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Direct oceanic emissions unlikely to account for the missing source of atmospheric carbonyl sulfide

Sinikka T. Lennartz¹, Christa A. Marandino¹, Marc von Hobe², Pau Cortes³, Birgit Quack¹, Rafel Simo³, Dennis Booge¹, Andrea Pozzer⁴, Tobias Steinhoff¹, Damian L. Arevalo-Martinez¹, Corinna Kloss², Astrid Bracher^{5,6}, Rüdiger Röttgers⁷, Elliot Atlas⁸, and Kirstin Krüger⁹

Correspondence to: Sinikka T. Lennartz (slennartz@geomar.de)

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Abstract. The climate active trace-gas carbonyl sulfide (OCS) is the most abundant sulfur gas in the atmosphere. A missing source in its atmospheric budget is currently suggested, resulting from an upward revision of the vegetation sink. Tropical oceanic emissions have been proposed to close the resulting gap in the atmospheric budget. We present a bottom-up approach including (i) new observations of OCS in surface waters of the tropical Atlantic, Pacific and Indian oceans and (ii) a further improved global box model to show that direct OCS emissions are unlikely to account for the missing source. The box model suggests an undersaturation of the surface water with respect to OCS integrated over the entire tropical ocean area and, further, global annual direct emissions of OCS well below that suggested by top-down estimates. In addition, we discuss the potential of indirect emission from CS₂ and dimethylsulfide (DMS) to account for the gap in the atmospheric budget. This bottom-up estimate of oceanic emissions has implications for using OCS as a proxy for global terrestrial CO₂ uptake, which is currently impeded by the inadequate quantification of atmospheric OCS sources and sinks.

1 Introduction

Carbonyl sulfide (OCS) is the most abundant reduced sulfur compound in the atmosphere. It enters the atmosphere either by direct emissions, e.g., from oceans, wetlands, anoxic soils or anthropogenic emissions, or indirectly via oxidation of the short-lived precursor gases dimethylsulfide (DMS) and carbon disulfide (CS₂) (Chin and Davis, 1993; Watts, 2000; Kettle, 2002). Both precursor gases are naturally produced in the oceans, and CS₂ has an additional anthropogenic source (Kettle, 2002; Stefels et al., 2007; Campbell et al., 2015). With direct and indirect marine emissions combined, the ocean is considered as the dominant source of atmospheric OCS (Chin and Davis, 1993; Watts, 2000; Kettle, 2002). The most important sink of atmospheric OCS is uptake by terrestrial vegetation (Brown and Bell, 1986; Protoschill-Krebs and Kesselmeier, 1992; Campbell et al., 2008) and oxic soils, while chemical loss by photolysis and reaction with the hydroxyl radical (OH) in the atmosphere are minor loss processes (Chin and Davis, 1993; Watts, 2000; Kettle, 2002). While tropospheric volume mixing ratios show a distinct annual cycle (Montzka et al., 2007), the interannual to decadal variation is low (Montzka et al., 2007; Kremser et al., 2015).

¹GEOMAR Helmholtz Centre for Ocean Research Kiel, Düsternbrooker Weg 20, 24105 Kiel, Germany

²Forschungszentrum Jülich GmbH, Institute of Energy and Climate Research (IEK-7), Wilhelm-Johnen-Strasse, 52425 Jülich, Germany

³Institut de Ciencies del Mar, CSIC, Pg. Maritim de la Barceloneta, 37-49, 08003 Barcelona, Catalonia, Spain

⁴Max Planck Institute for Chemistry, Hahn-Meitner-Weg 1, 55128 Mainz, Germany

⁵Alfred Wegener Institute Helmholtz Centre for Polar and Marine Research, Bussestrasse 24, 27570 Bremerhaven, Germany

⁶Institute of Environmental Physics, University of Bremen, 28334 Bremen, Germany

⁷Helmholtz-Zentrum Geesthacht, 21502 Geesthacht, Germany

⁸Rosenstiel School of Marine and Atmospheric Science, Miami, FL 33149, USA

⁹University of Oslo, Department of Geosciences, 0315 Oslo, Norway

Table 1. Missing source estimates derived from top-down approaches: the listed studies used an increased vegetation sink and an a priori direct and indirect ocean flux to estimate the magnitude of the missing source. Assigning the missing source to oceanic emissions results in the total ocean flux listed here. Fluxes are given in Gg S per year.

Reference	A priori ocean flux	Missing source	Total ocean flux
Suntharalingam et al. (2008)	235	230	465
Berry et al. (2013)	276	600	876
Kuai et al. (2015)	289	800	1089
Glatthor et al. (2015)	276	714	992

Accurate accounts of sources and sinks of atmospheric OCS are crucial for two reasons.

- First, OCS is climate-relevant because it influences the radiative budget of the Earth as a greenhouse gas and by contributing significant amounts of sulfur to the stratospheric aerosol layer (Crutzen, 1976; Brühl et al., 2012; Notholt et al., 2003; Turco et al., 1980) that exerts a cooling effect (Turco et al., 1980; Kremser et al., 2016). The two opposite effects are currently in balance (Brühl et al., 2012), but future changes in atmospheric circulation, as well as the magnitude and distribution of OCS sources and sinks, could change that. Hence, a better understanding of the tropospheric budget is needed to predict the effect of OCS in future climate scenarios (Kremser et al., 2016).
- Second, OCS has recently been suggested as a promising tool to constrain terrestrial CO2 uptake, i.e., gross primary production (GPP), as it is taken up by plants in a similar way as CO₂ (Asaf et al., 2013). GPP, a major global CO2 flux, can only be inferred from indirect methods, because the uptake of CO₂ occurs along with a concurrent release by respiration. Unlike CO₂, OCS is irreversibly degraded within the leaf. GPP can thus be estimated based on the uptake ratio of OCS and CO₂, from the leaf to regional scale (Asaf et al., 2013) or even global scale (Beer et al., 2010), under the condition that other sources are negligible or well quantified. The magnitude of terrestrial biogeochemical feedbacks on climate has been suggested to be similar to that of physical feedbacks (Arneth et al., 2010). In order to reduce existing uncertainties, it is thus crucial to better constrain single processes in the carbon cycle, especially GPP.

Nonetheless, current figures for tropospheric OCS sources and sinks carry large uncertainties (Kremser et al., 2016). While the budget has been previously considered closed (Kettle, 2002), a recent upward revision of the vegetation

sink (Sandoval-Soto et al., 2005; Suntharalingam et al., 2008; Berry et al., 2013) led to a gap, i.e., a missing source in the atmospheric budget of 230–800 Gg S per year (Suntharalingam et al., 2008; Berry et al., 2013; Kuai et al., 2015; Glatthor et al., 2015) (Table 1), with the most recent estimates at the higher end of the range. This revision of vegetation uptake was suggested to (i) take into account the different deposition velocities of CO₂ and OCS within the leaf and base it on GPP instead of net primary production (Sandoval-Soto et al., 2005) as well as (ii) to better reproduce observed seasonality of OCS mixing ratios in several atmospheric models (Berry et al., 2013; Kuai et al., 2015; Glatthor et al., 2015). Based on independent top-down approaches using MIPAS (Glatthor et al., 2015) and TES (Kuai et al., 2015) satellite observations, FTIR measurements (Wang et al., 2016), and NOAA ground-based time series stations and the HIPPO aircraft campaign (Berry et al., 2013; Kuai et al., 2015), the missing source of OCS was suggested to originate from the (tropical) ocean, most likely from the region of the Pacific warm pool. Other potential sources such as advection of air masses from Asia have been discussed (Glatthor et al., 2015) but not tested. If the ocean was to account for the missing source, the total top-down oceanic source strength would then be the a priori oceanic flux plus the missing source estimate of each inverse model simulation (Table 1). This addition would imply a 200-380 % increase in the a priori estimated oceanic source. If oceanic direct and indirect emissions were to account for the total missing source, an ocean source strength of $465-1089 \,\mathrm{Gg}\,\mathrm{S}\,\mathrm{yr}^{-1}$ would be required (Table 1).

OCS and its atmospheric precursors are naturally produced in the ocean. In the surface open ocean, OCS is present in the lower picomolar range, and has been measured on numerous cruises in the Atlantic (Ulshöfer et al., 1995; Flöck and Andreae, 1996; Ulshöfer and Andreae, 1998; von Hobe et al., 1999), including three latitudinal transects (Kettle et al., 2001; Xu et al., 2001), the Indian Ocean (Mihalopoulos et al., 1992), the Pacific Ocean (Weiss et al., 1995a) and the Southern Ocean (Staubes and Georgii, 1993). Measurements in tropical latitudes, where the missing source is assumed to be located, have previously been performed in the Indian Ocean (Mihalopoulos et al., 1992) and during the Atlantic transects (Kettle et al., 2001; Xu et al., 2001). OCS is produced photochemically from chromophoric dissolved organic matter (CDOM) (Andreae and Ferek, 2002; Ferek and Andreae, 1984) and by a not fully understood lightindependent production that has been suggested to be linked to radical formation (Flöck et al., 1997; Pos et al., 1998). Dissolved OCS is efficiently hydrolyzed to CO₂ and H₂S at a rate depending on pH and temperature (Elliott et al., 1989). CS₂ has been measured in the Pacific and Atlantic oceans in a range of $7.2-27.5 \,\mathrm{pmol}\,\mathrm{L}^{-1}$ (Xie et al., 1998) and during two Atlantic transects (summer and winter) in a range of $4-40 \,\mathrm{pmol}\,\mathrm{L}^{-1}$ (Xu, 2001). It is produced photochemically (Xie et al., 1998) and biologically (Xie et al., 1999), and no significant loss process other than air-sea gas exchange has

Table 2. Global oceanic emission estimates of OCS: direct ocean emission estimates of OCS from bottom-up approaches. Uncertainties are given in parentheses as in the original paper either as range or \pm standard deviation.

Reference	Emitted S as OCS (Gg S yr ⁻¹)
Extrapolated from measurements	
Rasmussen et al. (1982)	320 (±160)*
Ferek and Andreae (1983)	245*
Johnson and Harrison (1986)	110-210*
Mihalopoulos et al. (1992)	230 (110-210)*
Chin and Davis (1993)	160 (85-340)*
Weiss et al. (1995b)	-16 (-10-30)*
Ulshöfer and Andreae (1998)	41-80*
Watts (2000)	53 (±80)*
Xu et al. (2001)	53*
Model simulations	
Kettle (2002)	41 (±154)
Launois et al. (2015a)	813 (573–3997)
This study	130 (±80)

^{*} Units deviate from original paper, converted to Gg S for comparison.

been identified (Xie et al., 1998). DMS is present in the lower nanomolar range in the surface ocean and has been extensively studied in several campaigns, summarized in a climatology by Lana et al. (2011). DMS is biogenically produced and consumed in the surface ocean, as well as photo-oxidized and ventilated by air–sea exchange (Stefels et al., 2007).

Available bottom-up estimates of the global oceanic OCS fluxes from shipboard observations range from -16 to 320 Gg S yr⁻¹ (Table 2). However, the highest estimates were biased, because mainly summertime and daytime observations of water concentrations were considered. With the discovery of the seasonal oceanic sink of OCS during wintertime (Ulshöfer et al., 1995) and a pronounced diel cycle (Ferek and Andreae, 1984), direct oceanic emissions were corrected downwards.

Only recently, OCS emissions have been estimated with the biogeochemical ocean model NEMO-PISCES (Launois et al., 2015a) at a magnitude of 813 Gg S yr⁻¹, sufficient to account for the missing source. This oceanic emission inventory had been used to constrain GPP based on OCS on a global scale (Launois et al., 2015b). However, the oceanic OCS photoproduction in the ocean model included a parameterization for OCS photoproduction derived from an experiment in the North Sea (Uher and Andreae, 1997b), which might not be representative for the global ocean, as indicated by photoproduction constants that were an order of magnitude lower in the Atlantic ocean compared to the German Bight (Uher and Andreae, 1997a).

Here, we present new observations in all three tropical ocean basins, two of them measured with unprecedented pre-

cision and time resolution. Direct fluxes were inferred from continuous OCS measurements in the tropical Pacific and Indian oceans, covering a range of regimes with respect to CDOM content, ultraviolet (UV) radiation and sea surface temperature (SST). These observations are used to further constrain and validate a biogeochemical box model which had previously been shown to reproduce OCS concentration in the Atlantic Ocean reasonably well (von Hobe et al., 2001). The box model is now updated from its previous global application (Kettle, 2002) by adding and further developing the most recent process parameterizations to estimate the global source strength of direct OCS emissions. The emission estimate is further complemented by discussing the potential of indirect OCS emissions, i.e., the emissions of short-lived precursor gases CS2 and DMS, to account for the gap in the budget.

2 Methods

2.1 Study sites

Several cruises were conducted to measure the trace gases OCS (OASIS, TransPEGASO, ASTRA-OMZ) and CS₂ (TransPEGASO, ASTRA-OMZ). Cruise tracks are depicted in Fig. 1. The OASIS cruise onboard RV *SONNE I* to the Indian Ocean started from Port Louis, Mauritius, to Malé, Maldives, in July and August 2014, where mainly oligotrophic waters were encountered. TransPEGASO was an Atlantic transect starting in Gibraltar and leading to Buenos Aires, Argentina, and Punta Arenas, Chile. It took place in October and November 2014 and covered a variety of biogeochemical regimes. ASTRA-OMZ onboard RV *SONNE II* started in Guayaquil, Ecuador, and ended in Antofagasta, Chile, in October 2015. Although 2015 was an El Niño year, upwelling together with high biological production was still encountered during the cruise (Stramma et al., 2016).

2.2 Measurement setup for trace gases

OCS was measured during two cruises onboard the RV SONNE I (OASIS) and SONNE II (ASTRA-OMZ) with a continuous underway system similar to the one described in Arévalo-Marténez et al. (2013), at a measurement frequency of 1 Hz. The system consisted of a Weiss-type equilibrator, through which seawater is pumped from approximately 5 m below the surface with a flow of 3- $4 \,\mathrm{L}\,\mathrm{min}^{-1}$. The air from the equilibrator headspace was Nafion-dried and continuously pumped into an OCS analyzer (model DL-T-100, Los Gatos Research) that uses the off-axis integrated cavity output spectroscopy (OA-ICOS) technique. The instrument used onboard is a prototype of a commercial instrument (www.lgrinc.com/documents/ OCS Analyzer Datasheet.pdf), developed by Los Gatos Research (LGR) in collaboration with Forschungszentrum Jülich GmbH (Schrade, 2011). Data were averaged over

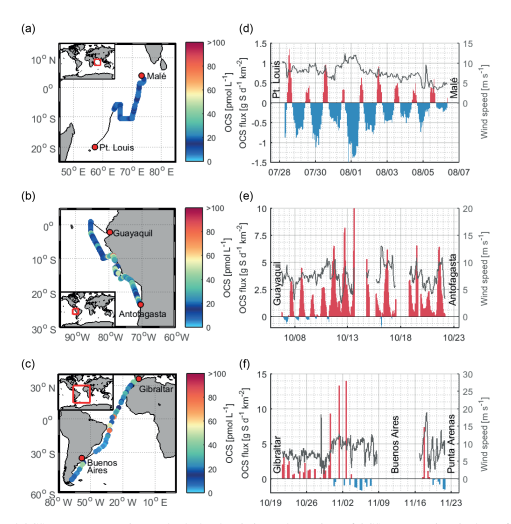


Figure 1. Observed OCS water concentrations and calculated emissions: observations of OCS concentrations in the surface ocean during the three cruises (a) OASIS, (b) ASTRA-OMZ, and (c) TransPEGASO as well as the corresponding emissions calculated based on the concentration gradient between water and marine boundary layer (d-f). Outgassing is indicated in red bars; oceanic uptake in blue bars. The grey line shows wind speed measured onboard the vessels. Flux data are shown with different scales on the y axes. Data gaps occurred during stays in port and territorial waters or during instrument tests.

2 min, achieving a precision of 15 ppt. OCS mixing ratios in the marine boundary layer (MBL) were determined by pumping outside air ca. 50 m from the ship's deck to the OCS analyzer (KNF Neuberger pump). A measurement cycle consisted of 50 min water sampling and 10 min air sampling, where the first 3 min after switching until stabilization of the signal were discarded.

Before and after the cruise the analyzer was calibrated over a range of concentrations using permeation devices. Both calibrations were consistent. However, during calibration the output of the internal spectral retrieval differed significantly from post-processing of the recorded spectra, which matched the known concentrations (this offset is not present in the commercial instruments). The calibration data were thus used to derive a correction function. After correction all data stayed within 5 % of the standards. The calibration scale of the permeation devices was 5 % below the NOAA scale. As

the OCS analyzer measured CO_2 simultaneously, and CO_2 standards were available during the cruise, drift of the instrument was tested by measuring CO_2 standard gases before and after the cruise and found to be less than 1% of the signal. Special care was taken to avoid contamination, and all materials used were tested for contamination before use.

During OASIS, the mirrors inside the cavity of the OCS analyzer were not completely clean, which led to a reduced signal. To correct the data, an attenuation factor was determined from simultaneous CO₂ measurements, because no OCS standard was available onboard, and OASIS data were corrected accordingly.

An independent quality check of the data was performed by comparing volume mixing ratios of the MBL from the OCS analyzer with samples from air canisters sampled during both cruises and measured independently (Schauffler et al., 1998; de Gouw et al., 2009). The calibrated (and attenuation corrected for OASIS) OA-ICOS data were on average 5 % lower than the air canister samples, which reflects the 5 % difference between the calibration at Forschungszentrum Jülich and the NOAA scale.

During ASTRA-OMZ, CS₂ was directly measured onboard within 1h of collection using a purge and trap system attached to a gas chromatograph and mass spectrometer (GC/MS; Agilent 7890A/Agilent 5975C; inert XL MSD with triple axis detector) running in single-ion mode. The discrete surface seawater samples (50 mL) were taken each hour to every 3 h from the same pump system as for continuous OCS measurements. CS₂ was stripped by purging with helium (70 mL min⁻¹) for 15 min. The gas stream was dried using a Nafion membrane dryer (Perma Pure) and CS2 was preconcentrated in a trap cooled with liquid nitrogen. After heating the trap with hot water, CS₂ was injected into the GC/MS. Retention time for CS_2 (m/z 76, 78) was 4.9 min. The analyzed data were calibrated each day using gravimetrically prepared liquid CS₂ standards in ethylene glycol. During purging, 500 µL gaseous deuterated DMS (d3-DMS) and isoprene (d5-isoprene) were added to each sample as an internal standard to account for possible sensitivity drift between calibrations.

During the TransPEGASO cruise onboard RV Hesperides, surface ocean OCS and CS2 were measured in discrete seawater samples by purge and trap and gas chromatography with mass spectrometry detection (GC-MSD). Samples were collected every day at 09:00 and 15:00 local time in glass bottles without headspace and analyzed within 1 h. Aliquots of 25 mL were withdrawn with a glass syringe and filtered through GF/F during injection into the purge and trap system (Stratum, Teledyne Tekmar). The water was heated to 30 °C and volatiles were stripped by bubbling with 40 mL min⁻¹ of ultrapure helium for 12 min and trapped in a U-shaped VO-CARB 9 trap at room temperature. After flash thermal desorption, volatiles were injected into an Agilent 5975T LTM GC-MSD equipped with an Agilent LTM DB-VRX column $(20 \text{ m} \times 0.18 \text{ mm OD} \times 1 \text{ } \mu\text{m})$ maintained at $30 \,^{\circ}\text{C}$. Retention times for OCS (m/z 60) and CS₂ (m/z 76) were 1.3 and 2.7 min, respectively. Peak quantification was achieved with respect to gaseous (OCS in N2) and liquid (CS2 in methanol and water) standards that were analyzed in the same way. Samples were run in duplicates. Detection limits were 1.8 pM (OCS) and 1.4 pM (CS₂), and precision was typically around 5%.

The systems are each calibrated against a standard, but they had not been directly intercompared. Still, our measurements are consistent with previous measurements using independent methods as discussed in Sects. 3.2.1 and 3.3.

2.3 Calculation of air-sea exchange

Fluxes F of all gases were calculated with Eq. (1):

$$F = k_{\mathbf{w}} \cdot \Delta C,\tag{1}$$

where $k_{\rm w}$ is the gas transfer velocity in water (i.e., physical constraints on exchange) and ΔC the air-sea concentration gradient (i.e., the chemical constraint on exchange). The airside transfer velocity (Liss and Slater, 1974) for OCS was calculated to be 7 orders of magnitude smaller and was therefore neglected. The concentration gradient was determined using the temperature dependent Henry constant (De Bruyn et al., 1995) and the measurements in the surface water and MBL for OASIS and ASTRA-OMZ. During TransPEGASO, no atmospheric volume mixing ratio was measured, and a value of 500 ppt was assumed (Montzka et al., 2007). As air volume mixing ratios of OCS vary over the course of a year, we performed a sensitivity test for a scenario of 450 and 550 ppt and found mean deviations of +7.8 and -7.8 %, respectively. The transfer velocity $k_{\rm w}$ was determined using a quadratic parameterization based on wind speed (Nightingale et al., 2000) which was directly measured onboard (10 min averages). Furthermore, k_w was corrected for OCS and CS₂ by scaling it with the Schmidt number calculated from the molar volume of the gases (Hayduk and Laudie, 1974). It should be noted that the choice of the parameterization for $k_{\rm w}$ has a non-negligible influence on the global emission estimate. Linear, quadratic and cubic parameterizations of k_w are available, with differences increasing at high wind speeds on the order of a factor of 2 (Lennartz et al., 2015; Wanninkhof et al., 2009). Evidence suggests that the air-sea exchange of insoluble gases such as CO2, OCS and CS2 follows a cubic relationship to wind speed because of bubble-mediated gas transfer (McGillis et al., 2001; Asher and Wanninkhof, 1998). However, this difference between soluble and nonsoluble gases is not always consistent (Miller et al., 2009), and too few data are available for a reliable parameterization at high wind speeds above $12 \,\mathrm{m \, s^{-1}}$, where the cubic and the quadratic parameterizations diverge the most. For reasons of consistency, e.g., for the fitted photoproduction p from previous studies, and the fact that most of the previous emission estimates were computed using a quadratic kw parameterization, we chose the same quadratic parameterization representing the mean range of observations (Nightingale et al., 2000). For a sensitivity test, we computed the global oceanic emission with a cubic relationship (McGillis et al., 2001), which results in an additional 40 Gg S per year as direct OCS emissions, leaving the missing source still unexplained. However, better constraints on the transfer velocity of insoluble gases would decrease the uncertainty in global oceanic emissions of marine trace gases.

2.4 Box model of OCS concentration in the surface ocean

A box model to simulate surface concentration of OCS is further developed from the latest version from von Hobe et al. (2003, termed vH2003), where concentrations along the tracks of five Atlantic cruises have been simulated and compared. The vH2003 model results from successful tests and validation to observations on several cruises to the Atlantic Ocean covering all seasons (i.e., Flöck and Andreae, 1996, in January 1994; Uher and Andreae, 1997a, in April/May 1992; von Hobe et al., 1999, in June/July 1997; Kettle et al., 2001, in September/October 1998). By comparing photoproduction rate constants of the five cruises to CDOM absorption, von Hobe et al. (2003) suggest a second-order process for photoproduction with the photoproduction rate constant being dependent on the absorption of CDOM in seawater.

In our approach, we test vH2003 along the cruise track of two cruises, include a new way of determining the photoproduction rate constant (see below) and apply it with global climatological input (termed L2016). Kettle (2000, 2002, termed K2000) applied a similar version of vH2003 globally, which included an optimized photoproduction constant from Atlantic transect cruise data, an optimized constant light-independent production and a linear regression to obtain CDOM from chlorophyll a. In comparison to K2000, we use (i) a new way of determining the photoproduction rate constant incorporating information from three ocean basins, (ii) the most recent parameterization of light-independent production available, and (iii) satellite observations for sea surface CDOM instead of an empirical relationship based on chlorophyll a.

Launois et al. (2015a) implemented parameterizations for light-independent production, hydrolysis and air—sea exchange similar to vH2003 in the 3-D global ocean model NEMO-PISCES. The main differences to the approach used here are the lack of accounting for mixing in L2016 (discussed in Sect. 3.2.2, which will theoretically lead to higher simulated concentrations in our case) and the application of a photoproduction rate constant in our model that incorporates information from three open ocean basins in contrast to one from a study in the North Sea (Launois et al., 2015a).

In L2016, the light-independent production term of OCS was parameterized depending on SST (K) and the absorption coefficient of CDOM at 350 nm wavelength, a_{350} (von Hobe et al., 2001) (Eq. 2).

$$\frac{dC_{OCS}}{dt} = a_{350} \times 10^{-6} \times \exp\left(55.8 - \frac{16200}{SST}\right)$$
 (2)

An overview on symbols and abbreviations used in equations in the following is provided in the Appendix. The parameterization for hydrolysis describes alkaline and acidic degradation of OCS by Reactions (R1) and (R2):

$$OCS + H_2O \rightarrow H_2S + CO_2, \tag{R1}$$

$$OCS + OH^{-} \rightarrow SH^{-} + CO_{2}. \tag{R2}$$

It was parameterized as a first-order kinetic reaction including the rate constant k_h according to Eqs. (3)–(5):

$$\frac{\mathrm{d}C_{\mathrm{OCS}}}{\mathrm{d}t} = [\mathrm{OCS}] \cdot k_{\mathrm{h}},\tag{3}$$

$$k_{\rm h} = \exp\left(24.3 - \frac{10450}{\rm SST}\right) + \exp\left(22.8 - \frac{6040}{\rm SST}\right) \cdot \frac{K}{a[H^+]},$$
(4)

$$-\log_{10}K = \frac{3046.7}{\text{SST}} + 3.7685 + 0.0035486 \cdot \sqrt{\text{SSS}},\tag{5}$$

where $a[H^+]$ is the proton activity and K the ion product of seawater (Dickinson and Riley, 1979).

Fluxes were calculated with Eq. (1) using the same parameterization for $k_{\rm w}$ as for the emission calculation from measurements described above.

Photoproduction was integrated over the mixed layer depth (MLD), assuming a constant concentration of OCS and CDOM throughout the mixed layer, with the photoproduction rate constant $p \pmod{J^{-1}}$, $a_{350} \pmod{m^{-1}}$ and UV radiation (W m⁻²) (Sikorski and Zika, 1993) (Eq. 6).

$$\frac{\mathrm{d}C_{\mathrm{OCS}}}{\mathrm{d}t} = \int_{-\mathrm{MLD}}^{0} p a_{350} \mathrm{UV} \mathrm{d}z \tag{6}$$

MLD was obtained from CTD (conductivity, temperature, depth) profiles and interpolated between these locations (Figs. S1, S2 in the Supplement). The photochemically active radiation that reaches the ocean surface was approximated by Eq. (7) (Najjar et al., 1995):

$$UV = 2.85 \times 10^{-4} \cdot I \cdot \cos^2 \theta, \tag{7}$$

with global radiation I (W m⁻²) and the zenith angle $\cos \theta$. The attenuated UV light intensity directly below the surface (Sikorski and Zika, 1993) down to the respective depth of the mixed layer was calculated in 1 m steps, taking into account attenuation by CDOM and pure seawater. As a simplification in this global approach, the box model did not resolve the whole wavelength spectrum, but rather used a_{350} and applied a photoproduction rate constant that takes into account the integrated spectrum. A similar approach had been tested and compared to a wavelength spectrum resolving version by von Hobe et al. (2003).

The rate coefficients for hydrolysis, light-independent production and air–sea exchange are all reasonably well constrained and parameterizations have been derived from dedicated laboratory and field experiments (hydrolysis, air–sea exchange) or from nighttime OCS observations in several regions assuming steady state (dark production;

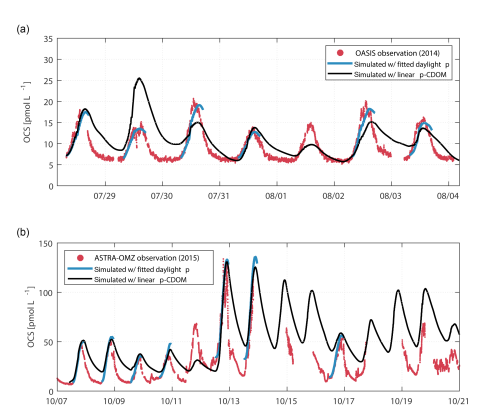


Figure 2. Box model simulations compared to observations: comparison of simulated OCS water concentrations against measurements from the OASIS cruise to the Indian Ocean (a) and the eastern Pacific Ocean during the ASTRA-OMZ cruise (b). Blue indicates OCS concentrations with a least-squares fit for the photoproduction rate constant p during daylight, fitted individually for days with homogeneous water masses (SST, a_{350}). Black shows the simulation including the p depending on a_{350} , obtained from linear regression of individually fitted p with a_{350} (r = 0.71). The time on the x axis is local time (GMT+5 during OASIS 2014, GMT-4 during ASTRA-OMZ 2015).

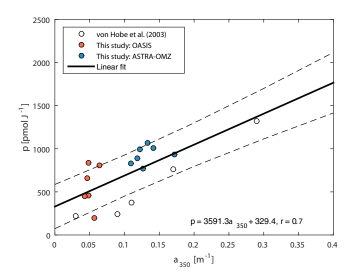


Figure 3. Dependence of photoproduction rate constant p on a_{350} including own fits for p (resulting in blue lines in Fig. 2) and fits from a similar study (von Hobe et al., 2003). Dashed lines indicate the 95 % confidence interval.

von Hobe et al., 2001). However, the photoproduction rate constant p is not well constrained and no generally applicable parameterization exists. In the study of von Hobe et al. (2003), a start was made in parameterizing p in terms of CDOM absorption, and they found this to be dependent on the exact model setup used with respect to wavelength integration and mixed layer treatment. To extend the p-CDOM relationship for other ocean basins, we use the two cruises OASIS and ASTRA-OMZ as case studies for parameter optimization of the photoproduction rate constant p. The photoproduction constant p in the case study simulations was fitted individually for periods of daylight $> 100 \,\mathrm{W m^{-2}}$ (Fig. 2, blue lines) with a Levenberg-Marquardt optimization routine in MatLab version 2015a (8.5.0) by minimizing residuals between simulated and hourly averaged measurements. Different starting values were tested to reduce the risk of the fitted p being a local minimum. Together with photoproduction rate constants obtained by a similar optimization procedure by von Hobe et al. (2003) (Table 2 therein, termed MLB STC), a relationship of the photoproduction constant p dependent on a_{350} was established (Fig. 3). The resulting linear relationship thus includes values from the Atlantic, Pacific and Indian oceans, making it a good approximation for

a globally valid dependence. For the global box model, p was calculated in every time step based on this relationship (r = 0.71, Eq. 8):

$$p = 3591.3 \cdot a_{350} + 329.4 \tag{8}$$

The scatter in Fig. 3 likely reflects the inhomogeneity of the water masses across the three oceanic basins considered, as CDOM absorbance is a valid proxy, but carries some uncertainty in the concentration of the actual precursor.

The model input for simulations of the cruises OASIS and ASTRA-OMZ consisted of measurements made during the respective cruise, including SST and SSS (MicroCAT SBE41) measured every minute, CDOM absorption coefficient (spectrophotometrically measured ca. every 3 h with a liquid capillary cell setup) and the ship's in situ measured meteorological data such as wind speed and global radiation averaged over 10 min (Figs. S1, S2, Tables S1, S2). Forcing data were linearly interpolated to the time step of integration of 2 min.

For the global box model, monthly global meteorological fields with a spatial resolution of $2.8^{\circ} \times 2.8^{\circ}$ were used (Table S3, Fig. S3). For global a_{350} at the sea surface, monthly climatological means for absorption due to gelbstoff and detritus a_{443} (gelbstoff representing CDOM) from the MODIS-Aqua satellite (all available data, 2002–2014) (NASA, 2014) were corrected to 350 nm with Eq. (9) (Fichot and Miller, 2010; Launois et al., 2015a):

$$a_{350} = a_{443} \cdot \exp(-0.02 \cdot (350 - 443)).$$
 (9)

SST, wind speed, and atmospheric pressure were obtained as monthly climatological means from the same period, i.e., 2002 to 2014, by ERA-Interim (Dee et al., 2011). A diel cycle of global radiation I was obtained by fitting the parable parameters a and b during time of the day t in Eq. (10) (Fig. S4),

$$I = -a \cdot t^2 + b,\tag{10}$$

to conditions of (i) x axis interceptions in the distance of the sunshine duration and (ii) the integral being the daily incoming energy by ERA-Interim (Dee et al., 2011). Monthly climatologies of mixed layer depths were used from the MI-MOC project (Schmidtko et al., 2013). For details of data sources please refer to Tables S1–S3 provided in the Supplement. The time step of the model was set to 120 min, which had been tested to result in negligible (< 3%) smoothing.

2.5 Assessing the indirect contribution of DMS with EMAC

Model outputs from ECHAM/MESSy Atmospheric Chemistry (EMAC) from the simulation RC1SDbase-10a of the ESCiMo project (Jöckel et al., 2016) are used to evaluate the contribution of DMS on the production of OCS. The model results were obtained with ECHAM5 version 5.3.02

and MESSy version 2.51, with a T42L90MA resolution (corresponding to a quadratic Gaussian grid of approx. 2.8 by 2.8° in latitude and longitude and 90 vertical hybrid pressure levels up to 0.01 hPa). The dynamics of the general circulation model were nudged by Newtonian relaxation towards ERA-Interim reanalysis data. DMS emissions were calculated with the AIRSEA submodel (Pozzer et al., 2006), which takes into account concentration of DMS in the atmosphere and in the ocean, following a two-layer conceptual model to calculate emissions (Liss and Slater, 1974). While atmospheric concentrations are estimated online by the model (with DMS oxidation), the oceanic concentrations are prescribed as monthly climatologies (Lana et al., 2011). It was shown that such an online calculation of emissions provides the most realistic results when compared to measurements compared to a fixed emission rate (Lennartz et al., 2015). The online-calculated concentrations of DMS and OH were been used to estimate the production of OCS. A production yield of 0.7 % was used for the reaction of DMS with OH (Barnes et al., 1994), using the reaction rate constant suggested by the International Union of Pure and Applied Chemistry (IUPAC) (Atkinson et al., 2004).

3 Results and discussion

3.1 Observations of OCS in the tropical ocean

OCS was measured in the surface ocean and MBL during three cruises in the tropics. Measurement locations (Fig. 1) include oligotrophic open ocean regions in the Indian Ocean (OASIS, 07-08/2014), open ocean and shelf areas in the eastern Pacific (ASTRA-OMZ, 10/2015) and a meridional transect in the Atlantic (TransPEGASO, 10-11/2014). In the Indian and Pacific oceans, continuous underway measurements provided the necessary temporal resolution to observe diel cycles of OCS concentrations in surface water. Dissolved OCS concentrations exhibited diel cycles with maxima 2 to 4h after local noon (Fig. 1), which are a consequence of photochemical production and removal by hydrolysis (Uher and Andreae, 1997a). OCS concentrations also varied spatially. Taking a_{350} as a proxy for CDOM content, we found that daily mean OCS concentrations were higher in CDOM-rich (Table 3, 28.3 ± 19.7 pmol OCS L⁻¹, a_{350} : 0.15 \pm 0.03 m⁻¹) than in CDOM-poor waters (Table 3, OASIS: $9.1\pm3.5 \text{ pmol OCS L}^{-1}$, a_{350} : $0.03\pm0.02 \text{ m}^{-1}$). Samples during TransPEGASO were measured with gas chromatography-mass spectrometry twice a day (around 08:00–10:00 and 15:00–17:00 local times). Therefore, the full diel cycles could not be reconstructed and potential variations of OCS with CDOM absorption were overlaid by diel variations. Nevertheless, the observed range of OCS concentrations in the Atlantic corresponds well to the observations from the eastern Pacific and Indian Ocean (Table 3) and is consistent with measurements from a previous Atlantic

Table 3. Average, standard deviation and range of parameters observed during the cruises OASIS (Indian Ocean, 2014), ASTRA-OMZ (Pacific Ocean, 2015) and TransPEGASO (Atlantic Ocean, 2014).

		Average (±SD)	Minimum	Maximum
OASIS (Indian Ocean)				
OCS sea surface concentration	$(\text{pmol } L^{-1})$	9.1 (±3.5)	5.1	20.7
OCS flux	$(g S d^{-1} km^{-2})$	$-0.25 (\pm 0.5)$	-1.6	1.5
SST	(°C)	$27.0 (\pm 1.4)$	22.2	32.0
Salinity	(-)	$34.9 (\pm 0.3)$	34.3	35.4
Wind speed	$(m s^{-1})$	$7.6 (\pm 2.1)$	0.2	14.5
$a_{\text{CDOM}}(350)$	(m^{-1})	$0.03~(\pm 0.02)$	n.d.	0.12
ASTRA-OMZ (Pacific Ocean)				
OCS sea surface concentration	$(\operatorname{pmol} L^{-1})$	28.3 (±19.7)	6.5	133.8
OCS flux	$(g S d^{-1} km^{-2})$	$1.5 (\pm 2.1)$	-1.5	19.9
CS ₂ sea surface concentration	$(\text{pmol } L^{-1})$	$17.8 (\pm 8.9)$	6.7	40.1
CS ₂ flux	$(g S d^{-1} km^{-2})$	$4.1 (\pm 3.2)$	0.2	14.4
SST	(°C)	$20.1 (\pm 2.9)$	15.6	26.9
Salinity	(-)	$35.0 \ (\pm 0.43)$	33.4	35.5
Wind speed	$(m s^{-1})$	$7.4 (\pm 2.0)$	0.3	15.5
$a_{\text{CDOM}}(350)$	(m^{-1})	$0.15~(\pm 0.03)$	0.1	0.24
TransPEGASO (Atlantic Ocean)				
OCS sea surface concentration	$(\operatorname{pmol} L^{-1})$	23.6 (±19.3)	2.6	78.3
OCS flux	$(g S d^{-1})$	$1.3 (\pm 3.5)$	-1.7	14.0
CS ₂ sea surface concentration	$(\text{pmol } L^{-1})$	$62.5 (\pm 42.1)$	23.2	154.8
CS ₂ flux	$(g S d^{-1} km^{-2})$	$13.7 (\pm 9.8)$	0.3	33.9
SST	(°C)	$22.6 (\pm 6.3)$	7.1	29.6
Salinity	(-)	$34.9 (\pm 2.6)$	28.4	38.1
Wind speed	$(m s^{-1})$	$7.4 (\pm 3.1)$	0.4	19.0
$a_{\text{CDOM}}(350)$	(m^{-1})	$0.13 \ (\pm 0.11)$	0.0023	0.45

meridional transect (AMT-7) cruise (Kettle et al., 2001) (1.3–112.0 pmol OCS L^{-1} , mean 21.7 pmol OCS L^{-1}).

Air—sea fluxes calculated from surface concentrations and mixing ratios of OCS as a function of wind speed generally follow the diel cycle of the surface ocean concentration. While supersaturation prevailed during the day, low night-time concentrations usually led to oceanic uptake of atmospheric OCS. OCS fluxes integrated over one day ranged from -0.024 to $-0.0002\,\mathrm{g\,S\,km^{-2}}$ in the open Indian Ocean and from 0.38 to 2.7 g S km $^{-2}$ in the coastal Pacific. During the observed periods, the ocean was a net sink of atmospheric OCS in the Indian Ocean, whereas it was a net source in the eastern Pacific. Although an assessment of net flux is difficult given the lower temporal resolution during TransPE-GASO, calculated emissions were in the same range as the ones measured in the Pacific and Indian Ocean.

The water masses encountered during the cruises to the Indian Ocean (OASIS) and eastern Pacific (ASTRA-OMZ), which are used to constrain the global box model, differ considerably with respect to the properties relevant for OCS cycling and, thus, span a large range of possible OCS variabil-

ity. The properties encountered during these two cruises encompass or exceed the ones of the Pacific warm pool (climatological averages, Table 4), which is where the location of the missing source has been hypothesized (Glatthor et al., 2015; Kuai et al., 2015). Both higher SST and lower wind speeds (Table 4) would decrease the OCS sea surface concentrations in the ocean, leading to decreased emissions to the atmosphere: higher SSTs favor a stronger degradation by hydrolysis (Elliott et al., 1989), and lower wind speeds decrease the transfer velocity k. Lower integrated daily radiation (SR in Table 4) in the Pacific warm pool also points to lower OCS production. Hence, our new OCS observations presented here likely span the range of emission variability in the tropics.

The observed concentrations and calculated emissions are approximately 1 order of magnitude lower than the annual mean surface concentrations and emissions simulated in the 3-D global ocean model NEMO-PISCES (Launois et al., 2015a).

Table 4. Comparison of water properties relevant for OCS production and consumption for the cruises OASIS (Indian Ocean, July–August 2014) and ASTRA-OMZ (eastern Pacific, October–November 2015) with the assumed source region in the Pacific warm pool (15° N–15° S, 120–180° E). Data from cruises are in situ measurements; the data for the Pacific warm pool were extracted from climatological monthly means from sources for the global model run as specified in the Supplement.

Parameter	OASIS	ASTRA-OMZ	Pacific warm pool
SST (°C)	27.0 ± 1.0	19.6 ± 2.6	28.9 ± 0.9
$SSS (g kg^{-1})$	35.0 ± 0.3	35.1 ± 0.3	34.5 ± 0.42
Wind speed $(m s^{-1})$	8.2 ± 1.7	7.5 ± 1.8	5.3 ± 0.4
$a_{350} (\mathrm{m}^{-1})$	0.039 ± 0.02	0.146 ± 0.02	0.050 ± 0.08
$I ({\rm W} {\rm m}^{-2})$	226.5 ± 303.0	196.4 ± 283.1	206.4 ± 286.6^{a}
$SR (J m^{-2})$	$1.9 \times 10^7 \pm 1.7 \times 10^6$	$1.6 \times 10^7 \pm 4.5 \times 10^6$	$8.9 \times 10^6 \pm 1.3 \times 10^6$
pH (-)	8.03 ± 0.01	_b	8.07 ± 0.01
MLD (m)	43.3 ± 15.8	18.9 ± 7.5	35.9 ± 14.1

 $^{^{}a}$ Calculated from an annual mean diurnal cycle based on ERA-Interim sunshine duration and flux. SR: surface radiation, daily integral. b Assumed pH = 8.15 for box model simulation.

3.2 A direct global oceanic emission estimate for OCS

The OCS observations from the Indian and Pacific Ocean were used to improve a box model for simulating OCS concentrations in the surface ocean (Kettle, 2002; Uher and Andreae, 1997b; von Hobe et al., 2003). With the a_{350} dependent photoproduction constant included, the model reproduced the diel pattern of OCS concentrations in the surface oceans for both cruises (Fig. 2, black lines). A slight overestimation of observed concentrations is present for the Indian Ocean cruise OASIS (observed mean concentration: $9.1 \pm 3.5 \, \text{pmol L}^{-1}$; simulated: $10.8 \pm 3.9 \, \text{pmol L}^{-1}$). This overestimation was more pronounced in the eastern Pacific (observed mean: $28.3 \pm 19.7 \,\mathrm{pmol}\,\mathrm{L}^{-1}$; simulated: $47.3 \pm 25.4 \,\mathrm{pmol}\,\mathrm{L}^{-1}$) and can largely be attributed to a lack of downward mixing inherent in the mixed layer box model due to the assumption of the OCS concentration being constant throughout the entire mixed layer.

Using the linear $p-a_{350}$ parameterization for the first time in a global model, the same box model as for the case studies is applied to estimate sea surface concentrations and fluxes of OCS on a global scale (Fig. 4). The OCS production is consistent with the global distribution of CDOM absorption (Fig. S5), with highest concentrations calculated for coastal regions and higher latitudes. Despite the photochemical hot spot in the tropics (30° N–30° S), degradation by hydrolysis prevents any accumulation of OCS in the surface water, as we calculated the lifetime due to hydrolysis to be only 7 h (Fig. S5). The simulated range of water concentrations is too low to sustain emissions in the tropics that could close the atmospheric budget of OCS (Fig. 4). With saturation ratios integrated over 1 year, the tropical ocean (30° N-30° S) is even undersaturated with respect to OCS, taking up 3.0 Gg S yr⁻¹. Globally, the integration over one year yields annual oceanic OCS emissions of 130 Gg S. Our results corroborate the upper limit of an earlier study that used an observation-derived emission inventory (Table 1) (Kettle, 2002) but which in-

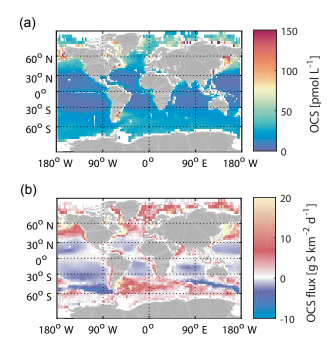


Figure 4. Annual mean of surface ocean concentrations of OCS simulated with the box model (a) and corresponding emissions (b).

cludes more process-oriented parameterizations as described in Sect. 2.4. Clearly, our results from both observations and modeling contradict the latest bottom-up emission estimate from the NEMO-PISCES model (Launois et al., 2015a), and do not support a hot spot of direct OCS emissions in the Pacific Warm Pool or the tropical oceans in general.

3.2.1 Comparison to previous ship-based measurements

The global simulation of OCS surface water concentrations generally reproduced the lower picomolar range of concen-

Table 5. Comparison of previous ship campaign measurements with corresponding month and approximate geolocation from the global box model in this study (L2016), taken either from figures or tables as provided in the original references. Note that the box model is based on input data from climatological means that do not fully represent the conditions encountered during the respective cruises. Only observational data with measurements of the full diel cycle were included for comparison. *n*: number of measurements.

References	Season	Region	$\begin{array}{c} \text{Mean OCS} \pm \text{SD} \\ \text{(pmol L}^{-1}) \end{array}$	n	$L2016$ mean $(pmol L^{-1})$
Mihalopoulos et al. (1992)		open Indian Ocean 20° N–37° S			
	Mar/May 1986	OCEAT II	19.9 ± 0.5^{a}	20	11.2 ± 6.3
	Jul 1987	OCEAT III	19.9 ± 1.0^{a}	14	17.7 ± 13.1
Staubes and Georgii (1993)	Nov-Dec 1990	Weddell Sea 40–72° S,72° W–24° E	109 ^b	126	66.6 ± 49.8
Ulshöfer et al. (1995)		North Atlantic Ocean			
	Apr/May 1992	47° N 20° W	14.9 ± 6.9	118	42.8 ± 11.3
	Jan 1994	48–50° N, 10–17° W	5.3 ± 1.6	120	8.9 ± 3.2
	Sep 1994	48–50° N, 10–17° W	19.0 ± 8.3	235	33.4 ± 3.5
Flöck and Andreae (1996)	Jan 1994	northeastern Atlantic Ocean 49° N, 12° W	6.7 (4–11)	120	9.6 ± 3.7
Ulshöfer and Andreae (1998)	Mar 1995	western Atlantic 32° N, 64° W	8.1 ± 7.0	323	15.8
von Hobe et al. (1999)	Jun/Jul 1997	northeastern Atlantic Ocean 30–40° N, 8–15° W	23.6 ± 16.0	940	30.5 ± 12.6
Kettle et al. (2001)	Sep/Oct 1998	Atlantic transect 50° N–60° S, 1–64° W	21.7 ± 19.1	783	22.9 ± 3.2
von Hobe et al. (2001)	Aug 1999	Sargasso Sea/BATS 32° N, 64° W	8.6 ± 2.8	518	8.1
Xu et al. (2001)	Oct/Nov 1997	Atlantic meridional transect 53° N–34° S, 25° W–20° E	14.8 ± 11.4	306	11.8 ± 12.7
	May/Jun 1998	Atlantic meridional transect 53° N–34° S, 25° W–20° E	18.1 ± 16.1	440	27.8 ± 47.9

^a Converted from $ng L^{-1}$ with a molar mass of OCS of 60.07 g. ^b Converted from $ng S L^{-1}$ with a molar mass of S of 32.1 g.

trations (Table 5), the seasonal pattern of higher concentrations during summer compared to winter (as, for example, in Ulshöfer et al., 1995) and the spatial pattern of higher concentrations in higher latitudes (e.g., Southern Ocean; Staubes and Georgii, 1993). Given that monthly means of a model simulation driven by climatological data of the input parameters are compared to cruise measurements, the absolute mean deviation of $6.9 \,\mathrm{pmol}\,\mathrm{L}^{-1}$ and the mean deviation of $3.7 \,\mathrm{pmol}\,\mathrm{L}^{-1}$ indicate an overall good reproduction of observations (differences between observation and model output were weighted to the number of observations in Table 4). It should be noted that, on average, the model overestimates OCS concentrations as indicated by the positive mean error, suggesting our emission estimate to be an upper limit to direct oceanic OCS emissions in most regions. The largest deviations from observations are found in the Southern Ocean (see Staubes and Georgii, 1993, in Table 4), where the model underestimated observations by 40 %. While there are several

explanations for this, i.e., a possible violation of the underlying assumption of a constant OCS production in regions with deep mixed layers such as the Southern Ocean, or the missing satellite data for CDOM during polar nights, it is a clear indication of the need for more observations from high latitudes. However, this underestimation does not interfere with our conclusion drawn for the tropical oceans, where the location of the missing source is derived from top-down approaches.

3.2.2 Uncertainties

Simulated concentrations and fluxes carry uncertainties from input parameters and process parameterizations. One major uncertainty associated with the mixed layer box model approach arises from the fact that it does not adequately account for downward mixing and vertical concentration gradients within the mixed layer. Under most circumstances,

and especially in the tropical open ocean, where hydrolysis greatly exceeds surface outgassing and low a_{350} makes photoproduction extend further down in the water column, the model tends to overestimate the real OCS concentrations, as was shown for our two cruises above. Therefore, we deem the fluxes from our global simulation to represent an upper limit of the true fluxes. Only at high latitudes would we expect more complex uncertainties, because hydrolysis at low temperatures is slow and only photoproduction and loss by outgassing are directly competing at the very surface.

Other uncertainties are associated with the calculation of the photoproduction rate. The wavelength of 443 nm combines the absorption of detritus and CDOM, which could have an impact especially in river plumes, where terrestrial material is transported into the ocean. As it is the CDOM that is important for photochemistry, assuming the 443 nm is purely CDOM would lead to an overestimation of photoproduction and therefore is a conservative estimate. It should also be noted that a single spectral slope from 443 to 350 nm in the global simulation is a simplification. Furthermore, using a wavelength integrated photoproduction rate constant instead of a wavelength-resolved approach, which would take global variations in the CDOM and light spectra into account, is an additional simplification. It has been shown that this does not lead to large differences regionally (von Hobe et al., 2003) but could, potentially, lead to variations globally. Our p-CDOM relationship is a first step for constraining this variability globally in one parameterization, as it incorporates photoproduction rate constants optimized to observations and thus accounting for differences in the light and CDOM spectra. More data from different regions can help to further constrain this relationship in future studies. Despite these simplifications, the simulated concentrations agree very well with previous observations (n > 4000, Table 4). To test the sensitivity of our box model to the photoproduction rate constant, we performed a sensitivity test with a photoproduction increased by a factor of 5 in the tropical region (30° N–30° S; note that this factor is considerably larger than the uncertainty in the p-CDOM relationship). This leads to an annual mean concentration of 35.1 pmol L^{-1} in the tropics (30° N–30° S), resulting in tropical direct emissions of 160 Gg S as OCS per year. The efficient hydrolysis in warm tropical waters prevents OCS concentrations from accumulating despite the high photoproduction and still results in emissions too low to account for the missing source.

With a mean error of $3.7\,\mathrm{pmol\,L^{-1}}$ in the OCS surface water concentrations added to (subtracted from) the modeled concentration and subsequent calculation of fluxes using annual climatologies for wind, pressure and SST (same data sources as global simulation forcing data), we calculate an uncertainty of $60\,\%$, which translates into a total uncertainty in the integrated global flux of $80\,\mathrm{Gg}\,\mathrm{S}\,\mathrm{yr}^{-1}$.

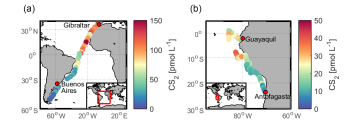


Figure 5. Measured concentration of CS_2 in surface waters during (a) ASTRA-OMZ in the eastern Pacific Ocean and (b) TransPE-GASO in the Atlantic Ocean.

3.3 Indirect OCS emissions by DMS and CS₂

A significant contribution to the OCS budget in the atmosphere results from oceanic emissions of DMS and CS2 that are partially converted to OCS on timescales of hours to days (Chin and Davis, 1993; Watts, 2000; Kettle, 2002). A yield of 0.7 % for OCS is used for the reaction of DMS with OH (Barnes et al., 1994), which results in a global oceanic source of DMS from OCS of $80(65-110) \,\mathrm{Gg} \,\mathrm{S} \,\mathrm{yr}^{-1}$ based on the procedure described in Sect. 2.5. The uncertainty range of 65–110 Gg S yr⁻¹ originated from the uncertainty in oceanic emissions, not the conversion factor. This conversion factor is much more uncertain, as the formation of OCS from DMS involves a complex multi-step reaction mechanism that is not fully understood. It has been shown in laboratory experiments that the presence of NO_x reduces the OCS yield considerably (Arsene et al., 2001), which would make our indirect emission estimate an upper limit. However, the yield was measured under laboratory conditions and may be different and more variable under natural conditions.

DMS emissions do not show a pronounced hot spot in the Pacific warm pool region, but as DMS transports much more sulfur across the air—sea interface than OCS, even low changes in the OCS yield could affect the atmospheric budget of OCS. As the spatial oceanic emission pattern of DMS does not reflect the spatial pattern of the assumed missing source, a locally specific tropospheric change in the conversion yield would be one potential way of bringing the patterns in agreement. While it is possible that the OCS yield could vary under certain conditions (e.g., it cannot be excluded that the low OH concentrations in the broader Pacific warm pool area as suggested by Rex et al., 2014, influence the yield), the (local) increase in the conversion factor would need to be on the order of a factor of 10–100.

For CS₂, the atmospheric reaction pathway producing OCS is better understood with a well-constrained molar conversion ratio of 0.81 (Chin and Davis, 1993). However, the global distribution of oceanic CS₂ concentration, and hence its emissions to the atmosphere, is poorly known. In our study, surface CS₂ concentrations (Fig. S6) were on average 17.8 \pm 8.9 pmol L⁻¹ during ASTRA-OMZ, and 62.5 \pm 42.1 pmol L⁻¹ during TransPEGASO (Table 3). The

latter values are higher than previously reported concentrations from the AMT-7 cruise in the central Atlantic (Kettle et al., 2001) $(10.9 \pm 15.2 \, \text{pmol L}^{-1})$. We extrapolate a weighted mean of the CS2 emissions from TransPEGASO $(n = 42, 13.7 \pm 9.8 \,\mathrm{g}\,\mathrm{S}\,\mathrm{d}^{-1}\,\mathrm{km}^{-2})$, ASTRA-OMZ $(n = 122, 13.7 \pm 9.8 \,\mathrm{g}\,\mathrm{S}\,\mathrm{d}^{-1}\,\mathrm{km}^{-2})$ $4.1 \pm 3.2 \,\mathrm{g}\,\mathrm{S}\,\mathrm{d}^{-1}\,\mathrm{km}^{-2}$) and AMT-7 (Kettle et al., 2001) $(n = 744, 1.6 \pm 1.8 \text{ g S d}^{-1} \text{ km}^{-2})$ in order to estimate CS₂derived OCS emissions from the global ocean. According to our extrapolation, 135 (7–260) Gg S yr⁻¹ enters the atmosphere as oceanic CS2 emissions converted to OCS. The uncertainty range of 7-260 Gg S yr⁻¹ results from extrapolating the highest and the lowest emissions encountered during the cruises to the global ocean. This number is at the highest end of the range for OCS emissions from globally simulated CS₂ oceanic concentrations (Kettle, 2000, 2002), as measured CS₂ concentrations from the cruises ASTRA-OMZ and TransPEGASO are higher than the simulated surface concentrations in Kettle (2000) for the respective month. However, the spatial pattern of higher concentrations and emissions in the tropical region in our measurements agrees well with the spatial pattern simulated in Kettle (2000). Nonetheless, even the extrapolation of the highest measurement would not close the budget for the three largest missing source estimates (Table 1).

For oceanic emission estimates used to constrain GPP, quantifying the seasonal cycle of the single contributors is essential. For example, high emissions during oceanic spring and fall blooms could mask OCS uptake by the terrestrial vegetation, and therefore neglecting them could lead to an underestimation of global GPP, with implications for the atmospheric and terrestrial carbon budget.

4 Conclusions and outlook

Considering the observational evidence and the modeled global emission estimate of $130\pm80\,\mathrm{Gg}\,\mathrm{S}\,\mathrm{yr}^{-1}$, direct OCS emissions from the oceans are too low to account for the missing atmospheric source. Together with indirect emissions, the oceanic source strength of OCS would add up to 345 Gg S yr⁻¹, compared to the 465–1089 Gg S yr⁻¹ required to balance the suggested increase in vegetation uptake. Direct and even additional indirect oceanic emissions of OCS are thus unlikely to balance the budget after the upward revision of the vegetation sink. Largest uncertainties are associated with the indirect emission estimates, especially in the conversion of DMS to OCS and the global source strength of CS₂.

As our study suggests, the search for an additional source of OCS to the atmosphere should include other sources than oceanic emissions alone. There are indications of other parts of the OCS budget being underestimated, such as domestic coal combustion (Du et al., 2016). Emissions of biomass burning and direct and indirect anthropogenic emissions have been considered in previous estimates (e.g., 315.5 Gg S yr⁻¹

in Berry et al., 2013, 224 Gg S yr⁻¹ in Kuai et al., 2015, and 219 Gg S yr⁻¹ in Glatthor et al., 2015), but a recent anthropogenic emission estimate by Lee and Brimblecombe (2016) increases this number to 598 Gg S yr⁻¹, which would already bring sources and sinks closer to agreement. They attribute the largest direct OCS emissions to biomass and biofuel burning, as well as pulp and paper manufacturing, and the largest CS₂ emissions to the rayon industry. Hence, a hot spot of anthropogenic emissions in the Asian continent might be a potential candidate, together with atmospheric transport, to produce atmospheric mixing ratios as observed by satellite.

A redistribution of the magnitude and seasonality of known sources and sinks could also bring top-down and bottom-up estimates closer together. For example, the general view of oxic soils as a sink for OCS has recently been challenged. Field (Maseyk et al., 2014; Billesbach et al., 2014) and incubation studies (Whelan et al., 2016) show that some oxic soils may shift from OCS uptake to emission depending on the temperature and water content. Furthermore, it has been speculated previously that vegetation uptake might not be solely responsible for the decrease in OCS mixing ratios in fall because of the temporal lag between CO₂ and OCS minimum (Montzka et al., 2007). The observed seasonality in mixing ratios is a superposition of the seasonality of all individual sources and sinks. These seasonalities are currently neglected or associated with a considerable uncertainty. An improved understanding of the seasonality of the individual sources and sinks could help to better constrain the gap in the atmospheric budget. First steps to resolve OCS seasonality in sources and sinks are currently being undertaken, e.g., in the case of anthropogenic emissions (Campbell et al., 2015).

All in all, better constraints on the seasonality and magnitude of the atmospheric OCS sources and sinks are critical for a better assessment of the role of this compound in climate and its application to quantify GPP on a global scale. This study confirms oceanic emission as the largest known single source of atmospheric OCS but shows that its magnitude is unlikely to balance the gap in the atmospheric OCS budget.

5 Data availability

All data, including OCS and CS₂ measurements in sea water and the marine boundary layer, as well as OCS model output, are available upon request from the authors (correspondence to S. T. Lennartz, slennartz@geomar.de or C.A. Marandino, cmarandino@geomar.de).

Appendix A: List of parameters

Symbol/abbreviation	Meaning
a ₃₅₀	absorption coefficient of CDOM at 350 nm
a	fitted parameter in diurnal cycle of I
b	fitted parameter in diurnal cycle of I
$c_{ m air}$	concentration in air
$C_{\rm OCS}$	concentration of OCS in water
F	gas flux
H	Henry constant
I	downwelling solar radiation
K	ion product of seawater
$k_{ m w}$	water-side transfer velocity in air-sea gas exchange
MLD	mixed layer depth
p	photoproduction rate constant
SSS	sea surface salinity
SST	sea surface temperature
Sc	Schmidt number
t	time
θ	zenith angle
u_{10}	wind speed at 10 m height
UV	ultraviolet radiation
z	depth

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