Assessing sources of uncertainty in formaldehyde air mass factors over tropical South America: Implications for top-down isoprene emission estimates

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[1] We use a nested-grid version of the GEOS-Chem chemistry transport model, constrained by isoprene emissions from the Model of Emissions of Gases and Aerosols from Nature (MEGAN), and the Lund-Potsdam-Jena General Ecosystem Simulator (LPJ-GUESS) bottom-up inventories, to evaluate the impact that surface isoprene emissions have on formaldehyde (HCHO) air-mass factors (AMFs) and vertical column densities (VCDs) over tropical South America during 2006, as observed by the Scanning Imaging Absorption Spectrometer for Atmospheric Chartography (SCIAMACHY) and Ozone Monitoring Instrument (OMI). Although the large-scale seasonal variability of monthly mean HCHO VCDs is typically unaffected by the choice of bottom-up inventory, large relative differences of up to $\pm 45\%$ in the HCHO VCD can occur for individual regions and months, but typically most VCD differences are of order $\pm 20\%$. These relative changes are comparable to those produced by other sources of uncertainty in the AMF including aerosols and surface albedo, but less than those from clouds. In a sensitivity test, we find that top-down annual isoprene emissions inferred from SCIAMACHY and OMI HCHO vertical columns can vary by as much as $\pm 30-50\%$ for each instrument respectively, depending on the region studied and the a priori isoprene emissions used. Our analysis suggests that the influence of the a priori isoprene emissions on HCHO AMFs and VCDs is therefore non-negligible and must be carefully considered when inferring top-down isoprene emissions estimates over this, or potentially any other, region.

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1. Introduction

[2] Formaldehyde (HCHO) is a key atmospheric constituent produced from the oxidation of volatile organic compounds (VOCs). While the oxidation of methane sustains low (subppbv) background levels, over continental regions, the

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oxidation of short-lived VOCs from anthropogenic, biogenic and pyrogenic sources, along with direct emissions from fires and industrial processes, can produce large (>5 ppbv) amounts of HCHO within the boundary layer. Since HCHO has a short atmospheric lifetime (approximately several hours) owing to photolysis and reaction with the hydroxyl radical (OH), it is a useful tracer for inferring surface emissions [Palmer et al., 2003]. Model studies indicate that isoprene, the dominant biogenic VOC emitted from plants [Guenther et al., 2006], is rapidly oxidized to form HCHO with a high yield and largely controls HCHO column variability over densely vegetated areas [Stavrakou et al., 2009a]. Yet although the importance of isoprene within climate is well established, its emissions are poorly quantified [Arneth et al., 2008]. Consequently, satellite measurements of HCHO columns have been used in several studies to provide top-down constraints on isoprene emissions [see, e.g., Stavrakou et al., 2009b]. Satellite-based estimates are particularly valuable over tropical ecosystems, where isoprene emissions are greatest and bottom-up inventories least well constrained [Barkley et al., 2008, 2009]. However, the errors associated with satellite derived emissions are large owing to significant uncertainties in the HCHO column

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retrieval itself, and also the selected chemistry-transport model (CTM) used to infer the estimates (e.g., owing to simplified parameterizations of isoprene oxidation chemistry).

[3] The HCHO vertical column retrieval consists of twosteps. First, HCHO slant column densities (SCDs) along the instruments line-of-sight are obtained through the spectral fitting of trace gas absorption cross-sections to measured solar UV backscatter spectra. Second, the slant columns are converted to vertical column densities (VCDs) after division by an air-mass factor (AMF = SCD/VCD) which accounts for the satellite viewing geometry, HCHO vertical distribution, surface reflectance, and atmospheric scattering by air-molecules, aerosols, and clouds [Palmer et al., 2001]. Since the AMF computation can significantly affect the accuracy of the final HCHO product, it is this aspect of the retrieval we focus on. In particular, we draw attention to the influence on the AMF of the HCHO vertical distribution, which is typically provided by a CTM driven by a specified bottom-up isoprene emission inventory - in most cases an application of the canopy-scale Model of Emissions of Gases and Aerosols from Nature (MEGAN), developed by Guenther et al. [2006]. Although the estimated AMF error due to uncertainties in the HCHO vertical profile is estimated to be only about 10-20% [Millet et al., 2006; Palmer et al., 2006; De Smedt et al., 2008], to our knowledge no-one has yet assessed this error component using model HCHO profiles based on more than one isoprene emission inventory. In this work, we attempt to address this oversight by assessing HCHO AMFs and VCDs over tropical South America, utilizing a recently developed GEOS-Chem CTM nested grid simulation of Amazonian biogenic emissions and tropospheric chemistry [Barkley et al., 2011]. The nested grid is unique in that it can be forced with isoprene emissions from MEGAN, or alternatively, from a leaf-scale algorithm which is coupled to the LPJ-GUESS dynamic vegetation model [Arneth et al., 2007]. Here we run GEOS-Chem with both inventories using the subsequent HCHO profiles to compute AMFs to apply to slant columns retrieved by the SCIAMACHY [Bovensmann et al., 1999] and OMI instruments [Levelt et al., 2006], in order to examine the relative changes in the resulting HCHO VCD distributions. Our goal is to determine the importance of the a priori isoprene emissions on the final HCHO VCDs, as compared with influence of other AMF inputs (e.g., aerosols). We stress that this work is not a full-error analysis of absolute column magnitudes, nor an inter-comparison of the two satellite products.

[4] In section 2 we provide a brief overview of the GEOS-Chem model, and describe the SCIAMACHY and OMI HCHO slant column retrievals and the computation of their AMFs. In section 3 we present the results of our sensitivity analysis, and then in section 4 we examine the impact of the different a priori isoprene inventories on subsequent top-down isoprene emission estimates. We conclude the paper in section 5.

2. Data and Methods

2.1. GEOS-Chem Chemistry Transport Model

[5] Since *Barkley et al.* [2011] describe the GEOS-Chem Amazon simulation in great detail, for brevity, here we only give a short description. The nest-grid has a horizontal resolution of $0.667^{\circ} \times 0.5^{\circ}$ (longitude × latitude), and 47 vertical levels extending from the surface to 0.01 hPa. The model is

driven using GEOS-5 meteorology [Rienecker et al., 2008], which is updated every 3–6 hours. Tracer mixing ratios from an off-line global $4^{\circ} \times 5^{\circ}$ simulation provide 3-hourly boundary conditions to the grid-edges. Based on a previous model evaluation [Barkley et al., 2011], we use an updated chemical mechanism [Paulot et al., 2009a, 2009b] to simulate O₃-NO_x-VOC-aerosol photochemistry. To quantify the effect of the input isoprene emissions on the HCHO AMFs and VCDs, we run GEOS-Chem with four emission scenarios for the year 2006 using two different variants of the MEGAN and LPJ-GUESS inventories taken from the study of *Barkley et al.* [2011], as summarized in Table 1 and shown in Figure S1 of the auxiliary material.¹ Substantial differences exist between the isoprene emissions predicted by these models owing to their different assignment of basal emission capacities, choice of driving algorithms and forcing meteorology, and their different vegetation distributions and leaf-area [Guenther et al., 2006; Arneth et al., 2007]. For example, MEGAN uses MODIS leaf-area applied to static vegetation maps where as LPJ-GUESS explicitly simulates vegetation foliage [Barkley et al., 2011]. MEGAN isoprene emissions tend to be highest along the Brazilian border with Peru and Bolivia owing to the presence of bamboo forest, where as LPJ-GUESS emissions are typically higher in the southeast owing to emissions from broad leaf rain green trees [Barkley et al., 2011]. Owing to a lack of observational with which to validate the simulated emissions during our study period and given the large uncertainties of tropical isoprene emissions, we have to assume each model is viable despite notable differences in their annual totals (Table 1). The choice of inventory can have a large impact on the simulated HCHO profiles; in extreme cases differences of up to 4 ppbv can exist within the lowest 3 km of the atmosphere (see, e.g., Figure S2).

2.2. SCIAMACHY and OMI HCHO Slant Columns

[6] SCIAMACHY is a passive UV-Vis-NIR grating spectrometer situated on board ESA's ENVISAT satellite [Bovensmann et al., 1999]. For the majority of its polar sunsynchronous orbit, SCIAMACHY makes measurements in an alternating limb and nadir sequence, crossing the equator at 10:00 local time (LT). In nadir mode, the ground swath has fixed dimensions of 960 \times 30 km² (across \times along track) with a nominal pixel size of $60 \times 30 \text{ km}^2$. Global coverage is achieved at the equator within about 6 days. HCHO slant columns are retrieved from nadir UV-spectra using differential optical absorption spectroscopy, as described in De Smedt et al. [2008]. A spectral fitting window of 328.5-346 nm is used to reduce fitting uncertainties due to a polarization anomaly at 350 nm, and a strong O₄ absorption band centered near 360 nm. Besides the fitting of the absorption cross sections of HCHO and other interfering gases, corrections for the Ring effect and a linear intensity offset are applied, along with a fifth order polynomial closure term. A reference sector adjustment is also made based on daily observations over the central Pacific Ocean (140-160°W) [De Smedt et al., 2008; De Smedt, 2011]. Mean (systematic) fitting uncertainties of a HCHO slant column measurement usually range between 30-150%.

¹Auxiliary materials are available in the HTML. doi:10.1029/2011JD016827.

Table 1. Description of the Air Mass Factor Sensitivity Simulations for the Year 2006

Scenario	Isoprene Emissions (Tg C)	Description			
SCIA* or OMI*	154	Default scenario for each instrument. Isoprene emissions calculated using a 5-layer canopy model and a combination of <i>Guenther et al.</i> [1999, 2006] algorithms ^a			
MEGAN'	98	As above, but with isoprene emissions scaled by 0.635 to match the monthly mean emissions from the study of <i>Müller et al.</i> [2008] ^b			
LPJ(GC)	75	Emissions based on the LPJ-GUESS model forced with GEOS-Chem's GEOS-5 meteorology			
LPJ(CRU)	90	Emissions based on the LPJ-GUESS model forced with its default CRU meteorology ^c			
BL	154	As the default scenario but with a non-local boundary layer mixing scheme employed in GEOS-Chem ^d			
ALB	154	As the default scenario but using the <i>Kleipool et al.</i> [2008] surface reflectances in the AMF computation			
AOD	154	As the default scenario but without an aerosol correction applied (i.e. $AOD = 0$) in the AMF computation			
CF	154	As the default scenario, but assuming a +0.1 cloud fraction error in the AMF computation ^e			
СТР	154	As the default scenario, but assuming a -60 hPa error in cloud top pressure in the AMF computation ^e			

^aThis (hybrid) emission scheme, which uses MEGAN v2.1 basal emission factors, is fully described in *Barkley et al.* [2011].

^bThe 0.635 scaling factor is based on both 2005 and 2006 emissions [see Barkley et al., 2011].

^cData from the Climate Research Unit of the University of East Anglia (http://www.cru.uea.ac.uk/).

^dThe non-local scheme includes 'local' mixing between adjacent model layers and, depending on the stability of the PBL, 'non-local' mixing due to turbulent eddies [*Lin and McElroy*, 2010].

^cEstimated uncertainties based on the study of *Acarreta et al.* [2004]; here we use a +0.1 error in cloud fraction to determine the likely maximum range of this effect on the AMFs and VCDs.

[7] The Dutch-Finnish Ozone Monitoring Instrument (OMI) [Levelt et al., 2006], is a nadir-viewing near-UV/ visible charged-couple device (CCD) spectrometer situated on board NASA's Aura satellite. OMI also orbits the Earth in a sun-synchronous polar orbit, but with an equator crossing time of 13:30 LT. OMI has a 114° field-of-view producing a 2600 km wide swath containing 60 cross-track pixels that range in size from 14×26 km² at nadir, to $28 \times$ 160 km² at the swath edges. HCHO slant columns are retrieved for each of the 60 cross track pixels, through a direct non-linear least squares fitting of spectral radiances within the interval 327.5-356.5 nm, as described in Chance [2002] and Kurosu et al. [2004]. The cross-sections of HCHO and other absorbers are fitted, along with a Ring effect correction, closure polynomials and optional spectral shift and squeeze parameters. The retrieval algorithm also includes dynamic calibration of solar and radiance wavelengths, an under-sampling correction, computation of common model residual spectrum, and a de-striping algorithm to minimize cross-track striping. Fitting uncertainties of a single HCHO slant column measurement typically range between 40-100% [Chance, 2002].

2.3. Calculation of SCIAMACHY and OMI AMFs

[8] To establish consistent AMFs for each instrument we construct look-up tables using monthly averaged HCHO profiles and aerosol optical depths from GEOS-Chem, appropriate to each instrument's overpass time (see Figures S3 and S4). These serve as input to the radiative transfer model LIDORT [*Spurr et al.*, 2001], which calculates scattering weights that represent the sensitivity of the backscattered radiance to the HCHO abundance at each altitude, but which also effectively decouples this

dependency from the profile shape [*Palmer et al.*, 2001]. We parameterize each look-up table as a function of location (i.e. surface pressure), solar zenith angle, cloud-top pressure, and scan angle. We account for partially cloudy scenes using the approach of *Martin et al.* [2002], which assumes the total AMF is the brightness-weighted average of the air mass factors for the clear (AMF_{clr}) and cloudy (AMF_{cld}) pixel sub-scenes:

$$AMF = \frac{AMF_{clr} \cdot R_{clr} \cdot (1-f) + AMF_{cld} \cdot R_{cld} \cdot f}{R_{clr} \cdot (1-f) + R_{cld} \cdot f}$$
(1)

where f is the cloud fractional coverage, and R_{cld} and R_{clr} are the sub-scene reflectivities calculated by LIDORT, given by:

$$R_{clr} = \frac{\pi I_{clr}}{I_0 \cos \theta_{SZA}},\tag{2}$$

$$R_{cld} = \frac{\pi I_{cld}}{I_0 \cos \theta_{SZA}},\tag{3}$$

with I_0 the solar irradiance at the top of the atmosphere perpendicular to the solar beam, θ_{SZA} the solar zenith angle, and I_{clr} and I_{clr} the clear and cloudy backscattered radiances. R_{clr} depends on wavelength, surface pressure, surface albedo and solar/viewing geometry; R_{cld} also depends on cloudtop pressure and cloud albedo [*Martin et al.*, 2002]. The AMFs are computed at 340 nm for SCIAMACHY and 328 nm for OMI; the different sensitivities of the two instruments for example scenes are shown in the auxiliary material Figure S5). A shorter wavelength is used for OMI to better represent the air mass of its wider fitting window; our tests indicate AMF differences due to the two



Figure 1. Monthly mean air mass factors (AMFs) for 2006, corresponding to 09:00–11:00 local time, for SCIAMACHY (first column) using a HCHO climatology based on the MEGAN bottom-up emission inventory. The remaining columns show the relative changes in the SCIAMACHY AMFs arising from the sensitivity simulations outlined in Table 1. The SCIAMACHY AMF data have been averaged onto the nested $0.667^{\circ} \times 0.5^{\circ}$ grid using observations with cloud cover \leq 40% and smoothed with a 9 \times 9 box-filter to remove noise. White-colored regions correspond to areas with no usable SCIAMACHY data. The overlain boxes (in black) correspond to the regional areas (north, west, east, southwest, and southeast) used in subsequent analyses (see section 3).

wavelengths alone are about 5%. Clouds are characterized as Lambertian reflectors with an albedo of 0.8 [*Chance*, 2002], with the cloud fraction and cloud-top pressure for each observation provided by the respective SCIAMACHY

FRESCO v5 [Koelemeijer et al., 2002], and OMI O_2-O_2 [Acarreta et al., 2004] cloud algorithms. Scenes with >40% cloud cover are removed from our analysis. For clear-sky conditions, the surface albedo (at ~360 nm) is taken from



Figure 2. Monthly mean air mass factors (AMFs) for 2006, corresponding to 12:00–15:00 local time, for OMI (first column) using a HCHO climatology based on the MEGAN bottom-up emission inventory. The remaining columns show the relative changes in the OMI AMFs arising from the sensitivity simulations outlined in Table 1. The OMI AMF data have been averaged onto the nested $0.667^{\circ} \times 0.5^{\circ}$ grid using observations with cloud cover $\leq 40\%$.

a climatological database derived from TOMS surface reflectance measurements [*Herman and Celarier*, 1997].

3. Results

3.1. Sensitivity to Surface Isoprene Emissions

[9] We define two scenarios: SCIA* and OMI* which use isoprene emissions based on MEGAN (see Table 1), as a baseline with which to compare the impact of the other bottom-up emissions on the AMFs and VCDs. AMFs based on this scenario range from 0.5–1.9 over land, and 0.7–2.2 over the oceans (see Figures 1 and 2). AMFs are lower over land owing to HCHO profile shapes that peak nearer the surface where scattering weights are smallest, in contrast to oceanic HCHO profiles that are more uniform with altitude (Figure S5) [see also *Palmer et al.*, 2001]. The corresponding monthly mean HCHO VCD distributions, shown in the first column of Figures 3 and 4, respectively, have a clear



Figure 3. Monthly mean HCHO vertical columns for 2006 corresponding to 09:00–11:00 local time, as observed by SCIAMACHY (first column) using AMFs based on the MEGAN bottom-up emission inventory. The remaining columns show the relative changes in the SCIAMACHY HCHO VCDs arising from the sensitivity simulations outlined in Table 1. The SCIAMACHY data have been averaged onto the nested $0.667^{\circ} \times 0.5^{\circ}$ grid using observations with cloud cover $\leq 40\%$ and smoothed with a 9 \times 9 box-filter to remove noise. White-colored regions correspond to areas with no usable SCIAMACHY data.

seasonal trend peaking in the dry-season in response to increased isoprene emissions and HCHO released from biomass burning [*Barkley et al.*, 2008]. The subsequent columns of Figures 3 and 4 which show the relative differences in the VCDs (or Δ VCD) arising from the sensitivity simulations outlined in Table 1, more importantly illustrate that the influence of the a priori isoprene emissions on the AMF computation, through their control on the HCHO

profiles in the input climatology, can result in non-negligible spatial and temporal changes in the HCHO VCD distributions (see also Table S1).

[10] For example, when the isoprene emissions of the SCIA* and OMI* simulations are scaled downward to match the MEGAN estimates of *Müller et al.* [2008] (scenario: MEGAN'), the HCHO columns of the climatology are mostly lowered thereby increasing the AMFs, particularly over



Figure 4. Monthly mean HCHO vertical columns for 2006 corresponding to 12:00–15:00 local time, as observed by OMI (first column) using AMFs based on the MEGAN bottom-up emission inventory. The remaining columns show the relative changes in the OMI HCHO VCDs arising from the sensitivity simulations outlined in Table 1. The OMI data have been averaged onto the nested $0.667^{\circ} \times 0.5^{\circ}$ grid using observations with cloud cover $\leq 40\%$.

eastern areas. This results in correspondingly lower HCHO VCDs across these regions by on average only 2–3%. However, the differences for individual grid cells can be larger, with Δ VCD ranging from approximately $\pm 10\%$ for both instruments.

[11] More dramatic changes in the HCHO VCDs are produced from the use of the alternative LPJ-GUESS emissions, which have very different spatial distributions to MEGAN (Figure S1). For instance, when GEOS-Chem's GEOS-5 meteorological fields are used to drive the LPJ-GUESS model (scenario: LPJ(GC)), the reduced emissions yield a significantly lower HCHO climatology in most areas, except in the east and southeast during March–June. As a consequence, in those regions and months the AMFs are smaller by on average 2–10%, and the VCDs larger by about 2–8%, than in the SCIA* and OMI* baseline simulations. Elsewhere, we typically find higher AMFs especially in western and southwestern areas during the dry season,



Figure 5. (left) Time series of the mean SCIAMACHY HCHO vertical columns (molecules cm⁻²) for different Amazon regions (defined in Figure 1). The black dots correspond to the SCIA* scenario whose AMFs were computed using a HCHO climatology based on the MEGAN bottom-up emission inventory (see section 2.1). The colored lines show the mean SCIAMACHY HCHO vertical columns determined using AMFs from the various sensitivity simulations as described in Table 1. (middle) Time series of the corresponding relative changes in the SCIAMACHY HCHO vertical columns (Δ VCD) arising from the AMF sensitivity simulations. (right) Time series of the percentage of grid cells that have a $|\Delta VCD|$ of between 10–20% ($F_{110-20\%}$).

except along the northwest coast where the AMFs are lower in the early part of the year (Figures 1 and 2). Overall, over land Δ VCD ranges from about $\pm 25\%$ for SCIAMACHY, and $\pm 45\%$ for OMI, though values exceeding $\pm 20\%$ are infrequent on an annual basis (see Table S1). That said, for specific regions systematic differences in the HCHO VCDs can occur. For example, over the southwest the number of grid cells that have a $|\Delta VCD|$ of between 10–20%



Figure 6. (left) Time series of the mean OMI HCHO vertical columns (molecules cm⁻²) for different Amazon regions (defined in Figure 1). The black dots correspond to the OMI* scenario whose AMFs were computed using a HCHO climatology based on the MEGAN bottom-up emission inventory (see section 2.1). The colored lines show the mean OMI HCHO vertical columns determined using AMFs from the various sensitivity simulations as described in Table 1. (middle) Time series of the corresponding relative changes in the OMI HCHO vertical columns (Δ VCD) arising from the AMF sensitivity simulations. (right) Time series of the percentage of grid cells that have a $|\Delta VCD|$ of between 10–20% ($F_{110-20\%}$).

(hereafter denoted $F_{110-20\%1}$) are annually 28% and 43% for SCIAMACHY and OMI, respectively. The seasonal variation in $F_{110-20\%1}$ for SCIAMACHY and OMI over different Amazon regions can also be large, as shown in Figures 5

and 6. For instance, over the southwest during August– October, OMI $F_{110-20\%}$ is as high as 50–60%, with 22% of grid cells having a $|\Delta VCD|$ of greater than 20% ($F_{>20\%}$) in October alone. [12] Similarly, when the LPJ-GUESS model is forced with its default CRU meteorology (scenario: LPJ(CRU)), the HCHO climatology also has higher HCHO columns in southeastern regions, but this time throughout most of the year and particularly during May–July. For both instruments this results in lower AMFs and thus higher VCDs over these areas (monthly $\overline{\Delta}$ VCD range from 2–15%). The range of Δ VCD is comparable to the LPJ(GC) scenario, typically ±25% for SCIAMACHY and ±40% for OMI, though the occurrence of such high values is rare. However, regional effects in Δ VCD are again evident, but this time most notably over the southeast during April–July when $F_{110-20\%l}$ is often greater than 40% for OMI (Figure 6), or during May when $F_{110-20\%l}$ is nearly 60% for SCIAMACHY (Figure 5).

3.2. Sensitivity to Other AMF Parameters

[13] Despite the clear impact of the isoprene emissions on the AMFs, examination of the HCHO VCD time series, shown in Figures 5 and 6, reveals that while the VCD magnitudes may change, observed seasonal trends are largely unaffected by the choice of bottom-up inventory, i.e. most of the temporal variability comes from the spectral fitting of the SCDs. The question is therefore: are such relative changes significant when compared with the impact of other AMF inputs, or for that matter, other model processes that affect the HCHO profile such as boundary layer (BL) mixing? To determine the response of the VCDs to these effects, we separately recomputed AMFs using the SCIA* and OMI* scenarios but with (a) no aerosol correction; (b) an alternative surface reflectance database derived from OMI [Kleipool et al., 2008]; (c) systematic errors of +0.1 in cloud fraction and -60 hPa in cloud-top pressure (typical uncertainties estimated by Acarreta et al. [2004]); and (d) with a GEOS-Chem HCHO climatology simulated with a more sophisticated non-local BL mixing scheme [Lin and McElroy, 2010], as opposed to a uniformly mixed BL in the default simulations.

[14] Owing to the relatively clean atmospheric conditions found over most undisturbed Amazonian regions, we find that the impact of switching off the aerosol correction (scenario: AOD) only has moderate effect on the SCIAMACHY and OMI AMFs and VCDs (assuming GEOS-Chem AODs are representative of the true aerosol conditions). We find differences in the HCHO VCDs of order $\pm 15\%$ at most, which are therefore comparable with the effects of the a priori isoprene emissions. The most noteworthy changes in the AMFs actually occur over the Atlantic due to the presence of transported Saharan dust, and over central Amazon during July–September owing to the presence of black and organic carbonaceous aerosol from biomass burning. As a further test, we recalculated the AMFs but with the black and organic carbon AODs increased everywhere by an arbitrary 50% in order to simulate a high aerosol loading from fires, but this had only a minor effect on the SCIAMACHY and OMI HCHO columns with ΔVCD mostly $\pm 3\%$. However, over biomass burning scenes the AMF can become highly sensitive to the relative vertical distribution of aerosols and HCHO [Fu et al., 2007]; aerosols underlying or coincident with the HCHO maximum typically increase measurement sensitivity [Gonzi et al., 2011]. To examine aerosol effects further, five additional simulations were conducted for the

default, LPJ(CRU) and LPJ(GC) scenarios, in which the black and organic AOD profiles were adjusted depending if their emissions both exceeded 10^9 molecules cm⁻² s⁻¹ and their individual maximum AOD occurred within the BL (see Figure S6 and auxiliary material for full details). Our results indicate that the HCHO VCD is not greatly affected if the aerosols are redistributed throughout and/or immediately above the BL ($|\Delta VCD| < 5\%$). Only when the AOD is distributed uniformly to higher altitudes (\sim 5 km) do substantial changes in the HCHO column occur. Aerosols at high altitudes shield the underlying HCHO, reducing the AMFs and increasing the HCHO VCDs by as much 10-50% over regions of active burning (Figures S7 and S8). The magnitude of ΔVCD is similar for both instruments but is more widespread for OMI since the modeled maximum AOD is more likely to reside in the deeper BL at its overpass. The impact of each aerosol simulation is generally consistent across the different isoprene emissions scenarios i.e. the HCHO profile over fire affected regions is more influenced by HCHO from burning vegetation than from oxidized biogenic emissions. The HCHO VCD is therefore potentially very sensitive to aerosols over areas of biomass burning, however, scenes affected by fires are often discarded when inferring top-down isoprene emissions [e.g., Barkley et al., 2008].

[15] Use of the new surface reflectances derived by Kleipool et al. [2008] also has a substantial impact on the AMFs and VCDs (scenario: ALB). The UV albedo over the rain forest is usually quite low (<0.1) at the relevant wavelengths computed for the SCIAMACHY (340 nm) and OMI (328 nm) fitting windows and comparison of the two albedo data sets reveals significant differences (see Figure S9). The resulting changes in the HCHO AMFs and VCDs arise from the different resolutions and observing times of the albedo data (TOMS: $1.25^{\circ} \times 1.0^{\circ}$ and 12:00 LT, OMI: $0.5^{\circ} \times 0.5^{\circ}$ and 13:30 LT), the use of surface reflectances at wavelengths more appropriate to each instrument's spectral fitting window (328 nm for OMI and 342 nm for SCIAMACHY), and the fact that the Kleipool et al. [2008] data has been derived using the mode or 1% cumulative probability threshold of OMI measurements of Lambertian Equivalent Reflectivity (LER), instead of the minimum LER as in the TOMS data (which tends to underestimate the overall reflectivity of an observed scene). Typically, a high surface albedo increase measurement sensitivity to a lower tropospheric absorber such as HCHO, since more solar radiation is reflected back to space [Palmer et al., 2001; De Smedt et al., 2008]. We find the AMFs are decreased where the newer albedo is lower than the Herman and Celarier [1997] values and higher where increased, thereby producing the converse effect in the HCHO vertical columns. Over land, ΔVCD ranges from about $\pm 30\%$ for SCIAMACHY, and $\pm 40\%$ for OMI, but annually values of $F_{>20\%}$ are low (<1%). In spite of this, there are clear seasonal variations in Δ VCD for some regions, e.g., in eastern and western areas where the mean difference is negative during May-July and positive outside this period (see Figures 5 and 6). Moreover, in eastern regions during October–March, $F_{110-20\%}$ often exceeds 20% for both instruments, with $F_{>20\%}$ nearly 10% for OMI during February-March.

[16] Previous studies have identified clouds as the main source of error on the AMF [e.g., *Millet et al.*, 2006]. We

Table 2. Estimated Bottom-Up and SCIAMACHY Top-Down Isoprene Emissions for 2006^a

Scenario ^b	Bottom-Up a Priori (Tg C)	Top-Down ^c a Posterior (Tg C)	Relative Difference (%)	Top-Down ^d a Posterior (Tg C)	Relative Difference (%)			
Land								
SCIA*	365	151	_	151	-			
LPJ(GC)	171	215	42	146	-3			
LPJ(CRU)	191	158	5	148	-2			
North Amazon								
SCIA*	40	16	_	16	-			
LPJ(GC)	23	22	38	18	13			
LPJ(CRU)	22	17	6	18	13			
Wast Amazon								
SCIA*	83	21	-	21	-			
LPJ(GC)	26	31	48	21	0			
LPJ(CRU)	28	24	14	21	0			
		East A	mazon					
SCIA*	71	40	-	40	-			
LPI(GC)	38	58	45	39	-3			
LPJ(CRU)	40	43	8	39	-3			
		Southwes	t Amazon					
SCIA*	72	19	-	19	-			
LPJ(GC)	12	24	26	14	-26			
LPJ(CRU)	26	22	16	18	-5			
		Southeas	t Amazon					
SCIA*	20	11	_	11	-			
LPJ(GC)	18	15	36	11	0			
LPJ(CRU)	20	15	36	12	9			

^aBoth the bottom-up and top-down isoprene emissions correspond to 09:00–11:00 local time coinciding with the overpass of SCIAMACHY (10:00). Relative differences calculated as 100% \times (scenario - SCIA*)/SCIA*.

^bRegions are shown in Figure 1 and scenarios described in Table 1.

^cEstimated using regression parameters from each individual scenario, as described in section 4.

^dEstimated using regression parameters from the default SCIA* scenario, as described in section 4.

also find the largest changes in the AMFs and VCDs by assigning a +0.1 error in cloud fraction (achieved by increasing f for each observation by 0.1 in equation (1) after being cloud filtered using its original value; scenario: CF). For SCIAMACHY, the subsequent AMFs are in general reduced (by about 4%) as less weight is placed on the clearsky sub-scene AMF, thereby increasing the VCDs over land by on average 8% (range: 55 to -41%). The largest decrease in the AMFs occur during January-April (i.e. the wet season) resulting in a widespread changes in ΔVCD , with $F_{10-20\%}$ often greater than 50% during these months. For OMI, the AMFs are also reduced in most continental areas increasing the HCHO VCDs by 4% (range: 70 to -73%), with $F_{110-20\%}$ varying between 15–30% over land (Figure 6). The only exception being in southern regions during April-August when the AMFs are increased by on average 5–21%, and the VCDs decreased by 4-13%. Increasing the cloud fraction increases the AMF in some regions because of the cloud albedo effect, whereby low-level clouds that reside underneath or with the peak of the HCHO profile increase measurement sensitivity [De Smedt, 2011]. A further simulation where we decreased f by 0.1 produced similar changes but of the opposite sign. Use of stricter cloud-filter (20%)

with a +0.1 increase in *f*, typically produces larger values of Δ VCD (not shown), with $F_{>20\%}$ varying between 10–30% for both instruments. By comparison, a 60 hPa systematic error in the cloud-top pressure (scenario: CTP) has a lesser but still significant impact on the AMFs and VCDs than a 0.1 error in cloud fraction. A higher altitude (i.e. lower) cloud-top pressure yields a lower AMF [*De Smedt et al.*, 2008], thus on average for both instruments we determine consistent decreases and increases in the AMFs and VCDs over land of about -6% and 6%, respectively, though for some months and locations the differences can be up to 20%.

[17] The choice of BL mixing scheme within GEOS-Chem can strongly influence near surface mixing ratios of isoprene and its oxidation products, including the vertical distribution of HCHO [Barkley et al., 2011]. The non-local mixing scheme can sometimes produces very different HCHO profile shapes compared with default full-mixing scheme (Figure S2), with the effect often more pronounced at the time of OMI's overpass. However, we find the effect of the non-local mixing scheme (scenario: BL) on the input HCHO climatology, results in only a moderate impact on the SCIAMACHY with \triangle AMFs of about \pm 3% and \triangle VCDs of $\pm 10\%$. For OMI, the differences are slightly larger with Δ AMFs ranging from 4 to -10% and Δ VCDs from 18 to -23%, though mostly $\Delta VCDs$ varies between $\pm 10\%$. The effects of BL mixing are therefore comparable with other AMF uncertainties, with the greater impact on OMI most likely reflecting a deeper and more often mixed BL at its overpass time.

4. Implications for Top-Down Isoprene Emissions Estimates

[18] To highlight what these VCD differences potentially imply for subsequent top-down isoprene emissions, as a sensitivity test, we estimate isoprene emissions from SCIAMACHY and OMI HCHO VCDs, computed using AMFs from the SCIA*, OMI*, and appropriate LPJ(GC) and LPJ(CRU) simulations. To infer the top-down estimates we adopt the approach of Palmer et al. [2003], whereby we linearly regress the model isoprene emissions and HCHO vertical columns to determine the gradient and intercept, which we transpose and apply to the corresponding OMI HCHO VCDs from each scenario [see also Barkley et al., 2008]. As rigorous estimates are not required in this instance, we neglect the effects of spatial smearing associated with delayed HCHO production, as well as other model uncertainties, but remove scenes influenced by fires using ATSR fire count data [Barkley et al., 2008]. Note, in a follow study we will present an ensemble of robust top-down estimates in which we pay strict attention to these error sources.

[19] Since differences in the top-down estimates occur due to changes in the retrieved HCHO VCDs, and the linear regression relationship obtained from each simulation we use two methods to infer top-down isoprene emissions. First, we run GEOS-Chem with each scenario's specified emissions, calculate the linear transfer function from the subsequent model HCHO columns and isoprene emissions, and then apply it to the observed HCHO VCDs, computed using AMFs corresponding to that simulation. Second, we use the gradient and intercept from our default SCIA* and OMI* simulations, and apply them to the observed HCHO VCDs determined using AMFs from the LPJ(GC) and LPJ(CRU)

Table 3. Estimated Bottom-Up and OMI Top-Down Isoprene Emissions for 2006^{a}

Scenario ^b	Bottom-Up a Priori (Tg C)	Top-Down ^c a Posterior (Tg C)	Relative Difference (%)	Top-Down ^d a Posterior (Tg C)	Relative Difference (%)
		La	nd		
OMI*	435	197	-	197	-
LPJ(GC)	200	234	19	192	-3
LPJ(CRU)	240	184	-7	197	0
		North 2	4mazon		
OMI*	43	20	-	21	-
LPJ(GC)	24	24	20	20	5
LPJ(CRU)	23	20	0	20	5
		West A	lmazon		
OMI*	95	28	-	28	-
LPJ(GC)	29	35	25	27	-4
LPJ(CRU)	32	29	4	28	0
		East A	mazon		
OMI*	77	51	-	51	-
LPJ(GC)	41	60	18	49	-4
LPJ(CRU)	43	46	-10	49	-4
		Southwes	t Amazon		
OMI*	98	29	-	29	-
LPJ(GC)	16	29	17	21	-28
LPJ(CRU)	37	29	0	27	-7
		Southeas	t Amazon		
OMI*	31	19	-	19	-
LPJ(GC)	32	27	42	20	5
LPJ(CRU)	53	23	21	23	21

^aBoth the bottom-up and top-down isoprene emissions correspond to 12:00–15:00 local time coinciding with the overpass of OMI (13:30). Relative differences calculated as $100\% \times (\text{scenario} - \text{OMI*})/\text{OMI*}$.

^bRegions are shown in Figure 1 and scenarios described in Table 1.

^cEstimated using regression parameters from each individual scenario, as described in section 4.

^dEstimated using regression parameters from the default OMI* scenario, as described in section 4.

scenarios. This latter approach ensures any differences in the inferred top-down emissions are due to changes in the observed HCHO VCDs alone. We show in Tables 2 and 3 the results of both methods, when applied to the observed SCIAMACHY and OMI HCHO VCDs.

[20] Focussing on OMI, we find using individual regression parameters results in large differences between the topdown estimates, as the annual isoprene emissions over land inferred from the OMI*, LPJ(GC) and LPJ(CRU) scenarios are 197, 234, and 184 Tg C respectively, at the time of the instruments's overpass (Table 3). However, for individual regions the differences in the top-down emissions can be much larger, e.g., in the southeast where the LPJ(GC) and LPJ(CRU) estimates are 21% and 42% higher than those determined from the OMI* scenario. When using the gradient and intercept from the default OMI* simulation alone, which eliminates compensating effects between the $\Delta VCDs$ and the regression parameters, we find for most regions the differences between the top-down estimates are smaller, as its only the only changes in the HCHO VCDs that matter. The only exception being the southwest region where the LPJ based estimates are 28% and 7% lower than the OMI* derived emission. These traits are also consistent in the topdown estimates determined from SCIAMACHY (Table 2).

For example, over land the annual isoprene emissions inferred in the SCIA*, LPJ(GC) and LPJ(CRU) scenarios are 151, 215, and 158 Tg C respectively when using the individual regression parameters, and 151, 146, and 148 Tg C using the SCIA* gradient and intercept (which typically brings the top-down estimates in closer agreement for most regions).

[21] Therefore, as this simple exercise illustrates, the use of two bottom-up inventories to compute SCIAMACHY and OMI AMFs and VCDs, can potentially lead to differences of up to $\pm 30-50\%$ in the top-down emission estimates, depending on the region studied and inference technique used. In spite of this, the top-down estimates generally agree more closely than the a priori inventories demonstrating the likely value of HCHO columns for constraining surface isoprene emissions.

5. Conclusions

[22] The AMF calculation is a critical step in the HCHO retrieval. Computing HCHO AMFs using a specified CTM and bottom-up isoprene emission inventory, ensures some degree of self-consistency for subsequent model-satellite comparisons and inference of top-down isoprene emission. However, we have shown that at least for tropical South America, the use of different isoprene emission inventories within the same CTM can cause non-negligible changes in the resulting HCHO VCD monthly distributions, and in some cases very large differences for individual locations and months. The size of these relative changes are at least comparable to those originating from other sources of uncertainty in the AMF calculation, such as aerosols or the surface albedo, but are potentially less than the relative changes associated with errors in cloud fractional coverage. While we find that large-scale seasonal trends are generally unaffected by the use of the two differing isoprene inventories, we do observe notable changes in the HCHO column magnitudes that will clearly impact on any inferred topdown emissions. Consistent with our previous work [Barkley et al., 2011], we advocate that satellite-based estimates of isoprene emissions over this region will be better characterized using two alternative bottom-up emission inventories to compute the HCHO AMFs and VCDs, than estimates based on a single emission scheme alone. Whether this is case for other areas, where isoprene emissions rates are significantly less than those from tropical vegetation, is unclear. However, we strongly advise that any future topdown estimates of isoprene emissions should properly assess this source of uncertainty. Furthermore, since satellite derived emissions are the only way to constrain isoprene fluxes from the entire Amazon, it is requirement that retrieved HCHO VCDs are well validated and their errors/ biases appropriately characterized. In situ measurements of the HCHO profile and aerosol distribution within the lower troposphere coincident with each satellite's overpass would help resolve the accuracy of calculated AMFs and VCDs, but over the Amazon such measurements are rare. Given the large uncertainties associated with this regions's isoprene emissions, oxidation chemistry and aerosol loading, we therefore emphasize the need for a dedicated aircraft campaign over the Amazon that could provide such observations, complemented by ground-based observations (e.g.,

lidar or DOAS), to give a critical assessment of the satellite HCHO columns where arguably the data is most useful.

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