A three year time-series of volatile organic iodocarbons in Bedford Basin, Nova Scotia :
 a Northwestern Atlantic fjord.

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- 7 Email: qshi@dal.ca 8
- 9 Abstract:
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We report weekly observations of volatile organic iodocarbons (CH₃I, CH₂ClI and CH₂I₂) over the 11 12 time-period May 2015 to December 2017 from 4 depths in Bedford Basin, a coastal fjord (70m 13 deep) on the Atlantic coast of Canada. The fjord is subject to winter-time mixing, seasonal 14 stratification and bloom dynamics, subsurface oxygen depletion, local input of freshwater and 15 occasional intrusions of higher density water from the adjacent continental shelf. Near-surface 16 concentrations showed strong seasonal and sub-seasonal variability which is compared with other 17 coastal time-series. The vertical variation of CH₂I₂ and CH₂CII within the upper 10m is consistent 18 with rapid photolysis of CH₂I₂. Average annual sea-to-air fluxes (46.7 nmol m⁻² day⁻¹) of total 19 volatile organic iodine were similar to those observed in other coastal and shelf time-series and 20 polyiodinated compounds contributed 80% of the total flux. Fluxes were subject to strong 21 interannual variability (a factor of two) mainly due to wind-speed variability. Near-surface net production of CH₃I averaged 1 pmol L⁻¹ day⁻¹ and was similar to rates in the English Channel but 22 23 an order of magnitude higher than in shallow waters of the Kiel Fjord, Germany, possibly due to 24 higher microbial degradation in the latter. The near-bottom (60 m) time-series showed evidence 25 for CH₃I production associated with organic matter degradation, and a possible "switch" from 26 production of $CH_{3}I$ via an alkylation pathway to production of $CH_{2}I_{2}$ by a haloform-type reaction. Near-bottom CH₃I production varied strongly between years but was generally ca. 20 times lower 27 28 than near-surface production.

Keywords: Iodocarbons, iodomethane, chloroiodomethane, diiodomethane, air-sea flux, time series

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4 1. Introduction

5 Volatile organic iodocarbons (VOIs) such as methyl iodide (CH₃I), chloroiodomethane (CH₂CII) 6 and diiodomethane (CH₂I₂) have a predominantly oceanic source and supply a significant amount 7 of iodine to the atmosphere (see review by Saiz-Lopez and Von Glasow, 2012). These gases, also referred to as VSLS (very short-lived halogenated substances) due to their reactivity and short 8 9 atmospheric lifetimes, have been implicated in supporting catalytic ozone destruction in the 10 troposphere (Davis et al., 1996; McFiggans et al., 2000) and, potentially in the lower stratosphere 11 (Solomon et al., 1994) as well as aerosol formation in the marine boundary layer (McFiggans et 12 al., 2000, 2004; O'Dowd et al., 2002). Recent modelling of atmospheric reactive iodine (IOx = IO+ I) as well as experimental studies (Carpenter et al., 2013; Jones et al., 2010; Mahajan et al., 2010) 13 14 suggest that the supply of volatile organoiodine represents <50% of the total sea-to-air delivery of reactive iodine, with most being supplied in the form of HOI and I₂. Nevertheless, the potential for 15 16 localized higher emissions coupled with their relatively long lifetimes (compared to I₂ and HOI) 17 allows the organic compounds to be a significant source of iodine to the free troposphere and even, 18 potentially, to the lower stratosphere in certain regions (Tegtmeier et al., 2013). Further, Mahajan 19 et. al (2012) noted a strong correlation of IO_x and CH₃I suggesting that the sources of CH₃I and 20 the shorter-lived precursors of IO_x are closely related or depend on similar variables.

CH₃I is the most abundant VOI species in the atmosphere (Yokouchi et al., 2011) because of its
longer lifetime (days) compared to CH₂ClI (hours) and CH₂I₂ (minutes) (Moessinger et al., 1998;
Rattigan et al., 1997). However, the total supply of organically-bound iodine to the atmosphere is

several times larger than the CH₃I supply alone (Carpenter et al., 2014) with the bulk of the 1 2 remainder transported in the form of CH₂I₂ and CH₂CII (additional iodocabons such as CH₃CH₂I, 3 CH₂BrI and CHI₃ are generally present in much lower concentration). Despite considerable 4 attention on the oceanic distribution and sea-to-air flux of these compounds, in particular CH₃I 5 (Ziska et al., 2013), it is not yet possible to apportion oceanic production of these compounds, 6 unequivocally, to specific mechanisms. Even for CH₃I, controversy remains, for example, as to 7 the relative importance of direct "biological" or "photochemical" production pathways with 8 experimental evidence reported for both, and correlation analysis generally being inconclusive, in 9 part because of the "snapshot" nature of most studies (Stemmler et al., 2014). Comparisons of 10 models to observed distributions have also proven ambiguous, with localized studies suggesting 11 predominance of a biological production pathway (Stemmler et al., 2013) but a global analysis 12 emphasising photochemical production as the dominant mechanism. This diversity of views has 13 been maintained through a variety of experimental studies (Amachi et al., 2001; Brownell et al., 14 2010; Hughes et al., 2011; Manley and delaCuesta, 1997; Moore and Tokarczyk, 1993; Moore and Zafiriou, 1994; Richter and Wallace, 2004; Shi et al., 2014a; Smythe-Wright et al., 2006). 15

For compounds other than CH₃I, similar uncertainty exists concerning production pathways, but with fewer underlying studies. Laboratory experiments have shown that the presence of dissolved iodide and dissolved organic matter can lead to production of these compounds in the dark (Martino et al., 2009). Fuse et al. (2003) and Martino et al. (2005) observed that CH₂CII could be produced by photolysis of CH₂I₂ in artificial and natural seawater. However detailed mechanisms and, especially, their relative importance in the field remain unclear.

Time-series observation can reveal processes and controlling factors underlying production and
loss of iodocarbons in the ocean and provide data for testing hypotheses and/or models. However,

only a very few long-term, time series observations of iodocarbons have been reported to date, all 1 2 from coastal water. Klick (1992) reported 13 months of weekly measurements of CH₂I₂ and 3 CH₂ClI from very shallow (3.5m) water in the Kattegat at the Swedish coast. Orlikowska and 4 Schulz-Bull (2009) reported a year of weekly data for CH₂ClI, CH₂I₂, CH₃I and C₂H₅I from a 5 nearshore (3m depth) site in the Baltic Sea. Archer et al. (2007) reported a seasonal study of 6 CH₂ClI, CH₂I₂, CH₃I, C₂H₅I, and CH₂BrI measured weekly at 4 depths (0-50m) in the western 7 English Channel from July 2002 to April 2004. Shi et al. (2014b) reported on the seasonal cycle of CH₃I from surface waters of the Kiel Fjord: a shallow (14 m), brackish water body in northern 8 9 Germany, which was sampled weekly for 2 years. Shimizu et. al (2017) presented a time-series of 10 vertical profiles (0-90m) of CH₂I₂, CH₂ClI, CH₃I, and C₂H₅I from the centre of Funka Bay, Japan, 11 which were measured every 2-4 weeks from March 2012 to December 2014.

12 Here, we report weekly observations of CH₃I, CH₂ClI and CH₂I₂ made over the time-period May 13 2015 to December 2017 at 4 depths (0-60m) in Bedford Basin: a coastal fjord on the east coast of 14 Canada. We report seasonal to interannual variability of the observed concentrations at different depths in the water column and compare our results with the other time-series. We report daily 15 16 average fluxes to the atmosphere and use a simple, time-varying mass-balance model for near-17 surface waters to estimate production rates and their variability. We discuss the observed 18 variability of both concentrations and production rates in the light of earlier studies, potentially 19 correlated variables and suggested production pathways.

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21 2. Methods

Time-series measurements of VOIs were carried out in the Bedford Basin (44.69 °N, -63.63 °E)
near Halifax, Canada. Bedford Basin is an 8 km long, 17 km² fjord with a maximum depth of 71m

and a total volume of 500 km³. The Bedford Basin is connected with continental shelf waters of 1 2 the Atlantic Ocean through "the Narrows" (a ca. 300 m wide and 20 m deep passage (Fig. 1)). The 3 Basin receives freshwater primarily from the Sackville River at its northwestern end, with a total average freshwater input of 5.41 m³ s⁻¹ (Buckley and Winters, 1992). The average near surface 4 5 salinity within the Basin is 29 which can be compared with salinities of >30 over the adjacent 6 Scotian Shelf. There are only relatively small horizontal gradients of near-surface salinity within the Bedford Basin itself (typically < 2 difference from close to the Sackville River mouth to the 7 8 Narrows).

9 Time series observations of physical, chemical and biological parameters have been recorded since 1992 (Li, 1998). Our halocarbon samples were collected weekly, in the center of the Bedford Basin, 10 at its deepest point (Fig. 1), between May 2015 and January 2018. Samples were collected with 11 12 10-L Niskin bottles attached to a rosette sampler at 1, 5, 10 and 60 m (10 m samples were collected 13 biweekly from May to September 2015). The upper three water samples covered the majority of 14 the euphotic zone. The 60m water sample was from typically stagnant, near-bottom water which 15 is renewed by vertical mixing events in late winter, and by occasional intrusions of higher-salinity 16 continental shelf water in both summer and winter. Chlorophyll a (Chla), dissolved oxygen, and 17 nutrients were measured weekly at the 4 depths as part of the Bedford Basin Monitoring Program (Details can be found in website: http://www.bio-iob.gc.ca/science/monitoring-monitorage/bbmp-18 19 pobb/bbmp-pobb-en.php). In addition to the Niskin bottle sampling, vertically continuous 20 measurements of temperature, salinity, dissolved oxygen and Chl *a* properties were measured with a CTD mounted on the rosette. Additional information concerning the measurements of supporting 21 22 physical and biological parameters can be found in the paper by Burt el al. (2013). Nutrients were measured using a Skalar SAN⁺⁺ autoanalyzer, the precisions of NO_2^- , NO_3^- and NH_4^+ were ± 0.01 23

1	μmol L ⁻¹ , ± 0.14 μmol L ⁻¹ and ± 0.14 μmol L ⁻¹ respectively. Chl <i>a</i> concentration was analyzed
2	using a fluorescence technique (Turner Design Model 10 Fluorometer) with the root mean square
3	error (RMSE) being 0.23 μ g L ⁻¹ . Dissolved oxygen concentrations were determined at 4 depths
4	using Winkler titration with a precision of \pm 0.5 µmol kg ⁻¹ . The resulting between-lab agreement
5	is approaching the specifications for repeat hydrography required by the Global Ocean Observing
6	System

7 (http://www.goosocean.org/components/com_oe/oe.php?task=download&id=35904&version=2.
8 0&lang=1&format=1).

9 The concentrations of iodomethane (CH₃I), chloroiodomethane (CH₂CII) and diiodomethane 10 (CH_2I_2) reported here, as well as of a number of other halocarbons (data not shown), were 11 measured using purge & trap gas chromatography with detection by both mass spectrometry (MS) and electron capture (ECD). All measurements were made using an Agilent Technologies gas 12 13 chromatograph (GC 7890B), equipped with a capillary column (RTX-VGC; 60 m; 1.4 µm coating, 14 column diameter: 0.25 mm; helium carrier gas 0.5 ml min⁻¹), together with an automated purge and trap system equipped with an autosampler (VSP4000 of IMT, Vohenstrauss, Germany). The 15 16 GC column was temperature programmed as follows: initial temperature 50 °C for 6 minutes, then 17 ramped to 150 °C at 6 °C min⁻¹; ramped to 200 °C at 10 °C min⁻¹. Water samples (10 ml) were stored in 20 ml vials equipped with an ultra-low-bleed septum, prior to purging with helium (20 18 19 ml min⁻¹ for 18 mins). Every sample was analysed in triplicate. The standard deviation of triplicate measurements (integrated peak area) was <10 % for CH₃I, <15 % for CH₂ClI and <20 % for CH₂I₂. 20

21 Calibration of the GC system for CH₃I, CH₂ClI and CH₂I₂ was performed using permeation tubes

22 (VICI, Houston, TX, USA) which were maintained at a constant temperature of 23 °C and weighed

23 every 2 weeks. Dilutions of the permeation tube effluent were made in ultra-high-purity N_2

(>99.995 %) with flow rates of 50 to 700 ml min⁻¹, and samples were injected into the purge and 1 2 trap system (VSP) through a 140 µl loop. Standard deviation of the peak area during these calibration runs was <5 % for CH₃I and CH₂ClI and <15 % for CH₂I₂. Overall the calibration 3 4 response varied by less than 15 % over the entire sampling period. 5 Throughout the paper, seasons are defined as follows: summer is June through August; fall is 6 September through November; winter is December through February and spring is March to May. 7 8 3. Results and discussion 9 3.1 Environmental Variables from the Bedford Basin 10 The vertical profiles of temperature, salinity, dissolved oxygen and fluorescence (Fig. 2) are wellmixed from top to bottom in late winter (Feb-Mar) as a result of wind-mixing and convection (Li, 11 12 2001). Temperature is marked by strong seasonality to depths of <30 m. Near-surface temperatures start to rise above winter values of 4 °C, and stratified conditions develop, around early April with 13 temperatures reaching ca. 21 °C by the end of August (Fig. 2a). 14 15 Salinity ranges from 23 to 31 through the entire water column, with the lowest salinities occurring very close to the surface (Fig. 2b and Fig. 3c). The near-surface stratification varied both seasonally 16 and between years, primarily in association with variability of precipitation and the discharge of 17 18 the Sackville River (source: Environment and Climate Change Canada; http://climate.weather.gc.ca/historical data/search historic data e.html). example, 19 For the salinity difference between 1m and 5m was >1 during much of the summer of 2015 (June to 20 21 September) and summer 2017 (June to August). In summer 2016, however, the salinity at 1m was close to that at 5m (difference < 0.3) (Fig. 3c). Occasional intrusions of more dense water from the 22

23 Scotian Shelf, results in increased salinity, especially of bottom waters. The intrusions are irregular

and tend to occur a few times per year, for instance in May 2016 at which time the salinity of
bottom water increased from 30.8 to 31.0 (Fig. 2b), and in early July 2017 when the salinity of
mid-depth water increased from 30.5 to 31 (see marked circle in Fig. 2b).

4 The dissolved oxygen time-series (Fig. 2c) shows the effect of temperature-dependent solubility 5 variations in surface waters as well as intrusions and late-winter vertical mixing in deeper water. 6 In surface water the highest O₂ concentrations occurred between March and April every year in association with lowest seawater temperature. The vertical gradient of O₂ concentration was, 7 generally, smallest towards the end of April as a result of vertical mixing. Sub-surface O₂ 8 9 concentrations (>30 m) generally decreased in summer due to respiration, with occasional interruptions of this O₂ decline (e.g. November 2016) as a consequence of shelf-water intrusions 10 11 which brought sudden increases in O₂ levels.

12 Fig. 3 depicts time-depth plots of the variation of chlorophyll *a*, total dissolved inorganic nitrogen $(DIN = [NH_4^+] + [NO_2^-] + [NO_3^-])$, salinity, precipitation, windspeed and solar irradiance in 13 14 Bedford Basin over the period of the VOIs sampling. The seasonal variations of chlorophyll a concentration in surface water (Fig. 3a) show that two blooms (spring and autumn) occur in surface 15 16 water. For example, in 2016, chlorophyll a increased rapidly from March to April (from 5 to 26 μg L⁻¹), and from September to October (from 10 to 28 μg L⁻¹). The vertical variation of 17 chlorophyll a (as determined from fluorescence measured on the CTD, see Fig. 2d) reached 12 µg 18 L^{-1} during the bloom period. Sub-surface (20- 40 m) fluorescence-derived chlorophyll *a* dropped 19 down to 4 μ g L⁻¹. In the near-bottom water chlorophyll *a* ranged between 0 and 2 μ g L⁻¹ during 20 21 the whole year and varied only slightly.

The seasonal variation of dissolved inorganic nitrogen(DIN) in surface water is plotted in Fig. 3b.
In winter, when chorophyll *a* levels are very low due to light limitation, DIN concentrations reach

ca. 12 μ mol L⁻¹ but are drawn down to low levels (< 1 μ mol L⁻¹) after the spring bloom. 1 2 Summertime chlorophyll *a* levels are moderate but variable (ca. 3 to 10 μ g L⁻¹), likely reflecting continuing nutrient input (e.g. from runoff and/or sewage treatment plants). The average 3 precipitation in Bedford Basin was 27.6 mm week⁻¹ in summer 2015 and was 16 mm week⁻¹ in 4 summer 2016 (Fig. 3d). Typically, strongest irradiance (data was downloaded from the CERES 5 6 FLASHFLUX system: https://power.larc.nasa.gov/cgi-bin/hirestimeser.cgi) occurs in June and 7 July (see Fig. 3f), and highest water temperatures are observed in August. 8 3.2 Variations of Iodocarbons Concentrations in Bedford Basin

9 Iodocarbon concentrations in surface water (1, 5 and 10 m) showed strong seasonality, with lowest concentrations from December through May (1.2 pmol L⁻¹ for CH₃I; 1.3 pmol L⁻¹ for CH₂CII and 10 0.3 pmol L⁻¹ for CH₂I₂). Concentrations start to increase in late May/ June, reaching levels as high 11 as 45 pmol L⁻¹ for CH₃I; 160 pmol L⁻¹ for CH₂ClI and ca. 80 pmol L⁻¹ for CH₂I₂ (with a single 12 peak of 500.5 pmol L⁻¹; Fig. 4). Near-surface, summertime concentrations of all three compounds 13 were characterized by a broad seasonal peak of 6-7 months duration (or shorter for CH₂I₂), on top 14 of which were superimposed ca. 3-4 peaks of ca. 1 month duration. The number, amplitude and 15 timing of these peaks varied amongst the three compounds with CH₃I, notably, showing only one 16 17 large peak in 2016 and four during the other two years of the time-series (Fig. 4a).

Concentrations at 60 m were almost always lower, and much less variable, ranging over the year from 1 to 9 pmol L⁻¹ for CH₃I (except the Fall/Winter 2015-2016, see below), 1 to 6 pmol L⁻¹ for CH₂CII and 0.4 to 18 pmol L⁻¹ for CH₂I₂ (Fig. 4d) respectively. Hence, the bottom water (60 m) concentrations of CH₂I₂ and CH₂CII were always much lower than in near-surface waters throughout the summers. The surface to deep concentration difference was smallest for CH₃I and showed interannual variability. Notably bottom water concentrations reached 26 pmol L⁻¹ and were even higher than in contemporary surface waters from September 2015 to March 2016 (Fig. 4d).
 Missing from the bottom water time-series, were the ca. 1 month duration variations seen in
 summertime surface water.

4 Inter-relations between the iodocarbons in surface seawater were examined with linear regression 5 of both weekly and monthly-averaged concentrations. The resulting correlations are shown in 6 Table 1. Using weekly data, significant correlations (i.e. p<0.05) were found between [CH₃I] and 7 [CH₂CII] at 1, 5 and 10 m depths with the strongest correlation (R= 0.7) at 10m. The only other significant correlation was between CH₂I₂ and CH₂ClI at 5 m. Use of monthly averaged values 8 9 gave stronger correlations. Once again, the significant correlations were between CH₃I and CH₂CII 10 (at 1, 5 and 10m depth) as well as between CH₂ClI and CH₂I₂ at 5 and 10m depth. Table 2 also 11 presents the correlations of iodocarbon concentrations with potentially related variables (discussed 12 in section 4.3).

Generally, the concentration of CH_2I_2 was higher than that of CH_2CII . The average ratio of CH_2I_2/CH_2CII within the top 10m of the water column over the summer months was 1.4. However, this ratio was significantly lower at 1m depth (average of 0.6) and increased with depth (1.5 at 5 m and 2.2 at 10 m, reaching values as high as 2.7 at 60 m).

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18 3.3 Sea-to-Air Flux

Using the concentrations of CH₃I, CH₂ClI and CH₂I₂ at 1m depth (Fig. 4) we estimated the sea-toair flux of VOIs (F) using the following equation and the parameterization of Nightingale et al.,
2000) for the water-side transfer velocity:

22
$$Flux = K(C_{aqu} - C_{air} \times H)$$
(1)

23
$$K_w = \left(\frac{s_c}{660}\right)^{-0.5} \left(0.222 \cdot u_{10}^2 + 0.333 \cdot u_{10}\right)$$
(2)

where u₁₀ is windspeed at 10 m height, Sc is the temperature-dependent Schmidt number, as
estimated by Groszko (1999) and Henry's Law constants (H) were from Moore et al. (1995). The
air-side resistance has been shown by Archer et al. (2007) to be significant for soluble gases such
as CH₂ClI and CH₂I₂. Hence K in equation 1 is calculated as follows (Liss and Slater, 1974):

$$\frac{1}{K} = \frac{1}{K_w} + \frac{1}{HK_a} \tag{3}$$

6 where the air-side transfer velocity was calculated according to Duce et al. (1991).

7
$$K_a = u_{10}/(770 + 45(MW)^{1/3})$$
 (4)

8 with MW being the molecular weight of the gas of interest.

9 Daily averaged wind speed was measured at the nearby Halifax Dockyard (Fig. 1) (source:
10 Environment and Climate Change Canada; <u>http://climate.weather.gc.ca/index_e.html</u>). Seawater
11 temperature and surface iodocarbon concentrations were interpolated linearly between the weekly
12 measurements in order to coincide with the wind speed data and generate daily flux estimates.

13 Following Archer et al. (2007) and Shimizu et al. (2017), we applied an atmospheric concentration of zero for calculating the flux of all three compounds. Rasmussen et al. (1982) reported an average 14 15 atmospheric mixing ratio of CH₃I of ca. 1 pptv for Cape Meares (45 °N) and Yokouchi et al. (2008) presented a mean concentration of 0.98 pptv for Cape Ochiishi (43.2 °N), with both sites sharing 16 a similar latitude to our sampling location (44.69 °N). If a mixing ratio of 1pptv had been used for 17 18 our calculations, the total annual flux of CH₃I would be reduced by only 5 %. Atmospheric mixing ratios of CH₂ClI and CH₂I₂ are generally lower (reviewed by Carpenter, 2003) so that any 19 overestimation of the fluxes of these compounds due to this assumption will certainly be negligible 20 21 (Archer et al., 2007).

Fig. 5 presents the weekly averaged combined flux of organically-bound iodine, F_{Iorg} , as stacked bar charts, where $F_{\text{Iorg}} = (F_{\text{CH3I}} + F_{\text{CH2CII}} + 2*F_{\text{CH2I2}})$. The calculated emissions of individual 1 compounds ranged from 0.9 to 39.2 nmol m⁻² d⁻¹ for CH₃I (annual average of 8.4 nmol m⁻² d⁻¹), 2 0.9 to 78.0 nmol m⁻² d⁻¹ for CH₂CII (annual average of 17.4 nmol m⁻² d⁻¹) and 0.3 to 78.0 nmol m⁻ 3 2 d⁻¹ for CH₂I₂ (annual average of 10.3 nmol m⁻² d⁻¹). Seasonal and annual average fluxes of the 4 individual compounds and of I_{org} are presented in Table 3. Clearly the sea-to-air flux is highest in 5 summer and fall and is dominated by the flux of the dihalomethanes rather than CH₃I.

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8 3.4 Net Production of CH₃I

9 We used a mass balance approach to estimate the production rate of CH₃I within the uppermost
10 10m of the water column, based on the concentration time-series (see also Shi et al., 2014b). Using
11 the annual cycle of average near-surface CH₃I concentrations (Fig. 4a), we examined the mass
12 balance of CH₃I for this fixed depth interval according to:

$$\Delta C = P_{net} - L_{sea-to-air} - L_{SN2} - L_{mix}$$
(5)

where ΔC is the daily change of the average CH₃Iconcentration in the near-surface seawater (0-10 14 m); P_{net} represents the net of gross production minus any additional, uncharacterized losses such 15 as microbial degradation; $L_{sea-to-air}$ is the sea-to-air flux (section 3.3) and L_{SN2} is the 'chemical' loss 16 due to nucleophilic substitution of Cl⁻ for I⁻ which was calculated based on reaction kinetics (Elliott 17 and Rowland, 1993; Jones and Carpenter, 2007) using the corresponding temperature, salinity and 18 mean concentration of CH₃I. L_{SN2} averaged 0.1 pmol L⁻¹ day⁻¹. L_{mix} is the loss due to downward 19 20 mixing and has been shown in several studies to be negligible compared with other loss terms e.g. (Richter and Wallace, 2004). The latter assumption will not always be valid in winter and, 21 especially, when mixed layers deepen to >10m. However most production of iodocarbons occurs 22 23 during summer and fall when there is strong stratification within the upper 10m (see Fig. 3c), and we excluded from our mass balance calculations the limited periods of time when density was
 uniform in this depth interval.

3 The net production rate of CH₃I over the 3-year time-series is shown in Fig. 6. The annual average production rate of CH₃I was 1.0 pmol L⁻¹ day⁻¹ (ranging from -1.6 to 8.5 pmol L⁻¹ day⁻¹). A 4 significant peak of P_{net} occurred from August to September in every year. The net production rate 5 of CH₃I in summer and fall averaged 1.6 pmol L⁻¹ day⁻¹ and was 5 times larger than wintertime 6 rates (ca. 0.3 pmol L⁻¹ day⁻¹). Net production rates for CH₂ICl and CH₂I₂ were also calculated 7 (results not shown), with typical summertime values being 3.2 and 1.3 pmol L^{-1} day⁻¹, respectively. 8 9 Photolytic loss can be very significant for these compounds and has not been estimated, so these P_{net} values represent the net of gross production minus uncharacterized losses including photolysis 10 11 as well as microbial degradation.

12

13 4. Discussion

In the following we discuss the Bedford Basin data in comparison with other studies that have reported concentrations of multiple iodocarbons and especially those that have reported time-series covering an annual cycle (see citations in the introduction). All of these time-series are from midlatitude (40-60°N) nearshore or continental shelf environments subject to strong seasonal variations of light, temperature and biological productivity. There are no reported time-series of seawater concentrations from low latitudes.

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21 4.1 Potential influence of nearshore and /or macroalgal sources

The potential of nearshore macroalgae to cause elevated coastal iodocarbon concentrations has
been mentioned in a number of studies (Giese et al., 1999; Manley and delaCuesta, 1997; Schall

et al., 1994). We investigated this in July 2017, by sampling at 5 nearshore sites around Bedford 1 2 Basin (Fig. 1) and comparing nearshore concentrations with values measured at the regular 3 sampling site in the center of the Basin (Table 4). The nearshore results were consistently within 4 one standard deviation of mean concentrations of VOIs measured at the center of Bedford Basin during July, indicating no significant difference. Klick (1992) also compared measurements on 5 6 samples collected directly over a rich bed of macroalgae with samples collected further away from 7 direct contact with macroalgae: whereas they observed significantly higher concentrations of 8 bromocarbons in proximity to the macroalgae, there was no difference observed for CH₂I₂ and 9 CH₂CII. Shimizu et al. (2017) sampled a number of nearshore regions around Funka Bay, including rocky shores with extensive macroalgae, and also found concentrations to be similar at both 10 11 nearshore and central Bay locations. We therefore conclude that any direct impact of macroalgae 12 on measured organoiodine levels is small, even in coastal regions, which lends strong support to the conclusion by Saiz-Lopez and Von Glasow (2012) that macroalgae are only a minor global 13 14 source of these compounds to the atmosphere.

15

16 4.2 Concentrations and relative abundance of iodocarbon compounds

The average concentration of total volatile organic iodine I_{org} (where $I_{org} = [CH_3I] + [CH_2CII] + 2[CH_2I_2]$) and the relative contributions of the different compounds to I_{org} from this and other studies is shown in Fig. 7. The combined concentrations of the three iodocarbons are highest but also show highest variability ($[I_{org}] = 25$ to 281 pmol L⁻¹) in summertime coastal waters (loosely defined here as within a few kms of land). Continental shelf waters have lower concentrations of I_{org} averaging 32 pmol L⁻¹, with open ocean waters having comparable or lower concentrations (average $I_{org} = 17$ pmol L⁻¹). Despite the differences in I_{org} concentration ranges evident for different regions in Fig. 7, a 1-way ANOVA showed no significant differences between the means
 for the three regions. However after pooling of shelf and open ocean results to make only two
 populations, a t-test did reveal a significant difference with coastal waters for at the 95%
 confidence level.

5 The distribution of I_{org} is contrary to the global distribution of CH₃I reported by Ziska et al. (2013) 6 who noted a tendency for the open ocean to have higher concentrations than coastal waters (their 7 definition of "coastal" was within 1 degree latitude or longitude of land and therefore much broader 8 than ours). As noted by Ziska et al. (2013), this may reflect higher CH₃I concentrations in tropical 9 and sub-tropical open ocean waters, as their general pattern was reversed in the Northern 10 Hemisphere. The coastal waters depicted in Fig. 7 are largely from mid-latitudes of the Northern 11 Hemisphere.

The relative contribution of the dihalomethanes to I_{org} also appears to vary between regions, with the ratio of dihalomethane-I to I_{org}, ([CH₂CII]+2*[CH₂I₂])/[I_{org}], averaging 0.71, 0.69 and 0.55 in coastal, shelf and open ocean waters, respectively. While an elevated contribution of dihalomethanes in coastal waters is consistent with the report by Jones et al. (2010), regional differences apparent in Fig. 7 are not significant at the 95% confidence level when tested with a 1way ANOVA.

Klick (1992), Jones et al. (2010) and Shimizu et al. (2017) reported concentrations of volatile
organic iodine in summertime coastal waters that are comparable to, or higher than those observed
in Bedford Basin (i.e. average I_{org} concentrations >100 pmol L⁻¹). Our results from Bedford Basin
correspond closely with concentrations and relative contributions reported by Shimizu et al. (2017)
for coastal water in Funka Bay, Japan (Fig. 7). In these coastal surface waters, the CH₂I₂
concentration and contribution was highest on average, followed by CH₂CII and the lowest was

1 CH₃I. In open ocean waters, the relative contribution of [CH₃I] to I_{org} appears higher, reaching 2 over 50% in some cases (see Fig. 7), with the contribution of CH₂I₂ generally being lower in the 3 open ocean than in coastal waters. However, once again, these apparent regional differences are 4 not significant at the 95% confidence level.

5 In laboratory studies, Fuse et al. (2003) demonstrated that relatively large amounts of CH_2I_2 and 6 I₂ together with smaller but still significant amounts of CH₂ClI and CHI₃ can be produced, 7 presumably abiotically, in dark incubations of (filtered) spent culture media with suspended bacterial cells and added [I⁻]. The CH₂I₂/CH₂ClI production ratio was ~35 and no mono-iodinated 8 9 CH₃I was produced in these experiments. The implication was that dissolved organic compounds within spent media were key to production of polyiodinated compounds. In the absence of spent 10 11 culture media, additions of oxaloacetic acid also resulted in formation of CH₂I₂ and CH₂ClI (with 12 a lower ratio of CH_2I_2/CH_2CII of ~10) suggesting that organic acids may be a substrate for their 13 formation. The mechanistic role of the suspended bacterial cells was not clear, however they may 14 have supplied haloperoxidases required for oxidation of I (see also Hill and Manley, 2009). Martino et. al (2009) demonstrated that, alternatively, oxidation of dissolved iodide to I₂ and HOI 15 16 by reaction with ozone (e.g. Garland et al., 1980) in filtered (0.2 µm) seawater containing natural levels of dissolved organic matter also resulted in formation of polyiodinated compounds (CH₂I₂, 17 CH₂ClI and CHI₃) with CH₂I₂/CH₂ClI production ratios ranging from 2 to 4. They suggested that 18 the yield of various iodocarbons depends on "the abundance and perhaps on the nature of the 19 20 organic substrate" which "can vary widely both temporally and spatially". We could not, however, 21 find any obvious relationship of near-surface iodocarbon concentrations with local measurements 22 of atmospheric ozone near Bedford Basin (results not shown).

We therefore suggest that the higher levels of CH₂I₂ observed in coastal waters, including Bedford 1 2 Basin, reflect a higher supply rate of HOI and/or I₂ and/or of the organic precursors suitable for 3 formation of polyiodinated compounds. Ultimately, reduction of seawater iodate to reduced forms, 4 such as iodide, likely supports the potential for organoiodine formation. Addition of iodide has 5 been identified, in short-term experiments, as a source for reactive iodine (iodine atoms) which 6 can stimulate photochemical formation of CH₃I (Moore and Zafiriou, 1994). However, Shi et. al 7 (2014b) found no positive correlation of [I⁻] with seasonal CH₃I production in a field study in Kiel 8 fjord, noting that background iodide levels may have been always sufficient to support production 9 of pM levels of CH₃I. Indeed, there is no *a priori* reason to expect a positive correlation of $[I^-]$ with 10 iodocarbon production if supply of more reactive iodine species (iodine atoms, HOI and/or I₂) is 11 the key, proximate control. The short-term formation rate of reactive species from a background 12 reservoir of I, and hence production of iodocarbons, may depend more on availability of 13 haloperoxidases, various oxidants and/or temperature-dependent kinetics. The conversion of I⁻ to 14 these reactive species could even lead to inverse correlations between I⁻ and iodocarbons.

Relatively small quantities of CH₂ClI were produced in several of the experiments cited above, 15 16 yet observations in Bedford Basin show average CH₂I₂/CH₂ClI ratios of 1.4 in the top 10 m of the 17 water column. Production ratios in these experiments vary, as noted above, but laboratory studies 18 have also shown that photolysis of CH₂I₂ can be an important source of CH₂ClI in surface waters 19 with a yield of 25 % to 35 % (Jones and Carpenter, 2005; Martino et al., 2005). We observed 20 significant correlation between [CH₂I₂] and [CH₂ClI] at 5 and 10 m depth (but not at 1m) (Table 21 1), which is consistent with a fraction of the CH_2I_2 production being transformed photochemically 22 (The lack of correlation at 1m may be due to the very rapid photolysis). However the correlation 23 may also reflect the original production ratio of the individual compounds (which laboratory

experiments suggest may be substrate-dependent). In both cases however, dissolved organic matter
 (DOM) quality and quantity (possibly associated with terrestrial supply) and/or elevated supply of
 I⁻ are likely to be underlying reason(s) for the high concentrations of dihalomethanes observed in
 Bedford Basin and other coastal waters.

5

6 4.3 Temporal variations of iodocarbons in near-surface water

7 The following discussion of temporal variability is separated into consideration of seasonal and8 interannual variations.

9 4.3.1 Seasonal Variations

All of the reported iodocarbon time-series showed strong seasonality, with minimum, sometimes 10 11 undetectable concentrations in winter, and higher concentrations in summer. Near-surface (0-10 12 m) concentrations of all three iodocarbons in Bedford Basin, including CH₃I, remained low until 13 mid-May to mid-June, with their subsequent increase coincident with initial warming of near-14 surface waters from wintertime minimum temperatures of ca. 1-2 $^{\circ}$ C (lag < 1 month; Fig. 8a). 15 Hence the initial appearance of all three iodocarbons occurred more than 3 months after the 16 seasonal increase in solar radiation, ca. 1-2 months after the Spring Bloom (Fig. 8d), after near-17 surface nitrate had been drawn down to low levels (Fig. 8e) and almost coincident with the seasonal temperature increase (Fig. 8a). 18

In the western English Channel (Archer et al., 2007), a gradual increase of CH₃I commenced in February, coincident with the seasonal increase in solar radiation. Summertime values remained high, with some higher-frequency variation, and then decreased in September/October. The increase of CH₂ClI and CH₂I₂ started later, in April, more or less coincident with both the Spring Bloom and initiation of near-surface warming from a wintertime minimum temperature of ca. 8 °C. Summertime values of CH₂ClI and CH₂I₂ showed periodic variations similar to those observed
 in Bedford Basin (section 3.2).

The lower temporal resolution of the study in Funka Bay (Shimizu et al., 2017), with sampling only every 1 or 2 months, precluded detailed examination of timing. A gradual increase in CH₃I appeared to start in March, during or towards the end of the Spring Bloom when surface water temperatures were still close to their wintertime minimum of -1 to 2.5 °C. The seasonal increase of CH₂I₂ and CH₂CII occurred later (May-June) at a time of rising water temperatures and low nutrient levels with concentrations remaining elevated through the summer and decreasing to wintertime levels in October.

The initial CH₃I increase at a shallow station in the Kiel Fjord (Shi et al., 2014b) occurred in March, and was closely linked in time to seasonal increases of solar radiation, temperature (winter minimum of 0 °C) as well as Chl *a* and the springtime drawdown of nitrate. Lagged correlation analysis showed similarly strong correlations of CH₃I with both temperature and solar radiation, with the annual cycle of CH₃I lagging temperature by ca. 1 month, however the very close correspondence of multiple seasonal cycles led the authors to note that "the use of correlation analysis to infer causality has likely reached its limit in this analysis".

The observation of a rapid increase in the production rate of I⁻ within phytoplankton cultures (diatoms and prymnesiophytes) when they enter stationary and, especially, senescent phases (Bluhm et al., 2011) is potentially relevant to the observed seasonality of iodocarbon formation. The reduction of iodate to iodide was suggested to be due to release of precursors, such as reduced sulphur species, to surrounding culture medium in association with a loss of membrane integrity by stressed cells or as a result of viral lysis. Hughes et al. (2011) also reported studies with cultures of *Prochlorococcus marinas* in which accumulation of CH₃I commenced when cultures became senescent. We note that significant iodocarbon accumulation in Bedford Basin was confined to
 summertime when DIN was depleted (see Fig. 3b) and when cells may have been stressed or
 subject to viral lysis, perhaps similar to later stages of batch culture experiments.

4 We therefore hypothesize that seasonal nitrate drawdown leads to increased supply of iodide to 5 surface waters which can, in turn, lead to increased formation of iodine atoms, HOI and I₂ as 6 precursors for iodocarbon formation by both photochemical and haloform reaction pathways 7 (Martino et al., 2009; Moore and Zafiriou, 1994). Whereas the supply of iodide may be one key control, it is likely that variations in light intensity and water temperature also contribute to the 8 9 overall seasonality of the production rate of CH₃I (e.g. through temperature influence on reaction 10 kinetics). For example, light can influence formation of CH₃I directly (e.g. Moore and Zafiriou, 11 1994; Richter and Wallace, 2004). Light can also influence iodocarbon production indirectly, for 12 example by producing oxidants such as H_2O_2 to promote oxidation of iodide by haloperoxidases 13 (Hill and Manley, 2009) or by altering the quality of dissolved organic matter. The time-series of 14 CH₂I₂ and CH₂ClI from very shallow (< 4 m), nearshore waters of the Kattegat, Sweden (Klick, 1992) and the Baltic Sea, Germany (Orlikowska and Schulz-Bull, 2009) showed peaks in April/ 15 16 May and again in September/October, with low concentrations throughout summer. This contrasts 17 with the deeper water columns of Bedford Basin, Funka Bay and the English Channel where 18 concentrations remain elevated throughout summer. This likely reflects dominance of photolytic 19 loss over production within very shallow water columns exposed to summertime light intensities 20 and long periods of daylight. Sub-surface production coupled with vertical mixing may explain 21 the summertime persistence in deeper water columns.

In addition to the broad seasonal variation, a number of maxima with duration of ca. 1 month, wereobserved, and appear similar to short-period fluctuations observed in the English Channel time-

series (Archer et al., 2007). There does not appear to be any consistent pattern linking the maxima
 and minima of the three compounds in the two studies, so that the underlying causes for these
 shorter period variations are unclear.

4

5 4.3.2 Interannual variability

6 The Bedford Basin time-series is unique in having high temporal resolution sampling (weekly) 7 over three annual cycles which allows interannual variability to be examined for the first time. The most obvious interannual difference was in the behavior of CH₃I. In particular, 2016 was markedly 8 9 different in that only a single peak was observed in late August, whereas the summers of 2015 and 10 2017 were marked by 3-4 quasi-periodic, multi-week maxima. As noted already, the English 11 Channel time-series of CH₃I did not exhibit this behavior. The shallow-water time-series of CH₃I 12 in the Kiel Fjord and coastal Baltic Sea (Orlikowska and Schulz-Bull, 2009; Shi et al., 2014b) also 13 did not exhibit this type of variability. Because the cause of the periodicity itself is not understood 14 or explained, discussion of reasons for its interannual variation must be highly speculative. One clear difference of 2016 relative to the other two years, was the lower summertime precipitation 15 16 and associated lack of near-surface salinity stratification. The temporal behavior of CH₃I in 2016 17 might therefore be related to altered near-surface mixing dynamics within Bedford Basin, or 18 alternatively, to decreased delivery of key precursors (e.g. DOM) from land via rivers and 19 wastewater.

20

21 4.4 Vertical distributions and subsurface temporal variability

Fig. 4 shows the near-surface concentration variations of the VOIs. For CH₃I, concentrations were almost always uniform between 1, 5 and 10 m. For CH₂ClI, the concentrations at 1 and 5m were

usually very similar (average difference -4.1 %; median -2.5 %), however concentrations at 10m
depth were noticeably lower for periods of time. For CH₂I₂, the highest concentrations were
observed at a depth of either 5 or 10 m, with concentrations at 5 m occasionally peaking at very
high levels (e.g. 250-350 pmol L⁻¹) for short periods (less than one week). Concentrations at 1 m
were almost always lower than at 5 m, with the percentage reduction relative to 5 m averaging 52 %
in summer. Concentrations at 10 m, on the other hand, were generally the same or higher as those
measured at 5 m (with the exception of the previously mentioned, short-lived peaks).

These results are consistent with earlier studies of vertical profiles in the open ocean (e.g. Moore and Tokarczyk, 1993; Yamamoto et al., 2001) as well as with model predictions (Jones et al., 2010; Martino et al., 2006). In particular, our results are consistent with the quantitative predictions of a mixed-layer model (Jones et al., 2010) that CH₂ClI would typically be near-uniform within the upper 6 m of the water column, whereas photolytic decay could remove up to 100 % of the CH₂I₂ over that depth range, depending on time of day and conditions.

14

15 4.5 Temporal variability in near-bottom water (60 m)

The time-series of VOIs in near-bottom waters (60 m) are presented in detail in Fig. 9a,b,c, with specific events labelled 1 through 9. Variability was generally of lower amplitude than in surface waters, except for CH₃I during the winter of 2015-2016 (Fig. 4d). From June to December 2015, [CH₃I] increased steadily (concentration change, $\Delta C = 20$ pmol L⁻¹) (event 1 to 3; Fig. 9a), exceeding surface water concentrations from October 2015 until the end of March 2016. No comparable increase was observed during 2016 (Fig. 9b), and a smaller increase ($\Delta C = 8$ pmol L⁻¹) was confined to the early summer of 2017 (Fig. 9c).

Concentrations of CH₂ClI remained almost constant at <5 pmol L⁻¹ throughout, with the notable 1 2 exception of abrupt (<1 week) increases in May and November 2016 (events 4 and 5; Fig. 9b) and 3 December 2017 (event 9; Fig. 9c). These increases ($\Delta C = 2-5 \text{ pmol } L^{-1}$) coincided with sudden 4 increases of salinity (and O₂) and reflect intrusion of saltier, near-surface waters from offshore, 5 rather than local production. The subsequent concentration declines reflect loss due to mixing or, 6 more likely, reaction and/or microbial degradation within the water column and sediments. The 7 same three intrusions also drove abrupt increases of CH_2I_2 with amplitude ca. 1.5 – 2 times higher than those for CH₂ClI, consistent with near-surface concentration ratios (see section 3.2). However, 8 9 CH₂I₂ also showed higher-amplitude variations unrelated to the bottom water intrusions (see below). 10

11 The increase of CH₃I from June through October 2015 (event 1 through 3; Fig. 9a), paralleled a 12 steady decline in oxygen, suggesting that production was linked to degradation of organic matter. The accumulation rate of ca. 0.06 pmol L⁻¹ day⁻¹ was 20 times smaller than typical P_{net} for CH₃I 13 14 in surface waters (see section 3.4). The increase appears consistent with results from short-term (3-day) incubation experiments with biogenic marine aggregates reported by Hughes et al. (2008) 15 16 in which concentrations of mono-iodinated iodocarbons, including CH₃I, increased but with no 17 corresponding increase in dihalogenated compounds. Their results suggested alkylation of 18 inorganic iodine or breakdown of higher molecular mass organohalogens as production pathways 19 and, following Amachi et al. (2001), they suggested that microbial degradation increased the 20 supply of precursors.

However, as O₂ concentrations declined further from October through late December (event 2 to
3; Fig. 9a), the concentration of CH₃I stabilized and CH₂I₂ concentration increased markedly from
2 to 12 pmol L⁻¹. From January through April 2016, CH₃I levels decreased (following event 3; Fig.

9a,b), in concert with increasing O₂ concentrations and decreasing salinity, reflecting the effects
of progressive vertical mixing with overlying waters which had lower CH₃I concentrations. Over
this same period, CH₂I₂ concentrations at 60 m remained almost constant (Fig. 9a, b) due to the
smaller vertical concentration gradient, until increasing due to an intrusion (event 4). This was
followed by a gradual decrease over summer months, in parallel with the seasonal decrease of O₂
(fig 9b).

The data from 2015 and 2017 are suggestive of a "switch" of production mechanism from an alkylation pathway producing mono-iodinated compounds (CH₃I) to a haloform-type reaction producing CH₂I₂. The "switch" took place in October 2015 (event 2; Fig. 9a), when oxygen concentrations dropped below 90 µmol kg⁻¹, although whether the switch was related to redox conditions in the water column or sediments, speciation and availability of iodine, or availability of suitable organic precursors and/or enzymes cannot be determined.

There was almost no near-bottom iodocarbon production during 2016 (Fig. 9b), and therefore no "switch", for reasons that are not clear given that O₂ declined through summer and fall, until interrupted by an intrusion in November (event 5; Fig. 9b). We speculate that the lack of production in near-bottom water might be linked to the relatively short period of CH₃I production in nearsurface waters during that year (fig 4a).

In 2017, there was moderate sub-surface production of CH_3I , associated with O_2 consumption (April through July; event 6 to 7), and, again, an apparent "switch" to CH_2I_2 production marked by plateauing of CH_3I concentrations (event 7) at the same time as CH_2I_2 concentrations started to increase (Fig. 9c). This was followed by a subsequent decrease to background levels over a period of about a month. The apparent "switch" in production took place earlier in the year and at higher O_2 concentrations (175 µmol kg⁻¹) than in 2015.

However close inspection of Fig. 2 shows that the plateauing of CH₃I in near-bottom waters (event 1 2 7) also coincided with the appearance of a mid-depth intrusion of saltier water (as denoted by the 3 31 salinity contour; see blue circled area in Fig. 2). The same period (between events 7 and 8) was 4 marked at 60m by a change from declining to increasing salinity, reduction in the rate of oxygen 5 concentration decline (Fig. 9c) and an increase in the rate of warming (not shown). The mid-depth 6 salinity maximum in Bedford Basin must reflect intrusion of saltier water from offshore. The 7 subsequent trends in temperature, salinity and dissolved oxygen at 60m, between events 7 and 8 (Fig. 9c), are consistent with mixing of pre-existing near-bottom water with this intrusion. It 8 9 therefore cannot be ruled out that mixing with this intrusion contributed in some way to the nearbottom increase of CH₂I₂, plateauing of CH₃I concentrations as well as the small but significant 10 increase in CH₂ClI (Δ C=1-2 pmol L⁻¹), and hence to the apparent "switch", which all occurred at 11 the same time. 12

At the end of this period, in mid-August (event 8), the rate of warming and salinity increase at 60 m decreased again and the rate of oxygen decline increased (Fig. 9c), suggesting that the intrusion's impact had lessened. At this time, CH₂I₂ and CH₃I concentrations started to return to background levels with estimated half-life, assuming 1st order kinetics, of ca. 65 days (CH₂I₂) and 14 days (CH₃I). A decrease in [CH₂CII] started about a month later with 1st order half-life of ca. 70 days. Similar rates of decline of [CH₂CII] and [CH₂I₂] were observed following sudden concentration increases associated with other intrusions, discussed earlier.

This very detailed discussion of the temporal variability emphasizes that a variety of physical and biogeochemical mechanisms can contribute to interannual, seasonal, and short-term variability of the three iodocarbons. The high amplitude variability observed in Bedford Basin could prove useful for validation of models representing complex iodine cycling as well as physical mixing dynamics. However, separation of multiple potential contributing factors and processes underlying
 temporal variability may require a data set with higher vertical resolution and, ideally, a seasonally resolved time-series of experiments (compare Shi et al., 2014a).

4

5 4.6 Sea-to-air fluxes

6 The temporal variation of the sea-to-air flux of I_{org} and the relative contribution from the three 7 iodocarbons are shown in Fig. 5 and Table 3. Similar to the findings of Archer et al. (2007), air-8 side resistance leads to significant reductions in calculated, annual average fluxes for CH₂ClI and 9 CH₂I₂ of 10% and 24%, respectively, relative to calculations when it is ignored. The study by 10 Shimizu et al. (2017) did not consider air-side resistance, so Table 5 presents both their original 11 reported fluxes, as well as fluxes adjusted for its likely impact based on our study and that of 12 Archer et al. (2007). The following discussion makes use of the adjusted fluxes.

Consistent with earlier time-series (excluding those from very shallow waters, see section 4.2), the sea-to-air flux of iodocarbons is generally highest in summer/fall. However, high wintertime fluxes are also possible, as seen in 2017 when there was a large efflux of CH_2I_2 (averaging 9.1 nmol m⁻² d⁻¹; Table 3), due to both strong winds and relatively high concentrations. The fluxes of CH_3I and CH_2CII , on the other hand, were always higher in summer/fall (ca. 3-5 times and 10 times higher, respectively). Similar findings were presented by Shimizu et al. (2017) with the total iodine flux in Funka Bay in summer being > 4 times that in winter.

Our estimated emissions of $CH_{3}I$ (8.4 nmol m⁻² d⁻¹, Table 5) are in the range calculated previously for coastal and continental shelf water in similar latitudes (11.9 and 7.7 nmol m⁻² d⁻¹; Archer et al., 2007; Shimizu et al., 2017 respectively). The average flux of $CH_{3}I$ reported by Jones et. al (2010), from the west of Ireland, was 4 times higher but based on a sampling period of only 1 month during summer. Sea-to-air fluxes of CH₂ClI from Funka Bay and English Channel were
similar to our calculated fluxes from Bedford Basin. However the highest variation is observed in
the annual averaged flux of CH₂I₂, ranging from 3.5 nmol m⁻² d⁻¹ (the west English Channel), 10.3
nmol m⁻² d⁻¹ (Bedford Basin) and 12.6 nmol m⁻² d⁻¹ (Funka Bay, Japan). The total, annual I_{org} seato-air flux from Bedford Basin averaged 46.7 nmol m⁻² d⁻¹, which was approximately 5 times larger
than the flux due to CH₃I alone. The total annual flux was similar between all three locations.

Fig. 5 and Table 3 show that the total I_{org} flux is subject to significant interannual variability, which
could not be assessed by the earlier studies. Notably, the I_{org} flux in 2016 was ca. 2 times smaller
than in 2015 and 2017. A comparison of wind-speeds and concentrations showed that the influence
of wind speed was dominant, due to winds during summer/fall of 2016 being 1-2 m s⁻¹ lower.

11

12 4.7 Production rate of CH₃I

The annual mean production rate (P_{net}) of CH₃I in this study, estimated using equation 1, was 1.0 13 pmol L⁻¹ day⁻¹ (ranging from -1.6 to 8.5pmol L⁻¹ day⁻¹, see section 3.4 and Fig. 6). This is 14 comparable with the global average production rate estimated by Stemmler et al. (2013) (1.64 15 pmol L⁻¹ day⁻¹), for which 70% was produced via a photochemical mechanism. Based on data 16 17 presented by Archer et al. (2007), the annual mean production rate of CH_3I in the western English Channel was ca. 2 pmol L⁻¹ day⁻¹ (range: -0.2 to 6 pmol L⁻¹ day⁻¹). Here it should be noted, that 18 their "minimum gross production rate" is equivalent to P_{net} in this study and in Shi et al. (2014a). 19 20 In contrast, Shi et al. (2014b) estimated a considerably lower annual mean net production rate in the Kiel Fjord of ca. 0.1 pmol L⁻¹ day⁻¹ (maximum of 0.8 pmol L⁻¹ day⁻¹). The maximum production 21 22 rates from the Kiel Fjord study were smaller as they were based on monthly average (and therefore 23 "smoothed") concentrations. However, Shi et al. (2014a) also conducted weekly incubation experiments which gave *in vitro* values of P_{net} which were closely comparable with the field-based
 estimates in Kiel Fjord.

3 The lower values of P_{net} in the Kiel Fjord compared with both Bedford Basin and the English 4 Channel must reflect either differences in gross production (e.g. due to differences in the supply 5 of precursors and reactants such as iodide) or differences in other, uncharacterized losses. Evidence 6 for a poorly characterized loss process, possibly microbial degradation, was in fact observed in the 7 Kiel Fjord incubation experiments (Shi et al., 2014a). On the other hand, incubation experiments conducted with additions of labelled methyl iodide (¹³CD₃I) to Bedford Basin surface waters (data 8 9 not shown) during the course of this study showed no such losses. We therefore hypothesize that the lower P_{net} in Kiel Fjord is a result of higher microbial degradation of CH₃I in that very shallow 10 11 (<12 m), nearshore environment.

12

13 5. Conclusions, Implications and Further Work

14 The 3-year time-series of weekly iodocarbon concentrations from Bedford Basin shows overall seasonality similar to that observed in coastal time-series from both the English Channel and Funka 15 16 Bay, Japan. There was no mid-summer minimum in the concentration of polyiodinated compounds 17 as observed in some time-series from very shallow water (<10m), which likely reflects dominance 18 of photolytic decay in such shallow water columns. Interannual variability in near-surface water 19 concentrations was particularly pronounced for CH₃I, with only a single, short-lived concentration 20 maximum observed in 2016, possibly as a result of anomalously low rainfall and consequently 21 reduced supply of terrestrial organic matter during that summer.

Based on the time-series as well as published lab studies, we hypothesize that seasonal, nearsurface production of iodocarbons is linked to accelerated reduction of iodate to iodide under post-

bloom conditions, following disappearance of nutrients and possibly also influenced by water
 temperature. The observed vertical variation of CH₂I₂ and CH₂ClI is consistent with the more rapid
 photolysis of CH₂I₂.

The average annual sea-to-air flux of total volatile organic iodine (46.7 nmol m⁻² d⁻¹) is almost identical to that observed in Funka Bay, Japan, and the English Channel. The polyiodinated compounds contributed ca. 80 % of the total flux which was similar to that in the other two timeseries and confirms that the sea-to-air flux of polyiodinated compounds dominates in coastal waters. The fluxes were variable on interannual timescales (factor of 2) as a result, mainly, of wind-speed variability.

10 The near-bottom water (60m) time-series was impacted by episodic intrusions of water from 11 offshore and showed evidence for CH₃I production associated with decay of organic matter, albeit 12 with a production rate more than an order of magnitude lower than in surface waters. The timeseries showed evidence for a possible "switch" from CH₃I production (e.g. by alkylation of organic 13 14 matter) to production of CH₂I₂ (e.g. by a haloform type reaction), after periods of about 1 month. The very high amplitude concentration variations encountered in Bedford Basin, coupled with its 15 16 relative accessibility for high-frequency sampling and constrained, yet variable, physical 17 exchanges make Bedford Basin a useful location to investigate iodine cycling. To-date, the 18 complexity of iodine biogeochemistry has hindered progress towards understanding the controls 19 on spatial and temporal fluxes of iodine between the ocean and atmosphere. We suggest that 20 progress can now be made through more comprehensive sampling (higher vertical resolution and 21 inclusion of inorganic iodine speciation measurement), coupled to a biogeochemical model of 22 Bedford Basin that includes iodine chemistry (e.g. Stemmler et al, 2013) and a time-series of 23 experimental studies conducted in the context of the time-series (see Shi et al., 2014a). In other

1	words, Bedford Basin may provide an ideal location and time-series upon which to base a multi-
2	investigator campaign to understand environmental controls on volatile iodine cycling and
3	improve its representation in models.
4	
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13	
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Figure 1. (a): Main sampling location (grey star) and near-shore sampling locations 1-5 (1: Tufts
cove; 2: Wrights cove; 3: Sackville; 4: Mill cove; 5: Fairview cove) in Bedford Basin; (b): two
layered flow in Halifax Harbour, adapted from Kerrigan et al. (2017); (c): horizontal circulation
of water in Halifax Harbour from Shan et al. (2011), using annual mean currents and velocities.



Figure 2. Seasonal patterns of environmental and biological variables in Bedford Basin from January 2015 to Dec 2017. (a) temperature; (b) salinity (grey contour line for S=30.5; white contour line for S=31). The blue circle (A) highlights a mid-depth intrusion. (c) dissolved oxygen; (d) chlorophyll fluorescence.



Figure 3. Seasonal variation of (a) chlorophyll *a*: 1 m (blue line) and 5 m (red line); (b) DIN;,(c) salinity in the surface layer: 1m (blue line), 5m (red line) and 10m (black line); (d) weekly precipitation and (e) daily averaged windspeed (shown here only for sampling days); (f) weekly average PAR from January 2015 to December 2017.





Figure 4. Seasonal variation of iodocarbons in the Bedford Basin at 1 m (blue line), 5 m (red line) and 10 m (green line) from May 2015 to December 2017: (a) CH₃I; (b) CH₂CII and (c) CH₂I₂. (d):





Figure 5 Weekly averages of daily sea-to-air flux estimates of I_{org} (see section 3.3): including relative contributions of individual compounds (blue: CH₃I, green: CH₂ClI and orange: CH₂I₂) and using the parameterization for transfer velocity (k_w) of Nightingale et al. (2000) and for the airside transfer velocity (k_a) of Duce et al. [1991].

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Figure 7. Contribution of Iodocarbons to total organic iodine (I_{org}) in surface seawater from
 different regions and studies:

6 1: Klick, 1992; 2: Orlikowski et al., 2015; 3: Jones et al., 2010;4: Jones et al. 2009; 5: Carpenter

7 et al., 2000; 6: Orlikowski & Schulz-Bull, 2009; 7: Shimizu et al.,2017; 8: this study; 9: Archer et

- 8 al., 2007; 10: Hepach et al., 2016; 11: Kurihara et al., 2010.





Figure 8. Annual cycle of (a) temperature, (b) salinity, (c) irradiance, (d) Chl *a*, (e) dissolved
inorganic nitrogen (DIN) and (f) dissolved oxygen for near-surface water (1-5 m) in Bedford Basin.
The black dashed line depicts the annual cycle of CH₃I. The figures present the monthly mean
values based on the data collected from May 2015 to December 2017.



1	Figure 9. Detailed time-series from near-bottom waters in (a) 2015, (b) 2016 and (c) 2017. For
2	each year, the upper panel shows variability of salinity (red dash-dot line) and dissolved oxygen
3	(black dashed line); the lower panel shows iodocarbons (CH ₃ I: blue stars; CH ₂ CII: red open circles
4	and CH ₂ I ₂ : green open squares). The vertical bold, dashed lines (1 to 9) represent special events
5	discussed in section 4.5.
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- Table 1. R² value (Pearson's correlation coefficients, p<0.05) for the individual iodocarbon data
- based on both weekly data and monthly average.

		weekly			monthly	
	CH₃I	CH ₂ ClI	CH_2I_2	CH₃I	CH₂CII	CH_2I_2
CH₃I (1m)	1.0			1.0		
CH₂Cll (1m)	0.4	1.0		0.7	1.0	
CH ₂ I ₂ (1m)	0.0	0.3	1.0	0.1	0.4	1.0
CH₃I (5m)	1.0			1.0		
CH₂ClI (5m)	0.4	1.0		0.5	1.0	
CH ₂ I ₂ (5m)	0.1	0.4	1.0	0.1	0.6	1.0
CH₃I (10m)	1.0			1.0		
CH₂Cll (10m)	0.7	1.0		0.7	1.0	
CH ₂ I ₂ (10m)	0.1	0.3	1.0	0.2	0.4	1.0

Table 2. R² value (pearson's correlation coefficients, p<0.05) for iodocarbons and potentially

			weekly			monthly	
	1m	CH₃I	CH₂ClI	CH_2I_2	CH₃I	CH₂ClI	CH ₂ I ₂
	SST	0.5	0.4	0.0	0.7	0.6	0.1
	SSS	0.1	0.0	0.0	0.1	0.1	0.0
	Oxygen	0.3	0.2	0.1	0.3	0.2	0.1
	Flu	0.0	0.0	0.0	0.1	0.0	0.0
	PAR	0.1	0.1	0.0	0.1	0.3	0.0
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2 relevant parameters based on both weekly data and monthly average.

year	season	CH₃I	CH ₂ CII	CH_2I_2	I_{org}
2015	Spring				
	Summer	13.9	29.3	15.4	74.0
	Fall	13.1	30.7	13.7	71.2
	Winter	3.6	3.0	8.6	23.7
	Annual	11.9	25.9	14.0	65.9
2016	Spring	3.4	11.8	16.5	48.2
	Summer	12.0	33.2	9.4	64.0
	Fall	7.2	9.4	1.8	20.2
	Winter	3.2	3.0	1.7	9.6
	Annual	6.6	14.4	8.2	37.3
2017	Spring	3.0	2.7	1.7	9.0
	Summer	7.8	22.1	12.1	54.1
	Fall	19.7	31.3	15.9	82.8
	winter	3.2	4.5	9.1	25.9
	Annual	8.5	15.9	10.1	44.7

Table 3. Seasonal variation of total sea-to-air fluxes of iodocarbons (nmol m⁻² d⁻¹). Highest flux

Table 4. Concentration (pmol L⁻¹) of iodocarbons measured at near shore locations around
Bedford Basin as well as at the regular sampling location on July 19, 2017 (1: Tufts cove, 2:
Wrights cove, 3: Sackville (rive), 4: Mill cove and 5: Fairview cove) (see Fig. 1a). "Centre" refers
to the regular sampling site for the weekly time-series.
*Centre Mean for July: the average (and std. deviation) of measurements at the regular, weekly

6 sampling location during the month of July 2017 (n=4).

	CH₃I	CH₂CII	CH_2I_2
¹ Tufts cove	5.8	35.5	8.8
² Wrights cove	6.7	20.6	12.5
³ Sackville	3.8	6.5	4.9
⁴ Mill cove	8.3	28.0	18.6
⁵ Fairview cove	6.2	26.3	6.4
Middle of Bedford Basin	6.1	37.6	6.3
*Centre Mean for July	5.9±0.9	58.5±25.2	26.1±32.0





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Table 5. Comparison of sea-to-air flux (nmol m⁻² d⁻¹) of total organic iodine from different studies.
English Channel is an average for 1 year; Funka Bay value is average over 3 years; Kiel Fjord is
average over 2 years but for CH₃I only; Bedford Basin (this study) is an average over 3 years.
Seasons as defined in this study (see section 2). For Funka Bay, values in parentheses represent
fluxes that have been adjusted from the original reported values to take into account effect of
air-side resistance, using correction factors of 12 and 28 % for CH₂ClI and CH₂I₂ respectively
(based on average effects reported in Archer et al. (2007) and this study; see section 4.6).

	English Channel		Funka Bay		Kiel Fjord		Bedford Basin	
	Archer et	al.(2007)	Shimizu	Shimizu et al. (2017)		Shi et al. (2014)		Study
Season	Total	%-CH₃I	Total	%-CH₃I	Total	%-CH₃I	Total	%-CH₃I
Spring			15.3(12.8)	33.8(40.3)	2.8		28.6	11.1
Summer			113.3(86.6)	4.7(6.2)	5.2		64.0	17.5
Fall			47.6(41.1)	32.5(37.6)	2.2		58.1	23.0
Winter			27.5(22.5)	22.3(27.2)	0.2		19.7	16.8
Annual	42.6	27.9	54.8(43.5)	14.0(17.6)	3.3		46.7	19.3