

Constraining the budget of atmospheric carbonyl sulfide using a 3-D chemical transport model

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Abstract.

Carbonyl sulfide (OCS) has emerged as a valuable proxy for photosynthetic uptake of carbon dioxide (CO₂) and is known to be important in the formation of aerosols in the stratosphere. However, uncertainties in the global OCS budget remain large, due mainly to three flux terms: vegetation and soil uptake, and oceanic emissions. Bottom-up estimates do not yield a closed budget, thought to be due to unaccounted-for tropical emissions of OCS. Here we present a simulation of atmospheric OCS over the period 2004-2018 using the TOMCAT 3-D chemical transport model aimed at better constraining some terms in the OCS budget. Vegetative uptake of OCS is estimated by scaling gross primary productivity (GPP) output from the Joint UK Land Environment Simulator (JULES), using the leaf relative uptake (LRU) approach. The remaining surface budget terms are taken from available literature flux inventories, and adequately scaled to bring the budget into balance.

The model is compared with limb-sounding satellite observations made by the Atmospheric Chemistry Experiment – Fourier Transform Spectrometer (ACE-FTS) and surface flask measurements from 14 National Oceanic and Atmospheric Administration – Earth System Research Laboratory (NOAA-ESRL) sites worldwide.

We find that calculating vegetative uptake using the LRU underestimates the surface seasonal cycle amplitude (SCA) in the NH mid and high latitudes, by approximately 37 ppt (35%). The inclusion of a large tropical source is able to balance the global budget, but further improvement to the SCA and phasing would likely require a flux inversion scheme.

Compared to co-located ACE-FTS OCS profiles between 5 km and 30 km, TOMCAT remains within 25 ppt (approximately 5% of mean tropospheric concentration) of the measurements throughout the majority of this region and lies within the standard deviation of these measurements. This provides confidence in the representation of atmospheric loss and surface fluxes of OCS in the model. Atmospheric sinks account for 154 Gg S of the annual budget, which is 10 – 50% larger than previous studies. Comparing the surface monthly anomalies from the NOAA-ESRL flask data at all sites to the model simulations shows a RMSE range of 3.3 – 25.8 ppt. We estimate the total biosphere uptake to be 951 Gg S which is in the range of recent inversion studies (893 – 1053 Gg S), but our terrestrial vegetation flux accounts for 629 Gg S of the annual budget, which is lower other

recent studies (657 – 756 Gg S). However, to close the budget, we compensate this with a large annual oceanic emission term of 689 Gg S, focused over the tropics, which is much larger than bottom-up estimates (285 Gg S). Hence, we agree with recent findings that missing OCS sources likely originate from the tropical region.

This work shows that satellite OCS profiles offer a good constraint on atmospheric sinks of OCS through the troposphere and stratosphere and are therefore useful for helping to improve surface budget terms. This work also shows that the LRU approach is a suitable alternative to mechanistic approaches to quantifying vegetative uptake and will be valuable in using OCS to estimate GPP going forward. Future work will utilise a formal inversion scheme to better quantify the OCS budget.

1 Introduction

Carbonyl sulfide (OCS) is the most abundant of all sulfur-containing gases in the atmosphere and is important due to its potential use as a proxy for the photosynthetic uptake of carbon dioxide (CO₂) by vegetation (Sandoval-Soto et al., 2005; Montzka et al., 2007; Campbell et al., 2008; Suntharalingam et al., 2008; Blonquist et al., 2011; Berry et al., 2013; Launois et al., 2015b). Furthermore, due to its oxidation in the stratosphere, OCS is the largest source of sulfuric acid in the stratospheric aerosol layer in times of low volcanic activity (Crutzen, 1976; Kremser et al., 2016). In the troposphere, OCS has a global mean mixing ratio (mole fraction) of approximately 480 parts per trillion (ppt) with a lifetime of approximately 2.5 years (Montzka et al., 2007). In the stratosphere the OCS mixing ratio declines strongly with increasing altitude and has a longer mean lifetime than the troposphere, of approximately 64 ± 21 years (Barkley et al., 2008), ranging from 54.1 ± 9.7 years in the sub-tropics to 103.4 ± 18.3 years in the Antarctic (Hannigan et al., 2022).

Observations of OCS by the Network for the Detection of Atmospheric Composition Change (NDACC) using ground-based solar-viewing Fourier Transform Interferometers (FTIR) show weak positive trends between 2009 and 2016 in the troposphere at most of the 22 measurement sites of $<1\%$ yr⁻¹ (Hannigan et al., 2022). This trend is matched stronger positive trends in the stratosphere above all sites, up to $1.93 \pm 0.26\%$ yr⁻¹, except low latitude sites that show a negative trend. Furthermore, a downturn in free tropospheric OCS concentration reveals a negative trend between 2016 and 2020 at all sites (Hannigan et al., 2022). Kremser et al. (2015) showed positive OCS trends between 2001 and 2015, determined from ground-based Fourier transform spectrometer total column measurements at 3 southern hemisphere (SH) sites (also used by Hannigan et al. (2022)) and driven by changes to the tropospheric column. In contrast, the National Oceanic and Atmospheric Administration - Earth System Research Laboratory (NOAA-ESRL) global monitoring network has 14 sites and shows no consistent trend in surface OCS at any one location during the period of 2000 to 2005 (Montzka et al., 2007). Additionally, Glatthor et al. (2017) concluded that the tropospheric OCS budget is balanced based on a global Michelson Interferometer for Passive Atmospheric Sounding (MIPAS) satellite dataset (2002-2012), and ground-based partial-column measurements at the Jungfraujoch (46.5°N, 8.0°E) showed no significant trend in the free troposphere between 2008 and 2015 (Lejeune et al., 2017).

One of the main source of atmospheric OCS is oceanic emission, with total estimates ranging from 230 to 992 Gg S yr⁻¹ (Kettle et al., 2002; Montzka et al., 2007; Suntharalingam et al., 2008; Berry et al., 2013; Glatthor et al., 2015; Kuai et al., 2015;

Launois et al., 2015a; Lennartz et al., 2021; Ma et al., 2021; Remaud et al., 2022). Oceanic emission has 3 main contributions: direct emission of OCS, oxidation of emitted dimethyl sulphide (DMS) and oxidation of emitted carbon disulphide (CS₂). Both light-dependent (photochemical) and light-independent production play a role in oceanic emission (Launois et al., 2015a), the former linked primarily to incident UV radiation at the sea surface and the latter far below the surface. Both are driven by biological production and are proportional to amounts of chromophoric dissolved organic matter (CDOM), especially at the surface, where it can act to absorb some of the available light (Lennartz et al., 2021). Furthermore, Lennartz et al. (2021) suggest the importance of direct ocean-emitted OCS and oxidized CS₂ exceeds that of oxidized DMS which accounts for only a small portion of the overall ocean borne OCS emissions.

Vegetative uptake is the most important sink of atmospheric OCS, and its magnitude is significantly more uncertain than the ocean flux, with estimates ranging from 210 to 2400 Gg S yr⁻¹ (Kettle et al., 2002; Sandoval-Soto et al., 2005; Suntharalingam et al., 2008; Berry et al., 2013; Glatthor et al., 2015; Kuai et al., 2015; Launois et al., 2015b; Kooijmans et al., 2021; Ma et al., 2021; Maignan et al., 2021; Remaud et al., 2022). OCS is consumed during the photosynthesis process, which proceeds along the same enzymatic pathways as CO₂ (Protoschill-Krebs et al., 1996). However, unlike for CO₂, this process is one-way due to the irreversible OCS hydrolysis reaction, catalysed by carbonic anhydrase (Protoschill-Krebs et al., 1996). OCS hydrolysis also occurs in soil, primarily catalysed by carbonic anhydrase contained in bacteria and fungi (Kesselmeier et al., 1999; Smith et al., 1999; Li et al., 2005; Seibt et al., 2006; Kato et al., 2008), as well as by other enzymes, such as nitrogenase, CO dehydrogenase and CS₂ hydrolase (Smith and Ferry, 2000; Masaki et al., 2021). Soil uptake is the second largest OCS sink, with an estimated annual net loss of 30 – 355 Gg S (Kettle et al., 2002; Montzka et al., 2007; Berry et al., 2013; Glatthor et al., 2015; Kuai et al., 2015; Kooijmans et al., 2021; Abadie et al., 2022). Other findings suggest that the seasonal variation in OCS soil uptake is relatively weak in boreal forest regions, but shows dependency on soil moisture (Sun et al., 2018). Soil has also been observed to act as an emitter of OCS in certain conditions, dependent such components as temperature, soil moisture, nitrogen content and incident solar radiation (Whelan et al., 2013; Maseyk et al., 2014; Spielmann et al., 2019; Kitz et al., 2020).

Chin and Davis (1993) presented one of the first attempts at quantifying the global OCS (and CS₂) budget terms, but these were subject to substantial uncertainties. However, multiple terms, such as atmospheric loss and volcanism, were subsequently used in the estimates presented by Watts (2000) and Kettle et al. (2002), the latter of which has been used as a benchmark for more recent studies (Montzka et al., 2007; Suntharalingam et al., 2008; Berry et al., 2013; Glatthor et al., 2015; Kuai et al., 2015). Analysis of flask and aircraft data spanning both hemispheres by Montzka et al. (2007) have offered the most significant updates since the aforementioned studies and suggests a vegetative sink (1115 Gg S yr⁻¹) up to five times larger than the estimate (240 Gg S yr⁻¹) presented by Kettle et al. (2002). Due to the negligible or weak-positive atmospheric OCS trend, this would suggest a larger source is required for balance. The general consensus is that this must originate in the tropical oceans, due to measurement peaks from satellite and aircraft observations (Glatthor et al., 2015; Kuai et al., 2015), as well as modelling estimates pointing to this region as an underestimated source (Berry et al., 2013; Launois et al., 2015a). There is opposition from (Lennartz et al., 2017), who estimate global oceanic emissions to be approximately 350 Gg S yr⁻¹, derived using a global

100 oceanic box model and measurements of surface waters, thus too low to account for this difference entirely. A recent study has also suggested there is an underestimation in previous gridded anthropogenic OCS flux inventories by 200 Gg S yr^{-1} , which could account for some of the deficit (Zumkehr et al., 2018), supported by measurements of OCS in firn air and ice core samples (Aydin et al., 2020). Top-down estimates by Ma et al.(2021), using an inversion scheme that assimilates surface flask observations, point to a tropical source of unknown origin but the inversion setup presented by Remaud et al.(2022) suggests
105 a large tropical OCS source of oceanic origin. Both studies downplay the likelihood if it being of exclusively oceanic origin, hence there is still substantial uncertainty in several of the global surface fluxes of OCS. These recent studies quantifying OCS flux inventories show less uncertainty than previous ones. However, to improve the inventories further, increased spatial coverage by ground-based and remote atmospheric OCS observations are required, as well as OCS flux measurements (Whelan et al., 2018).

110 In this study, we add a further model (TOMCAT) to those already employed to simulate global OCS distribution, with emphasis on a full vertical comparison extending through the troposphere and stratosphere (approximately 5 – 35 km). Three inventories of fluxes are used to drive the model in separate experiments, offering slightly different perspectives. Firstly, a control setup using those from Kettle et al.(2002) (the results of which are denoted TOMCAT_{CON}). Secondly an inventory using modified fluxes from Kettle et al. (2002) in addition to a vegetative uptake quantified using Gross Primary Productivity (GPP) in the
115 leaf relative uptake (LRU) approach (Campbell et al., 2008; Stimler et al., 2012; Asaf et al., 2013) and one from the literature, i.e. anthropogenic emissions from Zumkehr et al.(2018), TOMCAT_{OCS}. Finally, an inventory using newly available fluxes from recent literature, TOMCAT_{SOTA} (TOMCAT_{State-Of-The-Art}). Each inventory of OCS fluxes is used to drive the TOMCAT 3-D chemical transport model (CTM) over the time period 2004 – 2018, providing fresh insight into the magnitude and location of the fluxes of OCS and how this translates vertical information of OCS into improved understanding of both surface and
120 atmospheric fluxes. Furthermore, to evaluate misalignment in Southern Hemisphere (SH) stratospheric agreement between TOMCAT_{OCS} and satellite observations, additional simulations were performed to assess the influence of a hypothetical reduction in stratospheric photolysis would have on OCS distribution.

Sect. 2 summarises the data used for evaluating the model. The model setup and each flux inventory are described in Sect. 3. Results and comparisons with tropospheric and stratospheric satellite observations from the Atmospheric Chemistry
125 Experiment infrared Fourier transform spectrometer (ACE-FTS) instrument (Bernath, 2017; Boone et al., 2020) and measurements made by the NOAA-ESRL flask network (Montzka et al., 2007) are shown in Sect. 4, discussed further in Sect. 5 and concluding remarks are presented in Sect. 6.

2 Observations

2.1 Atmospheric Chemistry Experiment – Fourier Transform Spectrometer Observations

130 Onboard the SCISAT satellite, launched in August 2003, is the Atmospheric Chemistry Experiment infrared Fourier transform spectrometer (ACE-FTS), which operates in a solar occultation mode measuring radiation between 750 and 4400 cm^{-1} at a

spectral resolution of 0.02 cm^{-1} (Bernath et al., 2005; Bernath, 2017). Although the planned ACE mission duration was only two years, it now has a data record spanning 18 years. This longevity makes the ACE-FTS a valuable tool for measuring atmospheric trace gases and characterising their variability and trends. Atmospheric trace gas profiles are retrieved using a non-linear least squares global-fit approach on the measurement altitude grid (3 km vertical resolution), then interpolated on to a uniform 1 km grid. ACE-FTS is capable of measuring profiles for a number of trace gases, including OCS from 5 km (or cloud top) up to about 30 km. OCS is retrieved using microwindows of various widths between 2039.01 and 2057.52 cm^{-1} , including a band at 1950.10 cm^{-1} to minimise the impact of H_2O interference. Because the primary science mission of ACE-FTS is to measure atmospheric ozone distributions over Canada, the satellite's orbit is such that approximately 60% of all measurements are at latitudes poleward of $\pm 60^\circ$. However, over the course of a year measurements are taken over a wide range of latitudes, providing a wealth of data with which to validate global CTM simulations. For this study, ACE-FTS version 4.1 (hereafter ACE) retrieved profiles from February 2004 to December 2018 (approximately 98,000 profiles) (Boone et al., 2020) were used in the validation of the modelled TOMCAT OCS distribution. The version 4.1 retrievals incorporate a new instrumental line shape (Boone and Bernath, 2019) and utilise the 2016 High-Resolution TRANsmission molecular absorption database (HITRAN) data (Gordon et al., 2017). Systematic errors in OCS measurements occur as a result of contamination from other gases in the microwindow (and clouds), while random errors are induced by random fitting errors from the least squares analysis, both have generally been improved in the version 4.1 product, over version 3.6 (Boone et al., 2020).

2.2 NOAA-ESRL Flask Measurements

The surface OCS measurements described here are shown in Sect. 4; here we present a summary of the method of data collection (performed by the NOAA-ESRL network) and the site information (see Table 1). Flasks of ambient air have been collected approximately 1 to 5 times per month at 14 measurement sites across both hemispheres since early 2000. Measurements of the OCS concentrations within the flasks are made using gas chromatography and mass spectrometry at the NOAA-ESRL Boulder laboratories (Montzka et al., 2007). In this study, we use data from all the Halocarbons & other Atmospheric Trace Species (HATS) surface measurement sites for the purpose of validating the surface OCS concentrations from the TOMCAT model.

Table 1. NOAA-ESRL flask sampling site information for OCS measurements (Montzka et al., 2007).

Code	Name	Country	Latitude (°N)	Longitude (°E)	Elevation (metres)
ALT	Alert, Nunavut	Canada	82.5	-62.5	185
BRW	Utqiagvik (formerly Barrow), Alaska	United States	71.3	-156.6	11
CGO	Kennaook / Cape Grim, Tasmania	Australia	-40.7	144.7	94
HFM	Harvard Forest, Massachusetts	United States	42.5	-72.2	340
KUM	Cape Kumukahi, Hawaii	United States	19.6	-155.0	8
LEF	Park Falls, Wisconsin	United States	45.9	-90.3	472
MHD	Mace Head, County Galway	Republic of Ireland	53.3	-9.9	5
MLO	Mauna Loa, Hawaii	United States	19.5	-155.6	3397
NWR	Niwot Ridge, Colorado	United States	40.1	-105.6	3523
PSA	Palmer Station	Antarctica (United States)	-64.8	-64.1	10
SMO	Tutuila	American Samoa	-14.2	-170.6	42
SPO	South Pole	Antarctica (United States)	-90.0	-24.8	2810
SUM	Summit	Greenland	72.6	-38.4	3210
THD	Trinidad Head, California	United States	41.1	-124.2	107

3 Chemical transport modelling of OCS

160 3.1 TOMCAT Model Setup

We have used the TOMCAT 3-D off-line CTM (Chipperfield, 2006) to model atmospheric OCS. This model has been used in a wide range of studies, including recently to better constrain methane flux estimations (Wilson et al., 2016; Parker et al.,

2018), as a forward model for methane flux inversions (McNorton et al., 2018) and to investigate stratospheric ozone depletion (Claxton et al., 2019). In this work, TOMCAT is driven by meteorological reanalysis data (ERA-Interim) from the European
165 Centre for Medium-Range Weather Forecasts (ECMWF (Dee et al., 2011)). ERA-Interim convective mass fluxes are used following the scheme presented in Feng et al. (2011). The model distribution of OH is specified from pre-computed fields which vary monthly, but not inter-annually. The monthly distributions are taken from Spivakovsky et al. (2000) and scaled by a factor of 0.92 in accordance with Huijnen et al. (2010). The photolysis loss is based on precomputed rates from the full chemistry version of TOMCAT (Monks et al., 2017). Atmospheric OH loss accounts for approximately 120 – 130 Gg S yr⁻¹
170 (roughly 10% of the total OCS sink) of the TOMCAT_{OCS} budget and photolysis about a quarter of this, 2-3%. TOMCAT_{OCS} and TOMCAT_{SOTA} are spun for 10 years prior to 2004, and then run between 2004 and 2018 at a horizontal resolution of approximately 2.8° × 2.8° (T42 Gaussian grid), with 60 atmospheric layers from the surface up to 0.1 hPa, on a time-step of 6 hours. TOMCAT_{CON} is initialised using the distribution of TOMCAT_{OCS} at the end of the spin-up period, but is only run for just a single year, 2004, due to the negative trend and its purpose in this study as a point of reference, rather than a benchmark
175 for improvement. Surface flux fields of OCS were implemented within TOMCAT on a monthly 1° × 1° grid with varying inter-annual variability, depending on the inventory in use. These are mapped onto the model grid in a way that conserves local distributions and the total global flux. For comparison with ACE-FTS, the geopotential height output from the model is converted to altitude; this is done using the hypsometric equation at a reference pressure of 1000 hPa, and then interpolated on to the 1 km equidistant altitude grid used by ACE-FTS. Furthermore, the profiles outputted by TOMCAT are spatio-temporally
180 co-located with the ACE-FTS observations to provide a precise like-for-like comparison. Monthly mean surface concentrations are calculated from the flask observations made by the NOAA-ESRL network and compared with monthly mean TOMCAT output averaged across the time period, used for each respective setup.

3.2 Kettle Flux Inventory

The fluxes described in this section originate from the literature (Watts, 2000; Kettle et al., 2002; Suntharalingam et al., 2008),
185 and are used to run the control simulation, TOMCAT_{CON}. This model run is utilised as a comparison to the model driven by our new inventory of fluxes described in Sect. 3.3, TOMCAT_{OCS} and to TOMCAT_{SOTA}. TOMCAT_{CON} was initialised using OCS values in each grid box from TOMCAT_{OCS}, after 10 years (1994 – 2003) spin-up and run for only a single year (2004), due to the net negative budget from these fluxes of approximately -46 Gg S yr⁻¹.

Three of the six sources, used to simulate TOMCAT_{CON}, are oceanic: a direct OCS flux term, one due to oxidation of CS₂ and
190 one due to oxidation of DMS. These were converted to OCS emissions using molar conversion factors (Chin and Davis, 1993; Barnes et al., 1994). The OCS and CS₂ emission terms were quantified using a physio-chemical model, the main source being from photochemical production (Kettle et al., 2002). However, as DMS measurements are more abundant than OCS and CS₂, these were used to parameterize this flux (Kettle and Andreae, 2000). Anthropogenic OCS emissions consist of two factors, a direct term and one from the oxidation of CS₂, the latter being considerably larger. They are both calculated here using SO₂
195 fields from Watts (2000) due to the extensive datasets available and a relationship between the facilities that release SO₂ and

OCS, despite there being no direct chemical reaction (Kettle et al., 2002). The final source term is biomass burning scaled similarly to that in (Kettle et al., 2002), but varies according to the monthly climatology of (Duncan et al., 2003).

The three sink terms are an oceanic sink, soil uptake and a vegetative sink. The first was quantified using the same physio-chemical model used for the OCS and CS₂ source terms described above and covers the periods in the year where the direct oceanic emission of OCS flips from being a source to becoming a sink. These are focused mostly over extra-tropical open ocean regions and during each hemisphere's summer period. Gridded soil uptake was calculated by applying correction factors for temperature, ambient OCS and soil water content to a standardised uptake rate of 10 pmol m⁻² s⁻¹ (Kesselmeier et al., 1999). The monthly mean climatological data for the temperature and soil water content is taken from Sellers et al. (1995), where the soil water content is a percentage of saturation in the top 2 cm of soil. Anoxic soil emissions are neglected in this study. Finally, the vegetative uptake is calculated by employing a normalised difference vegetation index (NDVI) to scale net primary productivity (NPP) distribution from Fung et al. (1987). We also scale up this term to the quoted upper limit of 270 Gg S yr⁻¹ by Kettle et al. (2002). As mentioned in Section 3.1, removal of atmospheric OCS by OH loss and photolysis is also accounted for in the model.

The spatial distribution for the months of January, April, July and October for the vegetation and soil uptake and oceanic emissions used in TOMCAT_{CON} are presented in the supplement: Fig. S1, S2 and S3 respectively.

3.3 LRU approach and Modified Flux Inventory

TOMCAT_{OCS} uses an array of fluxes that is orientated around a calculated vegetative uptake term, F_{OCS}, using the LRU approach (Campbell et al., 2008; Stimler et al., 2012; Asaf et al., 2013). The calculation of F_{OCS} is explained in Section 3.3.1 and a summary of the accompanying fluxes is provided in Section 3.3.2, including the full budget in Table 2.

3.3.1 Calculating OCS Vegetative Uptake using Gross Primary Productivity

The new vegetative sink calculation, used in TOMCAT_{OCS}, differs fundamentally from the method described in Sect. 3.2, and used in the control model simulation, as the use of NPP has been shown to underestimate the seasonal amplitude in other modelling studies (Suntharalingam et al., 2008; Berry et al., 2013). Sandoval-Soto et al. (2005) suggested that using NPP to calculate OCS uptake would underestimate the global burden and therefore they recommend using gross primary productivity (GPP) as an alternative. Furthermore, they were the first to quantify deposition velocity ratios for CO₂ and OCS for different plant types, which previous studies had assumed to be equal.

$$F_{OCS} = GPP \frac{[OCS]}{[CO_2]} \times LRU, \quad (1)$$

Using Eq. (1) we calculated the vegetative flux of OCS (F_{OCS}), in units of Gg S yr⁻¹, by scaling GPP (Gg C yr⁻¹) using a leaf relative uptake (LRU) of 1.6; a mean value from gas-exchange measurements of 22 plant species (Stimler et al., 2012). LRU is the ratio of OCS assimilation rates to CO₂ at the leaf-scale, both normalised by their respective concentration, signified by the square brackets in Eq. (1), in **units of ppb**. Units of the calculations in TOMCAT_{OCS} are molecules cm⁻² s⁻¹, as this is the

typical for TOMCAT emissions. We then convert to Gg S yr^{-1} following simulation. Each time-step in the model a new F_{OCS} value is calculated, as $[\text{OCS}]$ is the concentration from the previous time-step, starting at a generic value of 500 ppt in 1994. The use of a constant LRU value has been found to contribute less to error in the calculation of F_{OCS} than differences in GPP
230 between models on a continental scale (Hilton et al., 2017). However, as there are available plant-function-type dependent LRU datasets, this will be considered for the future of this work to implement spatially varying LRU values (Seibt et al., 2010; Maignan et al., 2021).

The GPP flux used in our calculation, generated by the Joint UK Land Environment Simulator (JULES) model, applies the WATER and Global CHange (WATCH) Forcing Data methodology to ERA-Interim reanalysis (WFDEI) between 1979 and
235 2012 and uses Global Precipitation Climatology Centre (GPCC) precipitation data (Slevin et al., 2016). Only monthly data for 2010 was used, as the interannual variability in the amplitude of the GPP cycle is only about 1% (Chen et al., 2017). Monthly mean gridded CO_2 surface mixing ratios for 2010 from a TOMCAT simulation which assimilated surface flask observations of CO_2 are used for the CO_2 concentration (see Fig. S4 in the supplement) (Gloor et al., 2018). As we compare only monthly means at the surface and seasonal OCS to ACE-FTS, long-term inter-annual variability was not considered in the scope of this
240 work. Like the use of a single value of LRU, calculating F_{OCS} using inter-annually varying GPP, and CO_2 products will be considered in the future. Our resulting estimate of the mean global yearly value of F_{OCS} between 2004 and 2018 is 629 Gg S , which is nearly three times the value of Kettle et al. (2002) 240 Gg S , but is slightly under half that of the largest estimation of 1115 Gg S in Table 2 from Montzka et al. (2007), and over half that estimated by Launois et al. (2015b), 1335 Gg S yr^{-1} , using the ORCHIDEE land surface model. The spatial distribution of F_{OCS} for the months of January, April, July and October,
245 in 2010 only, is presented in the supplement: Fig. S5.

3.3.2 Balancing the OCS Budget

As the fluxes described in Section 3.2 are utilised in constructing the inventory for $\text{TOMCAT}_{\text{OCS}}$, with the exception of calculating the vegetative uptake and anthropogenic emissions (which are taken from Zumkehr et al. (2018)), the fluxes must be modified to suitably close the overall budget, which we assume to be in balance due to a negligible or weak trend in the
250 majority of the study period (Montzka et al., 2007; Kremser et al., 2015; Glatthor et al., 2017; Lejeune et al., 2017; Hannigan et al., 2022). As F_{OCS} is larger than the vegetative uptake term than that of Kettle et al. (2002), we scale up several of the emission terms described in Sect. 3.2, despite some of the difference was accounted for in the larger anthropogenic emissions. Furthermore, some of the fluxes were adjusted to better represent recent estimations in the literature, i.e., soil uptake.

$\text{TOMCAT}_{\text{OCS}}$ makes use of anthropogenic OCS emissions presented by Zumkehr et al. (2018). Like Kettle et al. (2002),
255 anthropogenic OCS emissions consist of two factors, a direct term and one from the oxidation of CS_2 , the latter being considerably larger. Eleven anthropogenic sources of OCS were quantified by Zumkehr et al. (2018) for the period 1980 – 2012, the largest contributions originating from residential and industrial coal usage and the rayon industry. Emission factors for each source are applied to country-scale industrial activity data, obtained from a wide range of sources, then gridded spatially and temporally based on a gridded proxy flux (Zumkehr et al., 2018).

260 Scaling OCS emitted from biomass burning (as described in Section 3.2) and anthropogenic sources was not considered suitable to balance increases to sink terms, as these are less uncertain than oceanic emissions. Furthermore, biomass burning is more focused in lower latitude agricultural regions and anthropogenic emissions tend to be focused over point sources, mostly in Asia.

The soil flux utilised for TOMCAT_{OCS} was calculated by Kettle et al. (2002) using the method described in Sect. 3.2 and
265 assumes a constant 500 ppt OCS ambient value in the scaling of the standardised uptake. Soil uptake was scaled by 2.5 times from 130 Gg S yr⁻¹ to 322 Gg S yr⁻¹ to bring it in line with literature findings that estimate soil uptake to be between 236 – 507 Gg S yr⁻¹ (Berry et al., 2013; Launois et al., 2015b; Ma et al., 2021; Remaud et al., 2022). These studies used different approaches; Berry et al. (2013) use a global carbon cycle model, SiB 3, to obtain a new estimate of soil uptake based on empirical data and a mechanistic understanding of the processes influencing OCS diffusion into soil. Launois et al. (2015b)
270 use H₂S soil deposition to infer OCS fluxes, as this is a by-product of the OCS hydrolysis reaction and therefore a proxy for OCS uptake. Ma et al., (2021) and Remaud et al. (2022) use inverse frameworks, the former estimate a combined vegetative and soil uptake of 1053 Gg S yr⁻¹ and the latter estimate a soil uptake of 236 Gg S yr⁻¹. Recent work using mechanistic soil uptake models (Ogée et al., 2016) suggest oxic soil uptake is lower than the estimates discussed here: Kooijmans et al. (2021) estimate an annual uptake of 89 Gg S yr⁻¹ and Abadie et al. (2022) estimates 126 Gg S yr⁻¹. These values do not yet align with
275 inversion studies, adding to the uncertainty in surface fluxes, especially in the tropics, that accounts for a large portion of terrestrial OCS uptake (Ma et al., 2021; Remaud et al., 2022).

Initial testing of our new fluxes in TOMCAT yielded low-biased simulated OCS concentrations at Northern Hemisphere (NH) NOAA-ESRL sites; ALT, BRW and MHD, but a seasonal cycle with appropriate amplitude (not shown), **the latter of which receives ocean air masses frequently**. To improve the agreement, the direct and indirect OCS ocean emissions arising from
280 DMS were increased by a factor of 2. These fluxes were chosen as their spatial distribution includes peaks in the Northern Atlantic and Pacific regions. When including the reduction implemented for these terms in the Southern Ocean, the global net increase for direct OCS and indirect OCS from DMS is roughly 10 Gg S yr⁻¹ and 7 Gg S yr⁻¹, respectively, which is relatively small compared to the changes to the vegetative and soil CS₂ fluxes.

Suntharalingam et al. (2008) recommend a reduction in direct OCS and indirect OCS emissions from DMS by 40% (as this
285 yielded the smallest root-mean-squared error in their analysis) in SH mid-latitude (ML) and high-latitude (HL, here defined as 60° – 90°) regions, due to the resulting improvements to the seasonal cycle at Antarctic NOAA-ESRL sites. They also implemented an enhanced OCS tropical ocean source that was aseasonal and uniform across the tropics. However, here we scale up the CS₂ source term to 439 Gg S yr⁻¹ to balance the increased vegetation and soil sink terms discussed above and bring the net budget to near balance. We scaled this flux not necessarily because it was suspected that CS₂ was the erroneous term
290 in the OCS budget, but because it is more realistic to add a flux that is focused spatially over the tropical region already (Kuai et al., 2015). The reason for this geographical distribution is that CS₂ emissions are proportional to temperature and incident solar radiation, hence why the tropics show the strongest emissions (Kettle et al., 2002). Bottom-up estimates of global annual direct and indirect oceanic emissions total approximately 285 – 345 Gg S yr⁻¹ (Lennartz et al., 2017, 2021). Hence, not enough

to account for the discrepancy in the global OCS budget. However, for the purposes of this study, we allocate the discrepancy
295 into oceanic emissions, due to the co-location of CS₂ emission fields over the tropics and this is the most suitable representation.
Using the flux inventory described here, TOMCAT_{OCS} simulations were carried out covering 2004 to 2018, initialised at 500
ppt in every grid-box and spun up for 10 years between 1994 and 2004. Average yearly burdens for 2004 to 2018 yield a
broadly closed OCS budget. Inter-annual variability in meteorology will have an impact on the model's ability to have a mean
closed budget over the full time period. The vegetative flux sits roughly in the middle of literature estimates, but the total sink
300 term (1144 Gg S yr⁻¹) is similar to larger estimates from Berry et al. (2013) and Ma et al. (2021), as seen in Table 2, as well as
estimates from Glatthor et al. (2015), Kuai et al. (2015) and Launois et al. (2015b), not shown in Table 2. Atmospheric
destruction, mainly in the form of tropospheric loss from OH and stratospheric photolysis reactions, account for approximately
154 Gg S yr⁻¹ removal, which is 25% larger than fields used in earlier studies in Table 2, of roughly 126 Gg S yr⁻¹, derived by
Watts (2000). The total oceanic emission has been increased 146% from the starting point of Kettle et al. (2002); the majority
305 of this increase is focused in the tropical region. With a global net annual emission of 1141 Gg S, roughly equal to that of our
sink terms, the model yields 14 years of broadly balanced OCS budget, with all terms broadly in line with the findings of recent
studies (Berry et al., 2013; Glatthor et al., 2015; Kuai et al., 2015; Launois et al., 2015b; Ma et al., 2021; Remaud et al., 2022).
The spatial distribution for the months of January, April, July and October for the adjusted soil uptake and oceanic emissions
used in TOMCAT_{OCS} are presented in the supplement: Fig. S6 and S7, respectively.

310

Table 2. Global OCS budgets (units Gg S yr⁻¹). Values for past studies are an average of the upper and lower limits stated in those studies, unless a value is stated exactly. Values for this study are an average between 2004 and 2018.

Source/Sink Process	Kettle et al. (2002)	Montzka et al. (2007)	Suntharalingham et al. (2008)	Berry et al. ^a (2013)	Ma et al. ^b (2021)	Remaud et al. (2022)	TOMCAT_{OCS} Fluxes
Vegetation	-238	-1115	-490	-738	-1053	-657	-629
Oxic Soil	-130	-127	-120	-355		-236	-322
Reaction with OH	-94	-96		-101	-101	-100	-122
Reaction with O(¹ D)	-11	-11	-130	0	0	0	0
Photolysis	-16	-16		0	-40	0	-32
Ocean	0	0	0	0	0	0	-39
<i>Total Sinks</i>	-489	-1365	-740	-1194	-1194	-993	-1144
Ocean (OCS)	41	40			40		
Ocean (CS ₂)	84		230	876	81	269	689
Ocean (DMS)	154	240			156		
<i>Total Ocean Emission</i>	279	280	230	876	277	269	689
Anthropogenic (OCS)	64	64		64	155		410
Anthropogenic (CS ₂)	116		180	116	188	398	
Anthropogenic (DMS)	1	0		1	6		0
<i>Total Anthropogenic Emission</i>	181	64	180	181	349	398	410
Biomass Burning	38	106	70	136	136	53	42
Other (mainly wetlands & anoxic soils)	26	66	25	0	0	0	0
<i>Total Sources</i>	523	516	505	1193	1187	720	1141
<i>Net Budget</i>	34	-849	-235	-2	-432	-273	-3

^aOcean emission term includes an additional photochemical oceanic flux of 600 Gg S.

^bPosterior estimates from the Su inversion are shown here. An ‘unknown’ term is used to quantify a missing source in the inversion and balance their budget.

315

3.4 State-of-the-art Flux Inventory

Emissions used in the TOMCAT_{SOTA} simulation use 5 unique fluxes, which vary monthly, and unlike those used in TOMCAT_{OCS} and TOMCAT_{CON} vary inter-annually. The 5 sectors in use here are vegetation and soil uptake, as well as

oceanic, anthropogenic and biomass burning emissions. Due to the biomass burning and anthropogenic emissions only being
320 available between 2010 and 2015, 2015 fluxes are repeated through 2016 – 2018. The same is done for 2010 fluxes for the
period of 2004 to 2009 for all 5 emission or sink fields.

The sink due to vegetation was derived by implementing the mechanistic OCS vegetative uptake model from Berry et al.
(2013), which calculates uptake based on leaf surface area, saturation and air pressure, into the land surface model ORCHIDEE,
which is explained in detail by Maignan et al. (2021). Additionally, in the same study, this approach was compared to the
325 LRU-GPP approach, used in the calculation of OCS in Section 3.3.1, by running the two in the LMDz6 CTM. This shows that
while the mechanistic approach works better on shorter time and smaller spatial scales, it is not adequate for global estimation,
unlike the LRU-GPP approach. Maignan et al. (2021) propose the idea of implementing soil fluxes into the ORCHIDEE land
surface model, simultaneously with vegetation, which would be a significant step forward in the capability of constraining and
quantifying surface uptake. The reason being: both uptake of OCS by soil and vegetation follows very similar enzymatic
330 pathways, catalysed by carbonic anhydrase (Protoschill-Krebs and Kesselmeier, 1992; Kesselmeier et al., 1999). Preliminary
work on implementing a mechanistic soil uptake model, originating from Ogée et al. (2016), into ORCHIDEE was used as the
soil flux in this work (Abadie et al., 2022). Estimates for each flux are $-532 \text{ Gg S yr}^{-1}$ and $-264 \text{ Gg S yr}^{-1}$ for vegetation and
soil respectively. Note, the vegetation estimate does not match that of Maignan et al. (2021) ($-756 \text{ Gg S yr}^{-1}$).

Oceanic emissions constitute two parts, direct OCS and indirect CS_2 emissions. Direct OCS is estimated using a global box
335 model and supplemented by measurements of OCS, where the former is developed from von Hobe et al. (2003) and further
improves the quantification of the photoproduction rate, parameterization of light-independent production and employs
satellite observations of CDOM for use in the model (Lennartz et al., 2017, 2021). Indirect emissions are estimated using CS_2
concentration measurements at the surface and converted using a molar conversion ratio of 0.81 (Chin and Davis, 1993;
Lennartz et al., 2017). Biomass burning emissions are estimated by Stinecipher et al. (2019) using the Global Fire Emissions
340 Database, version 4 (GFED4), and scaling CO emissions to OCS. GFED4 utilises six biomass burning categories: savanna and
grassland, boreal forests, temperate forests, tropical deforestation and degradation, peatland fires, and agricultural waste
burning. Their estimates for the period 1997-2016 total $60 \pm 37 \text{ Gg S yr}^{-1}$. Finally, anthropogenic emissions are from the study
by Zumkehr et al. (2018), as described in Section 3.3.2, which account for roughly 402 Gg S yr^{-1} of OCS emissions per year.
The spatial distribution for the months of January, April, July and October (2010 only) for the vegetation and soil uptake and
345 oceanic emissions used in $\text{TOMCAT}_{\text{SOTA}}$ are presented in the supplement: Fig. S8, S9 and S10 respectively.

4 Results

$\text{TOMCAT}_{\text{OCS}}$ and $\text{TOMCAT}_{\text{CON}}$ are compared with NOAA-ESRL surface flask monthly mean measurements and monthly
anomalies (monthly mean minus annual mean). $\text{TOMCAT}_{\text{SOTA}}$ is compared only to monthly anomalies, due to a negative trend
in the budget. All 3 models are co-located to the nearest grid box and altitude to the measurements. $\text{TOMCAT}_{\text{OCS}}$ is also co-
350 located and compared to ACE, which has approximately 98,000 profiles in the modelled time period, all of which are filtered

for outliers before analysis. As ACE measures the upper troposphere and stratosphere primarily, this region is less sensitive to surface processes, so we only compare TOMCAT_{OCS}. Furthermore, as TOMCAT_{CON} and TOMCAT_{SOTA} have a negative trend, this makes correcting for bias in both the troposphere and stratosphere and making a comparison throughout the entire profile challenging. An additional TOMCAT_{OCS} simulation with adjusted atmospheric photolysis for the year 2010 is presented and
355 used to test the suitability of this change to correct a negative model bias in the SH stratosphere.

4.1 Seasonality of Modelled OCS compared to Surface Flask Measurements

TOMCAT simulates OCS distributions down to the surface, where the majority of OCS fluxes occur; it is therefore important that the model performs well at this level. Figure 1 compares the NOAA-ESRL surface flask measurements (black) with TOMCAT_{OCS} (blocked-blue), and TOMCAT_{CON} (orange) simulations. As TOMCAT_{CON} was only run for 2004 (see Section
360 3.1), a blue dashed line representing 2004 of TOMCAT_{OCS} is also indicated (TOMCAT_{OCS} - 2004). Monthly standard deviation is calculated for each site and visualised using error bars associated with the observations and TOMCAT_{OCS}. The modelled vertical layer of TOMCAT_{OCS} and TOMCAT_{CON} closest to the altitude of the measurement site was used for closer comparison, because the bottom-most model layer does not necessarily correspond with the surface due to the relative coarseness of model grid boxes affecting the simulated surface topography.

365 Comparisons between TOMCAT_{OCS} and TOMCAT_{CON} are shown here to emphasise the improvements made by the flux inventory developed in this study, and TOMCAT_{SOTA} to present the latest bottom-up estimates of OCS fluxes. Generally, there is an improvement in RMSE across all the sites in Fig. 1, but in some cases, there is a degradation, which is mostly attributed to average concentration. Figure 2 presents the monthly anomalies for all model runs, including TOMCAT_{SOTA}, as well as the RMSE just for the seasonality (i.e., excluding influence of average concentration), and SCA values for all model
370 runs and the surface observations. Therefore, we can dissect the influence of changes to seasonality influencing RMSE, to some extent. For example, SMO shows a reduction in RMSE from TOMCAT_{CON} (27.6 ppt) to TOMCAT_{OCS} (19.3 ppt) of 30% in Figure 1, but also a reduction in RMSE of 28% in Figure 2. Thus, indicating not only has TOMCAT_{OCS} improved representation of average concentration, but the seasonality has also improved. The fluxes used to model TOMCAT_{OCS} reduces the RMSE from an annual mean of 24.3 ppt in TOMCAT_{CON} at all sites to 21.2 ppt (an error reduction of 12.5%). This improves
375 to 5.1 ppt (20.4%) if we are to exclude MHD, a particularly poorly represented site according to this metric.

The surface observations show that OCS concentrations peak in April or May in the NH (ranging from 505 to 540 ppt) and reach a minimum in September or October (ranging from 386 to 488 ppt), which is consistent in all 10 NH sites and resembles the seasonality of CO₂. Despite several of the NH sites being particularly far north (70 – 90 °N), photosynthesis is still the dominant driving flux, emphasising the strength of the OCS vegetative uptake signal. Phasing of the seasonal cycle in the SH
380 shifts several months earlier, peaking in February and troughs in August, driven by the seasonality of oceanic emissions.

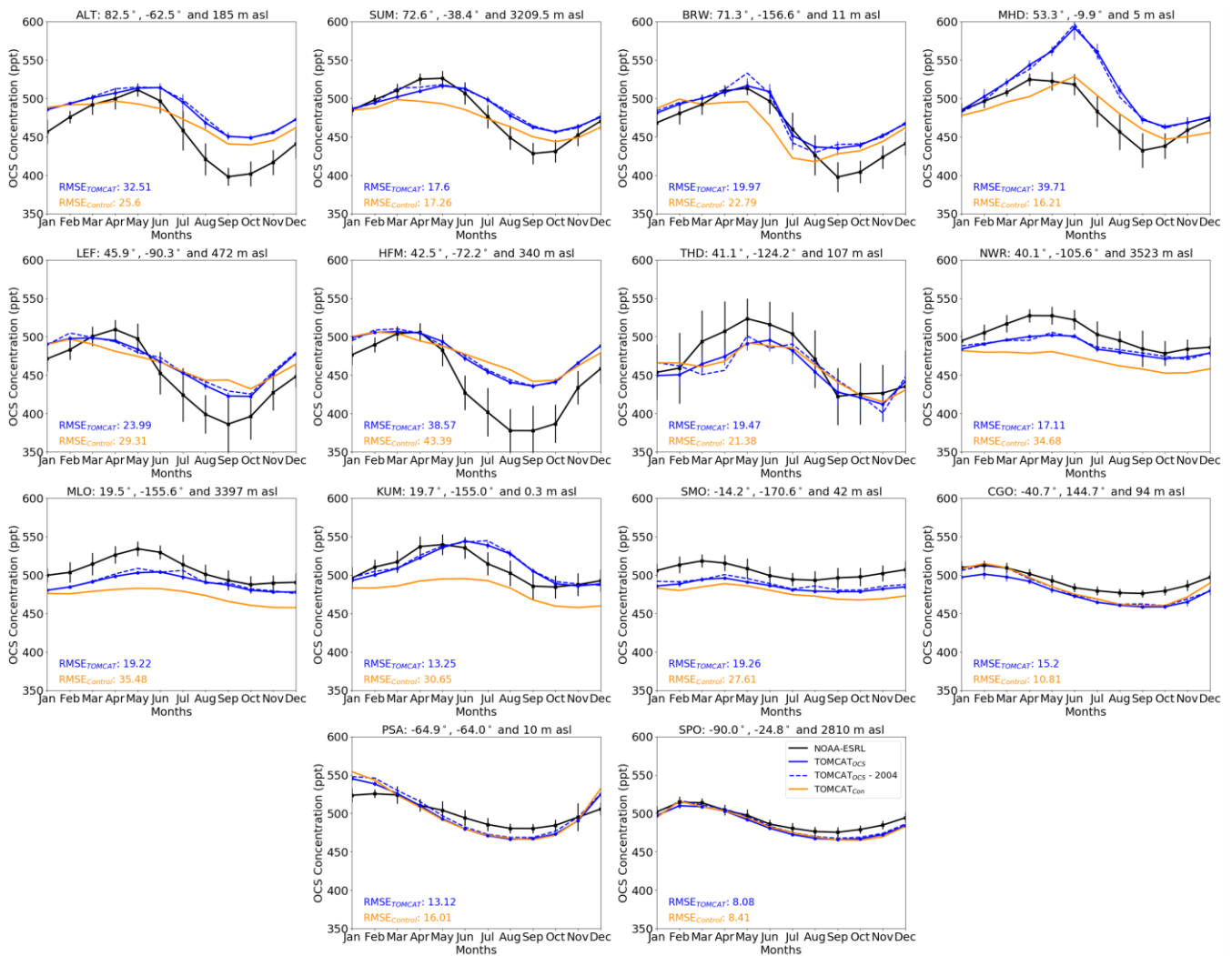


Figure 1. Monthly mean OCS concentration (in ppt) at NOAA-ESRL flask sites (black lines) compared with TOMCAT_{OCS} (blocked-blue line) for the 2004 to 2018 period. The dashed blue line is just 2004 for the TOMCAT_{OCS} dataset and is compared to TOMCAT_{CON} (orange line). Geographical location of each site is referenced in the titles. Altitude above sea level (asl) of the site is stated and nearest level in TOMCAT used for comparison. Error bars for both NOAA-ESRL and TOMCAT_{OCS} represent standard deviation. Units of RMSE are in ppt.

SCA presented in Figure 2 show an improvement from a mean absolute difference to the observations of ± 30.5 ppt (39.9%) in TOMCAT_{CON} and to ± 26.5 ppt (34.8%) in TOMCAT_{OCS}. These metrics suggest that the flux inventory used in TOMCAT_{OCS} offers an improvement in capturing seasonality and observation representation at the surface. The mean absolute difference in SCA of TOMCAT_{SOTA} compared to NOAA-ESRL is ± 43.7 ppt (57.2%).

Relative to the flask measurements, the SCA at the 8 NH continental measurement sites (top 8 plots in Fig. 2) is captured more successfully by TOMCAT_{OCS} than TOMCAT_{CON}. At all sites there is some improvement in TOMCAT_{OCS} SCA, except for BRW, which shows little change, but an improved RMSE (Fig. 2), and MHD which shows an overestimated SCA by 39.0%

in TOMCAT_{OCS}. SCA is underestimated in TOMCAT_{CON} output at all 8 of these sites by approximately 38.8 ppt on average.
395 TOMCAT_{OCS} improved this disparity to an absolute difference of 36.9 ppt (MHD is overestimated). TOMCAT_{SOTA} shows an absolute difference in SCA to NOAA-ESRL of 45.4 ppt, which improves substantially if LEF and HFM are to be ignored, to 26.9 ppt. Neglecting these same sites for TOMCAT_{OCS} and TOMCAT_{CON}, we see that TOMCAT_{SOTA} shows the best performance in SCA compared to NOAA-ESRL at the continental NH sites.

LEF and HFM are dense woodland sites and have particularly large SCA which can often be a challenge for models to simulate.
400 Here, we show SCAs from TOMCAT_{OCS} of 76 ppt at LEF and 71 ppt at HFM, compared to observed values of 123 ppt and 128 ppt, respectively. The underestimation in TOMCAT_{OCS} could potentially be attributed to using a single value for the LRU parameter globally, as this value is known to vary significantly between plant types (Stimler et al., 2012). The method of estimating OCS uptake using LRU, clearly underestimates OCS uptake over dense vegetation, as the model is likely to coarse to include the heavily depleted OCS concentration near the surface. As the GPP product has been compared to a spatially
405 gridded GPP from the FLUXNET network, including Harvard Forest and Park Falls, it is unlikely an underestimation in GPP in this geographical region (Slevin et al., 2016). TOMCAT_{SOTA} underestimates SCA at both sites by approximately 100 ppt, highlighted by RMSE values of 36.1 and 41.2 ppt respectively in Fig. 2.

ALT, SUM and BRW are located at high northern latitudes, where the landscape has significantly less vegetation and is more homogeneous than at LEF and HFM, although the seasonal cycle is still driven by typical NH processes. Phasing of the peak
410 and trough of the annual seasonal cycles at ALT, SUM and BRW are improved in TOMCAT_{OCS} output, compared to TOMCAT_{CON}, but RMSE and SCA are not improved significantly in Fig. 2. TOMCAT_{SOTA} improves SCA at ALT, BRW and MHD (9.9 ppt absolute difference to NOAA-ESRL SCA) compared to TOMCAT_{OCS} (39.4 absolute difference to NOAA-ESRL SCA). However, TOMCAT_{SOTA} exhibits larger RMSE values in Fig. 2 due to poor phasing of the seasonality. THD and MHD are both coastal sites and the misalignment in observed and modelled seasonal cycles is attributed to the impact from
415 the ocean fluxes in adjacent model grid boxes. Additionally, capturing the seasonal cycle of trace gases at a site such as MHD can be particularly challenging as there are significant seasonal changes in advected air masses. Berry et al. (2013) show overestimated peak concentration at MHD in their adjusted flux model runs, but posterior inversions by Ma et al. (2021) show a good alignment at this site, suggesting that the underlying cause is poorly represented surface fluxes.

At particularly high-altitude sites, NWR and MLO, Fig. 1 shows that TOMCAT_{OCS} underestimates the average concentration,
420 additionally showing no significant improvement in SCA from TOMCAT_{CON}. The measured SCA at SUM is 98 ppt (428 to 526 ppt), which is modelled relatively poorly by TOMCAT_{CON}, 54.8 ppt, and improved by TOMCAT_{OCS} to 60.1 ppt. A significant difference between SUM and the other two locations is that the topography in the grid boxes for NWR and MLO is very spatially variable; for example, MLO is a high volcano on a relatively small island in the Pacific Ocean. These results suggest the model underestimates OCS concentrations around 1 – 3 km above sea level and there is little improvement between
425 TOMCAT_{OCS} and TOMCAT_{CON}.

The two NH tropical sites, MLO and KUM, exhibit a seasonal cycle in the measurements similar to that of NH continental sites, with slightly different phasing and a reduced seasonal amplitude, which is due to the influence of oceanic processes.

Conversely, SMO, in the SH tropics, is more dominated by ocean processes and peaks earlier in the year. TOMCAT_{OCS} at MLO, KUM and SMO shows varying levels of agreement with the observations. The RMSE in Fig. 1 at KUM is reduced by 56.8% and improved SCA compared to TOMCAT_{CON} by 81%, from 17.7 ppt to 3.4 ppt. MLO and SMO average concentration is better represented by TOMCAT_{OCS}, both observing RMSE improvement of -45.8% and -30.2%, respectively. Also note in Fig. 1 that the seasonality is out of phase with the observations, peaking approximately 1 month too late, while KUM is 2 months late. A challenge in diagnosing the misalignment in the Hawaiian sites is their proximity to the ocean, as the 2.8° × 2.8° grid box is dominated by oceanic flux, as are all the boxes around it.

All four SH sites SMO, CGO, PSA and SPO show lower SCA in the observations than all NH sites, ranging from 25 ppt at SMO to 45 ppt at PSA, which is 80% and 65% less variation, respectively, than the forested site HFM. This emphasises the impact vegetation has on NH OCS seasonal cycle. Unlike SMO, the latter 3 sites are dominated much more by oceanic fluxes peaking in SH summer due to the association of phytoplankton growth with OCS emissions, driven by solar radiation. The average concentration of OCS is underestimated by TOMCAT_{OCS} and TOMCAT_{CON} at CGO, PSA and SPO. Seasonal amplitude is overestimated in all model runs for all three sites. TOMCAT_{OCS} overestimates the SCA at CGO, SPO and PSA, compared to the flask measurements by 5.1 ppt, 6.1 ppt and 33.6 ppt, respectively. In contrast, TOMCAT_{CON} overestimates by 18.2 ppt, 43.2 ppt and 11.8 ppt, respectively. This suggests the reduction in Southern Oceanic emissions in TOMCAT_{OCS} improves seasonality in OCS adequately but could be reduced further. TOMCAT_{SOTA} shows a much larger overestimation in seasonality.

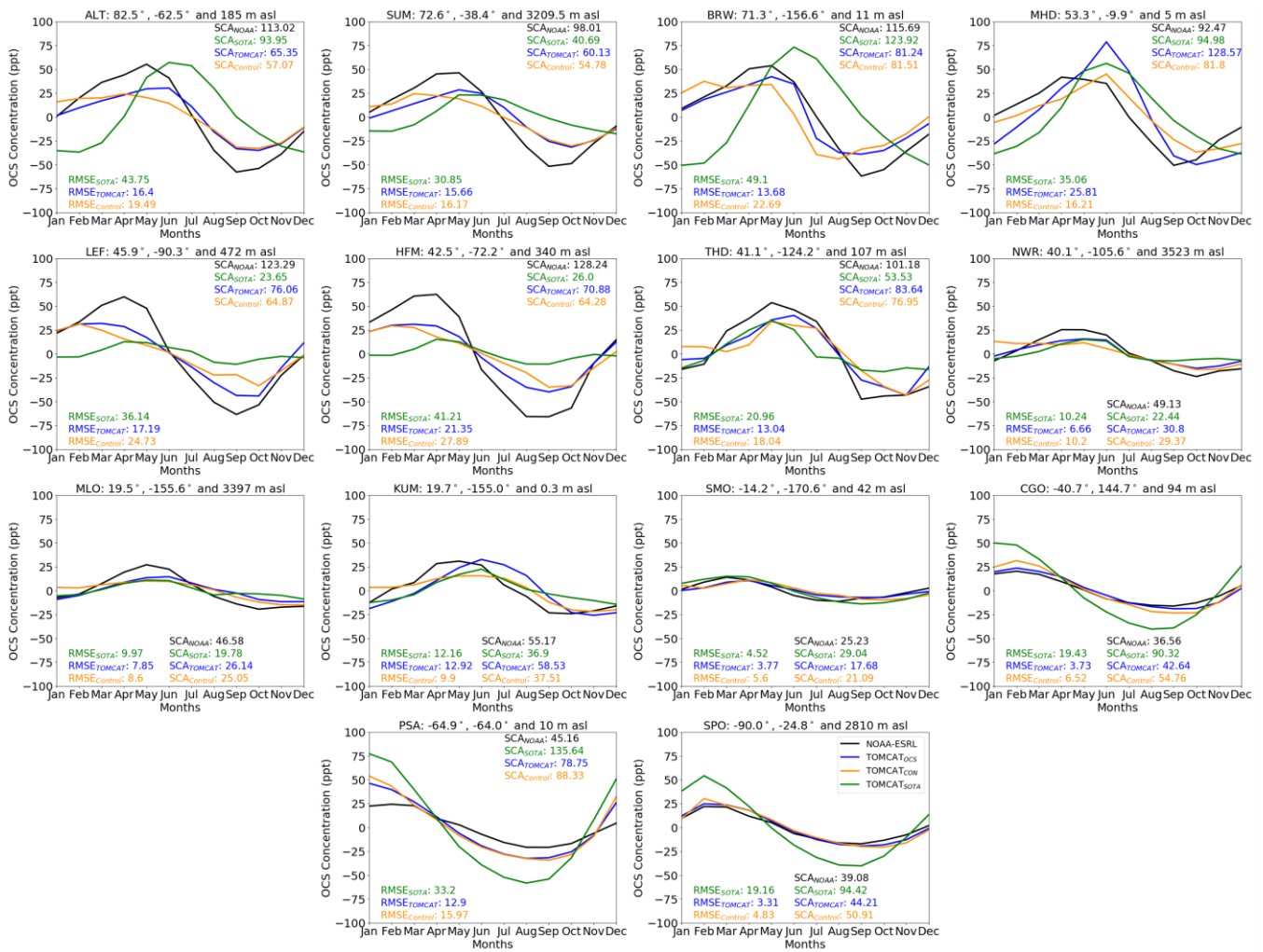


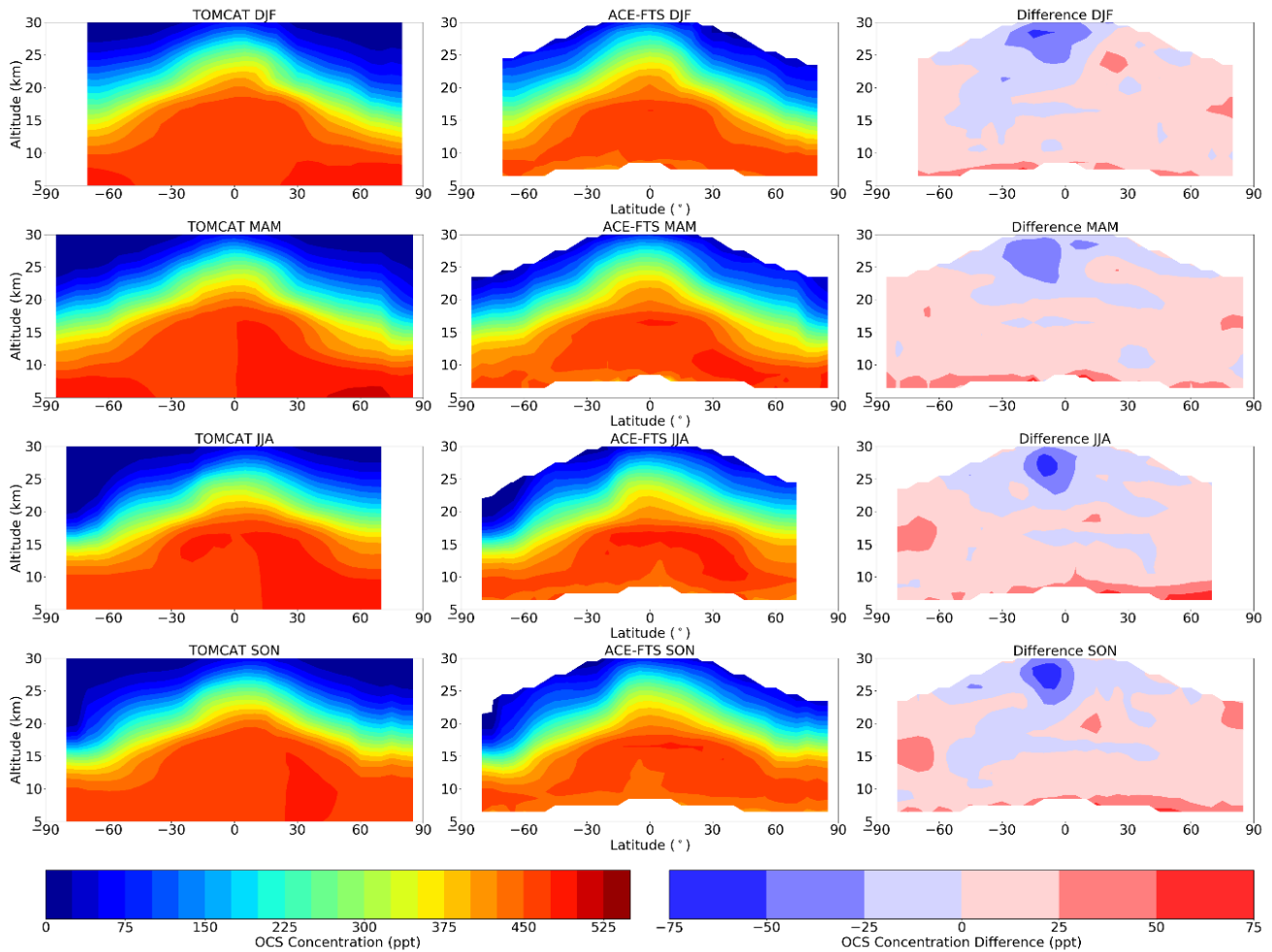
Figure 2. Monthly mean OCS anomalies (monthly mean less annual mean, in ppt) at NOAA-ESRL flask sites (black line) compared with TOMCAT_{OCS} (blue line), TOMCAT_{CON} (orange line) and TOMCAT_{SOTA} (green line) for the 2004 to 2018 period. Geographical location of each site is referenced in the titles. Altitude above sea level (asl) of the site is stated and nearest level in TOMCAT used for comparison. Units of RMSE and SCA are in ppt.

445
450 **4.2 Spatial Distribution of Modelled OCS compared to Satellite Observations**

Figure 3 shows the spatial distribution of atmospheric OCS obtained by averaging ACE profiles across all longitudes and in 5° latitude bins (central column), along with the TOMCAT_{OCS} profiles averaged in the same way (left column). The difference between the two datasets is shown in the right column (TOMCAT_{OCS} minus ACE). ACE-FTS is capable of measuring at altitudes between 6.5 km and 30.5 km, depending on latitude. Figure 3 shows that ACE tropospheric OCS mixing ratios (middle column) range from 425 to 500 ppt, peaking in the upper troposphere-lower stratosphere (UTLS) region, which extends from about 7 km in the NH ML and up to 17 km in the tropics. OCS values decline above and below the UTLS due to removal by vegetation and soil uptake at the surface and photochemistry in the stratosphere, leaving a peak in between which is

significantly more prevalent in March – May (MAM) and June – August (JJA). As there is relatively little photosynthesis in the December – February (DJF) and MAM periods, OCS builds up in the atmosphere, followed by net removal throughout
460 JJA and September – November (SON). Despite NH photosynthesis beginning slightly before JJA, there is a clear lag in removing OCS from the upper troposphere. The seasonal peak in OCS in the UTLS region only fully disappears in SON, suggesting there is roughly a 3-month delay on the influence of surface processes on the UTLS ambient mixing ratio. While this fluctuation is driven by seasonality in photosynthesis, the OCS peak is particularly large and extends lower in the atmosphere in the NH ML region, co-located with regions of year-round especially large anthropogenic emissions.

465 The tropopause height is captured adequately by TOMCAT_{OCS} (which is forced by ERA-Interim) and is visible in the homogeneity of the difference around the UTLS. TOMCAT_{OCS} agrees with ACE to within 25 ppt throughout most of the troposphere, which is about 5% of the average estimated atmospheric value of OCS (484 ppt) (Montzka et al., 2007). Similar to the seasonal pattern visible in ACE, the tropospheric OCS mixing ratio in TOMCAT_{OCS} peaks before the NH growing season, MAM. The maximum OCS concentration in TOMCAT_{OCS} can be seen below 6 km (around 50 – 70°N), peaking in
470 MAM, is larger than maximum OCS observed by ACE, and persists throughout most of the year. The overestimation in the NH ML is broadly contained within a discrepancy of 25 – 50 ppt from ACE, with the exception of a few anomalies in MAM and JJA, potentially attributed to underestimated or slower surface OCS uptake, or overestimated anthropogenic emissions in this region. The rate of removal of OCS is not quick enough in JJA to match the measurements exactly. This positive bias in the model below 10 km in the SH throughout DJF and MAM is probably unrelated to the NH positive model bias and would
475 likely be resolved by weaker oceanic emission in the SH, despite already having been reduced by 40% (Suntharalingam et al., 2008).

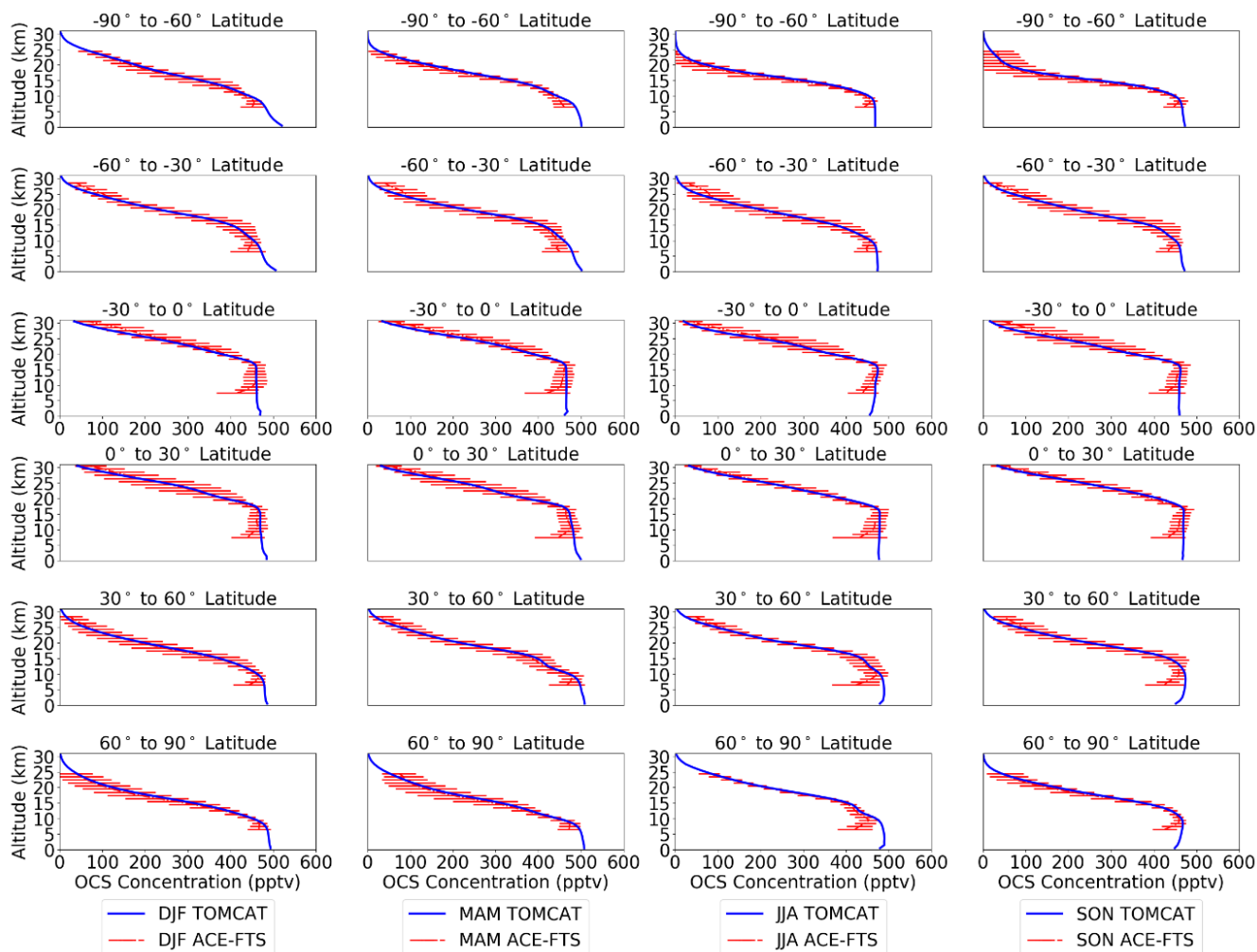


480 **Figure 3.** Seasonal zonal mean concentration (mixing ratio) of OCS (ppt) from TOMCAT_{OCS} (left), ACE (centre) and the difference between the two (TOMCAT_{OCS} minus ACE, right) for the period of 2004 to 2018. TOMCAT_{OCS} and ACE data averaged in 5-degree latitude bins and over all longitudes.

Differences between the model and observations in the stratosphere are broadly similar to those in the troposphere and are within ± 25 ppt. However, there is considerable model underestimation at 24.5 – 30.5 km between 0° and 30°S, of up to 65 ppt. This region shows a mean seasonal underestimation of between 24.6% and 18.6%, with a peak difference in JJA of 47.5% around 29.5 km at 10°S. As this feature does not follow the pattern of the inter-tropical convergence zone shifting with the
 485 hemispheric summertime period, it is unlikely that vertical fluxes are underestimated. The declining gradient in the stratosphere is steeper in the model than in ACE, which suggests that more OCS is being destroyed via photochemical processes in the model than in reality, which we examine in Section 4.3.

Figure 4 shows TOMCAT_{OCS} profiles (blue) in 30° latitude seasonal bins compared to ACE (red), including the standard deviation (shown as error bars) of ACE at each altitude. It is clear the model replicates the vertical structure of OCS, when

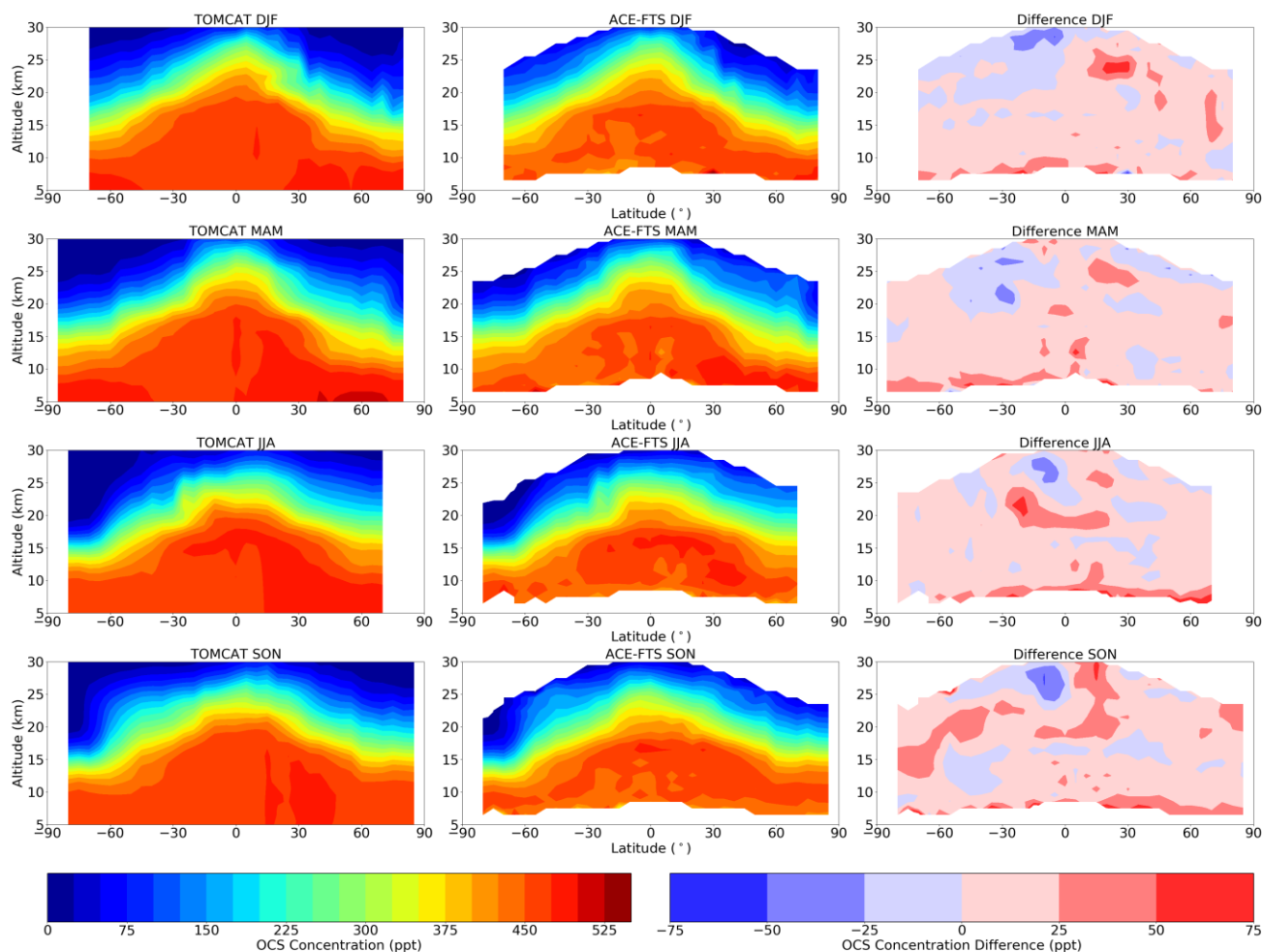
490 compared to observed OCS profiles from ACE-FTS. The negative discrepancy in the SH tropical stratosphere, visible in Fig. 3 and discussed above, can be seen in the third row of Fig. 4, as TOMCAT_{OCS} deviates from ACE from 20 km up to 30 km. However, as it remains within standard deviation of ACE throughout the entire profile, this suggests the upper atmospheric sinks are modelled moderately well by TOMCAT_{OCS}. This applies to most of the profiles compared in Fig. 4, in that the modelled TOMCAT_{OCS} profiles generally remain within the standard deviation of ACE. The positive model biases in both
 495 hemispheres below 10 km in Fig. 3 can be seen in Fig. 4, such that the trend at these altitudes in TOMCAT_{OCS} generally does not match ACE. Between 30°S and 90°N, ACE shows a depletion in OCS towards the surface from as high as 15 km – driven by surface uptake. Where a more neutral or increasing gradient between 90°S and 30°S is seen, as there is minimal vegetative uptake and a seasonal cycle strongly influenced by oceanic emission in this region (see Fig. 1).



500 **Figure 4.** Seasonal mean vertical profiles of OCS concentration (mixing ratio, ppt) from TOMCAT model output (blue) and ACE (red) for six different latitude regions. The error bars are standard deviation of ACE at each altitude level. All profiles are seasonal averages between the years 2004 to 2018.

4.3 Modelled OCS using reduced photochemical loss

The TOMCAT_{OCS} model setup described in Section 3.1 and 3.3 is used for this experiment in which just the year 2010 is run
505 with an adjusted atmospheric photolysis rate of 75%. The intensity of this simulation is to make a simple preliminary
assessment of if photolysis alone will correct the underestimation in the SH stratosphere. Figure 5 shows the modified version
of TOMCAT_{OCS} on the left, ACE measurements made in 2010 in the middle and the difference between the two (TOMCAT_{OCS}
less ACE). When compared with Figure 3, the reduced removal of OCS improves the differences between the model and
measurements above approximately 20 km, which is to be expected, as this is generally where photolysis is active. The regions
510 that exceed a difference of ± 25 ppt are limited to isolated pockets throughout the year and noticeably more in SON, in the
tropics around 20 – 30 km and the SH around the tropopause. Using only a year of data removes a lot of the smoothing we see
in Figure 3, which accounts for some of the differences in Figure 5. However, there are similar features between the two,
specifically persistent (but slightly reduced) underestimation in the model in the tropical SH. We also see an increase in positive
model bias, most obviously in the NH between 0 and 30 °N in DJF and SON. Overall, we find that differences between the
515 model and measurements were merely shifted by a reduction in photolysis and while a reduction by 25% improved these, it
introduced misalignment elsewhere. Further testing, including a simulation using a photolysis rate reduced by 50% exacerbates
these differences further (see Fig. S11), drawing the conclusion that transport or convection requires correction to fully resolve
these issues in the model.



520 **Figure 5.** Seasonal zonal mean concentration (mixing ratio) of OCS (ppt) from TOMCAT_{OCS} (left), only for the year 2010 and with a photolysis rate 0.75 times that of TOMCAT_{OCS}, ACE (centre) and the difference between the two (TOMCAT_{OCS} minus ACE, right) for the period of 2004 to 2018. TOMCAT_{OCS} and ACE data averaged in 5-degree latitude bins and over all longitudes.

5 Discussion

After a 10-year spin-up period, the TOMCAT_{OCS} simulations of atmospheric OCS concentrations and the vegetative flux, which are dependent on one another in the model, are in equilibrium between 2004 and 2018. By utilizing a GPP dataset, we estimate a mean yearly vegetative OCS uptake of 629 Gg S yr⁻¹, which is within the range and uncertainty of the magnitude of this flux from previous top-down studies (see Table 2), but slightly smaller than a bottom-up estimates by Kooijmans et al. (2021) and Maignan et al., 2021) (753 – 756 Gg S yr⁻¹). Our total vegetative and soil sinks agree with findings from Berry et al. (2013) and inversion studies by Kuai et al. (2015), Ma et al. (2021) and Remaud et al. (2022) approximately ±150 Gg S yr⁻¹ (15%). We balance the OCS budget by implementing an enlarged CS₂ emission source, for the exclusive reason that it is

focused over the tropics (Kettle et al., 2002), rather than it being a flux originating from oxidized CS₂. Bottom-up estimates recommend a constraint on global oceanic emission of OCS to approximately 285 – 350 Gg S yr⁻¹ (Lennartz et al., 2017, 2020, 2021), significantly lower than the fluxes required to balance our budget and thus bringing our tropical ocean estimate into question. It is clear that tropical fluxes are still uncertain, however inverse modelling of OCS fluxes shows that some
535 combination of a larger tropical oceanic source and vegetative sink resolves the budget and produces adequate model comparison with independent observations (Ma et al., 2021; Remaud et al., 2022).

TOMCAT_{OCS} output agrees with ACE-FTS profiles of OCS within 25 ppt throughout the majority of the observed atmosphere (approximately 5 km – 30 km), suggesting the sinks in the upper atmosphere are modelled well, with the exception of some discrepancies in the lower troposphere and the tropical lower stratosphere. Photochemical destruction is important in our
540 understanding of atmospheric OCS, and due to photolysis in the stratosphere, the model displays a declining vertical gradient above the tropopause. Our total estimate for this flux is 154 Gg S yr⁻¹, an upward revision of about 40% compared to the previous work of Kettle et al. (2002) and larger than all other estimates in Table 2. Comparison of TOMCAT_{OCS} with ACE profiles shows an excellent representation of the free troposphere, suggesting that we have found a suitable balance of fluxes at the surface, the spatial variability of which requires improvement. Overestimation in the NH ML region in JJA and SON
545 suggests that surface emissions could be slightly overestimated or that surface uptake does not initialise quick enough or strong enough to remove OCS from the atmosphere at the start of the growing season.

To assess the OCS surface seasonality modelled by TOMCAT_{OCS} we compare it to 2 other simulations: TOMCAT_{CON} and TOMCAT_{SOTA}. The vegetation, soil and ocean emission fields used to drive these models can be found in the supplementary material. When compared with surface flask observations, we show the OCS budget used to calculate TOMCAT_{OCS} reduces
550 RMSE compared to the control, TOMCAT_{CON}, at most sites by approximately 25%, as much as 57% at KUM, but degrading some RMSE, notably at MHD (see Fig. 1). We also show improvements in RMSE at NH continental sites, especially the forested sites of LEF and HFM, but there is still moderate underestimation in NH vegetative uptake. Comparing RMSE in the monthly anomalies between TOMCAT_{OCS} and TOMCAT_{CON} (see Fig. 2) shows that improved average concentration contributed significantly to improving RMSE in Fig. 1, as TOMCAT_{OCS} improves SCA by only 5% compared to
555 TOMCAT_{CON}.

The Hawaiian sites, MLO and KUM, show significantly improved RMSE, and some improvement in SCA and phasing, suggesting an enhancement in tropical oceanic emission is reasonable. The lack of OCS measurements in the tropics poses a challenge to both quantifying surface OCS exchange in this region both from a mechanistic perspective and from constraining inverted fluxes (Whelan et al., 2018; Ma et al., 2021; Remaud et al., 2022). While TOMCAT_{OCS} shows an adequate comparison
560 with tropical surface sites and a vertical comparison within the variability of ACE, we acknowledge that no attempt has been made in this work to experiment with reducing tropical surface OCS uptake, which has been suggested as an alternative solution to balance the OCS budget (Ma et al., 2021). Overestimation in TOMCAT_{OCS} SCA at SH sites CGO, PSA and SPO, indicates a reduction in oceanic emissions in this region is necessary, due to the limited continental landmass and associated uptake.

565 The method of estimating vegetative uptake using the LRU approach does have limitations, such as calculating OCS uptake
using a constant LRU value of 1.6 is not representative of reality. LRU values have shown to vary from approximately 1.0 to
4.0 based on different plant type and atmospheric conditions (Sandoval-Soto et al., 2005; Seibt et al., 2010; Stimler et al.,
2010, 2012). Our estimation of vegetative uptake in this work does not replicate OCS uptake universally and it is unclear if
570 this is due to localised differences in LRU or on the GPP fields themselves. While soil uptake has been scaled appropriately
according to the literature, the distribution is based on work by Kettle et al. (2002) and has since been updated, for example
by Ogée et al. (2016) and Abadie et al. (2022). When compared to bottom-up vegetation uptake estimates from the ORCHIDEE
model used to drive TOMCAT_{SOTA} (Fig. S8), the LRU approach shows similar spatial distribution, magnitude, and seasonality
(Fig. S5). There are notable differences in tropical location and magnitude year-round, i.e., South America in January and
Africa in April. These regions should have low impact on seasonality in the mid and high latitudes in the NH. In contrast,
575 comparing soil uptake used in TOMCAT_{OCS} vs. TOMCAT_{SOTA} (Fig. S6 vs. S9), also estimated using ORCHIDEE, these show
considerable differences which account for the different seasonal cycles in the NH, particularly ALT, SUM, BRW and MHD.
Figure S9 shows reasonably homogeneous distribution and seasonality compared to S6, which shows fair more annual
variability and spatial variation.

6 Conclusions

580 The 3-D chemical transport modelling setup in this study was used to compare three simulations, one utilising the LRU
approach to quantifying vegetative uptake and a series of scaled fluxes (TOMCAT_{OCS}), and two using bottom-up fluxes
originating from the literature; TOMCAT_{CON} and TOMCAT_{SOTA}. Where TOMCAT_{CON} uses fluxes estimated by Kettle et al.
(2002) and TOMCAT_{SOTA} uses a series of novel fluxes from recent literature (Lennartz et al., 2017; Zumkehr et al., 2018;
Stinecipher et al., 2019; Lennartz et al., 2020; Maignan et al., 2021). All simulations are compared with surface anomalies
585 from the NOAA-ESRL flask network and TOMCAT_{OCS} is compared ACE-FTS satellite observations. This study is novel in
the extended time period analysed and the quality of vertical comparison with the most recently available ACE-FTS satellite
measurements (version 4.1). Furthermore, we see an excellent comparison with ACE-FTS throughout most of the atmosphere,
which suggests the free troposphere and gradient above the UTLS is well represented by TOMCAT_{OCS}, and therefore so too
are the sources and sinks driving the model. Which shows promise for future OCS work using TOMCAT.
590 TOMCAT_{OCS} and TOMCAT_{CON} surface concentration is compared, and the former is shown to reduce root mean square error
(RMSE) compared to 14 NOAA-ESRL by 12.5% across all 14 sites, up to 20% when neglecting MHD. Further, surface
anomalies (monthly mean minus annual mean) are compared between all three model runs, yielding a RMSE that removes
annual mean and focuses solely on seasonality. TOMCAT_{OCS} reduces the RMSE in the anomalies by 18.7% and 52.4%
compared to TOMCAT_{CON} and TOMCAT_{SOTA}, respectively. Adequately modelling seasonality globally proved to be a
595 challenge and while TOMCAT_{OCS} performed best relative to NOAA-ESRL flask observations, the seasonal cycle amplitude
was still underestimated by 26.6 ppt. We have shown that the LRU approach for quantifying vegetative uptake yields similar

annual estimates (629 Gg S yr⁻¹) to mechanistic and inversion approaches (657 – 756 Gg S yr⁻¹) and resembles spatial variability (Maignan et al., 2021; Remaud et al., 2022). To suitably estimate a total biosphere uptake that reflects recent inversion studies (893 – 1053 Gg S yr⁻¹ (Remaud et al., 2022; Ma et al., 2021)) soil uptake is uplifted to by 2.5 times, yielding a combined total of 951 Gg S yr⁻¹. To bring the budget into balance we increase total net oceanic emissions to 650 Gg S yr⁻¹, from the starting point of 279 Gg S yr⁻¹ (Kettle et al., 2002). Overall, we draw similar to conclusions to other work, that the tropics are a suitable location for a compensatory source of OCS. Recommendations for advancing this work are below.

To improve the GPP-LRU approach **the following changes are necessary in future, such as** using inter-annually varying GPP and CO₂ mixing ratios, as well as a temporally and spatially resolved LRU. The latter of these **3** changes is challenging to achieve at a high resolution on a global scale, however, plant-functional-type-dependent datasets of LRU are available (Seibt et al., 2010; Whelan et al., 2018; Maignan et al., 2021), hence an initial step would be just to vary LRU based on ecosystem on a continental or ecosystem scale. Advances are being made in this area, with mechanistic and LRU approaches emerging that reduce uncertainty in OCS vegetative uptake (Kooijmans et al., 2021; Maignan et al., 2021). The use of an enhanced tropical ocean source to balance the budget is justified in this work by offering suitable satellite and surface observational comparisons. However, we acknowledge that oceanic emissions alone may not account for this discrepancy and based on TOMCAT_{OCS} performance at MLO and KUM, this is not a perfect solution for balancing the global OCS budget.

While we have shown that TOMCAT_{OCS} compares well with satellite observations, the region between the surface and approximately 6 km, which is not measured by ACE-FTS, could hold a lot of information useful in resolving surface fluxes. Measurements at the surface are sensitive to minor flux changes, although in the well-mixed mid to upper troposphere these spatial changes are less important. Validation of model output to ground-based Fourier Transform spectrometer column OCS measurements could improve our understanding and ability to model the lower troposphere. Furthermore, incorporating measurements with vertical information into an inversion scheme has been shown to improve the posterior OCS fluxes, specifically using HIPPO flight data (Ma et al., 2021). Therefore, further study following on from this work will be to derive an a posteriori set of fluxes using an inversion scheme based on an up-to-date prior, using surface observations and a dataset containing vertical information near the surface.

Code and Data Availability

Anthropogenic OCS emission data are available at <https://portal.neresc.gov/project/m2319/> (Campbell, 2022; Zumkehr et al., 2018). GPP dataset is available at <https://datashare.ed.ac.uk/handle/10283/2080> (Slevin et al., 2016). ACE-FTS data are available at <http://www.ace.uwaterloo.ca/data.php> (ACE-FTS, 2022). NOAA-ESRL surface flask measurements of OCS are available at <https://www.esrl.noaa.gov/gmd/dv/data/> (ESRL Global Monitoring Laboratory, 2022). Model data are available at <https://doi.org/10.5281/zenodo.6368542>.

Author Contribution

630 Model runs and data analysis were performed by MPCar, with support from RJP. JJH and MPChi designed the study. CJW provided the CO₂ model data. Control OCS emissions were provided by PS. TOMCAT model is maintained and updated by the team at the University of Leeds: MPChi, WF, CW and RJP. Manuscript was written by MPCar, with contributions from all co-authors.

Competing Interests

The authors declare that they have no conflict of interest.

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