Wide Discrepancies in the Magnitude and Direction of Modelled SIF in Response to Light Conditions

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Abstract:

Recent successes in passive remote sensing of far-red solar induced chlorophyll fluorescence (SIF) have spurred development and integration of canopy-level fluorescence models in global terrestrial biosphere models (TBMs) for climate and carbon cycle research. The interaction of fluorescence with photochemistry at the leaf- and canopy- scale provides opportunities to diagnose and constrain model simulations of photosynthesis and related processes, through direct comparison to and assimilation of tower, airborne, and satellite data. TBMs describe key processes relating to absorption of sunlight, leaf-level fluorescence emission, scattering and reabsorption throughout the canopy. Here, we analyze simulations from an ensemble of process-based TBM-SIF models (SiB3, SiB4, CLM4.5, CLM5.0, BETHY, ORCHIDEE, BEPS) at a subalpine evergreen needleleaf forest near Niwot Ridge, Colorado. These models are forced with tower observed meteorological data, and analyzed against continuous far-red SIF and gross primary productivity (GPP) partitioned eddy covariance data at diurnal and synoptic scales during the growing season (July-August 2017). Our primary objective is to summarize the site-level state of the art in TBM-SIF modeling over a relatively short time period (summer) when light, structure, and pigments are similar, setting the stage for regional- to global-scale analyses. We find that these models are generally well constrained in simulating photosynthetic yield, but show strongly divergent patterns in the simulation of absorbed photosynthetic active radiation (PAR), absolute GPP and fluorescence, quantum yields, and light response at leaf and canopy scale. This study highlights the need for mechanistic modeling of non-photochemical quenching in stressed and unstressed environments, and improved representation of light absorption (APAR), distribution of sunlit and shaded light, and radiative transfer from leaf to canopy scale.
Section 1: Introduction

Our ability to estimate and measure photosynthesis beyond the leaf scale is extremely limited. This inhibits the ability to evaluate the performance of terrestrial biosphere models (TBMs) that are designed to quantify the direct impact and feedbacks of the carbon cycle with climate change. Consequently, there are substantial uncertainties in estimating the gross primary production (GPP) response to environmental changes and carbon-climate feedback (Friedlingstein et al., 2014). Global, multi-scale remote sensing of solar induced fluorescence (SIF) may represent a major breakthrough in alleviating this deficiency (Mohammed et al, 2019). Spaceborne data indicate a linear relationship between SIF and GPP at large spatial (kilometer) and temporal (bi-weekly) scales (e.g., Sun et al., 2017) for several ecosystems, while ground-based measurements indicate a more non-linear relationship at leaf and canopy scales (Zhang et al., 2016; Gu et al., 2019; van der Tol et al., 2014; Magney et al., 2017, 2019a).

Chlorophyll fluorescence is re-emitted energy produced during the photosynthetic light reactions, in which a small fraction (roughly 2%) of photosynthetic active radiation (PAR) absorbed by chlorophyll is re-emitted at longer wavelengths (650-850 nm) as fluorescence. In ambient conditions, the emission of SIF represents a by-product of two primary de-excitation pathways, photochemical and nonphotochemical quenching (PQ, NPQ). Plants have evolved these regulatory mechanisms to prevent damage to photosynthetic machinery when the amount of absorbed radiation is greater than that which can be used to drive photochemistry. Chlorophyll fluorescence responds dynamically to changes in photochemistry and NPQ from instantaneous to hourly, daily, and seasonal timescales, as a function of changing environmental conditions and plant structural properties (Porcar-Castell et al., 2014; Demmig-Adams et al., 2012). SIF is fundamentally different than steady-state fluorescence yield typically measured at the leaf scale as it is sensitive to both changes in photochemistry as well as absorbed PAR (APAR, related to incident light, canopy structure, and biochemical content). The response of canopy SIF to APAR is well documented in deciduous and evergreen forests and cropping ecosystems (Yang et al., 2018; Badgley et al, 2017; Miao et al., 2018; Magney et al., 2019b; Li et al., 2020). More recently, Magney et al. (2019b) showed that seasonal changes in canopy SIF for cold climate evergreen...
systems is influenced by changes in needle physiology and photoprotective pigments (Magney et al., 2019b).

To properly account for these factors, process-based SIF models must represent these underlying non-linear biophysical and chemical processes. Several modeling groups have adapted TBMs to incorporate various SIF formalisms for the purpose of model evaluation, data assimilation, and improved model prediction (Lee et al., 2015; Koffi et al., 2015; Thum et al., 2017; Norton et al., 2019; Bacour et al., 2019; Raczka et al., 2019). With these goals in mind, TBM SIF modeling requires two important steps: (1) a representation of SIF at the needle/leaf scale that accounts for NPQ and photochemistry, and (2) canopy radiative transfer of SIF, which enables a comparison to large field-of-view observations (e.g. tower, satellites). The second step involves accounting for radiative transfer within the canopy and has typically relied on incorporating the Soil Canopy Observation Photosynthesis Energy model (SCOPE, van der Tol et al., 2009, 2014), which simulates chlorophyll fluorescence as a function of biophysics, canopy structure, environmental conditions, and sun/sensor geometries. This approach has been adopted by TBMs in various ways using different assumptions for fluorescence modeling and radiative transfer, as will be discussed in Section 2.

Typically, measuring chlorophyll fluorescence and competing pathways (PQ, NPQ) has been done at the leaf scale via pulse-amplitude modulation fluorescence (PAM, Schreiber et al., 1986). Recently, commercially available spectrometers have made it possible to measure SIF directly in the field at the leaf and canopy scale, and also enable the study of structural, environmental, and directional controls (Cogliati et al. 2015; Daumard et al. 2010; Migliavacca et al. 2017; Yang et al. 2015; Grossman et al., 2018; Gu et al., 2019b; Zhang et al., 2019). The use of field deployable instruments on eddy covariance towers has increased rapidly since 2014, providing coverage of multiple vegetation types across various climates around the world (Yang et al., 2018; Magney et al., 2019a,b; Parazoo et al., 2019). These data enable improved understanding of the relationship between SIF, GPP, APAR, and environmental effects at leaf to canopy scales. Novel tower-mounted spectrometer systems such as Fluospec2 (Yang et al., 2018) and Photospec (Grossman et al., 2018) have made it possible to monitor canopy SIF continuously in the field with high precision over multiple years providing opportunities for more direct comparison and evaluation.
of satellite data (Grossman 2018; Yang 2015, 2018; Magney et al., 2019). PhotoSpec offers the additional benefits (and challenge) of (a) precise field of view capable of resolving leaf-level SIF, and (b) canopy scanning at azimuth and elevation angles. These features enable SIF integration from leaf- to canopy- scales, and interpretation of directional variations of the emitted radiance.

Canopy scanning spectrometers such as PhotoSpec thus provide an opportunity to understand the physical processes that lead to a breakdown of SIF-GPP linearity at leaf to canopy scale (or conversely, emergence of linearity at increasing scale), and for detailed evaluation and diagnosis of TBM performance. This study provides a preliminary benchmarking site-level assessment for simulations of SIF within a TBM framework and across an ensemble of TBMs, with the primary purpose being an initial investigation into the response of modelled SIF and GPP to light during peak summer. We leverage continuous measurements of SIF and GPP at the Niwot Ridge US-NR1 Ameriflux flux tower in Colorado from June-July 2017 (Magney et al., 2019b) to (1) Benchmark TBM-SIF modeling, (2) Evaluate sensitivity to underlying processes and scaling techniques, (3) Identify strengths and weaknesses in current modeling strategies, and (4) Recommend strategies for models and observations.

The paper is organized as follows: Section 2 describes the seven TBM-SIF models (SiB3, SiB4, ORCHIDEE, BEPS, BETHY, CLM4.5, CLM5) which have recently been published or are in review, and provides more details on site level benchmarking observations. Section 3 summarizes results comparing modelled and predicted SIF and GPP at hourly and daily scales, as they relate to absorbed light, GPP and SIF yields, and quantum yields. Section 4 discusses results in more detail, including attribution of SIF magnitude and temporal phasing biases and sensitivities to absorbed light, and areas for improvement.

Section 2: Methods

2.1 Site: Niwot Ridge, Colorado

Our study focuses on an AmeriFlux (https://ameriflux.lbl.gov/) site in Niwot Ridge, Colorado, USA (US-NR1), where a tower-based eddy covariance system has been continuously measuring the net ecosystem exchange of carbon dioxide (NEE) over a high-elevation subalpine forest since 1999, and a spectrometer system that has been continuously monitoring SIF since June.
2017 (Grossman et al., 2018; Magney et al., 2019b). The 26-m tall tower is located in a high
elevation forest (3050 m asl) located in the Rocky Mountains of Colorado (Burns et al., 2015; Hu
et al., 2010; Monson et al., 2002) and consists primarily of the evergreen species of lodgepole
pine (*Pinus contorta*), Engelmann spruce (*Piceae engelmanii*), and subalpine fir (*Abies
lasiocarpa*). The mean annual temperature is 1.5°C and mean annual precipitation is 800 mm
(65% as snow). The forest is roughly 120 years old with a mean canopy height of 11.5 m, and a
leaf area index of 4.2 m^2 m^{-2}. More site-specific details can be found in Burns et al. (2015).

At Niwot Ridge, interannual variations in GPP are closely linked to winter snowfall amount, which
typically melts by early June, and summer precipitation, characterized by afternoon convective
thunderstorms triggered by upslope flow (Burns et al., 2015; Albert et al., 2017) and
climatological peak precipitation around 2 pm local time (Fig 1A). We note that our study period
of July-August 2017 is unusual for NR1 (relative to the 2015-2018 mean) in its bimodal
distribution of diurnal precipitation (morning and afternoon peaks), lower than normal afternoon
precipitation, cooler temperatures, and reduced vapor pressure deficit (Fig 1 A-C). The early
morning peak is due to a strong storm system that moved through from July 22-24 (Fig 1E), and
does not show up when these days are removed. This period also shows a decrease in incoming
shortwave relative to climatology despite lower precipitation (Fig 1D). We note that a second
storm passed through in early August. The combination of these two storms produced net
decreases in air temperature (Fig 1F), vapor pressure deficit (Fig 1G) and sunlight (Fig 1H) over a
two-week period from late July to early August.

2.2 Tower-Based Measurements: PAR, SIF, CO_2 Flux

2.2.1 Absorbed PAR

The site is equipped with two main upward-facing PAR sensors. The first (LICOR LI-190R),
mounted on the PhotoSpec telescope unit, provides an independent measurement of
direct/diffuse light and can be used to calibrate PhotoSpec (Grossman et al., 2018). The second
(SQ-500-SS; Apogee Instruments), mounted on the main flux tower, is part of a larger array of
upward- and downward-oriented PAR sensors above and below the canopy used for the
calculation of the fraction of PAR absorbed by the vegetation canopy (fAPAR). The two PAR
sensors show a similar diurnal pattern during July-August 2017 (Fig S1), including an afternoon dip and relatively smaller values overall compared to 2018 (the only other year with available PAR for comparison).

Full-spectrum quantum sensors (SQ-500-SS; Apogee Instruments) were new and factory-calibrated together just before installation. Above-canopy sensors (one up and one down-facing) were mounted on the main flux tower, and below-canopy sensors (six up and six down) were mounted at the 2 m height above ground on a shorter canopy-access towers. APAR was calculated for each pair of below-canopy relative to above-canopy sensors for every half-hour, then averaged among sensors over daylight hours to create a daytime average. We then estimate hourly APAR by multiplying hourly incoming PAR (measured and integrated from 400-700 nm) at the top of canopy (PAR) by the daytime average of fAPAR. Fig S2 shows the mean diurnal cycle for July-August 2017 for each sensor, and the across-sensor average, with APAR data collection beginning on July 13, 2017.

2.2.2 Tower Based Measurements of Solar Induced Chlorophyll Fluorescence (SIF)

SIF data has been collected from a scanning spectrometer (PhotoSpec) installed at the AmeriFlux US-NR1 tall tower since June 17, 2017. PhotoSpec sits atop the tower at 26 m above the ground and roughly 15 m above the forest canopy top, transferring reflected sunlight and SIF data collected from the needleleaf canopy through a tri-furcated optical cable to three spectrometers in a shed at the base of the tower. These spectrometers measure far-red fluorescence in the 745-758 nm retrieval window at high spectral resolution (FWHM = 0.3 nm) and with a 0.7 deg field of view (FOV), resulting in a 20-cm diameter footprint at nadir on top of the canopy. The far-red SIF data are then scaled to 740 nm for model intercomparison using the first principal component of the spectral shape in Magney et al., 2019a. Photospec scans from nadir to the horizon in 0.7 degrees steps at two azimuth directions, with a time resolution of ~20 s per measurement and complete scan time of 20 minutes. For this study, we aggregate scans across all azimuth and elevation angles into hourly, canopy level averages to benchmark model estimates of top of canopy (TOC) or canopy averaged SIF (BETHY only, see Sec 2.3.4.1) at diurnal and synoptic time scales. We refer the reader to Grossman et al. (2018) and Magney et al (2019b) for further details.
regarding PhotoSpec, implementation at US-NR1, and data filtering. A two-month data collection gap in fall of 2017 limits our model-data analysis to the 2017 growing season (July-August, 2017).

Diurnal composites of PhotoSpec SIF in 2017 show a late morning peak and afternoon dip (Fig S3A). The afternoon dip is consistent with decreased incoming shortwave, PAR and APAR (Figs S1 and S2, respectively). However, we note the retrieved signal from PhotoSpec is also affected by (1) viewing geometry, (2) fraction of sunlit vs shaded leaves (sun/shade fraction, i.e. the quantity of needles illuminated by incident sunlight) due to self-shading within the canopy, and (3) direct/diffuse fraction due to cloud cover. Structural and bidirectional effects lead to different SIF emission patterns depending on view angle and scanning patterns (Yang and van der Tol, 2018). The viewing geometry of PhotoSpec (as implemented at NR1 in 2017) causes a higher fraction of illuminated vegetation in the morning, which leads to a 2 to 3 hour offset in the timing of peak SIF (Fig S3A) and incoming far-red reflected radiance within the retrieval window (Fig S3B), from the peak zenith angle of the sun at noon (coinciding with the expected peak in PAR) to late morning. Normalizing SIF by far-red reflected radiance as relative SIF (SIF$_{rel}$, Fig S3C) and rescaling to SIF (Fig S3D) shifts the peak back to noon and preserved the afternoon dip (albeit with reduced magnitude). SIF$_{rel}$ helps to account for factors 1-3 listed above because it accounts for the amount of reflected radiation in the field of view of PhotoSpec, which is impacted by canopy structure, sun angle, and direct/diffuse light. SIF$_{rel}$ is likely a better approximation of SIF$_{yield}$ because we are normalizing for the exact amount of ‘illuminated’ canopy elements in each retrieval, whereas the APAR measurements are integrated for the entire canopy. As such, we expect SIF$_{rel}$ to have a strong seasonal change associated with downregulation of photosynthesis, and a more subtle diurnal change, as during mid-summer the SIF signal is primarily driven by light intensity.

It is important to note that the PhotoSpec system is highly sensitive to sun/shade fraction in the canopy (factor 2) due to the narrow FOV of the PhotoSpec telescoping lens. Increased afternoon cloud cover during summer causes diurnal asymmetry in incident PAR (Fig S1A). We examine this effect in more detail (Section 3) by analyzing SIF and GPP under clear and diffuse sky conditions using a threshold (0.5, top-of-canopy/top-of-atmosphere incoming shortwave radiation) similar to that used in Yang et al. (2017) and Yang et al. (2018).
NEE measurements are screened using $u_{\text{star}}$ filtering, and partitioned into gross primary production (GPP) and terrestrial ecosystem respiration components using the so-called nighttime method which is based on the relationship between NEE during the nighttime ($\text{PAR} < 50 \text{ umol m}^{-2} \text{ s}^{-1}$) and air temperature (Reichstein et al., 2005). Diurnal averages of GPP based on nighttime partitioning show similar diurnal structure to PAR and SIF including the afternoon dip and reduced overall magnitude compared to the 2015-2018 mean (Fig S4). Similar results are found using daytime light partitioning of NEE (Lasslop et al., 2010; Fig S4) and thus only nighttime partitioned GPP data are reported for the remainder of this study. All GPP estimates are processed as half hourly means, then gap filled and averaged hourly. Details on the flux measurements, data processing and quality control are provided in Burns et al. (2015).

## 2.3 Modeling Approach

### 2.3.1 TBM-SIF Overview

The parent TBMs are designed to simulate the exchanges of carbon, water, and energy between biosphere and atmosphere, from global to local scales depending on inputs from meteorological forcing, soil texture, and plant functional type. The addition of a fluorescence model that simulates SIF enables a direct comparison to remotely sensed observations for benchmarking, process diagnostics, and parameter/state optimization (data fusion) for improved GPP estimation. The TBM-SIF models analyzed here differ in ways too numerous to discuss. We refer the reader to the appropriate references in Section 2.3.4 for more detailed model descriptions. Instead, we focus on key differences affecting joint simulation of GPP and leaf/canopy level SIF at diurnal and synoptic scale, during the peak of summer. These differences, which are summarized in Table 1, include the representation of stomatal-conductance (all use Ball-Berry except CLM5.0, BEPS, and ORCHIDEE), canopy absorption of incoming radiation, limiting factors for photosynthesis ($V_{\text{cmax}}, \text{LAI}, \text{radiation}, \text{stress}$) and SIF ($k_N$), scaling and radiative transfer methods for transferring leaf-level SIF simulations to top of canopy, and parameter optimization. Further details on (a) photosynthetic structural formulation and parameter choice, (b)
representation of leaf level processes important to SIF ($k_N$ and $\phi_P$), and (c) leaf-to-canopy scaling approach ($SIF_{canopy}$) are provided in Sections 2.3.2 and 2.3.3.

2.3.2 Photosynthesis Models

All TBM-SIF models in this manuscript used enzyme-kinetic models to simulate leaf assimilation rate (gross photosynthesis) as limited by the efficiency of photosynthetic enzyme system, the amount of PAR captured by leaf chlorophyll, and the capacity of leaves to utilize end products of photosynthesis (Farquhar et al., 1980; Collatz et al., 1991, 1992; Sellers et al., 1996). However, there are important differences in the representation of (a) stomatal conductance that couples carbon/water cycles, and (b) limiting factors on carbon assimilation due to leaf physiology (maximum carboxylation capacity, Vcmax), radiation (APAR or fAPAR), canopy structure (LAI, leaf angle distribution), and stress (water supply and demand, temperature), that affect plant physiological processes and canopy radiative transfer. The underlying stomatal conductance models in the TBMs analyzed here are represented by the Ball-Berry family of empirical models rooted in the leaf gas exchange equation but with different representations of atmospheric demand (relative humidity or vapor pressure deficit), including the Ball-Berry model (Ball et al., 1987), the Ball-Berry-Leuning model (Leuning, 1995), the Yin-Stuik model (Yin and Struik, 2009), and the Medlyn model (Medlyn et al., 2011). These structural and parametric differences also influence calculated values such as the degree of light saturation (Section 2.3.3), which influence both the fluorescence and quantum yield as used by the fluorescence models. Differences in stomatal conductance, canopy type / radiation scheme, stress, Vcmax, and LAI are summarized in Table 1.

2.3.3 Fluorescence Modeling Approach

Following the general approach described in Lee et al. (2015) and van der Tol et al. (2014), the flux of total leaf-level emitted fluorescence, $SIF_{leaf}$, can be diagnosed using a light use efficiency framework analogous to the expression for photosynthesis (Monteith et al., 1972),

$$SIF_{leaf} = fAPAR \cdot PAR \cdot \phi_F$$

$$= APAR \cdot \phi_F$$  \hspace{1cm} \text{Equation 1}
where \( PAR \) and \( f\text{APAR} \) are defined in Section 2.2.1 but measured at leaf level, and \( \phi_F \) is the quantum yield of fluorescence, representing the number of photons emitted by fluorescence per absorbed photon. We note that photosystems I and II (PS1 and PSII, respectively) contribute to leaf level fluorescence but only PSII is considered in models analyzed here (with the exception of ORCHIDEE, Section 2.3.4.2). \( \phi_F \) is estimated as follows:

\[
\phi_F = \frac{k_F}{k_F + k_D + k_N} (1 - \phi_p)
\]

Equation 2

where \( k \) represents the rate coefficients for the different pathways for the transfer of energy from excited chlorophyll \( (k_F = \text{fluorescence}, k_D = \text{heat dissipation}, \text{and } k_N = \text{non-photochemical quenching}, \text{or NPQ}), \) and \( \phi_F \) is the quantum yield of electron transport (see Section 2.3.2). \( k_F \) is typically set to a constant value \((0.05)\) in models following van der Tol et al. (2014). \( k_D \) is also typically set to a constant value of 0.95, or temperature corrected in some cases (e.g., ORCHIDEE, CLM4.5, CLM5.0). \( k_N \) has a substantial and variable impact on energy partitioning at diurnal and seasonal scales which varies as a function of light saturation (e.g., Raczka et al., 2019; Porcar-Castell et al., 2011). Once leaf level emissions are known, an approach is needed estimate the total TOC fluorescence flux \((SIF_{\text{canopy}})\) for comparison to Photospec data. Leaf and canopy level fluorescence modeling is described in more detail in Section 2.3.3.1 and 2.3.3.2 below.

2.3.3.1 Leaf level SIF emission

The ‘quantum yield’ approach has been used to in SIF models to characterize the fraction of photons that are used for PQ, NPQ, or re-emitted as fluorescence (van der Tol 2014). It is important to note, that this does not translate into the actual amount of SIF emission leaving the leaf, but is used as an approximation. TBM-SIF models typically represent \( \phi_F \) using lake model formalism, which assumes large connectivity between photosynthetic units (Genty et al., 1989; van der Tol et al., 2014). \( \phi_F \) is expressed in terms of the degree of light saturation \((x)\), derived from the native photosynthesis module of the parent TBM and represents the balance between actual and potential electron transport rates, and the maximum photochemical yield under dark-acclimated conditions \((\phi_{P,max})\), which is derived from the fluorescence model and defined in terms of rate coefficients in Eq 2.
accounts for the ability of plants to dissipate excess energy as heat via NPQ through the regulation of xanthophyll cycle pigments (Demmig-Adams and Adams, 2006). NPQ can be represented as a sum of reversible ($k_R$) and sustain ($k_S$) components ($k_N = k_R + k_S$). $k_R$ accounts for the relatively fast (diurnal), reversible NPQ response to light. $k_S$ accounts for the relatively slow (seasonal), sustained NPQ response to light and other environmental factors. With the exception of CLM4.5, models do not typically account for $k_S$.

A significant challenge in fluorescence models is to find an appropriate relationship between $k_N$ and the degree of light saturation ($\chi$). The TBM-SIF models represent $k_N$ through an approach similar to the one used in SCOPE, which uses a parametric model of $k_N$ derived from PAM fluorometry measurements (van der Tol et al., 2014).

NPQ models can be classified as stressed (drought) and unstressed relative to water availability depending on the dataset from which empirical fits are derived. The unstressed model is ideal for irrigated systems such as crops, and the stressed model is more appropriate for water limited ecosystems such as Niwot Ridge. We examine each of these models using drought and unstressed models from van der Tol (2014), and a drought-based model from Flexas et al. (2002). These models use different empirical fits but are otherwise identical. In general, $k_N$ increases more rapidly with APAR (light saturation), and ramps up to a higher level, in the drought-based model compared to the unstressed model. Additionally, some models provide unique improvements such as dependence on environmental conditions (e.g., water stress vs no water stress in ORCHIDEE), and equations for reversible and sustained NPQ to represent the different time scales (minutes to seasonal) at which NPQ regulation occurs (e.g., CLM4.5) influenced by pigmentation changes in the leaf.

2.3.3.2 Leaf-to-Canopy scaling

The TBM-SIFs produce leaf-level fluorescence which needs to be converted to canopy-level fluorescence ($\text{SIF}_{\text{canopy}}$) to be directly compared to PhotoSpec and satellite observations. Leaf-to-canopy-level conversion of SIF requires a representation of canopy radiative transfer, which in general is too computationally expensive to include within the TBM models in this study, that are designed for global scale application. Therefore, most TBM models analyzed here account for canopy...
radiative transfer of SIF using some representation of SCOPE (van der Tol 2009a,b). The most commonly used approach is to run independent simulations of SIF from SCOPE to create an empirical conversion factor ($k_{740}$) between leaf and canopy level SIF that is a function of Vcmax (Lee et al., 2015). This conversion factor accounts for integration over the fluorescence emission spectrum, observation angle, and unit conversion. Model variations of this empirical approach, as well as additional approaches utilizing the full SCOPE model and a SCOPE emulator, are summarized below and in Table 1.

2.3.4 TBM-SIF Models

Here we provide a brief description of individual TBM-SIF models and within model experiments. We point out key differences in modeling of photosynthesis, fluorescence, and leaf-to-canopy scaling.

2.3.4.1 BETHY

The Biosphere Energy Transfer Hydrology (BETHY) model is the land surface component of the Carbon Cycle Data Assimilation System (CCDAS) developed to ingest a range of observational data for estimating GPP at global scale (Rayner et al., 2005; Kaminski et al., 2013; Koffi et al., 2012; Anav et al., 2015). Koffi et al. (2015) was the first to combine a process-based model of SIF with a global TBM. The native canopy radiative transfer and photosynthesis schemes of BETHY were effectively replaced with corresponding schemes and fluorescence model from SCOPE (Koffi et al., 2015), thus enabling spatially explicit simulation of GPP and SIF as a function of plant function type. This model was extended to include a module for prognostic leaf growth (Norton et al., 2018) and more recently adapted with a formal optimization algorithm for assimilating spaceborne SIF data (Norton et al., 2019). It has been updated for this study to accept hourly meteorological forcing. BETHY-SCOPE, denoted here as BETHY, remains the first and only global TBM-SIF model to simulate vertically integrated (1-D) fluorescence radiative transfer and energy balance.

We include three experiments to examine the impact of calibrating the $k_N$ model against PAM fluorometry data to different species: (1) BETHY-exp1 is adapted to unstressed cotton species (van der Tol et al., 2014), (2) BETHY-exp2 is adapted to drought stressed Mediterranean species...
(i.e., vineyard in controlled environment subjected to drought) including higher temperature correction (Flexas et al., 2002; van der Tol et al., 2014), (3) BETHY-exp3 is adapted to drought stressed Mediterranean species (Flexas et al., 2002).

We further leverage SCOPE enabled SIF modeling in BETHY (BETHY-exp3 specifically) to examine (a) leaf and canopy level SIF and quenching under sunlit and shaded leaves, and (b) SIF emissions at the top of canopy (SIF$_{\text{canopy}}$) versus the average emission within the canopy (SIF$_{\text{ave}}$), which accounts for the average emission from sunlit and shaded leaves. The latter analysis facilitates comparison to PhotoSpec, which observes the entire canopy.

An important caveat in the analysis of BETHY simulations is that, at the time of this writing, the prescribed met forcing at NR1 is only available for 2015. While this degrades comparison to diurnal and synoptic variation observed by PhotoSpec in 2017, we find that analysis of magnitude, light sensitivities, and within model experiments still provides useful insight for interpretation of other TBM-SIFs, and future modeling requirements in general.

2.3.4.2 ORCHIDEE

The Organizing Carbon and Hydrology In Dynamic Ecosystems (ORCHIDEE) model (Krinner et al., 2005) is the land surface component of the Earth System Model of Institut Pierre-Simon Laplace IPSL-CM, (Dufresne et al., 2013) involved in recent exercises of the Coupled Model Intercomparison Project (CMIP) established by the World Climate Research Programme (https://www.wcrp-climate.org/wgcm-cmip). Recently a mechanistic SIF observation operator was developed for ORCHIDEE to simulate the regulation of photosystem II $\phi_F$ at the leaf level using a novel parameterization of NPQ as a function of temperature, PAR, and normalized $\phi_F$. It emulates the radiative transfer of SIF to the top of the canopy using a parametric simplification of SCOPE. The details of the SIF modelling approach are provided in Bacour et al. (2019).

We include three experiments to examine the impact of water stress and parameter optimization (using OCO-2 SIF, see Section 2.4): (1) ORCHIDEE-exp1 is the standard configuration with default parameters, (2) ORCHIDEE-exp2 is the same as ORCHIDEE-exp1 with two key differences (a) water stress is applied to stomatal conductance, mesophyll conductance and to the photosynthetic
capacity, and (b) the tree height (12 m instead of 15 m) was set specifically for the NR1 site, (3) ORCHIDEE-exp3 is the same as ORCHIDEE-exp1 but includes OCO-2 optimized parameters.

2.3.4.3 BEPS

The Boreal Ecosystem Product Simulator (BEPS) is an enzyme kinetic two-leaf model for simulating carbon and water cycles for different plant functional types (Chen et al., 1999; Liu et al., 2003). BEPS uses a modified Ball-Berry stomatal conductance model (Leuning et al., 1995) and semi-analytical canopy radiative transfer. The canopy architecture is well considered in BEPS model, which has not only remote-sensed LAI but also the global map of the foliage clumping index. The fluorescence emission at the leaf level follows the approach of Lee et al (2015). SIF emission for sunlit and shaded leaves are separately simulated based on illumination and canopy geometry in BEPS. In addition, multiple scattering SIF is also simulated to account for the scattering process within the canopy. The scaling of leaf-level fluorescence emission to the canopy is based on a novel scheme for single-layer models which accounts for canopy scattering and extinction from sunlit and shaded leaves (Qiu et al., 2019). This scaling scheme is an effective approach to simulate the radiative transfer of SIF for a given canopy structure. We include two experiments similar to BETHY-exp1/2 in the calibration of the $k_N$ model against unstressed vs stressed species (BEPS-exp1 and BEPS-exp2, respectively).

2.3.4.4 CLM4.5

The Community Land Model version 4.5 (CLM4.5) provides a description of the biogeochemical profile spanning from the sub-surface bedrock to the top of the vegetation canopy. The fluorescence sub-model follows Raczka et al. (2019), in which the degree of light saturation is calculated from the potential and actual electron transport rate as determined from the photosynthesis model described above. $\phi_f$ is formulated as described in Equation 2 and $\phi_p$ is formulated as a function of the maximum $\phi_p$ under dark acclimated conditions and the degree of light saturation. CLM4.5 uses independent site-level SCOPE simulations that match the observed canopy characteristics and observed GPP at Niwot Ridge to calculate a leaf to canopy level conversion factor ($\kappa_{740}$) for estimating SIF$_{canopy}$. In CLM4.5, $\kappa_{740}$ is fitted to the modeled SCOPE data as a function of solar zenith angle (and implicitly $V_{cmax}$).
Similar to Raczka et al. (2019), here we examine three separate approaches to parameterize $k_N$. CLM4.5-exp1 only considers reversible NPQ ($k_R$), such that $k_N = k_R$, and the relationship between $k_R$ and the degree of light saturation is fitted to PAM fluorometry data based on Mediterranean shrubs (Flexas et al., 2002; Galmes et al., 2007). CLM4.5-exp2 parameterizes $k_R$ with PAM fluorometry from a Scots Pine forest (Porcar-Castell et al., 2011), and defines the rate coefficient in terms of both a reversible and sustained component ($k_N = k_R + k_S$). It has been found that sustained NPQ is important for cold climate evergreen conifer forests such as Niwot Ridge (Miguez et al., 2015; Magney et al., 2019b), and Raczka et al. (2019) found that representing both components provided improved simulations of seasonal SIF. CLM4.5-exp3 is similar to CLM4.5-exp3 but includes a seasonally varying representation of $k_R$. All model experiments use hand-tuned parameters specific to US-NR1 (Raczka et al., 2016).

2.3.4.5 CLM5.0

CLM version 5.0 (CLM5.0) is similar to CLM4.5 with respect to the implementation of the fluorescence sub-model, yet includes several important updates to the representation of photosynthesis from CLM4.5, including a prognostic calculation of Vcmax based upon leaf nitrogen and environmental conditions, revised nitrogen limitation scheme, Medlyn stomatal conductance model, and plant hydraulic water stress (Kennedy et al., 2019). To represent NPQ we use a single approach for $k_N$ (see CLM4.5-exp1), but examine three approaches for estimating $k_{\tau 40}$: (1) CLM5.0-exp1 uses $k_{\tau 40}$ as function of Vcmax following Lee et al (2015), (2) CLM5.0-exp2 follows the approach of CLM4.5, and (3) CLM5.0-exp3 adapts the approach proposed by Zeng et al. (2019) that estimates the fraction of total emitted SIF escaping the canopy by combining near-infrared reflectance of vegetation (NIRV) and fPAR.

2.3.4.6 SIB3

The Simple Biosphere Model version 3 (SIB3) involves the use of explicit biophysical mechanisms to directly calculate carbon assimilation by photosynthesis (Baker et al., 2003; 2008). SIB3 includes prognostic calculation of temperature, moisture, and trace gases in the canopy air space, but requires prescription of most structural properties including LAI. We examine two approaches for prescribing LAI: (1) SIB3-exp1 using values prescribed from MODIS, and (2) SIB3-
exp2 uses values observed at the study site (4.0 m² m⁻²). In general, the fluorescence sub-model follows the approach of Lee et al. (2015) except that $k_N$ is adapted to drought stressed species following van der Tol et al (2014).

2.3.4.7 SIB4

SIB4 (Haynes et al., 2019a,b) shares many similarities to SIB3 with respect to functional aspects of photosynthesis and fluorescence, however, SIB4 uses prognostic rather than prescribed phenology and LAI.

2.4 Data Assimilation

Details of the data assimilation protocols for BETHY and ORCHIDEE are provided in Norton et al. (2018) and Bacour et al. (2019), respectively. For the two models, an ensemble of parameters related to photosynthesis (including optimal Vcmax) and phenology were optimized for several plant functional types. Note that none of the assimilated pixels encompass the location of the US-NR1 tower. Also, in ORCHIDEE, the study site is treated as boreal needleleaf evergreen (ENF); as such, the ORCHIDEE-exp3 simulations in this study are based on parameters optimized against OCO-2 SIF data using an ensemble of worldwide ENF pixels. Note that for BETHY, each experiment uses the same set of optimized parameters whereas in ORCHIDEE the SIF simulations are performed separately for the standard parameters (ORCHIDEE-exp1/exp2) and optimized parameters (ORCHIDEE-exp3), thus providing a test of sensitivity to parameter optimization as discussed below.

2.5 Illumination Conditions

In order to gain insight into how SIF emissions and quantum yields vary with illumination, we further analyze Photospec and a subset of models with respect to (a) changes in incoming light and (b) self-shading within the canopy, respectively. For PhotoSpec, we analyze changes in canopy average SIF and SIF_rel under conditions of predominantly direct versus diffuse PAR, using the 0.5 threshold to distinguish between the two conditions (Section 2.2.2). For models we focus on emissions from sunlit vs shaded leaves. We analyze leaf- versus canopy-level SIF emissions (SIF_leaf and SIF_canopy) in CLM4.5-exp3, and leaf-level quantum yields ($\phi_f$, $\phi_p$, $\phi_N$) in BETHY-exp3. We further compare predictions of quantum yield at the top-of-canopy to canopy averages in...
BETHY-exp3. The motivation here is that top-of-canopy leaves see most of the sunlight, and thus should have different yields compared to shade adapted leaves lower in the canopy. This also provides a more direct comparison for PhotoSpec.

2.6 Modeling Protocol

Models are run for the period 2000-2018 (except BETHY, 2015 only) using identical, hourly, gap-filled meteorological observations. The primary hourly output fields analyzed are the top-of-canopy SIF ($\text{SIF}_{\text{canopy}}@740$ nm), GPP, $\phi_f$, $\phi_p$, and APAR. Model-observation comparisons are made for absolute and relative SIF, GPP, SIF$_{\text{yield}}$ ($\text{SIF}_{\text{canopy}}$/APAR) and GPP$_{\text{yield}}$ (GPP/APAR), sunlit versus shaded canopies ($\text{CLM4.5-exp3}$ and BETHY-exp3), and TOC versus canopy average SIF ($\text{SIF}_{\text{canopy}}$ versus SIF$_{\text{ave}}$, respectively, from BETHY-exp3). Quantum yields and within model experiments provide context to understand canopy integrated results. We focus our analysis on 8 am – 4 pm local time from July-August 2017 for comparison to available PhotoSpec and APAR data.

Models are controlled for meteorological forcing (meteorological data described in Burns et al., 2015) but other factors such as spin-up, land surface characteristics, parameter tuning, and model state, are not controlled for and are treated separately according to each model’s protocol. For example, CLM4.5 is better suited than others in prescribing observed vegetation characteristics at the study site. Three BETHY experiment and one ORCHIDEE experiment (ORCHIDEE-exp3) were preliminary optimized by assimilating independent Orbiting Carbon Observatory 2 (OCO-2) SIF data at the global scale (Section 2.4). We emphasize that our point here is not to identify the best model but to identify common patterns in model behavior through normalized SIF and deviation from observed behavior to identify areas requiring the most attention.

The results are organized around two parallel themes. The first theme addresses four key processes driving canopy-level fluorescence: (1) incoming illumination, (2) energy partitioning on incoming light between photochemistry, fluorescence, and NPQ, and (3) leaf-to-canopy emitted SIF, including linearity of yields at leaf and canopy scale. The second theme addresses sensitivity of these processes to environmental conditions at diurnal and synoptic scales. Here, synoptic
scale refers to the impact of day-to-day changes in weather, including two storm events which brought sustained cool, wet, and cloudy conditions from July 22-31 and then from August 6-10.

Section 3: Results

Incoming Illumination

Two key features dominate observed APAR variability: afternoon depression (Fig 2A) and reduction during two summer storms (Fig 2D). Both features are captured by models. More generally, models capture synoptic variability with high correlation (r > 0.8) and low across model spread (σ = 10%). The exception is BETHY, which is simulated outside our observation year (2015).

High model fidelity is expected given that observed PAR is prescribed, and it is promising that models show a consistent response to changes in illumination. The primary shortcoming is a systematic high bias in APAR magnitude (129%), with most models exceeding the upper range of observed APAR (as determined from the six within canopy PAR sensors, Fig S2), and high model spread. These errors are likely related to differences in predicted fAPAR. In the case of ORCHIDEE, high APAR is expected due to the big leaf assumption where all eaves are considered as opaque and fully absorbing.

Canopy Photosynthesis

Observed GPP shows a broad peak from mid-morning to early afternoon (~9 am – 1 pm local), followed by slight decrease until 4 pm (Fig 2B), consistent with afternoon cooling and reduced light availability (Fig 1B-D). The two month period under investigation is relatively flat with generally weak day-to-day variability (σ = 17%), but modest correlation with APAR (r = 0.61, Fig 2E). Some models capture the afternoon GPP depression, but all models strongly underestimate its magnitude, apparently independent of stomatal conductance formulation or more explicit accounting for plant hydraulic water stress such as in CLM5.0. Models are mostly uncorrelated with observed GPP at synoptic scale (r ranges from -0.2 to 0.36, highest value in SiB4), high biases, and show increased spread (in predicted magnitude) relative to APAR (143% +/- 23%).

While observed GPP$_{yield}$ is mostly stable over the diurnal cycle, most models (except BEPS) show a distinct midday minimum (Fig 3A). Half of the models show a similar midday minimum in photochemical quantum yield ($\phi_p$, Fig 4A), with the other half either increasing or decreasing in
the afternoon (CLM5.0 and SiB3/SiB4, respectively). The midday dip in yield is likely associated
with reduced photosynthetic efficiency at high light levels, as demonstrated by reductions in GPP,
GPP\(_{\text{yield}}\), \(\phi_p\) with APAR (Fig 5A, C, E).

Observed GPP\(_{\text{yield}}\) shows significant structure at synoptic temporal scale (Fig 3C), most notably
increased yield during the cool/rainy period (reduced heat and water stress), and decreased yield
in mid- to late- August (increased heat and water stress following the cooling pattern). In contrast
to predicted GPP, models show high fidelity in capturing the magnitude and variability of GPP\(_{\text{yield}}\)
at synoptic scale (\(r\) ranges from 0.35 – 0.69, highest values in CLM4.5 and CLM5.0). Individual
models are self-consistent in their predictions of GPP\(_{\text{yield}}\) and \(\phi_p\) at synoptic scale (\(r = 0.592 –
0.935\)) except for SiB3/SiB4 (\(r < 0.1\), Fig 4B).

**Canopy Fluorescence**

Observed SIF\(_{\text{canopy}}\) is strongly correlated with observed APAR at diurnal and synoptic scale (\(r =
0.77\)), with common features including afternoon depression and reduction during rainy periods
(Fig 2C & 2F). Observed PAR also feeds into the fluorescence sub-model and, unlike GPP, strongly
correlates with SIF\(_{\text{canopy}}\) at synoptic scale (\(r\) ranges from 0.58 to 0.92, highest values in ORCHIDEE).
However, we find a persistent positive model bias in SIF\(_{\text{canopy}}\) (170% +/- 45%) consistent with, but
not proportional in magnitude to, the APAR bias. We note that models are especially
oversensitive to APAR at high light levels (Fig 5D).

We investigate the high bias in SIF\(_{\text{canopy}}\) in more detail using BETHY-exp3 and CLM4.5-exp3.
Specifically, we examine leaf and canopy level SIF and quenching under sunlit and shaded leaves.
Analysis of quantum yields in BETHY-exp3 shows a reversal in the fractional amounts of absorbed
energy going to SIF and PQ vs NPQ in low- vs high-light conditions that is consistent with leaf level
data and theory (Porcar-Castell et al., 2014). More specifically, BETHY-exp3 predicts low \(\phi_F\) and
\(\phi_p\) and high \(\phi_N\) in sunlit leaves relative to shaded leaves, with more energy going to fluorescence
and photochemistry than to NPQ in shaded leaves, and more energy going to (shed off by) NPQ
in sunlit leaves (Fig S5). Likewise, total \(\phi_F\) shows decreasing values with increasing APAR in
BETHY-exp2/3 compared to BETHY-exp1, consistent with observed SIF\(_{\text{yield}}\) (Fig 5E-F), as \(\phi_N\) ramps
up to higher levels in the drought parameterized Kn model. Moreover, in stark contrast to SIF\(_{\text{yield}}\)
and $SIF_{\text{canopy}}$, $\phi_F$ does not show high values relative to other models (Fig 4D). These results point to an issue in BETHY with leaf to canopy scaling in needleleaf forests.

Analysis of CLM4.5-exp3 suggests several possible reasons for oversensitivity to APAR. First, we focus on emissions from sunlit/shaded portions of the canopy (Fig S6). CLM4.5-exp3 and PhotoSpec both show higher SIF under “high light” conditions (sunlit leaves and direct radiation, respectively) compared to “low light” conditions (shaded leaves and diffuse radiation, respectively), which is promising (Fig S6 A,D). Comparing the ratio of sunlit to shaded SIF in CLM4.5-exp3 to the ratio of direct to diffuse SIF in PhotoSpec (Fig S6 B,E) shows higher ratio in CLM4.5-exp3 on average. The difference peaks in midday, when sunlit leaf area is maximized (self-shading minimized) in CLM4.5 but no major difference in the amount of direct radiation, and decreases with increasing sun angle (morning and afternoon) and with increasing rainfall (in the afternoon on average, and during the rainy period in late July/early August), both of which increase the shaded fraction. As such, accounting for view angle and different illumination metrics for PhotoSpec and CLM4.5 (most comparable in morning, afternoon, and during rainy days) reduces, but does not entirely remove, the positive bias in high light conditions.

Second, the degree of light saturation ($x$, fraction of absorbed light not used in photosynthesis) is twice as high in the sunlit canopy in CLM4.5 (Fig S7), which leads to high fluorescence efficiency in sunlit leaves and low fluorescence efficiency in shaded leaves. While this produces high photochemistry in shaded leaves, it contributes a small fraction of SIF to the total canopy (~20%) despite higher fractions of shaded leaves (~2/3 at noon, Fig S6C) and thus sunlit leaves dominate $SIF_{\text{yield}}$ and $SIF_{\text{canopy}}$. Therefore, it seems likely that a model’s representation of canopy structure including the partitioning between sunlit/shaded canopy fraction has an important impact upon canopy SIF. Biases in the sunlit/shaded will likely propagate into the simulated value of canopy SIF. However, it’s important to know that the observed sunlit/shaded fraction from PhotoSpec is estimated as well, since it is currently no possible to determine the sun/shade fraction within PhotoSpec FOV.

Additionally, all formulations of CLM4.5 (and most models except BETHY) show lack of decline in $SIF_{\text{yield}}$ with APAR compared to measurements of absolute SIF (Fig 5E). For CLM4.5, the relationship between $SIF_{\text{yield}}$ and APAR depends upon the relationship between degree of light
saturation and reversible NPQ (Raczka et al., 2019). This suggests it is important to properly represent the NPQ response to environmental conditions when simulating SIF.

While most of the model bias is reduced in SIF\textsubscript{yield} (126%, mostly attributed to BETHY), the remaining signal, representing the dynamic response to synoptic conditions (e.g., Magney et al., 2019), is poorly represented in models (Fig 3D). Most models show zero to strongly negative correlation with observations at synoptic scale and only two models, ORCHIDEE-exp3 and CLM4.5-exp2, produce positive correlation (0.58 and 0.33, respectively). These are the only two models (prescribed with 2017 met forcing) that also capture a negative relationship between SIF\textsubscript{yield} and APAR (Fig 5E).

In general, predicted SIF\textsubscript{yield} is stable during our short study period and is uncorrelated with GPP\textsubscript{yield} (Fig 3; Fig S8). While this finding runs counter to observations of SIF\textsubscript{yield}, which shows a clear response during and following the storm event and varies linearly with observed GPP\textsubscript{yield} (r = 0.40), it is consistent with observations of SIF\textsubscript{rel} (grey line in Fig 3 and Fig S8A) which like most model predictions is stable and uncorrelated with GPP\textsubscript{yield}. We find it important to clarify the difference between SIF\textsubscript{yield} and SIF\textsubscript{rel} here, since these metrics represent different but equally important versions of reality. SIF\textsubscript{yield}, estimated as the ratio between absolute canopy SIF (SIF\textsubscript{canopy}) and APAR, is our best attempt to account for the effect of canopy absorbed light on the canopy integrated emission of SIF. However, factors such as observation angle, sunlit bias, and difference in footprint from APAR, necessitates our alternative calculation in SIF\textsubscript{rel}. While SIF\textsubscript{rel} removes model-observations differences in illumination, it confounds our interpretation of the relationship with GPP\textsubscript{yield}, which is derived from APAR. As such, we provide both results to be comprehensive, but note the temporal stability associated with SIF\textsubscript{rel} as the more physical interpretation of canopy yield for this short period of study.

Leaf-to-Canopy Scaling

Several methods have been proposed to transfer predicted leaf-level SIF emissions to the top of canopy. While leaf-to-canopy scaling enables efficient global scale simulation, the diversity of novel methods adds uncertainty to the canopy level estimate of SIF (in addition to aforementioned uncertainties in structure, APAR, photochemistry, fluorescence). These
differences are evident in comparison of Figures 3 and 4, in which yields are plotted on a similar scale.

At least at diurnal scale, there is some evidence that leaf and canopy emissions look more similar for models adopting simplified empirical scaling functions (SiB3, SiB4, CLM4.5, CLM5.0, BEPS) than for models that more explicitly account for radiative transfer (BETHY, ORCHIDEE). For the more explicit models, the diurnal cycle of $\phi_f$ is out of phase with SIF$_{yield}$, the former of which peaks in the afternoon and the latter of which peaks in the morning. This produces reasonable agreement to PhotoSpec in phase and magnitude between SIF$_{yield}$ and SIF$_{rel}$ for ORCHIDEE, but produces divergence in the magnitude of SIF$_{yield}$ for ORCHIDEE.

Model performance in leaf-to-canopy scaling is summarized in Figure S8. The only two models with a positive relationship between yields (Fig S8B) and between quenching terms (Fig S8C) are the two models with more explicit representation of radiative transfer (i.e., ORCHIDEE and BETHY). CLM4.5 is the only model with a positive relationship between yields, but not between quenching terms. SiB3/SiB4 are the only models with a positive relationship between quenching terms, but not between yields.

Finally, we note that PhotoSpec scans of leaf-level emissions are averaged and reported here as canopy averages, while model output is reported at the top of the canopy, which accounts for within-canopy radiative transfer, re-absorption of SIF, and shaded canopies, causing lower emissions compared to the canopy average. CLM4.5, for example, shows strong attenuation of SIF from leaf-level to TOC, decreasing by a factor of 2-3 at midday (Fig S7). The interpretation here is that the model bias in absolute SIF may actually be higher than reported here; however, we note that more quantitative information on the observed fraction of sunlit vs shaded leaves and comparative top-of-canopy SIF values for the same canopy elements are needed (to account for off-nadir SIF viewing) for more accurate determination of scaling between observed canopy and top-of-canopy SIF.

Within Model Experiments

In most cases, within model experiments produce improvements in some metrics and degradation across others (performance change is quantified by reporting correlation values in...
brackets). An important and unexpected result of this study is the impact of different levels of tuning to observations on our predictions. While this work represents a snapshot of the state-of-the-art in site-level TBM-SIF modeling, and we have taken great care to control for environmental conditions (most important being illumination), an important overall takeaway is for future model comparisons to make additional efforts to control for initial conditions and vegetation state.

The most basic example is tuning of LAI in SiB3. LAI, as prescribed by MODIS for SiB3-exp1 (~1.5), is on the low end for a subalpine evergreen forest, and consequently produces negative biases in APAR, GPP, SIF and SIF$_{yield}$. When prescribed according to tower observations in SiB3-exp2 (~4.0), the biases become positive (albeit on the lower end of the model ensemble), but produces degraded variation at synoptic scale for GPP (0.39 vs 0.19), SIF (0.87 vs .71) and SIF$_{yield}$ (0.09 vs -0.32).

Experiments in CLM4.5 comprise a higher level of hand tuning of vegetation structural and functional characteristics. Parameter tuning was imposed to match vegetation structure with site level measurements and consequently CLM4.5 produces overall low bias in yields. With respect to synoptic variation, NPQ experiments, tuned against the measured air temperature and a representative evergreen forest, produce improvements at synoptic scale for GPP (-0.01 vs 0.16), SIF (0.59 vs 0.86), and GPP$_{yield}$ (0.05 vs 0.63), but degradation in SIF$_{yield}$ (0.32 vs -0.25). Likewise, NPQ experiments in BETHY based on species information (calibration of $K_N$ against PAM fluorescence in stressed vs unstressed systems) shows improvement in the SIF$_{yield}$-APAR relationship for drought stressed models (BETHY-exp1 vs BETHY-exp2/3).

Experiments with ORCHIDEE demonstrate that errors in model parameters (such as $V_{cmax}$, LAImax, leaf age, or SLA) contribute to SIF and GPP uncertainty but can be alleviated by assimilation of OCO-2 SIF retrievals (ORCH-exp1/2 vs ORCH-exp3). Model optimization of parameters improves the functional link between SIF and GPP, thus reducing biases in APAR, GPP, and SIF$_{yield}$, and improving synoptic variation in SIF$_{yield}$ (-0.04 vs 0.58).

Section 4. Discussion
This study represents a first attempt to evaluate a controlled ensemble of TBM-SIF models against canopy integrated SIF observations to identify and attribute model-observation mismatches related to errors in canopy absorption of sunlight, photosynthesis, fluorescence, and leaf-to-canopy radiative transfer of fluorescence.

Different models match some observed parameters better than others (with respect to APAR and yield), but no model gets both APAR and SIF$_{\text{yield}}$ magnitude and/or sensitivities close to the observations. For example, BEPS closely matches the magnitude of APAR (Fig 2A), and BETHY captures the decline in SIF$_{\text{yield}}$ with APAR for NPQ quenching based on stressed species (Fig 5E), but both models overestimate observed yield by a factor of 2, hence SIF is overestimated (Fig 2). CLM4.5 correctly captures the diurnal SIF$_{\text{yield}}$ change, but overestimate APAR; in this case, SIF and SIF$_{\text{yield}}$ are overestimated. Importantly, models diverge strongly from each other and from observations in the magnitude of SIF$_{\text{yield}}$ and its decline with APAR (Fig 5E), partially reflecting model variability in $\phi_f$ (Fig 5F), but in general show a characteristic pattern of weak SIF$_{\text{yield}}$ decline with APAR. GPP$_{\text{yield}}$ shows higher agreement between models and with observations (Fig 5B), despite divergent $\phi_p$ (Fig 5C), which could be indication that the primary uncertainty is due to the representation of fluorescence and not the photosynthesis model.

Consequently, we find a strong linear and positive relationship between observed SIF$_{\text{yield}}$ and GPP$_{\text{yield}}$ for absolute SIF, which is underestimated on average by models (Fig 6A-B). In contrast, models show quite strong positive relationships between $\phi_f$ and $\phi_p$ (Fig 6C). Our study highlights an apparent challenge for models in transferring leaf level processes to canopy scale, and consequently, linking the proper canopy mechanistic SIF-GPP relationship at the leaf level.

The mismatch between multi-model simulations and tower-based observations of SIF and GPP at hourly and daily scales can be summarized as symptoms of five main factors: (1) PhotoSpec scan strategy, (2) radiative transfer of incoming PAR and impact on APAR and sunlit/shaded fraction, (3) representation of photosynthesis and sensitivity to water limitation especially during afternoon conditions, (4) representation of fluorescence and sensitivity to reversible NPQ response at Niwot Ridge, and (5) radiative transfer of fluorescence from leaf to canopy. Several persistent biases falling under these broad categories are discussed below.
Apples to Apples Comparison.

PhotoSpec is unique in its ability to scan entire canopies for signals that are largely hidden from nadir-oriented instruments. However, this creates unique challenges for interpretation of data and comparison to models. For example, the diurnal cycle of observed SIF is highly sensitive to view angle. PhotoSpec was set up in 2017 to scan back-and-forth between northwest and northeast view angles, but the instrument was slightly biased to the northwest, causing a low phase angle in the morning (more aligned with rising sun) and increased phase angle in the afternoon (more opposed to setting sun). As such, PhotoSpec observed predominantly illuminated canopies in the morning and shaded canopies in the afternoon (i.e., more shaded fraction), leading to the late morning peak in reflected radiance (Fig S3).

Moreover, Photospec scans specific locations at the top of the canopy from near nadir to view angles closer to the horizon (see Fig. S8 in Magney et al., 2019b), while models are currently configured to simulate top of canopy emission and simulated here as nadir viewing. The question becomes whether to retain nadir only data and sacrifice signal-to-noise, or to average over all elevation angles and risk aliasing view angle effects. This study, partly motivated by high agreement of canopy integrated SIF with spaceborne data from OCO-2 and TROPOMI (Magney et al., 2019b; Parazoo et al., 2019), has chosen the latter approach but with an attempt to minimize scan angle effects in SIF_{rel}. However, it is worth noting that swath sensors such as GOME-2 show high sensitivity to viewing angle especially under increasing illumination angles (Kohler et al., 2018; Joiner et al., in review). View angle effects are likely to be especially acute for PhotoSpec in the morning and afternoon with increasing anisotropy and changes in the illuminated field of view with sun and view angle. Other tower SIF instruments with a wide FOV (i.e. FluoSpec2; Yang et al., 2018) may more appropriately represent the TOC SIF emission, but also have difficulty disentangling the sunlit/shaded canopy components.

It is critical that model evaluation relative to measured SIF data and data assimilation studies properly account for the specificities of the instrument (viewing of the instrument, spectral band, time of the overpass for space-borne instruments), the representation of canopy emission, and correct observations for directional variations in SIF relative to observation geometry. Although normalizing SIF by reflected radiance partially alleviates scan angle effects, this highlights the...
need for models to get canopy structure, radiative transfer, and sunlit/shaded fraction correct, which feed all the way through to SIF and GPP. Further ground-based investigations of SIF anisotropy, sunlit/shade fraction, and vertical distribution (within canopy, canopy integrated, and top of canopy) with PhotoSpec may help to inform models on the physical aspects of the signal. Despite the issues we highlight in comparing observations to models, the potentially more interesting and important story here is with respect to model-model comparisons, which reveals wide divergence in response to light conditions and other factors, as discussed below.

**TBM SIF is too sensitive to APAR.**

Our results indicate a spectrum of SIF responses to APAR: TBMs are usually far too sensitive to APAR, observations of absolute SIF are less sensitive, and observations of relative SIF are least sensitive. The fact that relative SIF is the least sensitive is telling, as it reduces sensitivity to APAR and reveals a strong SIF response to changes in photochemical quenching. SIF models appear especially sensitive to sunlit leaves. In CLM4.5, SIF emissions from the sunlit portion of the canopy are a factor of 5 higher than emissions from shaded leaves, despite twice as fewer leaves in the sunlit canopy (Fig S6C). In CLM4.5, the combination of higher than average $\phi_0$ (Fig 5F) with higher fluorescence efficiency in the sunlit portion of the canopy, produce an increase in the magnitude and sensitivity to sunlit fraction, thus contributing to the high bias (factor of 3 higher than observed) and strong diurnal cycle (2-fold increase from morning to midday).

**Linearity of SIF and GPP yields.**

Observations show a positive but not significant linear relationship between $\text{SIF}_{\text{yield}}$ and $\text{GPP}_{\text{yield}}$ (Fig 6A, $r = 0.40$) at our study site. This is likely due to the short time period investigated here where there is relatively little change in $\text{SIF}_{\text{yield}}$ and $\text{GPP}_{\text{yield}}$ during peak summer. Only 3 of 7 models show a significant ($r > 0.35$) linear and positive slope ($r > 0.35$; ORCH-exp3, CLM4.5-exp3, and BETHY-exp3) between $\text{SIF}_{\text{yield}}$ and $\text{GPP}_{\text{yield}}$, while 5 models (except CLM5.0) show a significant positive slope between quantum yields ($\phi_f$ and $\phi_p$, Fig 6C). These regression plots of quantum yields, in turn, help explain the observed linearity of $\text{SIF}_{\text{yield}}$ vs. $\text{GPP}_{\text{yield}}$. At least in the case of Niwot Ridge, model (and presumably observed) $\phi_p$ stays within high light “NPQ-Phase” conditions, and generally doesn’t exceed the range in which decoupling of $\phi_f$ and $\phi_p$ ($\phi_p > 0.6$)
in low light “PQ-Phase’ conditions occurs (Porcar-Castell et al., 2014, cf Fig 9). BETHY-exp3, which
best captures the observed relationship in the canopy between SIF\text{yield} and GPP\text{yield}, is also the
only model that also shows a decline in SIF\text{yield} with APAR, as discussed below. These results are
likely to change when we expand the study to several years; however, the purpose of this study
was to provide an initial investigation into the response of modelled SIF and GPP to light during
peak summer.

### Insufficient decline in SIF\text{yield} with APAR.

In general, models show an insufficient decline in SIF\text{yield} with APAR, when compared to observed
SIF\text{yield} (Fig 5E). All models except SiB3 and SiB4 show some decline, with BETHY showing the best
agreement in slope magnitude. BETHY is the only model with full radiative transfer but this does
not appear to have a substantial impact on SIF\text{yield}, which has a similar (albeit suppressed) decline
with APAR as $\phi_f$ (Fig 5F). Within model experiments show little to no sensitivity of SIF\text{yield} or $\phi_f$
decline with APAR to water stress (e.g., ORCHIDEE) or prescribed LAI (e.g., SiB3), but high
sensitivity to the formulation of NPQ with respect to species calibration (e.g., BETHY) and
reversibility (e.g., CLM4.5).

Three CLM4.5 experiments demonstrate sensitivity to representation of NPQ variability at diurnal
and seasonal scales. The first simulation using the default NPQ parameterization from SCOPE
(CLM4.5-exp1, based on a 2-parameter fit to drought stressed Mediterranean species (Galmes et
al., 2007) produces the strongest decline in SIF\text{yield}. The second simulation, which includes a site-
specific NPQ formulation that accounts for $k_R$ and $k_S$ (CLM4.5-exp2), produces the weakest
decline. The third simulation with seasonally varying $k_R$ produces a slightly stronger decline. An
important point for this formulation is that $k_R$ is constrained by PAM fluorometry data at Hyytiala
(Scot Pine) and does not account for high light saturation values and summer drought conditions
that may be more typical of lower latitude sites such as Niwot Ridge. This could indicate that
parameterizing $k_R$ based upon similar PFTs may not be sufficient to properly characterize the NPQ
response for lower latitude sites such as Niwot Ridge.

Similar results are found in experiments with BETHY comparing stressed (drought) and
unstressed (relative to water availability) NPQ models at NR1 but controlling for $k_R$ (constant in
time in both cases, stronger negative $SIF_{yield}$ response to APAR in stressed model). In the
unstressed models of CLM4.5 and BETHY, the NPQ response to APAR becomes too low, causing
an oversensitivity of SIF to APAR and thus high SIF bias. The strongly regulated NPQ response of
the drought-based model enables more non-photochemical quenching at high light levels in
stressed ecosystems compared to typical unstressed plants. While this $k_{NPQ}$ model was
developed using drought-stressed plants, similar up-regulation of NPQ is expected to occur under
any condition where photosynthesis is limited and available excitation energy is high (e.g. cold
temperatures and high light, Sveshnikov et al., 2006, doi: 10.1093/treephys/26.3.325). Our
results thus emphasize the need for careful implementation of NPQ dynamics for simulating and
assimilating SIF in different light and stress environments (Raczk et al., 2019; Norton et al.,
2019).

Data assimilation reduces high bias. Assimilation of OCO-2 SIF in ORCHIDEE brings the magnitude
of both GPP and SIF in closer agreement with observations. This improvement is driven by
decreases in leaf photosynthetic capacity ($V_{cmax}$, $LAI_{max}$, leaf age, SLA, Bacour et al., 2019),
which decreases the magnitude (but not shape) of APAR closer to observed values (Fig 2), and
leads to improvements in $GPP_{yield}$ and $SIF_{yield}$ (Fig 3). Nevertheless, after the assimilation there
are still disagreements in $SIF_{yield}$ vs $GPP_{yield}$ relative to the measured quantities (Fig 6). For diurnal
and synoptic cycles, the assimilation effectively acts to scale the magnitude of SIF, GPP and APAR
(and related yields), but it does little to alter variability. Although data assimilation (i.e. calibrating
model parameters) is critical to improving modelled SIF and GPP, this should be done in
conjunction with improvements in the model formulation (as summarized in Section 5),
otherwise the estimated model parameters can be sub-optimal to compensate for the lack of
missing processes.

5. Conclusions/Recommendations

Our results reveal systematic biases across TBM-SIF models affecting leaf-to-canopy simulations
of APAR, GPP, and SIF. This highlights key areas where observing strategies and model
formulations can be improved:
1) Radiative transfer of incoming and absorbed PAR. The representation of incoming radiative transfer produces positive biases in APAR that leads to positive biases in GPP, both of which occur regardless of time of day. This is influenced by characterization of the canopy, leaf orientation and clumping, biochemical content, canopy layers, and leaf area, which dictates the sunlit/shaded fractions of the canopy. Furthermore, the combination of high APAR bias in models and high uncertainty in observed APAR highlights a need for more accurate and representative in situ measurements of APAR within the FOV of SIF observations and footprint of eddy covariance data.

2) Water stress impacts on photosynthesis. The underlying photosynthetic models fail to simulate the magnitude of depression of observed GPP in the afternoon, regardless of stomatal-conductance or water stress formulation. This likely results from the inability to account for afternoon water stress to properly restrict stomatal conductance and hence GPP and SIF. Additional effort is needed to characterize SIF and GPP sensitivity to increased atmospheric demand and/or reduced soil moisture.

3) Leaf Mechanism for Energy Partitioning. We provide evidence that many models fail to capture the correct reversible NPQ response to light saturation, leading to biases in SIF$^{\text{yield}}$ during high light conditions and especially with increasing moisture limitation at the end of summer. Further investigation using models such as BETHY and CLM is needed to better characterize sensitivity of NPQ formulations to PFT and environmental conditions.

4) Radiative transfer of SIF. SIF is emitted from the leaf level (sunlit shaded fractions of leaf level) and then is transferred to the top of canopy as a function of canopy structure (leaf geometry, canopy layers, leaf area). Despite high disagreement of BETHY-SCOPE with respect to the simulation of APAR and SIF magnitude, we recommend site level simulations using a similar framework where a radiative transfer model is coupled to a terrestrial biosphere model for more detailed investigation of sensitivity to canopy characteristics.

5) Observation strategy. The PhotoSpec scan strategy enables direct measurement of SIF emission at leaf-to-canopy scale, but requires off-nadir view angles that lead to changing fractions of sunlit and shaded canopies throughout the day as a function of sun angle. Further
work could be done using tower mounted instruments with a wider FOV that more accurately represent top of canopy emissions for comparison to model simulations, and to classify emissions from shaded vs sunlit canopies.

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Figure 1. Observed diurnal (A-D) and synoptic (E-H) precipitation (PPT), air temperature at 21 m (Tair), vapor pressure deficit (VPD), and downwelling shortwave (SWdown). Diurnal cycles are averaged over July-August, 2017. Synoptic cycles are plotted as 5-day averages from June 15 – Sep 15. Data from 2017 is shown in black and climatology (2015-2018) in grey. Typically, peak rainfall occurs in the afternoon at this site (A). A substantial rain event which occurred from DOY 203-205 is removed from the 2017 average to show the impact on diurnal variability and to demonstrate the dominance of the afternoon monsoon upon diurnal precipitation in summer.
Figure 2. Observed and simulated diurnal cycles of APAR, GPP and SIF. Diurnal cycles (A-C) are averaged over July-August, 2017. Synoptic cycles (E-F) are plotted as 5-day averages from June 15 – Sep 15. Observations are shown in black, with relative SIF (SIF$_{\text{canopy}}$ / far red reflected radiance) included in (C, F) in grey. The across model average (dashed black) represents the average of “best-case” model scenarios (solid lines; SiB3-exp2, SiB4, ORCHIDEE-exp3, CLM5.0-exp3, CLM4.5-exp3, BEPS-exp2, BETHY-exp3) with uncertainty bars indicating the across model 1 sigma uncertainty.
Figure 3. Same as Figure 2 except for SIF\textsubscript{yield} and GPP\textsubscript{yield}. Here, SIF\textsubscript{yield} = SIF\textsubscript{canopy} / APAR, and GPP\textsubscript{yield} = GPP / APAR.
**Figure 4.** Same as Figure 2, except for quantum yield of fluorescence ($\phi_F$) and photochemistry ($\phi_p$).
Figure 5. Observed and predicted change in GPP, SIF, and yields with APAR. Regression lines are shown for (A) GPP, (B) GPP<sub>yield</sub>, (C) photochemical quantum yield (ϕ<sub>p</sub>), (D) SIF<sub>canopy</sub>, (E) SIF<sub>yield</sub>, (F) fluorescence quantum yield (ϕ<sub>f</sub>), as a function of APAR, using daily mean (8 am – 4 pm local) values over the period July-August 2017. Observations are shown in solid black, individual models and experiments in color, the across model average in dashed black. Relative SIF is shown in grey in (D) and (E).
### Table 1. Summary of TBM-SIF models and within model experiments illustrating model components that may have led to differences in modeled SIF. These include a representation of stomatal-conductance (column 3), canopy absorption of incoming radiation (column 4), limiting factors for photosynthesis (Stress, Vcmax, LAI; columns 5-7) and SIF (kω; column 8), leaf-to-canopy scaling of SIF (column 9), and parameter optimization (column 10). The underlined model experiment was used for model intercomparison.

| Model | Model Experiments | Stomatal Conductance | Canopy Type / Radiation | SIF | Parameters of Fv/Fm | Leaf to Canopy Scaling | Parameter Optimization |
|-------|-------------------|----------------------|-------------------------|-----|------------------|----------------------|-----------------------|
| TBM   | adapted to offline SCOPE runs from Lee et al. (2015) | adapted to drought stressed Mediterranean species (Flexas et al., 2007) | Prognostic | Default | Default | Default | Default |
| BEPS  | Adapted to drought stressed Mediterranean species (Flexas et al., 2007) | Adapted to drought stressed Mediterranean species (Flexas et al., 2007) |  | Default | Default | Default | Default |
| ORCHIDEE | Adapted to drought stressed Mediterranean species (Flexas et al., 2007) | Adapted to drought stressed Mediterranean species (Flexas et al., 2007) |  | Default | Default | Default | Default |
| SiB3  | Adapted to drought stressed Mediterranean species (Flexas et al., 2007) | Adapted to drought stressed Mediterranean species (Flexas et al., 2007) |  | Default | Default | Default | Default |
| SiB4  | Adapted to drought stressed Mediterranean species (Flexas et al., 2007) | Adapted to drought stressed Mediterranean species (Flexas et al., 2007) |  | Default | Default | Default | Default |
| CLM5.0 | Adapted to drought stressed Mediterranean species (Flexas et al., 2007) | Adapted to drought stressed Mediterranean species (Flexas et al., 2007) |  | Default | Default | Default | Default |
| CLM4.5 | Adapted to drought stressed Mediterranean species (Flexas et al., 2007) | Adapted to drought stressed Mediterranean species (Flexas et al., 2007) |  | Default | Default | Default | Default |
| BEPS  | Adapted to drought stressed Mediterranean species (Flexas et al., 2007) | Adapted to drought stressed Mediterranean species (Flexas et al., 2007) |  | Default | Default | Default | Default |
| ORCHIDEE | Adapted to drought stressed Mediterranean species (Flexas et al., 2007) | Adapted to drought stressed Mediterranean species (Flexas et al., 2007) |  | Default | Default | Default | Default |