Automated, high frequency, on-line dimethyl sulfide measurements in natural waters using a novel "microslug" gas-liquid segmented flow method with chemiluminescence detection

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# 16 Abstract

17 Dimethyl sulfide (DMS) is the major biogenic volatile sulfur compound in surface seawater. Good quality DMS data with high temporal and spatial resolution is desirable 18 19 for understanding reduced sulfur biogeochemistry. Here we present a fully automated 20 and novel "microslug" gas-liquid segmented flow-chemiluminescence (MSSF-CL) based method for the continuous in-situ measurement of DMS in natural waters. 21 22 Samples were collected into a flow tank and DMS transferred from the aqueous phase 23 to the gas phase using a vario-directional coiled flow, in which microvolume liquid and 24 gas slugs were interspersed. The separated DMS was reacted with ozone in a reaction 25 cell for CL detection. The analytical process was automated, with a sample throughput of 6.6 h<sup>-1</sup>. Using MSSF for DMS separation was more effective and easily integrated 26 with CL detection compared with the commonly used bubbling approach. Key 27

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28 parameters of the proposed method were investigated. The linear range for the method was 0.05-500 nM ( $R^2 = 0.9984$ ) and the limit of detection (3 x S/N) was 0.015 nM, 29 which is comparable to the commonly used gas chromatography (GC) method and 30 sensitive enough for direct DMS measurement in typical aquatic environments. 31 Reproducibility and recovery were assessed by spiking natural water samples (river, 32 33 lake, reservoir and pond) with different concentrations of DMS (10, 20 and 50 nM), giving relative standard deviations (RSDs) <1.75% (n = 5) and recoveries of 94.4 – 34 35 107.8%. This fully automated system is reagent free, easy to assemble, simple to use, portable (weight ~5.1 kg) and can be left in the field for several hours of unattended 36 operation. The instrumentation can provide high quality DMS data for natural waters 37 with an environmentally relevant temporal resolution of ~9 min. 38

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#### 40 Keywords

Dimethyl sulfide; Chemiluminescence; Natural waters; Segmented flow; Automated;
On site analysis

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# 44 **1. Introduction**

Dimethyl sulfide (DMS) is a climatically active biogenic gas with an estimated 45 annual global emission of 28-31 Tg S a<sup>-1</sup>. Natural emissions account for approximately 46 78% of the total natural reduced sulfur global flux to the atmosphere [1,2] by transfer 47 from seawater, freshwater [3,4], soil [5,6] and plants [7]. After emission to the 48 atmosphere, DMS can be oxidized to SO<sub>2</sub>, which is a precursor of sulfate aerosol 49 50 particles that may act as cloud condensation nuclei (CCN) [8]. CCN are important for 51 climate because they affect the radiative properties of the atmosphere and clouds by 52 scattering solar radiation and influencing cloud microphysics and albedo [9,10].

53 DMS is volatile in natural waters and can be oxidized [11] and converted to other 54 sulfur compounds by microorganisms [12]. The transient nature of DMS means that in-55 situ analysis is essential. Currently, the most commonly used method for DMS 56 quantification is purge and trap gas chromatography (PT-GC) [13,14] coupled with 57 flame photometric [15] or mass spectrometric detection [16]. These GC based

techniques involve bulky instrumentation, require controlled laboratory settings and 58 have a relatively low sample throughput, which restricts the ability to make near-59 continuous measurements [17,18]. Techniques such as membrane inlet mass 60 spectrometry (MIMS) [19], equilibrator inlet proton transfer reaction mass 61 spectrometry (EI-PTRMS) [20-22] and atmospheric pressure chemical ionization-mass 62 63 spectrometry (AP-CIMS) [23,24] have become attractive for real-time DMS analysis 64 on research vessels. However, these devices are relatively heavy, fragile, expensive and 65 labor intensive to deploy on a ship.

An alternative strategy for measuring DMS is using gas phase chemiluminescence (CL) based on the chain reaction of DMS with ozone to form the sulfur monoxide radical (SO·), which then reacts with ozone to produce light with a wavelength maximum ( $\lambda_{max}$ ) at 370 nm [25,26]. The reaction is summarized in eq. (1) and eq. (2).

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$$DMS + \frac{1}{3}O_3 \xrightarrow{\text{chain}} SO \cdot$$
(1)

72 
$$SO \cdot +O_3 \rightarrow SO_2^* + O_2 \rightarrow SO_2 + hv$$
 (2)

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Green et al. [27] adapted a laboratory-based gas phase CL instrument for real-time 74 determination of DMS in marine samples. Air was bubbled through the sample to 75 76 transfer DMS from the aqueous phase to the gas phase. DMS and ozone mixed in a reaction chamber and the CL signal was recorded using a photomultiplier tube. A short-77 pass optical filter was used to reduce CL interference from other gases but this also 78 79 reduced the DMS signal by 89.7% and interference from methanethiol could not be 80 eliminated. Toda's group have pioneered the development of simple methods for the insitu measurement of DMS in seawater using gas phase CL in both sequential and 81 batchwise approaches [28-31]. DMS was vaporized and introduced into the CL reaction 82 83 cell by a physical shot or bubbling, while interferences from other gases were removed 84 either by adding a heavy metal agent to the sample or by using a soda lime column.

Here we present a fully automated microslug segmented flow-chemiluminescence
(MSSF-CL) system for the continuous measurement of DMS in natural waters. With

87 the proposed MSSF approach, nanomolar concentrations of DMS can be effectively transferred from the aqueous phase to the gas phase for CL detection. The whole 88 89 analytical procedure, including in-situ sampling, separation, CL quantification and 90 rinsing, was automated. This analytical system is easy to setup and operate, can be remotely operated and is light and portable (weight ~5.1 kg) and avoids the necessity 91 92 of using any reagents other than oxygen. The performance of the automated system was demonstrated by several hours of unattended, high temporal resolution DMS 93 94 measurement in the field.

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## 96 2. Experimental section

## 97 2.1. Reagents

A 1.0 mM DMS stock solution was prepared by diluting a DMS certified standard 98 (o2si, CA, USA) with methanol. The DMS stock solution was stored in a 20 mL glass 99 100 vial with an aluminum screw top cap and airtight silicon septum at -10 °C in the dark 101 to minimize evaporation. A 1.0 µM DMS working solution was prepared daily by 102 dilution of the stock solution with Milli-Q water. A 10 ppmv DMS gas standard cylinder (in nitrogen (N<sub>2</sub>), Sichuan Zhongce Biaowu Technology, Chengdu, China) was used for 103 calibration. The dilution of the DMS gas standard was achieved using a compressed N<sub>2</sub> 104 cylinder (≥99.999% purity, Sichuan Qiao Yuan Gas, Chengdu, China). Compressed N<sub>2</sub> 105 106 was also used as the gas source in the segmented flow line and the carrier gas to introduce DMS into the CL cell. An oxygen (O<sub>2</sub>) cylinder (>99.99% purity, Sichuan 107 108 Qiao Yuan Gas, Chengdu, China) was used as the source gas for ozone generation.

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#### 110 **2.2. Apparatus**

A peristaltic pump (YZ-15 pump head, BT50S driver, Lead Fluid Technology Co., Ltd., Baoding, China) was used for water sampling. A set of three-way solenoid valves (VAS101, Ristron, Jiashan, China) and a 9600-step syringe pump (PVS-100, Ristron, Jiashan, China) equipped with a 10 mL syringe (Hamilton, CA, USA) were used for handling the aqueous samples and water. Ozone was generated by an ozone generator (M1000, Tonglin Technology, Beijing, China) with a maximum output of 1 g h<sup>-1</sup>. The

ozone output was adjustable by changing the generator working power. Mass flow 117 controllers (S48 300/HMT, Horiba Metron Instruments, Beijing, China) were used to 118 119 regulate gas flow rates in the analytical system. A glass made gas-liquid separator (Sichuan Shubo, Chengdu, China) was used for phase separation after the MSSF and 120 the separated gas sample was injected into the CL detection system using an electrically 121 122 actuated 6-port injection valve (Valco Instruments, Houston, USA) and a PTFE holding coil (2.5 m x 3.175 mm i.d.). The CL detection system comprised a CL reaction cell (40 123 124 mm x 25 mm i.d.) and a photomultiplier tube (PMT; R3550P, Hamamatsu Photonics, Japan). The reaction cell was made of stainless steel and the inside wall was chromium-125 plated to enhance light reflection. The PMT was located in an aluminum housing (95 126 127 mm x 55 mm i.d.) sealed from external light sources. An optical convex lens (d=25 mm, f= 25.4 mm) was placed between the CL cell and the PMT to focus the light. The CL 128 129 signal was recorded in photon counting mode using a multifunctional photon signal 130 analyzer (Novaphoton Technology, Chengdu, China), with an integrated high voltage DC power supply for the PMT. The output from the detector was recorded in photon 131 132 counting units (p.c.u.) and all CL intensity data are reported as the integral of p.c.u. over 133 time. A schematic diagram of the CL system is shown in Fig. S-1.

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#### 135 **2.3. Analytical procedures**

A schematic of the MSSF-CL instrument for the determination of DMS is shown 136 in Fig. 1. Samples were collected by placing tubing with a 16 mesh (1.0 mm) nylon net 137 138 over the opening below the water surface. With a peristaltic pump (PP) and a set of 139 polyethersulfone (PES) filters (FT1, 50 mm x 100 µm; FT2, 50 mm x 10 µm; FT3, 50 140 mm x 0.8 µm), samples were continuously collected into a 50 mL sample flow tank at 141 200 mL min<sup>-1</sup>. The sample in the flow tank was either discharged to waste or held ready for analysis. 10 mL of sample was pulled into the syringe (SY) by the syringe pump 142 (SP) at 150 mL min<sup>-1</sup> and subsequently expelled to the T-junction (PP, 0.3 mm i.d.) at 143 2.0 mL min<sup>-1</sup>. Compressed N<sub>2</sub> regulated by the mass flow controller (MFC) was 144 delivered to the T-junction at a flow rate of 4.0 mL min<sup>-1</sup>. Segmented gas-liquid 145 146 microslugs formed as the gas and water mixed at the T-junction and these microslugs

entered a vario-directional flow coil (PP, 20 m x 1 mm i.d., see Fig. 3 (c)). DMS 147 transferred from the aqueous phase into the gas phase within the flow coil. The gas 148 149 sample was separated in the gas-liquid separator and then passed through a soda limepacked column that dried the gas stream and eliminated any potential signal 150 interferences. Sample gas was collected in a holding coil (PTFE, 2.5 m x 3.175 mm i.d.) 151 and a 6-port injection valve was switched periodically to pump the sample into the CL 152reaction cell at 400 mL min<sup>-1</sup>. Ozone was delivered continuously into the CL reaction 153 154 cell at 200 mL min<sup>-1</sup>. DMS reacted with ozone in the cell to produce a CL signal, which was detected and amplified by the PMT and recorded by the photon signal analyzer in 155 photon counting mode. Waste air was passed through an activated carbon column before 156 discharge to the ambient environment. The system was rinsed three times with 10 mL 157 of water, which was aspired into the syringe and expelled towards the MSSF-CL system. 158 The flow rate for both water and gas in the rinsing line was 150 mL min<sup>-1</sup>, resulting in 159 a 15 s period for a single washing cycle. The CL reaction cell and its connecting tubing 160 for DMS introduction were shielded from light by wrapping them with aluminum foil. 161 The photo of the proposed MSSF-CL analysis system was provided in Fig. S-2. 162 Windows based, self-programmed software written in C++ was used to control the 163 syringe pump, MFCs, solenoid valves and the injection valve. Details of the operation 164 165 of these control units are shown in Table S-3.

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Figure 1. A schematic diagram of the proposed MSSF-CL instrument for the determination of DMS. NT, nylon net;
FT 1-3, filter; PP, peristaltic pump; SV 1 and 2, three-way solenoid valves; SFT, sample flow tank; SY, syringe; TJ,
T-junction; MFC 1-3, mass flow controllers; OG, ozone generator; GLSFC, gas-liquid segmented flow coil; GLS,
gas-liquid separator; IV, 6-port injection valve; HC, holding coil; SLC, soda lime column; RC, chemiluminescence
reaction cell; PMT, photomultiplier tube; PSA, photon signal analyzer; AC, active carbon column.

# 175 **3. Results and discussion**

# 176 **3.1. Flow and mixing regime**

DMS must be effectively transferred from the aqueous phase into the gas phase prior to its introduction into the CL cell. In a coiled, gas-liquid segmented flow, centrifugal forces create a secondary flow and the liquid and gas slugs create two counter rotating vortices that cause asymmetrical micro-recirculation towards the main flow direction (see Fig. 2), resulting in increased mass transfer between the two phases [32]. We used a 20 nM DMS solution to compare our gas-liquid segmented flow system with the bubbling or 'purging' approach that is often used to transfer DMS from liquid

to gas phase (Fig. 3). DMS transfer from a 10 mL sample volume was 1.67-fold more 184 185 effective using the gas-liquid segmented flow approach because the microslugs are a more stable and homogeneous gas-liquid dispersion system. DMS transfer is enhanced 186 187 in the segmented flow compared to the bubbling approach because the surface area to volume ratio (gas-liquid contact area) is greater, the mass transfer diffusion distance is 188 shorter and there is intense relative motion between the two phases [32-34]. Moreover, 189 bubble films can form when air bubbles are introduced into the sample at higher speed, 190 191 which could result in an inferior and unstable CL signal.

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Figure 2. Effect of gas and liquid slugs moving through a coiled tube in (a) clockwise and (b) counterclockwisedirections.

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Figure 3. A comparison of (a) bubbling, (b) unidirectional segmented flow, and (c) vario-directional segmented flow for 20 nM DMS transfer. Bubbles were generated by introducing 20 mL of air through a quartz sand bubble stone. Both the unidirectional and vario-directional segmented flow setups used PP tubing (20 m x 0.79 mm i.d.) with 1 cm curve radius. Liquid and gas were delivered at 2.0 and 4.0 mL min<sup>-1</sup> respectively. Peak height recorded as photon counting units (p.c.u.).

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204 We also compared vario-directional and unidirectional segmented flow. Variodirectional flow was achieved by entwining tubing on two glass rods in alternating 205 206 clockwise and counterclockwise directions. The vario-directional flow gave a ~10% 207 higher response than the unidirectional flow (Fig. 3). This may be because the rate of 208 recirculation in liquid and gas slugs when the flow enters a coil is greater at the inner wall than at the outer wall. As the slugs move along the vario-directional segmented 209 flow channel, the asymmetrical recirculation switches periodically (see Fig. 2), thereby 210 211 increasing the relative motion between the two phases, resulting in enhanced DMS mass transfer. Note that the degree of this relative motion mainly depended on the size of the 212 liquid and gas slugs and the curvature radius [33,35]. 213

214 The geometry of the tubing (curve radius, R, and internal diameter, D) in the gas-liquid segmented flow coil significantly influences DMS mass transfer from the sample 215 216 microslugs (Fig. 3). A tighter coil radius enhances asymmetrical recirculation in the microslugs, whilst increasing the internal diameter increases the contact area between 217 the gas and liquid phases and the retention time of the microslugs in the flow coil, all 218 of which enhance the CL intensity (see Fig. 4). However, we did not observe any 219 significant enhancement in CL intensity when D > 1 mm and R < 1 cm were applied, 220 221 suggesting that complete mass transfer was achieved at D = 1.0 mm and R = 1.0 cm and 222 hence these values were used for all subsequent experiments. Flow tubing with larger internal diameter resulted in longer residence times (quantitative data for these 223 experiments are provided in Table S-4). The length of the segmented flow coil affected 224 225 the residence time of the microslugs in the tubing, which may have a positive 226 correlation on the efficiency of DMS mass transfer. We compared different length of tubing (5, 10, 20 and 30 m). The efficiency of DMS mass transfer increased while the 227 tubing length increased from 5 to 20 m, and kept constant thereafter, indicating a 228 229 complete DMS mass transfer may occurred.





231 Figure 4. Effect of segmented flow tubing curve radius and internal diameter on 20 nM DMS mass transfer in MSSF-

**3.2. T-junction geometry and gas / liquid flow rates** 

Microslug formation in the proposed method was achieved using a T-junction and 235 the size of the microslugs in the segmented flow were influenced by the dimensions of 236 the T-junction [36], the flow rates of the fluid [37] and the relative viscosity of the two 237 phases [38]. The effect of the T-junction inlet width and the gas flow rate were studied 238 239 while keeping the sample flow rate constant at 2.0 mL min<sup>-1</sup> (Fig. 5). By increasing the gas flow rate, the gas and liquid drop volume ratio  $(V_{gd}/V_{ld})$  also increased, generating 240 smaller liquid microslugs in the segmented flow. The total gas-liquid contact area was 241 242 increased and DMS mass transferring consequently enhanced, resulting in a higher DMS signal. However, at higher gas flow rates ( $> 4.0 - 6.0 \text{ mL min}^{-1}$ ), the DMS signal 243 levelled off or decreased (Fig. 5). This may be because a higher flow rate leads to the 244 use of a larger volume of gas, which is likely to dilute the DMS and ozone 245 concentrations in the CL reaction cell. Moreover, the retention time of the microslugs 246 247 in the segmented flow may be decreased at higher flows, resulting in reduction of DMS 248 mass transfer efficiency. Different T-junction inlet widths (0.3 mm, 0.5 mm and 1.0 mm) were also compared. Smaller drops were generated when using a narrower inlet at the 249 250 same flow rate, resulting in a higher DMS signal (Fig. 5). T-junction of inlet widths less than 0.3 mm was not investigated since it is unavailable. But the recovery for DMS 251 measurements by using 0.3 mm inlet widths T-junction at gas flow rate of 4.0 mL min<sup>-</sup> 252 <sup>1</sup> was 97.1% (n=3), indicating a complete DMS mass transfer. Consequently, the 253 optimum conditions for generating the segmented microslugs were gas and sample flow 254 rates of 4.0 and 2.0 mL min<sup>-1</sup> respectively through a 0.3 mm width T-junction. 255



Figure 5. Effects on CL intensity due to T-junction inlet width and gas flow rate through the coil. The segmented flow setup used PP tubing (20 m x 1.0 mm i.d.) with a 1 cm curve radius and vario-directional flow. Error bars represent  $\pm$  1 SD of triplicate measurements.

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#### 262 **3.3. Effect of salinity on CL detection**

It is important to be able to apply the MSSF-CL method to saline matrices in order 263 264 to study the biogeochemistry of DMS in natural waters. However, salt is often added to 265 aqueous samples to enhance the mass transfer of volatile compounds into the headspace by lowering their partition coefficient [6,39]. A 20 nM DMS sample was spiked with 266 varying concentrations of NaCl (to give sample salinities, expressed as m/v NaCl, in 267 268 the range 0 - 5%, m/v), and subjected to analysis by MSSF-CL. As shown in Table 1, 269 no significant signal variation was observed, i.e. all results were within the mean  $\pm 2$ SD (58,800  $\pm$  650; SD, standard deviation). Further evidence of the suitability of the 270 271 method for analyzing saline samples is shown in Fig. 6, which compares calibration 272 graphs for DMS (0 - 100 nM) in 0% and 3.5% NaCl. There is no significant difference  $(t_{calc} = 4.956; t_{tab} = 9.605)$  between the slopes of the two calibrations. 273

Sample salinity	CL intensity	RSD (n=3)	
(%, m/v)	(p.c.u.)	(%)	
0.0	59,200	1.3	
0.5	58,500	1.7	
1.0	58,700	1.6	
1.5	58,800	1.6	
2.0	59,000	1.5	
2.5	58,200	1.6	
3.0	58,500	1.7	
3.5	59,200	1.5	
4.0	58,800	1.4	
4.5	59,000	1.3	
5.0	58,600	1.7	

Table 1. Effect of sample salinity (as NaCl) for 20 nM DMS measurement by MSSF-CL.



279 Figure 6. Calibration graphs for DMS measurement by the proposed MSSF-CL method for samples with 0 and 3.5 %

280 (m/v, as NaCl) salinity. Error bars represent  $\pm 1$  SD of triplicate measurements.

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## **3.4. Effect of ozone flow rate and concentration on CL detection**

283 DMS and ozone were introduced into the CL cell through concentric tubes and the 284 CL reaction occurred in the center of the reaction cell. CL intensity depends on maximizing the emission within the cell window. We therefore investigated the effect 285 286 of different ozone concentrations on CL intensity by adjusting the O<sub>2</sub> input flow rate and the power supplied to the ozone generator. Lowering the O<sub>2</sub> flow rate enhanced the 287 288 CL signal due to a longer residence time in the cell and more efficient ozone production in the generator. If the O<sub>2</sub> flow rate dropped too low however, excess ozone was 289 290 produced, resulting in a quenching of the CL signal. The effect of ozone flow rate and 291 concentration on the DMS signal is shown in Table S-5, with a maximum CL intensity achieved when ozone was delivered into the reaction cell at 200 mL min<sup>-1</sup> with a 292 concentration of 6550 ppmv (with the ozone generator working at 40% of its maximum 293 294 output). Air was not used as an ozone source in this study due to unstable ozone production (RSD  $\geq 10.2\%$ , n=5) at low flow rates ( $\leq 250 \text{ mL min}^{-1}$ ). It should be noted 295 296 that the optimum flow rates of both ozone and the carrier gas, as well as the ozone 297 concentration, vary over a relatively wide range when different shapes and sizes of 298 reaction cell are used [27-31].

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## 300 **3.5. Interference study**

301 Certain compounds positively interfere with the DMS measurement by reacting 302 with ozone to produce a CL signal [27,29-31,40-42] and the effect of these compounds 303 at three concentrations was therefore investigated using the relative CL intensity, which was defined as the ratio of the CL intensity of the potential interferent with DMS and 304 305 the CL intensity of DMS alone. The results are shown in Table 2. 100 nM of ethene or 306 propene produces a CL signal equivalent to ~3.3-3.9 nM DMS. Ethene and propene are 307 not found in most natural waters and therefore interferences would be negligible [43]. 308 100 nM Dimethyl disulfide (DMDS) produces a CL signal equivalent to 2 nM DMS.

309 The concentration of DMDS in freshwaters is typically no more than 17% of the DMS concentration [44,45], suggesting a maximum interference of  $\sim 1\%$ . Methyl mercaptan 310 (CH<sub>3</sub>SH) is a biologically generated sulfur compound found in natural waters [46] and 311 is more volatile than DMS (Henry's law constant of 0.39 M atm<sup>-1</sup> and 0.56 M atm<sup>-1</sup> for 312 CH<sub>3</sub>SH and DMS respectively). Previous CL work has reported a comparable (or higher) 313 CL signal relative to DMS [13,29,30,31,47]. A column packed with soda lime was 314 introduced between the gas-liquid separator and the holding cell. The column 315 316 eliminated the CH<sub>3</sub>SH interference, dried the sample gas and had no detectable impact on the DMS signal. 317

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Table 2. Relative CL intensity (%CL =  $CL_{spiked}/CL_{DMS only}$ ) due to potential interference to the MSSF-CL signal from other compounds. Compounds were spiked into a 10 nM aqueous DMS

321 aqueous sample.

Compound	Relative CL intensity (%)		
Compound	1 nM	10 nM	100 nM
DMS	100.0		
Isoprene	100.1	100.7	109.0
Ethene	100.3	104.1	132.7
Propene	100.3	103.4	139.1
Hydrogen sulfide	100.0	100.1	100.8
Methyl mercaptan	100.4	100.4	101.2
Carbon disulfide	100.1	101.2	110.0
Dimethyl disulfide	100.5	102.7	120.0
Carbonyl sulfide	100.1	100.2	100.4

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## 323 **3.6. Analytical figures of merit**

Under optimum conditions, the linearity of the proposed MSSF-CL method for DMS determination was in the range 0.05-500 nM ( $R^2$ =0.9984). The limit of detection (LOD) calculated from three times the signal-to-noise ratio was 0.015 nM. The 327 reproducibility and recovery of the MSSF-CL method was investigated by analyzing four natural water matrices (river, lake, reservoir and pond; see Fig. 7 caption for further 328 329 matrix details) spiked with different concentrations of DMS (10, 20 and 50 nM). The 330 RSDs were  $\leq 1.8\%$  (n=5 for each set of measurements) and recoveries were 94.4-107.8%, indicating acceptable precision and accuracy for the analysis of natural water 331 332 samples. The complete analytical cycle (including rinsing) took 548 s, which provided a sample throughput of  $\sim 6.6 \text{ h}^{-1}$ . A comparison of the MSSF-CL method with purge and 333 334 trap gas chromatography with mass spectrometric detection (PT-GC-MS) demonstrated satisfactory agreement with minimal apparent bias (slope =  $1.042\pm0.018$ , intercept = -335  $0.159\pm0.096$ , R<sup>2</sup>=0.9947; see Fig. 7), which shows that the proposed method is robust 336 and can perform well for a broad variety of aqueous sample matrices. Analytical 337 conditions for the PT-GC-MS method are described in Method S-6 and the figures of 338 merit are given in Table S-7. 339



341 Figure 7. Comparison of DMS measurement in different freshwater samples by MSSF-CL and PT-GC-MS.

342 Regression equation (with 95% confidence intervals) follows PT-GC-MS measurement (nM) = 1.042 (±0.018) x

 $MSSF-CL measurement (nM) - 0.159 (\pm 0.096). Lake and pond samples were collected from East Lake (University of Electronic Science and Technology of China, Chengdu, China), river samples were collected from different sites along the Qingshui river (Chengdu, China) and reservoir samples were collected from Zipingpu and Tuanjie reservoir (Chengdu, China). A 50 mL plastic syringe equipped with a 0.22 µm membrane filter was used for sampling. The syringe was filled while under water to prevent headspace formation. Collected samples were stored in the dark at ~4 °C in an expanded polypropylene ice cooler box until analyzed.$ 

- 349
- 350 **3.7. Field analysis of freshwater samples**

The suitability of the proposed method for field deployment was evaluated by 351 352 (pseudo)continuous monitoring to determine DMS in East Lake (University of Electronic Science and Technology of China campus, China) over a 10 h period (66 353 samples) on the 17<sup>th</sup> May 2019. Samples were continuously collected at fixed position 354 at a depth of 50 cm, as described in section 2.3, and introduced into the MSSF-CL 355 system for DMS measurement (results shown in Fig. 8). The DMS concentration 356 increased steadily from 09:00 hrs, reaching a maximum of 7.37 nM at 14:00. A 357 358 significant drop was then observed, decreasing to 4.06 nM at 19:20.

The data in Fig. 7 and Fig. 8 are within the range of previous freshwater DMS observations [48]. The DMS observations follow a similar diurnal cycle that has been observed in other studies [27,49]. The proposed method is reagent free, portable (weight ~5.1 kg excluding the gas cylinders), simple to use and ideally suited for field analysis with good temporal resolution.

Compared with the recent reported CL based sequential and batchwise method for DMS field analysis [30, 31], the using of a novel microslug gas-liquid segmented flow for DMS phase transferring in this work was found to be highly effective and compatible with the whole automated measurement system that featured by its portability, ease of operation, and could be left in the field for several hours unattended operation.



Figure 8. Field analysis of DMS in freshwater by MSSF-CL. The MSSF-CL system was placed at fixed position on
a footbridge over the East Lake of University of Electronic Science and Technology of China campus, and samples
were continuously collected from 50 cm below the water surface and delivered into the MSSF-CL system for analysis.
DMS measurements were automatically carried out from 09:10 to 19:20 (local time) without interruption, providing
DMS data every ~9.1 mins. The red line is a ±30 min running average.

#### 378 **4. Conclusions**

379 DMS biogeochemistry has attracted significant attention in environmental studies 380 as a biologically-generated, climate-relevant sulfur compound. We have developed an 381 automated system based on gas-liquid segmented flow and gas phase CL detection for 382 the quantification of DMS in natural waters. DMS transfer from the aqueous phase to the gas phase using a vario-directional, microslug gas-liquid segmented manifold was 383 384 highly advantageous compared with the commonly-used bubble purging approach. 385 Sample throughput, including *in-situ* sampling, separation, detection and washing, was 386 6.6 h<sup>-1</sup>. The system is portable, reagent free, uses off-the-shelf components and fittings 387 for ease of assembly/disassembly and can be deployed unattended in the field. The

geometry and flow rates in the gas-liquid segmentation system are critical for optimum 388 performance, as are the flow rate and concentration of ozone in the reaction cell. Under 389 390 optimum operating conditions the linear range for DMS detection was 0.05-500 nM  $(R^2=0.9984)$ , the LOD (3 x S/N ratio) was 0.015 nM, RSDs were typically  $\leq 1.8\%$  (n=5) 391 and recoveries for spiked (10, 20 and 50 nM DMS) natural waters were 94.4-107.8%. 392 393 The analytical performance of the proposed method means that it can be applied to the continuous measurement of low level DMS concentrations in natural waters. Sample 394 395 throughput could be enhanced by the use of a multi parallel gas-liquid segmented flow manifold and/or tangential flow filtration. The multi parallel segmented flow manifold 396 would introduce samples into different parallel gas-liquid segmented flows at 397 398 prescribed time intervals and queue the sample gas prior to entering the CL cell. In-line 399 tangential flow filtration could be incorporated to enable longer-term deployments.

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# **CRediT authorship contribution statement**

Geng Leng: Conceptualization, Methodology, Formal analysis, Investigation, 402 403 Writing - original draft, Writing - review & editing, Visualization, Investigation, 404 Resources. Chao Feng Jin: Methodology, Writing - review & editing, Software, Visualization, Investigation. Thomas G. Bell: Writing - review & editing, Formal 405 analysis, Resources, Conceptualization. Simon J. Ussher: Project administration, 406 Writing - review & editing, Formal analysis, Resources, Conceptualization. Paul J. 407 Worsfold: Supervision, Conceptualization, Writing – review & editing, Formal 408 analysis, Resources. Wei-Yi Li: Validation, Writing - review & editing, 409 Conceptualization. 410

411

# 412 **Declaration of competing interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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# 562 Figure and table captions

564 Figure 1. A schematic diagram of the proposed MSSF-CL instrument for the

- determination of DMS. NT, nylon net; FT 1-3, filter; PP, peristaltic pump; SV 1 and 2,
- three-way solenoid valves; SFT, sample flow tank; SY, syringe; TJ, T-junction; MFC 1-
- 567 3, mass flow controllers; OG, ozone generator; GLSFC, gas-liquid segmented flow coil;

GLS, gas-liquid separator; IV, 6-port injection valve; HC, holding coil; SLC, soda lime
column; RC, chemiluminescence reaction cell; PMT, photomultiplier tube; PSA,
photon signal analyzer; AC, active carbon column.

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Figure 2. Effect of gas and liquid slugs moving through a coiled tube in (a) clockwiseand (b) counterclockwise directions.

574

Figure 3. A comparison of (a) bubbling, (b) unidirectional segmented flow, and (c) vario-directional segmented flow for 20 nM DMS transfer. Bubbles were generated by introducing 20 mL of air through a quartz sand bubble stone. Both the unidirectional and vario-directional segmented flow setups used PP tubing (20 m x 0.79 mm i.d.) with 1 cm curve radius. Liquid and gas were delivered at 2.0 and 4.0 mL min<sup>-1</sup> respectively. Peak height recorded as photon counting units (p.c.u.).

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Figure 4. Effect of segmented flow tubing curve radius and internal diameter on 20 nM DMS mass transfer in MSSF-CL method (coil tubing length = 20 m; liquid flow rate =  $2 \text{ mL min}^{-1}$ ; gas flow rate =  $4 \text{ mL min}^{-1}$ ).

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Figure 5. Effects on CL intensity due to T-junction inlet width and gas flow rate through the coil. The segmented flow setup used PP tubing (20 m x 1.0 mm i.d.) with a 1 cm curve radius and vario-directional flow. Error bars represent  $\pm$  1 SD of triplicate measurements.

590

Figure 6. Calibration graphs for DMS measurement by the proposed MSSF-CL method for samples with 0 and 3.5 % (m/v, as NaCl) salinity. Error bars represent  $\pm$  1 SD of triplicate measurements.

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595 Figure 7. Comparison of DMS measurement in different freshwater samples by MSSF-

596 CL and PT-GC-MS. Regression equation (with 95% confidence intervals) follows PT-

597 GC-MS measurement (nM) =  $1.042 (\pm 0.018)$  x MSSF-CL measurement (nM) - 0.159

598 (±0.096). Lake and pond samples were collected from East Lake (University of Electronic Science and Technology of China, Chengdu, China), river samples were 599 collected from different sites along the Qingshui river (Chengdu, China) and reservoir 600 601 samples were collected from Zipingpu and Tuanjie reservoir (Chengdu, China). A 50 mL plastic syringe equipped with a 0.22 µm membrane filter was used for sampling. 602 The syringe was filled while under water to prevent headspace formation. Collected 603 samples were stored in the dark at  $\sim 4$  °C in an expanded polypropylene ice cooler box 604 605 until analyzed.

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Figure 8. Field analysis of DMS in freshwater by MSSF-CL. The MSSF-CL system was placed at fixed position on a footbridge over the East Lake of University of Electronic Science and Technology of China campus, and samples were continuously collected from 50 cm below the water surface and delivered into the MSSF-CL system for analysis. DMS measurements were automatically carried out from 09:10 to 19:20 (local time) without interruption, providing DMS data every ~9.1 mins. The red line is  $a \pm 30$  min running average.

614

Table 1. Effect of sample salinity (as NaCl) for 20 nM DMS measurement by MSSF-CL.

617

Table 2. Relative CL intensity (% $CL = CL_{spiked}/CL_{DMSonly}$ ) due to potential interference to the MSSF-CL signal from other compounds. Compounds were spiked into a 10 nM aqueous DMS aqueous sample.