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J. Sommar, M.E. Andersson, H.-W. Jacobi. Circumpolar measurements of speciated mercury, ozone and carbon monoxide in the boundary layer of the Arctic Ocean. Atmospheric Chemistry and Physics, European Geosciences Union, 2010, 10 (11), pp.5031-5045. <10.5194/acp-10-5031-2010>. <isu00554371>

## HAL Id: insu-00554371 https://hal-insu.archives-ouvertes.fr/insu-00554371

Submitted on 27 Feb 2012

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# **Circumpolar measurements of speciated mercury, ozone and carbon monoxide in the boundary layer of the Arctic Ocean**

J. Sommar<sup>1</sup>, M. E. Andersson<sup>2</sup>, and H.-W. Jacobi<sup>3,4</sup>

<sup>1</sup>State key Laboratory of Environmental Geochemistry, Institute of Geochemistry, Chinese Academy of Sciences, Guiyang 550002, China

<sup>2</sup>Department of Chemistry, Göteborg University, 41296 Göteborg, Sweden

<sup>3</sup>CNRS/Université Joseph Fourier – Grenoble 1, Laboratoire de Glaciologie et Géophysique de l'Environnement, 54 Rue Molière, 38400 St Martin d'Hères, France

<sup>4</sup>Alfred Wegener Institute for Polar and Marine Research, Am Handelshafen 12, 27570 Bremerhaven, Germany

Received: 19 August 2009 – Published in Atmos. Chem. Phys. Discuss.: 5 October 2009 Revised: 10 May 2010 – Accepted: 10 May 2010 – Published: 1 June 2010

Abstract. Using the Swedish icebreaker Oden as a platform, continuous measurements of airborne mercury (gaseous elemental mercury (Hg<sup>0</sup>), divalent gaseous mercury species Hg<sup>II</sup>X<sub>2</sub>(g) (acronym RGM) and mercury attached to particles (PHg)) and some long-lived trace gases (carbon monoxide CO and ozone O<sub>3</sub>) were performed over the North Atlantic and the Arctic Ocean. The measurements were performed for nearly three months (July–September 2005) during the Beringia 2005 expedition (from Göteborg, Sweden via the proper Northwest Passage to the Beringia region Alaska – Chukchi Penninsula – Wrangel Island and in-turn via a northpolar transect to Longyearbyen, Spitsbergen). The Beringia 2005 expedition was the first time that these species have been measured during summer over the Arctic Ocean going from 60° to 90° N.

During the North Atlantic transect, concentration levels of  $Hg^0$ , CO and  $O_3$  were measured comparable to typical levels for the ambient mid-hemispheric average. However, a rapid increase of  $Hg^0$  in air and surface water was observed when entering the ice-covered waters of the Canadian Arctic archipelago. Large parts of the measured waters were supersaturated with respect to  $Hg^0$ , reflecting a strong disequilibrium. Heading through the sea ice of the Arctic Ocean, a fraction of the strong  $Hg^0$  pulse in the water was transferred with some time-delay into the air samples collected  $\sim 20$  m above sea level. Several episodes of elevated

Hg<sup>0</sup> in air were encountered along the sea ice route with higher mean concentration  $(1.81\pm0.43 \text{ ng m}^{-3})$  compared to the marine boundary layer over ice-free Arctic oceanic waters  $(1.55\pm0.21 \text{ ng m}^{-3})$ . In addition, the bulk of the variance in the temporal series of Hg<sup>0</sup> concentrations was observed during July. The Oden Hg<sup>0</sup> observations compare in this aspect very favourably with those at the coastal station Alert. Atmospheric boundary layer O3 mixing ratios decreased when initially sailing northward. In the Arctic, an O<sub>3</sub> minimum around 15-20 ppbV was observed during summer (July-August). Alongside the polar transect during the beginning of autumn, a steady trend of increasing O3 mixing ratios was measured returning to initial levels of the expedition (>30 ppbV). Ambient CO was fairly stable  $(84\pm12 \text{ ppbV})$ during the expedition. However, from the Beaufort Sea and moving onwards steadily increasing CO mixing ratios were observed (0.3 ppbV day<sup>-1</sup>). On a comparison with coeval archived CO and O<sub>3</sub> data from the Arctic coastal strip monitoring sites Barrow and Alert, the observations from Oden indicate these species to be homogeneously distributed over the Arctic Ocean. Neither correlated low ozone and Hg<sup>0</sup> events nor elevated concentrations of RGM and PHg were at any extent sampled, suggesting that atmospheric mercury deposition to the Arctic basin is low during the Polar summer and autumn.



*Correspondence to:* J. Sommar (jonas@mails.gyig.ac.cn)

#### 1 Introduction

The average residence time of gaseous elemental mercury  $(Hg^0)$  and carbon monoxide (CO) in the lower troposphere is sufficient to make its distribution homogeneous over each hemisphere. While the seasonal cycle of CO exhibits a summertime minimum largely due to chemical oxidation initiated by the hydroxyl radical OH (e.g. Holloway et al., 2000), the seasonality of Hg<sup>0</sup> is less obvious taking into consideration worldwide long-term observations (Kim et al., 2005). However, several background stations in the mid-latitudes north of 45° N (Kellerhals et al., 2003; Kim et al., 2005) report a wintertime Hg<sup>0</sup> maximum. This has been attributed to seasonal trends in anthropogenic emissions and/or meteorological conditions, atmospheric oxidation processes, the height of the atmospheric mixing layer (Kock et al., 2005) and the terrestrial carbon pool (Obrist, 2007). Unlike  $Hg^0$ and CO, the mixing ratio of ozone  $(O_3)$  in the lower troposphere reaches a maximum during summer. The Arctic atmospheric boundary layer Hg<sup>0</sup> and O<sub>3</sub> cycles derived from background observation sites in the European and American high Arctic (Schroeder et al., 1998; Berg et al., 2003; Helmig et al., 2007), however, exhibit large seasonal discrepancies from that of the mid-latitudes.

It is very likely that high level of neurotoxic mercury in the Arctic ecosystems is partially linked to rapid, near-complete depletion of Hg<sup>0</sup> (MDEs) in the atmospheric boundary layer occurring episodically during the polar spring (Steffen et al., 2008). Upon reaction with reactive bromine species such as bromine atoms (Donohoue et al., 2006) hundred(s) of tons of oxidised mercury (HgII) are produced and perennially deposited into the Arctic environment (Ariya et al., 2004; Banic et al., 2003; Dastoor et al., 2008; Skov et al., 2004). The relative magnitude of this sink is profound, resembling almost half of the annual atmospheric input within a few weeks (Outridge et al., 2008). To a yet not well quantified, but significant degree, a back-reduction of HgII to Hg<sup>0</sup> occurs resulting in a re-cycling of volatile mercury to the atmosphere (Schroeder et al., 2003). However, it cannot compensate for the total deposition, and a net assimilation into the food chain occurs. The Hg levels in the traditional food of indigenous people living in the Arctic pose a threat for human pre- and neonatal neurological development (Steffen et al., 2008). Arctic marine mammals such as beluga whales frequently contain total Hg levels well above Canadian Federal Consumption Guidelines (Lockhart et al., 2005). Again, the fate of surplus mercury deposited to the Arctic basin during polar spring is largely unknown with reference to transport and transformation. This issue has recently been discussed in modelling papers by Hedgecock et al. (2008) and Outridge et al. (2008) in favour of less net accumulation in the abiotic reservoirs. The Beringia 2005 expedition, taking place a few months after the period of elevated deposition of Hg to the Arctic Ocean, offered a unique opportunity to sample vast exposed, yet unexplored areas.

During the polar spring MDEs, Hg<sup>0</sup> is generally strongly correlated with O<sub>3</sub> (e.g. Ebinghaus et al., 2002; Eneroth et al., 2007). The loss of tropospheric O<sub>3</sub> and most probably Hg<sup>0</sup> in the Arctic is initiated by catalytic heterogeneous reactions involving reactive bromine gases that originate from refined sea salt reservoirs linked to sea ice (e.g. frost flowers (Kaleschke et al., 2004; Jacobi et al., 2006) or snow on sea ice (Simpson et al., 2005)) and is activated by gas-phase photochemistry: Young ice regions with open leads tend to promote the growth of salinity-enhanced ice crystals with large effective surface areas. Hönninger (2002), Brooks et al. (2006) and Sommar et al. (2007) reported ground-based measurement of elevated BrO concentration in concert with air masses depleted of  $Hg^0$  and  $O_3$ . Unlike  $Hg^0$  and  $O_3$ , CO is less susceptible to bromine radical initiated degradation and hence essentially remaining chemical inert during polar spring (Sumner et al., 2002). The spatial extent of MDEs may be approximated using satellite observations of BrO and ground-based observations (Garbarino et al., 2002; Hollwedel et al., 2004; Lahoutifard et al., 2006; Poulain et al., 2007b; Simpson et al., 2005). During the rest of the year the atmospheric reactions driving the cycling of Hg in the boundary layer is ambiguous, possibly involving HO<sub>x</sub> and NO<sub>v</sub> radicals (Calvert and Lindberg, 2005; Sommar et al., 2001; Sommar et al., 1997). Therefore, it is important to co-sample Hg<sup>0</sup> with other long-lived atmospheric species. Recently, Hg<sup>0</sup>/CO emission ratios from long-distance transported pollution episodes to remote observation sites have been employed to estimate the anthropogenic Hg<sup>0</sup> emissions from eastern Asia (Jaffe et al., 2005) and Europe (Slemr et al., 2006) as well as emissions from large scale forest fires (Ebinghaus et al., 2007; Sigler et al., 2003). Selected studies suggest a substantial underestimation of source strength of Hg<sup>0</sup> in the current inventories for East Asia and Europe respectively. We have already pointed out that the regimes of O<sub>3</sub> are similar with that of Hg<sup>0</sup> during MDEs, in addition O<sub>3</sub> is produced from degradation of CO by OH radicals, so the addition of O<sub>3</sub> measurements to the sampling programme was a matter of course.

A main objective was to estimate the spread of mercury in the Arctic environment and to investigate the lability of the deposited HgII compounds in condensed phases with respect to reduction. During the polar transect of the Arctic Ocean from Alaska to Spitsbergen, speciated Hgmeasurements were performed including gaseous divalent Hg species (RGM) and Hg attached to fine particles (PHg) in addition to that of  $Hg^0$ ,  $O_3$  and CO. PHg has turned out to be a regional tracer for combustion in Europe (Wängberg et al., 2003). This paper focuses on the air measurements alone and is a companion paper of those dealing with mercury in sea water (surface and deep-sea), ice, snow, and sediments collected in the Arctic Ocean during the Beringia 2005 expedition (Andersson et al., 2008b; Andersson, 2010). In Andersson et al. (2008b), the measurements of dissolved gaseous mercury (acronym DGM, practically ~Hg<sup>0</sup>(aq)) in

Species	Freq.	Start (yy-mm-dd hh:mm)	Stop (yy-mm-dd hh:mm)	Instrument
TGM, Hg <sup>0</sup>	$12  h^{-1}$	05-07-05 14:20	05-09-24 11:00	Tekran 2537A CVAFS <sup>a</sup>
RGM-PHg <sup>a</sup>	$4 \text{ day}^{-1}$	05-08-25 16:00	05-09-22 17:00	Front sampler+Tekran 2537A CVAFS
O <sub>3</sub>	$6  h^{-1}$	05-07-05 12:10	05-09-22 12:00	O <sub>3</sub> 41M Ansyco Environnement
CO	$120 \ h^{-1}$	05-07-06 10:00	05-09-24 16:20	AL 5001 AeroLaser

Table 1. Information concerning the air sampling protocol onboard IB Oden (4th deck).

<sup>a</sup> Front-sampler used during Leg 3.

the surface waters of the Arctic Ocean are thoroughly described. The DGM data reported will be capitalized on here in connection with the discussion of elevated Hg<sup>0</sup> observations in air. Elevated DGM concentrations observed in the surface water of the Arctic Ocean compared to adjacent oceans is presumably due to input by springtime MDEs and large riverine inputs of Hg<sup>II</sup> and dissolved organic carbon (DOC) forming photo-labile complexes as well as limited gas exchange with air due to the sea ice cover (Andersson et al., 2008b). Poulain et al. (2007a) demonstrated the continuous presence of active, cold resistant Hgreducing microbes in the Arctic Ocean, which do not depend on light. Based on relatively flat vertical profiles of total mercury (Hg-tot) from the Beaufort Sea and the Canadian Archipelago during summer, Outridge et al. (2008) suggested relative minor input from the atmosphere to the Arctic Ocean. St Louis et al. (2007) reported high levels of DGM  $(129\pm36 \text{ ng m}^{-3})$  and organo-mercurials in the surface waters near the Ellesmere Island during late spring.

#### 2 Experimental

#### 2.1 Sampling location

The measurements were conducted on board the icebreaker Oden. The IB Oden is a 108 m long and 31 m wide icebreaker, which has been rebuild to meet scientific demands. The ship is equipped with meteorological and oceanographic instrumentation. The cruise (track outlined in Fig. 1) was divided into three legs: 1. Göteborg, Sweden – Barrow, Alaska, 2. Barrow – Chukchi Peninsula – Wrangel Island – Barrow, 3. Barrow – Longyearbyen, Spitsbergen. Ambient air samples were continuously collected and analyzed using automatic instruments. The instruments were housed in a heated container laboratory at the 4th deck fronting the stem ( $\sim 20$  m above sea level). Further information is summarised in Table 1.

#### 2.2 Mercury measurements

A Tekran<sup>®</sup> (Model 2537A) gas-phase mercury vapour analyzer instrument was used to measure total gaseous mercury (TGM). TGM refers to gaseous elemental mercury (Hg<sup>0</sup>) and



**Fig. 1.** The spatial distribution of TGM,  $Hg^0$  in air (ng m<sup>-3</sup>) along the cruise track. The locations of coastal stations Alert (Nunavut, Canada) and Barrow (Alaska, USA) are included as well.

small contributions of other gaseous mercury species (<1%at ambient conditions) that may pass through the sampling line and be detected as Hg<sup>0</sup> (Temme et al., 2003). The instrument utilised two gold cartridges in parallel, with alternating operation modes (sampling and desorbing/analysing in mercury free Ar stream) on a predefined time base of 5 min. A 0.45 µm Teflon filter protected the sampling cartridges against contamination by particulate matter. Ambient air was sampled with a flow rate of  $1.5 \,\mathrm{L\,min^{-1}}$ . The exposure to marine air tends to permanently passivate the gold cartridges. Therefore, the cartridges were changed frequently. To regenerate the cartridges, they were ultrasonically cleaned in a Labosol<sup>®</sup> wash, repeatedly rinsed with Milli-Q water and dried in a mercury free stream of argon. The performance of the cartridges was routinely checked by injections from internal and external temperature controlled mercury sources. All concentrations are calculated to represent mass of Hg<sup>0</sup> per m<sup>3</sup> of air at STP (0 °C and 1013 hPa). The method detection limit was  $<0.15 \text{ ng m}^{-3}$  for a 5 min sample calculated as  $1 \times \sigma$ .

During the 3rd leg of the Beringia 2005 expedition, an automatic set-up was used to simultaneously measure RGM and PHg (Wängberg et al., 2008). The air to be sampled was pulled through a specially designed intake consisting of a Teflon coated elutriator with an acceleration jet with 2.5 µm cut-off and then into a 20 cm long KCl coated annular quartz glass denuder (URG-2000-30 K and URG-2000-30CD, University Research Glass Corp, Chapel Hill, NC, USA). A filter holder (made from quartz glass) was placed vertically downstream the KCl denuder. Fine particles ( $\leq 2.5 \,\mu m$ ) were collected on a quartz glass fibre filter of 20 mm diameter. The denuder and the particle filter were placed in separate ovens followed by a pyrolyser unit kept at 750 °C. The apparatus was assembled in a temperature controlled aluminium box fixed to the gun wale. The sample intake was located in the bottom of the aluminium box  $\sim 20$  m above sea level. The denuder was kept at 50 °C during sampling to protect the KCl coating from humidity. During analysis both the denuder and quartz filter were purged with mercury-free air. During the initial step of the analysis the particle filter was heated to 600 °C. PHg on the trap was desorbed and thermo-reduced to Hg<sup>0</sup> when passing the pyrolyser. In the next step, while the particle trap was maintained at high temperature, the denuder was heated to 540 °C. At 500 °C, RGM was desorbed from the KCl-matrix and totally decomposed to Hg<sup>0</sup> in the pyrolyser. The desorbed Hg<sup>0</sup> pulses were transferred via a thermally insulated FEP tubing temperature controlled at  $60\,^\circ\mathrm{C}$ to the Tekran<sup>®</sup> instrument (described above) for mercury detection. RGM and PHg samples were collected with a flow rate of  $8.5 \,\mathrm{L}\,\mathrm{min}^{-1}$  during 5 h effective sampling time with an intervening period of 1 h for analysis. The method detection limit (MDL) of RGM and PHg were 3 and  $1.5 \text{ pg m}^{-3}$ , respectively, calculated as  $1 \times \sigma$ .

In the following, no distinction will be made between the TGM and  $Hg^0(g)$  measurements performed. The acronym TGM,  $Hg^0$  used below is justified by comparing the precision of the Tekran instrument with the actual concentrations of RGM and PHg measured.

Measurement of DGM in surface water as thoroughly described in Andersson et al. (2008b) was performed using a automatic continuous equilibrium system with 10 min resolution (Andersson et al., 2008c)

#### 2.3 Ozone and carbon monoxide measurements

Surface level  $O_3$  mixing ratios were measured continuously using a commercial analyzer based on UV absorption ( $O_341M$ , Ansyco GmbH, Karlsruhe, Germany). The same instrument has been used before during cruises on the Atlantic and Arctic Oceans (Jacobi and Schrems, 1999; Jacobi et al., 2006). It was operated with an integration time of 10 min. Calibrations were made on a daily basis using an internal ozone generator. The detection limit was 1 ppbV at a sampling rate of 1 L min<sup>-1</sup>. CO mixing ratios were determined also using a commercial detector employing a vacuum-UV resonance fluorescence technique (AL 5001, Aero-Laser GmbH, Garmisch-Partenkirchen, Germany). Details of the instrument can be found elsewhere (Gerbig et al., 1999). Air samples were collected with a time resolution of 0.5 min. The instrument was calibrated daily using a certified CO/synthetic air mixture from cylinders. For zero measurements, ambient air was used after removal of CO by oxidation with Pd-catalyst (150 °C). The detection limit was <1 ppbV. The air inlet (FEP tubing, 10 m long) for both instruments was positioned in a protective housing (PFA material, Savillex Corp.) in the top of a pylon located 3 m above the 4th deck (approximately 22 m above sea level).

#### 2.4 Ancillary data

Meteorological and basic seawater parameters were measured by the ship's monitoring system. The air parameters were measured 30 m above the sea surface while the aquatic parameters were obtained from the bow-water intake located approximately 5 m beneath the sea surface.

Air parcel backward trajectories were calculated and converted to GIS line shape files using the TrajStat software (v. 1.1, Wang et al., 2009). A trajectory calculation module of Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) Model (Draxler and Rolph, 2003) included in TrajStat as an external process was using gridded meteorological data sets as input. These sets were retrieved from Global Data Assimilation System (GDAS) meteorological data archives (ftp://arlftp.arlhq.noaa.gov/pub/archives/ gdas1/) of the Air Resource Laboratory, National Oceanic and Atmospheric Administration (NOAA). Five-day backtrajectories arriving at 100-1500 m above sea level were generated every 6 h along the entire cruise track. The stability was verified by calculating trajectories at different starting heights. Selected trajectory shape files are depicted in Figs. 2-4.

Further data for 2005 regarding CO and O<sub>3</sub> measured at the Arctic coastal stations Alert, Canada and Barrow, Alaska are also presented as well as TGM data from Alert. CO data from Barrow were provided by Paul Novelli (NOAA/ESRL, personal communcation, 2009). O<sub>3</sub> and TGM data at Alert were provided by Doug Worthy (Environment Canada, personal communication, 2009) and Alexandra Steffen (Environment Canada, personal communication, 2010) respectively. Further data (O<sub>3</sub> at Barrow and CO at Alert) were retrieved from the WMO World Data Centre for Greenhouse Gases (gaw.kishou.go.jp/wdcgg.html). All data are available as 1-h averages.



**Fig. 2.** Cruise track illustrated by ship's position every 6 h (black stars) and for July plots of selected daily five-day back-trajectories starting (usually at 12:00 UTC) at 100–1500 m above sea level. The offset in starting elevation was chosen individually to produce a purposeful stable trajectory lacking of significant ground contact. The snow/ice coverage shown in the image does not represent the actual distribution during the cruise.

#### 2.5 Data analysis

The raw sampling data of  $Hg^0/TGM$ , CO and  $O_3$  was averaged with a common time base of 10 min. The composite data points were further investigated for abnormal high and/or low concentrations. Using a ship as sampling platform, occasional sampling of the plume of the internal engine is inevitable. It can be expected that in the combustion plume, which contain large amounts of nitrogen oxides and CO, the ratio between  $O_3$  and CO changes significantly due to the consumption of  $O_3$  by the following reaction:

$$NO + O_3 \rightarrow NO_2 + O_2 \tag{R1}$$

Thus, it is handy to screen the data by using the  $O_3/CO$  ratio. Mercury in crude oil is a catalyst's poison and has to be removed before processing the lighter fractions (Wilhelm and Spitz, 2003). However, the heavier fractions used for ship fuel may contain Hg in the ppb-range (Wilhelm, 2001). It is thus imperative to investigate and possibly screen the Hg<sup>0</sup> data to remove values influenced by emissions from the ship (Aspmo et al., 2006; Slemr et al., 1985). These rather frequent local pollution events exhibit an enhanced CO and low O<sub>3</sub> concentration. The exclusion of data points was, therefore, based on crossing a CO/O<sub>3</sub>-limit with certain conditions of the two variables and/or by wind direction (see below). To be able to examine the diurnal variability, the data points were lumped into 1-h bins representing the local time instead of UTC. A spatial plot of TGM, Hg<sup>0</sup> distribution along the



**Fig. 3.** Cruise track illustrated by ship's position every 6 h (black dots) and for the two first weeks of August plots of selected fiveday back-trajectories starting at 100-1,800 m above sea level. The offset in starting elevation was chosen individually to produce a purposeful stable trajectory lacking of significant ground contact. Concerning the TGM,  $Hg^0$ ; CO pollution event described in Sect. 3.1.2, trajectories with 6 h interval starting times are depicted (reddish coloured). The snow/ice coverage shown in the image does not represent the actual distribution during the cruise.

cruise track was generated using Ocean Data View (v. 3.0, Reiner Schlitzer, Alfred Wegener Institute for Polar and Marine Research, Bremerhaven, Germany).

#### 3 Results and discussion

#### 3.1 General observations

The time series of the concentrations of the trace gases TGM,  $Hg^0$ ; CO and O<sub>3</sub> in ambient air samples is shown in Fig. 5. Periods with passage through sea ice are indicated by the blue shading. Figure 6 provides a comparison of the ship-borne trace gas observations with the time series of GAW global stations Alert and Barrow. The spatial distribution of TGM,  $Hg^0$  is included in Fig. 1, where the location of Alert and Barrow is indicated. The time series for RGM and PHg during the polar transect (leg 3) is separately displayed in Fig. 7. Only a few PHg concentrations were above MDL, whereas the corresponding fraction for RGM was at 50%. The air samples generally comprised the pristine marine boundary layer (MBL), however, it also include several episodes of local and regional pollution, which has to be taken into consideration.



**Fig. 4.** Cruise track illustrated by ship's position every 6 h (black dots) and for August (weeks 3–5) and September (weeks 1–3) plots of selected daily five-day back-trajectories starting (usually at 12:00 UTC) at 300–2500 m above sea level. The offset in starting elevation was chosen individually to produce a purposeful stable trajectory lacking of significant ground contact. The snow/ice coverage shown in the image does not represent the actual distribution during the cruise.

#### 3.1.1 Pollution from internal combustion

The local pollution is derived from the plume of the ship's internal combustion engine. The NO<sub>x</sub>- and CO-containing plume titrates O<sub>3</sub> by thermal oxidation of NO (cf. Eq. 1). Hence, when sampling the ship's plume, the CO/O<sub>3</sub>-ratio increased significantly (from between 2 and 8 in background air to between 15 and a few hundreds). The sampling artefact occurred concomitantly with wind direction within a sector of  $180^{\circ}\pm60^{\circ}$  relative to the ship's bow or with wind speeds less than  $3 \text{ m s}^{-1}$ . The conceivable impact of the plume on the Hg<sup>0</sup>-levels measured was examined. For events with CO to  $O_3$  ratios larger than 15, the corresponding Hg<sup>0</sup>/TGM data was plotted vs. CO (Fig. 8). These parameters were only weakly correlated with no obvious trend. This was also the case for PHg and RGM. Hence, we conclude that the exhaust of the IB Oden does not affect the measurements of the mercury components. During the polar transect, IB Oden navigated in tandem with US-CGC Healy (WAGB-20). Occasionally, the plume from US-CGC Healy was intercepting the sampling intakes, likewise without indicating enhanced mercury levels.

In summary, the screening criteria finally used to reject CO and O<sub>3</sub> data possibly influenced by local pollution are:  $[CO]/[O_3] > 15$  and  $[O_3] < 8$  ppbV. A particular air pollution event on August 5th caused by an approaching bunker ship in open sea provided an elevated signal of TGM, Hg<sup>0</sup>. In this single case, Hg data were screened out. In Fig. 5, these screened data can be distinguished as open symbols (red circles for O<sub>3</sub> and black squares for CO respectively).

#### 3.1.2 Regional pollution

Some pollution episodes were encountered in a fjord and along the coast of the Chukchi Peninsula. These events during the first week of August are characterised by elevated TGM, Hg<sup>0</sup> and CO mixing ratios at mid-to-low O<sub>3</sub> concentrations (10-15 ppbV). Analysis of the corresponding back trajectories demonstrated that the air masses previously slightly passed over the Chukchi Peninsula but essentially had a marine fetch (cf. Fig. 4). The 2-3 August episode occurring in the fjord of Providence Bay was the most extensive one ( $\sim$ 7 h duration), where TGM, Hg<sup>0</sup> and CO showed maxima at  $\sim 5 \text{ ng m}^{-3}$  and  $\sim 225 \text{ ppbV}$ (~420 ppbV at 30 s resolution) respectively. A close-up TGM-CO scatter plot of this event in Fig. 9 shows a moderate linear correlation (statistically significant at >95% confidence). The enhancement of these gases represented by the slope  $\Delta$  (TGM, Hg<sup>0</sup>)/ $\Delta$ CO of 0.016 ng m<sup>-3</sup> ppbV<sup>-1</sup>  $(\sim 1.8 \times 10^{-5} \text{ mol mol}^{-1})$  is however well above the narrow range of  $(0.67 - 2.4) \times 10^{-7}$  mol mol<sup>-1</sup> hitherto reported for biomass burning (Ebinghaus et al., 2007) and also significantly higher than East Asian industrial emission ratios of  $\sim (5 \pm 2) \times 10^{-6} \,\mathrm{mol}\,\mathrm{mol}^{-1}$  reported by Weiss-Penzias et al. (2007). Since TGM, Hg<sup>0</sup> and CO were only moderately correlated during this episode, it is probable that the air plume was influenced by the mixing of changing backgrounds or was impacted by multiple sources (cf. Jaffe et al., 2005). Moreover, the mountainous and complex terrain settings (with peaks above 2000 m) around the fjord probably influenced the local meteorology and air flow.

#### 3.2 Observations of Arctic and North Atlantic background air

After removing data possibly influenced by internal combustion pollution, Table 2 was constructed showing the descriptive statistics of the measurements. The major trace gas data applying to merely the Arctic Ocean are summarized in Table 3. These sampled parameters have furthermore been divided into observations made over sea ice as well as into monthly information. Since the measurements were made during an extended period on a mobile platform covering major parts of the Arctic, they can be influenced by seasonal as well as spatial concentrations patterns. Table 3 also include a statistical summary of TGM observations at station Alert.



**Fig. 5.** 10-min averaged concentrations of atmospheric trace gases TGM,  $Hg^0$ ;  $O_3$ ; CO over the northern Atlantic and Arctic Ocean measured onboard the IB Oden during BERINGIA 2005. Passage through ice-covered sea is shaded in bluish colours. The names of some important geographical area discernible in Fig. 1 are indicated as well.

Table 2. Summary of the measurements after screening the raw data from local combustion.

Species	Ν	Mean	Min	Max	Median	SD	RSD
TGM, $Hg^0$ (ng m <sup>-3</sup> )	9061	1.73	1.00	5.24	1.67	0.36	0.21
$DGM (ng m^{-3})$	6138	42.4	4.9	207.4	37.7	25.9	0.61
O <sub>3</sub> (ppbV)	9793	21.4	8.0	47.3	21.0	5.5	0.27
CO (ppbV)	10000	84.4	50.3	223.8	84.0	11.6	0.14
RGM (pg m <sup><math>-3</math></sup> )	57	3.2	0.7 (ND)	10.0	3.1	1.7	0.53
$PHg (pg m^{-3})$	54	1.0	0.2 (ND)	3.2	0.9 (ND)	0.7	0.71

Before entering the Canadian Arctic (Labrador Sea into Baffin Bay) measurements represent conditions in the North Atlantic. Accordingly, O<sub>3</sub> mixing ratios between 30 and almost 50 ppbV were observed. In the Canadian Arctic, the airflow shifted from having a North Atlantic into a Polar fetch leading to a decrease of O<sub>3</sub> to around 15 to 20 ppbV (cf. Fig. 5). The drop in ambient O<sub>3</sub> levels may indicate air masses from remote, low-NOx-environments where more ozone destruction than ozone production occurs during summertime (Helmig et al., 2007). Henceforward, a steady increase in O<sub>3</sub> from 20 to 30 ppbV was observed during the 1month long polar transect from the Beaufort Sea to the Barents Sea. Trajectory simulations indicated that air masses were advected from long fetches predominantly within the Canadian and Siberian parts of Arctic Ocean and sometimes extending into continental Eurasia (Fig. 4). Besides these trends in the background  $O_3$  concentrations, the time series also indicates numerous short term reductions of  $O_3$ , when the mixing ratios dropped to 10 to 15 ppbV.

Compared to O<sub>3</sub> the variability in the CO mixing ratios was smaller. The average values ranged from around 80 ppbV during the first half of the cruise. During the second half of the expedition, we observed a statistically significant (>95% confidence) linear increase in the CO data (slope: 0.3 ppbV day<sup>-1</sup>, R = 0.50), until the values reached an average of around 110 ppbV during the second half of September. Nevertheless, even CO mixing ratios showed a short-term variability in the order of several tens of ppbV.

To distinguish between spatial and seasonal trends, Fig. 6 shows the ship-borne data together with measurements performed at the Arctic coastal stations Alert and Barrow during the same period. On two occasions dated in Fig. 6, IB Oden

**Table 3.** Summary of screened TGM,  $Hg^0$ ;  $O_3$  and CO data applicable to the Arctic Ocean (incl. Baffin Bay, Northwest Passages, Beaufort Sea, Chukchi Sea, and Greenland Sea). Excluded are thus data from the North Atlantic Ocean, Labrador Sea and Bering Sea (IHO, 1953). Descriptive statistics of TGM data from station Alert on Ellesmere Island are included as well.

Species	Overall			July		August		September		Sea Ice			Sea w/o ice						
	Ν	Mean	Med.	SD	Ν	Mean	SD	N	Mean	SD	N	Mean	SD	Ν	Mean	SD	N	Mean	SD
$ \begin{array}{c} \hline TGM, Hg^0 \\ (ng m^{-3}) \\ IB Oden \end{array} $	7864	1.72	1.67	0.35	1737	2.08	0.44	3686	1.62	0.24	2441	1.61	0.20	5443	1.81	0.41	2421	1.55	0.21
TGM (ng m <sup>-3</sup> ) Alert	1764	1.86	1.80	0.31	580	2.08	0.24	649	1.91	0.26	535	1.55	0.12						
O <sub>3</sub> (ppbV) CO (ppbV)	8491 9276	20.6 87.0	20.9 85.6	4.6 11.8	2435 2396	17.3 80.9	3.1 11.2	3,859 4060	20.4 84.3	3.4 9.4	2197 2820	24.5 96.2	4.7 9.9	5921 6215	21.1 87.3	4.8 10.4	2570 3061	19.4 86.5	3.9 14.3

![](_page_8_Figure_4.jpeg)

**Fig. 6.** Comparison of Oden (orange squares), Barrow (brown circles) and Alert (blue diamonds) observations for TGM,  $Hg^0$  (upper panel),  $O_3$  (middle panel) and for CO (lower panel). The time series from IB Oden depicted are comprised of 10-min averaged data while the others refer to hourly averaged data. Periods shaded in yellow resemble anchorage time of IB Oden outside Pt. Barrow.

![](_page_8_Figure_6.jpeg)

**Fig. 7.** Concentrations of airborne speciated mercury (TGM, Hg<sup>0</sup>; RGM and PHg, in lower panel) and some air parameters (temperature and relative humidity, in upper panel) measured along the transpolar route. Drawing the detection limits for RGM (green line) and PHg (yellow line) indicate the sampling data points below MDL.

anchored in the roadstead outside Barrow whereas Oden and Alert were separated by a shortest distance of  $\sim 800 \text{ km}$  during the cruise. Obviously, several of the trends in O<sub>3</sub> and CO concentrations are observed at the coastal stations as well as onboard. This concerns for example the relatively low values for O<sub>3</sub> and CO in July and August and the subsequent increase beginning in the second half of August. We conclude that the observed seasonal cycles of O<sub>3</sub> as well as CO are not restricted to the coastal sites, but also exist over the Arctic Ocean. Therefore, we suggest that the observations at Arctic coastal stations regarding O<sub>3</sub> and CO between July and September can be considered as representative for the

![](_page_9_Figure_1.jpeg)

**Fig. 8.** The relationship between TGM,  $Hg^0$  and CO during events of pollution from the ship's internal engine determined by a ratio of [CO]/[O<sub>3</sub>] larger than 15 (diamonds). Also included are the data screened according to Sect. 3.1.1 (triangles) but not comprised in the linear regression fit. The parameters are only weakly correlated implying that the ship's emissions are negligible with respect to Hg.

entire Arctic. Such a homogeneous distribution is also in good agreement with the relatively long chemical lifetimes of  $O_3$  and CO (between weeks and months, Helmig et al., 2007; Novelli et al., 1998) compared to average transport times into the Arctic of several days in summertime (Stohl, 2006).

The largest discrepancies between the Arctic coastal stations and the ship borne measurements in the long-term trends were observed for  $O_3$  at the beginning of the cruise. This clearly indicates the influence of the North Atlantic air masses during this part of the cruise, which contribute to elevated  $O_3$  mixing ratios compared to the Arctic. Interestingly, these air masses have no effect on the CO mixing ratios, which appear not to be lower over the Arctic Ocean compared to the North Atlantic.

The short-term variability in  $O_3$  and CO as observed in the ship borne measurements were in the case of  $O_3$  also apparent at the coastal stations. They might be to some extent obscured in the coastal CO data due to the longer averaging into 1-h averages compared to the 10-min averages collected onboard. Nevertheless, these short-term fluctuations are probably related to pollution events caused by continental sources (Novelli et al., 1998). It is interesting to note, that such pollution events were also observed in the second half of the cruise over the Arctic Ocean at relatively long distances from potential continental sources (Fig. 4). By contrast to the observations made in the Beringia region, these events exhibit a general lack of correlation between TGM, Hg<sup>0</sup> and CO.

By comparison with summer observations (median:  $1.67 \text{ ng m}^{-3}$ , July–September) from Alert, 1995–2002, Oden TGM, Hg<sup>0</sup> data exhibit an equivalent statistical median (Stef-

![](_page_9_Figure_7.jpeg)

**Fig. 9.** Scatterplot of TGM,  $Hg^0$  vs. CO observations during the 2–3 August pollution episode. The equation of the linear fit as well as the correlation coefficient are displayed.

fen et al. 2005). Decadal observations at Alert between 1995 and 2005 reveal no perennial statistically significant trend for the summer period. Analogous to Alert 2005 summer data, the magnitude of and variability in the Arctic MBL TGM, Hg<sup>0</sup> concentrations as observed onboard Oden were largest during the early phase of the season (Fig. 6). The variability in the latter data set is obviously higher, however the corresponding statistical means compare very favourably and are significantly higher (at >99% confidence level) than that of the North Atlantic observations (mean:  $1.73 \text{ ng m}^{-3}$ ). Upon entering the Canadian Arctic, the swift shift in ambient TGM, Hg<sup>0</sup> level qualitatively represents the reciprocal behaviour of O<sub>3</sub>. The increased concentration of TGM, Hg<sup>0</sup> observed in the early summer Arctic MBL is thus not caused by advection of polluted mid-latitudinal air masses but is according to back-trajectory analysis due to sources present in the Arctic basin. In a modelling study of bidirectional surface exchange in the Arctic, Dastoor et al. (2008) predicted that only in July a net re-emission of mercury prevails (averaged for years 2002-2004). Trajectory analysis for Hg<sup>0</sup> measurements at Zeppelin station during 2000–2008 indicates that the Arctic Ocean acts a source of Hg<sup>0</sup> to the atmosphere in July and August (Hirdman et al., 2009). In Table 3, the observed average TGM, Hg<sup>0</sup> concentration has been binned spatially into measurements over open sea and over sea ice (partial and total coverage, shaded in Fig. 6), respectively. The mean concentration of the first classification  $(1.55\pm0.21$  ng m<sup>-3</sup>) was lower (heteroscedastic t-test, p < 0.01) than that of the second  $(1.81\pm0.41 \text{ ng m}^{-3})$ . Also apparent in Table 3 is that no significant divergent distribution between ice and open water was observed for either  $O_3$  or CO. This agrees with the homogeneous distribution of both compounds between the Arctic Ocean and the coastal stations, which is mainly governed by large-scale transport and chemical processing in the atmosphere. However, concerning the corresponding DGM data, there was a

![](_page_10_Figure_2.jpeg)

**Fig. 10.** Diurnal variations of the  $O_3$ , CO and TGM,  $Hg^0$  concentrations summarised as in Table 2. The box assemblies include the mean, the upper and lower 95% confidence intervals and extremes (far outliers) for 1-h per bin diurnally integrated samples. The UTC time data has been recalculated to represent destination local time.

more striking difference between the averages of the two data bins (not shown in Table 3, see Andersson et al., 2008b). Our TGM, Hg<sup>0</sup> data set compares well with that of Aspmo et al. (2006) concerning higher TGM, Hg<sup>0</sup> concentrations over ice covered areas. The study by Aspmo et al. (2006) covering North Atlantic latitudes 54-85° in summer (June-August), where a step in ambient Hg<sup>0</sup> concentration occurred at 82° N when entering an area with elevated sea ice concentration. In the present study, the corresponding transition in the TGM,  $Hg^0$  level was encountered in the Baffin Bay  $\sim 70^\circ$  N. The study by Aspmo et al. (2006) also included measurements of airborne oxidised mercury (RGM and PHg). Jointly, low concentrations of RGM and PHg were measured. In Table 2, the RGM and PHg data tabulated include the observations below MDL. Replacing the values below MDL with a constant value of MDL/2 does not influence the overall statistics.

In Fig. 10, the diurnal variability of CO,  $O_3$  and TGM,  $Hg^0$  mixing ratios are shown in terms of box plots. The day length during air sampling was at least 22 h. In fact, more than 90% of the data was obtained during mid-night sun. No overall distinct diurnal pattern is discernible for ei-

Table 4. Over-all correlation matrix of trace gases and DGM.

Species	TGM, $Hg^0$	DGM	O <sub>3</sub>	CO
TGM, Hg <sup>0</sup>				
DGM	-0.035			
O <sub>3</sub>	-0.266	-0.027		
CO	-0.086	0.155	0.036	

ther CO, O<sub>3</sub>, TGM, Hg<sup>0</sup> or the transient mercury fractions RGM and PHg (not displayed). In general, diurnal cycles of  $O_3$  are small in polar regions (amplitude range 0-1.5 ppbV) and expected to decline with increasing latitude (Helmig et al., 2007). In contrast, during five consecutive days of the initial Atlantic transect diurnal features were observed for TGM (cf. Figs. 1 and 5). Such diurnal variations in TGM have previously been reported from ship-based measurements in more temperate latitudes of the northern hemisphere and explained to a large extent be accounted by Hg<sup>0</sup> evasion from surface water (Laurier et al., 2003). Strongly modulated features in TGM, Hg<sup>0</sup> were in the present study, however, above all confined to events with elevated DGM or as been reported in Sect. 3.1.2 to regional pollution events encountered at the Chukchi Peninsula. The former episodes of considerable interest for this study will be further investigated and discussed below.

# **3.2.1** The role of air-sea exchange of Hg<sup>0</sup> in the measured atmospheric Hg<sup>0</sup>/TGM concentrations

Table 4 shows the correlation matrix for the major trace gases measured including the dissolved gaseous mercury (DGM) in the surface water (Andersson et al., 2008b). All parameters were found to be only weakly inter-correlated. Furthermore, lagged cross-correlation analysis was applied on the subsets of the TGM, Hg<sup>0</sup>-DGM data involving periods of a high variability in both variables with high supersaturation of Hg<sup>0</sup> in the sea (such as the passage through certain parts of the Canadian archipelago, the discharge zone of Mackenzie river and the Arctic Ocean at  $\sim$ 86° N, cf. Fig. 11). By moving TGM, Hg<sup>0</sup> towards increasing time domain (typically 2-3 h), a significant improvement in the correlation from overall R = -0.035 to  $R \sim 0.4 - 0.6$  was obtained for several periods. In Fig. 11, the TGM, Hg<sup>0</sup>-DGM data are plotted together with time lags of 2 and 2.5 h for two episodes from different legs of the expedition. The physical rationale for a time delay in the TGM, Hg<sup>0</sup> signal is mass transport limitation in the sea-air transfer of the ensued strong supersaturation of DGM (the degree of supersaturation of DGM in the surface sea water (%) can be calculated according to:  $S = (DGM \times k_H / (TGM, Hg^0) - 1) \times 100$ , where  $k_H$  is the dimensionless Henry's law constant determined e.g. by Andersson et al., 2008a). For a sparingly soluble gas like  $Hg^{0}(g)$ 

the hydrodynamic barrier delaying the increasing DGM levels to be propagated into the air. The cases presented refer to a partially open sea (Fig. 11a) and an icebound sea (Fig. 11b). In the ice-covered oceanic surface waters such as in Fig. 11b, it is likely that elevated DGM concentrations were built up due to the restricted air exchange (Andersson et al., 2008b). On the other hand, the formation rate of DGM is dependant on biotic and abiotic reduction of  $Hg^{II}(aq)$  and a cap of seaice will attenuate photoreduction and retard DGM formation. The headway of IB Oden including the ice breaking presumably influences the gas-exchange of  $Hg^0$  positively.

The future predictions for the cycling of mercury in the Arctic are coupled with changes in the climate system. The global warming has and will have a profound and extensive impact on the biogeochemical cycle of Hg in the polar regions. The minimum sea ice extent of the Arctic Ocean has decreased at an average of  $(-7.4 \pm 2.4)\%$  per decade since 1979 while the corresponding decrease of mean annual extent of Arctic sea-ice is only at  $(-2.7\pm0.6)$  % per decade (Lemke et al., 2007). Simulations indicate abrupt future reduction reaching near ice-free September conditions by 2040 (Holland et al., 2006). As climate warming presumably will initially influence the magnitude and spatial extent of Arctic MDEs and boost photo-reduction of Hg<sup>II</sup>(aq) the efflux of Hg<sup>0</sup> from the Arctic Ocean to the atmosphere during summer is predicted to rapidly increase in the near future. The TGM, Hg<sup>0</sup>-DGM observations made in this study over the Arctic Ocean going from 60° to 90° N indicate that elevated concentrations are not only restricted to circumpolar marginal seas but also apply to the proper Arctic Ocean. Using a model developed by Nightingale et al. (2000), Andersson et al. (2008b) calculated a hypothetical average Hg<sup>0</sup> evasion of  $\sim$ 60 ng m<sup>-2</sup> day<sup>-1</sup> during the Beringia 2005 expedition with a maximum as high as  $\sim 2300 \text{ ng m}^{-2} \text{ day}^{-1}$ . St Louis et al. (2007) calculated a Hg<sup>0</sup> flux of  $130\pm30$  ng m<sup>-2</sup> day<sup>-1</sup> at ice break-up. In-turn Outridge et al. (2008) predict a 8-fold increase in summer Hg<sup>0</sup> flux for the future between 60° N and 80-90° N. The average Hg turnover time for the upper Arctic Ocean, which is at present  $\sim$ 5 years, may thus decrease.

## 3.2.2 The oxidation capacity of the Arctic MBL during summer

The low  $O_3$  concentrations in July and August agree well with the analysis of air mass transport from the Arctic Ocean to a coastal site in the Arctic (Hirdman et al., 2009). The detailed trajectory analysis for  $O_3$  measurements at Zeppelin station indicates that the Arctic Ocean constitutes a sink for  $O_3$  in spring- and summertime. While the destruction of  $O_3$ in April and May is related to ozone depletion events (ODEs) leading to the complete removal of  $O_3$  in the MBL (Simpson et al., 2007), the lower  $O_3$  concentrations in summer may be due to a combination of processes. The chemical destruction of  $O_3$  as a result of halogen activation from the abundant sea

![](_page_11_Figure_5.jpeg)

**Fig. 11.** The temporal air-sea distribution of  $Hg^0$  vapour for two specific destinations: (a) Mackenzie River discharge and (b) high Arctic Ocean (~86° N). Cross-correlation analysis revealed a maximised TGM,  $Hg^0$ -DGM covariance for time lags of 2 h and 2.5 h, respectively. Thus, the time-series of TGM,  $Hg^0$  and DGM has been shifted in (a) by 2 h and in (b) by 2.5 h. The maximum coefficient of correlation is 0.59 (n = 188) in case a and 0.54 (n = 307) in case b. The distribution of  $Hg^0$  between air and sea is in both cases in strong disequilibrium.

salt can continue, although it is necessarily less effective than in springtime since no ODEs have yet been observed in summer (Simpson et al., 2007). The low  $NO_x$ -environment like the Arctic can also contribute to a more efficient chemical  $O_3$ destruction during summer (Helmig et al., 2007). Both processes are related to the levels of incoming solar radiation, which start to decrease in late summer in the high latitudes. In addition, a more effective deposition to the open water areas compared to the snow-covered sea ice can reduce the  $O_3$  concentrations in the MBL (e.g. Gallagher et al., 2001; Helmig et al., 2009). In contrast, the absence of any significant diurnal variability for  $O_3$  indicates that in situ photochemical production in the Arctic MBL is low during this period of time.

The measurement of airborne  $Hg^0$ , RGM and PHg during the Polar transect of Leg 3 (Fig. 7) indicates that the Arctic Ocean MBL during summer does not promote significant chemical conversion of  $Hg^0$  and consequently constitutes a minor source of RGM and PHg. The apparent variability in TGM, Hg<sup>0</sup> concentration is as previously discussed source modulated by air-sea exchange and not correlated with either RGM or PHg. The speciated-Hg observations are in concert with those of Temme et al. (2003) and Aspmo et al. (2006) for the South Atlantic and the North Atlantic/Arctic Ocean respectively during Polar summer. Observations at temperate mid-latitudes indicate that the oxidising capacity of the MBL enriched with halide-abundant aerosols may induce a significant Hg<sup>0</sup> loss (Mason et al., 2003). However, the MBL of the Arctic Ocean during summer is distinguished by meteorological factors (high relative humidity, low visibility blocking solar radiation) not in favour of in-situ oxidation and in promotion of physical scavenging and removal of RGM and PHg. The lack of significant diurnal variability for RGM-PHg as well as the observed minor ratios RGM-PHg/Hg<sup>0</sup> of a few ‰ are obviously in support of a low in situ photochemical oxidation of  $\mathrm{Hg}^{0}$  in the Arctic MBL. We note that the temporal features of RGM (Hg<sup>II</sup>X<sub>2</sub>(g) species) in convolution with relative humidity (RH) are consistent with that of a sticky gas, which readily adsorbs onto surfaces and has a high water solubility (Fig. 7). The RGM-RH data exhibit a negative correlation with a correlation coefficient of 0.74.

In the case of CO, the summertime minimum can be attributed to the removal of CO by the reaction with hydroxyl radicals (Novelli et al., 1998); which is also more efficient during summer compared to the fall.

#### 4 Conclusions

For the first time we were able to perform measurements of airborne speciated-Hg together with O<sub>3</sub> and CO over major parts of the Arctic Ocean and adjacent seas during the polar summer period. The prevalent conditions of the Arctic MBL during the expedition resulted in relatively narrow statistical distributions of the trace gases (especially for CO). The largest relative variability was present in the TGM, Hg<sup>0</sup> data set during July, where generally higher concentrations were measured over the ice-capped sea. The results presented in this paper suggest the Arctic Ocean during summer to be a significant source of Hg<sup>0</sup> to the atmosphere. ODEs and MDEs, which occur regularly during polar spring, were not observed and, therefore, do not contribute to the removal of mercury and O<sub>3</sub> during summer. This agrees well with the seasonal cycle of BrO in high latitudes derived from remote sensing data (Hollwedel et al., 2004). BrO is one of the intermediate species generated during the ODEs and hence can be used as an indicator for such events. Nevertheless, O3 values lower than in the background air of the North Atlantic indicate an ongoing removal of O<sub>3</sub> from the Arctic MBL in summer probably due to a combination of chemical and physical processes. The good agreement between the ship-borne measurements of TGM, Hg<sup>0</sup>; O<sub>3</sub> and CO and the observations at Alert and Barrow indicate that the observed seasonal variation between July and September are representative for the entire Arctic. In terms of an abrupt future warming, the current cycling of Hg in the Arctic will be largely perturbed and sea-air interaction will be even more important in its atmospheric behaviour.

Acknowledgements. The Swedish Polar Secretariat is acknowledged for organising and including our project in the BERINGIA 2005 expedition. The Captain and crew of IB Oden are greatly appreciated for the support and help onboard. Special thanks go to Christian Temme (previously at G. K. S. S. Research Centre, Geesthacht, Germany) for kindly providing an analytical instrument for the expedition. The authors thank Paul Novelli (NOAA/ESRL), Alexandra Steffen (Environment Canada) and Doug Worthy (Environment Canada) for providing unpublished data from Barrow and Alert and the Japan Meteorological Agency for maintaining the WMO World Data Centre for Greenhouse Gases. The authors gratefully acknowledge Yaqiang Wang for producing and making the TrajStat software available, NOAA Air Resources Laboratory (ARL) for the provision of the HYSPLIT transport and dispersion model and READY website (www.arl.noaa.gov/ready.html) used in this publication. This research was partly supported by grants from the German Academic Exchange Service (DAAD) and the Swedish Foundation for International Cooperation in Research and Higher Education (STINT).

Edited by: K.-H. Kim

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