Relationships between photosynthesis and formaldehyde as a probe of isoprene emission

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Abstract

Atmospheric oxidation of isoprene emission from land plants affects radiative forcing of global climate change. There is an urgent need to understand the factors that control isoprene emission variability on large spatiotemporal scales but such direct observations of isoprene emission do not exist. Two readily available global-scale long-term observational-based datasets hold information about surface isoprene activity: gross primary productivity (GPP) and tropospheric formaldehyde column variability (HCHOv). We analyze multi-year seasonal linear correlations between observed GPP and HCHOv. The observed GPP-HCHOv correlation patterns are used to evaluate a global Earth system model that embeds three alternative leaf-level isoprene emission algorithms. GPP and HCHOv are decoupled in the summertime southeast US ($r=0.03$). In the Amazon, GPP-HCHOv are weakly correlated in March-April-May (MAM), correlated in June-July-August (JJA) and weakly anti-correlated in September-October-November (SON). Isoprene emission algorithms that include soil moisture dependence demonstrate greater skill in reproducing the observed interannual seasonal GPP-HCHOv correlations in the southeast US and the Amazon. In isoprene emission models that include soil moisture dependence, isoprene emission is correlated with photosynthesis and anti-correlated with HCHOv. In an isoprene emission model without soil moisture dependence, isoprene emission is anti-correlated with photosynthesis and correlated with HCHOv. Long-term monitoring of isoprene emission, soil moisture and meteorology is required in water-limited ecosystems to improve understanding of the factors controlling isoprene emission and its representation in global Earth system models.
1. Introduction

Isoprene emission, a by-product of photosynthesis, is fundamental in global chemistry-climate interactions. The global annual source strength is estimated at 0.5 Pg C per year (Guenter et al., 2006), which is of comparable magnitude to the present day total (anthropogenic and biogenic) annual source of methane (CH\textsubscript{4}) (Kirschke et al., 2013), and the net carbon dioxide (CO\textsubscript{2}) emission from land use change (Ciais et al., 2013). Isoprene emission rate depends upon ecosystem type, photosynthesis, temperature, and atmospheric CO\textsubscript{2}, and is therefore sensitive to changes in land cover and climate (Monson et al., 2007). In contrast to CH\textsubscript{4} and CO\textsubscript{2}, isoprene is highly reactive in the atmosphere with a lifetime of around only half an hour in the boundary layer. The atmospheric photo-oxidation of isoprene regulates the global budgets and variability of the major short-lived climate pollutants: tropospheric ozone (O\textsubscript{3}), CH\textsubscript{4} and secondary organic aerosol (Arneth et al., 2010; Carslaw et al., 2010). Large-scale perturbations to isoprene emission influence global climate change (Scott et al., 2014; Unger, 2014a). In Earth’s history, plant isoprene emission is recognized as an important terrestrial biogeochemical feedback that influences the global climate sensitivity (Beerling et al., 2007, 2011; Unger and Yue, 2014). Emerging research begins to quantify isoprene’s role as an anthropogenic climate forcing mechanism (Heald and Spracklen, 2015; Unger, 2014b). While short-term (hours to days) weather-related fluctuations in isoprene emission in the temperate zone are well understood (Guenter et al., 1995, 1991), many open questions remain as to the long-term (months to years) factors controlling isoprene emission. A complete understanding of isoprene emission on large spatiotemporal scales is imperative to allow reliable projections of future air quality and global
climate change, and to discern quantitatively the real-world effectiveness of mitigation strategies involving the short-lived climate pollutants.

Two readily available global observational-based datasets do hold information about isoprene emission variability: (i) gross primary productivity (GPP) and (ii) satellite tropospheric formaldehyde (HCHO) columns. GPP is the total amount of CO$_2$ removed from the atmosphere by plant photosynthesis. Isotopic labeling studies have shown that 70-90% of isoprene production is directly linked to photosynthesis that provides energy and precursors for isoprene biosynthesis in the chloroplast (Affek and Yakir, 2003; Delwiche and Sharkey, 1993; Karl et al., 2002). Precipitation controls photosynthesis in more than 40% of vegetated land (Beer et al., 2010). HCHO is a high-yield product of isoprene oxidation and has a lifetime of only a few hours against photolysis and oxidation by the hydroxyl radical (OH) during the day. Other HCHO sources include oxidation from CH$_4$, which provides a slowly varying background of HCHO, oxidation from other volatile organic compounds (VOCs), and direct emission from fires. Precipitation might affect HCHO indirectly by removing reactive carbon, nitrogen oxides and oxidants, thus dampening atmospheric photochemistry. Since isoprene emission frequently dominates the non-methane VOC budget over continental land, HCHO columns have been used as a direct proxy for inferring isoprene emissions (Barkley et al., 2008; Barkley et al., 2013; Fu et al., 2007; Millet et al., 2008; Palmer et al., 2003, 2006).

Neither GPP nor HCHO columns offer a perfect indicator of isoprene emission variability. In the case of GPP, incomplete coupling between isoprene emission and photosynthesis occurs due to the different temperature optimums of the processes, response to short-term drought and elevated
atmospheric CO$_2$, and onset time in the deciduous biome (Harrison et al., 2013). The optimal temperature for photosynthesis is around 25°C while isoprene emission has a higher thermal optimum of 35-40°C. In the case of HCHO columns, limitations in use as a direct proxy for isoprene include: (1) uncertainties associated with the HCHO vertical column retrieval (Barkley et al., 2012; Hewson et al., 2013), (2) distinguishing the component of the HCHO column produced solely from isoprene oxidation, and (3) uncertainties in isoprene oxidation chemistry.

Isoprene provides an intrinsic linkage between GPP and atmospheric HCHO. A recent study found a strong intra-seasonal correlation between satellite HCHO columns and canopy temperature but a weak correlation or even anti-correlation with GPP in 22 regions selected to minimize interference from fires (Foster et al., 2014). In that study, HCHO columns were assumed to be a direct proxy for surface isoprene emission. Soil moisture availability was not explicitly considered as a driving variable even though water availability and canopy temperature are tightly coupled through stomatal conductance and the canopy energy balance. Accounting for soil moisture dependence of isoprene emission decreases the global source strength by 25-30% (Muller et al., 2008; Unger et al., 2013).

Here, we investigate the multi-year (2005-2011) seasonal relationships between global observational datasets of FLUXNET-derived GPP and fire-screened satellite HCHO columns as a probe of isoprene emission on longer seasonal to interannual temporal scales. We assume that observed GPP and HCHO columns hold quantitative information about isoprene emission variability, but we do not assume that either is a direct proxy. The study proceeds in three steps. First, we calculate the covariance of the observational-based GPP and satellite HCHO columns
with key meteorological variables. Then, we compute the linear correlation between GPP and HCHO observations. Finally, we use the observed GPP-HCHO relationships to evaluate a global Earth system model that incorporates three alternative isoprene emission algorithms. The models are used to interpret the observed GPP-HCHO relationships. We focus our discussion on the major isoprene emitting source regions: the southeastern US [31 to 35°N; -94 to -79°E] and the Amazon [-15°S to 3°N, -76° to -54°E].

2. Methods

2.1 Observational and reanalysis datasets

In this study we apply datasets of observational-derived GPP, satellite-based tropospheric HCHO columns and meteorology reanalysis. The monthly-mean global GPP dataset is generated using data orientated diagnostic upscaling of site-level derived GPP from FLUXNET (Beer et al., 2010; Bonan et al., 2011; Jung et al., 2011) and is available for years 1982-2011 with native resolution of 0.5°×0.5° latitude by longitude. The main steps of the upscaling procedure are processing FLUXNET observational data and calculating GPP for each site, training model-tree-ensembles (MTEs) for each GPP using site-level explanatory variables, and applying the established MTEs using global gridded dataset of the same explanatory variables to obtain the global GPP estimates (Jung et al., 2011). Twenty-nine explanatory variables are used to train the MTE, including the fraction of absorbed photosynthetically active radiation (fAPAR), precipitation, temperature and other climate and land cover data (Jung et al., 2011). The uncertainties are mainly from but not limited to (1) measurement of eddy covariance fluxes
(Lasslop et al., 2008; Richardson et al., 2006), (2) the choice of explanatory variables (Jung et al., 2011), (3) gap filling and extrapolation to different environmental domains and temporal periods (Jung et al., 2009), (4) global gridded explanatory variables (Hicke, 2005; Zhao et al., 2006). The derived GPP in tropical and subtropical regions is less well constrained with observations and has larger uncertainties compared to the mid-latitudes (Beer et al., 2010; Jung et al., 2011).

The fire-screened monthly mean tropospheric HCHO vertical columns are retrieved by the Ozone Monitoring Instrument (OMI) over 2005-2013. We compute the fire-screened tropospheric HCHO vertical columns from retrieved slant columns provided in the official NASA OMI product (González Abad et al., 2015), in a three-step process. First, we apply our own reference sector correction to normalize the HCHO columns, on a daily basis. This is a standard technique used in many studies to remove retrieval biases (e.g., Barkley et al., 2013; González Abad et al., 2015; Marais et al., 2012). Here we compute the median OMI slant columns \( \Omega_{SM} \) in 1° latitude bins over the remote Pacific Ocean (140-160°W), and subtract this latitudinal bias from all retrieved slant columns \( \Omega_s \). We then re-normalize the vertical columns \( \Omega_v \) by adding a model HCHO latitudinal background \( \Omega_{VB} \), provided by the NASA ModelE2-YIBs simulation (described in Section 2.2), as follows:

\[
\Omega_v = \frac{\Omega_{SM}}{AMF} + \Omega_{VB}
\]  

(1)

where \( AMF \) is the air mass factor, defined as the ratio of the slant and vertical columns. Second, we generate AMF look-up tables using monthly averaged HCHO profiles from the global earth
system model NASA ModelE2-YIBs (three sub-versions, as described in Section 2.2), appropriate to the OMI’s overpass time. The AMF calculation is the same as that described in Barkley et al. (2013), with the exception that no aerosol correction is applied as model aerosol optical depth (AOD) profiles were not available. Third, we then apply the AMFs to the corrected slant columns, using Eq. (1), and average the resulting vertical columns onto a generic global 0.5°×0.5° latitude-longitude grid. We additionally filter the OMI data, excluding scenes with ≥40% cloud cover and that do not meet standard quality checks (González Abad et al., 2015); observations affected by the documented OMI row anomaly are also discarded. To remove biomass burning contamination from the data, we adopt the method devised by Barkley et al., (2013) which excludes fire-affected scenes using Advanced Along-Track Scanning Radiometer (AATSR) and Moderate Resolution Imaging Spectroradiometer (MODIS) active burning detections. Individual observations are discarded if a fire occurs in the 0.5° grid-cell in which it falls, or those immediately adjacent (within ± 2 grid-cells), of both the current or preceding day. The uncertainties on the gridded OMI vertical columns, mainly due to cloud contamination, the a priori modeled isoprene emissions and the HCHO vertical column retrieval, are estimated at 5-20% (Barkley et al, 2013). To ensure consistency in our satellite-model comparisons, the reference correction and AMFs are recomputed using HCHO profiles from the appropriate model simulation. In our subsequent analysis, we use the HCHO column variability (HCHOv), which is defined as the anomaly between local and zonal mean of the gridded fire-screened HCHO tropospheric column concentrations for each month, to explore its climatic covariance and relationship with GPP, and compare against the NASA ModelE2-YIBs output. There are two main limitations in using HCHOv as a proxy for isoprene emission: (1) HCHO from CH₄ oxidation is not strictly zonally uniform, thus HCHOv does not purely represent the influence of
the non-methane VOCs; (2) HCHO\textsubscript{v} is dominated by isoprene emission but their relationship is smeared by other VOCs such as biogenic terpenes and anthropogenic VOCs.

We use monthly-mean meteorological variables, including surface skin temperature ($T_s$), downward short wave radiation (SW), photosynthetically active radiation (PAR), and precipitation (P), from the NASA Modern Era Retrospective-Analysis for Research and Applications (MERRA) (Rienecker et al., 2011). The spatial resolution of the MERRA data is $0.5^\circ \times 0.667^\circ$ latitude by longitude and the temporal availability is 1979 to present.

All of the monthly average observational datasets are linearly interpolated to $2.0^\circ \times 2.5^\circ$ latitude by longitude spatial resolution.

2.2 Global Earth system model (NASA ModelE2-YIBs)

This study applies the NASA GISS ModelE2 global chemistry-climate model at $2^\circ \times 2.5^\circ$ latitude by longitude horizontal resolution with 40-vertical layers extending to 0.1 hPa (Schmidt et al., 2014). The Yale Interactive Terrestrial Biosphere Model (YIBs) is embedded inside NASA ModelE2 in a framework known as NASA ModelE2-YIBs. The global climate model provides the meteorological drivers for the vegetation biophysics. The land-surface hydrology submodel provides soil characteristics to the vegetation physiology in each grid cell. The model framework fully integrates the land biosphere-oxidant-aerosol system such that these components interact with each other and with the physics of the climate model at the 30-minute integration time step. The atmospheric composition model has been well tested against observations and compared
with other models e.g. (Koch et al., 2010; Myhre et al., 2013; Shindell et al., 2013a, 2013b; Stevenson et al., 2013).

The vegetation is described using eight plant functional types (PFTs): tundra, C3 and C4 grassland, shrub, deciduous, tropical rainforest, evergreen, and crop. Present-day vegetation cover fractions are derived from Moderate Resolution Imaging Spectroradiometer (MODIS) satellite data as used in the Community Land Model and converted to the eight PFTs here (Lawrence and Chase, 2007). Gridded spatially-varying PFT-specific leaf area index (LAI) is derived from Advanced Very High Resolution Radiometer (AVHRR) satellite data and linearly interpolated into daily values (Lawrence and Chase, 2007).

The canopy biophysical fluxes are computed using the well established Farquhar leaf model of photosynthesis (von Caemmerer and Farquhar, 1981; Farquhar et al., 1980) and the stomatal conductance model of Ball and Berry (Ball et al., 1987). The model vertically stratifies each canopy into an adaptive number of layers (typically 2-16) that distinguish LAI profiles for sunlit and shaded leaves (Friend and Kiang, 2005).

2.2.1 Isoprene emission algorithms

NASA ModelE2-YIBs incorporates two conceptually different leaf-level isoprene emission algorithms that are embedded within the exact same host simulation framework: (1) Y-PS: isoprene emission is calculated as a function of electron transport-limited photosynthesis, intercellular and atmospheric CO2 and canopy temperature (Unger et al., 2013) and (2) Y-
MEGAN: isoprene emission is calculated using empirical functions of canopy temperature and light commonly applied in The Model of Emissions of Gases and Aerosols from Nature (MEGAN) (Guenther et al., 1995). MEGAN is the most widely used system for estimating isoprene emissions from terrestrial ecosystems (Guenther et al., 2012). We test a third isoprene emission algorithm identical to Y-MEGAN but with an additional empirical multiplier to account for soil moisture availability (Y-MEGAN-SM).

In Y-PS, leaf-level isoprene emission is modeled as follows:

\[ I_{emis} = \varepsilon \cdot J_e \cdot \delta \cdot \tau \]  

(2)

where \( \varepsilon \) is the PFT-specific isoprene emission potential in units of fraction of electrons available for isoprene synthesis. \( J_e \) is the electron transport limited photosynthesis rate in units of \( \mu \text{mol m}^{-2} \text{[leaf]} \text{s}^{-1} \). \( J_e \) is a linear function of the incident photosynthetically active radiation (PAR) and the internal leaf CO\(_2\) concentration (\( C_i \)):

\[ J_e = a_{leaf} \cdot PAR \cdot \alpha_{qe} \cdot \frac{C_i - \Gamma^*}{C_i + 2\Gamma^*} \]  

(3)

where \( a_{leaf} \) is the leaf specific light absorptance, \( \alpha_{qe} \) is the intrinsic quantum efficiency for photosynthetic CO\(_2\) uptake in photosystem II (a product of the fraction of absorbed light that reaches photosystem II and the CO\(_2\) per absorbed photon), and \( \Gamma^* \) is the CO\(_2\) compensation point in the absence of non-photorespiratory respiration (Collatz et al., 1991).
The δ term in equation (2) translates the electron flux into isoprene equivalents given by equation (4) detailed in (Niinemets et al., 1999; Pacifico et al., 2011):

\[ \delta = \frac{C_l - I^*}{6(4.67C_l + 9.33I^*)} \]  

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

\[ \tau = \exp[0.1(T - T_{ref})] \]  

where T is leaf temperature in °C and \( T_{ref} \) is the leaf temperature under standard conditions (30°C).

In Y-MEGAN, leaf-level isoprene emission is modeled following:

\[ I_{emis} = E \cdot C_T \cdot C_L \]  

where E is the PFT-specific isoprene emission potential in units of µmolC m\(^{-2}\) s\(^{-1}\); \( C_T \) and \( C_L \) are defined as follows:

\[ C_T = \frac{\exp^{C_T(I_T - T_{Ks})}}{1 + \exp^{C_T(I_T - T_{M})}} \]  

\[ C_L = \frac{\exp^{C_L(I_L - T_{Ks})}}{1 + \exp^{C_L(I_L - T_{M})}} \]
and:

\[
C_L = \frac{\alpha C_{L1} \text{PAR}}{\sqrt{1 + \alpha^2 (\text{PAR})^2}}
\]  

\( T_K \) is the leaf temperature in Kelvin, \( T_{Ks} \) is the leaf temperature at standard conditions (=303 K), \( R \) is the ideal gas constant (=8.314 J K\(^{-1}\) mol\(^{-1}\)); \( C_{T1} \) (=95,000 J mol\(^{-1}\)), \( C_{T2} \) (=230,000 J mol\(^{-1}\)), \( T_M \) (=314 K), \( \alpha \) (=0.0027) and \( C_{L1} \) (=1.066) are empirical coefficients.

Y-PS and Y-MEGAN use identical PFT-specific isoprene emission potentials converted to the relevant units for \( \varepsilon \) (unitless) and \( E \) (\( \mu \text{molC m}^{-2} \text{s}^{-1} \)), presented here in units of \( \mu \text{gC g}^{-1} \text{h}^{-1} \): tundra=0, C3 grassland=16, C4 grassland=0, shrub=16, deciduous=45, tropical rainforest=24, evergreen=8, and crop=0 (Guenther, 2007; Lathiere et al., 2006). An additional multiplier to account for the long-term atmospheric CO\(_2\)-sensitivity of isoprene emission is applied to both isoprene models that is normalized to 1.0 for the present-day atmospheric CO\(_2\) levels used in this study (Arneth et al., 2007).

Y-MEGAN-SM is identical to Y-MEGAN but includes an additional multiplier to account for soil moisture availability following the approach used in the coupled photosynthesis-stomatal conductance vegetation biophysics submodel. The multiplier value is between 0 and 1 and 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 \( (s_{\text{wilt}}) \) (Porporato et al., 2001). The multiplier value is reduced linearly between the PFT-specific values...
of $s^*$ and $s_{\text{wilt}}$ based on the climate model’s soil water volumetric saturation in six soil layers. Values of $s^*$ and $s_{\text{wilt}}$ are documented in Unger et al. (2013).

The leaf-level isoprene emissions in each isoprene scheme are upscaled to the canopy level using the YIBs canopy vertical stratification and integration scheme (Unger et al., 2013). The canopy level isoprene fluxes are passed to the model’s atmosphere through the land-surface scheme on the 30-minute integration time step of the global climate model. Thus, the three isoprene emission algorithms ‘see’ the exact same PFT-specific isoprene emission potentials (basal rates), vegetation input data and meteorology, and apply the exact same upscaling from leaf to canopy.

In Y-PS, the light dependence occurs through the linkage to photosynthesis; in Y-MEGAN and Y-MEGAN-SM, isoprene emission is directly related to PAR. All three models are directly linked to canopy temperature. In Y-PS, soil moisture dependence occurs through the linkage to photosynthesis; Y-MEGAN has no direct soil moisture dependence but captures indirect effects through canopy temperature changes; and Y-MEGAN-SM has soil moisture dependence through the additional empirical multiplier.

### 2.2.2 Simulations

We perform three NASA ModelE2-YIBs simulations representative of present day (2000s) climatology for each of the isoprene emission schemes (Y-PS; Y-MEGAN; Y-MEGAN-SM). Decadal average (1996-2005) monthly-varying sea surface temperature and sea ice climatology from the HadSST2 dataset provide the physical climatic boundary conditions for the simulations (Rayner et al., 2006). The present day anthropogenic trace gas and aerosol emissions are
prescribed to year 2000 values from the inventory developed for IPCC AR5 (Lamarque et al., 2010). Atmospheric levels of long-lived greenhouse gases are prescribed to $\text{CO}_2 = 370$ ppmv, $\text{CH}_4 = 1733$ ppbv in Southern Hemisphere (SH) and 1814 ppbv in Northern Hemisphere (NH), $\text{N}_2\text{O} = 316$ ppbv. Integrations of eleven model years are completed for all control and sensitivity simulations; the first two years of the simulations are discarded as spin-up and the remaining nine years are used for analyses.

3. Results

The goal of this work is to investigate the large-scale observationally-derived climatic covariance and correlations in the photosynthesis-HCHO system, and to assess the models’ ability to reproduce these observationally-derived biosphere-atmosphere system sensitivities and to expose the implications for isoprene emission. Therefore, instead of direct comparison between simulated and observed GPP and HCHO columns, we conduct a multiple linear regression (MLR) analysis in Section 3.1 and a correlation analysis in Section 3.2, and use the observed climatic covariance and correlations to evaluate the NASA ModelE2-YIBs model embedded with three isoprene algorithms. The direct comparison results are included in the supplementary materials for reference: simulated and FLUXNET-derived GPP are of comparable absolute amounts (Figure S1); while simulated tropospheric HCHO columns are considerably higher than that obtained from the OMI retrieval by about a factor of 2 (Figure S2), which is likely due to the large uncertainties in the models as well as the satellite retrieval. In the following analysis, we apply a minimum threshold for GPP (monthly mean GPP > 0.01 g[C] m$^{-2}$ day$^{-1}$) to avoid
inclusion of meaningless noise, especially in boreal winter (December-January-February) when most NH regions have very low GPP and isoprene emissions.”

Using the exact same vegetation input data, meteorology and PFT-specific basal rates, the three isoprene algorithms give substantially different annual global isoprene emission strengths: 382 Tg[C] year\(^{-1}\) for Y-PS, 452 Tg[C] year\(^{-1}\) for Y-MEGAN and 263 Tg[C] year\(^{-1}\) for Y-MEGAN-SM. As shown in Fig. 1 (left column), isoprene emission in Y-MEGAN is lower in NH mid-latitudes than Y-PS, and is stronger in the tropics. Y-MEGAN-SM has lower isoprene flux than Y-MEGAN, especially in the dry subtropics in South America, Africa and Australia. Yet, the three OMI HCHO column datasets that use different AMFs for the three isoprene models show similar distribution (Fig. 1, right column). Further analysis of OMI HCHO column datasets, including the MLR of HCHO\(_v\) with meteorological variables in Section 3.1 and the observation correlation between GPP and HCHO\(_v\) in Section 3.2, show no difference among the three HCHO datasets. Therefore, in the following analyses, results shown are based on OMI-HCHO processed using Y-PS AMFs.

3.1 Meteorological drivers of GPP and HCHO\(_v\)

The regionally averaged meteorological variables \(T_s\), PAR, SW and P for the period 2005-2011 from MERRA reanalysis and the climate model NASA ModelE2-YIBs are summarized in Table 1. In MERRA, the average \(T_s\) values for March-April-May (MAM), June-July-August (JJA) and September-October-November (SON) in key regions are (in °C): 18.0±0.8, 26.8±0.5, 18.6±0.8 (southeast US); 23.5±0.5, 23.7±0.4, 25.3±0.6 (Amazon). Seasonal average \(T_s\) in southeast US in
JJA and in the Amazon in SON slightly exceed the photosynthetic thermal optimum (25°C). No vegetated region on the planet has a seasonal average $T_s$ that exceeds the thermal optimum of isoprene emission (35-40°C).

We perform a multiple linear regression analysis of FLUXNET-derived GPP and OMI-retrieved HCHOv against major meteorological variables to examine their climatic covariance and to determine the most important meteorological drivers in different regions and different seasons. Figure 2 shows the MLR results for monthly mean GPP (1982-2011) and HCHOv (2005-2013) in three seasons (MAM, JJA, SON) against $T_s$, PAR (SW for HCHOv), and $P$. MLR of GPP and HCHOv using 2005-2011 data (the overlapped time range) yield very similar results. A provocative implication is that the effects of decadal climate change (e.g. the rapid global rise in atmospheric CO$_2$ since 1982) do not appear to influence GPP’s and HCHOv’s seasonal climatic covariance in the contemporary period. The computed standardized partial regression coefficients ($\beta$-coefficients) represent the rate of change in the dependent variable for a unit change in the independent variable with all other independent variables held constant. The coefficients have been standardized in units of standard deviation, thus they can be directly compared with each other to determine the relative importance of the different driving variables. The standardized partial regression coefficients of GPP and HCHOv associated with $T_s$, PAR (SW for HCHOv) and $P$ are denoted by $\text{GPP}_\beta$-$T_s$, $\text{GPP}_\beta$-PAR, $\text{GPP}_\beta$-$P$ and $\text{HCHOv}_\beta$-$T_s$, $\text{HCHOv}_\beta$-SW, $\text{HCHOv}_\beta$-$P$. GPP $\beta$-coefficients are statistically significant ($p<0.05$) over most vegetated regions of the planet. HCHOv $\beta$-coefficients are not significant to 95% confidence level anywhere on the planet. It is not surprising that the FLUXNET-derived GPP climatic covariance results have high statistical significance values because this product is an
empirically upcaled dataset based on machine learning techniques (see Section 2.1) using a
large set of climatic and land cover explanatory variables, and the driving variables $T_s$, PAR and
$P$ used to determine the MLR in this study are an important subset of the original explanatory
variables. In contrast, remotely-sensed HCHO columns are relatively noisy due to the satellite
retrieval method (Palmer et al., 2001; De Smedt et al., 2008). Other reasons for the differences in
statistical significance include: satellite-based HCHO columns have many missing values due to
product quality control (e.g. contamination by clouds) and the biomass burning removal (see
Section 2.1); the GPP dataset has a longer record (1982-2011) than the HCHO dataset (2005-
2013); unlike GPP which has a simple near-parabolic relationship with $T_s$, HCHO dependence
on $T_s$ is more complex. For instance, increasing $T_s$ promotes isoprene emission and oxidation to
HCHO, but also accelerates the chemical destruction of HCHO (see supplementary and Fig. S4).

The regionally averaged $\beta$-coefficients over southeast US [31 to 35°N; -94 to -79°E] and the
Amazon [-15 to 3°N; -76 to -54°E] are summarized in Table 2. GPP is strongly positively related
to $T_s$ in the NH springtime and summertime high-latitudes (Fig. 2). In NH mid-latitudes in
summer, where $T_s$ values approach and/or exceed the photosynthetic thermal optimum,
sensitivity to $T_s$ decreases dramatically. In the southeast US, GPP$_\beta$$_T_s$ drops from 0.58 in
spring to 0.03 in summer. In NH subtropical and semi-arid regions, there is a marked
anticorrelation with $T_s$ in the NH summer (GPP$_\beta$$_T_s$ < -0.3). In contrast, HCHOv is generally
positively correlated with $T_s$ across all continents and seasons. The averaged HCHOv$_\beta$$_T_s$
values in the southeast US are 0.36, 0.31 and 0.53 in MAM, JJA and SON. In the Amazon, the
temperature dependence of both GPP and HCHOv are positive but weak.
GPP has a strong positive relationship with PAR in NH mid-latitudes (especially in SON) and in tropical continents in all seasons (Fig. 2). The spatial pattern of HCHOv dependence on SW is extremely patchy because HCHO can be both formed and destroyed by photolysis. In the southeast US, GPP_β_PAR are 0.44, 0.41, 0.51 in MAM, JJA and SON, whereas HCHOv_β_SW are -0.02, 0.16 and -0.18; the Amazon also shows relatively strong positive light dependence of GPP (0.46, 0.57, 0.17). In the Amazon, HCHOv displays no apparent relationship with SW in MAM and SON but a positive relationship in JJA (0.00, 0.31, 0.01).

The relationship between GPP and precipitation is always positive over heavily vegetated regions. GPP_β_P values tend to be weaker than GPP_β_Ts and GPP_β_PAR values in the NH middle to high latitudes, but much stronger in the tropical rainforest regions in MAM and SON. In the tropics, precipitation stimulates GPP significantly in MAM and SON (GPP_β_P=0.70 in MAM and 0.50 in SON). In contrast, there is no detectable relationship between precipitation and HCHOv in this region in MAM and JJA, but a strong anti-correlation in SON. Precipitation dampens local photochemistry by removing reactive carbon, nitrogen compounds and oxidants. Although wet deposition is not a major sink for HCHO due to the relatively low Henry’s Law coefficient, previous studies have found an anti-correlation with precipitation in highly polluted regions (Báez et al., 1993).

### 3.2 Observed GPP-HCHOv correlation

Figure 3 shows the Pearson’s correlation coefficient (r) between monthly mean observational GPP and HCHOv for each season calculated using the 2005-2011 data. We show results where
FLUXNET-GPP is greater than 0.01 g[C] m\(^{-2}\) day\(^{-1}\) for the latitude range -50°S to +50°N (except in boreal summer) because the satellite HCHO columns have known large biases in high-latitudes under limited-light conditions (De Smedt et al., 2008; Wittrock et al., 1997). The observed GPP-HCHO\(v\) correlation varies strongly with latitude and season. Regionally averaged seasonal correlation values for the southeast US and the Amazon are shown in Table 3. The southeast US shows a significant GPP-HCHO\(v\) coupling in transition seasons \((r=0.24\) in boreal spring and \(r=0.25\) in fall, \(p<0.05\)), which is likely driven by their covariance with temperature. In boreal summer, this positive correlation signal moves northward to NH high-latitudes where boreal forests emit terpenoids. GPP and HCHO\(v\) in the summertime southeast US are almost decoupled with a very weak anti-correlation signal \((r=-0.03)\). Similar decoupling or weak anti-correlation occurs in the tropics all the year round except in the Amazon in JJA \((r=0.33)\).

### 3.3 Model GPP-HCHO\(v\) correlation

We examine the simulated GPP-HCHO\(v\) correlations in NASA ModelE2-YIBs for the three isoprene emission algorithms: Y-PS, Y-MEGAN and Y-MEGAN-SM. Overall, the simulated GPP-HCHO\(v\) \(r\) values are stronger than the observed values everywhere on the planet. Generally, overestimates of GPP-HCHO\(v\) \(r\) values in the models may be due to over-simplified parameterizations of biogeochemical processes and photochemical oxidation mechanisms, missing, possibly important but highly uncertain, processes in the models, for instance nutrient availability, and the use of generic PFT-specific isoprene emission potentials. The three models successfully reproduce the GPP-HCHO\(v\) correlation pattern in the NH temperate spring and fall transition seasons that is likely driven by covariance with temperature (Fig. 4(a)). They broadly
capture the observed GPP-HCHOv spatial patterns in the tropics in MAM and SON, but not in JJA. The models’ overestimate of the positive correlation in southeast US in spring and fall may be because the algorithms do not include the delayed onset in spring or earlier shutdown of isoprene emission before senescence. Regionally averaged model correlation results for the southeast US and the Amazon are compared with the observational results in Table 3. In contrast to the observed GPP-HCHOv decoupling (no correlation) in the summertime southeast US, the models simulate anti-correlation but to different extents: \( r = -0.19 \) (Y-PS); \( r = -0.62 \) (Y-MEGAN); \( r = -0.37 \) (Y-MEGAN-SM). In the Amazon, Y-PS and Y-MEGAN-SM reproduce the observed GPP-HCHOv correlations in MAM and SON but are unable to reproduce the observed strong positive correlation there in JJA. Y-MEGAN fails to reproduce the seasonal observed GPP-HCHOv correlations in the Amazon; for this model, GPP-HCHOv are anti-correlated in JJA \( (r = -0.08) \) where observed GPP-HCHOv \( r = 0.33 \); and strongly anti-correlated in SON \( (r = -0.51) \) where observed GPP-HCHOv \( r = -0.09 \). In the Amazon in JJA, GPP is strongly related to PAR; similarly HCHOv is related to SW (Section 3.1).

Poor performance of all models in Amazon JJA may be due to the global climate model’s simulation of meteorology. Simulated \( T_s \) \( (26.4 \pm 0.3^\circ C) \) in the Amazon JJA is 2–3°C higher than the MERRA \( T_s \) \( (23.7 \pm 0.4^\circ C) \) and exceeds the GPP thermal optimum \( (25^\circ C) \). This temperature overestimation likely contributes to the non-real decoupling or weak anti-correlation between GPP and HCHOv in the three models.

To probe the underlying causes of the GPP-HCHOv relationships, we examine the model correlations between isoprene emission (ISOPe) and GPP, and between ISOPe and HCHOv.
shown in Fig. 4(b) and (c). Regionally averaged values for the southeast US and the Amazon are compared in Table 3. It is apparent that the GPP-HCHOv relationships are driven by different underlying causes contingent upon whether the isoprene emission algorithm includes soil moisture dependence. Focusing on southeast US, Y-PS indicates linear coupling between GPP and ISOPe ($r=0.94\pm0.07$), and only a weak or even anti-correlation between ISOPe and HCHOv in summertime southeast US ($r=-0.03\pm0.31$). In contrast, Y-MEGAN indicates strong coupling between ISOPe and HCHOv ($r=0.73\pm0.09$), but anti-correlation between GPP and ISOPe in the summer ($r=-0.39\pm0.23$).

In Y-PS, anti-correlation between GPP and HCHOv is determined by the anti-correlation between ISOPe and HCHOv. On interannual seasonal time scales, precipitation positively stimulates GPP but has no direct impact on HCHOv, which is predominantly controlled by temperature (see Supplement). Precipitation may dampen photochemistry by limiting OH and O($^1$D) concentration, thus may have an indirect impact on both formation and destruction of HCHO. Photochemical production and loss of HCHO strongly depend on temperature and light independent of isoprene emission (e.g. Seinfeld and Pandis, (2006); Fig. 2(b); Fig. S4). New research is showing that HCHO column variation reflects variation of OH production rather than isoprene emission variability, especially in low OH regions (Dr. L. Valin, Columbia University, personal communication). Furthermore, HCHO may be influenced by emission and oxidation of non-isoprene VOCs. In Y-MEGAN, the anti-correlation between GPP and ISOPe drives the GPP-HCHOv anti-correlation in this model under conditions when the thermal optimum of photosynthesis has been exceeded. Y-MEGAN-SM displays more Y-PS-like behavior, a correlation between GPP and ISOPe, but anti-correlation between ISOPe and HCHOv in
summertime southeast US and in MAM and SON in the Amazon. Since the only difference between Y-MEGAN-SM and Y-MEGAN is the soil moisture dependence of isoprene emission, this result suggests the importance of water availability as a control on the photosynthesis-ISOPe-HCHO system: all the three processes are strongly influenced by temperature, but the dependence on soil moisture determines the summertime covariance of photosynthesis and isoprene variability, which can over-ride their anti-correlation due to different thermal optima. The relative lack of sensitivity of HCHOv to water availability and precipitation leads to weaker correlation or even anti-correlation behavior between ISOPe and HCHOv.

4. Discussion and conclusions

We find that all three models reproduce the observed NH mid-latitude GPP-HCHOv strong correlation in spring and fall, but predict anti-correlation in summer when the observations suggest decoupling. The underlying causes for the predicted relationships are isoprene-algorithm-dependent. In the isoprene algorithms that account for soil moisture dependence (Y-PS and Y-MEGAN-SM), interannual seasonal isoprene emission variability is tightly linked to photosynthesis but anti-correlated with HCHO variability; the dependence on soil moisture determines the summertime covariance of isoprene emission and photosynthesis, which override their opposite response to high temperature. While in Y-MEGAN, isoprene emission is anti-correlated with photosynthesis at high temperatures due to their different thermal optima, and is strongly correlated with HCHO variability. These results suggest water availability could be an important driver of isoprene emission on intraseasonal to interannual time scales.
Multiple field experiments have studied the isoprene response to water deficit conditions on different time scales. Short-time mild drought stress on a time scale of a few days affects stomatal conductance and thus the rate of photosynthesis, while does not essentially diminish isoprene emission because photosynthetic electron transport is not inhibited (Fall and Monson, 1992; Niinemets, 2010). Several studies found increases in isoprene emission during the initial stages of mild drought conditions (e.g. Brilli et al., 2007; Pegoraro et al., 2004; Sharkey and Loreto, 1993). Severe drought or prolonged moderate drought conditions on time scales of weeks do result in significant reductions in isoprene emission that are presumably due to decreased leaf carbon availability following sustained reductions in photosynthetic rate (e.g. Brüggemann and Schnitzler, 2002; Funk et al., 2005; Sharkey and Loreto, 1993). Therefore, on the short time scales of a few days, there is a lag between isoprene emission and photosynthetic rate in response to water stress. On longer time scales (weeks to months), isoprene emission is tightly coupled with photosynthesis, both of which are limited by soil moisture deficit. Recent studies have shown the importance of water availability on photosynthesis on interannual scales: Jung et al. (2011) suggest the interannual variability of GPP in semi-arid to semi-humid regions is more sensitive to precipitation rather than temperature; Beer et al. (2010) find that GPP over 40% of the vegetated land is associated with precipitation. Therefore, despite the current lack of direct observations to constrain the soil moisture impact on the interannual variability of isoprene emission, we argue that water availability is likely to be a critical factor regulating isoprene emission on longer times.

This research raises more questions about long-term isoprene emission variability than it answers. Ground truthing of the findings is impeded by the lack of long-term isoprene emission
flux tower and meteorology measurements in water-limited ecosystems. However, our results do suggest that water availability may be an important driver of vegetation-chemistry-climate interactions under future global change. A corollary is that on longer time scales (seasonal, annual, decadal), GPP may be a more reliable indicator of surface isoprene emission than HCHOv. The soil moisture dependence of isoprene emission warrants further research. Long-term direct measurements of isoprene emission co-located with meteorological monitoring are essential to provide more information on the extent of water dependence of isoprene. Global Earth system models used to study long-term changes in isoprene emission should include soil moisture dependence. Currently, soil moisture is poorly represented in land-surface and climate models (Koster et al., 2009). The recent launch of the NASA Soil Moisture Active Passive instrument will produce global maps of soil moisture and was designed to help improve understanding of carbon and water cycles. Inadvertently, this dataset may also help improve understanding of isoprene emission and atmospheric chemistry.

Author contribution

N. U. and Y. Z. designed this study and developed the model code. Y. Z. performed the simulations and analysis. M. P. B. processed the satellite-based formaldehyde data. X. Y. contributed to the model development. Y. Z., N. U. and M. P. B. wrote the manuscript.

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Oinas, V., Olonso, A. O., Perlwitz, J. P., Puma, M. J., Putman, W. M., Rind, D., Romanou, A.,
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Spectroradiometer (MODIS) terrestrial primary production to the accuracy of meteorological
Table 1. Regionally averaged meteorological variables with standard deviation from MERRA reanalysis and NASA ModelE2-YIBs in the southeast US and the Amazon.

<table>
<thead>
<tr>
<th></th>
<th>Southeast US</th>
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<tbody>
<tr>
<td></td>
<td>T&lt;sub&gt;s&lt;/sub&gt; (°C)</td>
<td>PAR (W m&lt;sup&gt;-2&lt;/sup&gt;)</td>
<td>SW (W m&lt;sup&gt;-2&lt;/sup&gt;)</td>
<td>P (mm day&lt;sup&gt;-1&lt;/sup&gt;)</td>
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<tr>
<td><strong>MERRA</strong></td>
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</tr>
<tr>
<td>MAM</td>
<td>18.0 ± 0.8</td>
<td>110.1 ± 3.4</td>
<td>250.0 ± 8.3</td>
<td>2.6 ± 0.7</td>
</tr>
<tr>
<td>JJA</td>
<td>26.8 ± 0.5</td>
<td>108.5 ± 3.7</td>
<td>237.8 ± 9.0</td>
<td>4.8 ± 0.5</td>
</tr>
<tr>
<td>SON</td>
<td>18.6 ± 0.8</td>
<td>80.7 ± 4.3</td>
<td>182.3 ± 10.5</td>
<td>2.8 ± 0.7</td>
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<tr>
<td><strong>ModelE2-YIBs</strong></td>
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<tr>
<td>MAM</td>
<td>18.6 ± 0.8</td>
<td>106.8 ± 2.3</td>
<td>237.4 ± 5.0</td>
<td>4.2 ± 0.5</td>
</tr>
<tr>
<td>JJA</td>
<td>26.8 ± 0.4</td>
<td>118.7 ± 1.5</td>
<td>263.7 ± 3.4</td>
<td>4.5 ± 0.6</td>
</tr>
<tr>
<td>SON</td>
<td>20.5 ± 1.3</td>
<td>82.1 ± 1.9</td>
<td>182.4 ± 4.2</td>
<td>2.3 ± 0.7</td>
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<tr>
<td><strong>Amazon</strong></td>
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<tr>
<td><strong>MERRA</strong></td>
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<tr>
<td>MAM</td>
<td>23.5 ± 0.5</td>
<td>89.9 ± 2.7</td>
<td>193.4 ± 6.0</td>
<td>7.9 ± 0.4</td>
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<tr>
<td>JJA</td>
<td>23.7 ± 0.4</td>
<td>99.4 ± 3.1</td>
<td>219.9 ± 7.6</td>
<td>3.5 ± 0.5</td>
</tr>
<tr>
<td>SON</td>
<td>25.3 ± 0.6</td>
<td>103.3 ± 4.2</td>
<td>226.0 ± 9.7</td>
<td>4.9 ± 0.6</td>
</tr>
<tr>
<td><strong>ModelE2-YIBs</strong></td>
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<tr>
<td>MAM</td>
<td>26.4 ± 0.2</td>
<td>100.3 ± 0.8</td>
<td>222.8 ± 1.7</td>
<td>6.0 ± 0.3</td>
</tr>
<tr>
<td>JJA</td>
<td>26.4 ± 0.3</td>
<td>94.0 ± 0.9</td>
<td>208.9 ± 2.1</td>
<td>2.2 ± 0.2</td>
</tr>
<tr>
<td>SON</td>
<td>28.6 ± 0.4</td>
<td>102.6 ± 1.0</td>
<td>228.1 ± 2.1</td>
<td>3.3 ± 0.3</td>
</tr>
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</table>
Table 2. Regionally averaged MLR β-coefficients with standard deviation for GPP and HCHOv in the southeast US, defined as [31 to 35°N; -94 to -79°E] and the Amazon defined as [-15°S to 3°N, -76° to -54°E]. The covariance of GPP with $T_s$, PAR and precipitation (P) are denoted as $GPP_β_ T_s$, $GPP_β_ PAR$, $GPP_β_ P$; the covariance of HCHOv with $T_s$, SW and precipitation (P) are denoted as $HCHOv_β_ T_s$, $HCHOv_β_ SW$, $HCHOv_β_ P$. In MLR of OMI-HCHOv (a), (b) and (c), the OMI-HCHO columns are processed using model Y-PS, Y-MEGAN and Y-MEGAN-SM, respectively.

<table>
<thead>
<tr>
<th>Southeast US</th>
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<tbody>
<tr>
<td><strong>MLR of FLUXNET-GPP: 1982-2011</strong></td>
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<td></td>
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<tr>
<td><strong>MAM</strong></td>
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<tr>
<td><strong>JJA</strong></td>
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<td><strong>SON</strong></td>
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<tr>
<td><strong>MLR of OMI-HCHOv: 2005-2013</strong></td>
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<td><strong>MAM</strong></td>
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<td><strong>JJA</strong></td>
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<tr>
<td>MAM</td>
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<tr>
<td>JJA</td>
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<tr>
<td>SON</td>
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</table>

**MLR of OMI-HCHOv: 2005-2013**

<table>
<thead>
<tr>
<th></th>
<th>HCHOv_β_Ts</th>
<th>HCHOv_β_SW</th>
<th>HCHOv_β_P</th>
</tr>
</thead>
<tbody>
<tr>
<td>MAM</td>
<td>(a) 0.16 ± 0.27</td>
<td>(a) 0.00 ± 0.27</td>
<td>(a) -0.04 ± 0.29</td>
</tr>
<tr>
<td></td>
<td>(b) 0.16 ± 0.27</td>
<td>(b) 0.00 ± 0.27</td>
<td>(b) -0.05 ± 0.29</td>
</tr>
<tr>
<td></td>
<td>(c) 0.16 ± 0.27</td>
<td>(c) 0.00 ± 0.27</td>
<td>(c) -0.05 ± 0.29</td>
</tr>
<tr>
<td>JJA</td>
<td>(a) 0.18 ± 0.33</td>
<td>(a) 0.31 ± 0.54</td>
<td>(a) 0.03 ± 0.47</td>
</tr>
<tr>
<td></td>
<td>(b) 0.18 ± 0.33</td>
<td>(b) 0.31 ± 0.54</td>
<td>(b) 0.03 ± 0.47</td>
</tr>
<tr>
<td></td>
<td>(c) 0.18 ± 0.33</td>
<td>(c) 0.31 ± 0.53</td>
<td>(c) 0.03 ± 0.47</td>
</tr>
<tr>
<td>SON</td>
<td>(a) 0.03 ± 0.46</td>
<td>(a) 0.01 ± 0.52</td>
<td>(a) -0.31 ± 0.56</td>
</tr>
<tr>
<td></td>
<td>(b) 0.04 ± 0.46</td>
<td>(b) 0.01 ± 0.52</td>
<td>(b) -0.31 ± 0.56</td>
</tr>
<tr>
<td></td>
<td>(c) 0.03 ± 0.46</td>
<td>(c) 0.01 ± 0.52</td>
<td>(c) -0.31 ± 0.56</td>
</tr>
</tbody>
</table>
Table 3. Summary of regionally averaged observational and simulated seasonal correlation coefficients in the southeast US and the Amazon. In observational GPP-HCHOv (a), (b) and (c), the OMI-HCHO columns are processed using model Y-PS, Y-MEGAN and Y-MEGAN-SM, respectively.

<table>
<thead>
<tr>
<th>Southeast US</th>
<th>GPP-HCHOv</th>
<th>GPP-ISOPe</th>
<th>ISOPe-HCHOv</th>
</tr>
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<tbody>
<tr>
<td><strong>Observation</strong></td>
<td></td>
<td></td>
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</tr>
<tr>
<td><strong>MAM</strong></td>
<td>(a) 0.24 ± 0.10</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>(b) 0.24 ± 0.10</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>(c) 0.24 ± 0.10</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td><strong>JJA</strong></td>
<td>(a) -0.03 ± 0.10</td>
<td>-</td>
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<tr>
<td></td>
<td>(b) -0.03 ± 0.11</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>(c) -0.03 ± 0.10</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td><strong>SON</strong></td>
<td>(a) 0.25 ± 0.10</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>(b) 0.26 ± 0.10</td>
<td>-</td>
<td>-</td>
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<tr>
<td></td>
<td>(c) 0.26 ± 0.10</td>
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<td><strong>Y-PS</strong></td>
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<tr>
<td><strong>MAM</strong></td>
<td>0.86 ± 0.16</td>
<td>0.98 ± 0.01</td>
<td>0.88 ± 0.14</td>
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<tr>
<td><strong>JJA</strong></td>
<td>-0.19 ± 0.30</td>
<td>0.94 ± 0.07</td>
<td>-0.03 ± 0.31</td>
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<tr>
<td><strong>SON</strong></td>
<td>0.68 ± 0.22</td>
<td>0.97 ± 0.01</td>
<td>0.71 ± 0.20</td>
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<td>0.52 ± 0.26</td>
<td>0.69 ± 0.17</td>
<td>0.94 ± 0.05</td>
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<td>0.91 ± 0.11</td>
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<tr>
<td>JJA</td>
<td>-0.37 ± 0.22</td>
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<td>0.08 ± 0.35</td>
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<td>SON</td>
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<td>0.92 ± 0.02</td>
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<tr>
<td>JJA</td>
<td>(a) 0.33 ± 0.30</td>
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<tr>
<td>SON</td>
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<td><strong>Y-MEGAN-SM</strong></td>
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<td>-0.13 ± 0.49</td>
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**Figure captions**

**Figure 1.** Left column: simulated annual mean isoprene flux (mg[C] m\(^{-2}\) day\(^{-1}\)) in model Y-PS, Y-MEGAN and Y-MEGAN-SM. Right column: satellite-based HCHO columns (×10\(^{15}\) molecules cm\(^{-2}\)) from OMI processed using air-mass-factors from model Y-PS, Y-MEGAN and Y-MEGAN-SM.

**Figure 2(a).** The covariance of FLUXNET-GPP with monthly mean surface temperature (T\(_s\)), photosynthetically active radiation (PAR) and precipitation (P) in MAM (top), JJA (middle) and SON (bottom) from the MLR analysis. MLR is calculated using monthly mean data in 1982-2011. Significant regions (p<0.05) are shown with dotted shading.

**Figure 2(b).** The covariance of OMI-HCHO\(_v\) with monthly mean surface temperature (T\(_s\)), downward shortwave radiation (SW) and precipitation (P) in MAM (top), JJA (middle) and SON (bottom) from the MLR analysis. MLR is calculated using monthly mean data in 2005-2013. Significant regions (p<0.05) are shown with dotted shading.

**Figure 3.** Observed correlation between monthly mean FLUXNET-GPP and OMI-HCHO\(_v\) in four seasons: MAM, JJA, SON and DJF. Significant regions (p<0.05) are shown with dotted shading.

**Figure 4.** Simulated correlation between monthly mean (a) GPP and HCHO\(_v\), (b) GPP and ISOPe, (c) ISOPe and HCHO\(_v\) in MAM, JJA and SON using three isoprene algorithms: Y-PS, Y-MEGAN, Y-MEGAN-SM. Significant regions (p<0.05) are shown with dotted shading.
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(a) Model GPP-HCHOv correlation

Figure 4. Simulated correlation between monthly mean (a) GPP and HCHOv, (b) GPP and ISOPe, (c) ISOPe and HCHOv in MAM, JJA and SON using three isoprene algorithms: Y-PS, Y-MEGAN, Y-MEGAN-SM. Significant regions (p<0.05) are shown with dotted shading.
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