Mercury fluxes over an Australian alpine grassland and observation of nocturnal atmospheric mercury depletion events

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Abstract. Aerodynamic gradient measurements of the air–surface exchange of gaseous elemental mercury (GEM) were undertaken over a 40 ha alpine grassland in Australia’s Snowy Mountains region across a 3-week period during the late austral summer. Bi-directional GEM fluxes were observed throughout the study, with overall mean value of $0.2 \pm 14.5$ ng m$^{-2}$ h$^{-1}$ and mean nocturnal fluxes of $-1.5 \pm 7.8$ ng m$^{-2}$ h$^{-1}$ compared to diurnal fluxes of $1.8 \pm 18.6$ ng m$^{-2}$ h$^{-1}$. Deposition velocities ranged from $-2.2$ to $2.9$ cm s$^{-1}$, whilst ambient GEM concentrations throughout the study were $0.59 \pm 0.10$ ng m$^{-3}$. Cumulative GEM fluxes correlated well with 24 h running mean soil temperatures, and one precipitation event was shown to have a positive impact on diurnal emission fluxes. The underlying vegetation had largely senesced and showed little stomatal control on fluxes. Nocturnal atmospheric mercury depletion events (NAMDEs) were observed concomitant with O$_3$ depletion and dew formation under shallow, stable nocturnal boundary layers. A mass balance box model was able to reproduce ambient GEM concentration patterns during NAMDE and non-NAMDE nights without invoking chemical oxidation of GEM throughout the column, indicating a significant role of surface processes controlling deposition in these events. Surface deposition was enhanced under NAMDE nights, though uptake to dew likely represents less than one-fifth of this enhanced deposition. Instead, enhancement of the surface GEM gradient as a result of oxidation at the surface in the presence of dew is hypothesised to be responsible for a large portion of GEM depletion during these particular events. GEM emission pulses following nights with significant deposition provide evidence for the prompt recycling of 17% of deposited mercury, with the remaining portion retained in surface sinks. The long-term impacts of any sinks are however likely to be minimal, as cumulative GEM flux across the study period was close to zero.

1 Introduction

International support for a legally binding agreement on the control of mercury in the environment, beginning in 2003, recently culminated in the 2013 United Nations Environmental Programme (UNEP) Minamata Convention on Mercury (Kessler, 2013). This convention includes provisions for emissions reductions; technology sharing; public awareness; and enhanced mercury monitoring in human populations, wildlife and the environment. Such monitoring is essential in assessing the efficacy of emissions reduction measures and improving the understanding of the dynamics of the global mercury cycle (Selin, 2009). These dynamics – including methylation, oxidation/reduction, dry/wet deposition and emission/re-emission – all have implications for the transport and environmental legacy of previously emitted mercury (Pirrone et al., 2010). The burden of this legacy is likely to be felt to varying degrees by all nations as, due to its long atmospheric lifetime, mercury is known to be distributed across hemispheric scales (Lin and Pehkonen, 1999; Holmes et al., 2010), and environmental reactions to changes in atmospheric loading may take place on the order of years to centuries (Meili et al., 2003; Lindberg et al., 2007). These concerns point to the necessity of an enhanced research effort to better understand the complexities of mercury biogeochemical cycling (Jaffe et al., 2014).

Although atmospheric mercury is known to be transported across hemispheric scales, it is understood that it can undergo deposition to, and rapid re-emission from, terrestrial surfaces...
a process referred to as prompt recycling (Selin et al., 2008; Selin, 2009). The impact of this process has been estimated to contribute 256–1400 Mg a\(^{-1}\) to worldwide emission of mercury from terrestrial surfaces, based on 20 % prompt re-emission of deposited mercury (Selin et al., 2008; Kikuchi et al., 2013; Simone et al., 2014). Evidence for this process came from isotopic measurements (Hintelmann et al., 2002; Graydon et al., 2006), where newly deposited mercury was found to be more susceptible to re-emission to the atmosphere on a scale of days to months. Data from the Mercury Experiment to Assess Atmospheric Loadings in Canada and the US (METAALICUS) study suggest between 5 and 40 % of mercury newly deposited over terrestrial surfaces is promptly recycled (Hintelmann et al., 2002; Amyot et al., 2004; Selin et al., 2008). Further evidence has since been compiled during observations taken during atmospheric mercury depletion events (AMDEs), which are characterised by the rapid oxidation of gaseous elemental mercury (GEM) to the more reactive gaseous oxidised mercury (GOM) and subsequent deposition to the surface (Steffen et al., 2008, and references within). Flux studies by Lahoutifard et al. (2005) and Kirk et al. (2006) suggested net GEM emission from the surface following AMDEs, though others have seen very little subsequent GEM emission, or even net deposition to the surface (Brooks et al., 2006; Steffen et al., 2008). Further evidence for prompt recycling using surface GEM flux measurements is difficult to establish, as it is generally not possible to distinguish between mercury emitted from long-term stores and mercury re-emitted following atmospheric deposition (Gustin et al., 2006).

Despite the global nature of atmospheric mercury pollution both long-term monitoring and air–surface exchange research of this powerful neurotoxicant are weighted heavily towards industrialised countries in the Northern Hemisphere (United Nations Environmental Programme, 2013; Agnan et al., 2016). There is a lack of atmospheric mercury data taken within the Southern Hemisphere and in particular for Australia. Modelling efforts by Nelson et al. (2012) suggest that anthropogenic mercury emissions throughout Australia are around 10 to 20 Mg a\(^{-1}\), compared with 95 to 285 Mg a\(^{-1}\) emitted/re-emitted from natural sources. As patterns of wet and dry deposition of mercury are known to exhibit regional variability due to anthropogenic emissions (Pirrone et al., 2010), Australia provides a valuable opportunity for atmospheric mercury biogeochemical research as it is relatively unimpacted by anthropogenic sources. Atmospheric dry-deposition rates across remote regions of Australia were estimated by Nelson et al. (2009) to range between 20 and 70 µg m\(^{-2}\) a\(^{-1}\), based on background GEM pool concentrations of 1.2 ng m\(^{-3}\). Available empiricisms for such modelling efforts however are derived from Northern Hemisphere observations that may not apply to Australia’s unique ecosystems (Nelson et al., 2009, 2012; Edwards and Howard, 2013). Furthermore, recent evidence has shown that background atmospheric GEM pool concentrations in the Southern Hemisphere are considerably lower than those in the Northern Hemisphere (Slemr et al., 2015), which may mean that delivery of atmospheric mercury to ecosystems in the Southern Hemisphere is currently overestimated. A recent review of mercury air–surface exchange research by Agnan et al. (2016) found only one Australian mercury flux study (Edwards and Howard, 2013) and showed there was a general lack of flux data taken over vegetated surfaces globally.

This study was undertaken over a grassland in the Snowy Mountains region of Australia’s alpine country, during the austral summer. The site was chosen as it is the location of an ongoing OzFlux greenhouse gas net ecosystem exchange site. The micrometeorological aerodynamic gradient method was employed to obtain a high-time-resolution flux time series that did not disturb the natural ecosystem studied. This study furthers our understanding of Australian mercury cycling by providing additional background ambient and air–surface exchange data, and a comparison with similar existing flux datasets undertaken in alpine regions elsewhere. Further, we provide additional data on the process of prompt recycling observed during this study.

2 Methods

2.1 Site description

This study took place on Nimmo Plains (36°12′57″ S 148°33′10″ E, elevation 1340 m a.s.l.) in Australia’s Snowy Mountains region, bordering the eastern boundary of Kosciuszko National Park. Measurements were taken between 28 January and 18 February (DOY 28 to 49). Nimmo Plains is a ~40 ha region of reasonably level terrain (within 20 m elevation change) and the site of an ongoing OzFlux ecosystem research site. Vegetation immediately within the Kosciuszko National Park consists of dry sclerophyll forests that extend for a distance of at least 5 km. On the plain, vegetation consists predominantly of grasses of various Poa spp. that had largely senesced prior to this study. These grasses had been trampled and grazed by cattle to an average height of approximately 2 cm prior to the study taking place. Rainfall over the preceding 3 months (NDJ) was in the lowest 20th percentile since 1907, resulting in dry conditions. The plain is located 23 km to the north and 51 km west of the towns of Jindabyne (918 m a.s.l.; 2011 population: 1727) and Cooma (800 m a.s.l.; 2011 population: 6301) (Australian Bureau of Statistics, 2012). There are no known significant natural or anthropogenic mercury sources in the region.

2.2 Substrate characterisation

Substrates were characterised using samples taken in each of the four cardinal directions at distances 10, 30 and 50 m from the flux sampling tower. Surface vegetation, along with soils at depths of 0 to 2, 5 to 10 and 10 to 20 cm were sampled us-
ing clean equipment and stored in double-sealed plastic bags. Vegetation and soil samples were dried at 105 °C for 24 h, and soil samples were sieved into 2 to 19 mm (granules), 63 µm to 2 mm (sand) and < 63 µm (silt/clay) size fractions. Total mercury (THg) contents of soil and vegetation samples were determined in triplicate using a DMA-80 direct mercury analyser (Milestone Inc., Shelton, CT, USA) and US EPA Method 7473. Instrument precision was verified to within 5 % using five repetitions of 100 µL of a 100 ppb Hg standard between each run of 60 replicates. Accepted sample boat blank concentrations were < 0.002 ng THg. Soil organic carbon (SOC) and soil inorganic carbon (SIC) contents were determined according to the loss-on-ignition method described by Nelson and Sommers (1996). SIC values were multiplied by a correction factor of 1.36 under the assumption that it existed predominantly as carbonate (Bengtsson and Enell, 1986). Soil pH was determined using a 1 : 5 soil / water suspension and CyberScan pH 300 (Eutech Instruments, Vernon Hills, IL, USA).

2.3 Flux instrumentation and methods

Eddy covariance flux measurements of sensible and latent heat, carbon dioxide (CO2) and momentum were taken at a height of 2.42 m using a CSAT-3 sonic anemometer (Campbell Scientific, Logan, UT, USA) and LI-7200 closed-path, infrared gas analyser (Li-Cor Biosciences, Lincoln, NE, USA). Samples were collected at 20 Hz using data capture software developed in-house using LabVIEW (National Instruments, Austin, TX, USA). Post-processing of eddy covariance fluxes and quality control were undertaken using Li-Cor EddyPro 5.2.1. The convention of positive values representing fluxes upward from the surface is used herein. The planar fit method of Wilczak et al. (2001) was applied across the entire dataset in order to rotate sonic anemometer data into mean wind coordinates, and ogive analyses were prepared across the longest continuous measurement period (5.4 days) according to Foken et al. (2006) in order to determine an appropriate flux averaging period. Quality control flags were calculated for each averaging period using the scheme of Foken et al. (2004), and fluxes given a flag 2 were discarded from further analyses. Additionally, flux values beyond the mean ±3 standard deviations were deemed outliers and removed. Tests for significance were performed using Student’s two-sample t test, and use of the ± symbol hereafter signifies 1 standard deviation. Unless otherwise stated, significance is assessed at p < 0.05.

GEM fluxes were determined using the method and assumptions outlined in Edwards et al. (2005). Flux calculations were undertaken according to Eq. (1), where C(z2) represents GEM concentration at height z2, \( u_a \) friction velocity, \( k = 0.40 \) the von Kármán constant, \( d \) the zero-plane displacement height and \( \psi(z_1) \) the integrated universal similarity functions as given by Businger et al. (1971).

\[
F = \frac{k u_a (C(z_2) - C(z_1))}{\ln(z_2 - d) - \ln(z_1 - d)}
\]

(1)

GEM gradients were constructed from samples taken at two heights \( z_1 = 0.48 \text{ m and } z_2 = 0.73 \text{ m} \) and quantified using a Tekran 2537B (Tekran Instruments, Knoxville, TN, USA), with detection limit of 0.01 ng m\(^{-3}\) and reference volumes reported at 1 atm and 0 °C. Density corrections due to water vapour were undertaken according to Lee (2000). Sample air was drawn from the sample inlets through a 0.2 µm polytetrafluoroethylene (PTFE) filter and a single PTFE tube of length 14 m by a PTFE pump drawing at 10 L min\(^{-1}\). The 2537B sub-sampled from this flow through an additional 0.2 µm PTFE filter. Switching of sampling between the in-takes took place every 10 min (2 × 2537B samples) and was achieved using a PTFE solenoid valve controlled by the LabVIEW software. The random error in the gradient induced by the cumulative electronic and sampling delays following each switch of the solenoid was calculated to be 3 %. Calibration of the 2537B was undertaken 19 times throughout the study using the internal mercury permeation source. Verification of the permeation source was undertaken in the lab before and after the study using manual injection of mercury vapour to within 2 %.

2.4 Ancillary data

In addition to GEM fluxes, ambient GEM (at 3.1 m height) was quantified with a Tekran 2537A sampling at 1 L min\(^{-1}\) through an unheated 5 m PTFE tube and 2 × 0.2 µm PTFE filters. Calibration of the 2537A was undertaken every 23 h using the internal mercury permeation source, verified in the same manner as the 2537B. Comparison of the two 2537 instruments in the lab showed good agreement (< 4 % systematic difference) across a range of concentrations spanning 0 to 12 ng m\(^{-3}\). Ambient ozone (O\(_3\)) was sampled at the same height using a Thermo Scientific 49i (Thermo Fisher Scientific, Waltham, MA, USA) sampling through a separate, unheated 5 m PTFE tube. Incoming and outgoing solar and ter-restrial radiation were measured with a Campbell Scientific CNR1 net radiometer positioned at 1.5 m. Soil temperature was measured 2 cm beneath the surface with two Campbell Scientific TCAV temperature-averaging probes and soil volumetric water content at a depth of 5 cm with a Campbell Scientific CS615. Soil and radiation data were collected and stored using a Campbell Scientific CR3000 data logger.

Back trajectories and mixed-layer heights (MLHs) were modelled using NOAA’s Single Particle Lagrangian Integrated Trajectory Model (HYSPLIT; Draxler, 1999; Draxler and Hess, 1998; Stein et al., 2015) with GDAS 0.5° data as inputs. Aerodynamic (\( g_a \)) and bulk canopy (\( g_c \)) conductances were calculated using Eqs. (2) and (3), by rearranging the
Penman–Monteith equation (Monteith and Unsworth, 1990) in the same manner as Converse et al. (2010).

\[ g_a = \frac{\ln \left( \frac{z_d - \Psi_m}{z_o} \right) - \Psi_m}{k^2 u_r} \left[ \frac{\ln \left( \frac{z_d - \Psi_m}{z_o} \right) - \Psi_m}{k^2 u_r} \right]^{-1} \]  \tag{2} \\

\[ g_c = \left[ \left( \frac{\Delta T}{\gamma g_a} \right) \left( \frac{R_n - G}{\lambda E} - 1 \right) + \frac{\rho_a c_p D_a}{\gamma \lambda E} \right]^{-1} \]  \tag{3}

Here \( \Delta \) is the slope of the vapour pressure curve at temperature \( T \), \( \gamma \) the psychrometric constant, \( R_n \) net radiation, \( G \) soil heat flux, \( \lambda E \) latent heat flux, \( \rho_a \) air density, \( c_p \) specific heat of air at constant pressure, \( D_a \) vapour pressure deficit and \( z_0 \) the roughness length; the subscripts \( m \) and \( v \) represent momentum and water vapour, respectively.

We followed the method of Converse et al. (2014) and modelled dew depth on the basis of the surface energy balance methodology described in Jacobs et al. (2006). For each time step \( i \) of length \( \Delta t \), the depth of dew \( (D_{t+1}) \) was calculated using

\[ D_{t+1} = D_t + E_i \Delta t \quad \text{if} \quad D_{t+1} \geq 0 \]

\[ D_{t+1} = 0 \quad \text{if} \quad D_t + E_i \Delta t < 0. \]  \tag{4}

where \( E \) is the dew flux density calculated from Eqs. (4)–(7) in Jacobs et al. (2006), derived from the surface energy balance using Penman’s substitution. Surface temperatures were calculated from CNR1 outgoing long-wave-radiation data using the Stefan–Boltzmann equation and an emissivity of 0.95 (Öke, 1987).

A simple box model was employed in order to investigate the influence of observed nocturnal GEM deposition. This model took MLH from HYSPLIT outputs \( H \) and imposed a constant entrainment velocity \( w = 0.005 \, \text{m s}^{-1} \) between the mixed layer and free troposphere. No chemical production or destruction of GEM was incorporated in the model, as this would have led to speculation regarding unknown concentrations of important oxidative and reductive species. The model was started for each day at sundown (net shortwave \( < 20 \, \text{W m}^{-2} \)), and the GEM concentration in the free-tropospheric \( C_{\text{ft}} \) was set as the observed mean GEM concentration across the previous hour. The modelled GEM concentration \( C \) was then calculated using Eq. (5).

\[ C_{t+1} = C_t + (F_{t+1} + (C_{f_t} - C_t)w) \cdot \frac{\Delta t}{H} \]  \tag{5}

### 3 Results and discussion

#### 3.1 Observations and trends

#### 3.1.1 Site characterisation

Substrate sampling confirmed background (<100\( \mu \text{g kg}^{-1} \); Gustin et al., 2006) THg levels, with vegetation shown to have average concentrations of 18\( \pm \)3\( \mu \text{g kg}^{-1} \) (\( n = 12 \)). THg content in the soil was found to be 48\( \pm \)9\( \mu \text{g kg}^{-1} \) (\( n = 36 \)) and did not vary significantly with depth or size fraction. Soils were predominantly sandy, with silt and clay making up 7\% of mass below 2 mm, and mildly acidic (pH = 4.9). No soil material was found to have a diameter greater than 19 mm. SIC was spatially uniform both horizontally and vertically (6\% \( n = 36 \)), whereas SOC varied with depth, from 15\% (\( n = 12 \)) in the upper 2 cm to 8\% (\( n = 12 \)) in the 10–20 cm layer. Fibrous root matter was observed to extend to a depth of around 10 cm.

#### 3.1.2 Meteorological observations

Weather over the study period was dominated by high-pressure systems, punctuated by weak troughs that brought patchy cloud and minor showers (Fig. 1). Temperatures ranged between 2 and 28\( ^\circ \text{C} \), with an average overnight minimum of 6\( ^\circ \text{C} \) and average daily maximum of 22\( ^\circ \text{C} \). Winds were primarily from the higher, forested Kosciuszko range to the west, peaking at 8 m s\(^{-1} \), though nights were predominantly calm and stable. Following the calmest of these nights, fog events were witnessed and/or dew was observed. As no instrumental measurements of either fog or dew took place, such events were manually noted with the first observations of the day, taken at sunrise. These observations are noted in Fig. 1. Nights following which both dew modelling and observations at sunrise confirmed the existence of dew are hereby termed “dew nights” (\( n = 8 \)), and those where the existence of dew may or may not have been indicated from modelling but manual observations could not corroborate this are hereby “non-dew nights” (\( n = 10 \)). Measurements had failed during two of the nights, and therefore there were insufficient data to undertake dew modelling.

#### 3.1.3 GEM concentrations

GEM concentrations throughout the study were 0.59\( \pm \)0.10 ng m\(^{-3} \). Diurnal GEM concentrations were slightly higher than the mean and less variable at 0.63\( \pm \)0.08 ng m\(^{-3} \). Nocturnal GEM concentrations were 0.54\( \pm \)0.10 ng m\(^{-3} \). Wind direction and HYSPLIT analyses showed that there was no influence from significant GEM sources. These values are lower than annual mean sea-level measurements recently reported across the Southern Hemisphere (Slemr et al., 2015; Howard et al., 2017). Analysis of the systematic uncertainty of the 2537 system by Slemr et al. (2015) suggests this can be on the order of
0.1 ng m\(^{-3}\). With this uncertainty taken into consideration, GEM concentrations observed over the study were significantly \((p < 0.0001)\) lower than for all four sites operating in the same year (Cape Point, Amsterdam Island, Cape Grim, Troll; Slemr et al., 2015). This comparison is only against annual means for these sites and cannot account for any seasonal or regional variation. More recent observations of ambient GEM in Sydney \((33^\circ 45'55''S, 151^\circ 07'04''E;\) elevation: 59 m) over the period February–September 2015 gave a mean value of 0.64 ng m\(^{-3}\) (Sawyer, 2016).

Linear correlation with environmental correlates using all available data was generally stronger for ambient GEM than for GEM fluxes (Table 1). The strongest significant relationship was with specific humidity \((r = 0.51)\), followed by O\(_3\) \((r = 0.49)\) and air temperature \((r = 0.47)\). For diurnal data, correlation was maintained with specific humidity \((r = 0.52)\), though it decreased for O\(_3\) \((r = 0.37)\) and air temperature \((r = 0.26)\). Peak diurnal O\(_3\) followed a similar trend to peak MLH (Fig. 1b), whilst diurnal GEM changed very little, further confirming the absence of local sources. For nocturnal data, correlation with both O\(_3\) \((r = 0.41)\) and air temperature \((r = 0.50)\) were higher than for diurnal data. Nocturnal wind speed was also significantly correlated \((r = 0.56)\), as was specific humidity \((r = 0.60)\). Each of these variables showed significant decreases on some nights, though not on others. These decreases will be discussed further in Sect. 3.3.

### 3.2 GEM fluxes

#### 3.2.1 Quality control

Ogive analyses on wind, temperature, CO\(_2\) and H\(_2\)O showed that most scalar fluxes were convergent within 20 min and that low-frequency contributions at this site had little influence on fluxes, up to 150 min (Table 2). Though each scalar has its own unique sources and sinks (Foken et al., 2006), these results were taken to suggest that low-frequency contributions to GEM fluxes at this site were also minimal. The averaging period was set at 60 min, as this gave some smoothing to both flux and environmental data, allowing for greater comparison between the two that was not achievable using 20 min averaging. No further averaging or smoothing of GEM flux data was applied. Following the application of quality control protocols, GEM flux values were obtained for 87 % of the study period; 39 % of flux values fell below the theoretical detection limit (Eq. 8, Edwards et al., 2005). These were not removed as doing so would have resulted in an artificial increase of the observed mean (see also Fritsche et al., 2008a; Converse et al., 2010).

#### 3.2.2 Overview

Bi-directional GEM fluxes were observed during the study and were normally distributed with a mean of 0.2 ± 14.5 ng m\(^{-2}\) h\(^{-1}\) (Fig. 2). The range of GEM fluxes was between −52.9 and 54.7 ng m\(^{-2}\) h\(^{-1}\), with generally greater magnitudes throughout the day than at night. Mean nocturnal fluxes were significantly different to diurnal fluxes, at −1.5 ± 7.8 ng m\(^{-2}\) h\(^{-1}\) compared to 1.8 ± 18.6 ng m\(^{-2}\) h\(^{-1}\). Over the entirety of the study, the net cumulative GEM flux was close to zero (Fig. 3a) as net deposition to the surface seen in the first 7 days was balanced by net emission throughout the remaining period. Low biological productivity in the underlying vegetation, suggested by manual observations, was confirmed by observed CO\(_2\) fluxes that were largely positive with net cumulative CO\(_2\) flux across the period of 1.7 mol m\(^{-2}\) (Fig. 3a). Both GEM and CO\(_2\) fluxes were bi-directional and showed high variabil-
The mean deposition velocity calculated using all data was 0.0 ± 0.06 cm s⁻¹. The mean deposition velocity was significantly lower than that reported for Seebodenalp, Switzerland (Obrist et al., 2010). The mean deposition velocity ranged from 2.2 to 2.9 cm s⁻¹. Mean nocturnal and diurnal deposition velocities were opposite in sign (direction), though not significantly different at 0.1 cm s⁻¹.

Table 2. Results from ogive analyses. Definitions of cases are given in Foken et al. (2006) and here describe comparisons between 20 and 150 min covariance averaging.

<table>
<thead>
<tr>
<th>Case</th>
<th>w/u'</th>
<th>w/T'</th>
<th>w/CO₂'</th>
<th>w/H₂O'</th>
</tr>
</thead>
<tbody>
<tr>
<td>Case 1</td>
<td>82.2 % (37)</td>
<td>97.8 % (44)</td>
<td>93.3 % (42)</td>
<td>75.5 % (34)</td>
</tr>
<tr>
<td>Case 2</td>
<td>17.8 % (8)</td>
<td>2.2 % (1)</td>
<td>6.7 % (3)</td>
<td>17.8 % (8)</td>
</tr>
<tr>
<td>Case 3</td>
<td>0 % (0)</td>
<td>0 % (0)</td>
<td>6.7 % (3)</td>
<td></td>
</tr>
</tbody>
</table>

3.2.3 Comparison with other flux studies

Patterns of net diurnal emission and nocturnal deposition were reported by Fritsche et al. (2008b) and Converse et al. (2010) for summertime periods over subalpine grasslands in Fruebuel, Switzerland, and Big Meadows, USA (Table 3). The range of flux values observed here is larger than those reported by Fritsche et al. (2008b) yet lower than those reported by Converse et al. (2010). Determining the significance of these differences however is difficult, as flux averaging periods differed across each of the studies, as well as application of temporal smoothing. The mean deposition velocity is within the range of uncertainty of those reported at Fruebuel and Neustift, Austria (Fritsche et al., 2008b). Greater variation in deposition velocities was seen here, albeit with a similar range to those seen at Big Meadows (Converse et al., 2010). The mean deposition velocity was significantly lower than that reported for Seebodenalp, Switzerland (Obrist et al., 2006).

Comparability of deposition velocities was maintained despite a mean ambient GEM concentration between 50 and
Table 3. Mean GEM, GEM flux and deposition velocity values (±1 standard deviation, range in parentheses) for various studies undertaken over background THg substrate in sub-alpine grasslands during summer periods. Values taken from a Obrist et al. (2006), b Fritsche et al. (2008b) and c Converse et al. (2010). Values obtained using d modified Bowen ratio method and 30 min averaging, e aerodynamic gradient and 30 min averaging, f aerodynamic gradient and 20 min averaging, g aerodynamic gradient and 60 min averaging.

<table>
<thead>
<tr>
<th>Site</th>
<th>Ambient GEM ng m⁻³</th>
<th>GEM flux ng m⁻² h⁻¹</th>
<th>Deposition velocity cm⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Seebodenalp, Switzerland</td>
<td>1.65 ± 0.01</td>
<td>−1.9 ± 0.2d</td>
<td>0.03 ± 0.003</td>
</tr>
<tr>
<td>Freibuehl, Switzerland</td>
<td>1.20 ± 0.20</td>
<td>−4.3 (−27 to 14)e</td>
<td>0.10 ± 0.16</td>
</tr>
<tr>
<td>Neustift, Austria</td>
<td>1.22 ± 0.20</td>
<td>−2.1 (−41 to 26)e</td>
<td>0.05 ± 0.16</td>
</tr>
<tr>
<td>Big Meadows, United States</td>
<td>1.28 (0.98 to 1.50)</td>
<td>2.5 (−124.8 to 82.4)d</td>
<td>0.05 (−3.1 to 1.9)</td>
</tr>
<tr>
<td>Nimmo, Australia</td>
<td>0.59 ± 0.10</td>
<td>0.2 ± 14.5e</td>
<td>0.002 ± 0.7</td>
</tr>
<tr>
<td></td>
<td>(&lt; 0.01 to 0.91)</td>
<td>(−52.9 to 54.7)</td>
<td>(−2.2 to 2.9)</td>
</tr>
</tbody>
</table>

64 % lower than those reported at the Northern Hemisphere sites. It is not reported in these papers whether the underlying vegetation had senesced, and so the relative control on fluxes by vegetation may differ across sites. Converse et al. (2010) concluded that there was no relationship between their observed GEM fluxes and stomatal conductance. Seasonal studies by these authors (Fritsche et al., 2008a; Converse et al., 2010) as well as modelling studies such as Hartman et al. (2009) have shown increased deposition of GEM to the surface during vegetation growing seasons, and we therefore expect that deposition velocities at Nimmo would similarly change throughout the year as biological activity increases. Further discussion of deposition velocities across different ecosystems is given in Zhang et al. (2009). The agreement between deposition velocities obtained here and elsewhere over similar conditions provides evidence towards the robustness of such an approach to estimating GEM deposition.

3.2.4 Environmental correlates

Factors that have been identified as controls on GEM surface exchange in the literature include soil moisture (Gustin et al., 2006), precipitation (Song and Van Heyst, 2005), radiation (Carpi and Lindberg, 1997; Choi and Holsen, 2009), temperature (Gustin et al., 1997; Choi and Holsen, 2009), vegetation (Hanson et al., 1995; Stamenkovic and Gustin, 2009) and GEM/oxidant concentrations (Engle et al., 2005; Zhang et al., 2008). Linear correlations between GEM fluxes and environmental data were generally weak (Table 1). The strongest significant relationship using all available data was with latent heat flux ($r = 0.29$), followed closely by sensible heat flux ($r = 0.27$), net solar radiation ($r = 0.24$) and air temperature ($r = 0.21$). A similar result was seen for diurnal data, whilst for nocturnal data the only significant correlation was with CO$_2$ flux ($r = 0.26$).

Soil volumetric water content (VWC) was low (<0.07; wilting point: 0.08) for most of the study and had no significant linear relationship with GEM fluxes. The precipitation event on DOY 45 was the only to impact VWC, raising it from 0.06 to 0.08. The two largest maximum diurnal cumulative GEM emissions (cumulative GEM flux from sunrise to peak value) of 128.2 and 112.1 ng m⁻² occurred on the days following this event (Fig. 3a), as VWC receded down to 0.07. This is consistent with the laboratory results of Gustin and Stamenkovic (2005), who hypothesised that evaporation of soil water helps mobilise mercury adsorbed to soil matter upwards to the air–surface interface. In their study, a pulse of GEM emission was observed immediately after precipitation events, attributed to expulsion of soil GEM from within the pore space. Such an immediate pulse was not observed during DOY 45, likely due to the small amount of precipitation in this event and its minimal impact on the already-dry soil.

Solar radiation and temperature have been independently shown to influence GEM fluxes from soils via photo-reduction and volatilisation of stored mercury (Lindberg et al., 2007; Zhang et al., 2009), though their inherent interconnection makes it difficult to resolve relative influence in the field. Kikuchi et al. (2013) showed that inclusion of surface air temperature in parameterisations of soil GEM emissions gave closer reproduction of seasonal observations than with radiation alone. Shading by overlying vegetation may also reduce the influence of solar radiation on soil GEM emission (Carpi and Lindberg, 1998). Linear correlation between GEM flux and net radiation was similar to that with air temperature, though GEM flux showed low correlation with soil temperature. The low linear correlation between GEM flux and soil temperature compared to scalar fluxes of heat may suggest that turbulent atmospheric processes (quasi-laminar and turbulent diffusion), rather than in-soil processes, represent the larger influence on GEM fluxes. Zhu et al. (2015) found stronger relationships between aerodynamic variables and GEM fluxes when measured by micrometeorological methods than when measured with flux chambers, suggesting the choice of measurement may influence these relationships. Edwards and Howard (2013) however similarly observed stronger relationships with atmospheric variables and GEM fluxes measured using dynamic flux chambers over bare background THg soil. The stronger relationship with air temperature than with soil temperature...
only held across sub-diurnal timescales; comparing cumulative GEM flux to 24 h averaged temperature data (Fig. 3a) resulted in a much stronger correlation with soil temperature ($r = 0.62$, $p < 0.0001$, $n = 443$) than with air temperature ($r = 0.36$, $p < 0.0001$, $n = 443$).

Median diurnal composite CO$_2$ fluxes were almost consistently positive both during the day and at night (Fig. 4b), excepting a notable period between 1 and 3 h after sunrise during which median net CO$_2$ release switched to uptake before reverting back to release. Overnight GEM fluxes were close to zero with little variance relative to daytime (Fig. 4a); however during the day two peaks of GEM emission can be seen, the largest coinciding with solar noon, whilst the smaller peak occurred shortly after sunrise and coincides with an increase in bulk canopy conductance of water vapour. The cessation of this early pulse coincides with the switch to CO$_2$ uptake that may be related to uptake of GEM by the underlying vegetation. However, the correlation between the timing of the early GEM pulse and the brief increase in bulk canopy conductance suggests that this pulse is likely related to evaporation of dew. The bimodal pattern of GEM fluxes has been observed previously by others (e.g. Lindberg et al., 1998; Fritsche et al., 2008b; Bash and Miller, 2008; Zhu et al., 2015) and attributed to co-deposition of mercury with dew overnight, with subsequent re-volatilisation of GEM as the dew is evaporated from the surface. The linear correlation of GEM fluxes with latent heat fluxes in this study similarly provides evidence for this explanation of the bimodal GEM flux pattern.

3.3 Nocturnal atmospheric mercury depletion events

Both GEM and O$_3$ concentrations, as well as air temperature and wind speed, were intermittently observed to show significant decreases (Fig. 1b), twice to below instrument detection limits for GEM. These nocturnal atmospheric mercury depletion events (NAMDEs) occurred exclusively overnight, with concentrations returning to pre-depletion levels after sunrise. Depletion was most pronounced during the calm, stable dew nights. Patterns of depletion differ for GEM and O$_3$, with O$_3$ exhibiting initially rapid depletion followed by slower removal, mirroring decreases in atmospheric turbulence (linear correlation with $\sigma_w/\mu_s$ for dew nights; $r = 0.86$, $p < 0.0001$, $n = 86$). GEM concentrations show a more linear rate of depletion through to sunrise, when both GEM and O$_3$ levels increase with the turbulent breakup of the nocturnal boundary layer. These differences in patterns are particularly pro-

Figure 3. (a) Cumulative GEM and CO$_2$ fluxes across the study period, along with precipitation and 24 h running means of soil temperature at 2 cm depth. Colours match those of axis labels. Observations of dew and fog are denoted above by “D” and “F”, respectively. (b) 60 min GEM and CO$_2$ fluxes over the study period.

Figure 4. (a) Diel composite of GEM flux values. Black line represents median values, blue shading inter-quartile range (IQR), and blue lines 5th and 95th percentiles. Grey shading represents nocturnal periods. (b) As above for CO$_2$ fluxes (solid black line, green shading) and bulk canopy conductance for water vapour (broken black line, magenta shading).
nounced on DOY 45, 46 and 48 and are similar in character to the depletion events reported by Mao et al. (2008). Similar nocturnal depletion events have been reported elsewhere (e.g. Engle et al., 2010; Peleg et al., 2015; Fu et al., 2016; Howard et al., 2017) and differ from the better-known polar AMDEs (Steffen et al., 2008, and reference within), as the former take place in the absence of sunlight and photolytic reactions. HYSPLIT trajectories showed no distinct source pattern for NAMDE nights, suggesting that the observed phenomena are due to local interactions and not the result of long-range transport of depleted air masses such as those observed by Gauchard et al. (2005), Mastromonaco et al. (2016) and Moore et al. (2014).

Nocturnal composites of cumulative GEM flux, ambient GEM and modelled ambient GEM/dew were composed to further explore NAMDEs, demarcating between dew and non-dew nights. Figure 5 confirms that depletion of GEM occurred exclusively during dew nights, highlighting dew or fog as a potential sink of GEM. Field studies of mercury concentrations in fog water (e.g. Ritchie et al., 2006; Weiss-Penzias et al., 2012) are rare, though concentrations up to 435 ng L$^{-1}$ have been reported (Ritchie et al., 2006). When estimating conservatively using this concentration and a fog droplet density of 0.05 g m$^{-3}$, only 0.02 ng m$^{-2}$ of GEM depletion can be explained by fog uptake. Field measurements of mercury in dew are also limited, though they have generally shown that it represents only a small sink for atmospheric mercury. Malcolm and Keeler (2002), from the first such measurements, calculated values of mercury deposition that ranged between 1.2 and 9.6 ng m$^{-2}$ per dew event. THg concentrations similar to those from Malcolm and Keeler have since been reported by Engle et al. (2010) and Converse et al. (2014). Engle et al. (2010) reported GEM depletion events coincident with their dew mercury measurements and concluded, based on a mass balance approach, that mercury uptake to dew could explain <1 % of observed depletion. From the combined dew measurements of Malcolm and Keeler (2002), Engle et al. (2010) and Converse et al. (2014) a reasonable estimate of dew mercury concentration is $6.3 \pm 4.9$ ng L$^{-1}$ ($n = 27$). Applying this concentration to the median modelled dew depth (0.13 mm) gives a cumulative nocturnal deposition to dew of 0.82 ng m$^{-2}$.

Peleg et al. (2015) observed nocturnal depletion of GEM, concurrent with measurements of GOM and nitrate (NO$_3$). They showed evidence of NO$_3$ assisting in the oxidation of GEM to GOM overnight, noting a strong correlation between NO$_3$ and GOM, and weak correlations between GOM and other measured variables such as O$_3$ and wind speed. Mao et al. (2008), in chemical modelling of their observed depletion events, attributed 80 % of chemical oxidation of GEM to NO$_3$, whilst also considering oxidation by O$_3$ and OH.

The presence of cattle prior to the study may have provided an additional source of NO$_3$ at the surface (Schlegel et al., 2017), though without measurements of nitrogen species this remains speculative. Further, this source would be available during both dew and non-dew nights and cannot alone explain differences in the composite ambient GEM patterns. Higher relative humidity during dew nights relative to non-dew nights (Fig. 1) is likely to have a limiting influence on oxidation of GEM by NO$_3$ due to enhanced depletion of the radical (Geyer et al., 2001; Vrekoussis et al., 2004); Peleg et al. (2015) noted a significant anti-correlation between relative humidity and NO$_3$, as well as between relative humidity and GOM. They also highlighted possible evidence of GEM oxidation by O$_3$, albeit only at levels above 47 ppb, higher than the nocturnal concentrations observed here. Mao et al. (2008) note that O$_3$ oxidation represented the largest uncertainty in their modelling estimates, with differences in oxidation of 86 % attributable to varying rate constants from the literature.

In the absence of adequate measurements of potentially important chemical species, a simple box model was utilised in order to explore the relative influence of observed surface deposition (Eq. 5). The outputs from this model show that patterns of depletion and non-depletion can largely be explained by surface deposition, though these outputs are strongly dependent on the modelled mixed-layer height, which represents the largest uncertainty in the model. The median height during dew nights (50 m) was 58 % lower than for non-dew (120 m), consistent with surface stability measurements. Mao et al. (2008), in their modelling efforts, estimated that dry deposition to the surface could account for 70 % of their observed depletion events, whilst Fu et al.
(2016) investigated GEM depletion for a range of surface deposition and turbulent diffusivity values under a boundary layer of height 100 m. They showed that a deposition flux of 7.3 ng m\(^{-2}\) h\(^{-1}\) and turbulent diffusivity of 0.1 to 1 cm s\(^{-1}\) could reduce GEM concentrations within a stable boundary layer from 1.56 to 0 ng m\(^{-2}\). Turbulent diffusivity values during dew nights were similar to those seen by Fu et al. (2016) (median 2 cm s\(^{-1}\)), whilst deposition fluxes were generally smaller (median 3.1 ng m\(^{-2}\) h\(^{-1}\)). Free-tropospheric GEM concentrations however were also considerably smaller, as were modelled boundary layer heights.

GEM fluxes were significantly different between dew and non-dew nights, with the former showing consistent net deposition, whilst the latter showed fluxes not different from zero. The median cumulative nocturnal GEM flux for dew nights was 22.6 ng m\(^{-2}\) lower than for non-dew. Care must be taken here, as under low-turbulence conditions atmospheric gradients may be enhanced, leading to an overestimation of surface fluxes (Obrist et al., 2006). This however is the case for both dew and non-dew nights, where the integral turbulence characteristic is consistently < 1. This result points to an additional GEM sink at the surface that is present during dew nights only. The most likely candidate is chemical oxidation of GEM to the more reactive GOM, much as in the case of AMDEs, as both the solubility and deposition velocity of this form of mercury are higher than for GEM (Lindberg et al., 2007; Zhang et al., 2009). We hypothesise, based on the greater observed GEM deposition and results of the box model, that any oxidation is taking place largely at the surface, leading to an enhanced GEM gradient. Complex surface chemistry may be taking place in the presence of high humidity and liquid water, such as the enhanced oxidation of GEM by O\(_3\) observed by Snider et al. (2008). Other oxidation pathways are also possible, and we therefore recommend consideration of the chemical processes taking place at the surface in future investigations.

Once in oxidised form, mercury can be more readily taken into dew (Munthe, 1992; Pleuel and Munthe, 1995; Malcolm and Keeler, 2002), soil and vegetation (Andersson, 1979; Zhang et al., 2009) or vegetation via foliar uptake (Lindberg and Stratton, 1998; Stamenkovic and Gustin, 2009), where it can become associated with cuticular membranes (Mason et al., 1995). In their isotopic investigations, Hintelmann et al. (2002) observed that 66% of wet-deposited mercury was bound within vegetation, suggesting this may be a significant sink of GOM formed near the surface. Figure 5 shows that, following the significant deposition during dew nights, an emission pulse is observed shortly after sunrise. This initial pulse, with median value of 3.9 ng m\(^{-2}\), represents 17% of total nocturnal deposition. This pulse is not observed following non-dew nights, showing that the bimodal shape of GEM fluxes discussed in Sect. 3.2.4 is not the regular diel pattern. Instead, this suggests that co-deposition of mercury with dew overnight does play a role in the initial emission pulse as suggested by others; however the estimated 0.82 ng m\(^{-2}\) co-deposition represents only 4% of total mercury deposition. As such, we further hypothesise that the majority of this morning GEM emission pulse is due to prompt recycling of mercury, likely volatilised from GOM created near the surface and deposited overnight. This percentage of promptly recycled mercury is within the range observed by Hintelmann et al. (2002) and Amyot et al. (2004), with the remaining deposited mercury likely stored in vegetation and soils. The long-term impact of any additional sink is however likely to be minimal, as evidenced by the near-zero cumulative GEM flux over the study period.

4 Conclusions

GEM flux measurements were undertaken over a mid-latitude alpine grassland region of Australia during a 3-week period in the late austral summer. The micrometeorological aerodynamic gradient method was employed, providing high-time-resolution fluxes and facilitating evaluation of controlling factors. Both deposition and emission fluxes were observed, with a cumulative flux close to zero over the study period. Nocturnal GEM fluxes were \(-1.5 \pm 7.8\) ng m\(^{-2}\) h\(^{-1}\) compared to diurnal fluxes of \(1.8 \pm 18.6\) ng m\(^{-2}\) h\(^{-1}\); this equated to deposition velocities of 0.1 ± 0.4 cm s\(^{-1}\) and \(-0.1 \pm 0.9\) cm s\(^{-1}\), respectively. These deposition velocities are mostly within the range of uncertainty of others reported for alpine grasslands during summer periods in Europe and North America. This is an important result towards justification of applying known deposition velocities over similar surfaces in global atmospheric mercury models, as background atmospheric mercury pool concentrations in the Southern Hemisphere are now understood to be lower than previously believed. Worldwide, air–surface GEM exchange studies across multiple seasons are rare yet required in order to reduce uncertainty in chemical transport model parameters and resolve the importance of terrestrial systems as sources or sinks of atmospheric elemental mercury.

Direct linear correlations between GEM fluxes and other environmental parameters were generally weak, with measures of temperature, radiation and heat fluxes (both sensible and latent) showing the strongest relationships. Soil temperature did not prove to be strongly related to raw GEM flux values; however it showed a moderately strong relationship when averaged over a running 24 h period and compared to cumulative GEM flux. Soil moisture was below the wilting point for the majority of the study and had little observable impact on GEM fluxes; however during the drying period following one precipitation event diurnal GEM emission fluxes were enhanced. The vegetation at the site had largely senesced and showed little overall stomatal control on GEM fluxes. A bimodal pattern of diurnal GEM emission was observed, the larger peak of which coincided with solar noon, whilst the smaller coincided with a brief increase in bulk canopy conductance of water vapour shortly after sunrise.
Correlation between GEM and latent heat fluxes suggests that the latter pulse is attributable to release of co-deposited mercury as GEM during evaporation of dew.

Nocturnal atmospheric mercury depletion events are also reported here, concomitant with depletion of O3 and formation of dew, under calm, stable boundary layers. Modelling of dew depth confirmed manual observations that showed dew formation during NAMDEs only and that uptake of mercury to dew represents only 4% of nocturnal mercury deposition. Other researchers have also reported NAMDEs with no definitive explanation; however there is evidence to suggest that surface deposition plays a large role. Here we investigated the role of surface deposition using observed flux data in a simple mass balance box model. This model reproduced ambient GEM patterns for nights both with and without observed depletion, though with high uncertainty in the modelled boundary layer heights. GEM deposition fluxes were also enhanced during NAMDEs, leading us to hypothesise an enhancement of the near-surface GEM gradient due to oxidation of GEM to GOM and subsequent deposition. Early-morning GEM emission pulses representing 17% of nocturnal mercury deposition were observed following NAMDEs only. We further hypothesise that this pulse is due to prompt recycling of mercury deposited during these nights, with the remaining deposited mercury retained in vegetation and soil. As cumulative GEM fluxes over the 3-week period were close to zero, the influence of any additional sink would extend over reasonably short time periods. The growing literature on mercury emission and deposition behaviour is building greater evidence towards prompt recycling behaviour of GEM in the global mercury cycle, with significant impacts on our understanding of the legacy of anthropogenic perturbations to this cycle.

Data availability. Data are available upon request from the corresponding author.

Competing interests. The authors declare that they have no conflict of interest.

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References


D. Howard and G. C. Edwards: Mercury fluxes over an Australian alpine grassland


Kikuchi, T., Ikemoto, H., Takahashi, K., Hasome, H., and Ueda, H.: Parameterizing Soil Emission and Atmospheric Oxidation-Reduction in a Model of the Global Biogeochemical Cy-