-15 fully expanded leaves at ~ 2 m tall in the early reproductive phase 130 (VT) on 07/15; and an early senescent crop (~2 m tall) at the advanced reproductive development stage (R4) on 08/09. Canopy and leaf level measurements were taken along a 100-m north-south 131 direction transect in the middle of the field to minimize disturbance and to maintain 132 representativeness of the data. Hyperspectral reflectance (~1.5 nm Full Width Half Maximum; 133 134 FWHM) was obtained for vegetation at both leaf and canopy levels and on bare soil using an USB4000 Miniature Fiber Optic Spectrometer (Ocean Optics Inc., Dunedin, FL, USA) with a 135 bare fiber. In situ leaf reflectance observations were acquired directly adjacent to the adaxial leaf 136 137 surfaces. The leaves were excised on the next day, and a Li-Cor 1800-12 integrating sphere (Li-Cor, Lincoln, NE, USA) paired with a spectroradiometer (FieldSpec, ASD Inc., Boulder, CO, 138 139 USA) was utilized to determine transmittance from the leaf adaxial surfaces in the laboratory. At 140 the canopy level, reflectance spectra were acquired at eight different relative azimuth angles (ψ , 0° to 315° relative to the sun, at 45° increments) coupled with three different view zenith angles 141 $(\theta_{\rm v})$ -- 30°, 45°, 60° (obtained at 1.3, 0.75, 0.44 m above the canopy, respectively, to provide a 142 consistent center of the field of view). Nadir ($\theta_v = 0^\circ$; $\psi = 0^\circ$) observations were acquired above 143 144 the canopy at a height of approximately 1 m. This was accomplished by placing the fiber optics from a height-adjustable pole-mount, where a custom-made fixture was designed to position the 145 146 instrument at a desired view zenith angle and relative azimuth angle. Soil background reflectance was taken on bare soil also approximately 1 m above the surface at nadir. Measurements were 147 148 taken between local time 9 am to 4 pm, during which the solar zenith angle (θ_s) varied between 16.6° and 51.2° across the season. Crop LAI was also measured with a Li-Cor LAI-2000 plant 149 canopy analyzer (Li-Cor, Lincoln, NE, USA). More detailed information regarding field data 150 151 collections can be found in Cheng et al. (2010).

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2.2. Models and Simulation Methods

In this study, the canopy model ACRM (Kuusk, 1995a, b, 2001) was utilized to simulate 156 This RT canopy model is equipped with an enhanced Markov chain bidirectional gap 157 PRI. probability function that has been utilized in various studies using forward and inversion modes 158 to validate and/or to estimate plants biochemical properties at leaf and/or canopy level (Cheng et 159 al., 2010; Cheng et al., 2006; Fang et al., 2003; Houborg et al., 2009; Houborg et al., 2011). In 160 161 our previous study, in which ACRM successfully simulated PRI for a young corn canopy, ACRM was set to run in its forward mode utilizing in situ leaf and soil background spectra to 162 simulate canopy spectra at various viewing geometry (θ_{v} , ψ) configurations (Cheng et al., 163 2010). PRI was derived from the ACRM-simulated canopy reflectance spectra, and compared 164 165 with the PRI derived from *in situ* canopy reflectance spectra. In this study, we followed the 166 procedures presented in Cheng et al. (2010; 2011) and ran the model in two different modes: (i) with optical properties of sunlit leaves (only) in a single layer, or (ii) with both sunlit and shaded 167 168 leaves in two layers, where the shaded layer laid below the sunlit layer. PRI was then derived 169 from ACRM-simulated spectra and compared, as before, with PRI derived from *in situ* canopy 170 reflectance spectra for validation. Values of other essential input parameters for the model are 171 summarized in Table 1. These values came from either ancillary field measurements (e.g., LAI) 172 or were decided based on previous studies (Cheng et al., 2010; Cheng et al., 2006; Fang et al., 2003; Houborg et al., 2009; Jacquemoud, 1993; Kuusk, 2001; Zarco-Tejada et al., 2003). 173

174 In Cheng et al. (2010: 2011), a sensitivity analysis was performed on several canopy structure parameters to investigate their effects on PRI simulations. The important influence of 175 LAI on PRI simulations was reported. This study extends our progress and investigates how the 176 vertical distribution and partitioning of LAI between the sunlit upper and the shaded lower 177 canopy layers affect PRI simulations. In our earlier studies, when ACRM was set to run in the 178 one layer mode, the LAI of the upper canopy was assumed to represent the total LAI, or 100%, 179 such that the LAI fractions in upper/lower layers were 100% and 0% (i.e., 100/0). Likewise, 180 when ACRM was set to run in the two layer mode, the LAI fractions of the upper and lower 181 layers were assumed to equal half of the total LAI, (i.e., 50/50). In the current study, a sensitivity 182 analysis was performed by changing the ratio of sunlit upper/shaded lower layer LAI values in 183 10% increments from 100/0 (fully sunlit) to 10/90 (mostly shaded). 184

185

186 **3. RESULTS**

187 3.1. In situ Leaf and Canopy Observations

Leaf-level PRI values derived from *in situ* leaf reflectance are summarized in Figure 1 as 188 mean \pm standard error (SE). PRI for sunlit leaves consistently exhibited lower values than shaded 189 190 leaves on all three dates (ANOVA; n=60 for each day; 07/01, p<0.0001; 07/15, p=0.0003; 08/09, 191 p=0.001). Average PRI values varied from -0.009 to +0.005 for sunlit leaves, and consistently exhibited negative values in the afternoons throughout the growing season. In contrast, mean 192 PRI values for shaded leaves were always positive, varying from +0.002 to +0.022. PRI also 193 194 showed higher mean values in the morning (AM) than in the afternoon (PM) on the two dates dominated by green foliage before senescence, especially for sunlit leaves (n=30, p<0.001 for 195 both days). Among the three growth stages, PRI values were significantly higher for shaded 196

leaves (AM and PM) and sunlit leaves (AM) in the mature VT canopy (July 15) than on either
the early (V9) or later season (R4) growth stages (n=30, p<0.0001). At senescence, no clear
differences were observed between the morning and the afternoon (n=30, p=0.6 for sunlit;
p=0.48 for shaded leaves) but the pattern of higher PRI values for shaded vs. sunlit leaves was
maintained.

At the canopy level, in situ PRI values were plotted as mean \pm SE against viewing 202 geometry (θ_v, ψ) for the three growth stages in Figure 2. The pattern obtained at leaf level (Fig. 203 1) for higher PRI at the VT stage (July 15) was maintained at the canopy level (-0.02 to +0.01) 204 when viewed over a range of view angles (θ_v , 0° , 30° , 45° , and 60°). Lower PRI values (-0.03 to 205 -0.01) occurred at both early (Fig.2, 07/01) and late stages (Fig.2, 08/09), which were similar in 206 their PRI responses at the smaller view zenith angles $(0^{\circ}, n=48, p=0.44; 30^{\circ}, n=128, p=0.16; 45^{\circ}, n=128$ 207 208 n=128, p=0.06) as compared to mid-season VT stage. However, early and late growth stages were differentiated by PRI values obtained at the extreme view, $\theta_v = 60^\circ$ (early > late, n= 128, 209 p=0.0011). PRI values at all azimuth positions increased as a function of θ_v . For example, 210 increases in the mean PRI at the coldspot ($\psi = 180^{\circ}$) for the VT canopy were: -0.003±0.005 at 211 212 30° , 0.0 ± 0.003 at 45° , and $+0.009\pm0.006$ at 60° . This contrasts with the negative PRI obtained at nadir (-0.02±0.003), which would be interpreted as indicating greater physiological stress than 213 was determined at any other view. The general pattern exhibited for all measurement geometries 214 $(8 \ \psi \text{ at } 3 \ \theta_v)$ was for PRI values to be lower when θ_v was close to 0° and highest when $\psi \approx 180^\circ$ 215 at any θ_v , highlighting the dependence on viewing geometries. Together, these results (Figs. 1, 216 2) demonstrate the influence of diurnal and directional effects on PRI values retrieved from a 217 218 cornfield.

219 3.2. ACRM-simulated PRI

220 Reflectance spectra were simulated as output from the ACRM. PRI values were calculated from those and compared with in situ PRI for validation purposes in Figure 3, where 221 PRI values from both field measurements and simulations were plotted against θ_v and ψ (Fig. 3 222 a,c,e). ACRM-simulated PRI successfully captured the responses that *in situ* PRI exhibited to θ_v 223 and ψ (Fig. 3 a,c,e), producing lower values when ψ was close to 0° and higher values when ψ 224 was close to 180° at all θ_v on all three observation days. When the simulations were performed 225 226 with sunlit leaves only, considerable underestimations as compared to *in situ* PRI values were 227 observed (Fig. 3 a,c,e). The PRI underestimations were more pronounced on the young V9 crop (July 1) and the senescent R4 crop (August 9), but also occurred at smaller $\theta_v = 30^\circ$ for the 228 mature VT crop. On the contrary, the differences between simulated and *in situ* values were 229 230 much smaller, and in most cases not significant, when the simulation included both sunlit and shaded leaves (Figure 3a,c,e). Correlations between simulated vs. measured values are presented 231 232 in Fig. 3b,d,f (panels on the right). The fully mature VT canopy (July 15, Fig. 3d) exhibited the highest correlation between in situ values and simulations under both scenarios: when both 233 sunlit and shaded leaves were included (r = 0.87) and when only sunlit leaves were used (r = 0.87)234 0.84). The comparisons for the V9 canopy were also strong (r = 0.78, both sunlit and shaded 235 leaves; r = 0.80, sunlit leaves only), although the sunlit (only) set is clearly offset from the 1:1 236 line. Results were weaker in the senescent R4 stage (r = 0.65, both sunlit and shaded leaves; r =237 0.52, sunlit leaves only), with high variability-- especially for the sunlit dataset. Therefore, 238 simulated PRI using both sunlit and shaded leaves (Fig. 3) yielded better correspondence (closer 239 240 to the 1:1 line) on all three dates than simulations with only sunlit leaves. Statistics of comparisons between *in situ* and simulated PRI are summarized in Figure 4. For all three days, 241

when compared with *in situ* values, simulated PRI using both sunlit and shaded leaves (Fig. 4) generated significantly smaller root mean square error (RMSE) than simulations with sunlit leaves only (Fig. 4). Among the three days, simulated PRI showed the best agreement with *in situ* values at the VT stage, as evidenced by higher correlation coefficients (≥ 0.84) and smaller RMSEs (≤ 0.0096) when the canopy was mature (Fig. 4).

The performance of ACRM-simulated PRI was further examined by calculating the 247 difference from *in situ* values, which is summarized in Figure 5. The thick black line displayed in 248 Fig. 5 indicates no (zero) difference between *in situ* and simulated values. Simulations performed 249 250 using only sunlit leaves produced underestimations of field values, as shown by negative values (Fig. 5a). The largest underestimates (~ 0.03) were obtained at the two smaller view angles, nadir 251 (0°) and $\theta_{v}=30^{\circ}$. The only simulations with sunlit leaves alone that agreed with field 252 measurements occurred for $\theta_v = 60^\circ$ in the forward scattering direction ($\psi = 135^\circ$ to 270°), for the 253 fully green and mature mid-season VT canopy (Fig. 5a). When both sunlit and shaded leaves 254 were used in simulations, the differences were much closer to zero (Fig. 5b), providing better 255 256 agreements with in situ values under all observation and growth conditions. However, simulations done using both sunlit and shaded leaves for the VT canopy showed a small positive 257 258 bias for part of the ψ range at all θ_v , whereas the differences appeared to scatter around zero at the other two growth stages (Fig. 5b). 259

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3.3. Canopy Structure and PRI Simulations

The importance of taking optical properties of both sunlit and shaded leaves into account, as well as their relative proportions in the canopy, expressed as a canopy-level ratio, was further investigated using the mid-season VT canopy data. Figure 6 shows how the ACRM-simulated PRI values changed as a function of the sunlit/shaded canopy ratio and viewing geometry, for a 265 corn crop having LAI = 2.48. Columns with lighter shading indicate higher sunlit/shaded 266 fractions. In the modeling scheme, a higher sunlit/shaded canopy ratio describes a canopy that is dominated by sunlit leaves, and therefore, optical properties of sunlit leaves influence the 267 simulated canopy reflectances significantly more. On the contrary, a darker tone (Fig. 6) 268 indicates lower sunlit/shaded canopy ratios were used to simulate situations where shaded leaves 269 270 contributed more to the total canopy reflectance. Clearly, the PRI values obtained at any θ_v and ψ decrease as the sunlit/shaded canopy ratio favors more sunlit foliage (Fig. 6). A lower PRI 271 value would indicate greater environmental stress, and reduced LUE. 272 Therefore, for the same canopy LAI and growth stage, different inferences about LUE could be made based on the 273 observed PRI, depending on viewing geometry and the inherent canopy structure profile. These 274 results help explain why there have been so many confounding factors that influence the PRI of 275 276 canopies that have been reported by various researchers.

These simulations were also directly compared with *in situ* PRI values (Figure 7), to 277 reveal a linear shift away from the 1:1 line for the extreme cases, and displaying a general 278 279 underestimation for the full sun (100/0) case, especially at lower PRI values, and a general overestimation for the mostly shaded (20/80) case. Statistics for the correlation coefficients and 280 281 RMSEs are summarized in Figure 8, indicating the highest correlations paired with the lowest RMSE were associated with two groups in the mid-range (70/30, 60/40). The slope and offset of 282 the regression lines (Figure 9), show a consistent decline for the slope (parameter "a") as the 283 sunlit/shaded canopy ratio increasingly favored more shaded foliage, whereas the offset 284 (parameter "b") increased. Since the best agreement between simulations and *in situ* observations 285 was achieved for a 60/40 ratio, we can assume our field measurements were acquired at or near 286

the 60/40 sunlit/shaded canopy conditions. Thus, our original assumption of a 50/50 ratio was
not the optimal condition for the mature VT canopy in 2010.

289

290 4. Discussion

The PRI was developed to track the reversible changes in the photoprotective xanthophyll 291 292 cycle induced by light intensity changes through a diurnal cycle, (Gamon et al., 1992; Peñuelas et al., 1995). Subsequently, additional environmental stresses have been shown to influence the 293 pH of the chloroplast stroma, affecting the xanthophylls cycle and associated PRI values, such as 294 drought and cold temperatures (Demmig-Adams and Adams, 2000; Müller et al., 2001; Pfündel 295 and Bilger, 1994). Studies have also shown correlations between PRI and other 296 297 physiological/morphological changes, for instance, the carotenoids and chlorophyll ratio (Filella et al., 2004; Sims and Gamon, 2002). In our previous study, we successfully demonstrated that in 298 situ leaf optical properties coupled with ACRM could simulate PRI for a young, homogeneous 299 300 corn canopy, still growing and in the vegetative growth stage. Here, we extended our study to simulate PRI for a corn crop during three different growth stages during the 2010 growing 301 302 season.

303 4.1. In situ PRI at Leaf and Canopy Level

4.1. In suu I KI ui Leuj und Canopy Levei

First of all, the results presented here confirm our previous studies showing that shaded leaves captured in the coldspot of canopy directional reflectances have higher PRI values than sunlit leaves (Cheng et al., 2010; Gamon et al., 1990; Middleton et al., 2009; Peñuelas et al., 1995), and we extended those observations to examine the PRI responses through a growing season in the same experimental cornfield location as the previous study. Higher PRI values in 309 shaded foliage and canopy sectors indicate that the intensity of xanthophyll-regulated 310 photoprotection is lower than in sunlit leaves and canopy segments which are more likely to 311 experience high light stress and exhibit lower PRI values. Furthermore, previous studies have 312 shown correlations between leaf pigments (e.g. carotenoids/chlorophyll ratio) and PRI and the 313 changes in PRI values could be related to leaf development and aging during the growing season 314 (Garbulsky et al., 2011; Peñuelas et al., 2011).

Leaf level PRI for the "green" canopies (i.e., the V9 through VT growth stages) exhibited 315 lower PRI values (i.e., greater stress) during afternoons after several hours of high irradiance 316 exposure, than for mornings (Fig. 1). This pattern was also observed in 317 our V10 dataset acquired in the same field in 2008 (Cheng et al., (2010), although the 2008 values were much 318 higher, implying lower relative stress responses, which were very likely due to an abnormally 319 320 wet spring that year. Higher PRI values were also obtained in shaded leaves vs. sunlit leaves in both morning and afternoon observations in the chlorophyll-dominated growth stages. For the 321 senescent canopy, however, lower PRI values were found in shaded and sunlit leaves all day 322 323 These lower mean PRI values for the senescent growth stage discriminated between long. shaded and sunlit (shaded > sunlit) but not AM vs. PM due to high variability (Fig. 1). 324 Nevertheless, the importance of AM vs. PM observations in studying PRI vs. LUE or using PRI 325 to determine LUE needs to be emphasized. The daily PRI averages followed the expected 326 pattern: VT > V9 > R4, but the daily variation became large, as compared with either morning 327 328 or afternoon observations. On the other hand, considerable error in estimating daily PRI values would be incurred if only sunlit foliage was considered on any of the dates examined, but 329 especially for the mature, mid-season VT crop that had the largest sunlit vs. shaded PRI 330

difference. These findings should serve as a caution when utilizing daily average PRI values inmodel simulations of LUE at the ecosystem scale.

The canopy PRI observations showed substantial dependence on viewing geometry 333 (Fig. 2), similar to results reported in our previous study (Cheng et al., 2010). PRI values were 334 higher when the canopy was viewed at larger, oblique θ_v since more shaded foliage and less soil 335 336 background contamination was captured. Secondly, PRI exhibited higher values when ψ was at the coldspot, close to 180° where the shaded dominated the field of view, and lower values when 337 ψ was close to the hotspot at 0° (broadly including 45°, 315°) where it was associated with the 338 sunlit segment of the canopy (Fig. 2). This is a consistent pattern that has been observed in 339 multiple years under different conditions for the cornfield and in forests (Hall et al., 2008; Hilker 340 341 et al., 2008b; Huemmrich et al., 2009; Middleton et al., 2009). When utilizing spaceborne data, 342 the observations are not always acquired at nadir (e.g., EO-1 Hyperion, Terra/Aqua MODIS). 343 Therefore, this confounding effect needs to be addressed to retrieve meaningful information of plant physiological conditions from non-nadir as well as nadir PRI values. Previous studies 344 345 conducted at a Douglas fir forest in British Columbia, Canada, also reported that PRI exhibited 346 similar dependency to viewing geometry (Hall et al., 2008; Hilker et al., 2008b; Middleton et al., 2009), and are supported by a recent satellite study using off-nadir directional observations (Hall 347 348 et al., 2011; Hilker et al., 2011).

PRI values at both leaf and canopy levels expressed less variance (e.g., smaller SE) in the young, homogeneous, unstressed V9 canopy (Figs. 1,2), with more variability accruing through the season as the crop aged, weathered, and experienced various unfavorable environmental conditions.

353 4.2. Simulation Performance and Differences in Previous Study

ACRM has been shown to successfully simulate canopy PRI values and their 354 dependency on viewing geometry with the current 2010 data and with the previous 2008 data. 355 ACRM was able to deliver believable simulations when both sunlit and shaded leaves were used 356 (Fig. 3). However, when only sunlit leaves were included in the process, less agreement with 357 358 field measurements was achieved and comparisons to *in situ* measurements produced higher RMSEs due to underestimation. The agreement with *in situ* PRI values was better when the corn 359 crop was dominated by green foliage from the actively growing, early vegetative through the 360 361 mature, reproductive growth stages. When the corn crop approached the senescent stage, ACRM simulation was satisfactory, but agreed with *in situ* values the least well among our datasets. The 362 early senescent R4 crop, which had a lower leaf layer in the canopy comprised of brown (dead or 363 364 low chlorophyll) leaves coupled with a mixed green/brown upper leaf canopy layer, exhibited relatively low PRI values (and high stress) in general, especially in the sunlit layer. The 365 increasing complexity of the foliage distribution at this highly variable stage presents a challenge 366 367 for simulations.

In our previous study, we showed that using sunlit and shaded leaves in the ACRM 368 scheme can improve both the correlation and RMSE with in situ PRI values (Cheng et al., 2010). 369 By considering the results from two field studies (2008, 2010), we can conclude that the most 370 significant benefit of adding shaded leaves as the lower canopy layer in ACRM was to improve 371 372 RMSE relative to field observations (see Fig.4 and Cheng et al., 2010). When ACRM was run in the one layer mode using only the optical properties of sunlit leaves, the simulated PRI 373 showed satisfactory correlations with in situ values but had a significant offset, indicating an 374 375 underestimate that could be incorrectly interpreted as a higher than actual physiological stress

376 response. This is an important issue since misinterpretation of PRI values will lead to significant errors in LUE and GPP estimates. This point was emphasized by calculating the difference 377 between *in situ* and simulated PRI values for various viewing geometry and dates (Fig. 5) where 378 simulations performed with only sunlit leaves obviously produced most of the underestimations 379 as negative values (Fig. 5a). Small θ_v (nadir at 0° and 30°) had larger discrepancies than larger 380 off-nadir views $(45^{\circ}, 60^{\circ})$, as compared with measurements. This is consistent with our previous 381 382 study (Cheng et al., 2010), due in part to less soil background contamination at oblique angles. 383 Among the three dates, discrepancies between field observations and "sunlit only" simulations were the smallest for the mature VT canopy (July 15, 2010), especially notable for $\theta_v = 60^\circ$ (X), 384 suggesting that at this oblique angle, sunlit leaves might dominate the field of view for a fully 385 leafed out, green and erectophile canopy. We also note that the benefit of adding a shaded lower 386 387 leaf layer for the VT canopy in the ACRM scheme, while advantageous, was less than on the other dates. This may be because the fully mature crop exhibited more sunlit leaves, greater 388 canopy closure, and/or a well-developed vertical LAI profile. The latter factor has been shown 389 390 to be temporally variant based on the growth stages of corn canopies (Ciganda et al., 2008).

391

4.3. Sunlit/Shaded Canopy Ratio

We tested various cases of variable sunlit/shaded canopy ratios, using our VT mature canopy dataset, for which the ACRM-simulated PRI values (for a given θ_v and ψ) were expected to increase when the sunlit/shaded ratio changed from 100/0 to 10/90. Those simulations (Fig. 6) duplicated those from the earlier study (Cheng et al. 2010): (1) the highest PRI values occurred at the coldspot ($\psi = 180^\circ$) and the lowest at the hotspot ($\psi = 0^\circ$); and (2) the PRI values were higher when θ_v increased from 30° to 60°. Therefore, in the ACRM simulations, changes to the sunlit/shaded ratio affected the canopy PRI responses expected, but not the sensitivity to viewinggeometry.

However, changes in the canopy structure could affect PRI values, causing 400 underestimation or overestimation of "true" PRI values. When simulations were done with sunlit 401 leaves only (100/0 in Fig. 7), most of the data points fell below the 1:1 line, underestimating, 402 "true" field values. After adding optical properties of shaded leaves in the simulation, even for 403 the 80/20 case, the data points moved closer to the 1:1 line and generated a ~50% improvement 404 in RMSE (Fig. 7). On the other hand, when even more shaded leaves than sunlit leaves were 405 406 included in the simulation (e.g., 40/60 and 20/80, Fig. 7), the simulated PRI moved up and over the 1:1 line, and generated higher RMSE due to overestimation. For this dataset, the simulated 407 60/40 sunlit/shaded canopy ratio appeared to have the best agreement with field measurements, 408 409 and indicates that this was the likely field condition at that growth stage in 2010. Since these simulations used a homogeneous, fixed LAI (at 2.48), the sunlit dominated groups (e.g., 100/0 410 and 80/20) might indicate relatively more open canopies, made possible by longer stems and/or 411 412 wider rows that put space between the leaves (since the number of leaves per plant is fixed). Likewise, the extreme case for a mostly shaded 20/80 canopy has a more compact, closely 413 414 spaced leaf arrangement along a short stem, and/or a closed canopy in narrower rows.

Results summarized in Tab. 2 also confirm the importance of adding shaded leaves into the simulation scheme, since even when using a 80/20 ratio as the input, significant improvement in RMSE (~30% to 50%) can be achieved. For the mature VT canopy, even though the 60/40 ratio appeared to be optimal by generating the best agreement of the sunlit/shaded ratio to *in situ* measurements (r = 0.87; RMSE = 0.0045), the performance using 50/50 was still quite close (r = 0.87; RMSE = 0.0048). Furthermore, after finding that the 50/50 sunlit/shaded ratio was not the optimal value to generate the best simulations for the mature VT canopy, we investigated the issue for the other two dates in 2010 and one V10 dataset acquired on August 1, 2008 (Tab. 2). For all three of these other datasets, the 50/50 sunlit/shaded ratio did appear to be optimal for simulating canopy PRI, based on better statistical performances (higher correlation coefficients and lower RMSEs). Therefore, while additional canopy structure information might improve PRI values interpretation and simulation in a cornfield using ACRM, the 50/50 sunlit/shaded ratio will generate more than satisfactory results for most of the cases.

These results indicate that the sunlit/shaded ratio, a structure-based parameter, may 428 429 change within a growing season. Therefore, this sensitivity analysis highlights the importance of canopy structure in simulating and understanding PRI. The implication is that since different 430 vegetation types have different canopy structures (e.g., forests vs. crops vs. shrubs), our on-going 431 432 and future research will apply this modeling scheme to different vegetation functional types. More importantly, most approaches have assumed that the sunlit upper canopy is the major 433 contributor and regulator of GPP/NPP, and that either the shaded component can be largely 434 435 ignored or the whole system is assumed to operate in one mode (e.g., sunlit) for total canopy foliage amount defined by LAI. Our results suggest that only taking sunlit leaves into account 436 would lead to underestimation of canopy PRI values, implying greater than actual stress levels 437 and leading to underestimates of LUE and GPP. Previous studies have also recognized the 438 importance of separating sunlit and shaded leaves for modeling photosynthetic activities from 439 leaf to canopy level (Chen et al., 1999; De Pury and Farquhar, 1997; Wang and Leuning, 1998) 440 mostly due to the nonlinear response of leaf carbon assimilation to light intensity. Adding 441 spectral information about the shaded canopy foliage is critical for improving our understanding 442 443 about canopy physiological processes, and our ability to simulate PRI and related parameters.

Improvement in understanding PRI information will potentially reduce uncertainties in LUE
estimates using remote sensing observations and advance carbon uptake monitoring capabilities.

447 **5. SUMMARY**

In this study, we examined the capability of coupling *in situ* leaf optical properties and 448 ACRM to simulate canopy level PRI at various growth stages of a corn crop. ACRM-simulated 449 450 canopy PRI values were closer to field measurements when both sunlit and shaded leaves were utilized in the scheme. The performance of the model was greatly improved when the crop was 451 dominated by green foliage during the vegetative and mature reproductive stages. The least 452 453 satisfactory results were found when the corn crop reached the senescent stage. The significance 454 of taking both sunlit and shaded leaf segments into account for canopy PRI studies was presented. We further examined how variable sunlit/shaded canopy ratios affected the modeled 455 results. Simulated canopy PRI values increased as the contribution from the shaded fraction 456 457 increased (i.e., the sunlit/shaded ratio decreased). The analysis suggested that canopy structure information might be needed to improve simulations or to interpret PRI. These findings also 458 459 imply that canopy PRI investigations and simulations should be investigated for more plant functional types. 460

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679 FIGURE CAPTIONS

Figure 1. PRI values derived from *in situ* leaf reflectance for the sunlit (\Box) and shaded (\blacksquare)

leaves used as input data in the simulations and daily average (\blacksquare) on three field days in 2010.

682 Values are shown as mean \pm SE.

Figure 2. In situ canopy PRI values from field measurements are shown for nadir ($\theta_v=0^\circ$; $\psi=0^\circ$)

and for three additional view zenith angles ($\theta_v = 30^\circ, 45^\circ, 60^\circ$) which were coupled with eight

relative azimuth angles ($\psi=0^{\circ}$ to 315° with 45° increment) on July 1st (\blacksquare), July 15th (\blacksquare), and

686 August 9th (\Box) in 2010. Values are shown as mean \pm SE. The mature canopy was clearly

687 differentiated from early and late canopies, with higher PRI values at any θ_v . Early and late

growth stages were similar at $\theta_v = 30^\circ$ and 45° , but were differentiated at $\theta_v = 60^\circ$. These results

689 were used as validation data for simulations.

Figure 3. Comparisons and regressions between simulated and *in situ* PRI values on three days

during the 2010 growing season: (a)(b) July 1; (c)(d) July 15; and (e)(f) August 9. Simulations

692 were performed using either sunlit leaves only (\diamondsuit) or both sunlit and shaded leaves (\blacktriangle). Values

are shown as mean \pm SE. in (a)(c)(e). In general, simulations agreed with field observations

694 when both sunlit and shaded foliage were included.

Figure 4. Summary chart of statistics representing all data collected on the three 2010 field

dates, for simulations using either sunlit canopy only or both sunlit and shaded canopy sectors:

- 697 (a) correlation coefficient (r) and (b) root mean square error (RMSE) relating *in situ* and
- 698 simulated PRI values.

Figure 5. Differences between values for *in situ* versus simulated PRI plotted against viewing geometry (θ_v and ψ) for the three growth stages in 2010. The black dashed line indicates zero

difference between *in situ* and simulated values. Discrepancies indicate the error incurred insimulations.

Figure 6. Changes in PRI values when 2-layer simulations were performed with various 703 704 sunlit/shaded canopy ratios, where the upper layer is sunlit and the lower layer is shaded. Simulations were done using parameters from the mature and green VT canopy, LAI = 2.48 on 705 July 15, 2010. Six sunlit/shaded ratio cases were investigated, as shown in the label, represented 706 707 by increasingly darker grey tone as more shaded leaves are included. The nadir case is included in the top panel. 708 709 Figure 7. Correlations between *in situ* PRI measurements and PRI values simulated using 710 various sunlit/shaded canopy ratios, for the mature VT canopy on July 15, 2010. LAI = 2.48. Figure 8. Statistics for the correlation coefficient (r) and root mean square error (RMSE), 711 712 relating *in situ* PRI measurements and simulated PRI values across various sunlit/shaded canopy ratios. Based on the VT canopy (July 15, 2010; LAI = 2.48). 713 **Figure 9.** Parameters of the regression line (y=ax+b) relating *in situ* and simulated PRI values, 714 715 using various sunlit/shaded canopy ratios. Parameter "a" is the slope while "b" is the offset of the regression line. Based on the VT canopy (July 15, 2010; LAI = 2.48). 716 717

718 **TABLE CAPTIONS**

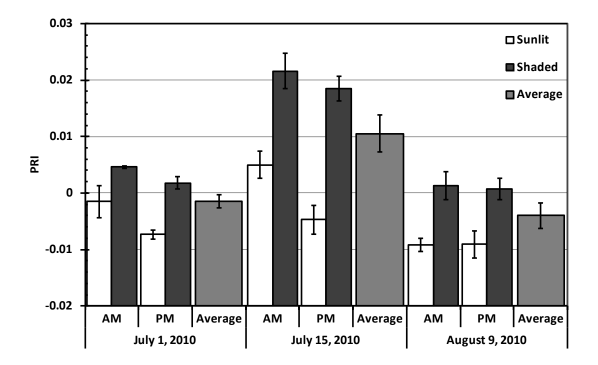
Table 1. Value or range of parameters used as input to ACRM in this study.

Table 2. Correlation coefficients (r) and root mean square errors (RMSE) relating *in situ* PRI

721 measurements and ACRM-simulated PRI values using various sunlit/shaded canopy ratios are

presented for three additional days: an early 2008 growth stage and two 2010 growth stages.

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724 725 **Figure 1.** PRI values derived from *in situ* leaf reflectance for the sunlit (\Box) and shaded (\blacksquare)

leaves used as input data in the simulations and daily average (■) on three field days in 2010. 726

727 Values are shown as mean \pm SE.

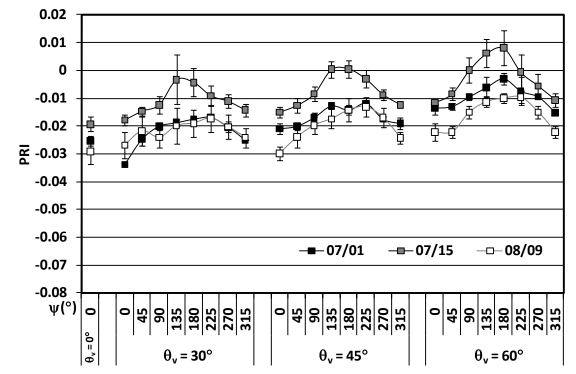


Figure 2. In situ canopy PRI values from field measurements are shown for nadir ($\theta_v=0^\circ$; $\psi=0^\circ$) and for three additional view zenith angles ($\theta_v=30^\circ,45^\circ,60^\circ$) which were coupled with eight relative azimuth angles ($\psi=0^\circ$ to 315° with 45° increment) on July 1st (\blacksquare), July 15th (\blacksquare), and August 9th (\Box) in 2010. Values are shown as mean ± SE. The mature canopy was clearly differentiated from early and late canopies, with higher PRI values at any θ_v . Early and late growth stages were similar at $\theta_v = 30^\circ$ and 45°, but were differentiated at $\theta_v = 60^\circ$. These results were used as validation data for simulations.

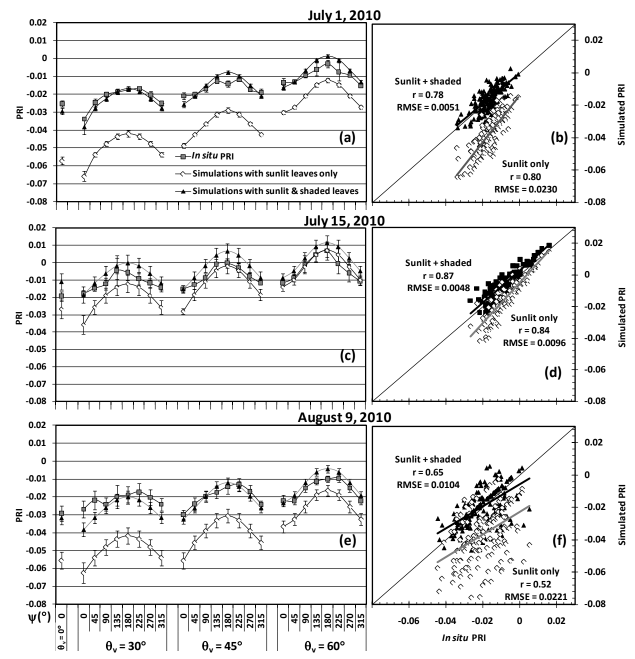
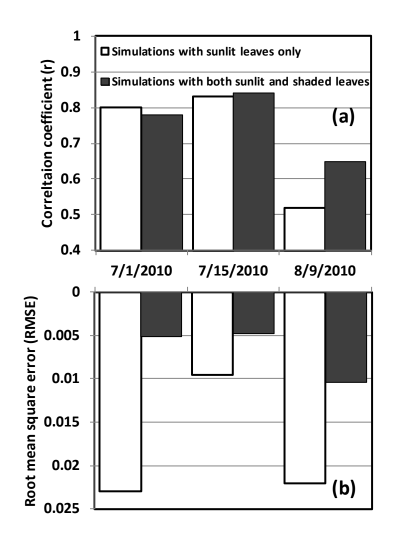
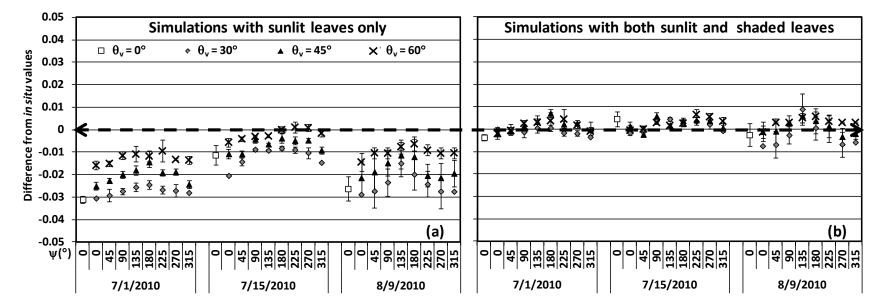




Figure 3. Comparisons and regressions between simulated and *in situ* PRI values on three days during the 2010 growing season: (a)(b) July 1; (c)(d) July 15; and (e)(f) August 9. Simulations were performed using either sunlit leaves only (\diamondsuit) or both sunlit and shaded leaves (\blacktriangle). Values are shown as mean \pm SE. in (a)(c)(e). In general, simulations agreed with field observations when both sunlit and shaded foliage were included.



- **Figure 4.** Summary chart of statistics representing all data collected on the three 2010 field
- 746 dates, for simulations using either sunlit canopy only or both sunlit and shaded canopy sectors:
- (a) correlation coefficient (r) and (b) root mean square error (RMSE) relating *in situ* and
- simulated PRI values.



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Figure 5. Differences between values for *in situ* versus simulated PRI plotted against viewing geometry (θ_v and ψ) for the three

growth stages in 2010. The black dashed line indicates zero difference between *in situ* and simulated values. Discrepancies indicate
 the error incurred in simulations.

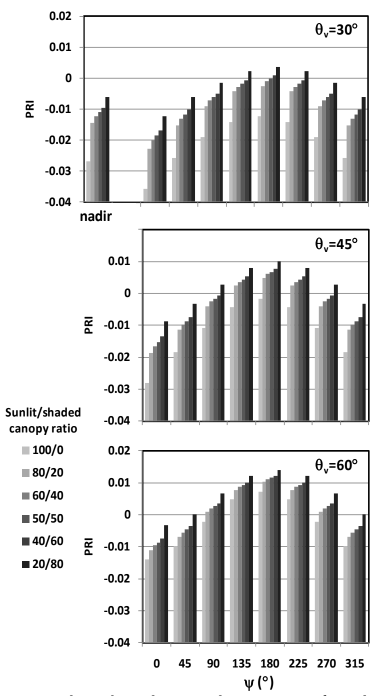


Figure 6. Changes in PRI values when 2-layer simulations were performed with various sunlit/shaded canopy ratios, where the upper layer is sunlit and the lower layer is shaded.

- 57 Simulations were done using parameters from the mature and green VT canopy, LAI = 2.48 on
- July 15, 2010. Six sunlit/shaded ratio cases were investigated, as shown in the label, represented
- by increasingly darker grey tone as more shaded leaves are included. The nadir case is included
- in the top panel.

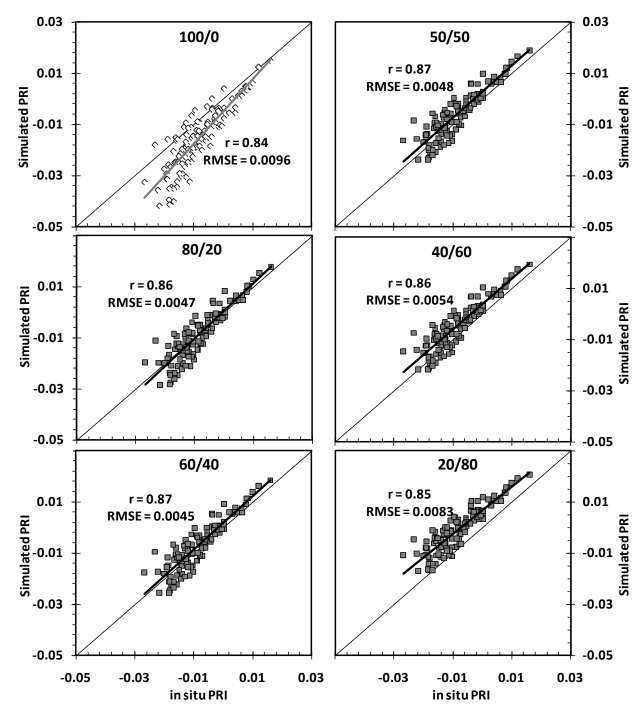




Figure 7. Correlations between in situ PRI measurements and PRI values simulated using 763 various sunlit/shaded canopy ratios, for the mature VT canopy on July 15, 2010. LAI = 2.48.

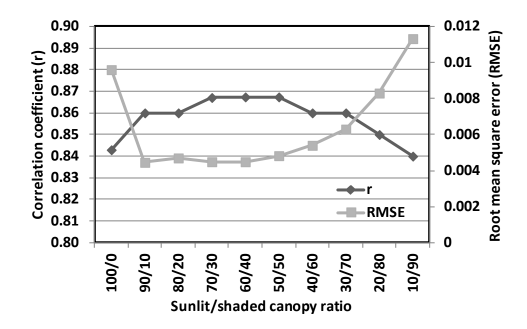
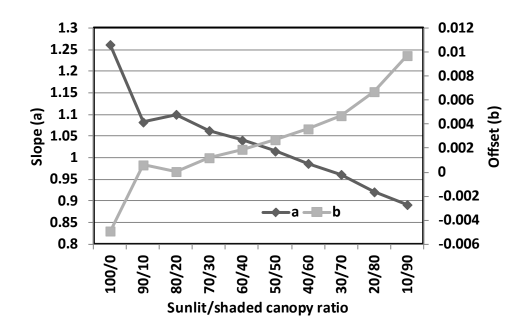


Figure 8. Statistics for the correlation coefficient (r) and root mean square error (RMSE),

relating *in situ* PRI measurements and simulated PRI values across various sunlit/shaded canopy ratios. Based on the VT canopy (July 15, 2010; LAI = 2.48).

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Figure 9. Parameters of the regression line (y=ax+b) relating *in situ* and simulated PRI values,

various sunlit/shaded canopy ratios. Parameter "a" is the slope while "b" is the offset of the

regression line. Based on the VT canopy (July 15, 2010; LAI = 2.48).

Date	July 1, 2010	July 15, 2010	August 9, 2010		
LAI	1.92	2.48	1.81		
Solar zenith angle (θ_s)	16.6° to 42.8°	18.1° to 45.3°	24.1° to 51.2°		
View zenith angle (θ_v)	0°, 30°, 45°, 60°				
Relative azimuth angle (ψ)	0° to 315° at 45° increments				
Relative leaf size	0.15				
Markov parameter	1.0				
Leaf angle distribution parameter	$\epsilon=0;\theta_m=0$				

Table 1. Value or range of parameters used as input to ACRM in this study.

Table 2. Correlation coefficients (r) and root mean square errors (RMSE) relating *in situ* PRI

measurements and ACRM-simulated PRI values using various sunlit/shaded canopy ratios are
 presented for three additional days: an early 2008 growth stage and two 2010 growth stages.

		2	2	0		0	<u> </u>
		100/0	80/20	60/40	50/50	40/60	20/80
August 1, 2008	r	0.71	0.80	0.85	0.86	0.84	0.82
August 1, 2008	RMSE	0.019	0.009	0.006	0.004	0.006	0.007
July 1, 2010	r	0.80	0.77	0.78	0.78	0.75	0.68
July 1, 2010	RMSE	0.023	0.010	0.007	0.005	0.006	0.007
August 0 2010	r	0.52	0.57	0.62	0.65	0.65	0.64
August 9,2010	RMSE	0.022	0.014	0.010	0.010	0.011	0.012

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