Circumpolar transport and air-surface exchange of atmospheric mercury at Ny-Ålesund (79° N), Svalbard, spring 2002


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Abstract. Mercury in different environmental compartments has been measured at Ny-Ålesund (78°54’ N, 11°53’ E) during an intensive campaign, 17 April to 14 May 2002. Time-resolved speciated determination of mercury in the atmosphere and snow was conducted at the Norwegian research station at the Zeppelin mountain, 474 m above the sea level, and at the Italian research facility Dirigibile Italia, 12 m above the sea level. Total Gaseous Mercury (TGM) was present in the range <0.1 to 2.2 ng m\(^{-3}\) during the campaign. Three mercury depletion events, identified as periods with decreased TGM concentrations, were observed. At the lower altitude, TGM concentrations following such events were found to exhibit both higher magnitude and larger variability in comparison to results from the Zeppelin station. Oxidised mercury species in air and fall-out with snow as well as mercury attached to particles were also measured and their concentrations were found to be anti-correlated with TGM in air. Concentrations of total Hg in snow (Hg-tot) showed a large (∼15×) increase in response to Gaseous Elemental Mercury Depletion Events (GEMDEs, range 1.5–76.5 ng L\(^{-1}\)). Solid evidence for photo-stimulated emissions of Hg\(^{0}\)(g) from the snow pack in conjunction to depletion events were obtained from gradient measurements as well as from flux chamber measurements. Steep diurnal concentration variations of Hg\(^{0}\)(aq) in surface seawater were also found to concur with changing solar radiation. The concentration of Hg\(^{0}\)(aq) in seawater was found to be in the range 12.2–70.4 pg L\(^{-1}\), which corresponds to supersaturation. Hence, the seawater surface constituted a source emitting elemental mercury. The concentrations of RGM (reactive gaseous mercury), Hg-p (particulate mercury), and BrO column densities (detected by DOAS) were very low except for a few individual samples during the major Hg\(^{0}\) depletion event. BrO vertical column densities obtained by the remote satellite ESR-2 and trajectory analysis indicate that the air masses exhibiting low Hg\(^{0}\) concentrations originated from areas with high BrO densities.

1 Introduction

Among the metals emitted to the atmosphere from natural, industrial and combustion processes, highly toxic mercury is unique as being mainly released in its gaseous elemental state (Hereafter, assigned Hg\(^{0}\)(g) or Gaseous Elemental Mercury, GEM). The semi-volatility of other environmentally relevant mercury compounds (e.g. HgCl\(_2\), CH\(_3\)HgCl) further underlines the atmospheric character of mercury. The latter oxidative state of mercury (e.g. mercuric compounds) has moderate residence times and a higher affinity to condensed phases. The apparently passive pollutant Hg\(^{0}\)(g) is distributed on a global scale, exhibiting intra-hemispherical and vertical uniformity with a gradient of ∼25% over the inter-hemispherical convergence zone (Schroeder and Munthe, 1998). Monitoring of mercury at global atmospheric watch mid-latitude stations does not currently reveal any consistent trend in tropospheric mercury load and a weak seasonality (Munthe and Palm, 2003). The seasonal cycle of boundary-layer airborne mercury in Polar Regions is however distinctly different from that observed at mid-latitude background sites (Ebinghaus et al., 2002) which has led to a considerable interest during the last 5 years. As tropospheric transport is a major channel for the distribution of mercury (Brosset, 1982; Lindqvist and Rodhe, 1985), the
During daytime, the self-reaction BrO+BrO is competitive to_various halogens, which can undergo photo dissociation (Adams et
al., 1997) and disproportionation to volatile diatomic elemental or inter-
halogens (catalysed process. Volatile Br(I) species can be absorbed by
fresh, partly frozen sea salt particles at low temperature (<253 K) more efficiently than to aqueous droplets. Ensuing are hydrolytic dark reactions such as hypobrominous-halide disproportionation to volatile diatomic elemental or inter-
alogens, which can undergo photo dissociation (Adams et al., 2002). Formation of Br2 is normally prevalent for reasons of thermo-chemical equilibria (Air-water distribution and aqueous complexation) and furthermore its actinic ab-
sorption extends out to longer wavelengths than Cl2 and BrCl. The relative importance of the Br2 photo-dissociation channel for the production of free halogen atoms is thus increased at high solar zenith angles. The multi-phase environ-
ment required for a “bromine explosion” (Ravishankara and Longfellow, 1999; Wennberg, 1999) suggests elevated concentrations especially close to relevant frozen surfaces, such as “frost flowers” (Rankin et al., 2002). Ozone is under such circumstances destroyed:

\[ \text{Br(g)} + \text{O}_2(g) \rightarrow \text{BrO(g)} + \text{O}_2(g) \]  

(1)

During daytime, the self-reaction BrO+BrO is competitive to photolysis only at high BrO levels. The latter processes reform Br(g) with unity quantum yield while photolysis and possible cross reactions (e.g. BrO+ Aldehydes/HO2) to vari-
ous extent produce reservoir compounds. As long as ozone is not completely depleted, most Br atoms react with ozone sustaining the BrO level to a certain extent.

Simultaneously with the modulation of the atmospheric ozone content, many other trace substances, among these Hg0(g) vary considerably indicating the influence of reactive halogen species (Schroeder et al., 1998). Hg0 deple-
tions (GEMDEs) cannot be quantitatively explained by the action of atomic chlorine in the gas-phase even if the reaction rate is set at diffusion-controlled limit (Sommar, 2001). A recent determination of the Hg+Cl reaction yielded a rate of \((1.0 \pm 0.2) \times 10^{-11} \text{ cm}^3 \text{s}^{-1}\) which corresponds to 10% of the diffusion controlled limit (Ariya et al., 2002). Concerning the reaction Hg0(g)+BrO(g), Höhninger (2002) estimated the lower limit for the reaction rate from field experiments to \(k_{\text{Hg}+\text{BrO}} \geq 3.7 \times 10^{-11} \text{ cm}^3 \text{s}^{-1}\). Raofie and Ariya (2003) report further constraints on \(k_{\text{Hg}+\text{BrO}} \leq 1 \times 10^{-13} \text{ cm}^3 \text{s}^{-1}\).

Boudries and Bottonen (2000) postulated from hydrocar-
on “clock” measurements that BrO radicals \((x=1)\) rather than bromine atoms \((x=0)\) oxidise atomic mercury to form molecular mercury products according to Reaction (2).

\[ \text{Hg}^0(g) + \text{BrO}_x(g) \rightarrow \text{Products} \]  

(2)

Assuming prompt formation of HgO(g), reaction (2, \(x=1\)) ap-
pears improbable using thermo-chemical arguments (Shepler and Pet-
terson, 2003; Sommar, 2001; Tosell, 2003). How-
ever, insertion product complexes like HgBrO are strongly bound and therefore feasible products (Balabanov and Peterson, 2003). Based on relative rate determinations of the homogeneous gas-phase reaction between Hg0+Br the following rate coefficient, \(k=(3.2 \pm 0.5) \times 10^{-12} \text{ cm}^3 \text{s}^{-1}\) was ob-
tained (Ariya et al., 2002) and the authors proposed that this reaction may explain Arctic Gas Elemental Mercury Deple-
tion Events (GEMDEs). In Malcolm et al. (2003) typical oxi-
dation rates of Hg0(g) in the MBL have been listed for a large number of oxidants also including BrO and ClO using previously unpublished kinetic data from Batelle Inc. The importance of the Hg0(g)+ClO(g) reaction in the perturbed Polar atmosphere and elsewhere suggested by Stutz et al. (2002) have less support in Malcolm et al. (2003). Likewise a theore-
tical prediction of the homogeneous rate coefficient for the reaction Hg0+ClO→HgO+Cl at elevated temperatures \((400° \text{K}<T<1800° \text{K}; 1 \text{ ~atm})\) by Xu et al. (2003) resulted in a relatively low value of \(\sim 5 \times 10^{-19} \text{ cm}^3 \text{s}^{-1}\) at 273°K (Arbe-
nius form: \(k \sim 2.29 \times 10^{-12} \exp(-4183/T) \text{ cm}^3 \text{s}^{-1}\)). Hg0(g) is currently categorised into the families of compounds (e.g. C2H2 and ozone) modulated by bromine species as a key ox-
idant during depletions (Bottonen et al., 2002; Lindberg et al., 2002). Although exhibiting disparate physico-chemical properties, the uniform temporal behaviour of GEM and O3(g) in connection with depletions in the surface layer is striking. We are aware that during the Antarctic summer, a persistence in elevated concentrations of molecular diva-
lent oxidative state mercury products in the gaseous phase (usually denoted RGM; for a discussion see Schroe-
der et al., 2003) has been reported and suggested to be controlled by HOx classes rather than reactive halogenated species (Sprovieri et al., 2002; Temme et al., 2003).

GEMDEs have been observed to occur annually often repetitively throughout maritime circumpolar stations: Alert, Ellesmere Island (Schroeder et al., 1998), Barrow, Alaska (Lindberg et al., 2002), Nord, Greenland (Skov et al., 2002), Neumayer, Queen Maud’s Land (Ebinghaus et al., 2002), Ny-Ålesund, Spitzbergen (Berg et al., 2003), Terra Nova Bay, Victoria Land (Sprovieri et al., 2002) and apparently in the Siberian Arctic (Golubeva et al., 2003; Steffen and Schroeder, 2002). At sub-polar latitudes, Höhninger (2002) in co-operation with Poissant and co-workers presents measurements of GEMDEs at 55°N surrounded by seawater ice in Hudson Bay. Stutz et al. (2002) propose that GEMDEs...
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take place over saline dry lakes in mid-latitudes. The observations made throughout the Arctic and Antarctica suggest that GEMDEs occur over large spatial scales and can thus be considered as analogues to ozone holes.

The atmosphere-surface partitioning behaviour appears however to be more complex possibly involving physico-chemical controls at the interface. While gradients of ozone are present in surface air/interstitial snow air suggesting a sink mechanism at the boundary between air, ice and water, a reversible red-ox balance may well be in control for mercury in snow which will strongly adjust its mobility. Hg^{II} deposited as a result of Hg^{0} depletion events is chemically (photo)labile toward reduction in the snow, hence to a certain extent transformed into Hg^{0}(aq) and ventilated to the atmosphere (Lalonde et al., 2002; Schroeder et al., 2003; Steffen et al., 2002). Moreover, GEMDEs correspond to short lived polar night ODEs that have been speculated to derive from local-driven chemistry (Foster et al., 2001). Since air-surface exchange and redox processes occurring in the condensed phase seem to be of prime importance, models describing only gas-phase processes cannot be used to evaluate atmospheric mercury budgets over snow-covered surfaces.

Being a high priority toxic element with diametrically opposite physico-chemical properties of the environmentally relevant oxidative states (e.g. Hg^{0}/Hg^{II}), efforts to estimate its atmospheric mass balance (e.g. perform specification/fractionation of airborne mercury in the field) during GEMDEs are of crucial importance. When comparing results of measurements of airborne mercury species during GEMDEs at different stations, it is evident that a closure of airborne mercury mass balance is not achieved. This is concordant with the strong tendency of RGM or mercury attached to existing particles (Hg-p) to undergo physical removal during transport. In terms of quantity and composition, the transient mercury groups discussed above tend in Hg^{0}(g)-depleted air masses to be site specific (Lindberg et al., 2002; Lu et al., 2003; Steffen et al., 2002; Temme et al., 2003). In some cases, the air masses are almost truly free from mercury while frequently the transient products are detected at elevated mixing ratios composing up to ~90% of the Total Gaseous Mercury (hereafter abridged TGM, for a definition see below) present in background air.

In this paper, we report and discuss results from the Norwegian high Arctic. The research area at Ny-Ålesund is located about 1300 km from the North Pole (cf. Fig. 1) and was established for long-term monitoring of airborne mercury in 1994. In 2000 high-time resolution measurement were started that allowed the identification of mercury depletion events (Berg et al., 2001, 2003). In addition, sampling of mercury in various other environmental compartments has been undertaken (Berg et al., 2001). Pongratz and Heumann (1998) have performed studies of the methylated mercury species (e.g. monomethylmercury, CH_{3}HgX; hereafter Methyl-Hg and dimethylmercury; (CH_{3})_{2}Hg) produced in the sea by different macro algae. They proposed that biogenic very toxic (CH_{3})_{2}Hg can be released from the sea surface and contribute to the polar atmospheric load of mercury (Pongratz and Heumann, 1999). Ozone destruction halogen catalysts XO have been found in mixing ratios up to a few tens of ppt during ODEs at this location since 1995 (X=Cl; e.g. Tuckermann et al., 1997; X=Br; e.g. Müller et al., 2002; X=I, e.g. Wittrock et al., 2000). ODEs at Ny-Ålesund coincide with elevated mixing ratios of XO essentially only for X=Br. The role of X=Cl is much less clear and X=I apparently lacking correlation. The corresponding integrated concentrations of atomic halogens have been estimated during ODEs (Cl\sim 10^{10} s cm^{-3} and Br\sim 10^{12} s cm^{-3}) using hydrocarbon “clock” measurements (Ramacher et al., 1999; Solberg et al., 1996). In a recent historical survey of the temporal and spatial occurrence of ODEs, Tarasick and Bottenheim (2002) report the incidence of ODEs an order of magnitude lower at Ny-Ålesund (together with e.g. Neumayer) compared to Alert. It has been suggested that the relative low frequency of ODEs at Ny-Ålesund is to some extent related to its local topography and the presence of open oceanic water (Lorenzen-Schmidt et al., 1998).

The purpose of this study was to provide further quantitative information on mercury in different environmental compartments including air, snow and seawater during polar spring and extend the database of GEMDEs. To contribute to the knowledge, we have conducted time-resolved specified determination of mercury in the atmosphere and snow at two altitudes. At the sea level, we focused on collecting data for gas-transfer calculations over snow and sea surfaces. In addition, this paper includes an analysis of meteorological and BrO data obtained by using multiple remote platforms (ground-based as well as satellite-based) and an assessment

Fig. 1. Some Arctic and sub-Arctic mercury sampling locations including long-time monitoring, and average distribution of 50% sea ice in April.
of how these factors influence the evolution of signatures in various samples collected during the campaign. Information needs/data gaps recognized at the first international Arctic atmospheric mercury research workshop 26–28 August 2002 will be addressed and discussed (Schroeder et al., 2003).

2 Experimental section

2.1 Sampling location

Experiments were performed during an intensive campaign (17 April to 14 May 2002) in the presence of 24-h daylight at three locations within the Ny-Ålesund International Research and Monitoring Facility. This establishment (78.9° N, 11.9° E) is situated at the southern shore of Kongsfjorden on the West Coast of Spitzbergen in the Norwegian high Arctic (Fig. 1) and is partly surrounded by mountains with peaks of 1000–1500 m altitude. It is situated about 15 km from the northern-most Barents Sea. An offshoot of the waning Gulf Current (West Spitzbergen current) causes ice-free conditions along the west coast of Spitzbergen during the entire year. Sea ice was in-turn dominant north of a border approximately demarcated by latitude ~80° N and longitude ~10° E (Fig. 1). When covered by ice, the local climate is however modulated to be more continental by the adjacent fjord. In connection with low-lying cloud cover, the presence of distant open water was discernible by noticeably darker skies over water. General large-scale circulation is steered by the Iceland low and the high-pressure systems over Greenland and the Arctic Ocean, forcing warm and humid air from the North Atlantic ocean along cyclone tracks to the Barents sea. The large differences in air temperature between air masses of Arctic or Atlantic origin cause large fluctuation in weather conditions at Ny-Ålesund. Positive phases of the North Atlantic and Arctic oscillations have recently created warming anomalies of circumpolar continental areas that partly are compensated by cooling over the northern North Atlantic. An overall descending tendency in the thickness and area extent of sea-ice is observed e.g. in the Arctic Ocean’s Atlantic layer. Owing to positive feedbacks, greenhouse warming has been proposed to be amplified in the Arctic yielding seasonally increased precipitation.

Samples of airborne mercury and mercury in snow were taken at the Global Atmospheric Watch monitoring observatory operated by the Norwegian Institute for Air Research (NILU) at Zeppelinfjellet (474-m a.s.l). From the Zeppelin mountain Station (ZS, 78°54.42'' N, 11°55.30'' E), there is a steep downhill slope to the north towards the settlement situated at 2-km distance. South of ZS is also a slope whereas there are higher mountain peaks in the east and west. This channels the local wind into either of the uphill directions, north-northwest and south-southeast (Beine et al., 2001). The station is typically positioned well above frequent ground inversion of the fjord valley (Braathen et al., 1990). Air and snow sampling were also carried out downhill at 12-m a.s.l. close to the Italian Consiglio Nazionale delle Ricerche (CNR) Base Arctica “Dirigibile Italia” (DI). The air sampling at this location comprised vertical gradient measurements at two heights and flux chamber measurements on the snow surface. The extensive snow pack in the experimental area is seasonal (usually melted by end of June) and has maritime characterisation with numerous layers of aged snow. Surface seawater samples were collected in Kongfjorden close to or offshore the settlement when manageable by jolly boat.

2.1.1 Sampling and analytical protocols

Airborne mercury

Ambient TGM

Ambient air point samples were collected and analysed with automised instruments. In addition to the stationary Tekran (Model 2537A) gas-phase mercury vapour analyser at Zeppelin, two other Tekran (Model 2537A) instruments were employed to measure TGM. The abbreviation TGM refers to gaseous elemental mercury (GEM) and small contributions of other gaseous mercury species (<1% at ambient conditions) that may pass through the sampling line and be detected as Hg⁰ (Schroeder et al., 1995; Temme et al., 2003). The TGM measurements were made according to the well-established method with a 0.45 µm front particle filter. The stationary instrument was also equipped with a heated sampling line.

Prior to moving to other sampling sites the instruments were calibrated using manual injections of known amounts of Hg⁰(g) from a Tekran 2505 mercury vapour source. The nominal settings of each Tekran’s internal permeation tube were verified within a few %. The internal calibration systems were further used on a daily basis for calibration during the campaign. The instruments were operated in continuous mode at 1.5 L min⁻¹ sample flow rate and with 5 min sampling intervals. The detector response to standards automatically invoked from the internal permeation source showed a high degree of stability from calibration to calibration. Succeeding this exercise, the instruments were in-turn intercalibrated for ambient air with their respective inlet tubes during ~40 h, spanning over a concentration range from <0.1 ng m⁻³ to ambient background.

Transient mercury fractions RGM and Hg-p

RGM and Hg-p were sampled by collection on KCl-coated denuders (Landis et al., 2002; Wängberg et al., 2003a; Xiao et al., 1997) and on filters (Berg et al., 2001; Wängberg et al., 2003b), respectively. An automated unit using sequential sampling of RGM and fine Hg-p (nominal cut-off 3 µm, f-Hg-p) was mounted on the roof of the
Zeppelin station ∼2 m above ground and connected through a heated line to a Tekran 2357A analyser. The transient mercury forms were concentrated and analysed with 1.5-h time step after thermal conversion to Hg\textsuperscript{0}(g), enabling separate quantification after transfer to the Tekran instrument. The temperatures and flows through the sampling components in quartz were controlled by a set of furnaces, pumps and fans programmed by a Setpoint Programmer. The programme was triggered by a signal from a Tekran 1120 Standard Addition Controller. Special attention was payed the reduction of the partitioning of mercury to the walls of the sampling inlet (Ariya et al., 2002; Landis et al., 2002; Sommar et al., 1997) by treating the surfaces with a perhalogenated carbon product. Based on 3σ of the sample blank, the Detection Limit (DL) of RGM and f-Hg-p was estimated to be ∼2 and ∼5 pg m\textsuperscript{−3}, respectively. The fraction of f-Hg-p was typically below the DL and this result will not further be commented. Manual denuders for RGM sampling with lower time resolution were also used. Two manual denuders were used in parallel. The denuders were contained in protective housings thermostatted to 40°C during sampling.

The two sampling systems were identical except that one of the samplers was equipped with Teflon grid inlet. Manual RGM sampling was conducted at ZS in the beginning of the campaign and later moved to DI. Total Hg-p (TPM) was collected at ZS using glass fibre filters according to Berg et al. (2001). Each filter sample was exposed during 1–3 days. All concentrations given are dimensioned to resemble mass of Hg\textsuperscript{0} m\textsuperscript{−3} at STP (0°C and 1013 hPa). Time reported is in UTC (Co-ordinated Universal Time).

2.1.2 Tropospheric BrO

Daily averaged BrO Vertical Column Densities obtained from the GOME instrument were used to interpret the mercury observations. The GOME is a UV/VIS double monochromator spectrometer with a spectral range of 240–793 nm and a resolution of 0.2 nm (λ<400 nm) and 0.4 nm (λ>400 nm), respectively. The instrument operates onboard ERS-2, which is on a sun-synchronous orbit since April 1995 (Burrows et al., 1999). BrO columns are retrieved from the measured radiances using the well-known Differential Optical Absorption Spectroscopy (DOAS) technique. Details on the BrO retrieval can be found in Richter et al. (1998). Ground-based zenith sky UV/VIS – DOAS measurements are also continuously performed at Ny-Ålesund (Wittrock et al., 1996). Since March 2002, the instrument has been upgraded to observe not only the zenith-sky, but also directions close to the horizon. By combining zenith-sky and horizon measurements, some information on the vertical distribution can be retrieved, and the sensitivity to boundary layer BrO explosions is much enhanced (Wittrock et al., 2003). Unfortunately, the instrument was not operational on 22 April, thereby missing part of the large depletion event observed during the campaign.

2.1.3 Mercury in snow

Snow samples were collected by the removal of the uppermost surface layer of the snow pack with a Teflon funnel on a daily basis and brought indoors where they were allowed to melt into acid cleaned 125 ml volume Teflon bottles. Moreover, no specific precipitation-segregated sampling was performed. Each sample was preserved by adding 0.5 ml HCl (Merck, Suprapur). All samples were analysed for total mercury (Hg-tot; Bloom and Crecelius, 1983) and a few for methyl-Hg (Lee et al., 1994).

2.1.4 Mercury in seawater

Surface seawater samples from Kongsfjorden were collected from a small boat or from a landing using an acid cleaned Teflon bottle of 1.3 L volume. The bottle consisted of a 1.0 m long tube with 4 cm inner diameter. The container was equipped with a glass frit in the bottom and had two functions, both sample collector and impinger. Each water sample was immediately transported to the nearby laboratory where 0.5 L of the sample was extracted for Dissolved Gaseous Mercury (hereafter denoted: DGM ≈Hg\textsuperscript{0}(aq)) during 60 min using purified inert gas (0.3 L min\textsuperscript{−1}). The extracted elemental mercury was collected on an Au-trap connected to the outlet at the top of the tube. The Au-trap was heated to 40°C to avoid condensation of moisture. The Au traps were analysed using the standard dual amalgamation and CVAFS detection technique. Detailed information about the procedure, its efficiency and reproducibility is found in Gårdfeldt et al. (2002).

In some cases part of the seawater was transferred into 125 ml volume acid cleaned Teflon bottles before DGM extraction. The samples were preserved with HCl (see Sect. 2.2.3.) and transferred to IVL in Sweden for analysis of Hg-tot.

2.1.5 Methods for mercury flux estimations

Gas-exchange model

A gas-exchange model (Wanninkhof, 1992) was used to estimate mercury evasion fluxes from the sea. The model requires TGM and DGM concentrations and water temperature and wind speed as the input parameters. Further parameterisation was made following the principles reported in Gårdfeldt et al. (2003).

Vertical gradient measurements and flux chamber measurements of TGM

Vertical gradient concentration measurements of TGM were performed at 20 and 180 cm above the snow surface. A teflonised three-way magnetic valve synchronised with the instrument’s preconcentration cycle was used...
enabling periodic alternate sampling from each level with a Tekran 2537A instrument.

Flux chamber measurements were made over the snow surface adjacent to the gradient measurements. The dynamic flux chamber (DFC) was made of FEP Teflon after a prototype from Oak Ridge National Laboratories (Kim and Lindberg, 1995). An external aluminium frame supported the rectangular, open-bottom chamber $60 \times 20 \times 20 \text{ cm}^3$. The flux was estimated using Eq. (3):

$$F = Q(x(C_{out} - C_{in})/A$$

where $Q$ is the total airflow rate pulled through the chamber by means of a pump connected to the chamber wall. The inlet air enters 20 cm above the snow surface through a port opposite to the pump side of the chamber. A is the snow surface covered by the chamber. $C_{out}$ and $C_{in}$ corresponds to the Hg concentration leaving and entering into the chamber, respectively. These two concentrations were measured in sequence using one Tekran 2537A operating in continuous mode with sampling intervals of 5 min. The measurements were initiated by a blank check where the chamber was placed on a FEP Teflon film. After some time when sufficiently low $C_{out}$ - $C_{in}$ values were obtained the measurement was started by putting the chamber on a flat snow plot. The blank procedure was repeated twice during the measurements after which the chamber was relocated on a fresh snow surface. The temperature in the ambient surface snow as well as that within the DFC was probed with embedded thermocouples in the snow. The sensors were connected to a mini meteorological logger (Skye Datahog 2 type SDL 5360).

2.1.6 Meteorological parameters

Meteorological parameters and surface ozone at ZS were measured by NILU (accessible via the Norwegian Institute for Air Research, see http://www.nilu.no/niluweb/services/zeppelin/). Extensive sets of meteorological data collected adjacent to DI and vertical profiles of ozone and temperature above Ny-Ålesund were kindly provided by the German Alfred-Wegener Institute for Polar and Marine Research. The temporal behaviour of TGM and ozone were investigated by calculating $(\Delta [X]/\Delta t)$ ratios. Relative removal rates were calculated using Eq. (4), where $[X]_1$ denotes the concentration of TGM at time $t_1$ etc.

$$\frac{\Delta \log[X]}{\Delta t} = \frac{\log([X]_2/[X]_1)}{t_2-t_1}$$

Back trajectories were obtained using the Hybrid single-particle lagrangian integrated trajectory (HYSPLIT) model. The calculations were made interactively using the FNL meteorological datasets, which is provided on the web by Air Resources Laboratory, National Oceanic & Atmospheric Administration after permission (http://www.arl.noaa.gov/ready/hysplit4.html). The stability of the back trajectories was verified by expanding the starting area or by calculating at different ending heights.

3 Results and discussion

3.1 Variability of local point ozone and local vertically resolved point TGM observations

Time-series of TGM measured at two altitudes are shown in Fig. 2c. Two depletion events (GEMDEs) are clearly depicted. The first and major episode was only measured from ZS, while the following GEMDEs were sampled at both ZS and DI. TGM dropped to undetectable levels in the first and
major event and below 0.5 and 0.9 ng m\(^{-3}\), respectively, during two minor events. The major depletion is characterised by a very fast decrease in TGM; within 24 h the concentration changed from \(\sim 1.6\) ng m\(^{-3}\) to \(< 10\) pg m\(^{-3}\). It is obvious that there also exist other significant spike-dip-like features in the TGM record in-between depletions. During the period between the polar sunrise 2002 and the start of the campaign, no major GEMDE was recorded with the stationary instrument at Zeppelinfjellet. As shown in Fig. 3a, the corresponding probability distribution of TGM is strongly left-skewed by events of low GEM influencing the arithmetical mean concentration (1.15 ng m\(^{-3}\)). The annual TGM mean at the site is 1.4–1.5 ng m\(^{-3}\) as derived from measurements made during recent years (Berg et al., 2001, 2003).

In addition to TGM, ozone was also measured at ZS. The complete ozone concentration-time series is shown in Fig. 2b while Fig. 4a displays the data confined to the major ODEs. The TGM and ozone data sets were compared as 709 paired hourly mean values. They were found to be correlated with maximum positive cross-correlation (R > 0.9) for a time lag of \(< 1\) h. For ambient air data measured by two autonomous instruments, the correlation is very high, but a similar resemblance has been obtained at other polar sites (Schroeder et al., 1997; Temme et al., 2003). By solely including the depletion periods even higher positive 1-h correlation coefficient was obtained (R \(\sim 0.98\)). The Zeppelin station is situated over or at least at the top of the marine boundary layer. The time series of the TGM measurements at 12 m a.s.l. and 474 m a.s.l. are altogether highly correlated as one can anticipate. Conversely to invariant gradients generally observed during GEMDE’s, considerable concentration difference between the stations was often observed successive to depletions (Fig. 2c). During the period 6–8 May the TGM values at DI exhibit a more than fivefold higher variance in comparison to that of ZS. The substantially larger variability in TGM data from DI is illustrated in Fig. 3, where the diurnal variation in TGM at DI and ZS is depicted in a Box and Whiskers diagram. The mean, the upper and lower 95% confidence intervals and extremes are shown for 15 min per bin diurnally integrated samples. The assembled data for DI shows a further distinct maximum around noon, which is influenced by a few days of TGM concentration peaks well above 2.0 ng m\(^{-3}\). Usually no time lag in the concentration variations was monitored. However during the observed depletions, the recovery phase appear to start earlier at the sea level. The vertical spatial gradient in TGM and its temporal development is likely to be due to re-emission of mercury from the snow surface, an effect that is enhanced by
restricted mixing of air on a micrometeorological scale. Similar seasonal and altitudinal ozone measurements have been performed by Lorenzen-Schmidt et al. (1998). In contrast to TGM, there is no evidence for emission from the snow-pack to the atmosphere concerning ozone.

3.2 Compositional change of mercury during and after incidence of GEM depletions

3.2.1 Airborne transient mercury fractions

Concentration-time series of RGM at two altitudes and TPM at ZS are shown in Figs. 2e and d, respectively. The concentrations measured were in the range <2–95 pg m\(^{-3}\) for RGM and <1–140 pg m\(^{-3}\) for TPM. Except for a few samples during GEMDEs the concentrations of these fractions generally were very low (median mixing ratios present in the sub-ppq (parts-per 10\(^{15}\)). The highest values were measured in connection to the first depletion. The sum of RGM and TPM corresponds to 16% of the TGM loss inferred. During GEMDEs year 2000, elevated levels of RGM and TPM also were observed in ozone-depleted air masses received at ZS. The corresponding concentrations extended to 12 and 47 pg m\(^{-3}\), respectively, but corresponded only to up to a few percent of TGM normally present in ambient air. The previous measurements also showed that RGM and TPM generally are low at ZS with concentrations often below 10 pg m\(^{-3}\) (Berg et al., 2001, 2003; Lu et al., 2003).

Within the week of 6 to 12 May, during the two minor GEM depletion events, matching RGM measurements at two altitudes were undertaken. The RGM measured at the higher altitude showed a modest variability around typical background concentrations while the parallel variance was significantly (F-test, \(P<0.01\)) larger at DI and the anti-correlation with TGM during GEMDEs more substantial.

3.2.2 Mercury in snow and seawater

The result from mercury analysis of snow samples, collected at two altitudes (ZS and DI), is shown in Fig. 2f. While Methyl-Hg was consistently undetectable (<0.06 ng L\(^{-1}\)) during the period, time series of total mercury (Hg-tot) presents a moderate maximum during the major GEMDE and becomes highly episodic during the minor events with peaks as high as 77 ng L\(^{-1}\). The strong peaks during the period 5 to 11 May are related to snowfall events. The highest values were obtained during the second depletion in conjunction to snowfall, which started in the late afternoon of 5 May. Two snow samples were collected that day, as shown in Fig. 2f. The ZS sample was collected before the snowfall started and contained small amounts of mercury. The DI sample was collected approximately one hour after the snowfall started and contained high amounts of mercury (~15-fold increase). Hence, this observation clearly indicates that the mercury arrived with the snow.

Some surface water samples from Kongsfjorden were collected during 10–11 May and analysed for Hg-tot. Quite high Hg-tot arithmetic mean concentrations were determined (2.1±1.5 (1\(\sigma\)) ng L\(^{-1}\), n=5). This value can be compared with 0.8 ng L\(^{-1}\) as reported as the corresponding mean for filtered water samples (0.45 \(\mu\)m pore size) collected at the same site during summer (Pongratz and Heumann, 1998). It is likely that the high concentrations observed here are due to deposition of snow enriched in mercury originating from GEMDEs. Only a few other Hg-tot data for Arctic waters have been published (Coquery et al., 1995, “Arctic coastal waters of northern Siberia”).

Surface water samples for analysis of dissolved gaseous mercury (DGM) were also collected at Kongsfjorden. Time series of the DGM concentration during 8–14 May is shown in Fig. 5. A strong diurnal concentration variation is displayed; ranging from 12 to 70 pg L\(^{-1}\). The variation in DGM is clearly linked to solar radiation, which suggests photolytic reduction of oxidised mercury species in the surface water. The DGM values correspond to a supersaturation of 40–480% as estimated using Henry’s Law coefficients valid at low water temperatures (Gärdfeldt et al., 2003).

3.2.3 Evasion of volatile mercury from snow and water surfaces

As discussed above in Sect. 3.1., the observations of diurnal patterns in the surface TGM concentration indicates a light-dependent evasional flux from condensed surfaces. These processes were further investigated by flux measurements using the DFC technique as well as vertical gradient measurements over snow plots outside DI. As shown in Fig. 6b, an accumulation of TGM was generally reached in the air passing through the DFC. This corresponds to midday peak emissions of up to 70 ng m\(^{-2}\) h\(^{-1}\) as calculated from Eq. (3). Elevated fluxes were observed contiguous to
enhanced deposition events during the two latter GEMDEs. The average DFC-flux obtained is $\sim 8 \text{ ng m}^{-2} \text{ h}^{-1}$. It should be noted that blank tests and repositioning of the chamber was concurrent with periods of enhanced mercury deposition. As shown in Fig. 6, a discernable diurnal variation in snow surface temperature exists with temperature maximum during mid-day. When directly sunlit the DFC creates a greenhouse effect that strongly influences the enclosed surface’s temperature. Gillis and Miller (2000) and Wällschläger et al. (1999) have examined the feasibility of DFCs in order to measure mercury gas exchange and found it significantly biased by physical constraints such as turnover rates and the degree of exposure to ambient wind. In papers by Lindberg and co-workers (Lindberg and Zhang, 2000; Zhang et al., 2002) it was established that a high-limit flux is attained when the enclosure is sufficiently ventilated. This limit steady-state is however not directly comparable either in time or in quantity with e.g. micrometeorological approaches. The highest emission rates are concurrent with surface snow temperatures inside the DFC around $0 \degree C$ indicating that reduction and emission occurs mainly in melting snow. Thus, these results may not be significant for ambient snow temperatures (i.e. $T<0 \degree C$).

The vertical two level TGM measurements over snow at DI generally displayed very small concentration differences ($<0.1–0.2 \text{ ng m}^{-3}$) during the period of sampling (3–13 May). They were frequently insignificant if taking the instrumental precision into consideration. Given the position of each intake, the invariant gradients are in accordance with results from Alert reported by Steffen et al. (2002). However, during one event, solid evidence for photo-stimulated emissions of Hg$^0(g)$ from the snow pack was obtained. Results from this period are shown in Fig. 7. The TGM gradients represents 9 h following a snowfall during which a large increase in Hg-tot in snow was observed in consecutive samples on 5 and 6 May (see Fig. 2f). The snow episode ended at ~09:30 the 6 May and was followed by calm winds, a stable temperature at $-10 \degree C$ and mostly overcast skies. During this period with low wind speeds ($<1–3 \text{ m s}^{-1}$ at 2 m above the ground) and strong pulses of short-wave direct solar radiation, a period of frequently significant TGM vertical gradients were observed. Multiple reflections and scattering of shortwave radiation between the surface and the atmosphere enhances the solar flux over the high albedo surface ($\geq 80\%$). Single TGM measurements yielded a difference of 30 to 50% between the lower and higher sampling heights. Rapid photo-induced production of GEM in the snow followed by gas-transfer from interstitial air to the overlying air is thus implied.

The magnitude of the corresponding fluxes during the period was estimated by applying a basic dispersion model (Olofsson et al., 2001). The upwind fetch of the gradient measurements, a snow-covered relatively flat plain without obstacles, was assumed to exhibit homogeneous source strength and surface properties. The contribution from sources at each level sampled ($z_1=1.8 \text{ m}$ and $z_2=0.2 \text{ m}$) was described by the Gaussian plume dispersion equation. If assuming neutral stability and an aerodynamic roughness length of 1.5 cm, the vertical concentration gradient observations are consistent with peak fluxes in the range 20–40 ng m$^{-2} \text{ h}^{-1}$. This interval compares satisfactorily with the figures derived from the present DFC measurements. The model predictions are also consistent with the low range emissions ($<10 \text{ ng m}^{-2} \text{ h}^{-1}$) obtained by the DFC and the observed absence of large TGM vertical gradients over snow.

Fig. 6. Comparison of (a) surface snow temperature within and outside the DFC, respectively, and (b) TGM concentrations in ambient air and air passing through the DFC, respectively.

Fig. 7. Vertical 5’ measurements of TGM at $z_1=1.8 \text{ m}$ and $z_2=0.2 \text{ m}$ over snow pack during 6 May following the major mercury fall-out episode in snow.
Table 1. TGM and ozone depletion temporal ascent and descent rates at 474 m a.s.l. during 2002 depletion events.

<table>
<thead>
<tr>
<th>Start (yy-mm-dd hh:mm)</th>
<th>Stop (yy-mm-dd hh:mm)</th>
<th>Start [Hg] (ng m(^{-3}))</th>
<th>Stop [Hg] (ng m(^{-3}))</th>
<th>Start Ozone (µg m(^{-3}))</th>
<th>Stop Ozone (µg m(^{-3}))</th>
<th>Wind Speed (m s(^{-1}))</th>
<th>Wind Direct. (deg)</th>
<th>Temp. (°C)</th>
<th>(\Delta \log[Hg]/\Delta t) (h(^{-1}))</th>
<th>(\Delta \log[O_3]/\Delta t) (h(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>02-04-20 22:00</td>
<td>02-04-21 00:00</td>
<td>1.47</td>
<td>0.43</td>
<td>81.7</td>
<td>24.3</td>
<td>304</td>
<td>11.9</td>
<td>-6.0</td>
<td>-0.27</td>
<td>-0.26</td>
</tr>
<tr>
<td>02-04-21 07:00</td>
<td>02-04-21 11:00</td>
<td>0.41</td>
<td>1.13</td>
<td>19.1</td>
<td>51.5</td>
<td>322</td>
<td>7.0</td>
<td>-14.4</td>
<td>0.11</td>
<td>0.11</td>
</tr>
<tr>
<td>02-04-21 13:00</td>
<td>02-04-21 15:00</td>
<td>0.94</td>
<td>0.25</td>
<td>46.7</td>
<td>17.3</td>
<td>313</td>
<td>6.3</td>
<td>-15.2</td>
<td>-0.29</td>
<td>-0.22</td>
</tr>
<tr>
<td>02-04-21 16:00</td>
<td>02-04-21 21:00</td>
<td>1.27</td>
<td>0.00</td>
<td>15.1</td>
<td>0.3</td>
<td>295</td>
<td>5.0</td>
<td>-16.3</td>
<td>-0.33</td>
<td>-0.34</td>
</tr>
<tr>
<td>02-04-22 00:00</td>
<td>02-04-22 00:00</td>
<td>0.005</td>
<td>0.12</td>
<td>0.5</td>
<td>4.7</td>
<td>345</td>
<td>2.5</td>
<td>-16.7</td>
<td>0.69</td>
<td>0.49</td>
</tr>
<tr>
<td>02-04-22 06:00</td>
<td>02-04-22 08:00</td>
<td>0.15</td>
<td>0.28</td>
<td>8.9</td>
<td>16.5</td>
<td>212</td>
<td>1.5</td>
<td>-16.1</td>
<td>0.14</td>
<td>0.13</td>
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<td>02-04-22 19:00</td>
<td>02-04-23 01:00</td>
<td>0.14</td>
<td>0.50</td>
<td>16.5</td>
<td>44.1</td>
<td>164</td>
<td>1.4</td>
<td>-17.7</td>
<td>0.09</td>
<td>0.07</td>
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<td>02-04-23 02:00</td>
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<td>0.57</td>
<td>0.73</td>
<td>40.9</td>
<td>49.9</td>
<td>151</td>
<td>1.7</td>
<td>-16.0</td>
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<td>02-05-03 08:00</td>
<td>02-05-04 08:00</td>
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<td>0.77</td>
<td>87.1</td>
<td>46.7</td>
<td>119</td>
<td>7.3</td>
<td>-6.9</td>
<td>-0.01</td>
<td>-0.01</td>
</tr>
<tr>
<td>02-05-04 19:00</td>
<td>02-05-04 21:00</td>
<td>1.12</td>
<td>0.74</td>
<td>55.3</td>
<td>39.1</td>
<td>139</td>
<td>2.0</td>
<td>-11.7</td>
<td>-0.09</td>
<td>-0.08</td>
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<td>02-05-05 04:00</td>
<td>02-05-05 06:00</td>
<td>0.72</td>
<td>0.55</td>
<td>39.7</td>
<td>26.7</td>
<td>135</td>
<td>4.7</td>
<td>-13.2</td>
<td>-0.06</td>
<td>-0.09</td>
</tr>
<tr>
<td>02-05-06 07:00</td>
<td>02-05-06 11:00</td>
<td>0.59</td>
<td>1.24</td>
<td>25.9</td>
<td>60.7</td>
<td>140</td>
<td>3.8</td>
<td>-12.6</td>
<td>0.08</td>
<td>0.09</td>
</tr>
<tr>
<td>02-05-10 09:00</td>
<td>02-05-10 15:00</td>
<td>0.93</td>
<td>1.47</td>
<td>41.3</td>
<td>64.7</td>
<td>288</td>
<td>4.5</td>
<td>-5.5</td>
<td>0.03</td>
<td>0.03</td>
</tr>
<tr>
<td>02-05-11 01:00</td>
<td>02-05-11 10:00</td>
<td>1.18</td>
<td>1.50</td>
<td>57.3</td>
<td>83.7</td>
<td>290</td>
<td>5.8</td>
<td>-5.9</td>
<td>0.01</td>
<td>0.02</td>
</tr>
</tbody>
</table>

TGM fluxes obtained at Alert during spring 2002 using DFC and micrometeorological methods yielded similar values. At Point Barrow, highly variable emission rates were observed by DFC during snowmelt 2002 ranging up to 0.6 µg m\(^{-3}\) h\(^{-1}\) (Schroeder et al., 2003).

The mechanism behind the formation of volatile mercury in the sunlit snow is not well understood. The reactions are presumably abiotic and involve reactants present on the ice crystal surface. In a study by Lalonde et al. (2003), it was indicated that newly deposited mercury was highly labile in snowpacks due to Hg(II) photoinduced reduction to volatile Hg\(^0\). The environmental feasibility of aqueous reduction mechanism of mercuric species was investigated and discussed by Gårdfeldt et al. (2003). It was concluded that Hg\(^0\) is not formed through consecutive one-electron reductions under aerated conditions. However, formation of Hg\(^0\) can occur for example via fragmentation of an organic ligand bound to Hg(II) under actinic radiation.

The seawater in Kongsfjorden was supersaturated in respect to DGM as mentioned above, which suggests reemission of mercury from the water surface as well. Gas-exchange model calculations yielded flux values from 0.1 to 7 ng m\(^{-2}\) h\(^{-1}\). It should, however, be mentioned that the bay was partly covered by ice which limited the wave field. The flux values obtained are therefore likely to be overestimated.

3.3 Surface depletions, occurrence of tropospheric BrO and the influence of transport

3.3.1 Local boundary-layer chemical and meteorological observations

Both ODEs and GEMDEs have recently been assumed to be a result of halogen chemistry as mentioned above. The occurrence of BrO is frequently used as a tracer for such processes. As can be seen in Fig. 2b, ground based DOAS BrO measurements at Ny-Ålesund revealed incidence of elevated BrO column densities simultaneous to the periods of low Hg\(^0\) and O\(_3\) surface levels. This initial signature hints that air parcels at this receptor may exhibit an augmented oxidation capacity. In order to evaluate the role of transport and transformation, various regimes of the TGM/O\(_3\) concentration-time series were evaluated. Features in composition change are generally reproduced for both gas-phase constituents but during May the patterns became increasingly less correlated. Events of concurrent significant variations in TGM and O\(_3\) concentration are given in Table 1 together with corresponding meteorological information. Shift in meteorological parameters is frequently encountered during the on-set of depletions and most utterly marked during the most intense depletion (see Fig. 4c–f). The on-set of the first depletion is associated with falling temperature and raising wind speed (ΔT≈−15 °C and Δu≈10 m s\(^{-1}\) during a few hours) and rapid shift in wind direction from SE to N-NW indicating intrusion of an Arctic airmass. It should be noticed that the local wind-direction during phases of the most distinct changes in TGM were strictly in-between WNW and NNW. This represents a sector almost entirely free from adjacent land masses (Fig. 1). In correspondence to observations made by Solberg et al. (1996) covering ODE’s at Ny-Ålesund during 1988–1993, the on-set of the present GEMDEs is associated with a cold boundary layer. Contrary to the suggestions given by Lu et al. (2001), depletions at Ny-Ålesund appear not to be restricted to stagnant synoptic situations (Solberg et al., 1996; Tuckermann et al., 1997). The major depletion episode indeed concurred under strong winds as shown in Fig. 4.

The temporal descents in TGM and O\(_3\) (Δ[X]/Δt) were during the on-set of the major GEMDE in the order of ∼0.5 ng m\(^{-3}\) h\(^{-1}\) and ∼30 µg m\(^{-3}\) h\(^{-1}\), respectively. The latter value exceeds the most rapid variation features of ODEs reported previously for this site. The very fast drop
may be interpreted as a combination of halogen chemistry and fast advection of an airmass extending to the surface that has already been affected by depletion for an extensive period of time. The relative variations of $O_3$ and TGM were evaluated by calculating $\Delta \log[X]/\Delta t$ ratios for the relevant period of time (see Table 1). It is evident that there exists a near 1:1 relationship between $O_3$ and TGM in most of the cases. This suggests a common meteorological influence by e.g. advection or turbulent dispersion. Nevertheless, due to the presence of moderately elevated BrO concentrations (a few ppt), bromine chemistry forming BrO must have occurred in the air mass. It is however not certain that ozone is a limiting factor.

3.3.2 Remote-sensing circumpolar BrO observations, trajectory and sea-ice intersect analysis

To further test whether the low TGM concentrations observed might be a result of transport of air masses already depleted from mercury, back trajectories were calculated and compared with daily average BrO vertical column density maps obtained from the GOME satellite. The results from this investigation regarding the first and the second depletion are shown in Fig. 8. The air received at Ny-Ålesund the 22 April (the day with maximum depletion) arrived via a north-north-west path as shown in Fig. 8a. The trajectory is of four days length and originates from the Western High Arctic Sea.

BrO vertical column densities mapped in Fig. 8a are from 18 April, the starting day of the trajectory. According to the GOME BrO maps related to the days in-between the 18 to 22 April the trajectory was not passing though areas with elevated BrO. Hence, this would suggest that the air mass with depleted mercury originated far from Ny-Ålesund. However, although not observed by the GOME satellite (BrO GOME data directly above Ny-Ålesund are not available from the 22 April) the ground based BrO measurements proved that the air mass actually contained elevated concentrations of BrO.

The climax of BrO concentrations concurrent with the very intrusion of the cold air mass indicates a sustained chemical oxidation of $Hg_0(g)$. This implies that the transition phase with active oxidation of GEM must have happened close to Spitzbergen. Conversely, physical removal of the majority of the airborne mercuric products onto snow and ice is a process requiring some time which therefore suggests that most of the GEM must have been oxidised quite far from Ny-Ålesund. Figure 8b shows a back trajectory ending at Ny-Ålesund at the 24 April. The GEM concentration was subsequently almost back to normal and as shown the air received does not originate from areas with elevated BrO. According to the back trajectories calculated there is no air transport between Ny-Ålesund and the area with high BrO column densities seen north west of Greenland. The next episode with low GEM concentrations occurred in the beginning of May, with a GEM minimum at the 5 May, see Fig. 8c. According to the BrO maps from this period, the low GEM concentration also seems to be a consequence of transport of an air mass depleted from mercury. However, considerable
Table 2. Observations of Hg-tot in snow for some Arctic coastal and marine sites (see Fig. 1 for some of the locations).

<table>
<thead>
<tr>
<th>Coastal or Marine Site</th>
<th>Hg-tot in surface snow (ng L⁻¹)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beaufort Sea Barrow (71°19' N, 156°37' W)</td>
<td>~90 (GEMDEs, May 2001) &gt;400 (Falling snow, GEMDE) ~300 (Surface snow, GEMDE)</td>
<td>Lindberg et al. (2002) Brooks et al. (2002)</td>
</tr>
<tr>
<td>Admiralty Bay (70°54' N, 155°49' W)</td>
<td>10±1 (04/28/96) 11±4 (04/27/96) 23±2 (04/28/96)</td>
<td>Garbarino et al. (2002)</td>
</tr>
<tr>
<td>Dease Inlet (71°06' N, 155°49' W)</td>
<td>23±2 (04/28/96)</td>
<td></td>
</tr>
<tr>
<td>Ice Station SHEBA</td>
<td>185 (light, GEMDEs)</td>
<td></td>
</tr>
<tr>
<td>Chukchi Sea Kasegaluk Lagoon (69°22' N, 163°05' W)</td>
<td>100±60 (04/26/96)</td>
<td>Garbarino et al. (2002)</td>
</tr>
<tr>
<td>(70°12' N, 162°10' W)</td>
<td>214±5 (04/26/96)</td>
<td></td>
</tr>
<tr>
<td>Eastern Canadian Arctic, Hudson Bay, and Greenland</td>
<td>25–160</td>
<td>Lu et al. (2001)</td>
</tr>
<tr>
<td>Hudson Bay Kuujjuarapik (55°30' N, 77°73' W)</td>
<td>82 (GEMDEs, April 2000)</td>
<td>Poissant et al. (2001)</td>
</tr>
<tr>
<td>Greenland Sea/Barents Sea Scoresbysund (~71° N, ~22° W)</td>
<td>&lt;2 (Spring 1997, 1998)</td>
<td>Lu et al. (2001)</td>
</tr>
<tr>
<td>Ny-Ålesund (78°54' N, 11°53' E)</td>
<td>10 (GEMDEs, April 2000) 38 (GEMDEs, April 2001) 20–77 (GEMDEs, April–May, 2002)</td>
<td>Berg et al. (2003) This paper</td>
</tr>
</tbody>
</table>

amounts of oxidised mercury were transported to the measurement area via snow, as mentioned above.

Following the approach of Temme et al. (2003), 3-day backward trajectories were interactively generated to describe diurnal (12-h interval separated) transport of air masses to Ny-Ålesund during the entire period. The transport denoted by twelve 6-h interval positions for each trajectory were indexed by the degree of sea ice coverage if such were available. The weight factors were obtained from the ice charts of the US-National Ice Center (http://www.natice.noaa.gov/index.htm). In Fig. 9, the concentration/density-time series of TGM, O₃ and BrO are spread above the surface of the summed-up 6-h resolved index (Σ ice coverage, base unit 10%). It gives an unambiguous signal that the onset of measured GEMDE’s coincides with the arrival of air masses with high degree of sea-ice transect.

The sea-ice transport index is surprisingly constant in the slow recovery phase of Hg₀(g) during the major GEMDE’s. The apparent recovery phase (22 April–) is associated with local transport over the barren interior of Spitzbergen Island that may well include mixing of air-masses (mixing-in from above with mercury-rich air from the free troposphere). Analogous to ODEs, the influence from local topography may thus obscure the perception of the scale of the phenomenon, making local GEMDEs at Ny-Ålesund display shorter duration and lower magnitude (Solberg et al., 1996).

4 Concluding remarks

The measurements made during spring 2002 strongly suggest that the observed GEMDEs were caused by transport of air already depleted from mercury rather than the action of solely atmospheric chemistry in the air at Ny-Ålesund. One of the findings proposing transport is that the GEMDEs occurred in conjunction with weather changes. The major depletion for example, was initiated when the wind direction changed from south to north and a cold air mass with wind speeds above 10 m s⁻¹ was received. Since some time
is needed for an air mass to become depleted from oxidised forms of mercury via deposition of RGM and the particulate fraction to ice and snow surfaces, the low content of airborne oxidised mercury constitutes an additional evidence for transport. During the major depletion the Total Airborne Mercury (TAM=Σ(GEM, RGM, TPM)) corresponded to less than 20% of the normal background value of TGM, indicating a >80% loss of airborne mercury. This is considerably lower in comparison to observations made at Nunavut and Alaskan sites (Steffen and Schroeder, 2002; Lindberg et al., 2002) where TAM values corresponding to 70–90% of the background TGM concentration have been observed. With marked referral to the publications by Brooks et al. (2002), Garbarino et al. (2002), Lindberg et al. (2002a) and Lu et al. (2001) respectively covering the American Arctic, Table 2 hints that the deposition pattern here appears to be stronger after Polar sunrise. The GOME BrO maps and back trajectories (Fig. 8) also support transport. The ground based BrO data indicate that elevated BrO concentrations can be sustained in an ozone depleted air mass. Low ozone values are large compared to BrO concentrations (a few ppt at most) possibly implying that ozone concentrations are just not the limiting factor for BrO formation.

As Hg(g) generally exhibit a uniform distribution in the polar lower atmosphere, its VCD below a height of

\[ \int_0^z c(z) \, dz. \]

can readily be estimated. The VCD of TAM lost during MDEs relative to an initial standard background atmosphere ought to be in the interval 0.1–1 µg m^{-2} (Banic et al. 2003; Lindberg et al. 2002). The specific atmospheric profile measurements of mercury at Ny-Ålesund during MDE 2002 imply the lost VCDs to be in the upper range. Assuming peak Hg-tot concentration in surface snow (~60 ng/L, density 0.4) derived from a single MDE, the conservation of mass is attained roughly in a depth interval of 5–10 cm. However, in order to quantitatively address the dynamic cycling of mercury at this site in a credible way, a more rigorous sampling protocol is required than that exercised during this campaign. This should at least involve temporally and spatially resolved speciated measurements in boundary layer troposphere as well as in the snow-pack. Still by observing a quick decline of Hg-tot in surface snow and a discrepancy in TGM between the ZI and DI stations in the aftermath of a GEMDE, we can conclude that a significant part of the oxidised mercury deposited onto snow is quickly re-emitted back to the atmosphere as elemental mercury. Here we report evidence for a similar process occurring in seawater. The concentration of DGM in the surface water in Kongsfjorden showed a pronounced daily variation well correlated with solar radiation. Hence, it seems like a significant part of the oxidised mercury deposited onto snow and water surfaces is quickly re-emitted back to the atmosphere as elemental mercury. The ratio between deposition and re-emission is an important parameter that determines the impact of mercury depletion in the Arctic environment. It is now clear that GEMDEs constitute a phenomenon of large geographical scale in Polar Regions. More research is needed to improve our understanding of this spectacular aspect of environmental mercury cycling.

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