1	Modeling photosynthesis of discontinuous plant canopies by linking
2	Geometric Optical Radiative Transfer model with biochemical processes
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Abstract

19 Modeling vegetation photosynthesis is essential for understanding carbon exchanges between terrestrial 20 ecosystems and the atmosphere. The radiative transfer process within plant canopies is one of the key 21 drivers that regulate canopy photosynthesis. Most vegetation cover consists of discrete plant crowns, of 22 which the physical observation departs from the underlying assumption of a homogenous and uniform medium in classic radiative transfer theory. Here we advance the Geometric Optical Radiative Transfer 23 24 (GORT) model to simulate photosynthesis activities for discontinuous plant canopies. We separate radiation 25 absorption into two components that are absorbed by sunlit and shaded leaves, and derive analytical solutions by integrating over the canopy layer. To model leaf-level and canopy-level photosynthesis, leaf 26 27 light absorption is then linked to the biochemical process of gas diffusion through leaf stomata. The canopy gap probability derived from GORT differs from classic radiative transfer theory, especially when the leaf 28 area index is high, due to leaf clumping effects. Tree characteristics such as tree density, crown shape, and 29 30 canopy length affect leaf clumping and regulate radiation interception. Modeled gross primary production 31 (GPP) for two deciduous forest stands could explain more than 80% of the variance of flux tower measurements at both near hourly and daily time scales. We demonstrate that ambient CO₂ concentrations 32 33 influence daytime vegetation photosynthesis, which needs to be considered in biogeochemical models. The 34 proposed model is complementary to classic radiative transfer theory and shows promise in modeling the 35 radiative transfer process and photosynthetic activities over discontinuous forest canopies.

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37 Key words: gross primary production; flux tower; carbon cycle; radiative transfer; carbon assimilation

38 1. Introduction

Terrestrial plants assimilate atmospheric carbon dioxide through photosynthesis (Keenan et al., 2013; 39 Myneni et al., 1997). The climate system, in turn, affects vegetation development and photosynthetic 40 activities (Broich et al., 2014; Xia et al., 2014; Yi et al., 2010). Photosynthesis, accompanied by exchanges 41 42 of heat, water vapor, and trace gases within the planetary boundary layer, modifies microclimates and local 43 environments and determines ecosystem functions and services (Peng et al., 2014; Xu et al., 2013). The 44 complex biosphere/atmosphere feedbacks are dynamic and interactive (Bonan, 2008; Heimann and 45 Reichstein, 2008), such that robust numerical models that simulate vegetation photosynthesis are required in terrestrial ecosystem models to understand the global carbon cycle (Cramer et al., 2001; Kucharik et al., 46 47 2006).

48

Vegetation photosynthesis activity is regulated by environmental factors, and the light environment within 49 50 plant canopies is one of the key drivers (Law et al., 2002; Pearcy and Sims, 1994). Biophysical models such 51 as Production Efficiency Models assume linear relationships between absorbed photosynthetically active radiation (APAR) and vegetation primary production (Field et al., 1995; Monteith, 1977; Potter et al., 1993; 52 53 Prince and Goward, 1995; Running et al., 2000). Because vegetation photosynthesis harvests solar radiation 54 by green chlorophyll, recent studies have attempted to quantify the fractions of APAR that are absorbed by 55 green chlorophyll (Zhang et al., 2014; Zhang et al., 2005). Physiologically, plants assimilate carbon dioxide 56 via the biochemical diffusion processes through stomata, numerous small pores on the leaf surfaces (Collatz 57 et al., 1991; Farguhar and Sharkey, 1982). Stomata can open and close in response to microenvironments, 58 thereby regulating plant carbon uptake (Bonan, 2002). Field physiological studies have accumulated 59 detailed information on the behavior of stomata under certain environmental conditions (Schulze et al., 1994), in which sunlight irradiance plays a vital role (Ball et al., 1987). In this domain, linking the physical 60 61 process of radiative transfer within plant canopies with the biochemical process of gas diffusion through leaf stomata is essential for accurate representation of vegetation photosynthesis. 62

64 Radiative transfer within a plant canopy is determined by many factors such as the partition of incoming 65 solar radiation, solar illumination geometry, terrain slope and aspects, canopy structure, leaf angle 66 distribution, and leaf and substrate spectral properties (Baldocchi et al., 1985; Fan et al., 2014; Schaaf et al., 67 1994). Classic radiative transfer theory assumes that plant leaves are randomly distributed in three-68 dimensional space within a homogeneous canopy layer (Goudriaan, 1977; Myneni et al., 1990). The canopy 69 radiative transfer process can be simply characterized by leaf area index (LAI) and leaf angle distribution 70 (LAD). Three-dimensional, multi-layer, and two-leaf radiative transfer models have been developed to 71 simulate leaf absorption of solar irradiance and canopy photosynthesis (Myneni, 1991; Pury and Farguhar, 72 1997; Ryu et al., 2011; Sellers, 1985). Although classic radiative transfer theory holds well for dense 73 vegetation canopies, most vegetation canopies, especially arboreal canopies, consist of discrete crowns in 74 reality (Yuan et al., 2013). Leaves are clumped within individual crowns, such that more sunlight penetrates 75 to understory layers and the ground surfaces (He et al., 2012; Ni-Meister et al., 2010). Tree crowns also cast 76 shadows on one another and on the background, resulting in self-shadowing effects as described by the 77 geometric-optical theory (Li and Strahler, 1992). Given natural differences in the radiative transfer process 78 between homogenous and discontinuous plant canopies, it is important to understand and account for the 79 influence of crown shape and tree structure on canopy radiation absorption and vegetation photosynthesis.

80

To address the radiative transfer process in discontinuous canopies, the Geometric-Optical Radiative-Transfer (GORT) model conceptually combines geometric optical principles for canopy structure and radiative transfer theory for volumetric scattering within canopy crowns (Li et al., 1995). The geometric optical method is used to characterize the process by which sunlight passes directly to the ground surface without reaching any canopy crowns. The radiative transfer principle is applied to model the probability of light penetration as it travels through crowns in the canopy. GORT has been used to model the physical aspects of discontinuous plant canopies such as gap fraction, radiation transmission, and bi-directional

reflectance (Ni et al., 1999; Ni et al., 1997; Xin et al., 2012), and has been validated under a variety of 88 89 environmental conditions (Liu et al., 2008). Recent efforts have been made to develop and evaluate a 90 simplified GORT model for the use in coupled global dynamic terrestrial ecosystem models (Ni-Meister et al., 2010; Yang et al., 2010). Despite these successful applications, the current version of the GORT model 91 92 does not have analytical solutions for radiation absorption by sunlit and shaded leaves, though previous 93 studies have tried to solve the process of multiple scattering between canopy and background in an iterative 94 manner (Song et al., 2009). However, sunlit and shaded leaves must be treated separately in photosynthesis 95 modeling because flux densities of photosynthetically active radiation (PAR) incident on leaf surfaces are 96 different (He et al., 2013). It is also necessary to integrate vertically over the canopy to derive mean PAR 97 absorbed by sunlit and shaded leaves because of the non-linear light attenuation within the canopy and the 98 non-linear dependence of leaf stomatal conductance on light absorption (Campbell and Norman, 1998).

99

The objectives of this study are to 1) advance the GORT model by providing analytical solutions to the radiation absorption of sunlit and shaded leaves and 2) link the radiative transfer process to biochemical processes to simulate leaf and canopy photosynthesis. We first describe the principles of our model and then perform model validation with eddy covariance data from two flux towers situated in the New England region of the United States.

105

106 2. Theoretical Basis

107 **2.1 Brief description of canopy gap probability modeled using GORT**

Gap probability, the probability of photons reaching a given canopy depth without being intercepted by canopy elements, is key to characterizing the radiation distribution within plant canopies. A detailed description for modeling the gap probability with GORT is described in previous studies (Li et al., 1995; Ni et al., 1999), and we summarize it briefly here because the concept of gap probability is necessary for understanding our subsequent work.







117 For homogeneous canopies, Beer's law describes the gap probability of sunlight penetration. For 118 discontinuous plant canopies, leaves are clumped within individual canopy crowns, forming an uneven 119 distribution of gap probabilities for beam radiation. GORT models tree crowns as a collection of ellipsoids 120 (Figure 1), of which the centers are randomly distributed between the upper and lower boundaries of the 121 canopy layer $(h_1 \text{ and } h_2)$. Each ellipsoid, or each canopy crown, is characterized by one-half of the vertical 122 crown length (b) and a horizontal crown radius (R). The total gap probability is modeled separately as the proportion of sunlight passing through the canopy layer without reaching any crown (hereafter referred to as 123 124 between-crown gaps) and the proportion of sunlight passing through crowns without being intercepted by 125 canopy leaves (hereafter referred to as within-crown gaps), such that:

 $P_{\text{gap}}(h,\theta_i) = P_{\text{gap}}(n=0|h,\theta_i) + P_{\text{gap}}(n>0|h,\theta_i)$ (1)

126 where $P_{gap}(h, \theta_i)$ is the gap probability for beam radiation at height h given an illumination zenith angle θ_i ,

127 $P_{gap}(n = 0 | h, \theta_i)$ is the between-crown gap, and $P_{gap}(n > 0 | h, \theta_i)$ is the within-crown gap.

128

129 The between-crown gap is modeled based on Boolean theory as an exponential function of crown numbers130 within a geometric volume that contains no crown centers:

$$P_{\text{gap}}(n=0|h,\theta_i) = e^{-\lambda_v V_{\Gamma}}$$
(2)

131 where λ_v is the tree density, and V_{Γ} is the beam projected cylinder volume with a radius *R* starting from the 132 canopy top and extending to height *h*.

133

Assuming that leaves are randomly distributed within each individual crown, the within-crown gap is modeled based on Beer's law as light penetration along the traveling path length, such that:

$$P_{\text{gap}}(n > 0 | h, \theta_i) = \int_0^\infty P(s | h, \theta_i) e^{-\tau(\theta_i)s} ds$$
(3)
136 where $\tau(\theta_i, \alpha) = k_b(\theta_i, \alpha) \cdot FAVD$, $FAVD$ is the foliage area volume density within a single crown, and
137 $k_b(\theta_i, \alpha)$ is the extinction coefficient for beam radiation given a specific solar illumination angle θ_i and leaf
138 distribution angle α . For a spherical leaf angle distribution, $k_b = \frac{0.5}{\cos(\theta_i)}$. $P(s | h, \theta_i)$ is the probability
139 distribution function associated with within-crown path length s .

140

141 The probability distribution of within-crown paths length can be solved in a convolutional manner:

$$P(s|h,\theta_i) = \int_h^{h_2} \sum_{n=1}^{n=\infty} P(s|n,z,h,\theta_i) P(n|z,h,\theta_i) dz$$
(4)

142 where $P(s|n, z, h, \theta_i)$ is the probability distribution of within-crown path length given that a solar ray enters 143 the crown at height *h* and angle θ_i , and $P(n|z, h, \theta_i)$ is the probability distribution of the numbers of crowns 144 intercepted by the solar ray incident at angle θ_i , entering crowns at height *z*, and then traveling to height *h*.

Diffuse radiation (i.e., the hemispherically isotropic radiation) can be treated as beam radiation from all directions in the upper hemisphere. The "openness" of discontinuous plant canopies to diffuse radiation on a horizontal plane is defined as:

$$K_{\text{open}}(h) = K_{\text{open}}(n = 0|h) + K_{\text{open}}(n > 0|h)$$

$$K_{\text{open}}(n = 0|h) = \frac{1}{\pi} \int_{0}^{2\pi} \int_{0}^{\frac{\pi}{2}} P_{\text{gap}}(n = 0|h, \theta_{i}) \sin(\theta_{i}) \cos(\theta_{i}) d\theta_{i} d\phi$$

$$= 2 \int_{0}^{\frac{\pi}{2}} P_{\text{gap}}(n = 0|h, \theta_{i}) \sin(\theta_{i}) \cos(\theta_{i}) d\theta_{i}$$

$$K_{\text{open}}(n > 0|h) = \frac{1}{\pi} \int_{0}^{2\pi} \int_{0}^{\frac{\pi}{2}} P_{\text{gap}}(n > 0|h, \theta_{i}) \sin(\theta_{i}) \cos(\theta_{i}) d\theta_{i} d\phi$$

$$= 2 \int_{0}^{\frac{\pi}{2}} P_{\text{gap}}(n > 0|h, \theta_{i}) \sin(\theta_{i}) \cos(\theta_{i}) d\theta_{i}$$

$$(7)$$

149 where $K_{\text{open}}(n = 0|h)$ and $K_{\text{open}}(n > 0|h)$ are between-crown and within-crown openness factors, 150 respectively. θ_i is the solar illumination angle, and ϕ is the azimuth angle.

151

152 **2.2 Sunlit and shaded leaf area index**

The gap probability describes the probability of beam radiation being intercepted by plant leaves, and hence determines the proportion of leaf areas that are sunlit. For a very thin layer, the reduction of total gap probability is due to leaf interception, of which the process still follows Beer's law:

$$P_{\text{gap}}(h - \delta h, \theta_i) = \exp(-k_b \delta LAI(h)) P_{\text{gap}}(h, \theta_i)$$
(8)

156 where k_b is the canopy extinction coefficient for beam irradiance, $\delta LAI(h)$ is the leaf area index within a 157 thin layer δh at height h, and $P_{gap}(h, \theta_i)$ is the gap probability modeled using GORT.

$$\exp(-k_b \delta LAI(h)) = 1 - k_b \delta LAI(h)$$

$$P_{gap}(h - \delta h, \theta_i) = P_{gap}(h, \theta_i) - P'_{gap}(h, \theta_i) \delta h$$
(10)

160 where $P'_{gap}(h, \theta_i)$ is the first derivative of gap probability $P_{gap}(h, \theta_i)$ with respect to height *h*.

161

162 Combining Equations (8), (9), and (10), we obtain:

$$\frac{P_{\text{gap}}'(h,\theta_i)}{P_{\text{gap}}(h,\theta_i)}\delta h = k_b \delta LAI(h)$$
(11)

163

164 For diffuse radiation, it can be derived in a similar manner:

$$\frac{K'_{\text{open}}(h)}{K_{\text{open}}(h)}\delta h = k_d \delta LAI(h)$$
(12)

165 where k_d is the extinction coefficient for diffuse irradiance, and $K'_{open}(h)$ is the first derivative of the 166 openness factor $K_{open}(h)$ with respect to height *h*.

167

The sunlit LAI at height h is the product of the probability of beam sunlight penetration to height h and the probability of sunlight being intercepted by the thin layer and divided by the ratio of leaf area projected on a horizontal surface (Campbell and Norman, 1998), such that:

$$\delta LAI_{Sun}(h,\theta_i) = \frac{P_{gap}(h,\theta_i) \left[1 - \exp(-k_b \delta LAI(h))\right]}{k_b}$$
(13)

171 where $\delta LAI_{Sun}(h, \theta_i)$ is the sunlit leaf area index within a thin layer δh at height h.

172

173 Substituting Equations (9) and (11) into Equation (13), we obtain:

$$\delta LAI_{Sun}(\theta_i) = \frac{P'_{gap}(h,\theta_i)}{k_b} \delta h$$
(14)

175 Sunlit LAI for the entire canopy at zenith angle θ is then obtained by integrating from the canopy top to 176 canopy bottom, such that:

$$LAI_{Sun}(\theta_{i}) = \int_{z_{1}}^{z_{2}} \frac{P_{gap}'(h,\theta_{i})}{k_{b}} dh = \frac{1 - P_{gap}(h = z_{1} | \theta_{i})}{k_{b}}$$
(15)

177 where $P_{\text{gap}}(h = z_2 | \theta_i)$ and $P_{\text{gap}}(h = z_1 | \theta_i)$ are the gap probabilities at the canopy top z_2 and canopy 178 bottom z_1 , respectively, whereas the gap probability at the canopy top is 1.

179

180 It is worth noting that our calculation of sunlit leaf area for discontinuous canopies is analogous to that for 181 homogeneous canopies, which is given as:

$$LAI_{Sun}^{*}(\theta_{i}) = \int_{0}^{LAI} \exp(-k_{b} \cdot L) \, dL = \frac{1 - \exp(-k_{b} \cdot LAI)}{k_{b}} \tag{16}$$

182 where $LAI_{Sun}^{*}(\theta_{i})$ is the sunlit leaf area for homogeneous canopies.

183

184 The shaded LAI is simply the remainder of the canopy LAI:

$$LAI_{Shd} = LAI - LAI_{Sun}$$
(17)

185

186 2.3 Analytical solutions for the scattering parameters of discontinuous canopies

187 Canopy scattering parameters such as directional-hemispherical reflectance and hemispherical-188 hemispherical reflectance (or black-sky albedo and white-sky albedo, respectively) can be obtained by 189 resolving the radiative transfer process or can be approximated using simple analytical solutions. For semi-190 infinite horizontally homogeneous media, Hapke's solutions of the proportion of unintercepted direct beam 191 $(t_0(h, \theta_i))$, hemispherical-hemispherical reflectance (R_{df}^{∞}) , directional-hemispherical reflectance (R_{df}^{∞}) ,

192 hemispherical-hemispherical transmittance (T_{ff}^{∞}) , and directional-hemispherical transmittance (T_{df}^{∞}) are

193 given as (Hapke, 1981):

$t_0(h,\theta_i) = e^{-\frac{\tau(\theta_i)h}{\mu_i}}$	(18)
$R_{ff}^{\infty} = \frac{1-\gamma}{1+\gamma}$	(19)
$R_{df}^{\infty}(\theta_i) = \frac{1-\gamma}{1+2\mu_i\gamma}$	(20)
$T_{ff}^{\infty}(h) = e^{-2\gamma\tau h}$	(21)
$T_{df}^{\infty}(h,\theta_{i}) = \frac{\sigma}{2} \frac{1+2\mu_{i}}{1-(2\mu_{i}\gamma)^{2}} [T_{ff}^{\infty}(h) - t_{0}(h,\theta_{i})]$	(22)

194 where σ is the single scattering albedo, $\tau = k(\theta_i) \frac{L_e}{H}$ is the projected foliage area volume density for the 195 plant canopy, L_e is the effective leaf area index, *H* is the depth of the canopy, θ_i is the solar illumination 196 angle, $\mu_i = \cos(\theta_i)$ and $\gamma = \sqrt{1 - \sigma}$.

197

Starting with surface energy balances, Ni (1998) derived the scattering parameters for a horizontally
homogeneous canopy layer with finite thickness as:

$t_{ff}(h) = T_{ff}^{\infty}(h) \frac{1 - (R_{ff}^{\infty})^2}{1 - (T_{ff}^{\infty}(h)R_{ff}^{\infty})^2}$	(23)
$\rho_{ff}(h) = R_{ff}^{\infty}(h) \frac{1 - (T_{ff}^{\infty}(h))^2}{1 - (T_{ff}^{\infty}(h)R_{ff}^{\infty})^2}$	(24)
$t_{df}(h,\theta_i) = T_{df}^{\infty}(h,\theta_i) - \rho_{ff}(h)[t_0(h,\theta_i)R_{df}^{\infty}(\theta_i) + T_{df}^{\infty}(h,\theta_i)R_{ff}^{\infty}]$	(25)
$\rho_{df}(h,\theta_i) = R_{df}^{\infty}(h) - t_{ff}(h)[t_0(h,\theta_i)R_{df}^{\infty}(\theta_i) + T_{df}^{\infty}(h,\theta_i)R_{ff}^{\infty}]$	(26)

where $t_{ff}(h)$, $\rho_{ff}(h)$, $t_{df}(h, \theta_i)$, and $\rho_{df}(h, \theta_i)$ are hemispherical-hemispherical transmittance, hemispherical-hemispherical reflectance, directional-hemispherical transmittance, and directionalhemispherical reflectance, respectively.

203

The scattering parameters for a discontinuous canopy can then be approximated as combinations of a homogeneous vegetation layer and a non-vegetated layer:

where $t'_{ff}(h)$, $\rho'_{ff}(h)$, $t'_{df}(h,\theta_i)$, and $\rho'_{df}(h,\theta_i)$ are hemispherical-hemispherical transmittance, hemispherical-hemispherical reflectance, directional-hemispherical transmittance, and directionalhemispherical reflectance, respectively. Note that our equations here are slightly different from those used by Ni et al. (1999) because between-crown gaps, within which light attenuation obeys Beer's law, are considered in the homogeneous vegetation layer.

211

The analytical approximation of the canopy reflectance for beam and diffuse radiation is the sum of three factors in radiative transfer: the incoming irradiance scattered by the canopy elements, the first-order scattered radiation from soil background, and the irradiance scattered back and forth between the canopy layer and background surface (Ni et al., 1999). Taking beam radiation as an example and assuming that the background surface is Lambertian, the incoming irradiance scattered by the canopy elements is ρ'_{df} , the first-order scattered radiance from soil background is $t'_{df}\rho_s t'_{ff}$, and the multiple scattering between the 218 canopy elements and soil background is $t'_{df} (\rho_s \rho'_{ff} \rho_s + \rho_s (\rho'_{ff} \rho_s)^2 + \rho_s (\rho'_{ff} \rho_s)^3 + \cdots) t'_{ff}$. The canopy

219 reflectance for beam irradiance can then be written as:

$$\rho_{cb} = \rho'_{df} + t'_{df} \left(\rho_s + \rho_s \rho'_{ff} \rho_s + \rho_s (\rho'_{ff} \rho_s)^2 + \rho_s (\rho'_{ff} \rho_s)^3 + \cdots \right) t'_{ff}$$

$$= \rho'_{df} + t'_{df} \frac{\rho_s}{1 - \rho_s \rho'_{ff}} t'_{ff}$$
(31)

220

221 The canopy reflectance for diffuse irradiance can be obtained similarly as:

$$\rho_{cd} = \rho'_{ff} + t'_{ff} \frac{\rho_s}{1 - \rho_s \rho'_{ff}} t'_{ff}$$
(32)

222

223 **2.4** Mean photosynthetically active radiation absorbed by sunlit and shaded leaves

Let I_0 be the flux density of incoming solar radiation on a horizontal plane at the top of the canopy and f_b be the fraction of incident beam radiation, the unintercepted beam and diffuse fluxes are then:

$$I_{b}(h,\theta_{i}) = P_{\text{gap}}(h,\theta_{i})(1-\rho_{cb})f_{b}I_{0}k_{b}$$

$$I_{d}(h) = K_{\text{open}}(h)(1-\rho_{cd})(1-f_{b})I_{0}k_{d}$$
(33)
(34)

where ρ_{cb} and ρ_{cd} are canopy reflectance for beam and diffuse irradiance, respectively; I_b and I_d are the unintercepted beam and diffuse fluxes, respectively; and k_b and k_d are canopy extinction coefficients for beam and diffuse irradiance, respectively.

229

The downward beam flux I_b is derived based on the assumption of black leaves, meaning that leaves absorb incident irradiance completely and do not transmit radiation (Bonan, 2002). To account for the effects of leaf scattering, the total beam I_{bt} (i.e., unintercepted beam and down scattered beam) and total diffuse I_{dt} (i.e., unintercepted diffuse and down scattered diffuse) irradiance can be modeled by introducing a factor of $\sqrt{1-\sigma}$ to extinction coefficients similar to the two-stream radiative transfer model (Sellers, 1985). As single scattering albedo increases, the effective extinction coefficient becomes smaller and more sunlight is

allowed to transmit through the canopy. That is:

$$I_{bt}(h,\theta_{i}) = P_{gap}(h,\theta_{i})^{\sqrt{1-\sigma}} (1-\rho_{cb}) f_{b} I_{0} \sqrt{1-\sigma} k_{b}$$

$$I_{dt}(h) = K_{open}(h)^{\sqrt{1-\sigma}} (1-\rho_{cd}) (1-f_{b}) I_{0} \sqrt{1-\sigma} k_{d}$$
(35)
(35)
(35)

where σ is the single scattering albedo of leaves. $\sigma = \rho_l + t_l$, where ρ_l and t_l are leaf reflectance and transmittance, respectively.

239

The total irradiance absorbed by the entire canopy per unit ground area consists of leaf absorption for both
beam and diffuse irradiance:

$$I_{c} = I_{cb} + I_{cd} = \int_{0}^{LAI} I_{bt}(h,\theta_{i}) dL + \int_{0}^{LAI} I_{dt}(h,\theta_{i}) dL$$
(37)

242

243 Substituting Equations (11), (12), (35), and (36) into Equation (37), we have:

$$I_{cb} = \int_{z_{1}}^{z_{2}} P_{\text{gap}}(h,\theta_{i})^{\sqrt{1-\sigma}} (1-\rho_{cb}) f_{b} I_{0} \sqrt{1-\sigma} \frac{P_{\text{gap}}'(h,\theta_{i})}{P_{\text{gap}}(h,\theta_{i})} dh$$

$$= \left(1-P_{\text{gap}}(h=z_{1}|\theta_{i})^{\sqrt{1-\sigma}}\right) (1-\rho_{cb}) f_{b} I_{0}$$

$$I_{cd} = \int_{z_{1}}^{z_{2}} K_{\text{open}}(h)^{\sqrt{1-\sigma}} (1-\rho_{cd}) (1-f_{b}) I_{0} \sqrt{1-\sigma} \frac{K_{\text{open}}'(h)}{K_{\text{open}}(h)} dh$$

$$= \left(1-K_{\text{open}}(h=z_{1})^{\sqrt{1-\sigma}}\right) (1-\rho_{cd}) (1-f_{b}) I_{0}$$
(38)
$$(38)$$

$$(38)$$

$$(38)$$

$$(39)$$

244

Irradiance absorbed by sunlit leaves per unit ground area is obtained as the sum of direct beam, downward
 scattered beam, and diffuse components:

$$I_{Sun} = I_{Sunb} + I_{Sunbs} + I_{Sund}$$

$$\tag{40}$$

248 Combining Equations (33), (35), (36), and (40), we have:

$$I_{Sunb} = \int_{z_{1}}^{z_{2}} (1-\sigma) f_{b} I_{0} \cdot P_{gap}'(h,\theta_{i}) dh = (1-\sigma) \left(1-P_{gap}(h=z_{1}|\theta_{i})\right) f_{b} I_{0}$$
(41)

$$I_{Sunbs} = \int_{z_{1}}^{z_{2}} \left[P_{gap}(h,\theta_{i})^{\sqrt{1-\sigma}} (1-\rho_{cb})\sqrt{1-\sigma} - P_{gap}(h,\theta_{i})(1-\sigma)\right] f_{b} I_{0} \cdot P_{gap}'(h,\theta_{i}) dh$$
(42)

$$= \left[\frac{\sqrt{1-\sigma}}{1+\sqrt{1-\sigma}} \left(1-P_{gap}(h=z_{1}|\theta_{i})^{1+\sqrt{1-\sigma}}\right)(1-\rho_{cb}) - \frac{(1-\sigma)}{2} \left(1-P_{gap}(h=z_{1}|\theta_{i})^{2}\right)\right] f_{b} I_{0}$$
(42)

$$I_{Sund} = \int_{z_{1}}^{z_{2}} K_{open}(h)^{\sqrt{1-\sigma}} (1-\rho_{cd})(1-f_{b}) I_{0} \sqrt{1-\sigma} \cdot K_{open}'(h) dh$$
(43)

$$= \frac{\sqrt{1-\sigma}}{1+\sqrt{1-\sigma}} \left(1-K_{open}(h=z_{1})^{1+\sqrt{1-\sigma}}\right)(1-\rho_{cd})(1-f_{b}) I_{0}$$
(43)

Note that σ is used instead of ρ_{cd} for the beam irradiance of sunlit leaves because sunlit leaves scatter direct beam sunlight only once.

251

The irradiance absorbed by shaded leaves per unit ground area is simply the difference between the total irradiance absorbed by the canopy and the irradiance absorbed by sunlit leaves:

$$I_{Shd} = I_c - I_{Sun} \tag{44}$$

254

255 The mean absorbed irradiance for sunlit and shaded canopy per leaf hemi-surface area is then:

$$Q_{Sun} = \frac{I_{Sun}}{LAI_{Sun}}$$

$$Q_{Shd} = \frac{I_{Shd}}{LAI_{Shd}}$$
(45)
(46)

257 **2.5 Modeling leaf photosynthesis and scaling up to canopy photosynthesis**

The biochemical process of carbon dioxide assimilation by leaves can be considered as a gas diffusion process through stomata. According to Fick's law, the process is described as:

$$A = g_c \cdot (C_a - C_i) \tag{47}$$

260 where *A* is the CO_2 assimilation rate, g_c is the stomatal conductance, and C_a and C_i are ambient and 261 intercellular CO_2 concentrations, respectively.

262

Field studies have firmly established the relationship between leaf stomatal conductance and environmental conditions. Jarvis and McNaughton (1986) successfully synthesize the response functions in a multipleconstraint model:

$$g_c = g_{cmax} \prod f(x_i) \tag{48}$$

266 where g_{cmax} is the maximum leaf stomatal conductance when environmental factors do not limit carbon 267 uptake and $f(x_i)$ are scalars that account for the influences of various environmental stresses on leaf 268 stomatal conductance.

269

Different formulas have been developed to describe the response functions of photosynthesis to environmental factors. Here, we consider three main limiting factors imposed by radiation, temperature, and water on vegetation photosynthesis. The equations developed for the dual-source dual-leaf (DSDL) model (Ding et al., 2014), Terrestrial Ecosystem Model (Raich et al., 1991), and Biome-BGC models (Running et al., 2004) are used to account for the influences of radiation, temperature, and vapor pressure deficit (VPD), respectively:

$$\int f(x_i) = f(Q) \cdot f(T) \cdot f(VPD)$$

(49)

$$f(Q) = \frac{k_{c} + k_{Q}}{k_{Q}} \cdot \frac{Q}{k_{Q} + Q}$$
(50)
$$f(T) = \frac{(T - T_{min})(T - T_{max})}{(T - T_{min})(T - T_{max}) - (T - T_{opt})^{2}}$$
(51)
$$f(VPD) = \frac{VPD_{max} - VPD}{VPD_{max} - VPD_{min}}$$
(52)

where k_c and k_q are the stress coefficients of PAR absorbed by plant leaves; Q is the mean APAR for sunlit 276 or shaded leaves per leaf hemi-surface area; T_{min} , T_{opt} , and T_{max} are the minimum, optimum, and 277 maximum temperature for photosynthetic activities, respectively; and VPD_{min} and VPD_{max} are the 278 minimum and maximum vapor pressure deficit, respectively. In the DSDL model, k_c and k_q are 500 W/m² 279 280 and 150 W/m², respectively. T_{min}, T_{opt}, and T_{max} are determined as 10 °C, 28 °C and 48 °C for C4 crops (Kalfas et al., 2011), and here we slightly lower their values to 0 °C, 25 °C, and 45 °C, respectively, for C3 281 plants. VPDmin and VPDmax are 0.65 kPa and 4.6 kPa for deciduous forests, respectively, in the Biome-282 283 BGC model (Heinsch et al., 2003).

284

Due to different PAR absorption by sunlit and shaded leaves, the stomatal conductance for sunlit and shaded
 leaves need to be calculated separately as:

$$g_{cSun} = g_{cmax} \cdot f(Q_{Sun}) \cdot f(T) \cdot f(VPD)$$

$$g_{cShd} = g_{cmax} \cdot f(Q_{Shd}) \cdot f(T) \cdot f(VPD)$$
(53)
(54)

where g_{cSun} and g_{cShd} are the stomatal conductance for sunlit and shaded leaves, respectively, and Q_{Sun} and Q_{Shd} are the mean PAR absorbed by sunlit and shaded leaves, respectively.

289

Given measured ambient CO_2 concentrations, the closure of the formulation (47) now requires the quantity of intercellular CO_2 concentrations. Katul et al. (2000) compared eight models and concluded that all reproduced the measured carbon assimilation rates well. Here, we employ Leuning's method (Leuning, 1995) to estimate the ratio of intercellular to ambient CO_2 concentrations as:

$$\frac{C_i}{C_a} = 1 - \frac{1 - \frac{\Gamma}{C_a}}{m_L} (1 + \frac{\text{VPD}}{\text{VPD}_0})$$
(55)

where VPD is the ambient vapor pressure deficit; VPD_0 is an empirical constant describing the species sensitivity to ambient vapor pressure deficit; Γ is the leaf CO_2 compensation point; C_a and C_i are ambient and intercellular CO2 concentrations, respectively; and m_L represents linear regression coefficients related to tree species. Calibrated values for model parameters are $m_L = 4.0$, $\Gamma = 40 \,\mu\text{mol/mol}$, and $VPD_0 =$ 30 kPa, respectively (Katul et al., 2000).

299

300 Given modeled carbon assimilation rates at the leaf level, the total rate of carbon assimilation at the canopy 301 level can be scaled up as:

$$GPP = A_{Sun} \cdot LAI_{Sun} + A_{Shd} \cdot LAI_{Shd}$$

where *GPP* is canopy gross primary production, A_{Sun} and A_{Shd} are leaf-level carbon assimilation rates for sunlit and shaded leaves, respectively, and LAI_{Sun} and LAI_{Shd} are the sunlit and shaded leaf area index.

305 3. Study materials and model parameterization

We studied two deciduous forest sites: Harvard Forest (US-Ha1) in Massachusetts and Bartlett Experimental Forest (US-Bar) in New Hampshire (Richardson et al., 2012). Basic information is briefly summarized in Table 1 for each site. Although plot layouts set up for the fieldwork did not match the exact footprints of flux towers (Yang et al., 2013), the measured tree structural attributes, such as tree density, are assumed to be representative of the two study sites.

311

312

(56)

Table 1. Site information as obtained from the AmeriFlux website unless notified.

	Site	Site name	Lat	Lon	Elevation	Canopy	Tree density	Dominant species
	code		(N)	(W)	(m)	height (m)	(trees/ha) ^a	
	US-	Harvard Forest	42.5378	72.1715	340	23.0	1020±72	red oak, red maple
	Ha1							
	US-	Bartlett Experimental	44.0646	71.2881	272	19.0	1432±67	American beech, red
	Bar	Forest						maple
314	^a data f	from Yao et al. (2011)					

316 Flux towers measure energy and material fluxes between ecosystem and the atmosphere continuously 317 (Baldocchi et al., 2001). Measured data are provided as standard Level 2 products in the AmeriFlux 318 database (http://ameriflux.ornl.gov/). The time steps of available data are half-hourly for US-Bar and hourly 319 for US-Ha1. The measurements we used include estimates of gross primary production (GPP) derived with 320 the eddy covariance technique (Baldocchi, 2003), and meteorological variables such as shortwave solar radiation, temperature, vapor pressure deficit, and canopy-scale CO₂ concentration. Raw measurements of 321 322 meteorological variables were used for analysis and missing values due to instrument malfunction or 323 unsuitable micrometeorological conditions were screened. However, we obtained GPP estimates from AmeriFlux Level 4 products if they were not delivered in Level 2 products. Extraterrestrial solar radiation 324 325 and solar zenith angle (i.e., the angle that the sun away from directly overhead) are calculated as a function 326 of geolocation (i.e., latitude and longitude), the day of year (DOY), and solar time of the day (Allen et al., 327 1998). If diffuse radiation is missing from the measurements, we implement Muneer's method to partition 328 global solar radiation into beam and diffuse components (Muneer, 2007):

$(1 - f_b) = 1.006 - 0.317K_t + 3.1241K_t^2$	$2.7616K_t^3 + 9.7166K_t^4 \tag{5}$	57)	
--	-------------------------------------	-----	--

329 where f_b is the proportion of beam radiation in global incoming radiation, and K_t is the hourly clearness 330 index. $K_t = I_0/I_e$, where I_0 is global solar radiation on the canopy top and I_e is the extraterrestrial solar 331 radiation.

332

333 We use typical parameter values from the literature for model parameterization. Because the spectral 334 signatures of vegetation leaves and soil background differ in the spectral bands of PAR and near infrared

335	(Table 2), we perform model simulations for these two discrete bands separately. Incident PAR is estimated
336	to account for 47.5% of incoming shortwave solar radiation, and the rest is attributed to the near infrared
337	band (Zhao et al., 2005). Maximum leaf stomatal conductance to H ₂ O is estimated as 5.5 mm/s for US-Bar
338	and 7.2 mm/s for US-Ha1 (Bonan, 2002; Ding et al., 2014), and they are translated to maximum leaf
339	stomatal conductance to CO ₂ assuming that the temperature is 20°C and the atmospheric pressure is 101.32
340	kPa (Pearcy et al., 1989). Heights for canopy top (z_2) were measured to be 23.0 m for US-Ha1 and 19.0 m
341	for US-Bar (Table 1), and heights for canopy bottom (z_1) were estimated as $z_1 = 0.15 z_z$. Canopy structure
342	in GORT is modeled with the ratios $H/b = 2.0$ and $b/R = 3.0$ (Strahler et al., 1999). Parameter values
343	defined for canopy structure are somewhat arbitrary but are identical to our previous modeling efforts (Liu
344	et al., 2008; Xin et al., 2012). The effects of tree structural parameters on model simulations are further
345	explored in our study by varying their values.

Table 2. The spectral signature of leaf and soil background.

Spectral bands	Leaf reflectance ^a	Leaf transmittance ^a	Soil reflectance ^b
Photosynthetic active radiation	0.10	0.05	0.23
Near infrared	0.45	0.25	0.32

^a data from Bonan (2002)

^b data from Myneni et al. (1995)

350

Model validation for vegetation photosynthesis is performed with time series data for 8 successive days and 351 352 for entire years. Based on AmeriFlux biological data, measured LAI were 4.7 \pm 0.2 on DOY 211 in 2004 at 353 the US-Bar site and 4.84 \pm 0.78 on DOY 234 in 2006 at the US-Ha1 site. Because field-measured LAI data 354 were insufficient to support model simulation for an entire calendar year, we obtained satellite-derived LAI 355 from the MODIS (Moderate Resolution Imaging Spectroradiometer) products (Myneni et al., 2002). The 356 standard MODIS products (MOD15A2) provide 8-day LAI estimates at 1000 m spatial resolution, and we 357 derived 8-day mean LAI for a 3 \times 3 pixel window centered at each site. We screened cloudy observations 358 based on the Quality Control data in MOD15A2 and applied double logistic equations to fit time series of 359 cloud-free LAI observations (Li et al., 2014; Zhang et al., 2003).

4. Results

362 **4.1 Gap probability**

The gap probabilities derived from the GORT model are shown in Figure 2. As the solar zenith angle increases, more beams of sunlight are intercepted by leaves and tree crowns, resulting in decreased gap probabilities for both between- and within-crown gaps. As LAI increases, within-crown gaps decrease but between-crown gaps remain the same. The physical explanation underlying is simple: tree leaves are clumped within each individual crown such that variations in LAI would not affect between-crown gaps, which are only a function of crown shape, canopy structure, and illumination geometry.

369



370

Figure 2: Canopy gap probabilities modeled using GORT with varied leaf area index. The total gaps are
 between-crown gaps plus within-crown gaps. Tree structure parameters for the US-Bar site are used in
 model simulation.

Figure 3 further compares the gap probabilities modeled using GORT and Beer's law. For both models, gap probabilities decrease as solar zenith angle increases (Figure 3a). Modeled gap probabilities are close when

377 canopy LAI is low. However, at high LAI, the total gap derived from GORT is considerably greater than 378 that modeled using Beer's law due to strong clumping effects. With an LAI of 4.0, the differences in gap 379 probabilities are as much as 0.3 at the nadir, and in this case, more sunlight is allowed to transmit to the 380 ground surface in GORT than in classic radiative transfer models. Modeled vertical structures of sunlight 381 penetration are also shown to be different between GORT and Beer's law (Figure 3b). The gap probability 382 modeled using Beer's law decreases exponentially as canopy depth increases, whereas the decrease in the 383 GORT-modeled gap probability follows an inverse sigmoidal curve. The reason behind this can be explained by the geometric factor: classic radiative transfer models assume that leaves are randomly 384 385 distributed within the canopy layer, but the GORT model assumes that leaves are randomly distributed 386 within individual crowns. Due to the ellipsoidal shape of tree crowns, there are simply more leaves in the 387 canopy center than near the canopy top and canopy bottom, where the gap probability decreases more slowly. 388



Figure 3: Comparisons between canopy gap probabilities modeled using GORT and Beer's law as a function of a) solar zenith angle and b) canopy depth. The canopy depth is defined as the distance from canopy top to a canopy height (h). Tree structure parameters for the US-Bar site are used in GORT simulation.

395

396 4.2 Model simulations over 8-day time periods

Figure 4 shows each component of the radiation regime at the US-Bar site. The diffuse radiation modeled using Muneer's method matches flux tower measurements and accounts for 69.1% of the variances (Figure 4a). Because diffuse radiation was not measured at the US-Ha1 site, Muneer's method was implemented to partition global radiation into diffuse and beam components for US-Ha1. Using the measured beam and diffuse radiation, we simulate net radiation with GORT as a linear combination of two discrete bands at PAR and near infrared. Modeled net radiation is highly correlated with measured values ($R^2=0.981$), demonstrating the ability of GORT to model radiation absorption at the US-Bar site.





Figure 4: Measured and modeled components of radiation in 8 successive days are shown for a) the partition of global solar radiation, b) surface radiation balance, c) modeled and measured diffuse radiation, and d) modeled and measured net radiation. Extraterrestrial radiation is derived following methods outlined in

Allen et al. (1998). Muneer's method is applied to model diffuse radiation. The GORT model is applied to
 model net radiation. Data are shown from the Day of Year 217 to 224 in 2004 for the US-Bar site.

411

412 Time series of each component for modeling canopy photosynthesis are shown in Figure 5. Given that total 413 LAI remains the same over the course of several days, modeled sunlit and shaded LAI have little day-to-day 414 variability and only vary as a function of solar zenith angle (Figure 5a). As solar zenith angle decreases, 415 sunlit LAI increases but shaded LAI decreases. Because sunlit leaves receive more illumination, they have 416 less radiation limitations on photosynthesis than shaded leaves (Figure 5b). Temperature limitation generally 417 decreases from morning until noon, while VPD limitation increases. Although the chemical process of 418 photosynthesis favors higher temperatures, leaf stomata tend to close to reduce water loss when atmospheric 419 dryness is high (Bonan, 2002). Because short-term canopy CO₂ concentrations vary with winds and 420 convection between the ecosystem and the atmosphere, the ambient CO₂ concentrations exhibit the greatest variation from day to day (Figure 5b), so do the modeled differences between ambient and intercellular CO₂ 421 422 concentrations.



Figure 5: Time series of components of the photosynthesis calculation shown for a) sunlit and shaded leaf area index, b) environmental limiting factors imposed by radiation absorption, temperature, and vapor pressure deficit, and c) CO_2 concentration. Data are shown from the Day of Year 217 to 224 in 2004 for the US-Bar site.

Figure 6 shows time series of measured and modeled GPP for two sites over eight successive days. GPP estimates match flux tower measurements well in terms of the phase and amplitude. Daily peak GPP from tower measurements are over $30.0 \ \mu mol \ CO_2 \ m^{-2} \ s^{-1}$ for both sites. It is also evident that modeled results can capture some subtle variations in GPP at the hourly time scale. However, GPP estimates are slightly higher on DOY 242 but lower on DOY 243 for US-Ha1. Note that we used Muneer's method for estimating the diffuse radiation in US-Ha1 because measurements were not available. Considering uncertainties from the partition of global solar radiation, results for both sites perform well in general.



Figure 6: Time series of modeled and measured GPP for 8 consecutive days at the sites (a) US-Bar and (b) US-Ha1. Data are half-hourly at the US-Bar site and hourly at the US-Ha1 site. Data are shown from DOY 217 to 224 in 2004 for US-Bar, and from DOY 241 to 224 in 2006 for US-Ha1. Negative GPP measurements are set to zero. Missing points in modeled GPP at the US-Ha1 site are due to missing measurement of canopy CO2 concentrations or other meteorological variables.

Figure 7 statistically compares measured and modeled GPP. Our model is able to explain 84.0% and 88.3% of the GPP variances for the US-Bar and US-Ha1 sites, respectively. The regression lines are close to the 1 : 1 lines, and GPP is only slightly overestimated for US-Bar and underestimated for US-Ha1. The root mean squared errors (RMSE) are 3.71 and 3.08 μ mol CO₂ m⁻² s⁻¹ for US-Bar and US-Ha1, respectively. The overall model performance is high considering that we did not attempt to perform model calibrations.

449



Figure 7: Regressions between modeled and measured GPP for 8 consecutive days at the sites (a) US-Bar and (b) US-Ha1. Data are from DOY 217 – 224 in 2004 for US-Bar and from DOY 241 to 224 in 2006 for US-Ha1. Only data during the photosynthetically active period (flux tower GPP > 0.5 μ mol CO₂ m⁻² s⁻¹) are included in the regression. The solid lines denote the 1 : 1 lines, and the dashed lines denote the regression lines.

456

450

457 **4.3 Model simulation over entire years**

LAI derived from satellite observations (Figure 8) are used as inputs to model daily GPP over an entire year in addition to the 8-day model simulations. The double logistic fitting lines are shown to reduce noises in time series of MODIS LAI due to the effects of clouds and solar and viewing geometry. Fitted LAI time series are slightly higher from June to August and lower from September to December in 2006 at the USHa1 sites, but match with field measurements in general. The differences are likely to be introduced by
mismatched observation footprints and uncertainties in satellite retrieval algorithms. The fitted time series of
MODIS LAI are used for subsequent model simulations.



Figure 8: Comparisons of field-measured and satellite-derived leaf area indexes (LAI) for the sites a) USBar in 2004 and b) US-Ha1 in 2006. The solid grey lines denote MODIS LAI as obtained from standard
MODIS FPAR/LAI products (MOD15A2). The solid black lines denote double logistic fitting lines that are
applied to MODIS LAI. The solid points denote the measured LAI as obtained from biological datasets
from AmeriFlux website.

472

473 Figure 9 presents time series of measured and modeled GPP at the US-Bar site. Modeled results capture the

- trend and subtle variations of measured GPP on a daily basis. Most of the dips in the GPP time series occur
- 475 on cloudy days when radiation is the main limiting factor for vegetation photosynthesis. GPP values at US-
- 476 Bar are slightly overestimated from DOY 100 to 150 in 2004 possibly due to overestimation of the LAI.
- 477 Statistically, modeled results can explain 79.5%, 89.7%, and 89.3% of the variance in daily GPP for the
- 478 years 2004, 2005, and 2006, respectively (Figure 10). Regression slopes are close to the 1 : 1 lines except in
- the year 2004 due to overestimated GPP in the early growing season. The RMSEs are 1.64, 1.31, and 1.56
- 480 gC m^{-2} day⁻¹ for 2004, 2005, and 2006, respectively.



Figure 9: Time series of modeled and measured daily GPP shown for (a) 2004, (b) 2005, and (c) 2006 at the US-Bar site. Model simulation is performed at a half-hourly time step. Measured and modeled half-hourly GPP are aggregated to generate daily time series with units converted from μ mol CO₂ m⁻² s⁻¹ to gC m⁻² day⁻¹. Occational negative GPP measurements are set to zeros. Missing points in modeled GPP time series are

486 due to missing measurements of meteorological variables during the daytime photosynthetically active 487 period (flux tower GPP > $0.5 \mu mol CO_2 m^{-2} s^{-1}$). 488



Figure 10: Regressions between modeled and measured daily GPP shown for (a) 2004, (b) 2005, and (c) 2006 at the US-Bar site. Only data during the photosynthetically active period (flux tower GPP > 0.5 gC m⁻² day⁻¹) are included in the regressions. The solid line denote the 1 : 1 lines, and the dashed lines denote the regression lines.

495 Because measurements of atmospheric CO₂ concentrations within the canopy are largely unavailable for 496 US-Ha1 (only approximately 41.4% of the measurements are valid for use), we do not aggregate hourly 497 results to daily sums but perform regression analysis using all available hourly data in Figure 11. For the US-Bar site, the R² value is 0.801 and the RMSE value is 4.31 μ mol CO₂ m⁻² s⁻¹. For the US-Ha1 site, the 498 499 correlation between modeled and measured GPP is strong with an R² value of 0.777 and an RMSE value of 6.49 μ umol CO₂ m⁻² s⁻¹. There were slight GPP underestimates when measured GPP values are high at the 500 501 US-Ha1 site, possibly due to empirical functions that we used in modeling diffuse radiation and leaf 502 photosynthesis. Table 3 lists major statistical results for our model performance, as evaluated using all 503 available hourly data at both sites. The model performance is consistent through time and is comparable to 504 the simulation of 8-day data (Figure 7), despite the fact that satellite-derived LAI instead of field 505 measurements were used for yearly simulation.



508 Figure 11: Regressions between modeled and measured GPP for all available hourly data at the sites of a) 509 US-Bar and b) US-Ha1 in 2006. Only data from the photosynthetically active period are included in the 510 regression. The solid line denotes the 1 : 1 line, and the dashed line denotes the regression line.

511

512 Table 3. The model performance at two study sites as evaluated using hourly data. Units for root mean 513 square error (RMSE) and mean bias error (Bias) are in μ mol CO₂ m⁻² s⁻¹.

Voor		US-Bar			US-Ha1	
Tear	R 2	RMSE	Bias	R 2	RMSE	Bias
2001				0.804	5.44	2.00
2002				0.729	6.75	3.09
2003				0.781	5.62	2.85
2004	0.784	4.28	1.01	0.737	6.39	1.85
2005	0.795	4.11	0.47	0.736	6.83	1.18
2006	0.801	4.31	1.06	0.777	6.49	2.28
2007				0.768	6.21	2.50
2008				0.689	7.34	3.10
2009				0.662	7.62	3.68
2010				0.752	6.55	0.35
2011				0.715	6.96	1.34

514

515

516 5. Discussion

517 5.1 Influence of CO₂ concentration on canopy photosynthesis

518 One important question is whether it is necessary to link radiative transfer with leaf stomatal conductance

519 for modeling photosynthesis, since some biogeochemical models such as Production Efficiency Models

simply assume that vegetation GPP/NPP is linearly related to canopy radiation absorption (Xin et al., 2013).
To understand the performance of Production Efficiency Models, we conduct linear regressions between
modeled APAR and measured GPP as shown in Figure 12. Indeed, canopy APAR is positively related to flux
tower GPP and explains 70.3% of its variance. The R² value increases slightly to 0.710 after accounting for
the influences of temperature and vapor pressure. The model performance here is comparable to results from
other studies that evaluate Production Efficiency Models (Chen et al., 2011; Sjöström et al., 2013; Xin et al.,
2015).

527



528

Figure 12: Regressions between modeled absorbed photosynthetic active radiation (APAR) and measured GPP. Half-hourly data are shown from DOY 217 – 224 in 2004 for US-Bar. The influences of temperature and vapor pressure deficit are modeled based on Equations (51) and (52). Only data during the photosynthetically active period are included in the regression. The dashed lines denote the regression lines.

However, there are strong partial correlations between canopy CO2 concentrations and GPP even after accounting for radiation absorption. Figure 13a shows the residual plot of GPP versus ambient CO2 concentrations when controlling on APAR. The slope is negative because the ambient CO2 concentration, as regulated by vegetation photosynthesis and respiration activities, is normally high during the nighttime but low during the daytime. The correlation coefficient is only -0.279, but it is statistically significant (p-value < 0.001) under a one-tailed partial correlation test. The data clearly allow rejection of the null hypothesis that ambient CO₂ concentration has no effects on canopy photosynthesis. This relationship holds even after considering the factors of temperature and vapor pressure deficit (Figure 13b). We therefore conclude that accounting for the influence of ambient CO₂ concentrations is essential for modeling daytime GPP at the half-hourly time scale.

544



545

Figure 13: Residual plots are shown for a) the partial correction between GPP and ambient CO₂ concentration (C_a) while controlling for the variable of *APAR* and b) the partial correction between GPP and C_a - C_i while controlling for the variable of *APAR* × $f(T) \times f(VPD)$.

550 **5.2 Clumping effects in the GORT model**

The clumping effects of leaves modeled using GORT influence canopy radiative transfer processes and are worthy of further examination. Chen et al. (1997) demonstrated that the net effects of leaf clumping could be modeled by introducing a clumping index. We derive the clumping index by inverting their functions (Zhao et al., 2011) as follows:

$$\Omega = \ln(P_{gap}) / \ln(P_{Beer}) = -\ln(P_{gap}) / k_b LAI$$

where Ω is the clumping index, P_{gap} is the gap probability modeled using GORT, $P_{Beer} = \exp(-k_b LAI)$ is the gap probability modeled using Beer's Law, k_b is the extinction coefficient, and *LAI* is the leaf area index.

559 The behavior of the derived clumping index shown in Figure 14 is intuitively interpretable. Leaves are more clumped when LAI is larger given constant tree structures. However, when LAI is constant but tree density 560 561 increases, leaves are distributed in a larger three-dimensional space, resulting in an increased clumping 562 index. Similarly, if the H/b ratio or b/R ratio decreases while other parameters are unchanged, the total 563 crown volume increases and leaves are less clumped. The sensitivity of the clumping index to the 564 illumination zenith angle varies when using different parameter sets. Our simulated results are in line with 565 the measured and modeled results in previous studies (Leblanc and Chen, 2001; Leblanc et al., 2002): the clumping indexes are insensitive to zenith angles in some forest stands and increase with zenith angles in 566 others. We do not attempt to derive clumping indexes at solar zenith angle greater than 85° when gap 567 568 fractions typically approach zeros. These results imply that tree structure strongly influences radiation 569 absorption and photosynthesis of canopies.

570

(58)



Figure 14: Derived clumping index as a function of solar zenith angle for varied canopy parameters. Tree parameters for US-Bar are used for GORT simulations. The default simulation is for a canopy composed of $H/b = 2.0, b/R = 3.0, \lambda = 1432$ trees/ha, and LAI = 2.0, and labeled curves are for the same case with only the labeled parameters varied.

577 **5.3 Assumptions and future improvements**

It is also necessary to review our model assumptions and identify possible avenues for future improvements. 578 First, we assume a spherical leaf angle distribution in the model simulations. However, most deciduous 579 580 forests have semi-horizontal leaf orientation (Bonan, 2002) and an assumption of planophile or plagiophile 581 LAD is likely to be more appropriate for temperate and boreal broadleaf forests (Pisek et al., 2013). Because 582 LAD influences the proportions of sunlit and shaded leaf areas, the way in which modeled canopy GPP 583 varies with LAD requires further exploration. Second, the substrate under the canopy layer is assumed to be 584 a Lambertian surface. Field studies have observed the effects of bi-directional reflectance distribution 585 function (BRDF) for soils (Liang and Townshend, 1996; Wang et al., 2010), and coupled soil and vegetation 586 model (Ni and Li, 2000; Verhoef and Bach, 2007) should be tested to understand the effects of soil BRDF on canopy photosynthesis. Third, we assume maximum constant leaf stomatal conductance over the 587

growing season. It is worth examining how optimal leaf stomatal conductance may evolve with leaf 588 589 development stages and long-term environmental changes (Keenan et al., 2013; Lammertsma et al., 2011). 590 Fourth, we use ellipsoids to describe tree crown shapes for deciduous broadleaf forests. Because many 591 evergreen needleleaf forests have conical crowns, future applications to areas with conifer forests may 592 require different treatment on crown shapes in the models. Finally, our linkage between radiative transfer 593 and biochemical processes is still empirical. We may need to test other mechanisms, for example, the 594 biochemical model based on the enzyme kinetics of rubisco and the regeneration of RuBP in response to 595 light absorption (Farguhar and Sharkey, 1982), in future studies.

596

597 6. Conclusion

598 We propose and validate a new model that links GORT with biochemical processes for modeling canopy 599 photosynthesis. Several main conclusions can be drawn from this study. First, the radiative transfer process 600 within the canopy is one of the key factors in modeling vegetation photosynthesis, and our proposed model 601 simulates canopy photosynthesis well. Modeled GPP robustly explained approximately 80% or more 602 variance in GPP measurements at both half-hourly and daily time scales. Second, tree structures influence 603 canopy gap probabilities and vegetation photosynthesis. Leaf clumping could vary as a function of tree 604 density, canopy depth, and crown shapes and affect canopy sunlight interception. Finally, ambient CO₂ 605 concentrations influence vegetation photosynthesis activities and should be included in biogeochemical 606 models.

607

Accurate modeling of vegetation photosynthesis is essential for improving our understanding of the global carbon cycle. The model we developed is complementary to classic radiative transfer models, especially in sparse and intermediate forest stands. Although more validation efforts are required, the GORTphotosynthesis model is promising in terms of simulating photosynthesis for discontinuous plant canopies.

613 Appendix A:

614 Table A1. Nomenclature

Symbols	Definition
$P_{\rm gap}(h, \theta_i)$	total gap probability for beam light passing through the canopy
$P_{\rm gan}(n=0 h,\theta_i)$	gap probability for beam light passing through the canopy without reaching any crowns
$P_{\text{gap}}(n > 0 h, \theta_i)$	gap probability for beam light passing through crowns without being intercepted by leaves
$P(s h,\theta_i)$	probability distribution function associated with within-crown path length
$P(s n, z, h, \theta_i)$	probability distribution of within-crown path length given that a solar ray enters the crown at
	height h and angle θ_i
$P(n z, h, \theta_i)$	probability distribution of the numbers of crowns intercepted by the solar ray incident at angle
	θ_i , entering crowns at height z, and then traveling to height h
$\lambda_{ m v}$	tree density (m ⁻²)
V_{Γ}	projected cylinder volume starting from the canopy top and extending to certain height
$\tau(\theta_i, \alpha)$	projected foliage area volume density (m ⁻)
$K_b(\theta_i, \alpha)$	extinction coefficient for diffuse rediction
K_d K (b)	canony openness factor to diffuse radiation
$K_{\text{open}}(n)$ $K_{\text{open}}(n-0 h)$	batwaan grown openness factor
$K_{\text{open}}(n = 0 h)$	within grown openness factor
$\Lambda_{\text{open}}(n > 0 n)$	within-crown openness factor the first derivative of conpress hability \mathbf{P}_{i} (b, 0) with respect to height
$P_{gap}(n, \sigma_i)$	the first derivative of gap probability $P_{\text{gap}}(n, \theta_i)$ with respect to height
$K_{\text{open}}(n)$	the first derivative of the openness factor $K_{open}(n)$ with respect to neight
$t_0(n, \theta_i)$	the proportion of unintercepted direct beam for semi-infinite homogeneous canopies
K_{ff}	directional homionhomion reflectence for somi infinite homogeneous cononios
R_{df}^{-}	directional-nemispherical reflectance for semi-infinite nomogeneous canopies
T_{ff}	hemispherical-hemispherical transmittance for semi-infinite homogeneous canopies
T_{df}^{∞}	directional-hemispherical transmittance for semi-infinite homogeneous canopies
$\rho_{ff}(h)$	hemispherical-hemispherical reflectance for homogeneous canopies with finite thickness
$\rho_{df}(h, \theta_i)$	directional-hemispherical reflectance for homogeneous canopies with finite thickness
$t_{ff}(h)$	hemispherical-hemispherical transmittance for homogeneous canopies with finite thickness
$t_{df}(h, \theta_i)$	directional-hemispherical transmittance for homogeneous canopies with finite thickness
$\rho_{ff}(h)$	hemispherical-hemispherical reflectance for discontinuous canopies
$\rho'_{df}(h, \theta_i)$	directional-hemispherical reflectance for discontinuous canopies
$t'_{ff}(h)$	hemispherical-hemispherical transmittance for discontinuous canopies
$t_{df}'(h, heta_i)$	directional-hemispherical transmittance for discontinuous canopies
$\delta LAI(h)$	leaf area index within a thin layer δh at height h
LAI	total leaf area index of the canopy
$LAI_{Sun}(\theta_i)$	sunlit leaf area index given a solar illumination angle θ_i
$LAI_{Shd}(\theta_i)$	shaded leaf area index given a solar illumination angle θ_i
$\Delta LAI_{Sun}(\theta_i)$	summer leaf area for nonlogeneous canoples given a solar multimation angle θ_i
θ_i	azimuth angle
φ	leaf single scattering albedo
ν	$\sqrt{1-\sigma}$
ι U;	$\cos(\theta_i)$
ρι	leaf reflectance
τ_l	leaf transmittance
ρ_s	soil reflectance
ρ_{cb}	canopy reflection coefficient for beam irradiance
ρ_{cd}	canopy reflection coefficient for diffuse irradiance

f_h	the fraction of incident beam radiation in total or global incoming solar radiation
$I_{h}(h,\theta_{i})$	unintercepted beam fluxes at canopy height h given a solar illumination angle θ_i
$I_d(h)$	unintercepted diffuse fluxes at canopy height h
$I_{ht}(h, \theta_i)$	unintercepted and down scattered beam fluxes
$I_{dt}(h)$	unintercepted and down scattered diffuse fluxes
I _c	total radiation absorbed by canopy elements
I _{ch}	beam radiation absorbed by canopy elements
Icd	diffuse radiation absorbed by canopy elements
Isun	total radiation absorbed by sunlit leaves
Isumb	beam radiation directly absorbed by sunlit leaves
Isumbs	down scattered beam radiation absorbed by sunlit leaves
Isund	diffuse radiation absorbed by sunlit leaves
Isum	total radiation absorbed by shaded leaves
O_{sum}	total radiation absorbed by sunlit leaves per leaf hemi-surface area
O_{shd}	total radiation absorbed by shaded leaves per leaf hemi-surface area
A	leaf-level CO ₂ assimilation rate
a _c	stomatal conductance
C_{α}	ambient CO ₂ concentrations
C _i	intercellular CO ₂ concentrations
q_{cSun}	stomatal conductance for sunlit leaves
g _{cshd}	stomatal conductance for shaded leaves
g _{cmar}	maximum leaf stomatal conductance when environmental factors do not limit carbon uptake
$f(x_i)$	scalars that account for the influences of environmental stresses on leaf stomatal conductance
f(0)	scalars that account for the influences of solar radiation on leaf stomatal conductance
f(T)	scalars that account for the influences of temperature on leaf stomatal conductance
f(VPD)	scalars that account for the influences of vapor pressure deficit on leaf stomatal conductance
k_c	stress coefficients of PAR absorbed by plant leaves for the temperature scalar
k_0	stress coefficients of PAR absorbed by plant leaves for the temperature scalar
T_{min}	minimum temperature for photosynthetic activities
Tmar	maximum temperature for photosynthetic activities
Tont	optimum temperature for photosynthetic activities
VPD	ambient vapor pressure deficit
VPDmin	minimum vapor pressure deficit
VPDmax	maximum vapor pressure deficit
VPDo	an empirical constant describing the species sensitivity to ambient vapor pressure deficit
Г	leaf CO ₂ compensation point
m_I	regression coefficient for ambient and intercellular CO_2 concentrations related to tree species
A_{Sun}	leaf-level CO ₂ assimilation rate for sunlit leaves
A _{shd}	leaf-level CO ₂ assimilation rate for shaded leaves
K _t	hourly clearness index
I_0	total or global incoming solar radiation on a horizontal plane at the canopy top
I _e	extraterrestrial solar radiation
Ω	foliage clumping index
P_{Beer}	gap probability for beam light passing through the canopy as modeled using Beer's Law
DUU	

616 Table A2. Values for model parameters

1 4010 1 12							
Symbols	Value	Units	Reference				
k _C	500	W / m ²	Ding et al. (2014)				
k_Q	150	W / m $^{\rm 2}$	Ding et al. (2014)				
T_{min}	0	$^{\circ}$ C	Kalfas et al. (2011)				
T_{max}	45	${}^{\mathbb{C}}$	Kalfas et al. (2011)				

T _{opt}	25	${}^{\mathfrak{C}}$	Kalfas et al. (2011)
VPD _{min}	0.65	kPa	Heinsch et al. (2003)
VPD _{max}	4.6	kPa	Heinsch et al. (2003)
VPD ₀	30	kPa	Katul et al. (2000)
Γ	40	µmol/mol	Katul et al. (2000)
m_L	4.0		Katul et al. (2000)

618 Author contribution

619 Qinchuan Xin developed the model code and performed the simulations. Qinchuan Xin designed the 620 experiments and Wenyu Li contributed to data analysis. Qinchuan Xin and Peng Gong prepared the 621 manuscript with contributions from all co-authors.

622

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