



Constraining 2010-2020 Amazonian carbon flux estimates with satellite solar-induced fluorescence (SIF)

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- 10 Abstract. Amazonia's Net Biome Exchange (NBE), the sum of biogenic and wildfire carbon fluxes, is a fundamental indicator of the state of its ecosystems. It also quantifies the magnitude and patterns of short- and long-term carbon dioxide sources and sinks but is poorly quantified and out of equilibrium (non-zero) due to both direct (deforestation) and indirect (climate-related) anthropogenic disturbance. Determining trends in Amazonia's carbon balance, shifts in carbon exchange pathways of NBE, and timescales of ecosystem sensitivity to disturbance requires reliable biogenic flux models that adequately
- 15 capture fluxes from diurnal to seasonal and annual timescales. Our study assimilates readily available observations and a derived solar-induced fluorescence (SIF) product to estimate hourly biogenic carbon dioxide (CO₂) fluxes (here in units of μmol CO₂ m⁻² s⁻¹) as Net Ecosystem Exchange (NEE), and its photosynthesis and respiration constituents, at 12 km resolution using four versions of the data-driven diagnostic Vegetation Photosynthesis and Respiration Model (VPRM). The VPRM versions are all calibrated with ground-based eddy flux data and vary based on whether (1) the photosynthesis term incorporates
- 20 SIF (VPRM_SIF) or traditional surface reflectance (VPRM_TRA) and (2) the respiration term is modified beyond a simple linear air temperature dependence (VPRM_SIFg; VPRM_TRG). We compare the VPRM versions with each other and with hourly fluxes from the bottom-up mechanistic Simple Biosphere 4 (SiB4 v4.2) model. We also use NASA's OCO-2 CO₂ column observations to optimize the VPRM and SiB4 models during the 2016 wet season which occurred at the tail of the 2015/2016 severe El Niño. The wet season 2016 case study suggests that relative to SiB4 and the SIF-based VPRMs, the
- 25 traditional VPRM versions can underestimate uptake by a factor of three. In addition, the VPRM_SIFg version better captures biogenic CO₂ fluxes at hourly to seasonal scales than all other VPRM versions in both anomalously wet and anomalously dry conditions. We also find that the VPRM_SIFg model and the independent bottom-up mechanistic hourly SiB4 model converge in NEE, although there are differences in the partitioning of the photosynthesis and respiration components. We further note that VPRM_SIFg describes greater spatial heterogeneity in carbon exchange throughout the Amazon. Despite the paucity of
- 30 OCO-2 CO₂ column observations (XCO₂) over the Amazon in the wet season, incorporating XCO₂ into the models significantly reduces near-field model-measurement mismatch at aircraft vertical profiling locations. Finally, a qualitative analysis of the unoptimized biogenic models from 2010-2020 agrees with the wet season 2016 case study, where the traditional





VPRM formulations significantly underestimate photosynthesis and respiration relative to VPRM-SIFg. Overall, the VPRM_SIFg biogenic flux model shows promise in its ability to capture Amazonian carbon fluxes across multiple timescale
and moisture regimes, suggesting its suitability for larger studies evaluating interannual and seasonal carbon trends in fire as well as the biogenic components of the region's NBE.

1 Introduction

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The terrestrial tropics dominate the interannual variability of the global carbon cycle (Piao et al., 2020). As the dominant component of the South American terrestrial tropics, the Amazon is a major contributor to both the long-term global terrestrial carbon balance and the inter-annual variations in the terrestrial biosphere's sequestration of anthropogenic CO₂ emissions

- (Phillips & Brienen, 2017; Gatti et al., 2014; Saleska et al., 2003). Covering about one-third of South America with about 80% as tropical rain forest, the tropical Amazon is being altered profoundly by human activity especially in recent decades (Andreae, 2019; Fu et al., 2013). Gatti et al. (2021) found that, from 2010–2018, most of the Amazon's carbon emissions come from the disturbed eastern Amazon, and most of those emissions are from fires. In the past two decades, 44% of carbon loss from the
- 45 Brazilian Amazon has been attributed to degradation and disturbance, and 56% to deforestation (Kruid et al., 2021). Recent work (Aragão et al. 2018) suggests that drought-related fires are increasing in importance relative to deforestation-related fires, but this statement was made under the assumption that protections of the Amazon against deforestation would remain in place. Following over a decade of decline, swaths of the Amazon have been illegally burned beginning with the 2019 dry-season; there was a three-fold increase in fire activity in 2019 compared to the previous year (Brando et al., 2020). Furthermore, the
- 50 fire severity is potentially exacerbated by increased ecosystem fragmentation: aggregate impacts of human activity reduce the buffer that would have been present in a healthier and more intact ecosystem, making the Amazon more vulnerable to fire impacts (Fu et al., 2013; Aragão et al., 2018; Alden et al., 2016; Brando et al., 2020).

The associated carbon response to large scale fire activity is complicated given that the Amazon is not evolutionarily adapted to fires: while fires have played a historical role in the region, they have almost all been started by humans, generally for agricultural purposes (Uhl et al., 1988; Brando et al., 2020). As such, it is uncertain how prior burn trauma impacts future

- ecosystem productivity (Trumbore et al., 2015). Yet the direct and indirect carbon effects of the fires ravaging the Amazon during the dry seasons, particularly along the south-eastern frontier termed the "Arc of Deforestation" (Fig. 1), are playing an increasing role in the region's Net Biome Exchange (NBE) (Gatti et al., 2021). With the dry-season length of southern Amazonia already increasing significantly by approximately 6.5 days per decade, conditions conducive to enhanced fire
- 60 seasons are further prolonged (Fu et al., 2013). Therefore, we are at a critical time for both the Amazonian and global carbon balance where a warmer, drier Amazon is poised to face additional pressures from the combined effects of both drought- and deforestation-related fires (Brando et al., 2020; Naus et al., 2022). Furthermore, under Brazil's volatile political regime, any re-established protections face an uncertain future (Brando et al., 2020; Naus et al., 2022; Gatti et al., 2023).





Despite the carbon response of the Amazon being a key variable in climate modeling (e.g., Cox et al., 2000, 2013), regional
surface flux estimates of CO₂ as well as the atmospheric measurements to constrain them have been historically sparse due to physical, economic, and political limitations. As a result, capturing Amazon carbon surface fluxes and their sensitivities has been a challenge (Liu et al., 2017). However, the past two decades have seen an increase in multi-platform measurement efforts with the potential to greatly improve our ability to quantify carbon cycling in the Amazonian biosphere. Ground-based carbon flux measurements through AmeriFlux and the Amazon Tall Tower Observatory (ATTO) provide important surface-based
ecosystem information at a high temporal resolution (e.g., Basso et al., 2023; Paca et al., 2022; Hayek et al., 2018). Aircraft observations have provided important constraints enabling in-depth assessments of land-atmosphere carbon exchange (e.g., Gatti et al., 2010, 2014, 2021; Gloor et al., 2012; van der Laan-Luijkx et al., 2015; Alden et al., 2016). Beginning in 2010, sustained semimonthly aircraft measurements have been conducted in the four corners of the Brazilian Amazon as an international effort among researchers, informing estimates of fire effects in the region (Gatti et al., 2021). Finally, total column
CO₂ retrievals and solar-induced fluorescence (SIF) from NASA's Orbiting Carbon Observatory (OCO-2; operational

September 2014) enable previously inferred features of the tropical biosphere's carbon response to climatological variability to be more accurately estimated at multiple spatiotemporal scales (Bloom et al., 2016).

With the Amazon becoming warmer and drier, the impacts of fires and other disturbances—both climate-related and directly through deforestation and other degradation—on carbon stock loss are compounded (Barkhordian et al., 2019; Kruid
et al., 2021). However, absent a consistent observationally constrained multi-decadal record extending into the present, where CO₂ surface fluxes from both photosynthesis and respiration are captured, our capacity to evaluate long-term shifts in carbon dynamics of the South American tropics is limited. Existing global flux products with multi-decadal temporal records such as Gross Primary Productivity (GPP) estimates from NASA's MODIS have been found to inadequately represent complex Amazonian ecosystems (Almeida et al., 2018). In recent years, the availability of global satellite-based SIF observations—a
more direct correlate to photosynthetic activity, including in the Amazon—has greatly improved our ability to capture the

- timing and magnitude of ecosystem carbon exchange (e.g., Doughty et al., 2019; Koren et al., 2018; Zhang et al. 2018a; Luus et al. 2017). To this end, our study uses SIF along with a suite of readily available ground-to-satellite observations to estimate hourly biogenic CO₂ fluxes (µmol CO₂ m⁻² s⁻¹) via an improved version of the Vegetation, Photosynthesis, and Respiration model (VPRM) (Mahadevan et al., 2008). The VPRM is a minimally parameterized data-driven diagnostic ecosystem light-
- 90 use efficiency model that has many advantages for exploring shifts in regional carbon dynamics and modes of carbon exchange, particularly in its translation of local ecosystem and satellite observations into spatially resolved fluxes of CO₂ as Net Ecosystem Exchange (NEE), GPP, and R_{eco} at fine spatial (here, 12 km) and temporal (hourly) scales. The VPRM has demonstrated effectiveness as a prior in diverse CO₂ source attribution studies at midlatitudes and the Arctic (e.g., Dayalu et al., 2018; Luus et al. 2017; Hilton et al., 2013; Matross et al., 2006). Furthermore, the region-specific approach of the VPRM
- 95 contrasts with subsets of existing global vegetation carbon flux models which, in data sparse regions such as the Amazon, can be insufficiently resolved (Dayalu et al., 2018; Luus et al., 2017; Hilton et al., 2013; Matross et al., 2006). The VPRM is calibrated with direct ecosystem observations when available for each land use category in the domain of interest. Traditionally,





satellite surface reflectance-based proxies such as the Enhanced Vegetation Index (EVI) and the Land Surface Water Index (LSWI) are incorporated into the VPRM; however, there are known issues with that approach, particularly pertaining to LSWI inadequately capturing the effects of water stress (Dayalu et al., 2018; Luus et al., 2017).

The SIF-based formulation of the VPRM has improved performance relative to the traditional VPRM (VPRM-TRA) when compared against aircraft and tower observations in extratropical settings (Luus et al., 2017). However, the SIF-based VPRM holds promise for the tropics as well: SIF is implicitly and significantly correlated to previously hard-to-quantify parameters like water stress (e.g., Mohammadi et al., 2022). Particularly in the Amazon, where moisture can be a strong control on carbon

- 105 cycling (Gatti et al., 2014), using SIF includes early stage ecosystem response to moisture availability in the VPRM in a way that traditional vegetation indices do not (Zhang et al., 2018a). Similar to Luus et al. (2017), our work adapts the VPRM formulation where the traditional MODIS-based satellite surface reflectance data is entirely replaced with SIF data derived from OCO-2 measurements (VPRM-SIF). In addition, our work further adapts the respiration term of the VPRM similarly to Winbourne et al. (2022) and Gourdji et al. (2022) where the amount of living biomass per pixel informs the autotrophic
- 110 respiration term. The adaptation of the VPRM respiration term addresses a tendency for the VPRM to systematically underestimate autotrophic respiration, at least in the extratropics (e.g., Dayalu et al., 2018, Gourdji et al., 2022). In the tropics, where respiration is a strong control on NBE, models that provide more accurate respiration parameterizations are needed. The biogenic carbon fluxes generated by SIF-based VPRM formulations provide the basis for this study's quantification and categorization of the spatial and temporal variations in the Amazon's carbon sink strength from 2010–2020.
- 115 In our study, we expand upon the latest research for the Amazon, leveraging observations where available, to construct, improve, and evaluate biosphere carbon fluxes. This is a critical first step to evaluating long-term anthropogenic impacts on and trends in the Amazon's carbon balance. We broadly define Amazonia's wet and dry seasons as six-month periods, with the wet season typically extending from December through May, and the dry season extending from June through November. We construct and evaluate multiple versions of the VPRM: the traditional version before and after modifying respiration
- 120 (VPRM_TRA, VPRM_TRG); and the SIF-based version before and after modifying respiration (VPRM_SIF, VPRM_SIFg). The VPRM CO₂ fluxes are generated for all hours from 2014 through September 2020. As an additional test, we compare all versions of the diagnostic VPRM to output from the process-based bottom-up Simple Biosphere 4 model (SiB4; Haynes et al., 2021), which to our knowledge is the only publicly available hourly partitioned biogenic carbon flux estimate for the Amazon that extends through at least 2018. The hourly VPRM fluxes also include two main events which, as suggested by Liu et al.
- (2017) and Brando et al. (2020), are increasingly representative of the Amazon's future conditions and impacts on the global carbon balance: the strong El Niño in 2015–2016, and the severe dry-season fires in 2019/2020. In this paper, we additionally evaluate model performance during March 2016 by constraining with OCO-2 land nadir/land glint (LNLG) total columns and evaluating against CO₂ observations from aircraft vertical profiles. The March 2016 case study was selected for three main reasons: (1) it falls well within the Amazon wet season such that the biosphere dominates land-atmosphere CO₂ exchange such
- 130 that model-measurement mismatch can largely be attributed to the biosphere flux model alone rather than entangling the influence of fires, and (2) it falls at the tail end of the severe 2015/2016 El Niño and more strongly illustrates potential strengths





and weaknesses in how the various models capture the impacts of temperature and water stress on Amazonian land-atmosphere carbon exchange; and (3) there was distinct and separable fire activity which provides an opportunity to evaluate the capacity of the biogenic flux models to separate NEE from fire fluxes. Figure 1 displays our study domain along with key modeled and observational data sets used in top-down constraints and evaluations.

2 Methods

2.1 Data Processing Tools

We used R v4.2.1(<u>https://cran.r-project.org/</u>) for all data processing and analysis. We used the Weather Research and Forecasting Model (WRF) version 3.8.1 to model the meteorology driving the Stochastic Time-Inverted Lagrangian Transport
(STILT) model. WRFv3.8.1 configuration is detailed in Table S1.

2.2 VPRM: Estimating Vegetation CO₂ Fluxes

VPRM CO₂ fluxes from vegetation are modelled as NEE, GPP, and R_{eco} and reported as μmol CO₂ m⁻² s⁻¹ (hourly, 12 km resolution) with the convention of negative fluxes indicating uptake from the atmosphere and positive fluxes indicating release
 to the atmosphere.

2.2.1 Model Overview

The traditional and SIF-based VPRM formulations—without (VPRM-TRA, VPRM-SIF) and with (VPRM-TRG, VPRM-SIFg) respiration term modifications—provide hourly fluxes of biogenic CO₂ NEE partitioned into GPP and R_{eco}. Equations 150 1(ab) and 2(ab) describe all VPRM formulations.

$$NEE_{VPRM_TRA} = -\left(\boldsymbol{\lambda} \times \mathbf{T}_{\text{scale}} \times \frac{1}{1 + \left(\frac{PAR}{PAR_0}\right)} \times PAR \times P_{\text{scale}} \times W_{\text{scale}} \times EVI\right) + (\boldsymbol{\alpha} \times T + \boldsymbol{\beta})$$
(1a)

$$NEE_{VPRM_TRG} = -\left(\boldsymbol{\lambda} \times \mathbf{T}_{\text{scale}} \times \frac{1}{1 + \left(\frac{PAR}{PAR_0}\right)} \times PAR \times P_{\text{scale}} \times W_{\text{scale}} \times EVI\right) + \left((\boldsymbol{\alpha} \times T + \boldsymbol{\beta}) + (\boldsymbol{\gamma} \times EVI)\right)$$
(1b)

$$NEE_{VPRM_SIF} = -\left(\boldsymbol{\lambda} \times T_{\text{scale}} \times \frac{1}{1 + \left(\frac{PAR}{PAR_0}\right)} \times PAR \times \frac{SIF}{\cos(SZA)}\right) + (\boldsymbol{\alpha} \times T + \boldsymbol{\beta})$$
(2a)

$$NEE_{VPRM_SIFg} = -\left(\boldsymbol{\lambda} \times T_{\text{scale}} \times \frac{1}{1 + \left(\frac{PAR}{PAR_0}\right)} \times PAR \times \frac{SIF}{\cos(SZA)}\right) + \left((\boldsymbol{\alpha} \times T + \boldsymbol{\beta}) + \left(\boldsymbol{\gamma} \times \frac{SIF}{\cos(SZA)}\right)\right)$$
(2b)





The bolded parameters λ (ecosystem light use efficiency, units: µmol CO₂ m⁻² s⁻¹/µmol Photosynthetically Active Radiation (PAR) m⁻² s⁻¹); PAR₀ (ecosystem half-saturation value of PAR, units: µmol m⁻² s⁻¹); α (ecosystem respiration temperature dependence, units: µmol CO₂ m⁻² s⁻¹°C⁻¹); β (µmol CO₂ m⁻² s⁻¹; flux-weighted mean size of the respiring carbon pools), and γ (µmol CO₂ m⁻² s⁻¹; sensitivity to water stress and above-ground biomass) are all optimized from non-linear least squares (NLS) fits to ground-based measurements (e.g., Mahadevan et al., 2008; Gourdji et al., 2022). Domain-wide hourly fluxes are then derived for each land use category present. Our domain land categories are based on the International Geosphere-Biosphere Programme (IGBP) and obtained from the MODIS MCD12Q1 0.05° gridded product. Calibration and

land classification details are provided in Sect. 2.2.2.

In all formulations, hourly photosynthetically active radiation (PAR, units: μ mol m⁻² s⁻¹; Eq. 3) drives the diurnal flux signal and T_{scale} (unitless; see Eq. 4) provides ecosystem temperature sensitivity. When scaling fluxes to the entire domain,

165 ERA5 meteorology (<u>https://registry.opendata.aws/ecmwf-era5/;</u> Accessed 9 Aug 2023) at 31 km horizontal resolution provides the hourly downwelling shortwave radiation (SW) used to derive diurnal PAR; 2-meter surface air temperature estimates (T_{air}) are used to calculate T_{scale} .

$$PAR = SW/0.505 \tag{3}$$

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$$T_{\text{scale}} = \frac{(T_{air} - T_{min}) \times (T_{air} - T_{max})}{\left[(T_{air} - T_{min}) \times (T_{air} - T_{max}) - (T_{air} - T_{opt})^2 \right]},$$
 (4)

As shown in Eq. 4, T_{scale} modifies the GPP term by weighting the differences between observed T_{air} and the ecosystemspecific requirements for minimum (T_{min}), maximum (T_{max}), and optimal (T_{opt}) temperatures for photosynthesis. T_{min} , T_{max} , and T_{opt} values are informed by literature values relevant to the tropics (Ma et al., 2017; Slot & Winter, 2017; Tan et al., 2017).

- 175 For all tropical ecosystems, T_{min} is set to 4°C. Following work by Slot & Winter (2017) we broadly divide our ecosystem classes into tropical dry and tropical wet ecosystems. Tropical dry ecosystems (pixels in domain with IGBP categories of woody savannas, savannas, and grasslands) are assigned a T_{max} of 41.8°C (SD=2.1°C) and T_{opt} of 30.7°C (SD=1.1). Tropical wet ecosystems (pixels in domain with IGBP category of evergreen broadleaf) are assigned a T_{max} of 40.1°C (SD=1.8°C) and T_{opt} of 29.8°C (SD=0.9°C).
- In the traditional formulations (Eqs. 1a and 1b) the unitless scalars P_{scale} (phenology; Eqs. 4a-c), W_{scale} (water stress; Eq. 5), and EVI (surface "greenness"; Eq. 6) modify GPP based on remotely sensed measurements of surface reflectance in the near infrared (ρ_{nir}), shortwave infrared (ρ_{swir}), blue (ρ_{blue}), and red (ρ_{red}) bands.

$$P_{\text{scale,non-evergreen}} = \frac{1 + \frac{\rho_{nir} - \rho_{swir}}{\rho_{nir} + \rho_{swir}}}{2}$$
(4a)
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$$P_{\text{scale,evergreen}} = 1$$
(4b)



$$P_{\text{scale,water}} = 0 \tag{4c}$$

$$W_{\text{scale}} = \frac{1 + \frac{\rho_{nir} - \rho_{swir}}{\rho_{nir} + \rho_{swir}}}{1 + m_{swir} - \rho_{swir}} \tag{5}$$

$$EVI = 2.5 \times \frac{\rho_{nir} - \rho_{red}}{\rho_{nir} + (6 \times \rho_{red} - 7.5 \times \rho_{blue}) + 1}$$
(6)

190 The SIF-based VPRM formulations (Eqs. 2a and 2b) entirely replace the $P_{scale} \times T_{scale} \times EVI$ quantity with instantaneous spatially contiguous SIF derived from OCO-2 measurements (4-day, 0.05° resolution as in Zhang et al., 2018a). However, as shown in Eq. 7, raw SIF has a PAR dependence (Zhang et al., 2018b; Zhang et al., 2020) which requires normalizing by top-of-canopy PAR—approximated as the cosine of the solar zenith angle (SZA) at the local OCO-2 overpass time of 1330h—before the diurnal PAR signal can be meaningfully introduced as written in Eqs. 2a-b.

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$$SIF = PAR \times fPAR_{chl} \times EF \tag{7}$$

In Eq. 7, *fPAR_{chl}* is the fraction of PAR absorbed by chlorophyll and EF is the fluorescence efficiency. Daily SZA data for each calibration site was obtained from the NASA Horizons portal (<u>https://ssd.jpl.nasa.gov/horizons/app.html</u>; Accessed
15 Feb 2021). When scaling to our domain, we use the solarPos package in R to calculate the average SZA at each pixel at 1330 local overpass time averaged over each four-day interval of the contiguous SIF product.

Finally, two of our VPRM formulations (Eqs. 1b and 2b) also incorporate a γ (µmol CO₂ m⁻² s⁻¹) term adapting recent modifications that better represent R_{eco} responses associated with biomass seasonality and/or water stress (Winbourne et al., 2022; Gourdji et al., 2022). In the tropics, Clark et al. (2013) note that temperature sensitivity of night-time plant respiration can have a significant control on aboveground biomass production. As shown in Eq. 7, Gourdji et al. (2022) apply a quadratic temperature formulation to the traditional VPRM for *R_{eco}* which allows for a nonlinear temperature response; furthermore, *EVI* and *W_{scale}* are explicitly incorporated to account for aboveground biomass seasonality and water stress:

$$R_{eco} = (\boldsymbol{\alpha}_1 \boldsymbol{T}_{air} + \boldsymbol{\beta}) + (\boldsymbol{\alpha}_2 T_{air}^2) + (\boldsymbol{\gamma} \boldsymbol{EVI}) + (\boldsymbol{\theta}_1 W_{scale}) + (\boldsymbol{\theta}_2 W_{scale} T_{air}) + (\boldsymbol{\theta}_3 W_{scale} T_{air}^2)$$
(8)

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However, given both the spatial and temporal limitations of our Amazon region calibration sites (see Sect. 2.1.2), applying the Gourdji et al. (2022) respiration parameterization risks overfitting Amazonian respiration to a small number of specific sites. Furthermore, Winbourne et al. (2022) found that the VPRM respiration term approached the traditional $\alpha \times T + \beta$ linear result during the growing season in a USA urban environment. In tropical environments, where growing season conditions last year-round, we therefore represent R_{eco} with the bolded terms in Eq. 8—that is, as a linear function of air temperature with an

additional γ term that uses *EVI* or *SIF/cos(SZA)* to incorporate the influence of aboveground biomass seasonality and water stress.

2.2.2 Domain Land Use Classification, Model Calibration, and Model Evaluation

- Prior to generating vegetation carbon fluxes for the entire study domain, the VPRM model is calibrated using NLS to 220 ground-based flux data collected for each major vegetation class. In our Amazonian study domain, ground-based data availability is an ongoing challenge; our calibration data was therefore restricted to publicly available data from the AmeriFlux and FluxNet eddy flux networks. Data from the eight available Amazon-relevant eddy flux sites span limited IGBP vegetation classes and subsets of time from 2001–2015 (Table 1, Fig. 2). With the exception of the tropical rainforest site Br-K67 (discussed later in this section), the eddy flux data is insufficient for separately fitting wet and dry seasons. Therefore, we
 - 225 perform single NLS fits to calibration data which result in constant VPRM parameters for each ecosystem type over the study period. While the calibration data set is not dense enough to span both the spatial and temporal extents of our study, we note that it does provide valuable and sufficient information to establish reasonable *a priori* vegetation CO₂ flux estimates that can be meaningfully constrained with OCO-2 CO₂ observations for two main reasons. First, approximately 95% of the land use in the study domain from 2010–2020 is encompassed by the four IGBP categories for which eddy flux data was available,
 - 230 namely: evergreen broadleaf (50%); savannas (20%); grasslands (20%), and woody savannas (5%) (Fig. 2b). Three of the eight sites represent the evergreen broadleaf ecosystems which is the most prevalent broad class in the domain. While we recognize the IGBP "evergreen broadleaf" categorization is an oversimplification of heterogeneous Amazonian evergreen ecosystems, we note the value in tuning of the VPRM to three separate evergreen broadleaf sites. Second, while the available eddy flux data can be offset up to thirteen years prior to the study period of 2014–2020, the major drivers of hourly ecosystem flux
 - variations are provided by T_{air} , PAR, surface reflectance indices, and SIF. That is, while the static ecosystem parameters of λ , PAR₀, α , β , and γ would benefit from tuning to eddy flux data seasonally and/or over the entire study period to reflect concurrent ecosystem states most accurately, the dominant source of real time variation is largely captured by T_{air} , PAR, surface reflectance indices, and SIF (Dayalu et al., 2018).
 - As shown in Table 1, each broad ecosystem type is assigned a representative calibration and evaluation site. The exception 240 is site Br-K34, which is the only representative of primary interior tropical rainforest and there was insufficient data to divide the data into calibration and evaluation years. As the remaining tropical rainforest sites (Br-K67, Br-K83) are impacted by higher degrees of disturbance and edge effects, comparison with Br-K34 was not appropriate. We therefore manually differentiate two categories of evergreen broadleaf/IGBP category 2: (1) western Amazon IGBP category 2 is designated as primary interior tropical rainforest and its pixels are represented by NLS parameters obtained from site Br-K34; (2) remaining
 - 245 IGBP category 2 primary tropical moist forest occurring in domain is represented by NLS parameters obtained from site Br-K67 (Fig. S1).

As noted earlier, the long record of Br-K67—spanning 2002–2011—enables assessing uncertainty associated with using annual fit VPRM parameters rather than seasonally fit. We therefore used the Br-K67 record to evaluate the impact of

parameter seasonality on model performance. Accounting for data gaps, a total of seven years of hourly Br-K67 observations were randomly separated into wet and dry seasons subsets for VPRM calibration (70% from each of wet and dry season data) and evaluation (30% from each of wet and dry season data). We then performed seasonal NLS fits for each VPRM version. We used the seasonal and annual fits in an assessment of model bias for the tropical rainforest/evergreen broadleaf class relative to observations.

Table 1. Eddy flux sites used for VPRM calibration and/or evaluation. *Br-K34 site represents interior relatively undisturbed evergreen broadleaf classes; K34 data is unvalidated due to insufficient representative data. ⁺⁺Br-PDG was the only member of IGBP 9 class and separate years were used for calibration (2003) and evaluation (2001–2002).

| Site | Coordinates | Data | Land Use (IGBP Category) | VPRM Role | Data Source |
|-------------------|-------------|-------|--|-------------------|-------------------|
| | | Years | | | |
| Br- | -9.82N, | 2003- | Seasonally flooded transitional | Evaluation (CST) | Restrepo-Coupe et |
| BAN | -50.1E | 2006 | forest/savanna (8/Woody Savannas) | | al. (2021) |
| Br- | -7.97N, | 2014- | Tropical dry forest/wet season cattle | Calibration | Antonino (2022) |
| CST | -38.4E | 2015 | grazing (8/Woody Savannas) | | |
| Br- | -10.8N, | 2001- | Pasture (10/Grasslands) | Calibration | Restrepo-Coupe et |
| FNS | -62.4E | 2002 | | | al. (2021) |
| Br- | -2.61N, | 2001- | Primary interior tropical rainforest | Calibration | Restrepo-Coupe et |
| K34* | -60.2E | 2002 | (2/Evergreen Broadleaf) | | al. (2021) |
| Br- | -2.89N, | 2002- | Primary tropical moist forest (2/ | Calibration | Saleska (2019) |
| K67 | -55.0E | 2011 | Evergreen Broadleaf) | | |
| Br- | -3.01N, | 2001- | Pasture/agriculture (10/Grasslands) | Evaluation (FNS) | Restrepo-Coupe et |
| K77 | -54.5E | 2005 | | | al. (2021) |
| Br- | -3.02N, | 2001- | Primary tropical moist forest, selective | Evaluation (K67) | Goulden (2019) |
| K83 | -55.0E | 2004 | logging (2/Evergreen Broadleaf) | | |
| Br- | -21.6N, | 2001- | Savanna (9/Savannas) | Calibration (PDG- | Restrepo-Coupe et |
| PDG ⁺⁺ | -47.6E | 2003 | | 2003); Evaluation | al. (2021) |
| | | | | (PDG-2001, 2002) | |

2.3 SiB4

- 260 The SiB4 model version 4.2 (SiB4) integrates land cover, phenology, dynamic carbon allocation, cascading carbon pools from live biomass to surface litter to soil organic matter, and NASA MERRA2 meteorology (Haynes et al., 2021). Unlike the VPRM it is a mechanistic model that is independent of satellite indices and site-specific tuning parameters (Haynes et al., 2019). We obtained SiB4 global hourly carbon fluxes partitioned into GPP and Reco at 0.5x0.5deg resolution from 2000–2018 from the Oak Ridge National Laboratory Distributed Active Archive Center (ORNL DAAC; Haynes et al., 2021). SiB4 fluxes 265 for each gridcell are disaggregated into one of 15 plant functional types (PFTs). We convert SiB4 fluxes per PFT to total fluxes
 - per grid cell using the provided area per grid cell. We then subset these grid cell total SiB4 fluxes over the VPRM domain (Fig. 1) and regrid to the VPRM model resolution using bilinear interpolation. Additional details and technical description of the SiB4 model v4.2 are provided in Haynes et al. (2020).

2.4 Regional Inversion Methodology

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In this study we apply the CarbonTracker-Lagrange regional inverse modeling framework Hu et al. (2019), Rastogi et al. (2021), Rastogi et al. (2021b). It uses a Bayesian inversion methodology with a geophysically-based simulation of satellite retrievals of column CO₂ profiles (Sec. 2.4.1), and sensitivities of the satellite observations to upstream surface fluxes (i.e. "footprints") derived from WRF-STILT (Sec. 2.4.2). The March 2016 period was extracted from a three-month inversion, spanning February 2016 through April 2016, to account for edge effects. On average, wet season OCO-2 data coverage was ~15% less than dry season coverage.

We use OCO-2 total column CO₂ (XCO₂) retrievals to constrain VPRM CO₂ flux estimates in a geostatistical Bayesian framework, where we write the posterior flux solution as:

$$s = s_a + QH^T (R + HQH^T)^{-1} (X_{CO2}^{ret} - X_{CO2}^{sim}),$$
(9)

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where X_{CO2}^{ret} is the $n \times 1$ vector of retrieved XCO₂. X_{CO2}^{sim} (Eq. 10) is the $n \times 1$ vector of simulated XCO₂. *H* (Eq. 11) is the column weighted sensitivity of X_{CO2} to surface fluxes upwind of the measurement, otherwise known as the column weighted footprint $[n \times m]$ or Jacobian. *s* is the posterior flux estimate $[1 \times m]$. *R* is the model-data mismatch covariance matrix $[n \times n]$, which ideally includes uncertainty due to geostatistical measurements, forward model, model representation, and boundary conditions. Given the difficulty in estimating these uncertainties, in our application, R is estimated from the uncertainty in XCO₂ using the methodology of Peiro et al. (2022) with some modifications, discussed next. *Q* is the prior flux error covariance matrix $[m \times m]$, and in our application is calculated using the standard deviation of the temporally averaged prior fluxes, a temporal correlation length of 7 days, and a spatial correlation length of 1000 km. s_a is the prior flux estimate $[1 \times m]$ from the VPRM or SiB4 flux models. In this study, 2644 retrievals (*n*) are used over the February to April 2016 time frame and 1266 land grid cells are optimized over the domain, hourly, for 2160 hours (m=2,734,560).

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Our methodology follows that for comparing satellite retrievals to simulated CO₂ columns from the Atmospheric CO₂ Observations from Space (ACOS) retrieval algorithm v10 (O'Dell et al., 2012). We follow the convolution method as detailed in Rastogi et al. (2021) to calculate the simulated XCO₂:

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$$X_{CO2}^{sim} = \sum_{i=1}^{N} w_i \Big[\alpha_i * \left(X_{CO2,i}^{bkg} + H_i (s_{bio} + s_{other}) \right) + (1 - \alpha_i) * X_{CO2,i}^{pri} \Big],$$
(10)

where N is the number of vertical levels. w_i is the pressure weighting function calculated as in Rastogi et al. (2021). α_i is the retrieval averaging kernel profile. $X_{CO2,i}^{bkg}$ is the CO₂ background profile from NOAA's CarbonTracker version CT2019B Jacobson et al. (2020). s_{bio} is the biospheric flux estimate from VPRM or SiB4 models. $X_{CO2,i}^{pri}$ is the OCO-2 prior CO₂ profile. H_i is the sensitivity function or footprint on discrete levels from WRF-STILT (Sec. 2.5.2):

(11)

$$H = \sum_{i=1}^{N} w_i \alpha_i H_i$$

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In this application, N is 17 levels. This accounts for 14 levels of WRF-STILT footprints, discussed in section 2.5.2, and 3 levels in the upper atmospheric column where the impact of surface fluxes are assumed to be negligible and the footprints are set to zero. Thus in the upper atmosphere, the X_{CO2}^{sim} calculation is dominated by the CarbonTracker background profile and OCO-2 prior CO₂ profile.

For this study, we assimilate land nadir and glint OCO-2 XCO₂ retrievals that are averaged to 2s along track (~13.5km) – rather than 10s averages – to better match the higher spatial resolution of our inversion domain (12 km) compared to global model resolutions. We average the OCO-2 XCO₂ 2s data following methodology from Peiro et al. (2022), where a 10s average dataset was used. However, for the application of 2s data over land we estimate the uncertainty in 2s XCO₂ (R) using a higher land correlation coefficient of 0.7, compared to 0.3 in Piero et al. (2022). In our application we also neglect transport model errors, which are difficult to quantify but can be further explored in the future using transport ensembles. Errors in WRF-STILT transport are discussed in Rastogi et al., (2021) in terms of calculating *X*^{sim}_{CO2} for OCO-2 XCO₂ retrievals in North America. They found that low partial column bias relative to independent vertical profile CO₂ data show that errors

315 in North America. They found that low partial column bias relative to independent vertical profile CO₂ data show that errors in WRF-STILT transport contribute very minimally to bias in X_{CO2}^{sim} .

2.4.2 Transport Model Framework

We use WRFv3.8.1 model transport fields to drive the STILT model dispersion and compute footprints. A footprint (H_i), in units of mixing ratio per unit flux (e.g., ppm µmol⁻¹ m⁻² s⁻¹), quantitatively describes how the surface fluxes upwind of a mixing ratio measurement location are influencing the measurement. For each OCO-2 XCO₂ retrieval location, footprints are computed at 14 individual levels throughout the column. These 14 footprints are then convolved with OCO-2's pressure weighting function and averaging kernel, and interpolated to STILT levels. This produces one column weighted footprint (H) per retrieval, which quantifies how the OCO-2 XCO₂ retrieval is influenced by surface fluxes upwind of the column. The column weighted footprint is not to be confused with a satellite footprint, which describes the area of earth reflecting the satellite signal. The computational domains of the two nested WRF grids are shown in Figure 1a. Model configuration details

325 satellite signal. The computationa are provided in Table S2.

STILT computes footprints by modeling the dispersion of 500 particles, 10 days back in time from each measurement location (in this case the x,y,z,t coordinate of an XCO₂ retrieval), and in 14 levels from 50 m to 14 km AGL. where each receptor is located at a column measurement latitude and longitude and on one of 14 levels (50m to 14km AGL). Back

330 trajectories computed by STILT are affected by both resolved wind velocities and parameterized sub-grid scale turbulent motions and convective fluxes. The footprint domain is outlined in Figure Error! Reference source not found.b, where

footprints are aggregated to 1 degree resolution at 1-hour intervals, compatible with the VPRM prior flux resolution used in the inversion.

2.5 Model Aircraft Vertical Profile Calculation

- We calculate prior and posterior modelled vertical profiles for SiB4 and each VPRM formulation for our March 2016 optimization period at locations roughly corresponding to the vertical profiling sites RBA and ALF displayed in Fig.1a. As the available OCO-2 receptors are not identical in space and time to locations of RBA and ALF profiling sites, a direct comparison of simulated prior and posterior CO₂ vertical profiles derived from convolving multi-level OCO-2 receptor footprints and vegetation flux models is not possible. In addition, RBA and ALF vertical profiles are typically obtained once or twice a month such that robust measured monthly averages for a single month are not available. To allow for direct comparison between modelled (prior and posterior) and measured vertical profiles, we construct pooled datasets that occasionally combine February and March 2016 measurements and/or modelled fields to develop a dataset that adequately represents a typical 2016 wet season month. Given the seasonal similarities across the Amazon in February and March, combining data across these months to create a representative "typical wet season month" is reasonable. Our method is detailed below.
- We first construct a pooled dataset of "typical wet season 2016" measured vertical profiles at each of RBA and ALF sites. For RBA, we combine all measurements obtained between February and March 2016 (2016-02-08 at 1630UTC; 2016-02-27 at 1645UTC; and 2016-03-17 at 1700UTC), resulting in three measurements at each of 17 vertical levels between 300 and 4500 m asl. For ALF, we combine all measurements obtained between February and March 2016 (2016-02-23 at 1630UTC; 2016-02-29 at 1540UTC; 2016-03-13 at 1600UTC; and 2016-03-30 at 1540 UTC), resulting in four measurements at each of
- 350 12 vertical levels between 450 and 4500 m asl. Second, we assess the availability of OCO-2 footprints in the vicinity of RBA and ALF sites from February to March 2016. Our goal was to obtain footprints sufficiently close to each profiling site to be representative of the near-field influences on the site, but also have a large enough bounding box so that at least two OCO-2 receptors and their footprints were present for transport uncertainty calculations. Figure 2a displays the final selected 5x5 km bounding boxes around each of RBA and ALF and the representative OCO-2 receptors. RBA and ALF simulated vertical
- 355 profiles are then derived from the OCO-2 receptor footprints bounded in each box (Fig. 2a). Third, we use NOAA's web-based HYSPLIT model to assess land surface influences on each of the measured vertical profile dates and compare them with the land surface influences on each of the selected OCO-2 receptors. Figures S2-S3 show that the land surfaces influencing both the simulated and actual profiles at RBA and ALF are comparable with the airmass trajectories representing typical seasonal prevailing winds in the February/March 2016 time frame and annually from 2010–2018; the average air mass trajectories are
- 360 displayed in Fig. 2a. Fourth, we pool all vertically resolved CT2019 background concentrations associated with each footprint file's endpoints. Next, we use the March 2016 prior and posterior hourly fluxes to estimate a "typical wet season 2016 month of fluxes" and convolve them with each of the six OCO-2 receptors in the RBA and ALF bounding boxes to obtain a spread of enhancements or depletions relative to the CT2019 background CO₂ from 10 days prior. Finally, we linearly interpolate all components on WRF-STILT vertical grids to each of the RBA and ALF vertical grids; calculate a month's worth of hourly

365 model-observation residuals at each vertical level; and extract means, 25^{th} , and 75^{th} percentiles. We bootstrap CT2019 background concentrations and vertical profile measurements at each vertical level to dimensions that enable merging with the month of hourly *footprint* × *flux* fields. Ultimately, vertical profiles of modelled and measured residuals for the simulated month typical of the 2016 wet season incorporate uncertainties in transport, background, vertical profile measurements, and flux fields.

370 3 Results and Discussion

3.1 VPRM Calibration and Evaluation

Site-specific values of tuning parameters are provided in Table S1. NLS calibration results are summarized in Table 2. Table 2 provides the root mean square error (RMSE) of predicted vs. measured NEE from the NLS fits at each calibration site, along with the interquartile range (IQR) of measured NEE which allows for normalizing RMSE across the varied ecosystems

- 375 present in the domain. Generally, relative to the traditional VPRM formulations, the VPRM-SIF formulations provide improved fits (calibration) and predictions (evaluation) to hourly NEE observations. Of the VPRM formulations with SIF, the SIFg formulation is a slight improvement over the standard SIF formulation in net. An interesting feature of the respiration parameterization—relative to the typical non-tropical VPRM formulations—is the estimate of negative values for α in the respiration temperature dependency for the tropical dry forest Br-CST and interior primary rainforest Br-K34 sites. This is
- 380 likely a response to the strength of temperature as a driver of NEE and respiration: strong decreases in respiration in certain tropical ecosystems have previously been observed (Clark et al., 2013; Gallup et al., 2021).

We note that Br-K83—the Br-K67 evaluation site—is a selectively logged disturbed forest site (Figueira et al., 2008). We also note that site Br-K77 is a highly disturbed pasture/cropland site with a history of forest conversion to cropland in the 1990s followed by burning for rice cultivation in 2001. Br-K77 is therefore limited in its representation of steady-state cropland/grassland mosaic ecosystems: its poor performance as an evaluation site for the FNS cropland/grassland mosaic reflects its disturbed ecological history in the limited years for which we have eddy flux data. Specifically, as we have K77 eddy flux data only for 2001–2005, we are comparing an unstable ecology with FNS which began its trajectory to cropland as far back as 1977 (Almeida et al., 2018).

Given the greater importance of respiration on the NEE (and NBE) signal in the tropics relative to the extra-tropics, we also examine differences in respiration parameterization across all VPRM formulations. We use night-time NEE as a proxy for respiration (i.e., absent confounding effects of daytime uptake) focusing on the hours from 20:00 LST to 03:00 LST. Overall, we find that annually and by season, the SIF-based VPRM formulations in general—and VPRM-SIFg in particular have consistently better agreement with observed (night-time) respiration than the traditional formulations (Fig. 3, Table S3). On average, dry season respiration is better constrained across all model versions. In the wet season, while all models tend to

395 underestimate respiration, VPRM-TRA displays the greatest bias. The underestimate in wet season night-time respiration also implies that the general underestimate of peak daytime wet-season drawdown in NEE occurs through underestimating GPP

rather than through over-estimating R_{eco} . Of the VPRM formulations, the SIF-based formulations have more instances of overestimating respiration, especially in savanna ecosystems (Table S3).

Diurnally, the SIF-based formulations consistently perform better than the traditional formulations when compared to 400 observations annually and seasonally (Figs. S2–S3). Notably, for evergreen broadleaf classes—the dominant domain vegetation type—peak uptake tends to be underestimated in both seasons but especially in the wet season. We discuss this further in Sect. 3.1.1.

405 Table 2. Root mean square errors and correlation coefficients (R²) from NLS fits to calibration sites and associated evaluation site predictions. For evaluation sites, the site used for calibration is provided in the Cal Site column. Calibration (model fitting) results with R²>0.5 are in *regular italics*. Evaluation (model predictions) with R² values >=0.5 are in *bold italics*. VPRM model version shown as one of TRA, TRG, SIF, SIFg. *K34 data is unvalidated due to insufficient representative data. +K77 ecological history suggests limitations in its representation of steady-state grasslands. ++PDG was the only member of IGBP 9 class and separate years were used for calibration (2003, PDG_a) and evaluation (2001–2002, PDG_b). The InterqIQR of measured NEE at each site is also provided to normalize RMSE and facilitate comparison across the varied ecosystem types.

| Site | Cal Site | IGBP | NEE _{IOR} | TRA | TRG | SIF | SIFg |
|-----------------------------------|-----------------------------------|------------------|--------------------|--------------|-------------|------------------------|------------------------|
| | | Ecosystem/Number | C C | RMSE (R^2) | $RMSE(R^2)$ | RMSE (R ²) | RMSE (R ²) |
| Br-BAN | BR-CST | Woody Sav/8 | 17.2 | 11.4 (0.29) | 11.2 (0.34) | 10.48 (0.42) | 10.39 (0.43) |
| Br-CST | | Woody Sav/8 | 4.95 | 3.88 (0.43) | 3.77 (0.45) | 3.60 (0.50) | 3.36 (0.60) |
| Br-FNS | | Grasslands/10 | 15.8 | 8.21 (0.44) | 8.18 (0.44) | 7.33 (0.60) | 7.31 (0.60) |
| Br-K34* | | Evergreen Bdlf/2 | 21.2 | 8.30 (0.61) | 8.20 (0.62) | 8.55 (0.59) | 8.28 (0.62) |
| Br-K67 | | Evergreen Bdlf/2 | 19.9 | 6.05 (0.69) | 5.58 (0.74) | 5.67 (0.74) | 5.30 (0.77) |
| Br-K77 ⁺ | BR-FNS | Grasslands/10 | 7.81 | 6.84 (0.08) | 6.63 (0.13) | 6.58 (0.15) | 6.57 (0.16) |
| Br-K83 | BR-K67 | Evergreen Bdlf/2 | 19.4 | 7.59 (0.60) | 7.28 (0.63) | 7.00 (0.65) | 6.79 (0.67) |
| Br-PDGa ⁺⁺ | | Savannas/9 | 8.33 | 3.89 (0.51) | 3.87 (0.51) | 3.93 (0.53) | 3.91 (0.53) |
| Br-PDG _b ⁺⁺ | BR-PDG _a ⁺⁺ | Savannas/9 | 7.46 | 3.98 (0.58) | 4.05 (0.55) | 4.01 (0.60) | 4.08 (0.58) |

3.1.2 Impact of Seasonality on VPRM Performance: Br-K67

parameters on modelled relative to observed NEE both in aggregate and diurnally.

Wu et al. (2017) explore causes of GPP variability in Amazonian rainforests, focusing on the relative importance of light versus water limitations at the Br-K67 site, using data spanning 2002–2011. Wu et al. (2017) estimate monthly and constant light use efficiency (LUE; CO₂ uptake per unit of PAR) directly from the ratio of observationally derived GPP to PAR under reference (i.e., "non-stressed") environmental conditions. Their resulting reference monthly LUE ranges from 0.02 to 0.03 µmol CO₂ (µmol photons)⁻¹ over the seven years of Br-K67 eddy flux data in contrast to a constant LUE of ~0.02 µmol CO₂
(µmol photons)⁻¹. Wu et al. (2017) show seasonality in monthly LUE, with peak dry season values lower than peak wet season by as much as 33%. In this section, we present our results examining (1) seasonal variations in the LUE (i.e, the λ term in VPRM) term across model versions and compared to literature values; and (2) impacts of seasonality in VPRM fitting

Figure 4a displays the monthly average LUE obtained from VPRM NLS fitting to all available Br-K67 data, with LUE estimated by Wu et al. (2017) included for reference. We note that all versions of the VPRM indicate a seasonal cycle in LUE whose timing is similar to Wu et al. (2017). However, while the traditional VPRM formulations estimate a peak dry season LUE that is 66% lower than the peak wet season LUE, the difference in SIF-based formulations is 39–46% lower in the dry season. The traditional VPRM versions estimate monthly LUE that is frequently an order of magnitude higher than those

- estimated by the SIF-based versions and in Wu et al. (2017). In contrast, both SIF-based VPRM formulations estimate LUE
 that is generally higher but comparable to Wu et al. (2017); differences are ascribed to the LUE estimation methodology.
 Overall, the LUE from SIF-based VPRM formulations have significantly better agreement with Wu et al. (2017); given the direct relation between LUE and GPP, our results provide higher confidence in SIF-based GPP estimates over the traditional VPRM GPP estimates. Seasonal variations in diurnally averaged modeled PAR are shown in Fig. 4b-c.
- Given the seasonality in tropical forest LUE, and their dominance in domain land cover type, we next test our assumption
 that (1) the VPRM environmental drivers (i.e., SIF, Tscale, Pscale, EVI, LSWI, T_{air}) explain more of the real-time variation in carbon flux than the tuning parameters, and (2) constant annual-based tuning parameters can reasonably be used across seasons and years. Wu et al. (2017) have previously shown that while environmental drivers indeed explain the most variability at shorter (hourly) timescales, they tend to explain progressively less at longer timescales (diurnal, monthly, annual) when the influence from intrinsic ecosystem variables begin to aggregate. However, Wu et al. (2017) use an LUE-based photosynthesis
 model that is adopted from the VPRM-TRA formulation as in Mahadevan et al. (2008).

Table 3. Impact of calibration parameter seasonality on NEE predictability. Calibration parameters used in evaluation are obtainedfrom the annual set where NLS fitting was conducted for entire Br-K67 data set (NEE_{ann}), the wet season where calibration was conductedfor a random 70% subset of Br-K67 wet season data (NEE_{wet}), and the dry season where calibration was conducted for a random 70% subsetof Br-K67 dry season data (NEE_{dry}). Mean Bias is reported for each of wet and dry seasons relative to hourly NEE observations in the 30%seasonal evaluation subsets. % Error (model relative to observations) and R² values are provided in parentheses.

| | | Mean Bias, μmol CO ₂ m ⁻² s ⁻¹ (% error, R ²) | | | |
|--------------|------------------------|--|------------------------------|--|--|
| Model | Calibration Parameters | NEEwet, VPRM VS. NEEwet, Obs | NEEdry, VPRM VS. NEEdry, Obs | | |
| 4 . | NEEann | 1.92 (36.4, 0.72) | -1.21 (18.9, 0.74) | | |
| PR/V FRA | NEEwet | 0.041 (0.788, 0.72) | -4.54 (70.9, 0.74) | | |
| 5 | NEE _{dry} | 2.81 (53.4, 0.72) | 0.015 (0.230, 0.74) | | |
| Ļ., | NEEann | 0.402 (7.64, 0.77) | -0.239 (3.73, 0.76) | | |
| VPRN TRG | NEEwet | 0.020 (0.373, 0.77) | -1.48 (23.0, 0.76) | | |
| | NEEdry | 0.456 (8.67, 0.77) | -0.007 (0.103, 0.77) | | |
| VPRM- SIF | NEEann | 0.451 (8.57, 0.71) | -0.326 (5.09, 0.77) | | |
| | NEEwet | 0.064 (1.22, 0.71) | -1.15 (17.9, 0.77) | | |
| | NEEdry | 0.591 (11.2, 0.77) | -0.021 (0.324, 0.77) | | |

| VPRM- SIFg | NEE _{ann} | 0.362 (6.88, 0.78) | -0.173 (2.71, 0.78) |
|---------------|--------------------|--------------------|---------------------|
| | NEEwet | 0.146 (2.79, 0.78) | -0.592 (9.24, 0.78) |
| | NEE _{dry} | 0.469 (8.92, 0.78) | -0.00 (0.013, 0.78) |

Table 3 summarizes the results from a test where the wet and dry season Br-K67 evaluation subsets (described in Sect. 2.2.2) were used to predict seasonal NEE from annual- and seasonal-fit calibration parameters (Table S1). Overall, while all models generally underestimated net uptake in the wet season and underestimated net release in the dry season, the VPRM-SIFg formulation performed consistently better across all calibration parameters with hourly errors ranging from 0-10%. As expected, the optimal model-observation fit across all model versions occurred when season-specific parameters were used (% error < 3). However, when seasonal NEE was predicted using calibration parameters tuned to the opposite season, VPRM-TRA performed the worst (% error: 53–71), followed by VPRM-TRG (% error: 8.7–23), VPRM-SIF (% error: 11–18) and
VPRM-SIFg (% error: 8.9–9.2). Finally, errors in seasonal NEE predictions from annual parameters smoothed the impacts of lower cross-season predictability; however, even with the cross-seasonal smoothing, the VPRM-TRA performed relatively poorly (% error: 19–36%), particularly when compared to VPRM-SIFg (% error: 2.7–6.9).

Figure 5 displays diurnal patterns in seasonal predictability. While the diurnal averaging by season smooths the hourly model-observation mismatch displayed in Table 3, the diurnal breakdown suggests improved respiration parameterization in 460 the SIF-based VPRM formulations.

While Br-K67 seasonal predictions using annually fit parameters smoothed the impacts of lower cross-season predictability, the cross-season results (Table 3, Fig. 5) combined with the order-of-magnitude higher LUEs (Fig. 4a), indicate the traditional VPRM formulations are driven by environmental variables that inadequately represent the dominant land cover type in the study domain. The exceptionally poor VPRM-TRA results also brings into question the reliability of the VPRM-

- 465 TRG formulation given that they both incorporate the same environmental drivers. We also note that across all model versions, model-observation mismatch was highest in the wet season which is intuitive given the added complexity of cloud cover and associated impacts on uncertainties in remotely sensed environmental drivers and light use efficiency. The results from the seasonality case study suggest that the SIF-based formulations in general—and the VPRM-SIFg formulation in particular—are better equipped at estimating biogenic carbon fluxes at both short and longer timescales. We explore these results further
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Of note, while there can be conflicting evidence on light-limitations and seasonal carbon uptake in the Amazon, several studies provide evidence for enhanced dry-season carbon uptake when cloud cover is reduced and sunlight is more readily available (Huete et al., 2006; Saleska et al., 2016; Doughty et al., 2019). But while the GPP constituent of NEE can be higher in the dry season, wet season aircraft observations tend to show higher uptake in net (Gatti et al., 2010, 2014, 2021).

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3.2 Wet Season 2016 Case Study: Biogenic Model Evaluation and Estimates of Fire Influence

via the wet season 2016 optimization case study in Sect. 3.2.

We evaluate the VPRM performance during March 2016, corresponding to the middle of the wet season immediately following the peak of the 2015/2016 severe El Niño, and compare with SiB4 performance. We select the March 2016 wet season, as it is a valuable natural experiment that demonstrates the relative importance of biogenic fluxes and fire fluxes. The 480 severe El Niño corresponded to one of the hottest periods recorded over Amazonia in the past century; as shown in Fig. 6a, the 2016 wet season was characterized by a dipole of extreme drought in the eastern Amazon and unusual wetting in the western Amazon (Silva Junior et al., 2019). The anomalously hot and dry conditions in portions of the Amazon were more conducive to drought-related fire activity; however, Silva Junior et al. (2019) found the peak in Amazon area affected by fires during the first quarter of 2016 was approximately 2% with fire anomalies concentrated in the northern state of Roraima. 485 Integrated over February and March 2016, Aqua satellite fire detects were concentrated at the northern and southeastern boundaries of the Amazon Basin (TerraBrasilis, 2024). The emissions strength of these fires (as fire radiative power) was not readily available, and could not be used to weight their relative importance. Dominance of fire emissions relative to biosphere fluxes varied by location; we use this feature in combination with drought index information and measured aircraft vertical profiles to evaluate biogenic flux models and estimate the relative contribution of the biogenic activity and fires to sub-regional 490 CO₂ signal.

3.2.1 Optimization Results for Biogenic Flux Models

Figure 6 displays $1^{\circ} \times 1^{\circ}$ prior (Fig. 6b) and posterior (Fig. 6c) fluxes, and their differences (Fig. 6d), as μ mol CO₂ m⁻² s⁻¹ averaged over the month of March 2016. The optimization suggests that all prior models in wet season 2016 tended to 495 underestimate carbon uptake in the western Amazon; the region of underestimated uptake corresponds to the anomalously wet region identified through the Self-calibrated Palmer Drought Severity Index (ScPDSI) by Jimenez-Muñoz et al. (2016) (Fig. 6a). Relative to OCO-2 total column observations, both traditional VPRM formulations tend to underestimate uptake throughout the Amazon and strongly underestimate uptake in the region corresponding to the wetter than normal conditions. However, incorporating SIF observations into the VPRM formulations enhances the model's ability to capture uptake in 500 anomalously dry conditions. We note the demonstrated drought sensitivity of SIF, providing confidence in its ability to

quantify impacts of water availability on carbon uptake (e.g. Mohammadi et al., 2022).

Overall, our optimization results during the differentially drought-impacted 2016 wet season suggest that SIF-based VPRM formulations are better able to capture carbon uptake during varying tropical light and moisture regimes than the traditional surface-reflectance based VPRM formulations. We also find that the posterior solutions of the diagnostic SIF-based

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VPRM formulations—and the SIFg formulation in particular—converge with that of the bottom-up process-based SiB4 model (Fig. 6d). We further note that the SIF-based VPRM formulations provide significantly higher spatial heterogeneity relative to the SiB4 model, suggesting improved parameterization of the complexity of Amazonian ecosystems.

3.2.2 Comparison of Prior and Posterior Models at Aircraft Vertical Profiling Locations

- 510 We evaluate the prior and posterior performance of the VPRM and SiB4 models at the RBA and ALF vertical profiling sites, focusing on altitudes where upwind biogenic fluxes dominated the CO₂ signal. After qualitatively accounting for likely presence of fire emissions, we examine the impact of biogenic model optimization through simulating vertical profiles, diurnal cycles, and total monthly fluxes. Methods to calculate vertical profiles and total monthly flux are described in Sects.2.6 and 2.7.
- 515 Overall, we find the SIF-based VPRM formulations and the SiB4 simulate vertical profiles with significantly better modelobservation agreement than the traditional VPRM formulations. We discuss the details of the analysis below, noting that the RBA profiling location alone provided observations at altitudes where the biosphere is dominant. Fire activity in the region upwind of ALF was determined to be substantial enough at all altitudes such that biogenic model evaluation—i.e., evaluation absent other significant and confounding sources of CO₂—would not be valid. Figures 7 and 8 display modelled and measured
- 520 vertical profiles and their 25th and 75th percentiles for a typical 2016 wet season month at RBA and ALF, where "typical" is defined using data across February and March 2016 (Sect. 2.6). At RBA, this period corresponds to surface influence from across the northeast and central Amazon Basin which is a strong modifier of the background CO₂ signal through evergreen broadleaf uptake (Figs. 2a, S4, S6). At ALF, however, the Amazon Basin has significantly less influence on the air masses; savanna and woody savanna ecosystems outside the Basin dominate the surface influences on the advected background air
- 525 (Figs. 2a, Figs. S5, S7). As indicated by the IQR of NEE in Table 2, savannas and woody savannas are weaker modifiers of the CO₂ signal relative to Amazon Basin evergreen broadleaf classes. In addition, during the first quarter (Q1; January–March) of the calendar year, the typical RBA influence region (Fig. S4) is ~14% deforested, unlike the ALF Q1 influence region (Fig. S5) which is ~23% deforested during the same period (Gatti et al., 2021). The footprints from the four OCO-2 receptors used to simulate ALF vertical profiles, and the back trajectories from measured profiling locations, indicate highest ALF influence
- 530 from the most heavily deforested regions (Fig. 2a, Figs. S4-S7). As previously noted, numerous upwind fire locations potentially impact ALF at all altitudes such that the biosphere models alone cannot sufficiently approximate the observed concentrations at any altitude. Table 4 summarizes these results as differences between measured CO₂ vertical profiles and the modelled background. At altitudes <2000 masl, ALF Q1 upwind surfaces result in lower background CO₂ signal modification relative to RBA Q1 upwind surfaces, due to a combination of lower ecosystem carbon flux modification and higher fire
- 535 emissions. At altitudes >2000 masl, the differences between ALF and RBA measurements are not significant, suggesting similar background influences above that altitude. At altitudes from ~900–1000 masl, CO/CO₂ ratios from Gatti et al. (2021) indicate fire plume influence dominating the signal at RBA and ALF.

The vertically resolved footprints in Figs. 7 and 8 and Table 4 suggest four distinct upwind CO₂ regimes influencing the RBA and ALF measured profiles: (1) biosphere flux dominance below 900 masl; (2) fire emission dominance between 900–

540 1000 masl (3) dilution of fire emissions by background air between 1000–2000 masl; and (4) background tropical marine Atlantic air dominance above 2000 masl. We note Gatti et al. (2021) surface influence regions are based on integrated back trajectories ranging from 300–3500 masl; while their upper threshold of 3500 masl defines tropical marine Atlantic background

air, their results were robust across background air designations from >1300 masl (typical regional planetary boundary layer height) to 3500 masl.

- 545 At RBA where the Amazon Basin biosphere dominates the signal more than at ALF, the optimization reduces the modelobservation mismatch for all flux models, particularly below 1000 masl. Generally, the traditional VPRM formulations are significantly more impacted by the optimization than the SIF-based VPRM and SiB4 models. The magnitude at which VPRM TRA and VPRM TRG underestimate CO₂ uptake across the RBA area of influence results in a model-observation a priori surface residual of over 10 ppm and a posterior surface residual of approximately 5 ppm (Fig. 7a-b). In contrast,
- VPRM SIF, VPRM SIFg, and SiB4 models are all similarly impacted by the optimization with a priori model-observation 550 surface residual of approximately 5 ppm, and a posterior model-observation surface residual of <2.5 ppm. At ALF, where the February-March 2016 upwind air masses tend to bypass a majority of the Amazon Basin and are potentially significantly influenced by fire activity at all altitudes, all prior and posterior flux models perform similarly with a typical model-observation residual of -3 to 3 ppm throughout the vertical column (Fig. 8a-e).

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Table 4. Difference between average measured CO₂ profiles and CT2019 background CO₂. Differences are based on averages pooled across February and March 2016. *1000 masl is approximate location of the peak of a fire plume.

| | CO _{2,OBS} – CO _{2,BG} (95% CI), ppm | | | |
|-----------|--|----------------------|----------------------|---------------------|
| | <900 masl | ~900–1000 masl* | 1200-<2000 masl | >2000 masl |
| RBA | -3.2 (-3.5, -2.9) | -0.36 (-0.80, 0.084) | -0.46 (-0.72, -0.17) | -1.1 (-1.3, -0.97) |
| ALF | -1.7 (-2.0, -1.3) | 2.5 (2.0, 3.0) | -0.97 (-1.4, -0.56) | -0.99 (-1.2, -0.82) |
| ALF - RBA | 1.6 (1.1, 2.1) | 2.8 (2.2, 3.5) | -0.52 (-1.0, -0.03) | -0.15 (-0.39, 0.10) |

3.3 Summary of VPRM SIFg Prior Model Decadal Performance

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Based on comparison with eddy flux data and results from the March 2016 optimization case study, the VPRM SIFg biogenic flux model shows promise in its ability to capture Amazonian carbon fluxes across multiple timescale and moisture regimes, suggesting its suitability for larger studies evaluating interannual and seasonal carbon trends in both the fire and biogenic components of the region's NBE. In this section, we summarize the decadal performance of the VPRM SIFg prior relative to VPRM TRG, VPRM TRA and SiB4 priors. We calculate average seasonal fluxes of the VPRM SIFg prior (as µmol CO₂ m⁻² s⁻²) from 2010–2020 (2010–2018 for SiB4) over Amazonia and compare with the corresponding average fluxes from the VPRM TRA, VPRM TRG, and SiB4 priors (Fig. 9).

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3.3.1 Comparison with Traditional VPRM Fluxes

Basin-wide, and across both seasons and all years, VPRM SIFg estimates more photosynthetic uptake than VPRM TRA and VPRM TRG in an overall pattern consistent with model validation and wet season 2016 optimization results. Across both

570 seasons, VPRM-SIFg estimates more respiration in the Br-K34 calibration region, and less respiration across the rest of the basin when compared to VPRM-TRA and VPRM-TRG.

Outside of the Basin, VPRM-SIFg universally estimates higher photosynthetic uptake and respiration release than the traditional VPRM models.

575 3.3.2 Comparison with SiB4 fluxes

When compared to SiB4, VPRM-SIFg estimates less dry season uptake and respiration in the western Amazon Basin corresponding to the regions calibrated by Br-K34. In the wet season, VPRM-SIFg generally estimates less uptake and respiration than SiB4.

Outside of the Basin, VPRM-SIFg generally estimates more photosynthesis and respiration in the dry season. In the wet season, higher photosynthesis and respiration relative to SiB4 is localized to woody savanna/savanna ecosystems (Br-CST, Br-BAN).

4 Conclusions

Compounded by the impacts of global climate change, the Amazon is experiencing unprecedented ecological disturbance through fires, deforestation, drought, and forest fragmentation. Multiple recent studies describe how the increasingly degraded

- 585 and disturbed Amazon has a lowered carbon sink capacity with significant impacts to regional and global carbon budgets. However, reliable biogenic flux models that can capture fluxes from hourly to annual timescales are needed to quantify trends in carbon sink strength; ecological health and recovery from disturbance; and estimates of emissions from increasingly prevalent fires.
- Assimilating observational CO₂ data from eddy flux sites, OCO-2 columns, and aircraft vertical profiles, we demonstrate 590 construct and evaluate the ability of four versions of the VPRM diagnostic light use efficiency model to capture biogenic carbon fluxes from hourly to seasonal scales. Of the VPRM versions evaluated, the respiration-modified VPRM_SIFg exhibits the least bias when compared to eddy flux observations, including observationally derived respiration estimates. In the tropics—where respiration exerts stronger controls on NBE than the extratropics—our work demonstrates the superior performance of the respiration-modified VPRM_SIFg diagnostic light use efficiency model relative to all other VPRM
- 595 versions. In addition, optimization results for March 2016—the middle of the wet season corresponding to the tail of the 2015–2016 severe El Niño—demonstrate (1) significant underestimate of net uptake by the traditional VPRM formulations relative to observations and VPRM-SIFg; (2) relative to the SiB4 model, VPRM_SIFg model describes more spatial heterogeneity in carbon exchange throughout the Amazon; and (3) convergence of flux estimates from two distinct methodologies namely, the diagnostic VPRM-SIFg and the process-based bottom-up SiB4 biogenic flux model. While the convergence of NEE in the
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(as GPP and Reco) warrant further exploration.

We also find that, despite the paucity of wet-season OCO-2 observations used in top-down constraints, modelled vertical profiles optimized with OCO-2 measurements compare significantly better with aircraft measurements than their unoptimized

distinct VPRM-SIFg and SiB4 flux estimation methodologies lends confidence to both models, differences in NEE partitioning

counterparts. We hypothesize that, given the drier than normal conditions experienced by the region influencing RBA during
 this time period, the sounding density during this wet season was higher than normal. Future work focusing on a range of wet
 season conditions will shed light on this possible anomaly.

The promising performance of the both the prior and posterior VPRM_SIFg model in the wet season 2016 case study provides confidence in its ability to capture interannual and seasonal trends in the biogenic carbon component of the Amazon's net biome exchange. We note that VPRM-SIFg would benefit from additional calibration sites, via inclusion of eddy flux data

610 from more recent time periods and/or through greater representation of interior primary moist evergreen broadleaf classes. Currently, Br-K34 is the only site representing the interior Amazon; additional eddy flux data for calibration and/or validation such as from Amazon Tall Tower Observatory (ATTO) would be beneficial. Future work will use the optimized VPRM-SIFg across both wet and dry seasons to evaluate inter-annual and seasonal trends, including assessing the changing role of fires in the net biome exchange of Amazonian carbon fluxes, and the changes in the pace of post-fire ecosystem recovery.

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Code availability

VPRM and inversion code are available upon request from the authors.

Data availability

CO2 vertical profiling data are available from https://doi.org/10.1594/PANGAEA.926834. VPRM fluxes, STILT footprints,
 and CT2019 background data for the March 2016 case study are available from https://doi.org/10.7910/DVN/PJ1EVC. Additional data available from authors upon request.

Supplement

The supplement related to this article is available online.

Author contributions

625 AD wrote the paper with input from all co-authors. AD constructed the VPRM. MM generated the WRF-STILT fields and performed the inversion with assistance from BR and JBM. LVG provided access to vertical profiling data sets and shapefiles.

Competing Interests

The contact author has declared that none of the authors has any competing interests.

Disclaimer

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References

650

- 640 Alden, C. B., Miller, J. B., Gatti, L. V., Gloor, M. M., Guan, K., Michalak, A. M., van der Laan-Luijkx, I. T., Touma, D., Andrews, A., Basso, L. S., Correia, C. S. C., Domingues, L. G., Joiner, J., Krol, M. C., Lyapustin, A. I., Peters, W., Shiga, Y. P., Thoning, K., van der Velde, I. R., van Leeuwen, T. T., Yadav, V., and Diffenbaugh, N. S.: Regional atmospheric CO₂ inversion reveals seasonal and geographic differences in Amazon net biome exchange, Glob. Change Biol., 22, 3427–3443, https://doi.org/10.1111/gcb.13305, 2016.
- 645 Almeida, C. T. de, Delgado, R. C., Galvão, L. S., Aragão, L. E. de O. C. e, and Ramos, M. C.: Improvements of the MODIS Gross Primary Productivity model based on a comprehensive uncertainty assessment over the Brazilian Amazonia, ISPRS Journal of Photogrammetry and Remote Sensing, 145, 268–283, https://doi.org/10.1016/j.isprsjprs.2018.07.016, 2018.

Andreae, M. O.: Emission of trace gases and aerosols from biomass burning—an updated assessment. Atmos. Chem.Phys., 19(13), 8523–8546, https://doi.org/10.5194/acp-19-8523-2019, 2019.

- Antonino, A.: AmeriFlux FLUXNET-1F BR-CST Caatinga Serra Talhada, Ver. 3-5, AmeriFlux AMP, (Dataset). https://doi.org/10.17190/AMF/1902820, 2022.
- Aragão, L. E. O. C., Anderson, L. O., Fonseca, M. G., Rosan, T. M., Vedovato, L. B., Wagner, F. H., Silva, C. V. J., Silva Junior, C. H. L., Arai, E., Aguiar, A. P., Barlow, J., Berenguer, E., Deeter, M. N., Domingues, L. G., Gatti, L., Gloor,
- 655 M., Malhi, Y., Marengo, J. A., Miller, J. B., Phillips, O. L., and Saatchi, S.: 21st Century drought-related fires counteract the decline of Amazon deforestation carbon emissions, Nature Communications, 9(1), 536, https://doi.org/10.1038/s41467-017-02771-y, 2018.
 - Barkhordarian, A., Saatchi, S. S., Behrangi, A., Loikith, P., and Mechoso, C.: A recent systematic increase in vapor pressure deficit over tropical South America, Sci. Rep., 9, 15331, https://doi.org/10.1038/s41598-019-51857-8, 2019.
- 660 Basso, L. S., Wilson, C., Chipperfield, M. P., Tejada, G., Cassol, H. L. G., Arai, E., Williams, M., Smallman, T. L., Peters, W., Naus, S., Miller, J. B., and Gloor, M.: Atmospheric CO₂ inversion reveals the Amazon as a minor carbon source caused by fire emissions, with forest uptake offsetting about half of these emissions, Atmos. Chem. Phys., 23(17), https://doi.org/10.5194/acp-23-9685-2023, 2023.

Bloom, A. A., Lauvaux, T., Worden, J., Yadav, V., Duren, R., Sander, S. P., and Schimel, D. S.: What are the greenhouse gas

- observing system requirements for reducing fundamental biogeochemical process uncertainty? Amazon wetland CH4
 emissions as a case study, Atmos. Chem. Phys., 16(23), 15199–15218, https://doi.org/10.5194/acp-16-15199-2016, 2016.
- Bowman, K. W., Liu, J., Bloom, A. A., Parazoo, N. C., Lee, M., Jiang, Z., Menemenlis, D., Gierach, M. M., Collatz, G. J.,
 Gurney, K. R., and Wunch, D.: Global and Brazilian carbon response to El Niño Modoki 2011–2010, Earth and Space
 Science, 4(10), 637–660, https://doi.org/10.1002/2016EA000204, 2017.

675

690

- Brando, P., Macedo, M., Silvério, D., Rattis, L., Paolucci, L., Alencar, A., Coe, M., and Amorim, C.: Amazon wildfires: Scenes from a foreseeable disaster, Flora, 268, 151609, https://doi.org/10.1016/j.flora.2020.151609, 2020.
- Clark, D. A., Clark, D. B., and Oberbauer. S. F.: Field-quantified responses of tropical rainforest aboveground productivity to increasing CO₂ and climatic stress, 1997–2009, J. Geophys. Res. Biogeosci., 118, 783–794, https://doi.org/10.1002/jgrg.20067, 2013.
- Cox, P., Pearson, D., Booth, B., Friedlingstein, P., Huntingford, C., Jones, C., and Luke, C.: Sensitivity of tropical carbon to climate change constrained by carbon dioxide variability, Nature, 494, https://doi.org/10.1038/nature11882, 2013.
- Cox, P., Betts, R., Jones, C., Spall, S., and Totterdell, I.: Acceleration of global warming due to carbon-cycle feedbacks in a coupled climate model, Nature, 408, 184–187, https://doi.org/10.1038/35041539, 2000.
- 680 Dayalu, A., Munger, J. W., Wofsy, S. C., Wang, Y., Nehrkorn, T., Zhao, Y., McElroy, M. B., Nielsen, C. P., and Luus, K.: Assessing biotic contributions to CO₂ fluxes in northern China using the Vegetation, Photosynthesis and Respiration Model (VPRM-China) and observations from 2005 to 2009, Biogeosciences, 15(21), 6713–6729, https://doi.org/10.5194/bg-15-6713-2018, 2018.
 - Doughty, R., Köhler, P., Frankenberg, C., Magney, T. S., Xiao, X., Qin, Y., Wu, X., and Moore, B: TROPOMI reveals dry-
- season increase of solar-induced chlorophyll fluorescence in the Amazon forest, Proc. Natl. Acad. Sci., 116(44), 22393–
 8, https://doi.org/10.1073/pnas.1908157116 2019, 2019.
 - Eldering, A., Wennberg, P. O., Crisp, D., Schimel, D. S., Gunson, M. R., Chatterjee, A., Liu, J., Schwandner, F. M., Sun, Y., O'Dell, C. W., Frankenberg, C., Taylor, T., Fisher, B., Osterman, G. B., Wunch, D., Hakkarainen, J., Tamminen, J., and Weir, B.: The Orbiting Carbon Observatory-2 early science investigations of regional carbon dioxide fluxes, Science, 358(6360), eaam5745, https://doi.org/10.1126/science.aam5745, 2017.
 - Figueira, A. M. e S., Miller, S. D., de Sousa, C. A. D., Menton, M. C., Maia, A. R., da Rocha, H. R., and Goulden, M. L.: Effects of selective logging on tropical forest tree growth, J. Geophys. Res., 113, G00B05, https://doi.org/10.1029/2007JG000577, 2008.
- Fu, R., Yin, L., Li, W., Arias, P. A., Dickinson, R. E., Huang, L., Chakraborty, S., Fernandes, K., Liebmann, B., Fisher, R.,
 and Myneni, R. B.: Increased dry-season length over southern Amazonia in recent decades and its implication for future climate projection, Proc. Natl. Acad. Sci., 110(45), 18110–18115, https://doi.org/10.1073/pnas.1302584110, 2013.
 - Gallup, S. M., Baker, I. T., Gallup, J. L., Restrepo-Coupe, N., Haynes, K. D., Geyer, N. M., and Denning, A. S.: Accurate simulation of both sensitivity and variability for Amazonian photosynthesis: Is it too much to ask? Journal of Advances in Modeling Earth Systems, 13, e2021MS002555, https://doi.org/10.1029/2021MS002555, 2021.
- 700 Gatti, L. V., Cunha, C. L., Marani, L., Cassol, H. L. G., Messias, C. G, Arai, E., Denning, A. S., Soler, L. S., Almeida, C., Setzer, A., Domingues, L. G., Basso, L. S., Miller, J. B., Gloor, M., Correia, C. S. C., Tejada, G., Neves, R. A. L., Rajao, R., Nunes, F., Filho, B. S. S., Schmitt, J., Nobre, C., Corrêa, S. M., Sanches, A. H., Aragão, L. E. O. C., Anderson, L., Von Randow, C., Crispim, S. P., Silva, F. M., and Machado, G. B. M.: Increased Amazon carbon emissions mainly from

705

decline in law enforcement, Nature, 621(7978), 318–323, https://doi.org/10.1038/s41586-023-06390-0. Epub 2023 Aug 23. PMID: 37612502, 2023.

- Gatti, L. V., Basso, L. S., Miller, J. B., Gloor, M., Gatti Domingues, L., Cassol, H. L. G., Tejada, G., Aragão, L. E. O. C., Nobre, C., Peters, W., Marani, L., Arai, E., Sanches, A. H., Corrêa, S. M., Anderson, L., Von Randow, C., Correia, C. S. C., Crispim, S. P., and Neves, R. A. L.: Amazonia as a carbon source linked to deforestation and climate change, Nature, 595(7867), 388–393, https://doi.org/10.1038/s41586-021-03629-6, 2021.
- Gatti, L. V., Gloor, M., Miller, J. B., Doughty, C. E., Malhi, Y., Domingues, L. G., Basso, L. S., Martinewski, A., Correia, C. S. C., Borges, V. F., Freitas, S., Braz, R., Anderson, L. O., Rocha, H., Grace, J., Phillips, O. L., and Lloyd, J.: Drought sensitivity of Amazonian carbon balance revealed by atmospheric measurements, Nature, 506(7486), 76–80, https://doi.org/10.1038/nature12957, 2014.
- Gatti, L. V., Miller, J. B., D'Amelio, M. T. S., Martinewski, A., Basso, L. S., Gloor, M. E., Wofsy, S., and Tans, P.: Vertical
 profiles of CO₂ above eastern Amazonia suggest a net carbon flux to the atmosphere and balanced biosphere between

2000 and 2009, Tellus B, 62, 581–594, https://doi.org/10.1111/j.1600-0889.2010.00484.x, 2010.

- Gloor, M., Gatti, L., Brienen, R., Feldpausch, T. R., Phillips, O. L., Miller, J., Ometto, J. P., Rocha, H., Baker, T., de Jong, B., Houghton, R. A., Malhi, Y., Aragão, L. E. O. C., Guyot, J.-L., Zhao, K., Jackson, R., Peylin, P., Sitch, S., Poulter, B., Lomas, M., Zaehle, S., Huntingford, C., Levy, P., and Lloyd, J.: The carbon balance of South America: a review of the status, decadal trends and main determinants, Biogeosciences, 9, 5407–5430, https://doi.org/10.5194/bg-9-5407-2012,
- status, decadal trends and main determinants, Biogeosciences, 9, 5407–5430, https://doi.org/10.5194/bg-9-5407-2012, 2012.
 - Goulden, M.: AmeriFlux BASE BR-Sa3 Santarem-Km83-Logged Forest, Ver. 3-5, AmeriFlux AMP, (Dataset). https://doi.org/10.17190/AMF/1245995, 2019.
 - Gourdji, S. M., Karion, A., Lopez-Coto, I., Ghosh, S., Mueller, K. L., Zhou, Y., Williams, C. A., Baker, I. T., Haynes, K. D.,
- and Whetstone, J. R.: A modified Vegetation Photosynthesis and Respiration Model (VPRM) for the Eastern USA and Canada, evaluated with comparison to atmospheric observations and other biospheric models, J. Geophys. Res. Biogeosciences, 127(1), https://doi.org/10.1029/2021JG006290, 2022.
 - Hayek, M. N., Longo, M., Wu, J., Smith, M. N., Restrepo-Coupe, N., Tapajós, R., da Silva, R., Fitzjarrald, D. R., Camargo,P. B., Hutyra, L. R., Alves, L. F., Daube, B., Munger, J. W., Wiedemann, K. T., Saleska, S. R., and Wofsy, S. C.: Carbon
- exchange in an Amazon Forest: from hours to years, Biogeosciences, 15(15), 4833–4848, https://doi.org/10.5194/bg-15-4833-2018, 2018.
 - Haynes, K., Baker, I., and Denning, S.: Simple Biosphere Model version 4.2 (SiB4) technical description. Mountain Scholar, Colorado State University, Fort Collins, CO, USA, https://hdl.handle.net/10217/200691, 2020.
- Haynes, K. D., I. T. Baker, and A. S. Denning: SiB4 Modeled Global 0.5-Degree Monthly Carbon Fluxes and Pools, 2000–
 2018. ORNL DAAC, Oak Ridge, Tennessee, USA, https://doi.org/10.3334/ORNLDAAC/1848, 2021.
 - Haynes, K. D., Baker, I. T., Denning, A. S., Stöckli, R., Schaefer, K., Lokupitiya, E. Y., and Haynes, J. M.: Representing grasslands using dynamic prognostic phenology based on biological growth stages: 1. Implementation in the Simple

750

770

Biosphere Model (SiB4), Journal of Advances in Modeling Earth Systems, 11(12), 4423–4439, https://doi.org/10.1029/2018MS001540, 2019.

- 740 Hilton, T. W., Davis, K. J., Keller, K., and Urban, N. M.: Improving North American terrestrial CO₂ flux diagnosis using spatial structure in land surface model residuals, Biogeosciences, 10(7), 4607–4625, https://doi.org/10.5194/bg-10-4607-2013, 2013.
- Hu, L., Andrews, A. E., Thoning, K. W., Sweeney, C., Miller, J. B., Michalak, A. M., Dlugokencky, E., Tans, P. P., Shiga, Y. P., Mountain, M., Nehrkorn, T., Montzka, S. A., McKain, K., Kofler, J., Trudeau, M., Michel, S. E., Biraud, S. C.,
 Fischer, M. L., Worthy, D. E., Vaughn, B. H., White, J. W., Yadav, V., Basu, S., and Van Der Velde, I. R.: Enhanced North American carbon uptake associated with El Niño, Science Advances, 5, https://doi.org/10.1126/sciadv.aaw0076, 2019.
 - Huete, A. R., Didan, K., Shimabukuro, Y. E., Ratana, P., Saleska, S. R., Hutyra, L. R., Yang, W., Nemani, R. R., and Myneni,
 R.: Amazon rainforests green-up with sunlight in dry season, Geophys. Res. Lett., 33(6), L06405, https://doi.org/10.1029/2005GL025583, 2006.
 - Jacobson, A. R., Schuldt, K. N., Miller, J. B., Oda, T., Tans, P., Arlyn Andrews, Mund, J., Ott, L., Collatz, G. J., Aalto, T., Afshar, S., Aikin, K., Aoki, S., Apadula, F., Baier, B., Bergamaschi, P., Beyersdorf, A., Biraud, S. C., Bollenbacher, A., Miroslaw Zimnoch. (2020). *CarbonTracker CT2019B*. NOAA Global Monitoring Laboratory. https://doi.org/10.25925/20201008
- Jiménez-Muñoz, J., Mattar, C., Barichivich, J. Santamaría-Artigas, A. Takahashi, K., Malhi, Y., Sobrino, J.A., and van der Schrier, G: Record-breaking warming and extreme drought in the Amazon rainforest during the course of El Niño 2015– Sci Rep 6, 33130 (2016). <u>https://doi.org/10.1038/srep33130</u>, 2016.
 - Koren, G., van Schaik, E., Araújo, A. C., Boersma, K. F., Gärtner, A., Killaars, L., Kooreman, M. L., Kruijt, B., van der Laan-Luijkx, I. T., von Randow, C., Smith, N. E., and Peters, W.: Widespread reduction in sun-induced fluorescence from the
- Amazon during the 2015/2016 El Niño, Philosophical Transactions of the Royal Society B: Biological Sciences, 373(1760), 20170408, https://doi.org/10.1098/rstb.2017.0408, 2018.
 - Kruid, S., Macedo, M. N., Gorelik, S. R., Walker, W., Moutinho, P., Brando, P. M., Castanho, A., Alencar, A., Baccini, A., and Coe, M. T.: Beyond deforestation: carbon emissions from land grabbing and forest degradation in the Brazilian Amazon, Frontiers in Forests and Global Change, 4, 645282, https://doi.org/10.3389/ffgc.2021.645282, 2021.
- 765 Liu, J., Wunch, D., Menemenlis, D., Frankenberg, C., Sun, Y., Bloom, A. A., Lee, M., Bowman, K. W., Schimel, D. S., Crisp, D., Gurney, K. R., Eldering, A., Jiang, Z., O'Dell, C. W., Gierach, M., and Parazoo, N. C.: Contrasting carbon cycle responses of the tropical continents to the 2015–2016 El Niño, Science, 358(6360), eaam5690, https://doi.org/10.1126/science.aam5690, 2017.

Luus, K. A., Commane, R., Parazoo, N. C., Benmergui, J., Euskirchen, E. S., Frankenberg, C., Joiner, J., Lindaas, J., Miller, C. E., Oechel, W. C., Zona, D., Wofsy, S., and Lin, J. C.: Tundra photosynthesis captured by satellite-observed solar-

775

induced chlorophyll fluorescence, Geophys. Res. Lett., 44(3), 1564–1573, https://doi.org/10.1002/2016GL070842, 2017.

- Ma, S., Osuna, J. L., Verfaillie, J., and Baldocchi, D. D.: Photosynthetic responses to temperature across leaf-canopyecosystem scales: a 15-year study in a Californian oak-grass savanna, Photosynthesis Research, 132(3), 277–291, https://doi.org/10.1007/s11120-017-0388-5, 2017.
- Mahadevan, P., Wofsy, S. C., Matross, D. M., Xiao, X., Dunn, A. L., Lin, J. C., Gerbig, C., Munger, J. W., Chow, V. Y., and Gottlieb, E. W.: A satellite-based biosphere parameterization for net ecosystem CO₂ exchange: Vegetation Photosynthesis and Respiration Model (VPRM), Global Biogeochemical Cycles, 22(2), https://doi.org/10.1029/2006GB002735, 2008.
- 780 Martin, S. T., Artaxo, P., Machado, L. A. T., Manzi, A. O., Souza, R. A. F., Schumacher, C., Wang, J., Andreae, M. O., Barbosa, H. M. J., Fan, J., Fisch, G., Goldstein, A. H., Guenther, A., Jimenez, J. L., Pöschl, U., Silva Dias, M. A., Smith, J. N., and Wendisch, M.: Introduction: observations and modeling of the Green Ocean Amazon (GoAmazon2014/5), Atmos. Chem. Phys. Discuss., 15(21), 30175–30210, https://doi.org/10.5194/acpd-15-30175-2015, 2015.
- Matross, D. M., Andrews, A., Pathmathevan, M., Gerbig, C., Lin, J. C., Wofsy, S. C., Daube, B. C., Gottlieb, E. W., Chow,
 V. Y., Lee, J. T., Zhao, C., Bakwin, P. S., Munger, J. W., and Hollinger, D. Y.: Estimating regional carbon exchange in
 New England and Quebec by combining atmospheric, ground-based and satellite data, Tellus B: Chemical and Physical
 Meteorology, 58(5), 344–358, https://doi.org/10.1111/j.1600-0889.2006.00206.x, 2006.
 - Mohammadi, K., Jiang, Y., and Wang, G., Flash drought early warning based on the trajectory of solar-induced chlorophyll fluorescence, Proc. Nat. Acad. Sci., 119(32), https://doi.org/10.1073/pnas.2202767119, 2022.
- Naus, S., Domingues, L. G., Krol, M., Luijkx, I. T., Gatti, L. V., Miller, J. B., Gloor, E., Basu, S., Correia, C., Koren, G., Worden, H. M., Flemming, J., Pétron, G., and Peters, W.: Sixteen years of MOPITT satellite data strongly constrain Amazon CO fire emissions, Atmos. Chem. Phys., 22(22), 14735–14750, https://doi.org/10.5194/acp-22-14735-2022, 2022.
- O'Dell, C. W., Connor, B., Bösch, H., O'Brien, D., Frankenberg, C., Castano, R., Christi, M., Eldering, D., Fisher, B., Gunson,
 M., McDuffie, J., Miller, C. E., Natraj, V., Oyafuso, F., Polonsky, I., Smyth, M., Taylor, T., Toon, G. C., Wennberg, P.
 O., and Wunch, D.: The ACOS CO₂ retrieval algorithm–Part 1: Description and evaluation against synthetic observations, Atmos. Meas. Tech., 5(1), 99–121, https://doi.org/10.5194/amt-5-99-2012, 2012.
 - Paca, V. H. d. M., Espinoza-Dávalos, G. E., da Silva, R., Tapajós, R., and dos Santos Gaspar, A. B: Remote sensing products validated by flux tower data in Amazon Rain Forest, Remote Sensing, 14(5), 1259, https://doi.org/10.3390/rs14051259, 2022.
- 800 2
 - Peiro, H., Crowell, S., Schuh, A., Baker, D. F., O'Dell, C., Jacobson, A. R., Chevallier, F., Liu, J., Eldering, A., Crisp, D., Deng, F., Weir, B., Basu, S., Johnson, M. S., Philip, S., and Baker, I., Four years of global carbon cycle observed from the Orbiting Carbon Observatory 2 (OCO-2) version 9 and in situ data and comparison to OCO-2 version 7, Atmos. Chem. Phys., 22(2), 1097–1130, https://doi.org/10.5194/acp-22-1097-2022, 2022.

- 805 Phillips, O. L., and Brienen, R. J. W.: Carbon uptake by mature Amazon forests has mitigated Amazon nations' carbon emissions, Carbon Balance and Management, 12(1), 1, https://doi.org/10.1186/s13021-016-0069-2, 2017.
 - Rastogi, B., Miller, J. B., Trudeau, M., Andrews, A. E., Hu, L., Mountain, M., Nehrkorn, T., Baier, B., McKain, K., Mund, J., Guan, K., and Alden, C. B.: Evaluating consistency between total column CO₂ retrievals from OCO-2 and the in situ network over North America: implications for carbon flux estimation, Atmos. Chem. Phys., 21(18), 14385–14401, https://doi.org/10.5194/acp-21-14385-2021, 2021.

810 https://doi.org/10.5194/acp-21-14385-2021, 2021.

- Rastogi, B., Miller, J., Trudeau, M., Andrews, A., Hu, L., Mountain, M., Nehrkorn, T., Baier, B., Mund, J. and Alden, C., (2021b), December. Constraints on OCO-2 retrievals to provide robust NEE estimates over North America. In AGU Fall Meeting Abstracts (Vol. 2021, pp. A12E-01).
- Restrepo-Coupe, N., da Rocha, H. R., Hutyra, L. R., de Araujo, A. C., Borma, L. S., Christoffersen, B., Cabral, O., de Camargo,
- P. B., Cardoso, F. L., Costa, A. C. L., Fitzjarrald, D. R., Goulden, M. L., Kruijt, B., Maia, J. M. F., Malhi, Y. S., Manzi, A. O., Miller, S. D., Nobre, A. D., von Randow, C., Abreu Safaj, L. D., Sakai, R. K., Tota, J., Wofsy, S. C., Zanchi, F. B., and Saleska, S. R.: LBA-ECO CD-32 Flux Tower Network Data Compilation, Brazilian Amazon: 1999-2006, V2. ORNL DAAC, Oak Ridge, Tennessee, USA, https://doi.org/10.3334/ORNLDAAC/1842, 2021.
- Saleska, S.: AmeriFlux BASE BR-Sa1 Santarem-Km67-Primary Forest, Ver. 5-5, AmeriFlux AMP, (Dataset), https://doi.org/10.17190/AMF/1245994, 2019.
 - Saleska, S. R., Wu, J., Guan, K., Araujo, A. C., Huete, A., Nobre, A. D., and Restrepo-Coupe, N.: Dry-season greening of Amazon forests, Nature, 531(7594), E4–E5, https://doi.org/10.1038/nature16457, 2016.
 - Saleska, S. R., Miller, S. D., Matross, D. M., Goulden, M. L., Wofsy, S. C., da Rocha, H. R., de Camargo, P. B., Crill, P., Daube, B. C., de Freitas, H. C., Hutyra, L., Keller, M., Kirchhoff, V., Menton, M., Munger, J. W., Pyle, E. H., Rice, A.
- H., and Silva, H., Carbon in Amazon Forests: unexpected seasonal fluxes and disturbance-induced losses, Science, 302(5650), 1554–1557, https://doi.org/10.1126/science.1091165, 2003.
 - Silva Junior, C. H. L., Anderson, L. O., Silva, A. L., Almeida, C. T., Dalagnol, R., Pletsch, M. A. J. S., Penha, T. V., Paloschi, R. A., and Aragão, L. E. O. C.: Corrigendum: fire responses to the 2010 and 2015/2016 Amazonian droughts, Frontiers in Earth Science, 7:97, https://doi.org/10.3389/feart.2019.00160, 2019.
- 830 Slot, M., and Winter, K.: In situ temperature response of photosynthesis of 42 tree and liana species in the canopy of two Panamanian lowland tropical forests with contrasting rainfall regimes, New Phytologist, 214(3), 1103–1117, https://doi.org/10.1111/nph.14469, 2017.
 - Tan, Z.-H., Zeng, J., Zhang, Y.-J., Slot, M., Gamo, M., Hirano, T., Kosugi, Y., da Rocha, H. R., Saleska, S. R., Goulden, M.L., Wofsy, S. C., Miller, S. D., Manzi, A. O., Nobre, A. D., de Camargo, P. B., and Restrepo-Coupe, N.: Optimum air
- temperature for tropical forest photosynthesis: mechanisms involved and implications for climate warming, Env. Res. Lett., 12(5), 054022, https://doi.org/10.1088/1748-9326/aa6f97, 2017.
 - TerraBrasilis: http://terrabrasilis.dpi.inpe.br/queimadas/bdqueimadas/#mapa. Data from Aqua Satellite February-March 2016. Accessed 5 March 2024, 2024.

840

860

- Trumbore, S., Brando, P., and Hartmann, H.: Forest health and global change, Science, 349(6250), 814–818, https://doi.org/10.1126/science.aac6759, 2015.
- Turner, A. J., Köhler, P., Magney, T. S., Frankenberg, C., Fung, I., and Cohen, R. C.: A double peak in the seasonality of California's photosynthesis as observed from space, Biogeosciences, 17(2), 405–422, https://doi.org/10.5194/bg-17-405-2020, 2020.
- Uhl, C., Kauffman, J. B., and Cummings, D. L.: Fire in the Venezuelan Amazon 2: Environmental Conditions Necessary for
- Forest Fires in the Evergreen Rainforest of Venezuela. Oikos, *53*(2), 176–184. <u>https://doi.org/10.2307/3566060</u>, 1988.
 van der Laan-Luijkx, I. T., van der Velde, I. R., Krol, M. C., Gatti, L. V., Domingues, L. G., Correia, C. S. C., Miller, J. B., Gloor, M., van Leeuwen, T. T., Kaiser, J. W., Wiedinmyer, C., Basu, S., Clerbaux C., and Peters, W.: Response of the Amazon carbon balance to the 2010 drought derived with CarbonTracker South America, Global Biogeochem. Cycles, 29, 1092–1108, https://doi.org/10.1002/2014GB005082, 2015.
- 850 Winbourne, J. B., Smith, I. A., Stoynova, H., Kohler, C., Gately, C. K., Logan, B. A., Reblin, J., Reinmann, A., Allen, D. W., and Hutyra, L. R.: Quantification of urban forest and grassland carbon fluxes using field measurements and a satellitebased model in Washington DC/Baltimore area, J. Geophys. Res.: Biogeosciences, 127(1), https://doi.org/10.1029/2021JG006568, 2022.
- Wu, J., Guan, K., Hayek, M., Restrepo-Coupe, N., Wiedemann, K. T., Xu, X., Wehr, R., Christoffersen, B. O., Miao, G., da
 Silva, R., de Araujo, A. C., Oliviera, R. C., Camargo, P. B., Monson, R. K., Huete, A. R., and Saleska, S. R.: Partitioning controls on Amazon forest photosynthesis between environmental and biotic factors at hourly to interannual timescales, Glob. Chang. Biol., 23(3),1240–1257, https://doi.org.10.1111/gcb.13509, Epub 2016 Oct 11. PMID: 27644012, 2017.
 - Zhang, Y., Joiner, J., Alemohammad, S. H., Zhou, S., and Gentine, P.: A global spatially contiguous solar-induced fluorescence (CSIF) dataset using neural networks, Biogeosciences, 15(19), 5779–5800, https://doi.org/10.5194/bg-15-5779-2018, 2018a.
 - Zhang, Y., Xiao, X., Wolf, S., Wu, J., Wu, X., Gioli, B., Wohlfahrt, G., Cescatti, A., van der Tol, C., Zhou, S., Gough, C. M., Gentine, P., Zhang, Y., Steinbrecher, R., and Ardö, J.: Spatio-temporal convergence of maximum daily light-use efficiency based on radiation absorption by canopy chlorophyl, Geophys. Res. Lett., 45(8), 3508–3519, https://doi.org/10.1029/2017GL076354, 2018b.
- 865 Zhang, Z., Zhang, Y., Zhang, Y., Gobron, N., Frankenberg, C., Wang, S., and Li, Z.: The potential of satellite FPAR product for GPP estimation: An indirect evaluation using solar-induced chlorophyll fluorescence, Rem. Sens. Environ., 111686, https://doi.org/10.1016/j.rse.2020.111686, 2020.

Figure 1. Study spatial domain with OCO-2 and aircraft observation locations. (a) WRF domain. The black diamonds indicate the semimonthly aircraft vertical profiling areas available for evaluation over the study period. Bounding boxes represent the VPRM (green, 12 km native resolution) and WRF meteorology domain nests (black: d01, 36km resolution; blue: d02, 12km resolution). The orange dashed line approximates the Arc of Deforestation. OCO-2 column measurements (nadir and glint land soundings, March 2016), thinned to 2s along-track, are provided as black points. (b) Vertically integrated footprints averaged for March 2016 optimization case study using 14 levels of WRF-STILT footprints within WRF d02 domain.

Figure 2. IGBP land use categories in domain and influences on measurements. (a) IGBP categories mapped to study domain, with eddy flux calibration/ validation sites and OCO-2 receptors overlaid. Vertical profiling sites RBA and ALF, their representative OCO-2 receptors, and their upwing influences are highlighted. (b) Land-use percentages in domain.

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Figure 3. VPRM Model-Observation (Night-time) Respiration residuals, annually and by season (μ mol CO2 m⁻² s⁻¹). Observed respiration is set to nighttime NEE where GPP = 0.

Figure 4. Light Use Efficiency and ERA5 modeled PAR at Br-K67. (a) Monthly and constant LUE estimated by VPRM and Wu et al. (2017); (b) modelled PAR.

Figure 5. Seasonal fits to Br-K67 and diurnal model performance. Left panels: wet season; right panels: dry season.

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Figure 6. Correlation of March 2016 regional drought patterns with differences between average monthly prior and posterior fluxes. (a) Self calibrated Palmer Drought Severity Index (scPDSI) from January through March 2016 exhibiting anomalously wet conditions in the western Amazon and anomalously dry conditions in eastern Amazon; (b-d) spatial patterns of average March 2016 fluxes for the (b) prior models, (c) posterior models, and (d) their differences. scPDSI data from Jiminez-Muñoz et al. (2016).

Figure 7. Modelled and measured vertical profiles for a typical month during 2016 wet season at RBA aircraft site. (ae) Prior and posterior residuals of modelled vertical profiles relative profiles; measured to (f) measured CO₂. Shaded regions are 25th and 75th percentiles of bootstrapped distributions.

Figure 8. Modelled and measured vertical profiles for a typical month during 2016 wet season at ALF aircraft site. (a-e) Prior and posterior residuals of modelled vertical profiles relative to measured profiles; (f) measured CO₂. Shaded regions are 25th and 75th percentiles of bootstrapped distributions.

Figure 9. Average GPP and R_{eco} flux differences between VPRM-SIFg relative to traditional VPRM formulations and SiB4. Averaging period is 2010–2020 (comparison with VPRM formulations) and 2010-2018 (comparison with SiB4). Top Panels: Dry season differences; Bottom Panels: Wet season differences. Blue (red) values indicate instances where SIFg estimates more (less) uptake and release than the comparison model.