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Photosynthesis-dependent isoprene emission from leaf to planet in a global carbon-chemistry-climate model

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Abstract. We describe the implementation of a biochemical model of isoprene emission that depends on the electron requirement for isoprene synthesis into the Farquhar– Ball–Berry leaf model of photosynthesis and stomatal conductance that is embedded within a global chemistry-climate simulation framework. The isoprene production is calculated as a function of electron transport-limited photosynthesis, intercellular and atmospheric carbon dioxide concentration, and canopy temperature. The vegetation biophysics module computes the photosynthetic uptake of carbon dioxide coupled with the transpiration of water vapor and the isoprene emission rate at the 30 min physical integration time

step of the global chemistry-climate model. In the model, the rate of carbon assimilation provides the dominant control on isoprene emission variability over canopy temperature. A control simulation representative of the present-day climatic state that uses 8 plant functional types (PFTs), prescribed phenology and generic PFT-specific isoprene emission potentials (fraction of electrons available for isoprene synthesis) reproduces 50 % of the variability across different ecosystems and seasons in a global database of 28 measured campaign-average fluxes. Compared to time-varying isoprene flux measurements at 9 select sites, the model authentically captures the observed variability in the 30 min

average diurnal cycle (R^2 = 64–96%) and simulates the flux magnitude to within a factor of 2. The control run yields a global isoprene source strength of 451 TgC yr⁻¹ that increases by 30 % in the artificial absence of plant water stress and by 55 % for potential natural vegetation.

1 Introduction

Global terrestrial gross primary productivity (GPP) is the total amount of carbon dioxide $(CO₂)$ removed from the atmosphere by plant photosynthesis. GPP is the largest flux in the global carbon cycle and is of fundamental importance to life on Earth as the basis for food and fiber. By absorbing an increasing amount of fossil fuel $CO₂$, GPP provides a critical ecosystem service of climate protection in the Anthropocene (Ballantyne et al., 2012). Estimates of GPP center around 120 PgC yr−¹ (Beer et al., 2010) although recent isotopic analysis supports a higher value of 150–175 PgC yr−¹ (Welp et al., 2011). Land ecosystems return to the atmosphere an estimated 1 % of GPP in the form of biogenic volatile organic compounds (BVOCs) (Guenther et al., 2012). This $> 1 \text{ PgC yr}^{-1}$ chemically reactive carbon flux is of comparable magnitude to the annual global net ecosystem production and an order of magnitude larger than the annual anthropogenic VOC source. The ecological and physiological roles of BVOCs are broad and range from abiotic and biotic stress functions to integrated components of carbon metabolism (Loreto and Schnitzler, 2010; Kesselmeier and Staudt, 1999). The dominant BVOC emitted is isoprene, amounting to half of the total annual flux of reactive carbon (Guenther et al., 2012). The biogenic isoprene flux is comparable to the total flux of methane including both biogenic and anthropogenic sources. The specific roles of isoprene emission are not fully understood but may be related to protection against heat and antioxidants. For instance, isoprene stabilizes chloroplastic membranes during high temperature events, allowing the plant's photosynthetic capacity to be maintained during rapid leaf temperature fluctuations caused by sun flecks in the canopy (Sharkey and Singsaas, 1995; Behnke et al., 2010). Isoprene reduces plant damage caused by ozone and reactive oxygen species (Vickers et al., 2009).

The chemical transformation of isoprene in the atmosphere has a profound effect on the distribution and variability of the key short-lived climate forcers: tropospheric ozone (O_3) , secondary organic aerosol, and methane (CH_4) (Fiore et al., 2012). Thus, the isoprene flux is of central importance to understanding interactions between atmospheric chemistry and climate. Indeed, isoprene emission may have played an important role in the Earth system climate sensitivity in past greenhouse worlds (Beerling et al., 2011). While it is qualitatively perceived that human and/or natural perturbations to isoprene emission may provide a powerful lever on regional climate and even trigger feedbacks to global climate (Pitman et al., 2012), there is a lack of quantitative understanding of this critical ecosystem-chemistryclimate linkage. The main challenges have been the accurate model representation of the isoprene emission response to the complex, sometimes, opposing, influences of global change (Monson et al., 2007) and the development of appropriate coupled global carbon-chemistry-climate modeling frameworks.

Isoprene emission rate depends strongly upon ecosystem type with broadleaf trees and shrubs exhibiting the strongest emission potentials. Isoprene is produced in the chloroplast from precursors formed during photosynthesis. Isotopic labeling shows that about 70–90 % of isoprene production is directly linked to photosynthesis, which provides the supply of energy, reducing power and carbon skeletons for inleaf biosynthesis (Delwiche and Sharkey, 1993; Karl et al., 2002; Affek and Yakir, 2003). The remaining isoprene production is associated with an older carbon source. Any single molecule of isoprene can include precursors from both newly assimilated and the older carbon source. In the laboratory and the field, the incomplete coupling to the photosynthetic flow is revealed in 4 ways: (1) the time lag between the onset of photosynthesis and isoprene emission, which is due to the effects of the growth environment on the expression of isoprene synthase e.g. (Kuzma and Fall, 1993; Sharkey and Loreto, 1993; Fuentes et al., 1999; Monson et al., 1994; Goldstein et al., 1998; Pressley et al., 2005); (2) the higher temperature optimum of isoprene production versus photosynthesis such that at high leaf temperatures the rates of isoprene production and photosynthesis are inversely correlated; (3) the initial short-term increase in isoprene emission under drought stress e.g. (Pegoraro et al., 2004, 2005, 2006). Under sustained drought stress, the isoprene emission rate does decrease according to the reduction in photosynthetic rate. Active upregulation of isoprene production during drought stress has been posited (Monson et al., 2007; Niinemets, 2010) but there remain unresolved issues regarding the isoprene emission response to drought (4) the isoprene emission rate has an inverse relationship in response to increasing atmospheric $CO₂$ concentration (opposite to that observed for the photosynthetic $CO₂$ assimilation rate), behavior known as the " $CO₂$ -inhibition effect" (Rosenstiel et al., 2003; Possell et al., 2004, 2005; Possell and Hewitt, 2011; Monson et al., 2007). Competition for carbon substrate has been offered as a mechanistic explanation for the CO2-inhibition effect (Rosenstiel et al., 2003; Wilkinson et al., 2009). Enhanced isoprene-related tolerance of heat and light stressed photosynthesis has been found at low but not high $CO₂$ concentrations (Way et al., 2011). It has been hypothesized that isoprene biosynthesis evolved in a low $CO₂$ climate state, which prompts the important question: what is the functional benefit of leaf isoprene production in a future high $CO₂$ world?

Global isoprene emission models have been developed for application in chemistry-climate studies (Guenther et al.,

1995, 2006; Lathiére et al., 2006; Arneth et al., 2007; Pacifico et al., 2011). The most widely used approach to simulate interactive isoprene emission is to modify ecosystem-specific basal emission rates under standard conditions, either at the leaf or canopy level, with empirical functions in the form of serial multipliers that describe the observed emission response to specific environmental controls (light, temperature, soil moisture) (Guenther et al., 1991, 1995, 2006). This approach has been applied within prescribed static vegetation frameworks, for instance in the complex global canopy environment Model of Emissions of Gases and Aerosols from Nature (MEGAN) (Guenther et al., 2006); and within dynamic global vegetation model (DGVM) frameworks, for instance in the Organizing Carbon and Hydrology in Dynamic EcosystEms model (ORCHIDEE) (Lathiére et al., 2006) and in the Community Land Model (Heald et al., 2008). Similarly, the empirical functions that describe isoprene emission response to light and temperature at the leaf or canopy level have been directly embedded within global chemistryclimate and global chemistry-transport models (CCMs and CTMs) using prescribed static vegetation e.g. (Wu et al., 2008; Shindell et al., 2006; Horowitz et al., 2007). Current generation global CCMs and CTMs usually neglect the response of isoprene emission to soil moisture. Precipitation controls photosynthesis in more than 40 % of vegetated land (Beer et al., 2010). It follows that water availability must play an important role in isoprene emission that will in turn affect the atmospheric composition and climate–air pollution interactions.

Medium to long-term changes of vegetation physiology and composition in response to global change drivers (climate change, $CO₂$, land use), and increasing awareness of the complex mutual feedbacks between isoprene emission and regional climate sensitivity, make it necessary to link isoprene emission directly to the biological processes that affect emissions. To address this urgent need, a leaf-level isoprene emission model that depends on the electron requirement for isoprene synthesis (Niinemets et al., 1999) has been modified for implementation within DGVM frameworks including the Lund Potsdam Jena General Ecosystem Simulator (LPJ-GUESS) (Arneth et al., 2007), and the Joint UK Land Environmental Simulator (JULES) (Pacifico et al., 2011). Isoprene emissions generated in this way have been used in chemistry-climate modeling studies. For instance LPJ-GUESS isoprene emissions were applied off-line in the UM_CAM model to quantify the effects of the $CO₂$ inhibition on future ozone predictions (Young et al., 2009), and JULES isoprene emissions have been applied on-line in the HadGEM2 Earth-system model to examine the sensitivity of isoprene emission to past and future global change and the implications for atmospheric chemistry (Pacifico et al., 2012).

Global modeling of carbon cycle-climate and chemistryclimate and interactions have evolved as entirely separate communities because $CO₂$ is chemically unreactive in the atmosphere and because of the vast differences in the system integration time scales. There are many existing DGVMs whose contemporary global carbon cycle simulations have been extensively evaluated (Sitch et al., 2008; Schwalm et al., 2010). These models do not typically include on-line atmospheric chemistry. On the other hand, global chemistry-climate models normally rely on off-line prescribed vegetation input data sets. Our objective is to implement the biochemical leaf isoprene production scheme of (Niinemets et al., 1999) into on-line vegetation biophysics integrated within a global chemistry-climate model framework. To achieve this objective, we introduce the Yale-E2 global carbon-chemistry-climate model that is built around the NASA Goddard Institute for Space Studies (GISS) Model-E2 global climate model and features advanced biogeochemistry-climate interactions under separate development at Yale University.

The major goals of this study are (i) to describe the model and (ii) to evaluate the global scale model performance in the present climate state. The modeling methodology and isoprene emission algorithm are detailed in Sect. 2. In Sect. 3, we specify the control and sensitivity global simulations that are carried out for this work. Section 4 summarizes the global model results and presents the model evaluation including: summary of simulated global GPP and isoprene emission (Sect. 4.1), isoprene emission sensitivity to GPP and canopy temperature (Sect. 4.1.1), evaluation of global GPP and isoprene emission (Sect. 4.2), FLUXNET-derived global GPP (Sect. 4.2.1), GPP and latent heat seasonal cycle at 6 benchmark sites (Sect. 4.2.2), global database of campaign-average isoprene flux measurements (Sect. 4.2.3), time-varying isoprene emission through campaign periods (Sect. 4.2.4), isoprene and GPP diurnal cycle (Sect. 4.2.5). Discussion and conclusions are presented in Sect. 5.

2 Methodology

2.1 Yale-E2 global carbon-chemistry-climate model

Yale-E2 is built around the new generation IPCC AR5 version NASA Model-E2 global climate model (Schmidt et al., 2006) and incorporates interactive terrestrial ecosystems, a dynamic carbon cycle module, and 2-way coupling between the on-line vegetation and atmospheric chemistry. The model has flexible horizontal and vertical resolution. In this study, we apply 2° × 2.5° latitude by longitude horizontal resolution with 40 vertical layers extending to 0.1 hPa. The vegetation submodel is embedded within the general circulation model that provides the key meteorological drivers for the vegetation physiology (Friend and Kiang, 2005). The landsurface hydrology submodel provides the grid cell level soil characteristics to the vegetation physiology. The well established gas-phase chemistry and aerosol modules are fully integrated, so that these components interact with each other

PFT	SimCONT fraction $(\%)$	SimSIB2 fraction (%)	$I_{\rm s}$ (µgC g ⁻¹ h ⁻¹)	\mathcal{E}
Tundra	5.5	8.1	θ	0.000
Grass	6.4	5.4	16	0.016
Shrub	3.2	8.2	16	0.055
Savanna	4.6	16.7	16	0.036
Deciduous	10.5	7.4	45	0.061
Evergreen	7.8	7.8	8	0.015
Rainforest	9.8	9.7	24	0.027
Crop	36.9	19.1	0	0.000

Table 1. PFT-specific fractional coverage of global vegetated land area in the standard and SiB2 data sets (%) and isoprene emission parameters used in Yale-E2 global carbon-chemistry-climate model. PFT-specific photosynthesis parameters are in Table A1.

and with the physics of the climate model (Bell et al., 2005; Shindell et al., 2006, 2013; Unger, 2011).

2.1.1 Vegetation structure

The vegetation is described using 8 plant functional types (PFTs): tundra, grass, shrub, savanna, deciduous, tropical rainforest, evergreen, and crop (Table 1). In this work, we apply two different vegetation cover data sets that have been converted to the 8 PFTs: (i) the standard atlas-based distribution in NASA Model-E2 (Matthews, 1983) and (ii) the Simple Biosphere Model II (SiB2) distribution based on the International Satellite Land Surface Climatology Project data initiative II (Loveland, 2009). The SiB2 data set provides its own crop cover. The standard data set consists of a map of the world's vegetation cover that would most likely exist in equilibrium with present-day climate and natural disturbance, in the absence of human activities (potential natural vegetation) onto which the crop fraction for each grid cell is overlaid. The crop fraction in each model grid cell is constructed from a harmonized gridded data set for the year 2000 (Hurtt et al., 2011). The crop cover is imposed by proportional decrease/increase of all the potential natural vegetation types in the grid cell fraction that is not occupied by crops and/or pasture. There is no right choice in how to implement the crop cover, but this approach is the most common treatment of crop cover in global climate modeling (de Noblet-Ducoudre et al., 2012). The fractional coverage of global vegetated land area by each PFT for the standard and SiB2 data sets is shown in Table 1 (bright and dark bare soil fractions not shown). In the standard data set, the fractional cover of crop PFT is about double that in SiB2 (36.9 % versus 19.1 %) because tropical and subtropical land that is classified as crop PFT in the standard data set is classified as savanna and shrub PFTs in SiB2.

Leaf area index (LAI) for each PFT is prescribed according to regular seasonal sinusoidal variation between PFTspecific minimum and maximum seasonal LAI values that is insensitive to climate drivers or carbon balances (Rosenzweig and Abramopoulos, 1997; Friend and Kiang, 2005). A complete mechanistic understanding of the processes that control the development and senescence of foliage is not yet available such that the current state of phenology modeling may even be considered qualitative (Migliavacca et al., 2012; Richardson et al., 2013). We have made some improvements to the phenology for this work. Firstly, we have implemented a simplified crop phenology using a global data set of crop planting and harvesting dates (Sacks et al., 2010). The dominant crop type in each model grid cell was identified using a published data set of global crop maps and areal coverage (Monfreda et al., 2008). Then, a global model input file of mean plant date and harvest date was constructed for the grid cell's dominant crop type. The plant and harvest dates are recycled every simulation year. Secondly, a parameterization for phenological control (frost hardening) on photosynthetic capacity (V_{cmax} , Sect. 2.1.1) has been added for the evergreen PFT. V_{cmax} is reduced in winter to protect against cold injury. Sensitivity to temperature determines the revival of photosynthetic capacity, hence the length of the growing season (Toivonen et al., 1991; Makela et al., 2004; Hanninen and Kramer, 2007). The onset of the growing season for the evergreen PFT can be delayed if temperature remains cold despite light being available.

The use of fixed canopy structures and phenology means that leaf mass is not driven by photosynthetic uptake of $CO₂$ and a closed carbon cycle is not simulated. However, this version of the isoprene emission model can respond to elevated $CO₂$ with regard to: $CO₂$ fertilization, reduced stomatal conductance/increased water-use efficiency, CO_2 -inhibition, the temperature and precipitation responses of photosynthesis, but not phenological timing because the simulation does not account for the coupling of photosynthetic uptake to variability in growth. Application of LAI that is insensitive to climate may dampen the simulated interannual variability of isoprene emission in this model version.

2.1.2 Canopy biophysics

Each model PFT fraction in the vegetated part of each grid cell represents a single canopy. The model vertically stratifies each canopy into diffuse and direct light levels, and LAI profiles using an adaptive number of layers (typically 2–16)

(Friend and Kiang, 2005). The well established Michealis– Menten leaf model of photosynthesis (Farquhar et al., 1980; von Caemmerer and Farquhar, 1981) and the stomatal conductance model of Ball and Berry (Collatz et al., 1991) is used to compute the biophysical fluxes at the leaf level in each canopy layer based on appropriate parameters for each of the 8 PFTs from (Friend and Kiang, 2005) and the Community Land Model (Oleson et al., 2010) with updates from (Bonan et al., 2011) (Table A1). This coupled photosynthesis/stomatal conductance leaf model has previously been widely used to project terrestrial biosphere responses to global change. We summarize the model briefly here for transparency and completeness. The photosynthesis model assumes that the rate of net $CO₂$ assimilation in the leaves of C3 plants is limited by one of three processes: (i) the capacity of the ribulose 1,5-bisphosphate (RuBP) carboxylaseoxygenase enzyme (Rubisco) to consume RuBP (Rubiscolimited photosynthesis); (ii) the capacity of the Calvin cycle and the thylakoid reactions to regenerate RuBP supported by electron transport (electron transport-limited photosynthesis); and (iii) the capacity of starch and sucrose synthesis to consume triose phosphates and regenerate inorganic phosphate for photo-phosphorylation (triose phosphate uselimited photosynthesis). Photosynthesis is electron transportlimited under low light conditions including overcast/cloudy conditions, at the start and end of the day, for shaded leaves and understory vegetation. The three processes are described as functions of the internal leaf $CO₂$ concentration (C_i) and/or the maximum carboxylation capacity at the optimal temperature, 25° C, (V_{cmax}) (Table A1). Leaf stomata control for the uptake of $CO₂$ versus the loss of water vapor (H₂O). In the model, the stomatal conductance of H_2O through the leaf cuticle is linearly related to the net rate of carbon assimilation and the relative humidity, and inversely related to the $CO₂$ concentration at the leaf surface. The coupled system of photosynthesis, stomatal conductance and $CO₂$ diffusive flux transport equations form a cubic in C_i that is solved analytically (Baldocchi, 1994). A simple but realistic representation of soil water stress is included in the vegetation biophysics following the approach of Porporato et al. (2001). The algorithm reflects the relationship between soil water amount and the extent of stomatal closure ranging from no water stress to the soil moisture stress onset point (s^*) through to the wilting point (s_{wilt}) . Stomatal conductance is reduced linearly between the PFT-specific values of s[∗] and swilt (Table 1) based on the climate model's soil water volumetric saturation in 6 soil layers.

The leaf-level carbon and water fluxes are scaled up to the canopy level by integrating over each canopy layer. The land-surface model uses its own internal adaptive time step from 5 s to 15 min depending on the conditions. The carbon and water fluxes computed in the land-surface scheme are integrated over the climate model time step (30 min) for exchange with the model's atmosphere.

2.1.3 Leaf isoprene production

A leaf-level isoprene emission model that describes the constitutive production as a function of the electron transportlimited photosynthesis rate, J_e , (Niinemets et al., 1999) has been integrated into the canopy biophysics scheme following modifications for global-scale modeling (Arneth et al., 2007). The leaf-level isoprene emission rate (I) in units of μ mol m⁻²[leaf] s⁻¹ is calculated as follows:

$$
I = J_e \cdot \beta \cdot \kappa \cdot \tau \cdot \varepsilon,\tag{1}
$$

where J_e is the electron transport limited photosynthesis rate in units of μ mol m⁻²[leaf] s⁻¹. J_e is a linear function of the incident photosynthetically active radiation (PAR) and the internal leaf $CO₂$ concentration (Ci):

$$
J_{\rm e} = a_{\rm leaf} \cdot \text{PAR} \cdot \alpha_{\rm qe} \cdot \frac{c_i - \Gamma^*}{c_i - 2\Gamma^*},\tag{2}
$$

where a_{leaf} is the leaf-specific light absorbance and α_{qe} is the intrinsic quantum efficiency for photosynthetic $CO₂$ uptake in the chlorophyll reaction system that absorbs PAR to drive the oxidation of water and the reduction of enzymes (photosystem II). α_{qe} is a product of the fraction of absorbed light that reaches photosystem II and the $CO₂$ per absorbed photon. Γ^* is the CO₂ concentration compensation point in the absence of non-photorespiratory respiration (Collatz et al., 1991).

The β term in Eq. (1) translates the electron flux into isoprene equivalents given by Eq. (3):

$$
\beta = \frac{C_i - \Gamma^*}{6(4.67C_i + 9.33\Gamma^*)}.
$$
\n(3)

A detailed description of the mechanistic origin of the coefficient values is given elsewhere (Niinemets et al., 1999; Pacifico et al., 2011).

The atmospheric CO_2 -inhibition is included via a simple parameterization (κ) :

$$
\kappa = \frac{C_{i_standard}}{C_i},\tag{4}
$$

where $C_{i_{\text{1-standard}}}$ is the leaf internal CO_2 concentration at standard atmospheric $CO₂$, which is chosen to be the year 2000 global average value (370 ppmv). Equation (4) mimics the observed response to both short-term and long-term changes in C_i (Wilkinson et al., 2009; Heald et al., 2009). For example, short-term reductions in C_i due to stomatal closure under drought conditions imply increases in κ , while the long-term effects of increasing (decreasing) atmospheric CO₂ imply decreases (increases) in κ .

The temperature relationship (τ) in the algorithm accounts for the difference in temperature optimum between photosynthesis and isoprene synthase:

$$
\tau = \exp[0.1(T - T_{\text{ref}})],\tag{5}
$$

where $T =$ leaf temperature and $T_{ref} =$ standard temperature condition (30 °C). The temperature optimum for isoprene synthase is about 40° C (Guenther et al., 1991). Equation (5) does not simulate a temperature optimum after which isoprene emission rate decreases with further increases in temperature. Such high temperature conditions in isoprene emitting biomes rarely occur in nature at large ecosystem scales. Canopy-scale temperatures of this magnitude may occur under severe drought stress conditions when transpiration is significantly reduced. Yale-E2 uses the canopy temperature (not air temperature) in Eq. (5), which represents a significant departure from other CCMs and CTMs that drive interactive isoprene emissions with surface air temperature. Yale-E2 intrinsically captures the effects of changing stomatal conductance on canopy energy balance, which affects the canopy temperature, and thus the isoprene emission rate. In future and past hot greenhouse worlds, plant photosynthesis may acclimate to the higher temperatures (Arneth et al., 2012). This plastic adjustment of photosynthesis will indirectly impact isoprene emission. Whether the temperature optimum for isoprene synthase will similarly shift in warmer climates is not known.

 ε is the PFT-specific fraction of electrons available for isoprene synthesis or isoprene emission potential and parallels the use of a PFT-specific leaf-level isoprene emission capacity in Guenther et al. (1991, 1995). To calculate ε , we make use of available generic PFT-specific standard (or basal) leaf isoprene emission rates (I_s) in $\mu g C g^{-1}$ [leaf] h⁻¹ based on recommendations from a wide range of observations (Table 1). Standard conditions are defined as: surface air temperature = 30 °C and PAR = 1000 µmol m⁻² s⁻¹. Again, we assume that atmospheric $CO₂$ concentration = 370 ppm under standard conditions. I_s must be converted to appropriate units of leaf area using the model values of specific leaf area (SLA), the amount of light-capturing surface area that is deployed with a given investment of dry mass in units of m²[leaf] g⁻¹[leaf]. SLA is intimately connected to the resource use economy of the plant (Milla and Reich, 2007). Then, ε is computed for each PFT by substituting $I = I_s$ into Eq. 1) under standard conditions. The crop and tundra PFTs are non-emitting for isoprene in this model although recently, isoprene emission has been observed from a tundra ecosystem (Potosnak et al., 2013). Deciduous, shrub and rainforest PFTs maintain the highest fraction of electrons available for isoprene synthesis (Table 1). In the current model, ε does not vary with time of day or season (Niinemets et al., 2010a, c) or through the canopy (Niinemets et al., 2010b). Many plant species in temperate and boreal ecosystems exhibit delayed onset of isoprene emission after the leaf development (Grinspoon et al., 1991; Hakola et al., 1998; Olofsson et al., 2005). Leaf age effects have been included in other global isoprene emission models using simplistic functions, for example involving growing degree-day temperature (Arneth et al., 2007; Guenther et al., 2006; Lathiére et al., 2006).

Scaling of isoprene emission from the leaf to the canopy uses the canopy vertical stratification and integration scheme as described in Sect. 2.1.2. The canopy-level isoprene fluxes are passed to the model's atmosphere through the landsurface scheme on the 30 min climate model time step.

3 Simulations

We apply Yale-E2 in an atmosphere-only configuration driven by sea surface temperatures and sea ice cover prescribed according to decadal average monthly varying observations for 1996–2005 from the HadSST2 data set (Rayner et al., 2006). Atmospheric $CO₂$ is prescribed to a uniform concentration of 370 ppmv representative of approximately year 2000 levels. We perform a control simulation ("SimCONT") that uses the standard atlas-based vegetation cover distribution in NASA Model-E2 (Matthews, 1983). In addition, we perform three sensitivity simulations described in Table 2. Sensitivity simulation "SimSiB2" uses an alternate vegetation cover data set based on the SiB2 distribution. We perform a simulation "SimH2O" in which the plant water stress function in the biophysics module is artificially switched off to quantify the effects of water limitation on isoprene emission rate. In order to explore the extent of human land cover change impacts on isoprene, a simulation "SimPNV" is run that uses the potential natural vegetation cover data set (i.e. without the crop PFT superimposed). The control and sensitivity simulations are each run for 10 model years that are averaged for the evaluation analyses. The model version applied in this study outputs the total isoprene (and GPP) in each grid cell and does not characterize PFT-specific attribution. The model's carbon fluxes (GPP and isoprene) are present-day climatologies and do not pertain to any specific meteorological year.

4 Results

4.1 Simulated global GPP and isoprene emission

The global annual average isoprene emission and GPP flux for each simulation are reported in Table 2. Global GPP in SimCONT and SimSiB2 $(124-126$ PgC yr⁻¹) is in reasonable agreement with current understanding of the contemporary carbon cycle budget. The global isoprene source in Sim-CONT and SimSiB2 $(451-498 \text{ TgC yr}^{-1})$ is consistent with previous global estimates of 400–700 TgC yr⁻¹ (Guenther et al., 2006). This good agreement is because the model applies widely used generic PFT-specific I_s values to derive the PFT-specific fraction of electrons for isoprene synthesis (ε) as previously alluded (Arneth et al., 2008). The spatial distribution of annual and seasonal average GPP and isoprene emission are shown in Fig. 1a, b, respectively. The major isoprene emitting biome is tropical rainforest that provides a perennial reactive carbon flux to the atmosphere, but the

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Table 2. Summary of global annual average carbon fluxes in the model runs. In total 10 model run years are included in the annual average for each simulation. All runs use 1996–2005 average monthly varying SSTs and sea ice. Uncertainty values are based on the standard error relative to internal climate model variability ($n = 10$ yr).

Simulation	Land cover	Description and purpose	GPP $(PgCyr^{-1})$	Isoprene $(TgCyr^{-1})$
SimCONT	Standard, Matthews (1983)	Present Climate Control	124 ± 2	451 ± 5
SimSiB2	SiB2, Loveland (2009)	Test impact of alternate vegetation cover data set	$126 + 2$	498 ± 5
SimH ₂ O	Standard, Matthews (1983)	Test impact of water stress function in vegetation biophysics	155 ± 1	592 ± 3
SimPNV	Standard $+$ no human alteration, Matthews (1983)	Test impact of potential natural vegetation	$137 + 2$	701 ± 10

Fig. 1a. Spatial distribution of annual and seasonal average modeled GPP in $gC m^{-2}$ day⁻¹ in the control run SimCONT.

Fig. 1b. Spatial distribution of annual and seasonal average modeled isoprene emission in mgC m⁻² h⁻¹ in the control run SimCONT.

growing season isoprene emission in the eastern US rivals the tropical emission in magnitude.

Globally, the isoprene source strength is 10 % higher in SimCONT than SimSiB2 whereas GPP is only ∼1 % higher. The different response sensitivity of isoprene and GPP to the vegetation cover data set arises because of the differences in the classification of crop PFT between data sets, and all PFTs assimilate carbon but the crop PFT is non-emitting for isoprene. Previous studies have suggested an even higher sensitivity of the global isoprene source (20–30 %) to the model's vegetation cover data sets for a given algorithm (Guenther et al., 2006; Pfister et al., 2008; Arneth et al., 2011). In the artificial absence of plant water stress, GPP increases by 25 % with a similar increase in global isoprene emission (∼30 %). Similarly, the MEGAN model suggested that accounting for soil moisture stress in the present climate state leads to a 20–30 % reduction in the global isoprene source strength (Müller et al., 2008). The response to potential natural vegetation cover is more dramatic: the global isoprene source increases by 55 % whereas GPP increases by only 10 %. The strong sensitivity of isoprene to potential natural vegetation cover suggests significantly higher isoprene emission in pre-human disturbance conditions and large variability in isoprene emission due to human land cover change. In the most recent community assessment of anthropogenic ozone radiative forcing led by the Atmospheric Chemistry and Climate Model Intercomparison Project, only 4 of 15 participating state-of-the-science global chemistry-climate models included climate-sensitive isoprene emission (Young et al., 2013). Those 4 models all projected a small increase in isoprene emission from preindustrial to present-day in response to temperature change. 9 out of 15 models prescribed isoprene emission and used the same off-line input data for preindustrial and present-day.

There are substantial regional differences in isoprene emission between the sensitivity and control simulations (Fig. 2). Annual average isoprene emission in Europe and the southeastern US is smaller in SimCONT than SimSiB2 by ~0.5–1.5 mgC m⁻² h⁻¹. In the tropical biome, SimCONT simulates higher annual average isoprene emission than Sim-SiB2 by \sim 0.3–1 mgC m⁻² h⁻¹. The presence of plant water stress affects isoprene in all emitting biomes. In the artificial absence of plant water stress, large increases in the annual average isoprene emission (\sim 1 mgC m⁻² h⁻¹) occur in the tropics, southeastern US, and the boreal region. Potential natural vegetation cover implies additional isoprene emission $\sim 2 \,\text{mgC} \,\text{m}^{-2} \,\text{h}^{-1}$ on the annual average) in the eastern US, Europe and the tropics on land that has zero emission capacity in the standard cover data set.

4.1.1 Isoprene emission sensitivity to GPP and canopy temperature

We have applied multiple linear regression to compute the standardized regression coefficients between isoprene emis- \mathbf{r}

Fig. 2. Difference in simulated annual average isoprene emission relative to the control simulation for an alternative vegetation cover map (top), in the artificial absence of water stress (middle), and for potential natural vegetation cover (bottom). Units are mgC m⁻² h⁻¹.

Figure 2. Different and **Figure 2.** Different and **Figure 2.** The different driving variables (Fig. 3). The leaf-level isoprene emission rate is linearly related ression to compute the two the electron transport-infinited photosynthesis rate, J_e between isoprene emission follows an sion and the key drivers (GPP and canopy temperature) in boreal summer for SimCONT and SimH2O (Fig. 3). The coefficients represent the partial regression coefficients in units of standard deviation, such that they can be directly compared with each other to determine the relato the electron transport-limited photosynthesis rate, J_e

Fig. 3. Standardized regression coefficients between isoprene emission and the key drivers GPP (left column) and canopy temperature (right column). Results shown are averages for June-July-August in SimCONT (top row), SimH2O (middle row), and the difference [SimH2O – SimCONT] (bottom row).

Latitude (°)

Figure 4. Seasonal zonal average GPP simulated in SimCONT and SimSiB2 compared to the FLUXNET-derived dataset (gC/m² /day). and SimSiB2 compared to the FLUXNET-derived data set **Fig. 4.** Seasonal zonal average GPP simulated in SimCONT $(gC m^{-2} day^{-1})$.

exponential relationship with temperature (Eq. 5). In the control run (SimCONT), the standardized regression coefficient is about a factor of 2 higher for GPP than canopy tem-

 \mathbb{F} \mathbb{F} \mathbb{F} \mathbb{F} over canopy temperature in controlling isoprene emission $\begin{array}{c|c|c|c|c|c} \hline \end{array}$ \mathcal{U} (i.e. \mathcal{U} and \mathcal{U} ariability in most regions of the world in the model. In the absence of water stress (SimH2O), the standardized regres-
 \mathbb{R}^d \mathbb{R}^d absence of water stress (SimH2O), the standardized regresperature everywhere except the most severely water-limited ecosystems (e.g. Indian subcontinent). Thus, GPP dominates sion coefficients for canopy temperature increase by around 30 % while the standardized regression coefficients for GPP decrease by around 30 %, relative to the control run (Sim-CONT). However, GPP still exerts the dominant control over canopy temperature in controlling isoprene emission variability in this model. The presence of plant water stress reduces the sensitivity of isoprene emission to canopy temperature because drought conditions are frequently associated with high temperatures that result in a decrease in GPP and therefore isoprene emission. Two important corollaries are that (1) variability in isoprene emission is largely determined by the same factors that determine GPP variability (temperature, precipitation, the availability of water and nutrients, disturbance) and (2) isoprene emission algorithms that do not account for the impacts of water availability may overestimate the sensitivity of isoprene emission to canopy (and/or surface air) temperature. Since a global observational data set of isoprene emission is not available, we are unable to

explore these relationships at large regional scales in the real world.

4.2 Evaluation of global GPP and isoprene emission

Direct large-scale measurements of photosynthesis and isoprene emission do not exist. The FLUXNET network of tower sites continuously measures the $CO₂$ net ecosystem exchange (NEE) flux between the biosphere and atmosphere using the eddy covariance technique (Baldocchi et al., 2001). NEE is the imbalance between photosynthesis and ecosystem respiration. Photosynthesis is zero at night, which allows for a temperature-based predictor of the respiration that can be used to extrapolate GPP from the day time NEE. Isoprene fluxes have been measured during short campaigns at several FLUXNET sites. Our approach to evaluate the global model performance is to compare the simulated GPP and isoprene emission rate to available above-canopy flux tower measurements of the carbon fluxes. Recent research has shown that space-based observations of formaldehyde (HCHO) columns may be a useful proxy of surface isoprene emissions across broader ecosystem and regional scales (Palmer et al., 2003; Barkley et al., 2009). We elect not to use this approach here because of the inherent limitation that an a priori isoprene emission model is necessary to interpret the HCHO satellite columns (Barkley et al., 2012).

Because of the different spatial and temporal scales of the model output versus the observations, comparing to point measurements from flux towers represents an extremely stringent performance test and a number of caveats must be emphasized. Firstly, the model GPP and isoprene emission are extracted for the single grid cell ($2° \times 2.5°$ latitude by longitude) in which the flux tower site is located. Our rationale is that, while the typical eddy covariance footprint is small (\sim 1 km²), the trace gas flux variability at the flux towers is often representative over much larger spatial scales because of the spatial coherence of climate anomalies (Ciais et al., 2005). Secondly, the model parameters are not tuned in any way to the local site vegetation properties. The model grid cell level output is based entirely on the vegetation structure and soil properties in Yale-E2 and not the flux tower site data. Finally, the point measurements are obtained at flux towers during specific meteorological periods. The model output represents 10 yr average climatological GPP and isoprene emission rates (and meteorological variables) at the site locations, which do not necessarily reflect the exact local weather conditions that occurred during the observation period. Since the climate model has not been forced to the observed meteorology during the measurement periods, the model cannot be expected to reproduce day-today variability in the isoprene emission and GPP. Specifically for isoprene, the measurements are canopy exchange fluxes whereas the current model configuration does not include isoprene loss through the canopy. However, canopy loss is likely less than 10 % of the total emission (Karl et al., 2004; Rinne et al., 2012). Based on the above caveats, we posit that the most important diagnostic quantities in this model/measurement comparison are: zonal average and seasonal cycle for GPP flux, and campaign-average variability across different ecosystems and diurnal cycle for isoprene emission. Focusing on these diagnostic quantities does provide for valuable insights into the global climate model's strengths and weaknesses. Typical measurement uncertainties associated with the eddy flux technique are about ± 30 %.

4.2.1 FLUXNET-derived global GPP

We compare the model simulated GPP (SimCONT and Sim-SiB2) to a global GPP data set that has been generated by data orientated diagnostic upscaling of site-derived GPP from FLUXNET (Beer et al., 2010; Jung et al., 2011; Bonan et al., 2011). The model runs reproduce the seasonal zonal average variability in the FLUXNET-derived data set with remarkable fidelity (Fig. 4). Based on linear regression, the model runs capture ≥ 89 % of the variability in the zonal seasonal average FLUXNET-derived data set. The model runs perform best in the summer and winter seasons (\geq 93 % of the variability). SimSiB2 demonstrates slightly superior performance over SimCONT. The Northern Hemisphere (NH) June-July-August maximum at 50◦–60◦ N in the deciduous and crop biomes is well reproduced by the model (to within 10 %). In March-April-May and September-October-November, the model underestimates tropical GPP and overestimates GPP in the NH mid-latitudes between 35◦–60◦ N by around 10 %. Overall, the comparison results provide confidence in the model's ability to simulate the magnitude and zonal average variability in GPP.

4.2.2 GPP and latent heat seasonal cycle at 6 benchmark sites

A benchmarking system of seasonal FLUXNET data at select sites has been constructed for evaluation of the seasonal cycle of carbon and water biophysical fluxes in global modeling frameworks (Blyth et al., 2011). The characteristics of the 6 benchmark sites that are relevant for isoprene emission are described in Table 3. At the Hyytiala, Harvard and Santarem sites, the dominant vegetation PFT in the model grid cell is consistent with the FLUXNET tower site. At Morgan Monroe and Tharandt, the dominant model PFT in the grid cell is crop versus deciduous and evergreen, respectively at the FLUXNET tower sites. The Kaamanen FLUXNET tower site is wetland/woody savanna in reality whereas the standard cover is predominantly grass PFT in this model grid cell, and the SiB2 cover is a mixture of evergreen, deciduous and shrub PFTs.

The climatological models capture the GPP seasonality at the 6 sites reasonably well (Fig. 5a). We provide root mean square bias (RMSE) as the diagnostic metric (Table 3). All model climatologies overpredict at both NH high latitude

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Table 3. Summary of FLUXNET benchmark site characteristics, model vegetation cover fractions for the grid cell in which the site is located and model performance in terms of RMSE range across the model runs. Td = tundra; $G =$ grass; Sh = shrub; S = savanna; D = deciduous; $T =$ tropical rainforest; $E =$ evergreen; and $C =$ crop.

		Site characteristics		Vegetation cover fraction (%)		RMSE	
Name	Data year	Ecosystem Type	Location	SimCONT	SimSiB2	GPP	Latent Heat
						(umol m ² s ⁻¹)	(Wm^{-2})
Kaamanen	2002	Wetland/woody savanna	69.1° N, 27.3° E	G:D 80/10	E:Sh:D:Td:S:C 40/31/17/9/3	$1.3 - 1.5$	$7.4 - 9.0$
Hyytiala	2000	Evergreen needleleaf	61.8° N, 24.3° E	E:D:C 35/34/31	E: C:D 75/19/5	$1.4 - 1.7$	$12.7 - 14.4$
Morgan Monroe	2002	Deciduous broadleaf	39.3° N, 86.4° W	C:D 51/49	C:D 67/31	$1.7 - 1.8$	$20.5 - 21.6$
Harvard	1999	Deciduous broadleaf	42.5 \degree N, 72.2 \degree W	D: C.97/3	$D: E: C$ 79/16/5	$1.9 - 2.6$	$24.0 - 25.6$
Tharandt	1999	Evergreen needleleaf	51.0° N, 13.6° E	C.E.D 51/25/24	$C: D: E$ 82/15/2	$2.2 - 3.4$	$14.4 - 16.4$
Santarem Km67	2003	Tropical evergreen broadleaf	2.9° S, 55.0° W	$T:$ C 94/6	$T: S: E$ 87/7/3	$0.4 - 1.0$	$6.1 - 10.5$

GPP fluxes at 6 FLUXNET sites (μ molC m⁻² s⁻¹). **Fig. 5a.** Comparison of eddy-derived and modeled monthly average

Fig. 5b. Comparison of measured and modeled latent heat flux at 6 FLUXNET sites (W m⁻²).

sites (Kaamanen and Hyytiala) and underpredict at NH midlatitude sites (Morgan Monroe, Harvard and Tharandt). The model demonstrates skillful ability to reproduce the GPP flux at the tropical site (Santarem). SimCONT and SimSiB2 capture the observed GPP reduction in the dry season that is over predicted in the artificial absence of water stress (SimH2O). The model's ability to reproduce the magnitude of summer GPP at Harvard is improved in the absence of water stress (SimH2O) but that configuration gives too high GPP in the spring and fall.

Previous analyses of FLUXNET measurements at a wide range of sites found a strong linkage between carbon gain and water loss for all biomes integrated over the year (Law et al., 2002). Figure 5b compares the modeled and measured latent heat (evaporative) fluxes at each of the 6 FLUXNET sites. RMSE values are shown in Table 4. Yale-E2 satisfactorily captures the seasonal cycle in latent heat flux at all sites, except at Harvard. Again, artificially turning off the water stress function does appear to improve the model's latent heat simulation at the Harvard site. The model's high values of latent heat outside of the growing season are due to the grid cell having 25 % water coverage.

The model/measurement comparison of the GPP seasonal cycle at the 6 benchmark sites permits reasonable confidence in the model's biophysics for the simulation of isoprene emission. Based on the model GPP biases alone, the model may be expected to overestimate isoprene emission at NH high latitudes, underestimate at NH mid-latitudes, and perform well in the tropics.

Table 4. Description of the 28 campaign-average above-canopy isoprene flux measurements collected between 1995–2010 across a wide range of ecosystem types, regions and seasons that comprise the benchmark global database. Codes for model vegetation fractions are: Td = tundra; G = grass; Sh = shrub; S = savanna; D = deciduous; T = tropical rainforest; E = evergreen; and C = crop.

Measurement sites				Model grid cell vegetation fractions		
Identifier	Ecosystem	Date	Location	Reference	Standard	SiB2
Santarem	Rainforest	07/2000	2.9° S, 55° W	Rinne et al. (2002)	T: 94	T/Sv/E: 87/8/4
Manaus	Rainforest	07/2001	2.35° S, 60° W	Kuhn et al. (2007)	T: 99	T/Sv: 89/5
Manaus	Rainforest	09/2004	2.6° S, 60° W	Karl et al. (2007)	T: 99	T/Sv: 89/5
Costa Rica	Rainforest	04/2003	10° N, 84° W	Karl et al. (2004)	T/C: 82/18	T/C: 74/14
Costa Rica	Tropical lowland	10/1999	10° N, 84° W	Geron et al. (2002)	T/C: 82/18	T/C: 74/14
Malaysia	Rainforest	11/2003	3° N, 102° E	Saito et al. (2008)	T/C: 79/21	$C/T/D$: 45/27/16
Borneo	Rainforest	04/2008	5° N, 118° E	Langford et al. (2010a)	T/C: 87/13	T/E: 78/15
Borneo	Rainforest	$06 - 07/2008$	5° N, 118° E	Langford et al. (2010a)	T/C: 87/13	T/E: 78/15
Congo	Mixed tropical	03/1996	2° N, 16° E	Serca et al. (2001)	T/C: 93/7	T/95
Congo	Mixed tropical	11/1996	2° N, 16° E	Serca et al. (2001)	T/C: 93/7	T/95
Congo	Mixed tropical	11/1996	4° N, 18° E	Greenberg et al. (1999)	T/Sv/C: 72/18/10	T/Sv: 76/19
South Africa	Savanna	02/2001	25° S. 31° E	Harley et al. (2003)	C/Sh/Sv: 64/22/14	$C/D/T$: 66/14/18
Mongolia	Grassland	$06 - 07 - 08/2002$	44° N, 116° E	Bai et al. (2006)	C/Sh: 97/2	G/C: 84/10
Mongolia	Grassland	09/2002	44° N. 116° E	Bai et al. (2006)	C/Sh: 97/2	G/C: 84/10
SW China	Plantation	$02 - 03/2002$	21° N, 101° E	Baker et al. (2005)	T: 100	T/D/E: 43/27/27
SW China	Plantation	07/2002	21° N. 101° E	Baker et al. (2005)	T: 100	T/D/E: 43/27/27
Germany	Deciduous	07/2001-2002	50° N, 11° E	Graus et al. (2006)	C/D : 48/47	C/D: 84/15
NW Germany	Deciduous	07/2003	50° N, 6° E	Spirig et al. (2005)	C/D: 52/48	C/D: 85/14
Germany	Deciduous	08/1995	50° N, 8° E	Steinbrecher et al. (2000)	D/C: 59/41	C/D : 74/24
Sweden	Plantation	07/2001	58° N, 12° E	Olofsson et al. (2005)	$E/C/D$: 49/35/16	C: 97
Finland	Boreal	08/2001	61° N, 24° E	Spirig et al. (2004)	E/D/C: 35/33/32	E/C: 75/19
N Wisconsin	Deciduous	07/1993	46° N, 90° W	Isebrands et al. (1999)	D/C/Sv: 43/37/19	C: 86
UMBS	Deciduous	06-07-08/2000-2002	46° N, 85° W	Pressley et al. (2005)	D/E/C: 57/29/13	$E/D/C$: 36/33/31
UMBS	Deciduous	08/1998	46N, 85W	Westberg et al. (2001)	D/E/C: 57/29/13	$E/D/C$: 36/33/31
Harvard Forest	Deciduous	$06 - 07 - 08/1995$	45° N, 72° W	Goldstein et al. (1998)	D/C: 92/8	D/C : 64/26
Canada	Boreal	07/1994	42° N, 72° W	Pattey et al. (1999)	C/E/Sv: 67/16/10	C/E: 70/13
Canada	B oreal	09/1994	54° N, 105° W	Pattey et al. (1999)	C/E/Sv: 67/16/10	C/E: 70/13
London	Urban	10/2006	54° N, 105° W	Langford et al. (2010b)	D/C: 57/43	C: 98

4.2.3 Global database of campaign-average isoprene flux measurements

Our goal is to assess the model's ability to capture the isoprene emission variability across a wide range of ecosystem types and seasons. We have assembled a global database of campaign-average above-canopy isoprene flux measurements obtained between 1995–2010 (Table 4). The measurements are sorted by ecosystem type. The dominant vegetation fraction in the model grid cells for both the standard and SiB2 vegetation cover data sets corresponds to the local ecosystem at the measurement towers for all sites in the database except for South Africa and SW China. Table 5 compares the model simulated isoprene emission in Sim-CONT, SimH2O and SimSiB2 to the observations in the benchmark database (shown graphically in Fig. 6 for Sim-CONT and SimH2O results). The model results are monthly or seasonal averages that correspond to the time period during which the observations were made. The campaign periods are typically shorter than one month and as such the observations tend to refer to shorter averaging periods than in the model output. In some cases the published observational campaign-average is for mid-day average or daytime average as indicated in Table 5. Evaluation against this global database suggests possible systematic biases in the model: under prediction in the summer time Canadian boreal but over prediction in the Scandinavian boreal. The model performs well in simulating the magnitude of isoprene mission in the tropical biome but appears to simulate either a lack of seasonality or reverse seasonality compared to that observed at the non-Amazon tropical rainforest sites. The ORCHIDEE model isoprene emission has been compared to a limited subset of this global database (Canada, UMBS and Santarem) (Lathiére et al., 2006). Similar to Yale-E2, ORCHIDEE reproduced the tropical flux magnitude well but underpredicted at the Canada and UMBS sites. The presence of water stress has the largest impact on the magnitude of isoprene emission rate in the North American deciduous sites. Based on linear regression, the control run SimCONT is able to reproduce 50 % of the variability across different ecosystems and seasons in the global database. SimH2O and SimSiB2 capture 64 % and 40 % of the variability in the global database, respectively. The model/measurement discrepancies in the isoprene emission magnitude are most likely due to the model's assignment of PFT-specific emission potentials (fraction of electrons available for isoprene synthesis, ε) that in reality

Table 5. Comparison of the global database and simulated isoprene emission. Units are mgC m⁻² h⁻¹. The model values are either monthly or seasonal means to be consistent with the observation time period. The averaging period for the observations is indicated where the information is available. [MDA = mid-day average; DTA = daytime average; CA = campaign average; DTMX = daytime maximum].

vary greatly for a single plant species (Goldstein et al., 1998) and across plant species lumped into each PFT class e.g. (Rinne et al., 2009). For example, the deciduous PFT includes species that are strong isoprene emitters (willows, aspen) and others that do not emit isoprene (birches).

4.2.4 Time-varying isoprene emission through campaign periods

We examine the time-varying performance at 9 select measurement sites described in Table 6. Some of the sites are part of the global database of campaign-average fluxes (Table 3). There are 4 tropical sites on 2 different continents, 4 temperate broadleaf sites on 2 different continents, and 1 temperate mixed site in Europe. At 3 of the temperate sites, measurements are available across multiple years: Harvard Forest (1995 and 2007); UMBS (2000, 2001, 2002, 2003, 2005); and Belgium (2009–2011). These three sites provide measurements that span an entire growing season. The remaining 6 sites offer measurements over periods of a few days to several weeks. The sites in southern France (La Verdière and Montmeyan) occur in the same model grid cell. We focus on analysis of the control run SimCONT and the sensitivity simulation SimH2O. Sub-daily model output is generated at half-hourly resolution (the physical time step of the climate model itself). The model isoprene emissions are 10 yr climatological averages for present-day, thus cannot be expected to reproduce the day-to-day weather-related variability in the observations.

In this section, we assess qualitatively the model/measurement comparison of isoprene emission, surface air temperature (SAT) and downward shortwave radiation (SW) across the entire campaign period for the 6 sites with measurement periods shorter than a growing season. The goal is to identify possible model biases in isoprene emission magnitude before a more quantitative assessment of the average diurnal cycle is executed in the next section. Results for tropical sites are shown in Fig. 7a–f and results for the temperate sites are shown in

Fig. 6. Scatter plot of the simulated isoprene emissions against measurements from the global above-canopy flux database sorted by ecosystem type (Table 5). "Plantation" measurements are lumped into "crop" here. Units are mgC m⁻² h⁻¹. Values and temporal averaging periods are detailed in Table 5.

Fig. 7a. (a–f) Time evolution of isoprene emission flux, surface air temperature (SAT) and downward shortwave (SW) radiation at the tropical measurement sites. Measurements are shown in black new appear measurement sites. Heasurements are shown in stated (where data is available). Model climatological results are averages of 10 simulation years in SimCONT (red) and SimH2O (blue). **(a)** Costa Rica during 17 April–4 May, 2003.

Fig. 8a–e. Results for Harvard Forest (2007) have been split into three separate time periods. Meteorological data are not available for all sites. Measured isoprene fluxes are highly variable during the day at both tropical and temperate sites reflecting the local weather conditions. The model

Fig. 7b. (a–f) Time evolution of isoprene emission flux, surface air temperature (SAT) and downward shortwave (SW) radiation at the tropical measurement sites. Measurements are shown in black (where data is available). Model climatological results are averages of 10 simulation years in SimCONT (red) and SimH2O (blue). **(b)** Manaus during 17–27 September 2004.

Fig. 7c. (a–f) Time evolution of isoprene emission flux, surface air temperature (SAT) and downward shortwave (SW) radiation at the tropical measurement sites. Measurements are shown in black (where data is available). Model climatological results are averages of 10 simulation years in SimCONT (red) and SimH2O (blue). **(c)** wet season Santarem during 17–21 April 2001.

reproduces the SW with striking fidelity at both the tropical sites and temperate sites (where data is available). The SAT simulation is generally weaker and the discrepancies are not related to a particular biome. In Costa Rica, the model underestimates the diurnal range in temperature. In Manaus, La Verdière and Montmeyan, the model captures accurately the daytime maximum SAT but is not able to simulate the nighttime minimum (the nighttime SAT is too warm in the

the tropical measurement sites. Measurements are shown in black **Fig. 7d. (a–f)** Time evolution of isoprene emission flux, surface air temperature (SAT) and downward shortwave (SW) radiation at (where data is available). Model climatological results are averages of 10 simulation years in SimCONT (red) and SimH2O (blue). **(d)** dry season Santarem during 22 October–6 November 2003.

Fig. 7e. (a–f) Time evolution of isoprene emission flux, surface air temperature (SAT) and downward shortwave (SW) radiation at the tropical measurement sites. Measurements are shown in black (where data is available). Model climatological results are averages of 10 simulation years in SimCONT (red) and SimH2O (blue). **(e)** wet season Borneo during 21 April–9 May 2008.

model versus the observations). In Borneo the model's SAT diurnal range appears reasonable but there is a consistent ∼5 ◦C positive bias. Only at the Harvard Forest site does the model accurately reproduce the observed SAT throughout the measurement campaign. It must be emphasized that the

Fig. 7f. (a–f) Time evolution of isoprene emission flux, surface air temperature (SAT) and downward shortwave (SW) radiation at the tropical measurement sites. Measurements are shown in black (where data is available). Model climatological results are averages of 10 simulation years in SimCONT (red) and SimH2O (blue).**(f)** early dry season Borneo during 22 June–22 July 2008.

model's climatological temperature and radiation variables refer to average values over the $2° \times 2.5°$ grid cell. SAT is expected to exhibit much higher sub-grid scale variability than downward SW.

The model demonstrates significant skill in closely reproducing the time-varying isoprene emission in the dry season tropics at the Manaus, Santarem and Borneo (early-dry) sites. However, there is an obvious model overestimate at the Costa Rica, Santarem and Borneo sites in the wet season. A large part of the discrepancy is likely related to variation in the emission potential (ε) across different tropical forests. Some of the discrepancy at the Santarem site in the wet season may be related to the lack of leaf age effects in the model (i.e. no seasonal variation in ε). For instance, low HCHO columns over Amazonia observed from space during the wet season have been ascribed to leaf flushing prior to the dry season (Barkley et al., 2009).

As expected, at the wet season tropical sites, SimCONT and SimH2O yield similar climatological fluxes. At the dry season tropical sites, especially Santarem and Borneo, Sim-CONT reduces to about half the value in SimH2O and more closely matches the observations lending some confidence in the model's simulation of water-stressed photosynthesis and isoprene emission in this region, previously indicated in Fig. 5a. Some of the model overestimate in Borneo may be related to the climate model's high bias in SAT. At the temperate sites in the growing season, the isoprene emission magnitude is well captured in the model (within a factor of 2 of the observations). At Montmeyan (June) both model

Fig. 8a. (a-e) Time evolution of isoprene emission flux, surface air temperature (SAT) and downward shortwave (SW) radiation at the temperate measurement sites. Measurements are shown in black (where data is available). Model climatological results are averages of 10 simulation years in SIMCONT (red) and SimH2O (blue). **(a)** La Verdière during 21 June–6 July 2000.

Fig. 8b. (a–e) Time evolution of isoprene emission flux, surface air temperature (SAT) and downward shortwave (SW) radiation at the temperate measurement sites. Measurements are shown in black (where data is available). Model climatological results are averages of 10 simulation years in SIMCONT (red) and SimH2O (blue). **(b)** Montmeyan during 11–27 June 2001.

runs perform well, whereas at La Verdière (June–July), Sim-CONT reproduces the measured isoprene flux more closely than SimH2O, indicative of increases in water stress later into the growing season. At the Harvard Forest site, the measured fluxes tend to lie between SimCONT and SimH2O.

Fig. 8c. (a-e) Time evolution of isoprene emission flux, surface air temperature (SAT) and downward shortwave (SW) radiation at the temperate measurement sites. Measurements are shown in black (where data is available). Model climatological results are averages of 10 simulation years in SIMCONT (red) and SimH2O (blue). **(c)** Harvard Forest during 13–22 June 2007.

Fig. 8d. (a–e) Time evolution of isoprene emission flux, surface air temperature (SAT) and downward shortwave (SW) radiation at the temperate measurement sites. Measurements are shown in black (where data is available). Model climatological results are averages of 10 simulation years in SIMCONT (red) and SimH2O (blue). **(d)** Harvard Forest during 20–31 July 2007.

The model reproduces the decrease in emission magnitude in early September at this site.

Fig. 8e. (a–e) Time evolution of isoprene emission flux, surface air temperature (SAT) and downward shortwave (SW) radiation at the temperate measurement sites. Measurements are shown in black (where data is available). Model climatological results are averages of 10 simulation years in SIMCONT (red) and SimH2O (blue). **(e)** Harvard Forest during 31 August–6 September 2007.

4.2.5 Isoprene and GPP diurnal cycle

We constructed campaign average diurnal cycles at 30 min resolution for isoprene emission at the 9 measurement sites (1 h resolution for Harvard Forest in 1995) and GPP at 2 temperate sites where suitable data is available (UMBS and Harvard Forest). The measurement days included in the averaging are indicated in Table 6. At UMBS and Belgium, the isoprene (and GPP at UMBS) average diurnal cycle includes the data for all available years. The interannual variability in the measured isoprene diurnal cycle (30–100 %) is larger than for the GPP diurnal cycle $(< 5\%)$ (not shown).

Comparison between measured and modeled isoprene average diurnal cycles at the tropical and temperate sites are shown in Fig. 9a and b, and GPP average diurnal cycle at 2 temperate sites in Fig. 9c. Model/measurement evaluation statistics including the correlation coefficient for linear regression and root mean square error (RMSE) are indicated in Tables 7 and 8 along with a comparison of the measured and modeled maximum average diurnal emission. The measured diurnal average profiles show large fluctuations during the middle of the day that are not seen in the smoother 10 year average model climatologies. The 5 yr average diurnal average at UMBS does not exhibit fluctuations to the same extent as the other sites.

The model simulates the magnitude of the average diurnal cycle in the dry season tropics and the growing season temperate zone (except at Belgium) to within a factor of 2, which is comparable to the performance of higher spatial resolution models driven with observed meteorology that are used in air

Fig. 9a. (a–c) Comparison between measured and modeled isoprene emission and GPP average diurnal cycles for the time periods indicated in Table 6. Measurements are black lines. Error bars on the measurements represent ± 1 standard deviation. Model results are 10 yr climatological averages for SimCONT (red) and SimH2O (blue). The dotted lines represent model uncertainty due to internal variability in the climate model computed as ± 1 standard deviation for $n = 10$ model run years. (a) isoprene emission tropical sites (wet season results are shown in the left column and dry season results are shown in the right column)

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quality modeling for the development of environmental policy e.g. (Warneke et al., 2010; Arneth et al., 2011). Based on the RMSE (Table 7), the control run SimCONT performs better than SimH2O in simulating the emission magnitude, especially at UMBS, Harvard Forest and the seasonally dry tropics. The large model/measurement discrepancy (∼factor of 5) at Belgium is because the model classifies the grid cell as deciduous PFT whereas in reality the measurement site is mixed temperate composed of coniferous (Douglas fir) and deciduous (beech). The fraction of electrons available for isoprene synthesis (ε) is a factor of 4 higher for the deciduous PFT than the evergreen PFT (Table 1). In reality, the contribution to the isoprene flux at the Belgium site from coniferous versus deciduous is small but beech is non-emitting for isoprene. We retain the Belgium site in this study because it illustrates the dependency of the model's simulation on the vegetation data set and structure (Schurgers et al., 2011; Arneth et al., 2011).

The simulated isoprene average diurnal cycle is lower at all sites (except Manaus) in SimCONT than SimH2O following the expected GPP response to the presence of water stress. In dry season Manaus, isoprene emission is higher in the

Fig. 9b. (a–c) Comparison between measured and modeled isoprene emission and GPP average diurnal cycles for the time periods indicated in Table 6. Measurements are black lines. Error bars on the measurements represent ± 1 standard deviation. Model results are 10 yr climatological averages for SimCONT (red) and SimH2O (blue). The dotted lines represent model uncertainty due to internal variability in the climate model computed as ± 1 standard deviation for $n = 10$ model run years. **(b)** isoprene mission temperate sites.

water-stressed control run because drought-induced stomatal closure drives higher canopy temperatures that stimulate the isoprene emission. At the North American sites (Harvard Forest and UMBS), the isoprene emission rate is a factor of 3 lower in SimCONT than SimH2O. At these sites, the measured average diurnal cycle lies between the SimCONT and SimH2O climatologies. Interestingly, SimH2O authentically reproduces the GPP average diurnal cycle (Fig. 8c) at both sites (RMSE ~0.1 gC m⁻² h⁻¹, $R^2 > 0.94$). SimCONT GPP average diurnal cycle is too low by a factor of 2–3. It may be tempting to conclude that the water stress function in the control run is too strong at these locations, either because of errors in the climate model's soil moisture simulation (too dry), and/or because of the model's assignment of PFTspecific stress onset and wilting points. Discrepancies in the model's PFT-specific photosynthesis and isoprene emission parameters provide another possible explanation. For example, using a local emission potential (I_s) of 100 µgC g⁻¹ h⁻¹ for the deciduous PFT, more than double the generic value applied in Yale-E2 (Table 1), isoprene emission simulated in LPJ-GUESS is within 5–10 % of measured values at the UMBS site but only about half the magnitude of the measured values at the Harvard Forest 1995 site (Arneth et al., 2007). Increasing I_s to 160 μ gC g⁻¹ h⁻¹ for the Harvard For-

Fig. 9c. (a–c) Comparison between measured and modeled isoprene emission and GPP average diurnal cycles for the time periods indicated in Table 6. Measurements are black lines. Error bars on the measurements represent ± 1 standard deviation. Model results ~ 10 such all $\sim t$ denoted a variability in the climate model computed as ~ 10 standard computed as ~ 10 such as ~ 1000 are 10 yr climatological averages for SimCONT (red) and SimH2O (blue). The dotted lines represent model uncertainty due to internal variability in the climate model computed as ± 1 standard deviation for $n = 10$ model run years. **(c)** GPP temperate sites.

est site in LPJ-GUESS improved the agreement with observations to within 20 %. In contrast, the JULES model used identical I_s values for temperate broadleaf (45 µgC g⁻¹ h⁻¹) and tropical $(24 \mu g C g^{-1} h^{-1})$ as those applied in this study, but overestimated the isoprene emission flux at 6 of the 9 sites (UMBS, Harvard Forest, LV, Montmeyan, Manaus and Santarem) with the maximum discrepancy at La Verdiere (+236 %). In JULES, model isoprene emissions in the middle of the day and in the afternoons are generally higher than observed at all test sites.

Figure 9a and b suggest that the model isoprene emission ramps up too quickly in the morning hours compared to measurements at both tropical and temperate sites. The observed rate of emission decrease in the early evening is better captured by the model. The existence of a circadian rhythm in the PFT-specific fraction of electrons available for isoprene synthesis (ε) provides one possible explanation, such that ε is lower in the morning hours during the first onset of

Site	Linear regression coefficient for 30 min data		RMSE (mgC m ⁻² h ⁻¹)			Maximum in average diurnal cycle	
	SimCONT	SimH2O	SimCONT	SimH2O	Measurement	SimCONT	SimH ₂ O
Costa Rica	0.89	0.90	3.03	3.14	1.32	7.31	7.65
Manaus	0.93	0.93	1.78	1.11	6.41	8.68	6.94
Santarem (wet)	0.82	0.81	3.09	3.20	0.38	6.93	7.00
Santarem (dry)	0.92	0.95	1.44	2.32	3.41	6.13	8.17
Borneo/OP3	0.96	0.96	2.57	3.04	1.65	6.78	7.77
Harvard Forest (1995)	0.67	0.90	1.33	2.62	7.40	3.37	10.23
Harvard Forest (2007)	0.74	0.95	0.81	3.23	5.21	3.37	10.23
UMBS(5yrs)	0.64	0.89	1.21	2.44	4.07	1.71	8.61
La Verdière	0.82	0.90	2.43	4.01	4.01	7.89	11.24
Montmeyan	0.81	0.84	1.78	2.48	8.95	9.40	11.41
Belgium (3 yr)	0.87	0.88	4.22	4.52	0.77	9.25	9.77

Table 7. Statistical performance of model simulated isoprene average diurnal cycles at 9 sites in the tropical and temperate zones.

Table 8. Statistical performance of model simulated GPP average diurnal cycles at 2 temperate sites.

Site	Linear regression coefficient for 30 min data		RMSE ($gC m^{-2} h^{-1}$)		Maximum in average diurnal cycle		
	SimCONT	SimH2O	SimCONT	SimH2O	Measurement	SimCONT	SimH2O
Harvard Forest (2007)	0.97	0.97	0.34	0.09	l.08	0.45	0.95
UMBS $(5\gamma rs)$	0.79	0.94	0.36	0.10	LO2	0.37	0.99

photosynthesis (Hewitt et al., 2011). We will explore the use of a diurnally varying ε in future site-level work.

Based on linear regression (Table 7), the model demonstrates significant skill in capturing the observed variability in the 30 min average diurnal cycle across all sites (64–96 %). The model performance is best in the tropics $(>80\%)$ where there is little difference between SimCONT and SimH2O. In the temperate zone, SimH2O demonstrates superior performance over SIMCONT (e.g. > 90 % versus 70–80 %) in reproducing the diurnal average variability, especially at Harvard Forest and UMBS. Essentially, the model is able to reproduce all of the observed variability in the average diurnal cycle at Santarem (dry), Manaus and Borneo. SimH2O reproduces the diurnal variability better than SimCONT at Harvard Forest (2007) and UMBS (all years). Despite the model's overestimate of emission magnitude at Belgium (because model PFT is deciduous in this location versus mixed temperate at the actual site), the model captures > 87 % of the measured variability in average diurnal cycle.

5 Discussion and conclusions

Isoprene emission is a quintessential quantity in chemistryclimate interactions. In this study, we have implemented a biologically realistic photosynthesis-dependent isoprene emission scheme into a global chemistry-climate model framework. The model provides a new tool that will allow us to improve understanding of the multiple interactions between atmospheric chemistry, land ecosystem physiology and climate at regional and global scales that manifest through BVOC emissions. Such improvements are essential to provide adequate assessment of the climate and air quality benefits of mitigation strategies involving the short-lived climate forcers (Arneth et al., 2009).

Overall, based on comparison with above canopy flux measurements, the model provides an authentic representation of isoprene emission diurnal variability in tropical and temperate ecosystems but is less successful in reproducing the magnitude due to the assignment of PFT-specific isoprene emission potentials, and possibly biases in the climate model's internal meteorology. Current generation global climate models are limited to 5–20 PFTs. To improve simulation of isoprene emission magnitude in global models, more measurements of the plant-to-plant species level variability in isoprene emission (the isoprene emission potential) are needed as well as a way to represent the greater species diversity in models. In addition to accounting for leaf age effects, improvements in simulating isoprene variability may be achieved by including climate-sensitive phenology, variable atmospheric surface $CO₂$ concentrations and the effects of ozone on plant physiology.

In future work, we will use the model to assess the impacts of the photosynthesis-dependent isoprene emission on atmospheric chemical composition and the impacts of global change on the isoprene emission. We will explore the

PFT	$V_{\rm cmax}$ (μ mol m ⁻² s ⁻¹)	Nleaf $(g[N]m^{-2}$ [leaf])	s^*	Swilt	SLA $(m^2$ [leaf] kg ⁻¹ [leaf])
Tundra	21	1.60	0.50	0.30	2.25
Grass	26	3.27	0.30	0.10	11.7
Shrub	17	2.38	0.40	0.22	3.25
Savanna	30	3.12	0.65	0.22	5.1
Deciduous	30	1.07	0.50	0.29	8.3
Evergreen	42	1.80	0.50	0.25	5.9
Rainforest	54	2.70	0.60	0.29	9.9
Crop	30	2.50	0.45	0.27	6.36

Table A1. PFT-specific photosynthesis parameters used in Yale-E2 global carbon-chemistry-climate model.

implementation of a leaf demography scheme (Caldararu et al., 2012) that will allow us to account for the effects of leaf age on the isoprene emission, which has implications for atmospheric chemistry, for instance, timing of the seasonal transition from VOC to NO_x -limited ozone production. We will explore daily and seasonal average isoprene emission model performance using an off-line version of the vegetation model that is driven by meteorology from the GMAO Modern Era-Retrospective Analysis (Rienecker et al., 2011).

We will extend the model to include a photosynthesisbased monoterpene algorithm (Schurgers et al., 2009). A major advantage of the vegetation biophysics in Yale-E2 is the high temporal resolution (30 min) that will allow additional treatment of monoterpene short-term storage in the mesophyll not considered in any previous studies. The short-term storage may be important in simulating a correct monoterpene emission diurnal cycle.

This study has adopted an isoprene emission model that depends on the electron requirement for isoprene synthesis (Niinemets et al., 1999) necessitating an empirical parameterization of the $CO₂$ -inhibition effect. Recently, a simple conceptual model of isoprene emission based on regulation of metabolism in the chloroplast has been proposed (Harrison et al., 2013). This formulation does not require empirical parameterization of the $CO₂$ -inhibition effect. The model needs further testing and development before application at the global scale but does hold significant promise of an even more faithful isoprene emission representation for the new generation Earth-system models.

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