The contribution of oceanic halocarbons to marine and 1 free troposphere air over the tropical West Pacific 2 3 S. Fuhlbrügge¹, B. Quack¹, S. Tegtmeier¹, E. Atlas², H. Hepach¹, Q. Shi³, S. Raimund¹, 4 K. Krüger⁴ 5 [1] GEOMAR | Helmholtz Centre for Ocean Research Kiel 6 [2] Rosenstiel School for Marine and Atmospheric Sciences, Miami, Florida 7 [3] Department of Oceanography - Dalhousie University, Halifax, Canada 8 [4] University of Oslo, Oslo, Norway 9 Correspondence to: K. Krüger (kirstin.krueger@geo.uio.no) 10 11 12 **Response to Editor** 13 We would like to thank the Editor for his suggestions and efforts to clarify the manuscript. 14 Below you find our answers to your specific points (*italic*). 15 16 1. Editor: 17 Dear authors, 18 Thanks for your explanations. They have made clear, where I think the problem lies. 19 The budget in the MABL is well documented. I would appreciate to indeed use your new 20 figure B with the Daily contributions in %/d indicated on the figure. Although confusing the 21 ODR etc terms are probably a matter of style and could stay. 22 23 Authors: 24 We thank the editor that he agreed to keep the ODR terms. According to the editors 25 suggestion we will exchange Figure 9 in the manuscript with Figure B from the former reply 26 to the editor report:

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Figure B: Average budgets of the Oceanic Delivery (OD, blue), Chemical Loss (CL, red),
Advective Delivery (AD, green) and Convective Loss (COL, orange) of CHBr₃, CH₂Br₂ and
CH₃I in the Marine Atmospheric Boundary Layer (MABL).

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- 35 2. *Editor*:

36 What I think is not resolved is how you arrive at 60% oceanic contribution to FT bromoform

mixing ratios, starting from 45% in the MABL. You do not explain how you get to that number.

Bromoform: 60% oceanic contribution to FT air cannot be kept as a conclusion, to my

40 *opinion.* 45% as in the MABL air is an upper limit for the FT air. That should be explained.

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42 Authors:

We agree with the reviewer that this part is confusing. Although the results in Figure 11 will change now (see your comment no. **3**), we like to clarify how this occurred:

45 In Section 5.3.2 we describe the results of Figure 11 as: "FT mixing ratios from VMR_{ODR} are

- 46 on average 0.3 ppt lower than observations from R/A FALCON and thus explain about 60 %
- 47 of the observed FT mixing ratios."



Figure 11 from the manuscript: Mean FT mixing ratios (solid lines) and 1 standard deviation (shaded areas) from in-situ and flask observations on R/A FALCON (Obsv., black) versus simulated mean FT mixing ratios from MABL air (MABL, red) and oceanic emissions (Ocean, blue) observed by R/V SONNE. R/A FALCON in-situ observations have been adjusted for CH₃I (Obsv.*, dashed black) according to measurement deviations during the meetings of R/V SONNE and R/A FALCON (Table 2; Section 4.3).

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Free troposphere mixing ratios in Figure 11 are based on observations by R/A FALCON 50 (black line) and on *simulations* (red and blue line) using Flexpart trajectories and MABL 51 VMRs as input. Starting with 45 % in the MABL for bromoform, the mean ratio of simulated 52 free troposphere mixing ratios from the oceanic emissions (blue line) and from MABL 53 observations (red line) is 45 % as well. The 60 % describe the mean ratio between *simulated* 54 free troposphere mixing ratios from oceanic emissions (blue line) and observations by R/A 55 FALCON (black line). Since for bromoform the observed mixing ratios (black line) are on 56 average lower than the simulated mixing ratios from observed MABL air (red line), the ratio 57 between Ocean (blue line) and Observation (black line) has to be higher than 45 %. 58

59 In a short way (for bromoform, Figure 11):

60 Ratio Ocean / MABL = 45 %

- 61 Ratio **Ocean** / **Observation** = 60 %
- 62

63 Since the results in the new Figure 11 changed with the revised method (Editor comment 3),

64 we suggest removing the 60 % and the idea behind it in the abstract and the conclusion and

replace it by the 45 %, resulting from the ODR calculations and our new results:

66 Abstract:

- 67 "Up to 45 % of the accumulated bromoform in the FT above the region originates from the
- local South China Sea area, while dibromomethane is largely advected from distant source
- regions and the local ocean only contributes 20 %."
- 70 Section 5.3.2, removing the sentence (new since reply to editor report 2):
- "FT mixing ratios from VMR_{ODR} are on average 0.3 ppt lower than observations from R/A
- 72 FALCON and thus explain about 60 % of the observed FT mixing ratios."
- 73 Discussion, 1^{st} sentence, change to:

"Oceanic emissions of CHBr3 from the South China Sea contribute on average 45 % to the
simulated FT mixing ratios (Figure 11). Simulated FT mixing ratios from MABL

- observations and observed FT mixing ratios agree quite well up to 10 km height."
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- 78
- 79 *3. Editor:*

After several reads I come to the conclusion that there is serious flaw in your vertical mixing
computation method.

Air with any given mixing ratio at MABL level can not be transformed to air with a higher 82 mixing ratio when mixed up into the free troposphere. FT-air, which has supposedly smaller 83 background mixing ratios, will dilute more and more. Yet, when looking at figure 11 the 84 mixing ratios for bromoform increase towards the top of the troposphere, relative to the mid 85 level altitudes (actually seen for all three compounds). How can this "accumulation" 86 happen? I think this is a numerical artifact of the rather unorthodox method to compute the 87 vertical mixing, as described in section 2.4.3, esp equation 13. Using a trajectory model and 88 a pressure profile. I think this is an oversimplified paternoster model of vertical mixing. As if 89 MABL air is injected directly, eg into a layer at 12 km height, expands and then the clean FT 90 air is, in compensation, pressed down. There is no entrainment, no turbulent mixing assumed, 91 which in reality will dilute MABL air considerably during transport. 92

- By the way: The observations don't show this enrichment at the top of the troposphere, no surprise for a substance with a surface source and sink in the FT.
- 95 Please expand on the proper value of your vertical mixing calculations. And its caveats. I
- consider this a major revision, since I don't believe this can be done in 1-2 sentences.

97 Authors:

After intensively re-checking our method again, we agree with the Editor that Eq. 13 in our
method has a conceptual flaw. Thus we have revised the method and Eq. 13 to simulate the
VMRs in the free troposphere.

In the manuscript the VMR of a compound in a specific height was computed after Eq. 13, with r_{MABLi} as the amount of MABL air relative to the total available air volume at a specific height (e.g. 0.5 - 1.0 km height) at run *i* and VMR_{MABLi} as the specific VMR including chemical decay since observed in the MABL:

$$VMR_{FT} = r_{MABL_1} \cdot VMR_{MABL_1} + r_{MABL_2} \cdot VMR_{MABL_2} + \dots + r_{MABL_i} \cdot VMR_{MABL_i}$$
(Eq. 13)

In our revised method the VMR in the free troposphere at height z are now computed after the following equation 13A:

$$VMR_{FT}(z) = \sum_{i=1}^{n} \begin{pmatrix} r_{MABL}(t(i), z) \cdot VMR_{MABL}(CL(t(i))) \\ + \\ (1 - r_{MABL}(t(i), z)) \cdot VMR_{FT}(t(i-1), z) \cdot CL(COL) \end{pmatrix}$$
(Eq. 13A)

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109 Here, *n* is the number of runs for each of the 80 profiles, according to the residence time of the trajectories in the MABL and the total runtime of the trajectories. For example a 110 trajectory residence time of 7 hours in the MABL in combination with a total trajectory 111 runtime of 16 days leads to n = (24 / 7) * 16 = 54. r_{MABL} gives the volume ratio of MABL air 112 parcels at time t and height z to the total volume of a specific height layer. VMR_{MABL} gives 113 the compounds mixing ratio in the air parcels including chemical degradation (CL) since the 114 air was observed in the MABL. Since we use a mean tropospheric lifetime, CL and thus also 115 VMR_{MABL}(CL) are independent from the height z. For i = 0 we set VMR_{FT} = 0 to start with 0 116 ppt background in the free troposphere, followed by *n* times iteration of $VMR_{FT}(i-1)$. 117

In contrast to the old formula where all air masses at a specific height were mixed at once, now the trajectories contribute during each step *i* to the previously computed VMR_{FT} at step i - 1.

The results from Eq. 13A are given in Figure 11-revised below. Please note that there is more variability in the observed VMR profiles now as we found a small code bug in calculating the mean and standard deviation of FALCON measurements.



Figure 11-revised: VMR_{FTS} computed after Eq. 13A, assuming the trajectories and meteorology during the time of the cruise to be representative for the region.

We agree with the Editor that our method is a simplified model of vertical mixing since we 125 have no entrainment of different air masses or turbulent mixing of the air parcels along the 126 trajectory. We also agree with the Editor that air with any given mixing ratio cannot be 127 128 transformed to air with a higher mixing ratio when mixed up with air that has supposedly smaller background mixing ratios. This is true for our method (old and revised) as well. All 129 computed mean free tropospheric mixing ratios are lower compared to the initial MABL 130 mixing ratio of each compound. An increase of the mixing ratios above 10 km height relative 131 to the lower free troposphere has to be ascribed to the convective outflow. Indeed this 132 convective outflow was observed during several flights of R/A FALCON along the northern 133 coast of Borneo (Sala et al., 2014), which is shown for methyl iodide exemplarily (Figure 1). 134



Figure 1: Two exemplary observations for convective outflow observed by R/A FALCON during SHIVA campaign and also modelled by a global CH_3I approach with FLEXPART. Data taken from Tegtmeier et al. (2013).

We agree with the Editor that the convective outflow is quite pronounced in our FLEXPART 135 ERA-Interim simulations, since it is not reflected in the mean of the aircraft observations as 136 well. However, this is not caused by our method to compute the mixing ratios, in particular 137 Eq. 13/13A, but due to the meteorology (with active ongoing convection) during the 138 trajectory releases used in our approach and above the South China Sea (SCS). Since a high 139 percentage of the trajectories reveal this convective outflow (Figure 10 from manuscript) and 140 the simulated entrainment of air in the upper troposphere is given by the same trajectories 141 using the same meteorology, the intensity of the convective outflow in our method is 142 overestimated compared to the aircraft measurements. However, we are able in particular 143 with Eq.13A to simulate observed free troposphere mixing ratios up 10 km (bromoform) and 144 8 km (dibromomethane) in the area from MABL observations with a simplified approach. 145

To better address the simplicity and caveats of our method we add a more detailed discussion
of it in Section 2.4.3 and Section 5.3.3, particular emphasizing the lack of turbulent mixing,
entrainment, and steady meteorology.

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151 *4. Editor:*

Figure 8 shows that 80% of trajectories left the box in 10 days. At the same time the authors state that the steady state of the vmr vertical profile is reached after ca 10 days. I suspect that the profiles reach steady state because after 10 days most of the trajectories left your bounding box. How long are the trajectories actually run? Can the trajectories go higher than 13km? What happens to air if carried out of the Fig 8 box? What would happen to the profiles if the Fig 8 box was twice as big, or spanning the whole northern hemisphere?

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159 Authors:

The editor is right. Indeed the size of the box influences the VMR_{FT}s of the compounds as 160 well as the time until a steady state of the VMR_{FT}s is reached. The effect of the box size in 161 the old method (Eq. 13) and the revised one (Eq. 13A) as well as higher altitude calculations 162 are given in the supplement figures below (Fig. S1-S4). The simulation of free troposphere 163 entrainment is in a simple way controlled by the size of the box (e.g. SCS). A smaller box 164 leads to less entrainment, as trajectories are not considered anymore once they leave the box, 165 while a larger box leads to increased entrainment, as the trajectories are longer considered. 166 This leads to an enhancement of the convective outflow, since the entrained air shows a 167

168 strong convective outflow as well (due to the same meteorology in the method). Thus the 169 reliability of the computed profiles decreases with altitude which has to be mentioned.

Using Eq. 13 without a box, a steady state couldn't be reached within the runtime of the trajectories of 16 days. In contrast, using Eq. 13A the difference of VMR_{FT} between the last two steps of the iteration (i = n-1 and i = n) is less than 1 % within the 16 days runtime.

Thus we suggest removing the box approach from the manuscript since it is an adjustment of the entraining air in the free troposphere that can only be guessed in our simulation. Instead we suggest presenting Figure 11-revised and explain the pronounced convective outflow in the simulation and discuss the assumptions of our method as described above. Removing the box concept would also contribute to the readability of the manuscript.

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179 *5. Editor:*

180 As written before, I believe the material is very worth publishing,

Thanks again for your positive evaluation and for your efforts to improve the paper. In case the Editor agrees with our revised method and suggestions, we will update the abstract, results and discussion in the manuscript with the above given changes and present the changed manuscript to the Editor once more.

185

Supplement Figures



Figure S-1: VMR_{FT}s computed after old method (Eq. 13) up to 22 km height with horizontal box (South China Sea).



Figure S-2: VMR_{FT}s computed after old method (Eq. 13) up to 22 km height without horizontal box, assuming the trajectories and meteorology to be representative for the entire region.



Figure S-3: VMR_{FT}s computed after new method (Eq. 13A) up to 22 km height with horizontal box (SCS). *Note: FALCON observations have been corrected*.



Figure S-4: VMR_{FT}s computed after new method (Eq. 13A) up to 22 km height without horizontal box, assuming the trajectories and meteorology to be representative for the entire region. *Note: FALCON observations have been corrected*.

193 **References**

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- 204

205 Abstract

Emissions of halogenated very short lived substances (VSLS) from the oceans contribute to 206 the atmospheric halogen budget and affect tropospheric and stratospheric ozone. Here we 207 investigate the contribution of natural oceanic VSLS emissions to the Marine Atmospheric 208 Boundary Layer (MABL) and their transport into the Free Troposphere (FT) over the tropical 209 West Pacific. The study concentrates on bromoform, dibromomethane and methyl iodide 210 measured on ship and air craft during the SHIVA (Stratospheric Ozone: Halogen Impacts in a 211 Varying Atmosphere) campaign in the South China and Sulu Seas in November 2011. 212 Elevated oceanic concentrations for bromoform, dibromomethane and methyl iodide of on 213 average 19.9, 5.0 and 3.8 pmol L⁻¹ in particular close to Singapore and at the coast of Borneo 214 with high corresponding oceanic emissions of 1486, 405 and 433 pmol $m^{-2} h^{-1}$, respectively, 215 characterize this tropical region as a strong source of these compounds. Atmospheric mixing 216 ratios in the MABL were unexpectedly relatively low with 2.08, 1.17 and 0.39 ppt for 217 bromoform, dibromomethane and methyl iodide. We use meteorological and chemical ship 218 and aircraft observations, FLEXPART trajectory calculations and source-loss estimates to 219 identify the oceanic VSLS contribution to the MABL and to the FT. Our results show that the 220 well-ventilated MABL and intense convection led to the low atmospheric mixing ratios in the 221 222 MABL despite the high oceanic emissions. Up to 45 % of the accumulated bromoform in the FT above the region originates from the local South China Sea area, while dibromomethane 223 224 is largely advected from distant source regions and the local ocean only contributes 20 %.Most of the accumulated bromoform in the FT above the region originates from the local 225 South China Sea area (up to 60 %), while dibromomethane is largely advected from distant 226 source regions. The accumulated methyl iodide in the FT is higher than can be explained with 227 local contributions. Possible reasons, uncertainties and consequences of our observations and 228 model estimates are discussed. 229

230

231 **1. Introduction**

Halogens play an important role for atmospheric chemical processes. Chlorine, bromine and iodine radicals destroy ozone in the stratosphere (e.g. Solomon, 1999) and also affect tropospheric chemistry (e.g. Saiz-Lopez and von Glasow, 2012). Halogens are released following the photochemical breakdown of organic anthropogenic and natural trace gases. A large number of very short lived brominated and iodinated organic substances, originating from macro algae, seaweed, phytoplankton and other marine biota, are emitted from tropical oceans and coastal regions to the atmosphere (Gschwend et al., 1985;Carpenter and Liss,

2000; Quack and Wallace, 2003; Quack et al., 2007; Liu et al., 2013). In particular, marine 239 emissions of bromoform (CHBr₃), dibromomethane (CH₂Br₂) and methyl iodide (CH₃I) are 240 major contributors of bromine and iodine to the atmosphere (Montzka and Reimann, 2011). 241 Annually averaged mean tropical lifetimes of these halogenated very short-lived substances 242 (VSLS) in the boundary layer are 15 (range: 13 - 17) days for CHBr₃, 94 (84 - 114) days for 243 CH_2Br_2 and 4 (3.8 – 4.3) days for CH_3I . The mean tropospheric lifetimes of these compounds 244 at 10 km height are 17 (16 - 18) days, 150 (144 - 155) days, respectively 3.5 (3.4 - 3.6) days 245 (Carpenter et al., 2014). Climate change could strongly affect marine biota and thereby 246 247 halogen sources and the oceanic emission strength (Hughes et al., 2012;Leedham et al., 2013;Hepach et al., 2014). 248

Aircraft measurements from Dix et al. (2013) suggest that the halogen-driven ozone loss in 249 the Free Troposphere (FT) is currently underestimated. In particular, elevated amounts of the 250 iodine oxide free radical (IO) in the FT over the Central Pacific indicate that iodine may have 251 a larger effect on the FT ozone budget than currently estimated by chemical models. 252 Coinciding with this study, Tegtmeier et al. (2013) projected a higher CH₃I delivery to the 253 Upper Troposphere / Lower Stratosphere (UTLS) over the tropical West Pacific than 254 previously reported, using an observation based emission climatology by Ziska et al. (2013). 255 256 Recent studies reported significant contributions of bromine and iodine to the total rate of tropospheric and stratospheric ozone loss (e.g. von Glasow et al., 2004; Yang et al., 257 2005;Saiz-Lopez et al., 2014;Yang et al., 2014;Hossaini et al., 2015). Deep tropical 258 convective events (Aschmann et al., 2011;Tegtmeier et al., 2013;Carpenter et al., 2014) as 259 well as tropical cyclones, i.e. typhoons (Tegtmeier et al., 2012), are projected to transport 260 VSLS rapidly from the ocean surface to the upper tropical tropopause layer. Despite the 261 importance of halogens on tropospheric and stratospheric ozone chemistry, halogen sources 262 and transport ways are still not fully understood. While the tropical West Pacific comprises 263 strong VSLS source regions (Krüger and Quack, 2013), only low mean atmospheric mixing 264 ratios were observed during ship campaigns in 1994 and 2009 (Yokouchi et al., 1997;Quack 265 and Suess, 1999) and in 2010 (Quack et al., 2011;Brinckmann et al., 2012). None of these 266 previous studies investigated the contribution of oceanic VSLS emissions to the marine 267 atmospheric boundary layer (MABL) and to the FT in this hot spot region with large oceanic 268 sources and strong convective activity. 269

The SHIVA ('Stratospheric Ozone: Halogen Impacts in a Varying Atmosphere') ship, aircraft and ground-based campaign during November and December 2011 in the Southern South China and Sulu Seas investigated oceanic emission strengths of marine VSLS, as well as their atmospheric transport and chemical transformation from the ocean surface to the upper
troposphere. For more details about the SHIVA campaign see the ACP special issue
(http://www.atmos-chem-phys.net/special_issue306.html).

In this study, we present campaign data from the research vessel (R/V) SONNE and the 276 research aircraft (R/A) FALCON. We identify the contribution of oceanic emissions to the 277 MABL and their exchange into the FT applying in-situ observations, trajectory calculations 278 and source-loss estimates. The results are crucial for a better process understanding and for 279 chemical transport model validation (Hossaini et al., 2013;Aschmann and Sinnhuber, 2013). 280 An overview of the data and the methods used in this study is given in Chapter 2. Chapter 3 281 provides results from the meteorological observations along the cruise. Chapter 4 compares 282 atmospheric VSLS measurements derived on R/V SONNE and R/A FALCON. The 283 contribution of the oceanic emissions to the MABL and FT air is investigated and discussed 284 in Chapter 5. Finally, a summary of the results is given in Chapter 6. 285

286

287 2. Data and Methods

288 **2.1. Ship and aircraft campaigns**

The R/V SONNE cruise started on November 15, 2011 in Singapore and ended on November 289 290 29, 2011 in Manila, Philippines (Figure 1). The ship crossed the southwestern South China Sea towards the northwestern coast of Borneo from November 16 – 19, 2011. From 291 November 19 - 23, 2011 the ship headed northeast along the northern coast of Borneo 292 towards the Sulu Sea. Two diurnal stations took place on November 18, 2011 at 2.4° N / 293 110.6° E and on November 22, 2011 at 6.0° N / 114.8° E. Two meetings between ship and 294 aircraft were carried out on November 19 and 21, 2011, where R/A FALCON passed R/V 295 SONNE within a distance of about 100 m several times to simultaneously measure the same 296 air masses. On November 24, 2011 the ship entered the Sulu Sea, and after 4 days transect, 297 R/V SONNE reached the Philippine coast. 298

16 measurement flights were carried out with R/A FALCON between November 16 and December 11, 2011 as part of the SHIVA campaign to investigate halogenated VSLS from the surface up to 13 km altitude over the South China and Sulu Seas. Observations were performed between 1° N and 8° N, as well as 100° E and 122° E, from Miri, Borneo (Malaysia) as the aircraft base. A detailed description of the VSLS measurements and flight tracks can be found in Sala et al. (2014).

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306 2.2. Meteorological observations during SHIVA

307 2.2.1. Measurements on board R/V SONNE

Meteorological parameters (temperature, air pressure, humidity and wind) were recorded at 308 20 m height every second. A 10 minute running mean of this data is used for this study. An 309 optical disdrometer ('ODM-470') measured the amount and intensity of precipitation during 310 the cruise at 15 m height every minute (see Supplementary Material for further details). To 311 obtain atmospheric profiles of air temperature, relative humidity and wind from the surface to 312 the stratosphere 67 GRAW DFM-09 and 6 GRAW DFM-97 radiosondes were launched 313 every 6 hours at standard UTC times (0, 6, 12, 18) from the working deck of R/V SONNE at 314 315 about 2 m above sea level. At the 24 h stations, the launch frequency was increased to 2-3hours to analyse short term diel variations of the atmospheric boundary layer. The radiosonde 316 data was integrated in near real time into the Global Telecommunication System (GTS) to 317 improve meteorological reanalyses such as ERA-Interim (Dee et al., 2011), which is used as 318 input data for the trajectory calculations (Section 5). 319

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321 **2.2.2. Marine atmospheric boundary layer**

322 The MABL is the atmospheric surface layer above the ocean in which trace gas emissions are mixed vertically by convection and turbulence on a short time scale of about an hour (Stull, 323 324 1988;Seibert et al., 2000). The upper boundary of the MABL is either indicated by a stable layer e.g. a temperature inversion or by a significant reduction in air moisture. Determination 325 of the MABL height can be achieved by theoretical approaches, e.g. using critical Bulk 326 Richardson number (Troen and Mahrt, 1986; Vogelezang and Holtslag, 1996; Sorensen, 1998) 327 or by practical approaches summarized in Seibert et al. (2000). An increase with height of the 328 virtual potential temperature, the temperature an air parcel would acquire if adiabatically 329 brought to standard surface pressure with regard to the humidity of the air parcel, identifies 330 the base of the stable layer, which is typically found between 100 m and 3 km altitude (Stull, 331 1988). In this study, we use the height of the base of the stable layer increased by half of the 332 stable layer depth as the definition for the MABL height. The height of the MABL is 333 determined from the atmospheric profiles measured by radiosondes launched on board the 334 ship, as described in detail by Fuhlbrügge et al. (2013). 335

336

337 2.3. VSLS measurements and flux calculation

338 VSLS in marine surface air and sea water were sampled synchronously on R/V SONNE 339 along the cruise track. From these data the oceanic emissions of the compounds during the

- 340 SHIVA campaign were calculated (Section 2.3.3). Additionally, VSLS were measured in the
- MABL and the FT by R/A FALCON (Sala et al., 2014;Tegtmeier et al., 2013).
- 342

343 2.3.1. Atmospheric samples

Air samples were taken 3 hourly along the cruise track, and 1 - 2 hourly during the 24 hour 344 stations on R/V SONNE resulting in a total of 195 samples during the cruise. The air was 345 pressurized to 2 atm in pre-cleaned stainless steel canisters with a metal bellows pump. The 346 samples were analyzed within 6 months after the cruise at the Rosenstiel School for Marine 347 and Atmospheric Sciences (RSMAS, Miami, Florida) according to Schauffler et al. (1999) 348 with an instrumental precision of ~5 %. Further details of the analysis are described in 349 Montzka et al. (2003) and Fuhlbrügge et al. (2013). On R/A FALCON ambient air was 350 analysed in situ by a GhOST-MS (Gas Chromatograph for the Observation of Stratospheric 351 Tracers – coupled with a Mass Spectrometer) by the Goethe University of Frankfurt (GUF). 352 Additionally 700 ml glass flasks were filled with ambient air to a pressure of 2.5 bar with the 353 R/A FALCON whole air sampler (WASP) and analysed within 48 hours by a ground-based 354 gas chromatography - mass spectrometry (GC/MS) instrument (Agilent 6973) of the 355 University of East Anglia (Worton et al., 2008). During the flights GhOST measurements 356 were conducted approximately every 5 minutes with a sampling time of 1 minute, while 357 WASP samples were taken every 3 - 15 minutes with a sampling time of 2 minutes. Further 358 details on the instrumental precision and intercalibration on R/A FALCON are given in Sala 359 et al. (2014). Given that the ground-based GC/MS investigated only brominated compounds, 360 CH₃I data is not available from WASP. Measurements from R/V SONNE and R/A FALCON 361 were both calibrated with NOAA standards. 362

363

364 **2.3.2. Water samples**

VSLS sea water samples were taken 3 hourly from the moon pool of R/V SONNE at a depth of 5 m from a continuously working water pump. Measurements were interrupted between November 16, 00 UTC to November 17, 2011 12 UTC due to permission issues in the southwest South China Sea. The water samples were analysed on board with a purge and trap system, attached to a gas chromatograph with mass spectrometric detection in single-ion mode and a precision of 10 % determined from duplicates. The method is described in detail by Hepach et al. (2014)

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373 **2.3.3. Sea – air flux**

The sea – air flux (*F*) of CHBr₃, CH₂Br₂ and CH₃I is calculated with k_w the concentration gradient, and Δc the concentration gradient between the water and atmospheric equilibrium concentrations (Eq. 1). For the determination of k_w , the wind speed-based parameterization of Nightingale et al. (2000) was used and a Schmidt number (*Sc*) correction to the carbon dioxide derived transfer coefficient k_{CO_2} after Quack and Wallace (2003) was applied for the three gases (Eq. 2).

380

$$F = k_w \cdot \Delta c \tag{Eq. 1}$$

381

$$k_w = k_{CO_2} \cdot \frac{Sc^{-\frac{1}{2}}}{600}$$
 (Eq. 2)

382

Details on measuring the air – sea concentration gradient are further described in Hepach et
al. (2014) and references therein.

385

386 2.4. Oceanic VSLS contribution to the MABL and FT

387 **2.4.1. Trajectory calculations**

The air mass transport from the surface to the FT was calculated with the Lagrangian Particle 388 Dispersion Model FLEXPART from the Norwegian Institute for Air Research in the 389 Department of Atmospheric and Climate Research (Stohl et al., 2005). The model has been 390 extensively evaluated in earlier studies (Stohl et al., 1998;Stohl and Trickl, 1999) and 391 includes parameterizations for turbulence in the atmospheric boundary layer and the FT as 392 well as moist convection (Stohl and Thomson, 1999;Forster et al., 2007). Meteorological 393 input fields are retrieved from the ECMWF (European Centre for Medium-Range Weather 394 Forecasts) assimilation reanalysis product ERA-Interim (Dee et al., 2011) with a horizontal 395 resolution of 1° x 1° and 60 vertical model levels. The ship-based 6 hourly radiosonde 396 measurements were assimilated into the ERA-Interim data (Section 2.2.1) and provide air 397 temperature, horizontal and vertical wind, boundary layer height, specific humidity, as well 398 as convective and large scale precipitation. For the trajectory analysis, 80 release points were 399 defined along the cruise track. Time and position of these release events are synchronized 400 401 with the water and air samples (Section 2.3). At each event, 10,000 trajectories were launched from the ocean surface within a time frame of \pm 30 minutes and an area of ~400 m². 402

404 **2.4.2. VSLS source-loss estimate in the MABL**

The time scales of air mass transport derived from FLEXPART together with the oceanic 405 emissions and chemical losses of the VSLS are used for a mass balance source-loss estimate 406 over the South China and Sulu Seas. For each release event, a box given by the in-situ height 407 of the MABL and by the horizontal area of the trajectory releases (~400 m² centred on the 408 measurement location) is defined. The MABL source-loss estimate is based on the 409 assumption of a constant VSL S mixing ratio (given by the atmospheric measurements), a 410 constant sea – air flux, the chemical loss rate, and a VSLS homogeneous distribution with the 411 412 box during each release.

The Oceanic Delivery (OD) is given as the contribution of VSLS sea – air flux (in mol per 413 day) to the total amount of the VSLS in the box (in mol) in percentage per day. The loss of 414 MABL air to the FT caused by vertical transport, denoted here COnvective Loss (COL), is 415 calculated from the mean residence time of the FLEXPART trajectories in the observed 416 MABL during each release and is given as a negative number in percentage per day. COL 417 equals the loss of VSLS from the MABL to the FT. The Chemical Loss (CL), in the form of 418 reaction with OH and photolysis, is estimated in percentage per day (negative quantity) and is 419 based on the tropical MABL lifetime estimates of 15 days for CHBr₃, 94 days for CH₂Br₂ and 420 4 days for CH₃I (Carpenter et al., 2014). 421

Relating the delivery of VSLS from the ocean to the MABL (*OD*) and the loss of MABL air
containing VSLS to the FT (*COL*) results in an Oceanic Delivery Ratio (*ODR*) (Eq. 3):

424

$$ODR = \frac{OD [\%d^{-1}]}{-COL [\%d^{-1}]} = \frac{Sea - Air flux contribution [\%d^{-1}]}{Loss of MABL air to the FT [\%d^{-1}]}$$
(Eq. 3)

425

Similarly, the Chemical Loss in the MABL (*CL*) related to the MABL VSLS loss into the FT
(*COL*) leads to a Chemical Loss Ratio (*CLR*) (Eq. 4):

428

$$CLR = \frac{CL [\%d^{-1}]}{-COL [\%d^{-1}]} = \frac{Loss through chemistry [\%d^{-1}]}{Loss of MABL air to the FT [\%d^{-1}]}$$
(Eq. 4)

429

The oceanic delivery, chemical loss and loss to the FT must be balanced by advective transport of air masses in and out of the box. We define the change of the VSLS through advective transport as Advective Delivery (AD) in percentage per day (Eq. 5). Additionally, we define the ratio of change in VSLS caused by advection (*AD*) to the loss of VSLS out of
the MABL to the FT as Advective Delivery Ratio (*ADR*) in Eq. 6:

435

$$AD = -COL - CL - OD \tag{Eq. 5}$$

$$ADR = \frac{AD [\%d^{-1}]}{-COL [\%d^{-1}]} = 1 - CLR - ODR$$
(Eq. 6)

436

Note that for the VSLS within the MABL box, *COL* and *CL* are loss processes and given as negative numbers while *OD* and *AD* (besides very few exceptions for the latter) are source processes and given as positive numbers. In order to derive the ratios, we divided *CL*, *OD* and *AD* by -COL and therefore end up with negative ratios for the loss process and positive ratios for the source processes.

In a final step, we relate the source-loss ratios (*ODR*, *CLR* and *ADR*) to the MABL VSLS volume mixing ratio (*VMR_{MABL}*) in the box (Eq. 7 – 9), in order to estimate VSLS newly supplied from oceanic delivery (*VMR_{ODR}*), lost by chemical processes (*VMR_{CLR}*) and supplied by advective transport (*VMR_{ADR}*).

446

$$VMR_{ODR} = ODR \cdot VMR_{MABL}$$
(Eq. 7)

$$VMR_{CLR} = CLR \cdot VMR_{MABL}$$
(Eq. 8)

$$VMR_{ADR} = ADR \cdot VMR_{MABL}$$
(Eq. 9)

447

448 2.4.3. Oceanic and MABL VSLS contribution to the FT

We use a simplified approach to calculate the mean contribution of boundary layer air masses 449 450 observed from various oceanic regions in the South China Sea on the ship, and the oceanic compounds therein, to the FT above the South China and Sulu Seas. The contribution is 451 determined as a function of time and altitude based on the distribution of the trajectories 452 released at each measurement location along the ship track. According to R/A FALCON 453 observations and our trajectory calculations we assume a well-mixed FT within 5° S - 20° N, 454 100° E – 125° E. Observations on R/V SONNE, on the other hand, are characterized by large 455 variability and are considered to be representative for the area along the cruise track where 456 the VSLS were measured in the water and atmosphere. We constrain our calculations to this 457 area and define 80 vertical columns along the cruise track. Each column extends horizontally 458 over the area given by the starting points of the trajectories (20 m x 20 m centred on the 459 measurement location) and vertically from the sea surface up to the highest point of R/A 460

FALCON observations around 13 km altitude. For each of the 80 columns along the cruise 461 track, 10,000 trajectories were launched and assigned an identical MABL air parcel 462 containing air with the VSLS mixing ratios observed on R/V SONNE during the time of the 463 trajectory release. The volume of the air parcel is given by the in-situ height of the MABL 464 and the horizontal extent of the release box (20 m x 20 m) divided by 10,000 trajectories. The 465 transport of the MABL air parcels is specified by the trajectories, assuming that no mixing 466 occurs between the parcels during the transport. Chemical loss of the VSLS in each air parcel 467 is taken into account through chemical degradation according to their specific tropospheric 468 lifetimes. The VSLS mixing ratios in the FT from the aircraft measurements are considered 469 representative for the whole South China Sea area. Thus we average over the volume and 470 mixing ratios of all trajectories within the South China Sea area, independent of their exact 471 horizontal location. Due to the decreasing density of air in the atmosphere with height, the 472 volume of the MABL air parcels expands along the trajectories with increasing altitude. The 473 expanding MABL air parcels take up an increasing fraction of air within the FT column, 474 which is taken into account in our calculations using density profiles from our radiosonde 475 476 measurements.

We calculate the contribution of oceanic compounds to the FT for 25 layers of 500 m height 477 intervals between 0.5 km and 13 km altitude within the column above the measurement 478 479 location. For each layer, the ratio r_{MABL} of the volume of the MABL air parcels with the VSLS mixing ratio VMR_{MABL} to the whole air volume of the layer is calculated. The ratio of 480 advected FT air with a mixing ratio VMR_{AFT} to the whole air volume of the layer is r_{AFT} , 481 respectively, with $r_{MABL} + r_{AFT} = 1$. In our simulation, the FT air with a mixing ratio VMR_{FT} 482 observed by R/A FALCON at a specific height is composed of the MABL air parcels and of 483 the advected FT air parcels (Eq. 10): 484

485

$$r_{MABL} \cdot VMR_{MABL} + r_{AFT} \cdot VMR_{AFT} = (r_{MABL} + r_{AFT}) \cdot VMR_{FT}$$
(Eq. 10)

486

487 The relative contribution C_{MABL} of VSLS observed in the MABL to the VSLS observed in the 488 FT is computed in altitude steps of 500 m (Eq. 11):

489

 $C_{MABL}[\%] = 100 \cdot (r_{MABL} \cdot VMR_{MABL}) / VMR_{FT}$ (Eq. 11)

491 The oceanic contribution C_{ODR} of the South China Sea emissions to the VSLS in the FT is 492 computed after Eq. 12:

493

$$C_{ODR}[\%] = 100 \cdot (r_{MABL} \cdot VMR_{ODR}) / VMR_{FT}$$
(Eq. 12)

494

The simplified approach also allows deriving mean VSLS mixing ratios accumulated in the FT from both MABL VSLS and oceanic emissions. The FT VSLS mixing ratios are simulated for each of the 80 columns by initiating a new trajectory release event using same meteorological conditions and VSLS MABL observations, when the former MABL air has been transported into the FT, according to the specific residence time in the MABL. The initial FT background mixing ratios are 0 ppt for each VSLS. The accumulated mean mixing ratio of a compound at a specific height is then computed after Eq. 13:

502

$$VMR_{MFT} = r_{MABL_1} \cdot VMR_{MABL_1} + r_{MABL_2} \cdot VMR_{MABL_2} + \dots + r_{MABL_i} \cdot VMR_{MABL_i}$$
(Eq. 13)

503

Here, VMR_{MFT} is the modelled accumulated FT mixing ratio, r_{MABL_i} is the ratio of MABL air 504 505 parcels in 20 m x 20 m x 500 m layers between 0.5 km and 13 km altitude to the total volume of each layer, VMR_{MABLi} is the mixing ratio in the MABL air parcels including chemical 506 degradation since release from the MABL, and *i* is the number of initiated runs per release. A 507 steady state for the compounds is reached, when variations in their mixing ratios vary less 508 than 1 % between two initiated runs. For CHBr₃ the steady state is reached after 11.0 ± 2.1 d 509 (mean $\pm \sigma$), 11.8 \pm 2.4 d for CH₂Br₂ and 8.0 \pm 1.4 d for CH₃I. The modelled overall mean FT 510 mixing ratio in the South China Sea is derived as the mean from the 80 individually 511 calculated FT mixing ratios determined along the cruise. The oceanic contribution to the FT 512 compounds is calculated with VMR_{ODR} from Eq. 7 inserted as VMR_{MABL} in Eq. 13. 513

514

3. Meteorological conditions in the MABL and the FT

516 **3.1. Meteorology along the ship cruise**

Moderate to fresh trade winds were dominating the South China and Sulu Seas during the cruise (Figure 1a-b), indicated by the overall mean wind direction of northeast $(50^{\circ} - 60^{\circ})$ and a mean wind speed of $5.5 \pm 2.9 \text{ ms}^{-1}$. The wind observations reveal two different air mass origins. Between November 15 and 19, 2011 a gentle mean wind speed of $3.7 \pm 1.8 \text{ ms}^{-1}$ with a *northern* wind direction was observed, influenced by a weak low pressure system (not

shown here) over the central South China Sea moving southwest and passing the ship 522 position on 17.11.2011. During November 20 - 29, 2011 the wind direction changed to 523 *northeast* and the mean wind speed increased to moderate $6.4 \pm 3.0 \text{ ms}^{-1}$. A comparison 524 between 6 hourly ERA-Interim wind and a 6 hourly averaged mean of the observed wind on 525 R/V SONNE reveals an underestimation of the wind speed by ERA-Interim along the cruise 526 track by $1.6 \pm 1.4 \text{ ms}^{-1}$ on average (not shown here). The mean deviation of the wind 527 direction between reanalysis and observation is 2 ± 37 degree. Reanalysis and observed wind 528 speeds correlate with R = 0.76 and the wind directions with R = 0.86, reflecting a good 529 530 overall agreement between ship observation and ERA-Interim winds. With an observed mean surface air temperature (SAT) of 28.2 \pm 0.8 °C and a mean SST of 29.1 \pm 0.5 °C the SAT is 531 on average 1.0 ± 0.7 °C below the SST, which benefits convection of surface air (Figure 2). 532 Indeed enhanced convective activity and pronounced precipitation events have been observed 533 during the cruise (S-Figure 1). Figure 3a shows the time series of the relative humidity 534 measured by the radiosondes launched on R/V SONNE from the surface up to the mean 535 height of the cold point tropopause at 17 km. Elevated humidity is found on average up to 536 about 6 km, which implies a distinct transport of water vapour to the mid troposphere during 537 the cruise by deep convection or advection of humid air from a nearby convective cell. 538

539

3.2. Marine atmospheric boundary layer

Higher SSTs than SATs (Figure 2) cause unstable atmospheric conditions (negative values) 541 between the surface and about 50 - 100 m height (Figure 3b). Surface air is heated by warmer 542 surface waters and is enriched with humidity both benefiting moist convection. The stability 543 of the atmosphere increases above 420 ± 120 m and indicates the upper limit of the MABL at 544 this altitude range derived from radiosonde data (Figure 3b). The MABL height given by 545 ERA-Interim along the cruise track is with 560 ± 130 m systematically higher (not shown), 546 but still within the upper range of the MABL height derived from the radiosonde 547 measurements. The unstable conditions of the MABL and the increase of the atmospheric 548 stability above the MABL reflect the characteristics of a convective, well-ventilated tropical 549 boundary layer. In contrast to cold oceanic upwelling regions with a stable and isolated 550 MABL (Fuhlbrügge et al., 2013;Fuhlbrügge et al., 2015), the vertical gradient of the relative 551 humidity measured by the radiosondes (Section 3.1) and the height of the MABL do not 552 coincide. This is caused by increased mixing through and above the MABL by turbulence 553 and convection, which leads to the convective, well-ventilated MABL. 554

4. Atmospheric VSLS over the South China and Sulu Seas

557 4.1. Atmospheric surface observations on R/V SONNE

Overall, the three VSLS show a similar pattern of atmospheric mixing ratios along the cruise 558 track with lower atmospheric surface abundances before 21 November 2011 and higher 559 concentrations afterwards, which can be attributed to a change in air mass origin (Figure 1). 560 A decrease from 3.4 to 1.2 ppt of CHBr₃ occurs at the beginning of the cruise (Figure 4a) 561 when the ship left Singapore and the coast of the Malaysian Peninsula. On November 16 -562 19, 2011, when the ship passed the southern South China Sea, lower mixing ratios (\pm 563 standard deviation 1σ) of 1.2 ± 0.3 ppt prevail and also the lowest mixing ratios for CHBr₃ 564 during the whole cruise of 0.8 ppt are observed. At the coast of Borneo and the Philippines, 565 the average mixing ratio of CHBr₃ increases to 2.3 ± 1.4 ppt. The overall mean CHBr₃ 566 mixing ratio during the cruise is 2.1 ± 1.4 ppt (Table 1) and therefore higher than earlier 567 reported CHBr₃ observations of 1.2 ppt in January – March 1994 (Yokouchi et al., 1997), 1.1 568 ppt in September 1994 (Quack and Suess, 1999) and 1.5 ppt in June – July 2009 (Nadzir et 569 al., 2014) further offshore in the South China Sea. The higher atmospheric mixing ratios 570 during the R/V SONNE cruise in November 2011 in contrast to the lower mixing ratios in 571 these previous studies may point to stronger local sources, strong seasonal or interannual 572 573 variations, or even to long-term changes. CH_2Br_2 shows a mean mixing ratio of 1.2 ± 0.2 ppt (Table 1). Yokouchi et al. (1997) observed a lower mean atmospheric mixing ratio of 0.8 ppt 574 and Nadzir et al. (2014) of 1.0 ppt in the South China Sea. An increase of the CH₂Br₂ mixing 575 ratios from 1.0 ± 0.1 ppt to 1.3 ± 0.2 ppt is observed after November 21, 2011 coinciding 576 with an increase of the CH₃I concentrations from primarily 0.3 ± 0.0 ppt to 0.4 ± 0.1 ppt 577 (Figure 4a). The highest mixing ratio of CH₃I was detected in the southwestern Sulu Sea on 578 November 25, 2011 with 0.8 ppt. The overall mean atmospheric mixing ratio for CH₃I, of 0.4 579 \pm 0.1 ppt (Table 1) is lower than the mean of 0.6 ppt observed by Yokouchi et al. (1997). 580

The concentration ratio of CH_2Br_2 and $CHBr_3$ (Figure 4b) has been used as an indicator of relative distance to the oceanic source, where a ratio of 0.1 was observed crossing strong coastal source regions (Yokouchi et al., 2005;Carpenter et al., 2003). The ten times elevated CHBr₃ has a much shorter lifetime, thus degrading more rapidly than CH_2Br_2 , which increases the ratio during transport. Overall, the mean concentration ratio of CH_2Br_2 and CHBr₃ is 0.6 ± 0.2 , which suggests that predominantly older air masses are advected over the South China Sea.

588

589 4.2. Oceanic surface concentrations and emissions from R/V SONNE

VSLS in the surface sea water along the cruise track show highly variable distributions 590 (Figure 4c and Table 1). Oceanic CHBr₃ surface concentrations range from 2.8 – 136.9 pmol 591 L^{-1} with a mean of 19.9 pmol L^{-1} during the cruise, while CH₂Br₂ concentrations range from 592 2.4 - 21.8 pmol L⁻¹ with a mean of 5.0 pmol L⁻¹. CHBr₃ and CH₂Br₂ have similar distribution 593 patterns in the sampling region with near shore areas showing typically elevated 594 concentrations. CH₃I concentrations range from 0.6 - 18.8 pmol L⁻¹ with a mean of 3.8 pmol 595 L^{-1} and show a different distribution along the ship track which might be ascribed to 596 additional photochemical production of CH₃I in the surface waters (e.g. Manley and Dastoor, 597 1988; Manley and de la Cuesta, 1997; Richter and Wallace, 2004). 598

High levels of all VSLS are found in waters close to the Malaysian Peninsula, especially in 599 the Singapore Strait on November 16, 2011, possibly showing an anthropogenic influence on 600 the VSLS concentrations. VSLS concentrations decrease rapidly when the cruise track leads 601 to open ocean waters. Along the west coast (November 19 - 23, 2011) and northeast coast of 602 Borneo (November 25, 2011), bromocarbon concentrations are elevated, and especially 603 CHBr₃ concentrations increase in waters with lower salinities, indicating an influence by 604 river run off. Elevated CHBr₃ concentrations are often found close to coasts with riverine 605 inputs caused by natural sources and industrial and municipal effluents (see Quack and 606 Wallace, 2003; Fuhlbrügge et al., 2013 and references therein). 607

Oceanic emissions were calculated from synchronized measurements of sea water concentrations and atmospheric mixing ratios, sea surface temperatures and wind speeds, measured on the ship (Section 2.3.3). The overall VSLS distribution along the ship track is opposite for the oceanic and atmospheric measurements (Figure 4a-d). While the sea water concentrations of VSLS generally decrease towards the Sulu Sea, the atmospheric mixing ratios increase, leading to a generally lower concentration gradient of the compounds between sea water and air in the Sulu Sea (not shown here).

Coinciding low VSLS atmospheric background concentrations, high SSTs, elevated oceanic 615 VSLS concentrations and high wind speeds, lead to high emissions of VSLS for the South 616 China and Sulu Seas (Figure 4d) of $1486 \pm 1718 \text{ pmol m}^{-2} \text{ h}^{-1}$ for CHBr₃, $405 \pm 349 \text{ pmol m}^{-1}$ 617 2 h⁻¹ for CH₂Br₂ and 433 ± 482 pmol m⁻² h⁻¹ for CH₃I. In particular, CHBr₃ fluxes are very 618 high and thus confirm elevated coastal fluxes from previous campaigns in tropical source 619 regions (Quack et al., 2007). They often exceed 2000 pmol $m^{-2} hr^{-1}$ in the coastal areas and 620 are sometimes higher than 6000 pmol $m^{-2} hr^{-1}$, as in the Singapore Strait on November 15, 621 2011 and on November 22, 2011 at the northwest coast of Borneo, which was also an area of 622 strong convection (Figures 1, 4b). 623

4.3. VSLS intercomparison: R/A FALCON and R/V SONNE

The two profiles of the bromocarbon mixing ratios and the profile for CH₃I from the surface 626 to 13 km altitude as observed on R/A FALCON with the GhOST and WASP instruments 627 (Sala et al., 2014;Tegtmeier et al., 2013) are shown in Figure 5. Mean CHBr₃ mixing ratios 628 are 1.43 ppt (GhOST) and 1.90 ppt (WASP) in the MABL (0 - 450 m, determined from 629 meteorological aircraft observations similarly as for the radiosondes, Section 2.2.2) and 0.56 630 ppt (GhOST) and 1.17 ppt (WASP) in the FT (0.45 km - 13 km, Table 1). The GhOST 631 mixing ratios in the MABL are lower than those observed on R/V SONNE (2.08 ppt). A very 632 good agreement of the measurements is given for the longer lived CH₂Br₂ with 1.17 ppt (R/V 633 SONNE), 1.19 ppt (GhOST) and 1.15 ppt (WASP). CH₃I mixing ratios measured by GhOST 634 are 0.59 ± 0.30 ppt within the MABL of 450 m height, which is about 0.2 ppt higher than the 635 values from R/V SONNE. Above the MABL, the average mixing ratio of CH₃I decreases to 636 0.26 ± 0.11 ppt (Figure 5). 637

638 CHBr₃ and CH₂Br₂ concentrations in the MABL correlate with R = 0.83 for all instruments 639 (Figure 6). CHBr₃ and CH₃I concentrations correlate with R = 0.55 and CH₂Br₂ and CH₃I 640 with R = 0.66; all correlations are significant at 99 %. Even higher correlations are found if 641 only measurements on R/V SONNE are taken into account with R = 0.92 for CHBr₃ and 642 CH₂Br₂, R = 0.64 for CHBr₃ and CH₃I, and R = 0.77 for CH₂Br₂ and CH₃I.

Comparison of R/A FALCON and R/V SONNE data are obtained from their meetings on 643 November, 19 and 21, 2011 (Table 2), when aircraft and ship passed each other within 100 m 644 distance several times, measuring the same air masses. During both meetings, deviations 645 between the GhOST and WASP instruments on the aircraft are larger for the bromocarbons 646 than the deviation between the WASP and the ship measurements. According to Sala et al. 647 (2014) the agreement between the GhOST and WASP instruments are within the expected 648 uncertainty range of both instruments which is then assumed to be also valid for the ship 649 measurements (this study). The good agreement between WASP and ship data might be 650 caused by the same sampling and analysis method, both using stainless steel canisters and 651 subsequent analysis with GC/MS, while GhOST measures in-situ with a different resolution. 652 Since GhOST and WASP measurements together cover a larger spatial area and higher 653 temporal resolution, a mean of both measurements is used in the following for computations 654 in the free troposphere. For CH₃I significantly higher mixing ratios were measured during the 655 meetings between ship and aircraft (Table 2). Whether this offset is systematic for the 656 different methods, needs further investigation. 657

659

5. Air mass and VSLS transport from the surface to the free troposphere

5.1. Timescales and intensity of vertical transport

Forward trajectories computed with FLEXPART starting at sea level along the cruise track 662 yield an average MABL residence time of 7.8 ± 3.5 hr before the trajectories enter the FT 663 (Figure 7), reflecting a relatively fast exchange due to the convective well ventilated MABL 664 (Figure 3). The trajectories generally show a strong contribution of surface air masses to the 665 FT, despite some exceptions during November 18 - 22, 2011 (Figure 7). Most intense and 666 rapid transport of MABL air masses up to 13 km height occurs on November 17 and 23, 667 2011. To estimate the loss of air masses out of the South China Sea area between 5° S – 20° 668 N and 100° E – 125° E (Section 2.4.3) we determine the loss of trajectories out of this area 669 after their release (Figure 8). After 4 days, 88 % of all trajectories released along the cruise 670 track are still within this defined area of the South China Sea, and 31 % after 10 days. 671

- 672
- 673

5.2. Contribution of oceanic emissions to VSLS in the MABL

From the sea – air fluxes (Section 4.2) and the residence times of the surface trajectories in
the MABL (Section 5.1), the Oceanic Delivery (OD) and the COnvective Loss (COL) were
computed (Table 3) using the method described in Section 2.4.2.

Based on the OD and the COL, the Oceanic Delivery Ratio (ODR) is calculated in order to 678 characterize the relative contribution of the local oceanic emissions compared to the loss of 679 MABL air into the FT (Table 3, Figure 9). The average ODR during the cruise is 0.45 ± 0.55 680 for CHBr₃, which means that the loss from the MABL to the FT is balanced to 45 % by 681 oceanic emissions along the cruise track. The ODR for CH_2Br_2 is 0.20 ± 0.21 and for CH_3I 682 0.74 ± 1.05 , respectively, suggesting that the major amount of CH₃I originates from nearby 683 sources. Similarly to the ODR the CL is related to the COL to derive the Chemical Loss Ratio 684 (CLR) for the VSLS, which is 0.03 ± 0.01 for CHBr₃, 0.01 ± 0.00 for CH₂Br₂ and 0.09 ± 0.04 685 for CH₃I. When compared to the other source and loss processes, the chemical loss appears 686 negligible for all three gases. The ratio of the advective delivery (ADR) is 0.58 ± 0.55 for 687 CHBr₃, 0.80 ± 0.21 for CH₂Br₂ and 0.35 ± 1.02 , implying that most of the observed CH₂Br₂ 688 (80%) in the MABL is advected from other source regions. Applying the ODR to the 689 observed mixing ratios in the MABL gives an estimate of the VSLS originating from the 690 local oceanic emissions (VMR_{ODR}, Table 4). The local ocean emits a concentration that 691

equates to 0.89 ± 1.12 ppt CHBr₃, 0.25 ± 0.26 ppt CH₂Br₂ and 0.28 ± 0.40 ppt CH₃I in the MABL. The average transport from the MABL to the FT (Flux_{MABL-FT}), computed from the MABL concentrations and the trajectory residence time in the MABL, is 4240 ± 1889 pmol m⁻² hr⁻¹ for CHBr₃, 2419 ± 929 pmol m⁻² hr⁻¹ for CH₂Br₂ and 865 ± 373 pmol m⁻² hr⁻¹ for CH₃I. Calculations with the ERA-Interim MABL height, which is on average 140 m higher than the one derived from the radiosondes, leads to similar estimates (S-Table 1).

Since the wind is a driving factor for oceanic emissions and advection of VSLS, changes in 698 wind speed are assumed to affect atmospheric VSLS mixing ratios in the MABL during this 699 700 cruise. Significant correlations are found between wind speed and the observed mixing ratios of all three VSLS in the MABL with correlation coefficients of R = 0.55 (CHBr₃), R = 0.57701 (CH_2Br_2) and R = 0.56 (CH₃I), respectively. Mixing ratios that originate from oceanic 702 emissions (VMR_{ODR}) correlate significantly to the wind speed with R = 0.52, R = 0.72 and R 703 = 0.62, respectively. On the opposite, VMR_{ADR} , which is calculated as the residual from 704 VMR_{ODR}, is negatively correlated to the wind speed with R = -0.21, R = -0.32 and R = -0.53. 705 The correlations reveal that the contribution of oceanic emissions to MABL VSLS increase 706 707 for higher wind speeds, while the advective contribution decreases.

708

709 5.3. Oceanic contribution to the FT

710 5.3.1. Identification of VSLS MABL air in the FT

With a simplified approach (method description in Section 2.4.3) we are able to estimate the contribution of MABL air and regional marine sources observed on R/V SONNE to the FT. Individual MABL air masses during the cruise show the strongest contribution to the FT air within the lower 3 km of the atmosphere during the first 6 days after release (Figure 10a-f). Once the MABL air is spread in the FT column, a decrease of the contribution with time is due to the transport of trajectories out of the predefined South China and Sulu Seas area and their chemical loss (Figure 8).

The average contribution of VSLS concentrations in the MABL air to the FT concentrations (c_{MABL}) is generally highest for CHBr₃, with about 5 – 10 % above 3 km height within 10 days (coloured contours in Figure 10a), followed by CH₂Br₂ (2 – 5 % Figure 10b) and only 0 – 3 % for the short-lived CH₃I (Figure 10c). The chemical degradation of CH₃I, according to its short tropospheric lifetime of 3.5 days leads to a rapid decrease of the contribution 2 days after release.

To identify the contribution of the oceanic emissions to the FT VSLS during the cruise, the VMR_{ODR} of each compound is used as the initial mixing ratio in the MABL air mass. For CHBr₃ and CH₂Br₂ the local emissions contribute only up to 6 % and 2 % to the FT concentrations (c_{ODR} , Figure 10d-e) compared to the 10 %, respectively 5 %, of c_{MABL} . In contrast, the contribution of the local oceanic emissions of CH₃I (Figure 10f) is almost similar to the contribution of the observed MABL concentrations (Figure 10c).

730

731 5.3.2. Accumulated VSLS in the free troposphere

By simulating a steady transport of MABL air masses into the FT, mean accumulated VSLS mixing ratios in the FT along and during the cruise were computed (Figure 11) as described in 2.4.3. The simulated FT mixing ratios of CHBr₃ and CH₂Br₂ from the observed MABL (VMR_{MABL}) decrease on average from 1.7 and 1.1 ppt at 0.5 km height to 0.6 and 0.6 ppt at 7 km height and increase again above 8 km up to 0.8 (CHBr₃) and 0.8 ppt (CH₂Br₂). Simulated CH₃I shows a decrease from 0.20 ppt at 0.5 km to 0.05 ppt at 3 km. Above this altitude, the simulated mixing ratios of CH₃I are almost constant with 0.05 ppt.

739 To estimate the accumulated FT mixing ratios solely from oceanic emissions, the VMR_{ODR} is used as the initial MABL mixing ratio (Figure 11). The simulated FT mixing ratios using 740 either VMR_{MABL} or VMR_{ODR} as input reveal a similar vertical pattern, since both simulations 741 are based on the same meteorology and trajectories. While FT mixing ratios based on 742 VMR_{MABL} and VMR_{ODR} are similar for CH₃I (due to the large oceanic contribution to the 743 744 MABL mixing ratios), FT mixing ratios from VMR_{ODR} are on average ~0.5 ppt lower for CHBr₃ and ~0.6 ppt CH₂Br₂ than from VMR_{MABL}. Comparing the simulated VMR_{MABL} FT 745 mixing ratios with the observed FT mixing ratios from R/A FALCON reveals stronger 746 vertical variations for the simulations in contrast to the observations. CHBr₃ is overestimated 747 in the VMR_{MABL} simulation between 0 and 4 km altitude, as well as above 12 km. FT mixing 748 ratios from VMR_{ODR} are on average 0.3 ppt lower than observations from R/A FALCON and 749 thus explain about 60 % of the observed FT mixing ratios. Simulated CH₂Br₂ in the FT based 750 on VMR_{MABL} underestimates the observed mixing ratios between 6 and 10 km height. In 751 particular, the maximum between 7 and 9 km height is not reflected in the simulations. 752 However, observations of both bromocarbons are within 1σ of the FT simulations with 753 754 MABL air. The CH₃I simulations show a distinct underestimation of the observed FT mixing ratios. Adjusting the R/A FALCON values by the identified offset to R/V SONNE (Section 755 4.3 and Table 2) reveals a better agreement between observed and simulated FT mixing 756 ratios (Figure 11). 757

758

759 **5.3.3. Discussion**

760 Oceanic emissions of CHBr₃ from the South China Sea contribute on average 45 % to the simulated FT mixing ratios (Figure 11). Simulated FT mixing ratios from MABL 761 observations and observed FT mixing ratios agree quite well up to 10 km height. Oceanic 762 emissions of CHBr₃ from the South China Sea contribute on average 60 % to the FT mixing 763 ratios observed on the aircraft (Figure 11), while simulations based on the MABL mixing 764 ratios reflect the observations in the FT quite well. Thus, we assume that the observed MABL 765 mixing ratios of CHBr₃ are representative for the South China and Sulu Seas. In case of 766 CH₂Br₂ the underestimation of simulated FT mixing ratios between 6 and 10 km using 767 observed MABL mixing ratios may be due to higher background concentrations of CH₂Br₂ in 768 the mid to upper troposphere given the long atmospheric lifetime of 150 days at these levels. 769

On average, 45 % and 20 % of CHBr₃ and CH_2Br_2 abundances, respectively, in the MABL observed by the ship originate from local oceanic emissions along the ship track. Thus advection from stronger source regions, possibly along the coast (for CHBr₃) and from the West Pacific (CH₂Br₂), are necessary to explain the mixing ratios.

In contrast to CHBr₃ and CH₂Br₂, the simulated mixing ratios of CH₃I in the FT are strongly underestimated no matter whether observed MABL mixing ratios or oceanic emissions are used. The offset between the simulated and observed FT CH₃I could be caused by additional strong sources of CH₃I in the South China Sea area. Furthermore, modelling or measurement uncertainties may add to this offset.

The simulations use constant atmospheric lifetimes for each compound and neglect variations 779 with altitude which could impact the simulated abundances. However, the altitude variations 780 of the CH₃I lifetime in the MABL and FT are around 0.5 days (Carpenter et al., 2014) and 781 thus impacts on the simulated abundances are quite small. Therefore it seems unlikely that 782 the lifetime estimate causes a large underestimation of the FT CH₃I. Additional uncertainties 783 may arise from cloud induced effects on photolysis rates (Tie et al., 2003) and OH levels (e.g. 784 Tie et al., 2003; Rex et al., 2014) impacting the VSLS lifetimes. Deficiencies in the 785 meteorological input fields and the FLEXPART model, in particular in the boundary layer 786 and in the convection parameterizations would affect all compounds and their contribution to 787 the FT concentrations in a similar way and thus seems to be unlikely as well. Ship and 788 aircraft measurements revealed a possible instrumental offset for CH₃I (Section 4.3). When 789 we adjust the observations for the offset of CH₃I between R/V SONNE and R/A FALCON 790 the simulated and observed FT mixing ratios match better. Thus, an instrumental offset 791 causing, at least partially, the calculated discrepancy for CH₃I appears likely (Section 2.3.1). 792

Another explanation for the elevated CH_3I in the FT is advection of fresh air with elevated 793 CH₃I mixing ratios in the FT from e.g. South East Asia or the Philippines. These areas are 794 known to comprise strong sources for atmospheric CH₃I from e.g. rice plantations (Redeker 795 et al., 2003;Lee-Taylor and Redeker, 2005). In combination with convective activity over 796 land, which is common in this area (Hendon and Woodberry, 1993), the high observed FT 797 mixing ratios of CH₃I could be explained, despite the low oceanic contribution during the 798 cruise. The low observed MABL mixing ratios of CH₃I on R/V SONNE may thus also not be 799 representative for the area. Yokouchi et al. (1997) observed higher atmospheric CH₃I mixing 800 801 ratios in the South China Sea.

Finally, the method of our simplified approach includes uncertainties as well. Different 802 parameterizations for the transfer coefficient k_w such as Liss and Merlivat (1986), which is at 803 the lower end of reported parameterizations, and Wanninkhof and McGillis (1999), which is 804 at the higher end, are discussed in Lennartz et al. (2015). Both lead to a reduction of the 805 oceanic contribution to the atmospheric mixing ratios at the observed average moderate wind 806 speeds (~6 ms⁻¹) when applied to our data. Nonetheless, the general conclusion that local 807 oceanic sources of CHBr₃ and CH₃I significantly contribute to MABL mixing ratios remains 808 for the cruise. In times of possible higher wind speeds (>10 ms⁻¹), which are likely for this 809 810 region, the flux variations between the different parameterizations as well as the oceanic contribution to atmospheric abundances, would increase. Since observational studies 811 quantifying the oceanic contribution to atmospheric abundances of VSLS are quite rare, it is 812 difficult to evaluate our findings at the moment and more studies for different oceanic 813 regimes should be carried out to validate our results. 814

815

816 **6. Summary**

The contribution of oceanic VSLS emissions to marine atmospheric boundary layer (MABL) 817 and free troposphere (FT) air during the SHIVA campaign in November 2011 in the South 818 China and Sulu Seas was investigated in this study. Meteorological parameters were 819 measured near the ocean surface and in the troposphere by regular radiosonde launches on 820 R/V SONNE during the cruise. Oceanic VSLS emissions were determined from simultaneous 821 atmospheric observations and sea surface water concentrations. The transport from the 822 surface through the MABL into the FT was computed with the trajectory model FLEXPART. 823 The ship campaign was dominated by north-easterly winds with a characteristic moderate 824 mean wind speed of 5.5 ms⁻¹. The radiosonde launches revealed a convective, well-ventilated, 825 weakly developed MABL with an average height of 420 ± 120 m during the cruise. 800,000 826

forward trajectories, launched from the ocean surface along the cruise track, show a rapid 827 exchange of MABL air with the FT within 7.8 hrs. The observations on R/V SONNE reveal 828 high mean ocean surface concentrations and emissions for CHBr₃ (19.94 pmol L^{-1} and 1486 829 pmol $m^{-2} hr^{-1}$), CH_2Br_2 (4.99 pmol L^{-1} and 405 pmol $m^{-2} hr^{-1}$) and CH_3I (3.82 pmol L^{-1} and 830 433 pmol $m^{-2} hr^{-1}$) in comparison to other oceanic source regions. Atmospheric mixing ratios 831 in the MABL, on the other hand, are relatively low with mean values of 2.08 ppt CHBr₃, 1.17 832 ppt CH₂Br₂, and 0.39 ppt CH₃I. The contribution of the oceanic VSLS emissions to their 833 MABL concentrations was evaluated by simple source-loss estimates, resulting in an Oceanic 834 Delivery Ratio (ODR). The ODR for CHBr₃ is 0.45, revealing that up to 45 % of CHBr₃ 835 mixing ratios in the MABL above the marginal seas originated, on average, from local 836 oceanic sources, while 74 % of CH₃I and only 20 % of CH₂Br₂ originates from the local 837 ocean. This indicates that the long-lived CH₂Br₂ is largely advected in the MABL. 838

We extend our analysis to the FT using VSLS observations from R/A FALCON above the 839 South China Sea. A single MABL air release contributes up to 28 % (CHBr₃), 12 % (CH₂Br₂) 840 and 5 % (CH₃I) to the FT mixing ratio (Section 5.3.1). The mean contributions of the local 841 oceanic VSLS to the FT within this MABL air release are up to 13 % (CHBr₃), 3 % (CH₂Br₂) 842 and 4 % (CH₃I). In order to estimate if the accumulated contributions from the single MABL 843 air releases are sufficient to explain the accumulated VSLS mixing ratios observed in the FT, 844 a steady transport of observed MABL air masses and oceanic emissions into the FT above the 845 South China Sea was simulated. The simulations for CHBr₃ based on the volume mixing 846 ratios in the MABL (VMR_{MABL}) reflect the observed mixing ratios in the FT, while the 847 simulations based on the local oceanic emissions (VMR_{ODR}) explained about 60 %. In the 848 MABL, the local oceanic emissions along the cruise track also explain half of the CHBr₃. 849 Thus, we conclude that the observed mixing ratios of CHBr₃ in the MABL are influenced by 850 stronger, possibly coastal sources, in the region. 851

CH₂Br₂ in the FT, simulated from observed MABL mixing ratios, shows a good agreement 852 between observations and simulations. However, slightly lower simulated mixing ratios in the 853 mid to upper troposphere compared to the observations indicate an accumulation of CH₂Br₂ 854 at these levels due to its longer atmospheric lifetime compared to CHBr₃. CH₃I in the FT is 855 underestimated in the simulations, using both the observed MABL mixing ratios and the 856 oceanic emissions. Even addressing an unresolved offset between the ship and aircraft data 857 leads to an underestimation of CH₃I in the FT, which points to additional CH₃I sources, e.g. 858 rice plantations in the region together with pronounced convection. 859

Our investigations show how oceanic emissions of VSLS in a strong oceanic source region 860 contribute to the observed atmospheric mixing ratios in the MABL. Furthermore, the 861 contributions of these atmospheric mixing ratios, and the local oceanic VSLS, to the observed 862 VSLS in the FT above this source region are derived. The results reveal strong links between 863 oceanic emissions, atmospheric mixing ratios, MABL conditions and prevailing convective 864 activity in the troposphere. The methods should be applied to other oceanic regions to derive 865 a better process understanding of the contributions of air-sea gas exchange on atmospheric 866 abundances. For the detection of future climate change effects on ocean surface trace gas 867 868 emissions and their influence on atmospheric chemistry and composition it is important to study the complex interplay between oceanic sources and emissions, meteorology, 869 atmospheric mixing ratios, and transport to the upper atmosphere. 870

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Figure 1: ERA-Interim mean wind field November 15 - 30, 2011 (arrows) and 10 minute running mean of wind speed observed on R/V SONNE as the cruise track. The black squares show the ships position at 00 UTC each day.



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Figure 2 a-b: Time series of (a) wind speed (blue) and wind direction (orange) and (b) surface air temperature (SAT, orange) and sea surface temperature (SST, blue) on the left scale, as observed on R/V SONNE. The temperature difference of SAT and SST (Δ T) is given on the right scale in (b). A Δ T of 0 K is given by dashed line. The shaded areas (grey) in the background show the 24 h stations. The data are averaged by a 10-minute running mean.



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Figure 3: (a) Relative humidity by radiosondes up to 17 km height, the mean cold point tropopause level. The dashed lines and the two numbers above the figure indicate the two 24 h stations. (b) Virtual potential temperature gradient as indicator for atmospheric stability (red for stable, white for neutral and blue for unstable) with MABL height from radiosondes (black curve) and from ERA-Interim (blue curve). The y axis is non-linear. The lower 1 km is enlarged to display the stability around the MABL height. The vertical lines and the two numbers above the figures indicate the two 24 h stations.



Figure 4 a-d: (a) Atmospheric mixing ratios of CHBr₃ (blue), CH_2Br_2 (dark grey) and CH_3I (red) measured on R/V SONNE. (b) Concentration ratio of CH_2Br_2 and $CHBr_3$ on R/V SONNE. (c) Water concentrations of CH_3I , $CHBr_3$ and CH_2Br_2 measured on R/V SONNE. (d) Calculated emissions of $CHBr_3$, CH_2Br_2 and CH_3I from atmospheric and water samples measured on R/V SONNE. The two shaded areas (light grey) in the background show the 24 h stations. Y-axis for (c) and (d) are non-linear.

Table 1: Mean ± standard deviation and range of atmospheric mixing ratios observed on R/V SONNE (195 data points) and R/A FALCON (GhOST-MS with 513 and WASP GC/MS with 202 data points) in the MABL and the FT, water concentrations observed on R/V SONNE and the computed sea – air fluxes. MABL and FT mixing ratios on R/A FALCON are adopted from Sala et al. (2014) and Tegtmeier et al. (2013). The R/A FALCON MABL height was analysed to be 450 m (Sala et al., 2014).

				CHBr ₃	CH ₂ Br ₂	CH ₃ I
	R/V SONNE			2.08 ± 1.36	1.17 ± 0.19	0.39 ± 0.09
				[0.79 - 5.07]	[0.71 - 1.98]	[0.19 - 0.78]
Atmosph. mixing ratios [ppt	R/A FALCON	GhOST	MABL	1.43 ± 0.53	1.19 ± 0.21	0.59 ± 0.30
				[0.42 - 3.42]	[0.58 - 1.89]	[0.29 - 3.23]
			FT	0.56 ± 0.17	0.87 ± 0.12	0.26 ± 0.11
				[0.16 - 2.15]	[0.56 - 1.54]	[0.08 - 0.80]
		WASP	MABL	1.90 ± 0.55	1.15 ± 0.14	/
				[0.99 - 3.78]	[0.85 - 1.59]	7
			FT	1.17 ± 0.50	0.88 ± 0.14	/
			11	[0.43 - 3.22]	[0.46 - 1.36]	/
Water concentrations			ntrations	19.94 ± 17.90	4.99 ± 2.59	3.82 ± 2.43
$[pmol L^{-1}]$]	[2.80 – 136.91]	[2.43 – 21.82]	[0.55 - 18.84]
Sea – air flux			flux	1486 ± 1718	405 ± 349	$\overline{433 \pm 482}$
$[pmol m^{-2} h^{-1}]$			$^{2} h^{-1}$]	[-8-13149]	[16 - 2210]	[13 - 2980]





Figure 5: Vertical distribution $CHBr_3$ (blue), CH_2Br_2 (grey) and CH_3I (red) mixing ratios measured in-situ by GhOST (diamonds) and with flasks by WASP (circles) on R/A FALCON. CH_3I was only measured in-situ by GhOST. The lower 2 km are non-linear displayed.



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Figure 6: Correlation of bromoform and CH_2Br_2 (upper left), bromoform and CH_3I (upper right), and CH_2Br_2 and CH_3I (lower left) from GhOST and WASP for all heights (ALL) and only within the MABL (MABL) and from R/V SONNE.

Table 2: Mean atmospheric mixing ratios of CHBr₃, CH_2Br_2 and CH_3I observed on R/V SONNE and R/A FALCON during two case studies on 19.11.2011 at 3.2° N and 112.5° E and on 21.11.2011 at 4.6° N and 113.0° E. During the two meetings two (one) measurements have been taken by R/V SONNE, 20 (5) measurements on R/A FALCON by GhOST and 17

1113 (21) by WASP.

		CHBr ₃ [ppt]	CH ₂ Br ₂ [ppt]	CH ₃ I [ppt]
November 19, 2011	R/V SONNE	1.37	1.37 0.99	
	R/A FALCON: GhOST / WASP	1.02 / 1.37	0.94 / 1.03	0.45 / -
2011	R/V SONNE	2.05	1.08	0.28
November 21,	R/A FALCON: GhOST / WASP	1.63 / 2.00	1.31 / 1.08	0.82 / -

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Figure 7: Forward trajectory runs along the cruise track with FLEXPART using ERA-Interim data. The black contour lines show the mean amount of trajectories (in %) reaching this height within the specific time (colour shading). The white line indicates the radiosonde MABL height.



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Figure 8: Horizontal distribution, altitude and amount of trajectories over time during the cruise. The red box represents the South China and Sulu Seas area. The lower right plot shows the amount of trajectories that remain in the box with time from all trajectory releases.

1127 Table 3: Mean ± standard deviation of Oceanic Delivery (OD), COnvective Loss (COL),

1128	Chemical Loss (CL), Advective Delivery (AD), Oceanic Delivery Ratio (ODR), Chemical
1129	Loss Ratio (CLR), and Advective Delivery Ratio (ADR) for CHBr ₃ , CH ₂ Br ₂ and CH ₃ I.

	OD	COL	CL	AD	ODB		
	[% d ⁻¹]	UDK	CLK	ADK			
	116.4	-307.6		198.2	0.45	-0.03	0.58
HBr ₃	±	±	-7.1	±	±	±	±
CF	163.6	124.3		199.7	0.55	0.01	0.55
~	54.2	-307.6		254.6	0.20	-0.00	0.80
CH ₂ Br ₂	±	±	-1.2	±	±	±	±
	66.7	124.3		131.9	0.21	0.00	0.21
	166.5	-307.6		165.2	0.74	-0.09	0.35
CH ₃ I	±	±	-24.0	±	±	±	±
	185.8	124.3		242.3	1.05	0.04	1.02



Figure 9: Average budgets of the Oceanic Delivery (OD, blue), Chemical Loss (CL, red),
Advective Delivery (AD, green) and Convective Loss (COL, orange) of CHBr₃, CH₂Br₂ and
CH₃I in the Marine Atmospheric Boundary Layer (MABL).Budgets of the Oceanic Delivery
Ratio (ODR, blue), Chemical Loss Ratio (CLR, red) and Advective Delivery Ratio (ADR,
green) of CHBr₃, CH₂Br₂ and CH₃I.

Table 4: Mean \pm standard deviation of observed Volume Mixing Ratios in the MABL on R/V SONNE (VMR_{MABL}) versus the amount of VMR originating from oceanic emissions (VMR_{ODR}), chemically degraded according to the specific lifetime (VMR_{CLR}), originating from advection (VMR_{ADR}) and the Flux from the MABL into the FT (Flux_{MABL-FT}) for CHBr₃, CH₂Br₂ and CH₃I.

	VMR _{MABL}	VMR _{ODR}	VMR _{CLR}	VMR _{ADR}	Flux _{MABL-FT}
	[ppt]	[ppt]	[ppt]	[ppt]	$[pmol m^{-2} hr^{1}]$
	2.08	0.89	-0.06	1.18	4240
HBr ₃	±	±	±	±	±
CI	1.36	1.12	0.04	1.20	1889
5	1.17	0.25	-0.01	0.92	2419
2Br	±	±	土	±	±
СН	0.19	0.26	0.00	0.27	929
	0.39	0.28	-0.04	0.13	865
H ₃ I	±	±	±	±	±
D D	0.09	0.40	0.02	0.37	373

1144

- 1146 Table 5: Correlation coefficients between wind speed and VSLS MABL mixing ratios
- 1147 (VMR_{MABL}), the Oceanic Delivery (OD), the COnvective Loss to the FT (COL), the
- 1148 Advective Delivery (AD), computed as the residual of OD, and the mixing ratios originating
- from the OD (VMR_{ODR}) and from the AD (VMR_{ADR}). Bold numbers are significant at the 95
- 1150 % (p-value).

Wind speed	CHBr ₃	CH ₂ Br ₂	CH ₃ I
VMR _{MABL}	0.55	0.57	0.56
OD	0.31	0.48	0.52
COL		-0.33	
AD	-0.46	0.56	-0.57
VMR _{ODR}	0.52	0.72	0.62
VMR _{ADR}	-0.17	-0.31	-0.49



Figure 10: Mean MABL air contribution (a - c) and oceanic contribution (d - f) to observed FT mixing ratios observed by R/A FALCON for three VSLS. The black contour lines show the mean portion of MABL air masses in the FT [%], the colours show the oceanic contribution to the observed compounds in the FT at specific height and day after release [%] including chemical degradation, the loss out of the South China Sea area with time and the vertical density driven extension of MABL air masses. The scale of the coloured contour is logarithmic.



Figure 11: Mean FT mixing ratios (solid lines) and 1 standard deviation (shaded areas) from in-situ and flask observations on R/A FALCON (Obsv., black) versus simulated mean FT mixing ratios from MABL air (MABL, red) and oceanic emissions (Ocean, blue) observed by R/V SONNE. R/A FALCON in-situ observations have been adjusted for CH₃I (Obsv.*, dashed black) according to measurement deviations during the meetings of R/V SONNE and R/A FALCON (Table 2; Section 4.3).