# Halocarbon emissions and sources in the equatorial Atlantic Cold Tongue

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#### 1 Abstract

2 Halocarbons from oceanic sources contribute to halogens in the troposphere, and can be transported into the stratosphere where they take part in ozone depletion. This paper presents 3 4 distribution and sources in the equatorial Atlantic from June and July 2011 of the four 5 compounds bromoform (CHBr<sub>3</sub>), dibromomethane (CH<sub>2</sub>Br<sub>2</sub>), methyl iodide (CH<sub>3</sub>I) and 6 diiodomethane (CH<sub>2</sub>I<sub>2</sub>). Enhanced biological production during the Atlantic Cold Tongue 7 (ACT) season, indicated by phytoplankton pigment concentrations, led to elevated concentrations of CHBr<sub>3</sub> of up to 44.7 pmol  $L^{-1}$  and up to 9.2 pmol  $L^{-1}$  for CH<sub>2</sub>Br<sub>2</sub> in surface 8 water, which is comparable to other tropical upwelling systems. While both compounds 9 correlated very well with each other in the surface water, CH<sub>2</sub>Br<sub>2</sub> was often more elevated in 10 greater depth than CHBr<sub>3</sub>, which showed maxima in the vicinity of the deep chlorophyll 11 12 maximum. The deeper maximum of CH<sub>2</sub>Br<sub>2</sub> indicates an additional source in comparison to CHBr<sub>3</sub> or a slower degradation of CH<sub>2</sub>Br<sub>2</sub>. Concentrations of CH<sub>3</sub>I of up to 12.8 pmol  $L^{-1}$  in 13 14 the surface water were measured. In contrary to expectations of a predominantly 15 photochemical source in the tropical ocean, its distribution was mostly in agreement with biological parameters, indicating a biological source. CH<sub>2</sub>I<sub>2</sub> was very low in the near surface 16 water with maximum concentrations of only 3.7 pmol L<sup>-1</sup>. CH<sub>2</sub>I<sub>2</sub> showed distinct maxima in 17 deeper waters similar to CH<sub>2</sub>Br<sub>2</sub>. For the first time, diapycnal fluxes of the four halocarbons 18 19 from the upper thermocline into and out of the mixed layer were determined. These fluxes were low in comparison to the halocarbon sea-to-air fluxes. This indicates that despite the 20 21 observed maximum concentrations at depth, production in the surface mixed layer is the main 22 oceanic source for all four compounds and one of the main driving factors of their emissions 23 into the atmosphere in the ACT-region. The calculated production rates of the compounds in the mixed layer are  $34 \pm 65$  pmol m<sup>-3</sup> h<sup>-1</sup> for CHBr<sub>3</sub>,  $10 \pm 12$  pmol m<sup>-3</sup> h<sup>-1</sup> for CH<sub>2</sub>Br<sub>2</sub>,  $21 \pm 24$ 24 pmol m<sup>-3</sup> h<sup>-1</sup> for CH<sub>3</sub>I and 384  $\pm$  318 pmol m<sup>-3</sup> h<sup>-1</sup> for CH<sub>2</sub>I<sub>2</sub> determined from 13 depth 25 26 profiles.

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#### 28 **1** Introduction

Oceanic upwelling regions where cold nutrient rich water is brought to the surface are connected to enhanced primary production and elevated halocarbon production, especially of bromoform (CHBr<sub>3</sub>) and dibromomethane (CH<sub>2</sub>Br<sub>2</sub>) (Quack et al., 2007a; Carpenter et al., 2009; Raimund et al., 2011; Hepach et al., 2014). Photochemical formation (Moore and Zafiriou, 1994; Richter and Wallace, 2004) with a possible involvement of organic precursors is an important source for methyl iodide (CH<sub>3</sub>I). An abiotic formation pathway for

1 halocarbons involving ozone has been found for diiodomethane  $(CH_2I_2)$  in the laboratory 2 (Martino et al., 2009). But, its production is generally suggested to be biotic, occurring likely 3 through different species of phytoplankton than are involved in the production of CHBr<sub>3</sub> and 4 CH<sub>2</sub>Br<sub>2</sub> (Moore et al., 1996; Orlikowska and Schulz-Bull, 2009). Additionally, bacterial 5 involvement in the formation of halocarbons e.g. CH<sub>3</sub>I and CH<sub>2</sub>I<sub>2</sub> has been observed in the 6 field and the laboratory (Manley and Dastoor, 1988; Amachi et al., 2001; Fuse et al., 2003; 7 Amachi, 2008). Large uncertainties regarding the production of halocarbons in the ocean 8 remain. Depth profiles of the different compounds provide insight into the processes 9 participating in their cycling. Elevated concentrations of CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub> at the bottom of 10 the mixed layer and below, often close to the chlorophyll a (Chl a) subsurface maximum, are 11 a common feature in the water column (Yamamoto et al., 2001; Quack et al., 2004; Liu et al., 12 2013a), and are attributed to enhanced production by phytoplankton. While occasionally  $CH_3I$ 13 maxima close to the Chl a maximum were observed as well (Moore and Groszko, 1999; 14 Wang et al., 2009), Happell and Wallace (1996) ascribed surface maxima in several oceanic 15 regions including the equatorial Atlantic to a predominantly photochemical source. Rapid 16 photolysis and biogenic sources in the deep Chl a maximum are suggested to determine the 17 depth distribution of CH<sub>2</sub>I<sub>2</sub> concentrations (Moore and Tokarczyk, 1993; Yamamoto et al., 18 2001; Carpenter et al., 2007; Kurihara et al., 2010). The complex interactions between the 19 sources (biogenic and non-biogenic production), sinks (hydrolysis, photolysis, chlorine substitution and air-sea gas exchange), advection, and turbulent mixing in and out of the 20 21 mixed layer (diapycnal fluxes), which determine the water concentrations of these 22 compounds, are still sparsely investigated.

23 Once they are produced in the ocean, halocarbons can be transported from the oceanic mixed 24 layer into the troposphere via air-sea gas transfer. CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub> are the largest 25 contributors to atmospheric organic bromine from the ocean (Penkett et al., 1985; Schauffler 26 et al., 1998; Hossaini et al., 2012). Marine CH<sub>3</sub>I is the most abundant organoiodine in the 27 troposphere, while the very short lived CH<sub>2</sub>I<sub>2</sub> and CH<sub>2</sub>CII contribute potentially as much 28 organic iodine (Saiz-Lopez et al., 2012). Significant amounts of halocarbons and their 29 degradation products can be carried into the stratosphere (Solomon et al., 1994; Hossaini et 30 al., 2010; Aschmann et al., 2011), especially in the tropical regions where surface air can be 31 transported very rapidly into the tropical tropopause layer by tropical deep convection 32 (Tegtmeier et al., 2012; Tegtmeier et al., 2013). The short-lived brominated and iodinated 33 halocarbons produced in the equatorial region may hence play an important role for 34 stratospheric halogens.

This paper characterizes the distribution of CHBr<sub>3</sub>, CH<sub>2</sub>Br<sub>2</sub>, CH<sub>3</sub>I, and CH<sub>2</sub>I<sub>2</sub> in the surface 1 2 water and the water column of the equatorial Atlantic Cold Tongue (ACT) for the first time. 3 The ACT is a known feature in the equatorial region, which is characterized by intensive 4 cooling of SSTs. This cooling is also associated with phytoplankton blooms (Grodsky et al., 5 2008) as potential source for halocarbons. CHBr<sub>3</sub>, CH<sub>2</sub>Br<sub>2</sub>, CH<sub>3</sub>I and CH<sub>2</sub>I<sub>2</sub> represent the most 6 important carriers of organic halogens into the troposphere, which have important 7 implications for atmospheric chemistry and are poorly characterized in the ACT region. We 8 therefore aim to provide more insight into the biological and physical processes contributing 9 to the mixed layer budget of halocarbons in the equatorial Atlantic. Sea-to-air fluxes and, for 10 the first time, diapycnal fluxes from the upper thermocline are calculated as sources and sinks 11 for the mixed layer. Phytoplankton groups (obtained from pigment concentrations) are 12 evaluated as potential sources of these four compounds. Additionally, surface water 13 halocarbons are correlated to meta data such as temperature, salinity and global radiation to 14 understand their distribution further. Finally, we estimate production rates for the mixed layer 15 of the ACT region.

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#### 17 2 Methods

18 Cruise MSM18/3 onboard the RV Maria S. Merian took place from June 21 to July 21 2011. 19 One goal of the campaign was the characterization of the Atlantic equatorial upwelling with 20 regard to halocarbon emissions and their sources. RV Maria S. Merian started in Mindelo 21 (Sao Vicente, Cape Verde) at 16.9° N and 25.0° W, and finished in Libreville (Gabon) at 22 0.4° N and 13.4° E with several transects across the equator. The ship entered the ACT 23 several times. Measurements of halocarbons and phytoplankton pigments were conducted in 24 surface water along the cruise track, and at 13 stations (Figure 1). Samples for dissolved 25 halocarbons from sea surface water were taken from a continuously working pump in the 26 ships moon pool at a depth of about 6.5 m every 3 h. Deep water samples were taken from up 27 to eight different depths per station between 10 and 700 m from 12 L Niskin bottles attached 28 to a 24-bottle-rosette with a CTD (Conductivity Temperature Depth). Halocarbon stations 1 – 4 were located at the first meridional transect across the ACT at  $15^{\circ}$  W, stations 5 – 7 at the 29 second transect at  $10^{\circ}$  W, 8 - 10 were located at the third section at around  $5^{\circ}$  W, and the last 30 31 three stations 11 - 13 were taken during the last section at  $0^{\circ}$  E (Figure 1). Water temperature 32 and salinity were recorded with a thermosalinograph. Air pressure and wind speed were 33 derived from sensors in 30 m height, averaged in 10 min intervals, and wind speed was 34 corrected to 10 m. Global radiation was measured onboard in 19.5 m height with sensors (SMS-1 combined system from MesSen Nord, Germany) measuring downward incoming
 global radiation (GS, shortwave) and infrared radiation (IR, long-wave).

#### 3 2.1 Sampling and analysis of halocarbons in seawater

A purge and trap system attached to a gas chromatograph with mass spectrometric detection (GC-MS) in single ion mode was used to analyze 50 mL water samples for dissolved halocarbons. Volumetrically prepared standards in methanol were used for quantification. Precision lay within 3 % for CHBr<sub>3</sub>, 6 % for CH<sub>2</sub>Br<sub>2</sub>, 15 % for CH<sub>3</sub>I and 20 % for CH<sub>2</sub>I<sub>2</sub> determined from duplicates. For a detailed description see Hepach et al. (2014).

# 9 2.2 Phytoplankton pigment analysis and continuous measurement of 10 chlorophyll *a*

11 Water samples were filtered onto GF/F filters, shock-frozen in liquid nitrogen and stored 12 at -80 °C. Pigments listed in Table 1 of Taylor et al. (2011) were analyzed using a HPLC 13 technique according to Barlow et al. (1997) as described in Taylor et al. (2011). Surface 14 pigment data were already used in a study by Bracher et al. (2015). All pigment data are 15 already published and available from PANGAEA (http://doi.pangaea.de/10.1594/PANGAEA.848586). For interpretation of the pigment data, 16 17 CHEMTAX® (Mackey et al., 1996) was used, and initiated with the pigment ratio matrix 18 proposed by Veldhuis and Kraay (2004) for the subtropical Atlantic Ocean. The following 19 phytoplankton groups were evaluated: diatoms, Synechococcus-type, Prochlorococcus HL (high light adapted) and Prochlorococcus LL (low light adapted), dinoflagellates, 20 21 haptophytes, pelagophytes, cryptophytes and prasinophytes.

22 10-min-averaged continuous surface maximum fluorescence measured by a microFlu-chl 23 fluorometer from TriOS located in the ships moon pool was used to derive continuous total 24 Chl a (TChl a) concentrations along the underway transect. This is based on the assumption 25 that active fluorescence F is correlated to the amount of available TChl a (Kolber and 26 Falkowski, 1993). The method to convert fluorescence to TChl a is described in detail in 27 Taylor et al. (2011). Mean conversion factors specific for each zone were determined for 28 collocated F and HPLC-TChl a (the sum of monovinyl Chl a, divinyl Chl a and 29 Chlorophyllide *a*; the latter is mainly formed as artefact of the former two during the 30 extraction process and therefore included in the calculation) measurements. A linear 31 regression of r = 0.83 (p < 0.01, n = 89) was observed between surface HPLC-derived TChl a 32 and F-derived TChl a, which indicates the robustness of the conversion of F to TChl a. The 33 high depth resolved chlorophyll profiles were derived from fluorescence values obtained from

- 1 a Dr. Haardt fluoremeter mounted to the CTD and calibrated with collocated HPLC-derived
- 2 TChl *a* concentrations at six depths of each profile according to Fujiki et al. (2011).

#### 3 **2.3Correlation analysis of halocarbons**

4 Different parameters were correlated to surface water halocarbons. Physical influences were 5 investigated with 10 min averages of sea surface temperature (SST), sea surface salinity 6 (SSS), global radiation and wind speed, and a relationship with location was explored using 7 latitude. Biological parameters used for correlations were TChl a, and the abundances of all 8 phytoplankton groups. Since most of the data sets were not normally distributed and common 9 transformations into normal distributions were not possible, the Spearman's rank correlation 10 coefficient  $r_s$  was applied. All correlations with p < 0.05 were regarded as significant. 11 Correlation analysis of the entire depth profile dataset using the Spearman's rank coefficient

did not allow for drawing specific conclusions due to the complexity of the data set. Hence, the mixed influences on water column halocarbon concentrations were examined with principal component analysis (PCA) using MATLAB®. PCA analyzes the collective variance of a dataset including several variables. The PCA has the advantage to simplify a complex data set and find similarities. Concentrations of all four halocarbons, all phytoplankton groups, the TChl *a*, density, temperature, and salinity were included.

#### 18 2.4 Mixed layer depth

Mixed layer depths  $z_{ML}$  were determined using the method introduced by Kara et al. (2000). It proved to be closest to the visually determined  $z_{ML}$  from the temperature, salinity and density profiles. The mixed layer of each CTD profile was calculated as the depth where the temperature from the reference depth in the upper well-mixed temperature region was reduced by a threshold value of 0.8 °C.

#### 24 **2.5Calculation of sea-to-air fluxes of halocarbons**

25 The air-sea gas exchange parameterization of Nightingale et al. (2000) was applied to 26 calculate sea-to-air fluxes  $F_{as}$  of halocarbons (equation 1). Schmidt number corrections as 27 reported by Quack and Wallace (2003) were applied to determine the compound specific 28 transfer coefficient  $k_w$ . The air-sea concentration gradient was computed from sea surface 29 water measurements and mean atmospheric mixing ratios  $c_{atm}$  of 2.50 ppt for CHBr<sub>3</sub>, 1.20 ppt 30 for CH<sub>2</sub>Br<sub>2</sub>, and 0.50 ppt for CH<sub>3</sub>I determined from 10 atmospheric data points during 31 MSM18/3, and atmospheric mixing ratios of 0.01 ppt for CH<sub>2</sub>I<sub>2</sub> as reported by Jones et al. 32 (2010) for the tropical Atlantic. Henry's law constants H of Moore and co-workers (Moore et

al., 1995a; Moore et al., 1995b) were used to obtain the equilibrium concentrations  $c_{atm}/H$ .

$$1 F_{as} = k_w \cdot \left(c_w - \frac{c_{atm}}{H}\right) (1)$$

#### 2 **2.6Calculation of diapycnal fluxes of halocarbons**

To estimate the halocarbon transport perpendicular to the stratification, equation 2 was used with  $F_{dia}$  as the diapycnal flux in mol m<sup>-2</sup> s<sup>-1</sup>,  $\rho$  as the seawater density in kg m<sup>-3</sup>,  $\Delta c$  being the diapycnal gradient of the concentration in mol kg<sup>-1</sup>, and  $K_{dia}$  as the diapycnal diffusion coefficient in m<sup>2</sup> s<sup>-1</sup>.

$$7 F_{dia} = \rho \cdot K_{dia} \cdot \Delta c (2)$$

8 In the equatorial near surface water, molecular and double diffusion are negligible compared 9 to turbulent mixing.  $K_{dia}$  from turbulent mixing can be estimated from measurements of the 10 velocity microstructure (turbulent motions on length scales of centimeters to meters). During 11 MSM18/3, velocity microstructure profiling was performed immediately before or after 12 taking halocarbon profiles, so that local and pointwise in time estimates of the diapycnal flux 13 resulted from the combination of the two profiles via equation 2. The microstructure profiler 14 (MSS) was a loosely tethered MSS90 equipped with airfoil shear probes, manufactured by 15 Sea & Sun Technology. In order to calculate  $K_{dia}$  from velocity fluctuations measured by the 16 MSS, first the average spectrum of vertical shear for a depth interval of typically 10 to 50 m 17 was calculated and integrated to get an estimate of the average dissipation rate of turbulent kinetic energy (epsilon in W kg<sup>-1</sup>). Equation 3, first proposed by Osborn (1980) allows to 18 deduce  $K_{dia}$ , with y a function of the mixing efficiency and N the buoyancy frequency for the 19 20 chosen depth interval.

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$$K_{dia} = \gamma \cdot \frac{\varepsilon}{N^2}$$
 (3)

y was chosen to be 0.2 following Hummels et al. (2013) for the tropical Atlantic. A more detailed description of the method to derive  $K_{dia}$  and diapycnal fluxes below the mixed layer can be found in Schafstall et al. (2010), Hummels et al. (2013), and Schlundt et al. (2014).

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#### **3** Physical and biological characteristics of the investigation area

#### 27 **3.1 Oceanographic description**

The equatorial Atlantic is described by a complex current system. The surface is characterized by the westward South Equatorial Current (SEC), which spreads between 3° N and 15° S and reaches as deep as 100 m, but has shallow mixed layers close to the equator (Tomczak and

1 Godfrey, 2005). The Equatorial Undercurrent (EUC) can be found below the SEC (Molinari, 2 1982), and is a narrow band between 2° N and 2° S flowing towards the east while reducing 3 speed. It carries mostly water with characteristics of deeper tropical surface water (TSW) and of shallower central water. TSW around and north of the equator is characterized by high 4 5 temperatures and comparably low salinities due to enhanced precipitation (Tsuchiya et al., 6 1992). While the core of the EUC in the west is at 100 m, its position in the east follows the 7 seasonal vertical migration of the thermocline (Stramma and Schott, 1999). In agreement with 8 this, the mixed layer depth was shallow and ranged only between surface and 49 m with a 9 mean of 28 m during MSM18/3. The mixed layer was also exposed to diurnal variability. During daytime, it was shallower due to warmer air temperatures and more stratification. At 10 night, when the air temperature and SSTs cool, water mixes further down. The shallowest 11 12 mixed layers were found between 0° N and 3° S in agreement with the location of the EUC. 13 The Atlantic Cold Tongue (ACT) is a known feature in the equatorial region where SSTs between 20° and 5° W can drop by 5 – 7 °C during May to September (Weingartner and 14 15 Weisberg, 1991). Many uncertainties remain with respect to the exact mechanisms that lead to the development of the ACT. Jouanno et al. (2011) suggested that the strong increase of the 16 17 westward SEC associated with the ITCZ (Philander and Pacanowski, 1986), and the 18 maximum shear above the core of the underlying EUC lead to the low SSTs, confirmed later 19 by microstructure measurements (Hummels et al., 2013; Schlundt et al., 2014). Although the shear is maximal at 0° E, maximum cooling appears at 10° W due to the stronger stratification 20 21 in the eastern basin of the equatorial Atlantic. SSTs during MSM18/3 of mean (range) 24.4 22 (22.1 - 29.0) °C and SSSs of 35.7 (34.5 - 36.3) were measured in the investigated region 23 (Table 1, Figure 2). Generally, high SSTs and low SSSs of less than 35.5 in the TSW were 24 observed north of the equator. Lower SSTs and higher SSSs were measured in the South 25 except for the 10° W section where these low SSTs and high SSSs were also found north of 26 the equator. Maximum SSTs around the equator of 28.5 °C were found at 3° N and 20° W, while the lowest SSTs of 22.1 °C were located at 1° N and 10° W (Figure 1, Figure 2, Table 27 28 1).

#### 29 **3.2 Biological description**

The cooling of SSTs in the ACT region is usually accompanied by a phytoplankton bloom. Grodsky et al. (2008) found a seasonal peak of TChl *a* of 0.60  $\mu$ g L<sup>-1</sup> in boreal summer. In comparison, surface TChl *a* during MSM18/3 reached values as high as 1.20  $\mu$ g L<sup>-1</sup> around 0.8° N and 0° E (Figure 2c). Very high TChl *a* concentrations above 1.00  $\mu$ g L<sup>-1</sup> were also measured from the continuous fluorescence sensor around 10° W, coincidentally with the most intense cooling. The three hourly HPLC measurements of up to 0.99  $\mu$ g L<sup>-1</sup> generally also agree with the high TChl *a* maximum values measured with the fluorescence sensor (Fig. 2, Table 1). Additionally, nitrate and phosphate were significantly anticorrelated with SST (not shown), hence the upwelled water of the EUC was connected to enhanced biological production.

6 The most abundant phytoplankton group in the ACT were *chrysophytes* in both surface water 7 and depth profiles during MSM18/3 (Figure 2a). Chrysophytes, golden algae with flagellar 8 hairs, are thought to be mostly common in freshwater (Round, 1986). Nevertheless, they have 9 been previously shown to be also the most abundant phytoplankton group in several regions 10 of the Atlantic ocean, including the lower latitudes around the equator (Kirkham et al., 2011). This group correlated significantly with SST ( $r_s = -0.45$ ) and SSS ( $r_s = 0.48$ ) (Table 2), it 11 hence seems to be associated with the upwelling water of the EUC. In the surface water, 12 chlorophytes and Prochlorococcus HL correlated positively with SST ( $r_s = 0.13$ , not 13 significant, and  $r_s = 0.44$ , significant) and negatively with SSS ( $r_s = -0.15$ , not significant, and 14  $r_s = -0.39$ , significant). They were associated with warmer and less salty water masses than 15 chrysophytes, dinoflagellates and haptophytes. Thus, they were found predominantly north of 16 17 the equator. Prochlorococcus HL dominate among the species occurring from the surface 18 down to 50 m. Prochlorococcus LL, only observed in deeper layers (not shown here), were 19 the most abundant group from about 75 m downwards in the water column. These results are 20 in agreement with Johnson et al. (2006), where it was shown that *Prochlorococcus* dominate 21 in oligotrophic tropical waters, especially where nutrient concentrations are low at high 22 temperatures (between 15° S and 15° N of the Atlantic Ocean).

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#### 24 **4 Results**

#### 25 **4.1 Surface water**

#### 26 4.1.1 CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub>

27 Large regional variations were observed for the bromocarbons, especially for CHBr<sub>3</sub> in surface water of the tropical Atlantic with a mean of 12.9 (1.8 - 44.7) pmol L<sup>-1</sup>, and of 3.7 28 (0.9 - 9.2) pmol L<sup>-1</sup> for CH<sub>2</sub>Br<sub>2</sub> (Figure 2, Table 1). Concentrations from the underway 29 30 measurements and from the shallowest profile depths (<10m) were included in the evaluation 31 of the surface water concentrations. The observed values are in agreement with data from the 32 tropical oligotrophic Atlantic north of 16° N and the Mauritanian upwelling ranging between 1.0 and 43.6 for CHBr<sub>3</sub> and 0.6 - 9.4 pmol L<sup>-1</sup> for CH<sub>2</sub>Br<sub>2</sub> with the largest values close to the 33 34 coast and the upwelling (Quack et al., 2007a; Carpenter et al., 2009; Hepach et al., 2014).

Quack et al. (2004) observed lower CHBr<sub>3</sub> of 2.3 pmol  $L^{-1}$  and CH<sub>2</sub>Br<sub>2</sub> of 0.2 pmol  $L^{-1}$  at 1  $10^{\circ}$  N through the tropical Atlantic in boreal fall and values of 12.8 and 5.3 pmol L<sup>-1</sup> for 2 CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub> at the equator in agreement with our study. Values of up to 10 pmol L<sup>-1</sup> 3 (CHBr<sub>3</sub>) and 3 pmol  $L^{-1}$  (CH<sub>2</sub>Br<sub>2</sub>) near the equator were reported by Liu et al. (2013b). The 4 5 latter study covers the region during October and November, indicating that the equatorial 6 Atlantic seems to be a larger source for bromocarbons during the intense cooling in the 7 summer months. Both compounds show the same pattern in surface water throughout the 8 MSM18/3 cruise with hot spots slightly south of the equator.

9 The very good correlation between CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub> is in agreement with studies from several regions, mostly attributed to related sources for both compounds from macro- and 10 11 microalgae (Nightingale et al., 1995; Moore et al., 1996; Schall et al., 1997; Laturnus, 2001; 12 Quack et al., 2007b; Karlsson et al., 2008). Significant correlations to SST, SSS and TChl a 13 were found for CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub>, while very low insignificant correlations were observed 14 with the 10 min averaged global radiation values (Table 2). The strongest correlations were found to *Prochlorococcus* HL with  $r_s = -0.70$  for CHBr<sub>3</sub> and -0.57 for CH<sub>2</sub>Br<sub>2</sub>, and to 15 chrysophytes with  $r_s = 0.43$ , and  $r_s = 0.41$ , respectively. 16

#### 17 **4.1.2 CH<sub>3</sub>I and CH<sub>2</sub>I<sub>2</sub>**

The second highest mean sea surface water concentration was observed for  $CH_3I$  of 5.5 (1.5 – 18 12.8) pmol  $L^{-1}$  (Figure 2, Table 1), which is in the range of earlier studies. These studies were 19 widely spread in the region from 20° S to 25° N between the coasts of South America and 20 Africa with values between 0 and 36.5 pmol  $L^{-1}$  (Happell and Wallace, 1996; Schall et al., 21 1997; Richter and Wallace, 2004; Jones et al., 2010; Hepach et al., 2014). 7.1 to 16.4 pmol  $L^{-1}$ 22 were detected in the vicinity of our investigated region (Richter and Wallace, 2004). CH<sub>2</sub>I<sub>2</sub> 23 was characterized by the lowest sea surface water concentrations of 1.1 (0.3 – 3.7) pmol  $L^{-1}$ 24 25 during MSM18/3. Literature reports of CH<sub>2</sub>I<sub>2</sub> in the tropical Atlantic are very sparse: Schall et al. (1997) report on average three times higher values of 3.4 (2.1 – 6.8) pmol  $L^{-1}$  in the 26 tropical Atlantic, while Jones et al. (2010) measured a five times higher mean of 5.8 (0.9 and 27 17.1) pmol  $L^{-1}$  (reported in Ziska et al. (2013)) in the northern tropical Atlantic. 28

Similar to CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub>, sea surface CH<sub>3</sub>I was significantly anticorrelated with SST ( $r_s = -0.42$ ) and not correlated with global radiation (Table 2). In contrast to the bromocarbons, correlations were neither found to SSS, nor to latitude. Additionally, sea surface CH<sub>3</sub>I correlated to biomass indicators (TChl *a*:  $r_s = 0.36$ ). The regional distribution of CH<sub>3</sub>I often followed qualitatively that of *haptophytes* ( $r_s = 0.39$ ) with the most elevated concentrations south of the equator. Positive correlations were also found to *dinoflagellates*  1  $(r_s = 0.29)$  and *chrysophytes*  $(r_s = 0.26)$ . A weak, but significant anticorrelation was observed 2 to wind speed  $(r_s = -0.22)$ . In contrast to the other three halocarbons, CH<sub>2</sub>I<sub>2</sub> was positively 3 correlated with SST  $(r_s = 0.33)$ , and elevated concentrations were observed mostly north of 4 the equator. A weak negative correlation of CH<sub>2</sub>I<sub>2</sub> was found with global radiation 5  $(r_s = -0.25)$ , indicating higher sea surface CH<sub>2</sub>I<sub>2</sub> during the night time and lower 6 concentrations during the day. CH<sub>2</sub>I<sub>2</sub> correlated both with *chlorophytes*  $(r_s = 0.32)$  and 7 *Prochlorococcus* HL  $(r_s = 0.27)$ .

#### 8 4.2 Water column

#### 9 4.2.1 $CHBr_3$ and $CH_2Br_2$

10 CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub> showed maxima at the surface, in the of the mixed layer and below it 11 (Figure 3, Table 3). The highest deep maximum concentrations of both CHBr<sub>3</sub> (up to 19.2 12 pmol L<sup>-1</sup>) and CH<sub>2</sub>Br<sub>2</sub> (up to 10.6 pmol L<sup>-1</sup>) were observed in profile 4. At stations where 13 CHBr<sub>3</sub> was most elevated at the surface (profiles 2, 7, 12, 13), much higher overall CHBr<sub>3</sub> 14 concentrations of up to 35.0 pmol L<sup>-1</sup> were measured. CH<sub>2</sub>Br<sub>2</sub> only reached maximum values 15 of up to 6.6 pmol L<sup>-1</sup> in the surface (profiles 2, 7).

In contrast to surface water, CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub> were distributed differently in the water 16 17 column with CH<sub>2</sub>Br<sub>2</sub> being elevated 10 m below CHBr<sub>3</sub> in several profiles (Figure 3e). This can also be seen in the T-S diagrams of these compounds (Figure 4a, b): while the most 18 elevated CHBr<sub>3</sub> was observed in the density layers between 1024 and 1025 kg m<sup>-3</sup> (shallower 19 central water of the EUC), CH<sub>2</sub>Br<sub>2</sub> was often also elevated in the denser, deeper layers below 20 21 30 m (Table 3). The maxima of both compounds were mostly in the vicinity of the TChl a 22 maximum. Results of the PCA (Figure 5) also show the dissimilarity of CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub> at 23 depth: while the variance of CHBr<sub>3</sub> seems comparable to salinity and several phytoplankton 24 groups such as *chrysophytes*, CH<sub>2</sub>Br<sub>2</sub> shows many similarities with the distribution of CH<sub>2</sub>I<sub>2</sub> 25 in the water column.

#### 26 4.2.2 CH<sub>3</sub>I and CH<sub>2</sub>I<sub>2</sub>

- In agreement with  $CHBr_3$  and  $CH_2Br_2$ ,  $CH_3I$  was elevated in the surface (three profiles 4, 6, 7)
- 28 (Table 4, Figure 3b) with values of up to 12.8 pmol  $L^{-1}$ , and also elevated in the deeper layers
- 29 in and below the mixed layer (Figure 3f), reaching up to 8.5 pmol  $L^{-1}$ . Most maxima of  $CH_3I$
- 30 were observed closer to the surface within the mixed layer (Figure 4d). The PCA of  $CH_3I$
- 31 revealed that its variance was similar to the variance of *dinoflagellates* and temperature32 (Figure 5).
- 33  $CH_2I_2$  was always depleted in the surface. Maxima of  $CH_2I_2$  were found in different depths, 34 sometimes associated with the TChl *a* maximum (Figure 3f), and mostly below the mixed

1 layer (Figure 3j). The maxima in deeper depths appeared concurrently with the deeper  $CH_2Br_2$ 2 maxima (Figure 4), which is also expressed in the PCA (Figure 5). Values were generally 3 much higher in deeper depths with e.g. 13.8 pmol L<sup>-1</sup> between 60 and 100 m at profile 5. The 4 highest concentrations of the whole cruise of 16.0 pmol L<sup>-1</sup> (profile 1) were found between 30 5 and 60 m. Concentrations of only up to 12.0 pmol L<sup>-1</sup> were found between 0 and 30 m (profile 6 b) (Table 4).

7 4.3 Fluxes

#### 8 4.3.1 CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub>

Sea-to-air fluxes of CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub> of 644 (-146 – 4285) and 187 (-3 – 762) pmol m<sup>-2</sup> h<sup>-1</sup> 9 10 during MSM18/3 were larger during the first two western NS-transects of the cruise which 11 were characterized by higher seawater concentrations, as well as higher wind speeds (Table 1, Figure 6). Carpenter et al. (2009) and Hepach et al. (2014) reported -150 and 3504 12 pmol  $m^{-2} h^{-1}$  CHBr<sub>3</sub> fluxes as well as of 5 – 917 for CH<sub>2</sub>Br<sub>2</sub> from the Cape Verde and 13 14 Mauritanian upwelling region. The lower fluxes in the equatorial region are a result of the lower wind speeds measured during MSM18/3, ranging from 0.3 - 11.1 with a mean of 6.1 15 m s<sup>-1</sup>, and the lower concentration gradients in comparison to Carpenter et al. (2009). Quack 16 et al. (2004) reported CHBr<sub>3</sub> fluxes from the equatorial Atlantic of 2700 ( $\pm$  800) pmol m<sup>-2</sup> h<sup>-1</sup>, 17 18 which compare well to this study.

19 Diapycnal fluxes are the fluxes of halocarbons that diffuse out or into the mixed layer from 20 below the thermocline. Maxima within the mixed layer will lead to fluxes towards the 21 thermocline, while maxima below the mixed layer will result in a flux of halocarbon-22 molecules into the mixed layer. Diapycnal fluxes of halocarbons were generally low although 23 the EUC can lead to enhanced mixing. This is due to the comparably small concentration 24 gradients of the halocarbons. Diapycnal fluxes were 80 (CHBr<sub>3</sub>) to 200 times (CH<sub>2</sub>Br<sub>2</sub>) lower 25 than sea-to-air fluxes (Table 5). They acted both as a source and a sink for halocarbons in the mixed layer. At eight stations, CHBr<sub>3</sub> was diffusing into the mixed layer, providing on 26 average 5 (0 – 14) pmol m<sup>-2</sup> h<sup>-1</sup> from below to the mixed layer budget of CHBr<sub>3</sub>. On the other 27 hand, on average 30 (2 – 125) pmol  $m^{-2} h^{-1}$  were diffusing out of the mixed layer, which is the 28 29 highest flux to the thermocline of all four halocarbons, as a result of its large concentration gradients across the bottom of the mixed layer. Diapycnal fluxes of CH<sub>2</sub>Br<sub>2</sub> were generally 30 31 lower than for CHBr<sub>3</sub> due to its lower concentration gradients. Its fluxes into the mixed layer from eight profiles were on average 3 (0 - 8) pmol m<sup>-2</sup> h<sup>-1</sup>, while the diapycnal flux reduced 32 the mixed layer budget of CH<sub>2</sub>Br<sub>2</sub> by 2 (0 – 8) pmol  $m^{-2} h^{-1}$  at the remaining five stations. 33

#### 1 4.3.2 CH<sub>3</sub>I and CH<sub>2</sub>I<sub>2</sub>

CH<sub>3</sub>I sea-to-air fluxes were on average 425 (34 – 1300) pmol  $m^{-2} h^{-1}$  during the cruise. During 2 the eastern NS-transects, fluxes were elevated at several locations mostly during daytime in 3 4 contrast to the bromocarbons, in accordance to a larger concentration gradient of CH<sub>3</sub>I in that 5 region (Table 1, Figure 6). The fluxes are only half of the sea-to-air fluxes from the equatorial Atlantic region reported by Richter and Wallace (2004) of  $958 \pm 750$  pmol m<sup>-2</sup> h<sup>-1</sup> and a fifth 6 of the fluxes reported from Jones et al. (2010) of on average 2154 pmol m<sup>-2</sup> h<sup>-1</sup> from the Cape 7 Verde and Mauritanian upwelling region. But, they were two times larger than the fluxes of 8 Hepach et al. (2014) of on average 246 pmol m<sup>-2</sup> h<sup>-1</sup>. CH<sub>2</sub>I<sub>2</sub> fluxes were generally larger in the 9 beginning of the cruise where higher wind speeds and higher surface water concentrations 10 existed. Only few studies have published sea-to-air fluxes of CH<sub>2</sub>I<sub>2</sub> from the tropical ocean. 11 CH<sub>2</sub>I<sub>2</sub> emissions calculated for MSM18/3 are with only 82 (3 – 382) pmol  $m^{-2} h^{-1}$  very low in 12 comparison to mean fluxes reported by Jones et al. (2010) of on average 541 -13 688 pmol m<sup>-2</sup> h<sup>-1</sup>, which are the result of higher oceanic CH<sub>2</sub>I<sub>2</sub> (Jones et al., 2010). 14

Similar to the bromocarbons, diapycnal fluxes of CH<sub>3</sub>I and CH<sub>2</sub>I<sub>2</sub> were generally lower (117 15 16 and 7 times, respectively) than sea-to-air fluxes (Table 5). Due to the larger CH<sub>3</sub>I 17 concentrations in the mixed layer compared to the upper thermocline, diapycnal fluxes of 5 (1 -13) pmol m<sup>-2</sup> h<sup>-1</sup> were mostly acting as a sink for the mixed layer budget. Only at three 18 stations, 2 (1 - 5) pmol m<sup>-2</sup> h<sup>-1</sup> were transported into the mixed layer. Diapycnal fluxes of 19  $CH_2I_2$  acted mostly as source for the mixed layer, providing on average 12 (0 – 20 21 39) pmol  $m^{-2} h^{-1}$  due to its much higher concentrations in the water below. This represents the highest halocarbon flux of the four compounds into the mixed layer. The diapycnal flux of 22  $CH_2I_2$  of 2 (0 – 4) pmol m<sup>-2</sup> h<sup>-1</sup> out of the mixed layer was only observed at three stations. 23

24

#### 25 **5** Discussion

#### 26 **5.1 Surface water distribution**

#### 27 5.1.1 CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub>

The equatorial Atlantic is a source of CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub> to the atmosphere during the ACT season, and the correlations of their water concentrations to biogenic parameters indicate biological formation. CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub> correlated significantly, but weakly with TChl *a*, which is not an unusual feature (Abrahamsson et al., 2004a; Carpenter et al., 2009; Liu et al., 2011; Hepach et al., 2014). It has been suggested that CHBr<sub>3</sub> is not produced directly from phytoplankton, but rather from dissolved organic matter (DOM) present in sea water (Lin and Manley, 2012). This was more closely investigated in laboratory experiments by Liu et al.

1 (2015), who suggested that the weak in-situ correlations of bromocarbons with Chl a are a 2 result of this indirect production pathway. The correlation with certain phytoplankton groups 3 may then be caused by the production of phytoplankton-specific DOM. The very negative correlations of bromocarbons with SST and positive correlations with SSS indicate a 4 5 relationship of bromocarbon abundance with processes within the cold and nutrient-rich 6 upwelled water of the EUC (section 3.2), supported by the T-S diagrams (Figure 4). Weak, but significant negative correlations with latitude ( $r_s = -0.38$  for CHBr<sub>3</sub> and  $r_s = -0.18$  for 7 8 CH<sub>2</sub>Br<sub>2</sub>) and maximum values of the bromocarbons between 2 and 3° S, where EUC water 9 reaches the surface, underline this hypothesis. Although the correlation analysis of 10 halocarbons with phytoplankton groups cannot directly resolve production and loss processes 11 by algal activity, it is still an indicator for possible involvement of these species in halocarbon 12 production. Bromocarbon production might exceed loss processes, which leads to the 13 observed statistical link of CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub> to chrysophytes. Chrysophytes are to our 14 knowledge not yet among observed halocarbon producers in incubation and field studies. The 15 strong negative correlations of Prochlorococcus HL with CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub> have been 16 observed previously (Hepach et al., 2014). These significant negative correlations can be 17 explained by the large abundance of Prochlorococcus in warm water while bromocarbons on 18 the other hand are more correlated with the cooler water of the EUC, which is richer in 19 nutrients and chrysophytes, haptophytes and dinoflagellates.

#### 20 5.1.2 CH<sub>3</sub>I and CH<sub>2</sub>I<sub>2</sub>

21 CH<sub>3</sub>I concentrations and wind speed were weakly anticorrelated during MSM18/3. Richter 22 (2004) interprets this as depletion of the surface concentrations, when air-sea fluxes exceed 23 the production rate during high wind speed. There are two production mechanisms suggested for CH<sub>3</sub>I. Previous studies (Richter and Wallace, 2004; Jones et al., 2010) have attributed 24 25 CH<sub>3</sub>I in the tropical ocean mainly to photochemical formation based on the observations of 26 Moore and Zafiriou (1994). In contrast to these studies, indications for biological formation of 27 CH<sub>3</sub>I were found in the ACT region during our study. CH<sub>3</sub>I showed a weak negative 28 correlation with SST, significant correlations with the biologically produced CHBr<sub>3</sub> and 29 CH<sub>2</sub>Br<sub>2</sub> (Table 2) and with TChl *a* as biomass indicator, and no correlation to global radiation. 30 These imply a relationship with the biologically active upwelled water. Elevated concentrations of CH<sub>3</sub>I were found between 10° and 5° W during midday (see CH<sub>3</sub>I in 31 32 comparison to global radiation in Figure 2), which could be a result of photochemical 33 formation. Thus we suggest that photochemistry and biological production likely both played 34 a role during MSM18/3. Haptophytes correlated most significantly of the phytoplankton

1 groups with CH<sub>3</sub>I and have already been shown to produce CH<sub>3</sub>I both in the laboratory (Itoh 2 et al., 1997; Manley and de la Cuesta, 1997; Scarratt and Moore, 1998; Smythe-Wright et al., 3 2010) and in the field (Abrahamsson et al., 2004b). Correlations during MSM18/3 4 additionally indicate a possible involvement of *dinoflagellates* and *chrysophytes* in the 5 production of methyl iodide (Table 2). The importance of oceanic CH<sub>3</sub>I production by 6 Prochlorochcoccus is a matter of dispute. Brownell et al. (2010) report it to be a minor 7 source, in contrast to both Smythe-Wright et al. (2006) and Hughes et al. (2010, 2011). No 8 evidence of involvement of *Prochlorococcus* HL was found during MSM18/3.

9 The very low sea surface concentrations of  $CH_2I_2$  with lowest concentrations during the day 10 can be explained by its fast photolysis (few minutes lifetime in surface sea water) (Jones and 11 Carpenter, 2005; Martino et al., 2005). Although  $CH_2I_2$  is generally assumed to be of biogenic 12 origin in the open ocean (Moore and Tokarczyk, 1993; Yamamoto et al., 2001; Orlikowska 13 and Schulz-Bull, 2009; Hopkins et al., 2013), great uncertainties remain as to which species 14 are involved in its production. During MSM18/3, indications were found for different source 15 species than of the other three compounds (*chlorophytes* and *Prochlorococcus* HL).

#### 16 **5.2 Water column distribution**

17 Halocarbon maxima in the TChl a maximum, attributed to their biological production, are 18 often observed from polar to tropical regions (Moore and Tokarczyk, 1993; Moore and 19 Groszko, 1999; Yamamoto et al., 2001; Quack et al., 2004; Carpenter et al., 2007; Hughes et 20 al., 2009). In contrast, photochemical formation of  $CH_3I$  can lead to surface maxima (Happell 21 and Wallace, 1996). During MSM18/3, maxima of halocarbons were not always found in the 22 TChl a maximum. This does not contradict their biological production, as the location of the 23 TChl a maximum is not necessarily the location of highest biomass or primary production, but 24 rather reflects the photoadaption capability of the predominant phytoplankton groups 25 (Claustre and Marty, 1995). Unfortunately, neither biomass nor primary production was 26 measured during the cruise. Additionally, halocarbons could be produced by phytoplankton 27 groups that are not in the maximum of the biomass distribution in the water column, and the 28 location of the halocarbon maximum might be more determined from their sink processes 29 than from their production. Surprisingly, the time of day, influencing sink and production 30 processes, seemed to play a minor role for the shape of the profiles for all four compounds 31 (see the location of the CTD stations in Fig. 2).

#### 32 5.2.1 CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub>

33 In contrast to their similar occurrence in the surface,  $CHBr_3$  and  $CH_2Br_2$  showed different 34 distributions in the water column (Figure 5). Strong indications for biological sources of

1 CHBr<sub>3</sub> exist in the PCA, and *chrysophytes* as potential source group are in agreement to the 2 surface water observations (Table 2, Figure 5). Maximum CH<sub>2</sub>Br<sub>2</sub> concentrations were 3 occasionally found below the CHBr<sub>3</sub> maxima, which have already been observed in the 4 Mauritanian upwelling (Quack et al., 2007b). The deeper maxima may be either due to an 5 additional source of CH<sub>2</sub>Br<sub>2</sub> such as the biologically mediated conversion of CHBr<sub>3</sub> (Hughes 6 et al., 2013) or to a faster degradation of CHBr<sub>3</sub> than of CH<sub>2</sub>Br<sub>2</sub> at depth. Sinks for CHBr<sub>3</sub> and 7 CH<sub>2</sub>Br<sub>2</sub> in tropical surface waters include very slow hydrolysis (hundreds to thousands of 8 years) (Mabey and Mill, 1978) and slow halogen substitution (5 years) (Geen, 1992). 9 Photolysis, which has been suggested to be faster for CHBr<sub>3</sub> (9 years with a mixed layer of 10 100 m for CHBr<sub>3</sub>) than for CH<sub>2</sub>Br<sub>2</sub> (Carpenter et al., 2009) would be of more significance in 11 the surface layer. A faster degradation of CHBr<sub>3</sub> in greater depths is also somewhat contrary 12 to the observed very fast bacterial degradation of CH<sub>2</sub>Br<sub>2</sub> with a half-live of 2 days (Goodwin 13 et al., 1998). An additional source for CH<sub>2</sub>Br<sub>2</sub> that involves CHBr<sub>3</sub> therefore seems more 14 plausible. At four of the 13 stations, indications for the additional source were found. There, 15 maximum CH<sub>2</sub>Br<sub>2</sub> concentrations were found below CHBr<sub>3</sub>, which could be the result of its 16 faster conversion to CH<sub>2</sub>Br<sub>2</sub> than its production. CH<sub>2</sub>Br<sub>2</sub> in denser water is also co-located 17 with *Prochlorococcus* LL, which might be involved in the CHBr<sub>3</sub>-conversion.

#### 18 **5.2.2 CH<sub>3</sub>I and CH<sub>2</sub>I<sub>2</sub>**

19 CH<sub>3</sub>I was usually elevated in the top 30 m of the water column apart from three profiles, 20 where maximum concentrations were found between 30 and 60 m. The surface maxima, as 21 seen in the T-S diagram (Figure 4), support the photochemical formation of CH<sub>3</sub>I (Happell 22 and Wallace, 1996). Deeper maxima could also arise if the sea-to-air flux exceeds the 23 photochemical production. However, the low wind speed during the cruise (section 3), the 24 relationship with biological parameters, and the partly co-located maxima with the other three 25 biogenic halocarbons (Figure 3, Figure 5) also point to a direct production of CH<sub>3</sub>I from 26 phytoplankton. These include *dinoflagellates* as indicated by the correlations and the PCA 27 (Figure 5).

CH<sub>2</sub>I<sub>2</sub> was always depleted in the surface with respect to the underlying water column as a result of its strong photolysis (Jones and Carpenter, 2005; Martino et al., 2006). It was frequently elevated below the TChl *a* maximum and below the base of the mixed layer (Figure 3) in contrast to previous studies (Moore and Tokarczyk, 1993; Yamamoto et al., 2001). The similarity in its distribution to  $CH_2Br_2$  (Figure 4, Figure 5) could indicate similar production and sink processes at depth. Bacterial formation of  $CH_2I_2$  (Fuse et al., 2003; Amachi et al., 2005) in the upper thermocline could also be an additional source for this 1 compound. Alternatively,  $CH_2I_2$  may not degrade as quickly as  $CHBr_3$  and  $CH_3I$  in greater 2 depths, which would lead to its accumulation below the mixed layer.

#### **5.3 Factors contributing to halocarbon emissions from the mixed layer**

4 Halocarbon emissions into the atmosphere depend strongly on the mixed layer budget of these 5 compounds, which is determined by their sources and sinks. It is unclear, where the main 6 halocarbon production occurs. It has been suggested that it takes mainly place in the 7 subsurface TChl a maximum (Quack et al., 2004; Martino et al., 2006), whereas other model 8 studies assume production of e.g. CHBr<sub>3</sub> to be coupled to primary production in the whole 9 water column (Hense and Quack, 2009). Assuming production of halocarbons takes place 10 mainly in the TChl a maximum, which is often located below the mixed layer, diapycnal 11 fluxes from below the thermocline will be the most important source for mixed layer 12 halocarbons.

#### 13 **5.3.1** Transport and loss processes in the mixed layer

14 To evaluate the significance of halocarbon production below the mixed layer for emissions 15 into the atmosphere, production, loss and transport processes have to be considered. The 16 diapycnal fluxes of the four halocarbons were calculated from 13 halocarbon profiles and 17 parallel measurements of eddy diffusivity (section 4.3). The data are characterized by a low 18 depth resolution of the halocarbons within the water column and a short validity of the 19 diffusion coefficients, which make the diapycnal fluxes subject to some uncertainties. Given 20 that the depth profiles measured during MSM18/3 agree well to previous studies from the 21 tropical ocean (Yamamoto et al., 2001; Quack et al., 2004), a general idea of the significance 22 of diapycnal fluxes for the mixed layer budget of halocarbons can be obtained. The chemical 23 loss rates are estimated from published data which include hydrolysis, halogen substitution 24 and photolysis. The half-lives of CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub> due to hydrolysis are hundreds to 25 thousands of years (Mabey and Mill, 1978), while for CH<sub>3</sub>I, the half-life due to hydrolysis 26 ranges from 1600 days at 25 °C to 4000 days at 5 °C (Elliott and Rowland, 1995). The half-27 life of CHBr<sub>3</sub> with respect to photolysis is 9 years assuming a mixed layer depth of 100 m and 28 is potentially slower for  $CH_2Br_2$  (Carpenter and Liss, 2000) halogen-substitution is 5 years in 29 warm waters (Geen, 1992). Liu et al. (2011) calculated the half-life of CHBr<sub>3</sub> due to 30 photolysis in a coastal mixed layer of 5 m to be only 82 days. Mixed layers during MSM18/3 31 were from down to 49 m, photolysis of bromocarbons in the mixed layer will lead to half-32 lives of several months. Sea-to-air flux is the most significant sink for CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub> 33 from the mixed layer. Mean half-lives of 8 days were calculated for both compounds during 34 MSM18/3, based on the fluxes (section 4.3.1) and the mixed layer depths during the cruise

1 (Table 3). We consider a very short time scale of 1 h for our budget calculations due to the 2 validity of the diapycnal flux coefficients, while the general findings of our calculations are 3 also valid for a longer time scale. As the sink from the mixed layer due to sea-to-air fluxes is a 4 magnitude larger than the other mentioned sinks, we will neglect them in our estimates for 5 CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub> as they do not play a large role. Photolysis of CH<sub>3</sub>I is very slow in 6 comparison to halide substitution (Zika et al., 1984). The latter is suggested to be an important 7 sink in the tropical ocean during low wind speeds (Jones and Carpenter, 2007), while large 8 wind speeds favor sea-to-air fluxes as main sink (mean half-life of 8 days during MSM18/3). 9 All three sink processes are included in our budget estimates using the rates published by 10 Elliott and Rowland (1993). For CH<sub>2</sub>I<sub>2</sub>, photolysis is the most significant sink in surface water 11 (Jones and Carpenter, 2005). In our calculations, losses of CH<sub>2</sub>I<sub>2</sub> due to photolysis were 12 calculated according to Martino et al. (2006) with a photon flux calculated from the NASA COART model (Jin et al., 2006), a TChl *a* concentration of  $0.4 \,\mu g \, L^{-1}$ , absolute quantum 13 14 yields from Martino et al. (2006), and absorption cross sections determined by Jones and 15 Carpenter (2005).

#### 16 **5.3.2** Mixed layer budget of halocarbons during MSM18/3

In the following section, the results of the halocarbon budget calculations are presented. The total mixed layer concentrations were calculated at every station considering a water column with a volume of 1 x 1 x  $z_{ML}$  m<sup>3</sup>. Assuming that halocarbons are only produced below the mixed layer, the following relationship (equation 4) is valid for the steady state concentration  $C_{hal}$ , with  $F_{dia}$  and  $F_{adv}$  as the source terms from diapycnal fluxes and advection, while  $S_{as}$ (Figure 6) and  $S_{ch}$  represent the loss terms sea-to-air flux and chemical sinks as described in the previous section:

24 
$$C_{hal} = F_{dia} + F_{adv} - S_{as} - S_{ch}$$
 (4)

25  $S_{as}$  is the main sink term for CHBr<sub>3</sub>, CH<sub>2</sub>Br<sub>2</sub> and CH<sub>3</sub>I during MSM18/3 (Table 6). On the 26 short time scales considered here, diapycnal fluxes of CH<sub>3</sub>I, which can reduce the mixed layer by around 5 pmol per hour (Table 5), compete with the loss due to chloride substitution ( $S_{ch}$ ). 27 For CH<sub>2</sub>I<sub>2</sub>,  $S_{ch}$  (photolysis) is about 10 times higher than  $S_{as}$ , and reduces the mixed layer 28 29 budget by 24 % after 1 h. In total, diapycnal fluxes ( $F_{dia}$ ) into the mixed layer were not 30 sufficient to account for the losses of all four compounds from the mixed layer (Table 6). The 31 discrepancies with respect to the total mixed layer are 169 (CH<sub>2</sub>Br<sub>2</sub>), 255 (CH<sub>3</sub>I), 269 (CHBr<sub>3</sub>) to 8382 (CH<sub>2</sub>I<sub>2</sub>) pmol h<sup>-1</sup>, which are small compared to the total amount of 32 33 halocarbons in the mixed layer (CHBr<sub>3</sub> - 0.17 %, CH<sub>2</sub>Br<sub>2</sub> - 0.19 %, CH<sub>3</sub>I - 0.34 %, CH<sub>2</sub>I<sub>2</sub> -34 13.11 %). Possible reasons for the observed discrepancies are evaluated in the following.

Advection of the missing halocarbons,  $F_{adv}$ , likely does not play a large role for CH<sub>2</sub>Br<sub>2</sub>, CH<sub>3</sub>I 1 2 and CH<sub>2</sub>I<sub>2</sub>, since mean mixed layer concentrations of these compounds were rather 3 homogeneous in the whole region. Thus, only for CHBr<sub>3</sub>, with more variable concentrations, 4 advection may transport significant amounts from one location to another. In addition, 5 halocarbon maxima were found within the mixed layer, which may either result from a mixed layer that is not well mixed or halocarbon production is faster than mixing in the mixed layer. 6 7 According to the temperature and salinity profiles during the whole cruise (Figure 3), the 8 mixed layer was very well mixed. Consequently, production in the mixed layer is the most 9 likely process balancing the missing halocarbons (Table 6) as diapycnal fluxes and advection play minor roles. The maxima that occasionally evolve in the mixed layer suggest that 10 11 production of halocarbons is rapid, but may vary with depth. The mixed layer production 12 term, here called  $P_{ML}$ , has to be included in the budget calculation of equation 4:

13 
$$C_{hal} = F_{dig} + F_{ady} - S_{as} - S_{ch} + P_{MI}$$
 (5)

The relative production of halocarbons in the mixed layer is likely largest for CH<sub>2</sub>I<sub>2</sub>, because 14 15 its largest discrepancy arises from its rapid photolysis (up to 24 % loss in 1 h) (Table 6). This 16 is in agreement to earlier studies investigating macroalgal production, proposing larger release 17 rates of CH<sub>2</sub>I<sub>2</sub> than of CHBr<sub>3</sub>, CH<sub>2</sub>Br<sub>2</sub> and CH<sub>3</sub>I (Klick and Abrahamsson, 1992; Carpenter et 18 al., 2000).

#### 19 5.3.3 Production rates of halocarbons

20 From the budget calculations, described in the previous section, potential production rates  $P_{ML}$ 21 for the mixed layer are determined for each station. The mean production rates show large 22 standard deviations (Table 7), including the variability and uncertainties in the estimated 23 production rates. Production rates are  $34 \pm 65$  (CHBr<sub>3</sub>),  $10 \pm 12$  (CH<sub>2</sub>Br<sub>2</sub>),  $21 \pm 24$  (CH<sub>3</sub>I), and  $384 \pm 318$  pmol m<sup>-3</sup> h<sup>-1</sup> (CH<sub>2</sub>I<sub>2</sub>). These are the first estimated production rates of CHBr<sub>3</sub> 24 and CH<sub>2</sub>Br<sub>2</sub> for tropical phytoplankton species. For comparison to other studies, the 25 production rates from this study are converted to rates per µg TChl a (reported in Tables 3 26 and 4), which results in mean ( $\pm$  standard deviation) production rates of 2.5 x 10<sup>-3</sup>  $\pm$  4.5 x 10<sup>-3</sup> 27 (CHBr<sub>3</sub>), 8.4 x  $10^{-4} \pm 1.0$  x  $10^{-3}$  (CH<sub>2</sub>Br<sub>2</sub>), 2.2 x  $10^{-3} \pm 3.0$  x  $10^{-3}$  (CH<sub>3</sub>I) and 3.3 x  $10^{-2} \pm 3.3$  x 28  $10^{-2} \text{ pmol } [\mu \text{g TChl } a]^{-1} \text{ h}^{-1} (\text{CH}_2\text{I}_2).$ 29

#### 30 5.3.4 Comparison to previously reported rates – CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub>

Tokarczyk and Moore (1994) and Hughes et al. (2013) determined production rates from 31 polar algae in laboratory studies ranging between 2 x  $10^{-3}$  and 2.1 x  $10^{-2}$  pmol [µg Chl a]<sup>-1</sup> h<sup>-1</sup> 32 33 on average for CHBr<sub>3</sub>, depending on the growth phase, which is in the range of our calculated rates. Production rates for CH<sub>2</sub>Br<sub>2</sub> of on average  $2.1 - 4.2 \times 10^{-3}$  pmol [µg Chl a]<sup>-1</sup> h<sup>-1</sup> were 34

1 much higher than the ones calculated in our study (Tokarczyk and Moore, 1994). Karlsson et al. (2008) published production rates of 2.6 – 9.3 x  $10^{-2}$  pmol [µg Chl a]<sup>-1</sup> h<sup>-1</sup> for CHBr<sub>3</sub> 2 (depending on the time of day) and 5 x  $10^{-4}$  – 3.6 x  $10^{-3}$  pmol [µg Chl a]<sup>-1</sup> h<sup>-1</sup> for CH<sub>2</sub>Br<sub>2</sub> from 3 4 an in situ study in the Baltic Sea during a cyanobacterial bloom. Liu et al. (2011) calculated 417 (CHBr<sub>3</sub>) and 258 pmol m<sup>-3</sup> h<sup>-1</sup> (CH<sub>2</sub>Br<sub>2</sub>) for the subtropical and temperate eastern US 5 6 coast, which are tenfold higher than the production rates determined from our study (Table 7). 7 The differences between these studies and ours may have several origins. Taking an average 8 production rate for the total mixed layer during MSM18/3 does not take a potential variable 9 production with depth into account. Second, the different production rates determined in the monocultural studies (Tokarczyk and Moore, 1994; Hughes et al., 2013) show large variations 10 between different types of microalgae. Third, the indirect estimates during MSM18/3 are 11 12 afflicted by the uncertainties in the individual budget terms, which are also expressed in the 13 large standard deviations.

#### 14 5.3.5 Comparison to previously reported rates – CH<sub>3</sub>I and CH<sub>2</sub>I<sub>2</sub>

Production rates of CH<sub>3</sub>I determined from *Prochlorococcus* vary significantly from 5.8 x 10<sup>-4</sup> 15 to 9.4 x  $10^{-2}$  pmol [µg Chl a]<sup>-1</sup> h<sup>-1</sup>(Smythe-Wright et al., 2006; Brownell et al., 2010). Hughes 16 17 et al. (2011) suggested this variability to be caused by different cell states, e.g. healthier cells producing less CH<sub>3</sub>I. While Scarratt and Moore (1999) determined rates from 8.3 x  $10^{-3} - 5.0$ 18 x  $10^{-2}$  pmol [µg Chl a]<sup>-1</sup> h<sup>-1</sup> from a red microalgal species, Karlsson et al. (2008) reported a 19 rate of 1.0 x  $10^{-2}$  pmol CH<sub>3</sub>I [µg Chl a]<sup>-1</sup> h<sup>-1</sup> from a cyanobacterial bloom in the Baltic Sea, 20 21 which is at the higher end of the range mentioned here. Our estimates lie well within these 22 cited ranges of phytoplankton production rates and are thus a reasonable assumption for the 23 CH<sub>3</sub>I production strength of tropical algae (see section 5.1.2).

24 In contrast to the other three halocarbons, very few studies have actually determined 25 production rates of CH<sub>2</sub>I<sub>2</sub> from phytoplankton. CH<sub>2</sub>I<sub>2</sub> was shown to be produced in 26 comparatively larger concentrations than other halocarbons, but generally from fewer species 27 (six polar and temperate *diatom* species were tested, of which only two produced CH<sub>2</sub>I<sub>2</sub>) 28 (Moore et al., 1996). Martino et al. (2006) assumed a theoretical production rate of 17,000 pmol m<sup>-3</sup> h<sup>-1</sup> in the tropical equatorial Atlantic. These were calculated from previously 29 30 reported CH<sub>2</sub>ClI fluxes based on the assumption that CH<sub>2</sub>ClI is mainly formed during the photolysis of CH<sub>2</sub>I<sub>2</sub> and that CH<sub>2</sub>I<sub>2</sub> is only produced in the TChl *a* maximum. This rate 31 32 appears very large in comparison to our estimate and in comparison to the production rates of 33 the other halocarbons. We showed evidence that  $CH_2I_2$  is not only produced within the TChl a 34 maximum but in the whole mixed layer, thus, lower average production rates seem more 1 plausible.  $CH_2I_2$  together with  $CH_2CII$  have been suggested to be equally important carriers of 2 organoiodine into the troposphere (Saiz-Lopez et al., 2012), hence it is important to determine 3 specific phytoplankton production rates of  $CH_2I_2$  in future studies.

4 Our calculated production rates of CHBr<sub>3</sub>, CH<sub>2</sub>Br<sub>2</sub> and CH<sub>3</sub>I lie well within the ranges of 5 several laboratory and field studies of mostly temperate and polar algae, suggesting 6 production from tropical algae to be similarly significant. CH<sub>2</sub>I<sub>2</sub> was shown to be produced in 7 larger rates than the other three compounds, but very rapid photolysis leads to lower sea 8 surface concentrations of this compound. However, considering the large ranges in reported 9 production rates of CHBr<sub>3</sub>, CH<sub>2</sub>Br<sub>2</sub>, CH<sub>3</sub>I and the lack of studies concentrating on CH<sub>2</sub>I<sub>2</sub>, more incubation experiments are severely needed to constrain in situ production rates of 10 11 tropical algae. This information is crucial to evaluate the significance and contribution of the 12 tropical ocean with respect to halogen transport into the troposphere, and finally into the 13 stratosphere. Understanding the fate of halocarbons within the water column is an important 14 task to estimate their distribution and emissions from the future ocean.

15

#### 16 6 Summary and conclusions

17 Increased biological production during the Atlantic Cold Tongue (ACT) caused elevated CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub> concentrations of up to 44.7 pmol  $L^{-1}$  and up to 9.2 pmol  $L^{-1}$  within the 18 19 equatorial surface water with comparable concentrations to other tropical upwelling systems. 20 Both compounds showed similar distributions and maxima in the region where the Equatorial 21 Undercurrent (EUC) influences the surface water between 2° and 3° S with cooler water and 22 elevated nutrients. Chrysophytes, the dominating phytoplankton group in the equatorial 23 surface water, were likely involved in the bromocarbon production. In contrast to their similar 24 surface water occurrence, CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub> showed different distributions in the water 25 column. While CHBr<sub>3</sub> was mostly elevated in shallower layers in close proximity to the TChl 26 a maximum, CH<sub>2</sub>Br<sub>2</sub> frequently showed maxima in deeper water likely caused by an 27 additional source.

In contrast to other tropical Atlantic regions, correlations of  $CH_3I$  with  $CHBr_3$  and with biological parameters indicate biogenic formation of  $CH_3I$  during the ACT. Moderate  $CH_3I$ concentrations of up to 12.8 pmol L<sup>-1</sup> were measured in the surface water.  $CH_2I_2$  surface water and mixed layer concentrations were lowest due to its strong photolysis with maximum values of only 3.7 pmol L<sup>-1</sup>.  $CH_2I_2$  maxima below the mixed layer, suggest similar formation pathways to  $CH_2Br_2$  possibly tied to heterotrophic activities below the layers of maximum production.

1 Sea-to-air fluxes were the most important sink from the mixed layer of CHBr<sub>3</sub>, CH<sub>2</sub>Br<sub>2</sub> and 2  $CH_3I$ , while photolysis was the main sink for  $CH_2I_2$ . For the first time, halocarbon turbulent 3 fluxes from and into the mixed layer were calculated using microstructure measurements and 4 halocarbon concentration gradients in the water column. The significance of these diapycnal 5 fluxes as a source for mixed layer halocarbons, suggested by halocarbon maxima below the 6 mixed layer, was evaluated in comparison to sea-to-air fluxes and other sinks. All sinks of 7 halocarbons from the mixed layer were much larger than the diapycnal supply into the mixed 8 layer. Hence, halocarbon production in the entire mixed layer is the most important factor 9 contributing to marine emissions of these compounds.

Production rates of halocarbons were estimated from 13 profiles for the tropical mixed layer. Calucalted production rates varied between the stations and were:  $34 \pm 65$  pmol m<sup>-3</sup> h<sup>-1</sup> for CHBr<sub>3</sub>,  $10 \pm 12$  pmol m<sup>-3</sup> h<sup>-1</sup> for CH<sub>2</sub>Br<sub>2</sub>,  $21 \pm 24$  pmol m<sup>-3</sup> h<sup>-1</sup> for CH<sub>3</sub>I and  $384 \pm 318$  pmol m<sup>-3</sup> h<sup>-1</sup> for CH<sub>2</sub>I<sub>2</sub> with large variability between the different stations. These are generally in the range of rates reported from both monocultural and in situ incubation studies for CHBr<sub>3</sub>, CH<sub>2</sub>Br<sub>2</sub> and CH<sub>3</sub>I, while CH<sub>2</sub>I<sub>2</sub> seems to be emitted in larger concentrations from phytoplankton.

17 Our results show the need to conduct more process-related studies in the field. The first 18 consideration of diapycnal mixing revealed that maximum concentrations in the vicinity of 19 the TChl *a* maximum are insignificant for the mixed layer budget. Investigating the exact 20 mechanisms of formation, degradation and transport of halocarbons in the water column 21 remains an important task toward understanding current and future emissions of these 22 compounds. Understanding the actual processes that contribute to their concentrations and 23 distribution within the water column is crucial to predict their emissions. We therefore 24 suggest further mono-cultural incubation studies to determine species-dependent production 25 and consumption rates. Temporally resolved in situ incubations in different depths within the 26 water column in combination with diapycnal flux measurements will help to explain the 27 profile shapes. Further halocarbon emission studies in the tropical ocean in different seasons 28 are crucial to evaluate their importance for the stratospheric halogen loading in a global 29 perspective.

30

#### 31 Acknowledgements

We thank the chief scientist of the cruise MSM18/3 Arne Körtzinger, as well as the captain, the crew and the scientific crew of RV *Maria S. Merian* for all of their help. The authors acknowledge Sonja Wiegmann for pigment analysis, Bettina Taylor for CHEMTAX

1 calculations, and Martina Lohmann for nutrient measurements. We thank Björn Fiedler for 2 providing the fluorescence sensor data. We also appreciate the helpful input of Christa 3 Marandino. Additionally, the authors acknowledge NASA for providing satellite MODIS-4 Aqua data of June and July 2011. This work was part of the German research project 5 SOPRAN II and III (grant no. FKZ 03F0611A and 03F0662A) funded by the 6 Bundesministerium für Bildung und Forschung (BMBF), and was also supported by the EU 7 project SHIVA (grant no. FP7-ENV-2007-1-226224) and by the HGF Innovative Network 8 Fund (PHYTOOPTICS project).

### 2 Tables

3 Table 1. Mean (minimum – maximum) values of physical parameters (sea surface temperature (SST), sea surface salinity (SSS), and wind

4 speed), surface biomass proxies (TChl a-H: TChl a from HPLC measurements, TChl a-F: TChl a determined from the continuously

- 5 measuring fluorescence sensor), and sea surface concentrations, as well as sea-to-air fluxes of the four halocarbons CHBr<sub>3</sub>, CH<sub>2</sub>Br<sub>2</sub>, CH<sub>3</sub>I, and
- 6 CH<sub>2</sub>I<sub>2</sub> during the cruise MSM18/3.

Para- meter	SST	SSS	Wind speed	Bion pro.	nass xies	Halocarbons								
						СН	Br <sub>3</sub>	CH	$_2Br_2$	CH	I <sub>3</sub> I	СН	$I_2I_2$	
				TChl <i>a</i> -H	TChl <i>a</i> -F	Con- centrations	Sea-to-air fluxes	Con- centrations	Sea-to-air fluxes	Con- centrations	Sea-to-air fluxes	Con- centrations	Sea-to-air fluxes	
Unit	[° C]		[m s <sup>-1</sup> ]	[µg	L <sup>-1</sup> ]	[pmol L <sup>-1</sup> ]	$[pmol m^{-2} h^{-1}]$	[pmol L <sup>-1</sup> ]	$[pmol m^{-2} h^{-1}]$	[pmol L <sup>-1</sup> ]	$[pmol m^{-2} h^{-1}]$	[pmol L <sup>-1</sup> ]	$[pmol m^{-2} \\ h^{-1}]$	
Mean	24.4	35.7	6.1	0.51	0.44	12.9	644	3.7	187	5.5	425	1.1	82	
Min	22.1	34.5	0.3	0.10	0.06	1.8	-146	0.9	-3	1.5	34	0.3	3	
Max	29.0	36.3	11.1	0.99	1.20	44.7	4285	9.2	762	12.8	1300	3.7	382	

	CHBr <sub>3</sub>	CH <sub>2</sub> Br <sub>2</sub>	CH <sub>3</sub> I	CH <sub>2</sub> I <sub>2</sub>	SST	Salinity	Global radiation	Latitude	Wind speed	Chlorophyll a + Div a	Chlorophytes	Chrysophytes	Dinoflagellates	Haptophytes
Prochlorococcus (HL)	-0.70	-0.57	-0.21	0.27	0.44	-0.39	-0.20	0.49	0.26	-0.01	0.34	-0.28	-0.14	-0.33
Haptophytes	0.34	0.37	0.39	-0.25	-0.58	0.34	0.16	-0.21	-0.34	0.57	-0.18	0.37	0.53	
Dinoflagellates	0.22	0.22	0.29	-0.02	-0.50	0.10	-0.14	-0.33	-0.37	0.72	0.09	0.40		
Chrysophytes	0.43	0.41	0.26	0.13	-0.45	0.48	-0.28	-0.15	-0.15	0.71	0.22			
Chlorophytes	-0.29	-0.26	-0.15	0.32	0.13	-0.15	-0.26	0.25	-0.05	0.11				
TChl a	0.23	0.27	0.36	0.04	-0.58	0.35	-0.22	-0.13	-0.27					
Wind speed	-0.18	-0.16	-0.22	0.20	0.56	-0.06	0.12	0.04						
Latitude	-0.38	-0.18	0.03	0.12	0.10	-0.20	-0.08							
Global radiation	0.05	0.04	-0.09	-0.25	0.19	-0.09								

1 Table 2. Spearman's rank correlation coefficients  $r_s$  of halocarbons with different physical parameters and phytoplankton species measured in

surface water. Numbers printed in bold are regarded as significant with p < 0.05.

SSS	0.48	0.41	-0.09	-0.04	-0.42
SST	-0.46	-0.46	-0.42	0.33	
$CH_2I_2$	0.07	0.09	-0.04		
CH <sub>3</sub> I	0.50	0.62			
CH <sub>2</sub> Br <sub>2</sub>	0.90				

2 Table 3. Concentrations of CHBr<sub>3</sub>, CH<sub>2</sub>Br<sub>2</sub> and TChl *a* (from HPLC measurements) averaged over different depths at every CTD station (1 –

3 13), as well as the mixed layer depth. If a range is not given, only one measurement point exists. Bold numbers indicate the depth of

4 maximum concentrations at this station.

		0 – 30 m			31 – 60 m	1	61 – 100 m			
<i>z<sub>ML</sub></i> [ <b>m</b> ]	Concent [pmo]	trations   L <sup>-1</sup> ]	TChl <i>a</i> [µg L <sup>-1</sup> ]	Concen [pmo	trations l L <sup>-1</sup> ]	TChl <i>a</i> [µg L <sup>-1</sup> ]	Concer [pmo	$trations$ of $L^{1}$ ]	TChl <i>a</i> [µg L <sup>-1</sup> ]	
_	CHBr <sub>3</sub>	CH <sub>2</sub> Br <sub>2</sub>		CHBr <sub>3</sub>	CH <sub>2</sub> Br <sub>2</sub>		CHBr <sub>3</sub>	CH <sub>2</sub> Br <sub>2</sub>		

1	34	5.4 (3.2 - 6.5)	1.7 (1.3 - 2.1)	0.60 (0.52 - 0.69)	5.8 (3.7 - 7.9)	3.0 (1.8 - 4.2)	0.59 (0.53 - 0.65)	2.1	1.1	
2	16	30.2 (25.4 - 35.0)	6.5 (6.4 - 6.6)	0.92 (0.76 - 1.07)	9.0 (7.6 - 10.3)	5.2 (5.1 - 5.4)	0.86 (0.74 - 0.97)	2.4 (1.2 - 4.6)	1.8 (0.8 - 3.6)	0.20 (0.10 - 0.30)
3	37	6.8 (6.2 - 7.4)	3.9 (3.6 - 4.2)	0.80 (0.75 - 0.86)	3.0 (2.6 - 3.2)	2.4 (2.4 - 2.5)	0.65 (0.51 - 0.80)	2.3 (2.2 - 2.5)	2.3 (2.3 - 2.3)	0.18
4	14	12.5 (5.8 - 19.2)	7.2 (3.8 - 10.6)	0.56 (0.26 - 0.86)	5.9 (4.8 - 6.9)	3.1 (3.0 - 3.2)	0.80 (0.79 - 0.81)	2.6 (2.0 - 3.2)	2.5 (1.8 - 3.2)	0.19 (0.13 - 0.26)
5	49	14.0 (13.6 - 14.4)	4.2 (4.0 - 4.3)	0.34 (0.28 - 0.39)	11.7	4.8	0.58	7.6 (6.6 - 8.5)	7.4 (6.1 - 8.6)	0.39 (0.24 - 0.53)
6	12	13.4 (12.5 - 14.3)	5.0 (3.8 - 6.3)	0.99	5.4 (5.1 - 5.7)	4.8 (4.7 - 4.8)	0.30 (0.17 - 0.43)	4.9 (4.7 - 5.1)	4.6 (4.6 - 4.7)	0.10 (0.04 - 0.17)
7		11.2 (8.8 - 13.7)	4.6 (3.5 - 4.6)	0.71 (0.65 - 0.76)	3.7 (2.5 - 4.9)	3.4 (2.5 - 4.2)	0.46 (0.44 - 0.48)	3.1 (2.9 - 3.4)	3.0 (2.9 - 3.1)	0.11 (0.06 - 0.17)
8	45	5.0 (4.7 - 5.3)	1.0 (0.6 - 1.4)	0.34 (0.31 - 0.38)	7.0 (5.7 - 8.3)	2.5 (1.9 - 3.2)	0.51 (0.47 - 0.58)	1.1	1.5	0.51
9	21	3.6 (2.7 - 4.5)	1.8 (1.6 - 2.0)	0.75 (0.64 - 0.85)	8.9 (7.4 - 10.3)	4.2 (3.9 - 4.6)	0.77 (0.68 - 0.85)	5.4 (4.5 - 6.3)	3.2 (2.6 - 3.7)	0.24 (0.17 - 0.32)

10	10	5.2 (4.9 - 5.5)	2.6 (2.3 - 2.8)	0.50 (0.41 - 0.59)	8.9 (8.3 - 9.5)	3.8 (3.7 - 4.0)	0.62 (0.51 - 0.73)	3.5 (3.1 - 3.9)	2.5 (2.4 - 2.6)	0.47 (0.32 - 0.62)
11	24	6.0 (4.1 - 7.9)	2.5 (1.8 - 3.3)	0.46 (0.42 - 0.49)	13.1	4.3	0.82	4.0 (2.5 - 6.8)	4.0 (2.8 - 6.0)	0.23 (0.04 - 0.44)
12	35	18.1 (16.4 - 19.8)	5.8 (5.6 - 6.1)	0.77 (0.76 - 0.79)	11.6 (9.1 - 14.1)	6.3 (5.4 - 7.1)	0.70 (0.68 - 0.72)	5.3 (4.7 - 6.0)	5.5 (5.3 - 5.8)	0.25
13	41	11.6 (6.9 - 16.4)	3.5 (2.5 - 4.4)	0.55 (0.51 - 0.58	8.9 (8.3 - 9.5)	4.6 (3.0 - 5.6)	0.16 (0 - 0.48)	5.9 (3.3 - 7.6)	5.2 (4.1 - 5.7)	0.12 (0 - 0.30)

Table 4. Concentrations of  $CH_3I$ ,  $CH_2I_2$  and the sum of TChl *a* averaged over different depths at every CTD station (1 - 13), as well as the mixed layer depth. If a range is not given, only one measurement point exists. Bold numbers indicate the depth of maximum concentrations at this station.

		0 – 30 m			30 – 60 m	I	60 – 100 m			
 <i>z<sub>ML</sub></i> [ <b>m</b> ]	Concen [pmo	trations l L <sup>-1</sup> ]	TChl <i>a</i> [µg L <sup>-1</sup> ]	Concentrations [pmol L <sup>-1</sup> ]		TChl <i>a</i> [µg L <sup>-1</sup> ]	Concentrations [pmol L <sup>-1</sup> ]		TChl <i>a</i> [µg L <sup>-1</sup> ]	
	CH <sub>3</sub> I	$CH_2I_2$		CH <sub>3</sub> I	$CH_2I_2$		CH <sub>3</sub> I	CH <sub>2</sub> I <sub>2</sub>		

1	34	2.7 (2.1 - 3.4)	4.5 (1.2 - 6.8)	0.60 (0.52 - 0.69)	2.5 (1.8 - 3.2)	9.9 (3.9 - 16.0)	0.59 (0.53 - 0.65)	0.2	1.7	
2	16	2.8 (0.4 - 5.2)	4.8 (1.7 - 8.0)	0.92 (0.76 - 1.07)	3.1 (2.7 - 3.6)	12.2 (11.5 - 12.9)	0.86 (0.74 - 0.97)	0.6 (0.1 - 1.3)	2.0 (0.7 - 4.3)	0.20 (0.10 - 0.30)
3	37	8.5 (8.4 - 8.5)	4.1 (1.7 - 6.4)	0.80 (0.75 - 0.86)	2.6 (1.0 - 3.5)	4.6 (4.3 - 4.9)	0.65 (0.51 - 0.80)	0.7 (0.4 - 1.1)	3.3 (2.3 - 4.4)	0.18
4	14	6.1 (5.5 - 6.6)	7.0	0.56 (0.26 - 0.86)	4.6 (4.6 - 4.7)	2.3 (2.2 - 2.4)	0.80 (0.79 - 0.81)	0.8 (0.7 - 0.9)	1.0 (0.7 - 1.3)	0.19 (0.13 - 0.26)
5	49	5.4	0.6 (0.5 - 0.7)	0.34 (0.28 - 0.39)	4.5	4.9	0.58	2.4 (1.9 - 3.0)	10.5 (7.1 - 13.8)	0.39 (0.24 - 0.53)
6	12	10.4 (8.0 - 12.8)	6.9 (1.8 - 12.0)	0.99	1.6 (1.5 - 1.7)	4.0 (3.1 - 4.8)	0.30 (0.17 - 0.43)	1.4 (1.0 - 1.7)	2.4 (1.7 - 3.1)	0.10 (0.04 - 0.17)
7		4.1 (3.4 - 4.8)	2.3 (1.2 - 3.4)	0.71 (0.65 - 0.76)	1.3 (1.2 - 1.3)	4.7 (3.3 - 6.1)	0.46 (0.44 - 0.48)	0.9 (0.6 - 1.2)	2.0 (1.5 - 2.7)	0.11 (0.06 - 0.17)
8	45	0.2 (0.1 - 0.4)	0.3 (0.3 - 0.3)	0.34 (0.31 - 0.38)	4.7 (3.0 - 7.0)	1.2 (0.5 - 1.9)	0.51 (0.47 - 0.58)	0.0	2.4	0.51
9	21	4.4 (4.1 - 4.8)	1.3 (1.2 - 1.5)	0.75 (0.64 - 0.85)	5.3 (3.4 - 7.3)	6.2 (4.5 - 8.0)	0.77 (0.68 - 0.85)	1.3 (1.3 - 1.3)	2.9 (2.3 - 3.6)	0.24 (0.17 - 0.32)

10	10	4.5 (3.6 - 5.5)	0.5 (0.4 - 0.6)	0.50 (0.41 - 0.59)	4.9 (4.2 - 5.7)	1.3 (0.9 - 1.7)	0.62 (0.51 - 0.73)	0.8 (0.7 - 0.9)	3.4 (2.6 - 4.1)	0.47 (0.32 - 0.62)
11	24	3.8 (2.9 - 4.6)	0.4	0.46 (0.42 - 0.49)	4.4	2.3	0.82	1.7 (1.0 - 2.3)	1.7 (0.6 - 3.2)	0.23 (0.04 - 0.44)
12	35	7.0 (6.8 - 7.1)	1.2 (0.3 - 2.2)	0.77 (0.76 - 0.79)	2.7	4.1 (3.8 - 4.3)	0.70 (0.68 - 0.72)	2.0	2.7 (1.6 - 3.8)	0.25
13	41	5.1 (4.3 - 5.9)	1.5 (0.8 - 2.1)	0.55 (0.51 - 0.58	3.8 (2.0 - 5.6)	5.9 (3.9 - 7.4)	0.16 (0 - 0.48)	1.0 (0.1 - 2.0)	3.4 (1.0 - 4.8)	0.12 (0 - 0.30)

Table 5. Diapycnal and sea-to-air fluxes at every CTD station for the four halocarbons. Positive fluxes in bold provide the mixed layer with
the corresponding halocarbon, while negative fluxes indicate losses from the mixed layer.

CTD station	CHBr <sub>3</sub>		CH	Br <sub>2</sub>	CH	I <sub>3</sub> I	$CH_2I_2$		
	Diapycnal flux	Sea-to-air flux							
	$[pmol m^{-2} h^{-1}]$								
1	14	14	8	-27	5	-119	39	-64	

2	-125	-3651	-8	-689	-13	-44	29	-199
3	0	-184	1	-195	-6	-703	7	-129
4	8	-241	4	-265	-1	-671	3	
5	-3	-893	4	-275	-2		9	-45
6	5	-590	7	-185	-13	-988	27	-121
7								
8	-2	-110	-0	-25	-1	-4	0	-22
9	3	-57	1	-64	1	-337	3	-88
10	2	-45	-2	-83	-6	-300	-1	-30
11	4	-248	1	-136	1	-316	0	-24
12	-4	-1208	-1	-357	-2	-583	-0	-20
13	1	-837	0	-231	-3	-446	-4	-54

Table 6. Total mixed layer budget of each halocarbon, potential sinks and sources (box size  $1 \times 1 \times z_{ML} \text{ m}^3$ ). The upper four rows indicate cases where diapycnal fluxes act as sources, while the lower four rows summarize the budget for the cases where the diapycnal fluxes were sinks for the mixed layer budget. "Other sinks" is halogen substitution for CH<sub>3</sub>I and photolysis in case of CH<sub>2</sub>I<sub>2</sub>. The negative numbers indicate sinks for the budget.

	Compound	<b>z</b> <sub>ML</sub>	Total ML budget	Air-sea fluxes (S <sub>as</sub> )	Diapycnal fluxes (F <sub>dia</sub> )	Other sinks (S <sub>ch</sub> )	Total after 1 h	Difference
Unit		[m]	[pmol]	[pmol h <sup>-1</sup> ]	[pmol h <sup>-1</sup> ]	[pmol h <sup>-1</sup> ]	[pmol]	[pmol]
Diapycnal fluxes as source	CHBr <sub>3</sub>	24	157543	-274	5		157274	-269
	$CH_2Br_2$	29	90058	-172	3		89889	-169
	CH <sub>3</sub> I	26	75263	-257	2	0	75004	-255
	$CH_2I_2$	28	63947	-78	13	-8317	55565	-8382
Diapycnal fluxes as sink	CHBr <sub>3</sub>	36	417098	-1186	-30		415882	-1216
	$CH_2Br_2$	27	99604	-236	-2		99366	-238
	CH <sub>3</sub> I	29	137560	-420	-5	0	137135	-425

CH212 29 100387 -33 -2 -4977 101373 -3014		$CH_2I_2$	29	106587	-35	-2	-4977	101573	-5014
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2 Table 7. Theoretical mean production rate of the four halocarbons in the equatorial mixed layer with the standard deviation.

Compound	Compound Production rate		Production rate per TChl <i>a</i>	
	$[pmol m^{-3} h^{-1}]$	$[pmol m^{-3} h^{-1}]$	$[\text{pmol} [\mu \text{g TChl } a]^{-1} \text{h}^{-1}$	
CHBr <sub>3</sub>	34	65	2.5 x 10 <sup>-3</sup>	
CH <sub>2</sub> Br <sub>2</sub>	10	12	8.5 x 10 <sup>-4</sup>	
CH <sub>3</sub> I	21	24	2.2 x 10 <sup>-3</sup>	
$CH_2I_2$	384	318	3.3 x 10 <sup>-2</sup>	

## 1 Figures



Figure 1. Cruise track with SST in °C (small box) and the section (large box) were halocarbons were sampled in both the sea surface and during CTD stations (numbered circles), plotted on monthly average Chl *a* for July 2011 derived from mapped level 3 MODIS Aqua Data.



Figure 2. a) Species composition (HL – high light, LL – low light), b) SST and salinity during
the cruise, c) TChl *a* from underway fluorescence sensor measurements and global radiation,
e) CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub> in surface sea water, and e) CH<sub>3</sub>I and CH<sub>2</sub>I<sub>2</sub> surface sea water
concentrations. The top numbers mark the CTD stations.



Figure 3. Selected CTD profiles (top – down: profiles 7, 9 and 10, see Figure 1 for the location) of CHBr<sub>3</sub>, CH<sub>2</sub>Br<sub>2</sub>, CH<sub>3</sub>I, and CH<sub>2</sub>I<sub>2</sub> in a – b), e – f), and i – j), along with temperature, salinity, and density (c, g and k), as well as TChl *a* in d), h), and l), and the mixed layer depth as black dashed line at the same stations.

2



1

Figure 4. a – d) Temperature-Salinity (T-S) plots for halocarbons (in pmol L<sup>-1</sup>) and e – f) phytoplankton species (in  $\mu$ g Chl *a* L<sup>-1</sup>). Square markers indicate surface values of halocarbons from underway measurements, circles are depth measurements from CTD profile, and the lines indicate the potential density – 1000.



Figure 5. Principal component analysis (PCA) of halocarbon and phytoplankton species
composition data, as well as temperature, salinity, and density for the 13 CTD stations during
MSM18/3.





Figure 6. Wind speed during the cruise and sea-to-air fluxes calculated with sea surface water
concentrations and mean atmospheric halocarbon data a) CHBr<sub>3</sub> and CH<sub>2</sub>Br<sub>2</sub> and b) CH<sub>3</sub>I and

4 CH<sub>2</sub>I<sub>2</sub>. Numbers on the top indicate CTD stations.

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