A model study of global mercury deposition from biomass burning

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Abstract

Mercury emissions from biomass burning are not well characterized and can differ significantly from year to year. This study utilizes three recent biomass burning inventories (FINNv1.0, GFEDv3.1 and GFASv1.0) and the global Hg chemistry model, ECHMERIT, to investigate the annual variation of Hg emissions, and the geographical distribution and magnitude of the resulting Hg deposition fluxes. The roles of the Hg/CO enhancement ratio, the emission plume injection height, the Hg0 oxidation mechanism and lifetime, and the inventory chosen, and the uncertainties with each

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were considered. The greatest uncertainties in the total Hg deposition were found to be associated with the Hg/CO enhancement ratio and the emission inventory employed. Deposition flux distributions proved to be more sensitive to the emission inventory and the oxidation mechanism chosen, than all the other model parameterizations. Over 75% of Hg emitted from biomass burning is deposited to the world’s oceans, with the highest fluxes predicted in the North Atlantic and the highest total deposition in the North Pacific. The net effect of biomass burning is to liberate Hg from lower latitudes and disperse it towards higher latitudes where it is eventually deposited.

1 Introduction

Mercury pollution is a global threat to human and ecosystem health, since its elemental form (Gaseous Elemental Mercury (GEM) or Hg$_{(g)}^0$), which makes up the major part of atmospheric emissions and re-emissions, can be transported far away from its emission source, (natural or anthropogenic, AMAP/UNEP, Driscoll et al., Pirrone and Mason). Due to increased interest in trying to constrain the global budget of Hg as it cycles between environmental compartments, increased attention has also been given to Biomass Burning (BB) emissions. Friedli et al. estimated Hg emissions from BB by combining outputs from global carbon emission models with Hg enhancement ratios and found that globally 675 (±240) Mg yr$^{-1}$, averaged over the period 1997-2006, is emitted from BB. As this figure is approximately one third of the yearly anthropogenic emissions of Hg to the atmosphere, it is clear that BB plays an important role in the Hg biogeochemical cycle. As controls on anthropogenic Hg emissions become stricter, proportionally the role of BB will increase, possibly substantially if the instances and extent of wildfires increases in a changing climate. It should also be noted that the location of Hg emissions from BB is very different from the location of anthropogenic emissions, with the exception of artisanal and small scale gold mining. Mercury from BB is almost all emitted as Hg$_{(g)}^0$, with a small fraction associated with the soot from the fires. Elemental mercury has an estimated lifetime of between 8
months and 1 year\textsuperscript{2,8,9} and therefore can be deposited to ecosystems very distant from fire locations.

Atmospheric Hg\textsubscript{0} can be oxidized to Hg\textsubscript{II}, which is subsequently removed by both wet and dry deposition. A part of the Hg\textsubscript{II} that is deposited may be methylated within ecosystems and it is this form of Hg which can enter the food web and is toxic to living organisms. The recent Minamata Convention (http://www.mercuryconvention.org/) is aimed at reducing the anthropogenic impact on the global Hg biogeochemical cycle\textsuperscript{10}. However, the natural Hg cycle is already significantly perturbed; it is estimated that there is five times the Hg in the present day atmosphere than was present in pre-industrial times\textsuperscript{2,11}. The legacy of past emissions will most likely continue to influence the global biogeochemical cycle of Hg for decades to come\textsuperscript{12}, and fires will play an important role in the continued cycling of Hg between environmental compartments. The primary objective of the study is to simulate the magnitude and geographical location of the Hg deposition flux that result from BB.

Three recent BB inventories, FINNv1.0\textsuperscript{13}, GFEDv3.1\textsuperscript{14} and GFASv1.0\textsuperscript{15}, referred to simply as FINN, GFED and GFAS hereafter, have been used to simulate Hg emissions from fires over the period 2006–2010. Hg emissions have been calculated as a function of CO emissions and the deposition flux distribution of Hg from BB has been simulated, using the global on-line chemical transport Hg model ECHMERIT\textsuperscript{16,17}.

2 Methodology

2.1 The Global Biomass Burning Inventories

The details of the three inventories used, FINN, GFAS, and GFED and how they were compiled can be found in the literature\textsuperscript{13–15}. All three inventories are based on the imagery obtained from the MODIS instruments on-board the NASA Terra and Aqua satellites; however they differ in the way in which the data are filtered or processed. GFED makes use of the burned area retrieval, FINN uses an active fire data product, while GFAS uses fire
radiative power retrievals (the algorithm for which is based on active fire detection). Further differences in the inventories concern the land cover maps used, and the details concerning fuel load and fuel consumption. A detailed comparison and description of the inventories can be found in Andela et al.\textsuperscript{18}.

Over the period 2003–2011 three inventories agree fairly well on the annual average CO emissions\textsuperscript{18}. The inventories identify the same regional BB hot spots caused by tropical deforestation in South America, fires in African savannas, forest fires in South-East Asia and seasonal wildfires in Northern Hemisphere boreal regions. However, the regional differences in CO emissions between FINN, GFAS and GFED are substantial. GFAS has the highest values for areas with low burning intensity such as dry savannas. Conversely for high burning intensity fires, GFED has higher emissions. The different approaches in compiling the inventories is apparent from the relatively high emission estimate of GFAS and FINN in some areas of the world (Africa, South-East Asia and northern Brazil), whereas GFED is tuned particularly to capture large scale deforestation in central Brazil. GFED thus has higher emissions in the Southern Hemisphere than FINN and GFAS. For boreal forests GFAS and GFED emission estimates are considerably higher than FINN, see Andela et al.\textsuperscript{18}.

The GFAS and GFED inventories were obtained from the Emissions of Atmospheric Compounds and Compilation of Ancillary Data (ECCAD) Global Emissions InitiAtive (GEIA) portal\textsuperscript{19}, while the FINN inventory was obtained from the Atmospheric Chemistry Division of National Center for Atmospheric Research (NCAR)\textsuperscript{20}. For the purpose of model validation the most recent anthropogenic Hg emission inventory from AMAP/UNEP (reference year 2010) was used\textsuperscript{1}, as described in the SI.

### 2.2 Model set-up

The global Hg chemical transport model ECHMERIT\textsuperscript{16,17} is based on the fifth generation General Circulation Model ECHAM5\textsuperscript{21,22}. ECHMERIT was run using T42 horizontal resolution (roughly 2.8° by 2.8° at the equator) and 19 vertical levels up to 10 hPa. The increase
in atmospheric Hg concentration resulting from BB were estimated as in Friedli et al.\textsuperscript{6}, using an Enhancement Ratio (ER), defined as,
\[ \text{ER} = \frac{\Delta [\text{Hg}]}{\Delta [\text{CO}]} \]
where \( \Delta [\text{Hg}] \) is the sum of all Hg species in excess of background, and \( \Delta [\text{CO}] \) is the difference between the plume and background CO concentration\textsuperscript{6}. The global average ER (ER\textsubscript{av}) reported by Friedli et al.\textsuperscript{6} is \( 1.54 \times 10^{-7} \). This value was used in most of the simulations, however a number of simulations were run in which the ER\textsubscript{av} was substituted by biome specific ERs as described in the SI. The GFAS and GFED emissions were mapped on the ECHMERIT T42 grid using the mass conserving remapping tool included in the Climate Data Operators (CDO)\textsuperscript{23}. The NCAR ACD Fortran pre-processor program, \texttt{Fire.Emis}, was used to interpolate the FINN inventory on to the ECHMERIT grid\textsuperscript{24}. The monthly average emissions were calculated for the FINN and GFAS inventories to be compatible with the GFED inventory.

With the exception of the simulations performed for model validation purposes, all simulations were performed using Hg emissions from BB only.

\section*{2.3 The simulations performed}

Base case simulations used the \( \text{O}_3/\text{OH} \) oxidation mechanism, however there is some uncertainty over the atmospheric Hg oxidation pathway\textsuperscript{25–27}, therefore simulations were performed using a Br based oxidation mechanism to assess how the oxidation mechanism influences the deposition flux fields. Further simulations were performed introducing the BB emissions into different model levels, and combinations of levels. Five year simulations (2006–2010) were performed to investigate long-term differences between the inventories, while single year simulations were performed to investigate how deposition patterns varied from year to year.

In the case of the single year simulations, since these were aimed at assessing the direct deposition of Hg, the mechanism by which a fraction of deposited Hg is rapidly re-emitted from terrestrial, snow/ice and water surfaces\textsuperscript{28} was switched off, in all other simulations
re-emission was included. Single year simulations were continued beyond 12 months without further emissions until at least 95% of the emitted Hg had been deposited. This took a further 9 to 12 months. Finally, simulations to investigate the differences in emission and deposition fields when using biome/land-cover based ERs were performed. A summary of the simulations performed can be found in Table S1.

3 Results

Although the primary aim of this study is to identify the areas most impacted by Hg emissions from BB, and to see how greatly these differ from one BB inventory to another, the first simulations were performed using Hg emissions from all sources. The runs were performed for the year 2010, using each BB inventory for the Base mechanism, and GFED for the Br oxidation mechanism. GFED was also used for simulations using a fixed lifetime against oxidation ("pseudo" oxidation mechanism). The results from these simulations were compared to available measurement data, and a statistical summary of the comparison for gas phase Hg and for Hg wet deposition can be found in table 1. Maps of the comparisons are included in the SI in figures S1–S3.

Table 1: Comparison of the Base, Br and fixed lifetime simulations with global observations for 2010

<table>
<thead>
<tr>
<th></th>
<th>FINN</th>
<th>GFAS</th>
<th>GFED</th>
<th>GFED Br-Oxdn</th>
<th>GFED 12-m fixed</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Base</td>
<td>Base</td>
<td>Base</td>
<td>Base Br-Oxdn</td>
<td>12-m fixed</td>
</tr>
<tr>
<td>TGM</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intercept</td>
<td>0.34</td>
<td>0.31</td>
<td>0.34</td>
<td>0.18</td>
<td>0.34</td>
</tr>
<tr>
<td>Slope</td>
<td>0.68</td>
<td>0.69</td>
<td>0.67</td>
<td>0.72</td>
<td>0.70</td>
</tr>
<tr>
<td>Pearson’s r</td>
<td>0.77</td>
<td>0.78</td>
<td>0.77</td>
<td>0.75</td>
<td>0.76</td>
</tr>
<tr>
<td>NRMSE(%)</td>
<td>14.6</td>
<td>14.4</td>
<td>14.3</td>
<td>16.7</td>
<td>15.5</td>
</tr>
<tr>
<td>Wet Dep</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intercept</td>
<td>9.26</td>
<td>9.05</td>
<td>9.11</td>
<td>10.4</td>
<td>7.15</td>
</tr>
<tr>
<td>Slope</td>
<td>0.32</td>
<td>0.31</td>
<td>0.31</td>
<td>0.22</td>
<td>0.19</td>
</tr>
<tr>
<td>Pearson’s r</td>
<td>0.21</td>
<td>0.21</td>
<td>0.21</td>
<td>0.14</td>
<td>0.17</td>
</tr>
<tr>
<td>NRMSE(%)</td>
<td>19.1</td>
<td>18.6</td>
<td>18.8</td>
<td>19.6</td>
<td>13.9</td>
</tr>
</tbody>
</table>

The comparison between the different model versions and observations all yield similar
results, which are reasonable for Total Gaseous Mercury (TGM, the sum of gas phase ele-
mental and oxidized Hg species), and less good for Hg wet deposition. Interestingly simply
assuming a fixed atmospheric lifetime for Hg does not give results that are significantly worse
than when a more detailed chemical mechanism is employed. However it should be pointed
out that for the year 2010 almost all the observations are from the northern hemisphere, and
this may not be the case when southern hemisphere sites are taken into account. (Currently
the Global Mercury Observation System project is performing Hg monitoring at a number
of sites in the southern hemisphere (www.gmos.eu)).

3.1 Geographical distribution and seasonality of emissions

The temporal and spatial distribution of the Hg emissions is dictated by the distribution of
CO emissions because of the way they have been calculated. The differences between the
inventories, in terms of CO emissions, are described elsewhere\textsuperscript{13–15,18,29}.

Although the annual average Hg emitted between 2006–2010 is similar: 678, 603 and 600 Mg
for FINN, GFAS and GFED respectively, there are significant interannual differences and
noticeable variations in the latitudinal distribution (see Figure 1).

The highest year to year variability is seen in the GFED inventory. While the FINN and
GFED inventories have similar temporal profiles and are reasonably correlated (r = 0.9).
The GFAS inventory shows a markedly different temporal profile, Figure 1(a) (r = 0.2 and
0.5, with FINN and GFED respectively). The decreasing trend in emissions over time seen
in the GFAS inventory is also at odds with the other two inventories.

The latitudinal profiles of the emissions, for the period 2006-2010, while similar, do have
noticeable differences (Figure 1(c)). The GFED inventory has significantly higher emissions
at around 10°S (6.4 g km\(^{-2}\) y\(^{-1}\)), whereas the FINN inventory shows a much higher peak at
around 20°N (4.2 g km\(^{-2}\) y\(^{-1}\)). The FINN inventory also lacks the peaks at 7°N and at 65°N
which are evident in the GFAS and GFED inventories. In terms of the latitudinal profile
Figure 1: Annual trends and averaged latitudinal profiles of mercury emissions ((a) and (c)) and deposition ((b) and (d)). Figure (b) excludes 2006 due to low re-emissions, see section 3.2.1.
the GFAS and GFED inventories show the highest correlation \((r = 0.9)\). The global Hg emission spatial and seasonal distributions are shown in Figure 2, as is the distribution of the emissions between source regions.

Figure 2: Geographical (left), seasonal (center, DJF - December January February, MAM - March April May etc.) and regional (right) distribution of mercury emissions. Annual averages over the 2006–2010 period. The regions are, following the nomenclature used in van der Werf et al.\(^{14}\), (Boreal North America (BONA), Temperate North America (TENA), Central America (CEAM), Northern Hemisphere South America (NHSA), Southern Hemisphere South America (SHSA), Europe (EURO), Middle East (MIDE), Northern Hemisphere Africa (NHAF), Southern Hemisphere Africa (SHAF), Boreal Asia (BOAS), Central Asia (CEAS), Southeast Asia (SEAS), Equatorial Asia (EQAS) and Australia (AUST)
3.2 Hg deposition

3.2.1 Five year simulations

Figure 3 shows the geographical distribution of the annual total deposition (wet plus dry) due to BB averaged over the last four years of the 5 year simulation period, (to avoid the first year where re-emission is lower). Not surprisingly, high emissions combined with high precipitation downwind of emission source regions gives rise to high deposition fluxes. Figure 3 also shows that while BB emissions are terrestrial, most of the Hg deposition occurs over the oceans. The yearly Hg deposition totals using each inventory follow the emission totals (but also include deposition of re-remitted Hg), (see figures 1(a) and 1(b)). The emissions latitudinal profile have well defined peaks and a distinct cut-off at the southern and northern limits of vegetation (Fig. 1(c)). The deposition profile, due to the lifetime of Hg in the atmosphere, shows far less pronounced peaks, a broader profile, and never reaches zero, at any latitude, Fig. 1(d). Looking at the southern hemisphere, almost all emissions are between the equator and 30°S, even at 50°S the Hg deposition is still 40% of that seen in the high Hg deposition regions. This latitudinal distribution of the Hg is almost independent of the BB emissions inventory used, indicating that atmospheric transport determines to a great extent the Hg deposition flux distribution.

Another way to illustrate the importance of atmospheric transport on the simulated deposition fields is to compare the spatial correlation (R) of the emission and the deposition fields, Table 2. The values reported were calculated using the horizontal pattern correlation method\textsuperscript{30,31}. The highest correlation for the emission inventories is found between GFAS and GFED (R = 0.68), the lowest between FINN and GFED (R = 0.38). The value of R varies from year to year (Table 2), reflecting differences in the approaches used to compile the inventories, which are discussed by Andela et al.\textsuperscript{18}. Higher spatial correlations (R very close to 1) are found for the simulated Hg deposition fields, due to the effect of the Hg\textsubscript{0}(g) atmospheric lifetime, and hence transport, which smooths the variations seen in the emissions.
Figure 3: Geographical distribution of the total mercury deposition (wet + dry) that result from BB. Annual averages over the 2007–2010 period.
Table 2: Spatial correlations (R) between the emissions inventories and the simulated deposition fields

<table>
<thead>
<tr>
<th>Year</th>
<th>FINN-GFAS</th>
<th>FINN-GFED</th>
<th>GFAS-GFED</th>
<th>FINN-GFAS Depos</th>
<th>FINN-GFED Depos</th>
<th>GFAS-GFED Depos</th>
</tr>
</thead>
<tbody>
<tr>
<td>2006</td>
<td>0.47</td>
<td>0.33</td>
<td>0.82</td>
<td>1.00</td>
<td>0.97</td>
<td>0.98</td>
</tr>
<tr>
<td>2007</td>
<td>0.42</td>
<td>0.35</td>
<td>0.66</td>
<td>1.00</td>
<td>0.98</td>
<td>0.99</td>
</tr>
<tr>
<td>2008</td>
<td>0.30</td>
<td>0.31</td>
<td>0.56</td>
<td>0.99</td>
<td>0.99</td>
<td>0.99</td>
</tr>
<tr>
<td>2009</td>
<td>0.29</td>
<td>0.24</td>
<td>0.47</td>
<td>0.99</td>
<td>0.99</td>
<td>0.99</td>
</tr>
<tr>
<td>2010</td>
<td>0.31</td>
<td>0.37</td>
<td>0.46</td>
<td>0.99</td>
<td>0.99</td>
<td>0.99</td>
</tr>
<tr>
<td>2006-10</td>
<td>0.42</td>
<td>0.38</td>
<td>0.68</td>
<td>0.99</td>
<td>0.98</td>
<td>0.99</td>
</tr>
</tbody>
</table>

The net effect of BB in tropical regions is essentially to cycle Hg from the tropics to mid-latitudes and to a lesser extent to high latitudes (see Figure S4). Northern boreal BB directly impacts mid- and high latitudes.

### 3.3 Overall and yearly deposition comparison

To compare the deposition fields simulated using the three inventories, maps of agreement which highlight similarities and differences in geographically resolved datasets can be used. Model cells in which the Hg deposition was greater than the average plus one standard deviation ($\mu + \sigma$) were identified for each BB inventory simulation. These cells were mapped to see how consistent the extremes of the deposition distribution is between the simulations. The detailed pseudo-language algorithm used to generate such maps is presented in the SI. Figure 4 shows all of the areas where the deposition is greater than $\mu + \sigma$, for the 5 year (Base) simulations. The color of the cells denotes the level of agreement between the inventories. The high Hg deposition regions on which all the inventories agree represents roughly 15% of the Earth’s surface, and as the map makes quite clear, most of these regions are over the tropical and northern oceans. The map (Figure 4) shows the agreement between the FINN and GFAS simulations (gray plus orange cells), and there are relatively few cells where these inventories are the only ones to predict high Hg deposition (red and yellow cells). In contrast the simulations performed using the GFED inventory show a difference in the
Figure 4: Agreement map of Hg deposition fields obtained from GFAS, GFED and FINN for the five year simulation. The map shows the areas where deposition is $>\mu+\sigma$. Primary colors (red, blue and yellow) represent non-agreement between inventories, green, purple and brown indicate agreement between two of the inventories and gray indicates agreement between all three. The numbers refer to the number of cells in common between the simulations using the different inventories (The whole globe is represented by 8192 cells)
prediction of regions of high Hg deposition, and particularly in the southern hemisphere, and
to the southern edge of the region where all three inventories agree.

All of the inventories have an emission peak at roughly 10°S but while that of FINN and
GFAS is \( \approx 4.5-5 \text{ g km}^{-2} \text{ y}^{-1} \), that in GFED is \( \approx 7 \text{ g km}^{-2} \text{ y}^{-1} \). This accounts for the large
number of cells in the southern hemisphere where the simulation performed with GFED
predicts high Hg deposition values. Interestingly, at around 20°N there is a peak in the
FINN inventory that is more than twice as high as the corresponding values in GFED and
GFAS. Given the relatively few areas where only the simulations with the FINN inventory
predict high Hg deposition, this peak in emissions seems to affect the results relatively little,
suggesting that at certain latitudes, differences (in magnitude and precise location) in the
inventories have a negligible influence on the simulation results. (See Figure 1(c)). However
in the case with the simulation performed using GFED, the magnitude and location of the
emissions are much more important. Anthropogenic emissions in the southern hemisphere
are low compared to the northern hemisphere, therefore the contribution to atmospheric
Hg from BB is relatively more important in this region. From these results it appears
that precision in the magnitude and location of BB emissions in the southern hemisphere is
particularly necessary.

ECHMERIT has also been run for individual years (2006–2010) using each of the emission
inventories. As above for the 5 year simulation, agreement maps for deposition greater than
\( \mu + \sigma \) for each inventory have been prepared to compare the results from each inventory
for each year. The simulations predict the same total global area of high deposition (\( \approx 14-16\% \)) each year, and also that these areas are consistent from year to year. While the FINN
and GFAS inventories give similar results, the GFED inventory consistently predicts higher
deposition in the Southern hemisphere. This is true for each of the single year simulations
as it was for the 5 year simulation. Since the major BB source in south of the Equator is
the Amazon, this may well reflects the fact that GFED is “tuned” to capture large scale
deforestation in this region\(^{18}\) (Figure S5).
3.4 Sensitivity Studies

The emission inventories used for these studies have their own intrinsic uncertainties\textsuperscript{13–15,32}. Biome specific emission factors, assumptions concerning the oxidation mechanism and the atmospheric lifetime of Hg, and also the height (model layer) at which the BB emissions are introduced into the atmosphere are all potential sources of uncertainty. Some of these uncertainties are common to all BB studies, such as plume modeling, injection height, diurnal variation of fire intensity, fire areas and enhancement ratios\textsuperscript{6,33}. To investigate the impact of the parameterizations on the Hg deposition fields a number of sensitivity runs were performed (see Table S1).

3.4.1 Enhancement ratio

The most critical of all the assumptions made concerning Hg emissions from BB is the Enhancement Ratio (ER). Using $\text{ER}_{av}$ makes the modeling studies themselves more simple, and also avoids making a series of interconnected assumptions concerning ERs, vegetation types and their distribution, each of which could potentially introduce further errors into the model. The two major uncertainties when attempting to use a biome dependent approach to Hg emissions from BB are knowledge of the distribution of vegetation types, and the ER associated with a given type of vegetation, which may vary with location. There have been relatively few determinations of Hg concentrations in BB plumes for specific vegetation types\textsuperscript{6}. The ERs reported vary significantly for most vegetation types and can differ by more than an order of magnitude for a given biome. This is most likely due to a combination of factors including soil Hg content, fire intensity and fire location. Two biome specific ER simulations were performed using the GFED inventory. The first, $\text{ER}_{\text{coarse}}$, was calculated using the vegetation type characterization published in Friedli et al.\textsuperscript{6}, whereas the second set, $\text{ER}_{\text{fine}}$, was obtained following a more detailed characterization methodology as described in the SI. Using $\text{ER}_{av}$ (leads to Hg emissions from BB of 599.4±104.6 Mg, whereas $\text{ER}_{\text{coarse}}$ and $\text{ER}_{\text{fine}}$ give 447.9±81.2 Mg and 301.9±114.0 Mg respectively, see Table S2. Not only does
the emission total change with ER calculation method, but so does interannual variability
(from 17% to 38%) and the spatial correlation pattern (see R in Table S2). Nearly all of the
difference is due to the distinction between savanna and tropical forest in Africa. The ER for
savanna, at $0.28 \times 10^{-7}$ is less than 20% of ER$_{av}$ ($1.54 \times 10^{-7}$), and even though the tropical
forest ER is higher than the global average ($2.05 \times 10^{-7}$) this does not compensate for the
decrease in Hg emissions from the vast savanna regions of Africa. In comparison the higher
ratio of forest to savanna in South America means that overall there is little change in the
total Hg emissions for this region. The simulations using the more detailed ER estimates
show a decreased spatial correlation for Hg deposition with respect to the simulations using
an average ER, see Tables 2, 3 and S2. Although the magnitude of Hg emission, and therefore
also the magnitude of the Hg deposition flux, is different using the specific ERs, the impact
on the geographical distribution of the deposition is limited.

3.4.2 Injection height

The height at which emissions from BB are introduced into the model can have a significant
impact on pollutant transport. Some recent studies have shown that boreal fire emissions can
be lofted above the boundary layer$^{34,35}$. A long term study of the CALIOP (Cloud-Aerosol
Lidar with Orthogonal Polarization) profiles over South-Western Russia and Eastern Eu-
rope for 2006–2008 showed that as much as 50% of the BB plumes were above the boundary
layer$^{36}$. A detailed review of injection heights and plume rise models can be found in Ichoku
et al.$^{33}$. Simulations were performed in which the emissions were added to different model
levels up to approximately 2000 m. Further simulations, one in which the emissions were
distributed uniformly throughout the lower levels of the model, and a second with a pre-
scribed latitudinally dependent vertical distribution, were performed$^{37,38}$. Comparing the
Hg deposition patterns obtained in these experiments to the base case reveals a very high
correlation, R $\approx$1, see table 3. The atmospheric lifetime of Hg$^{0}_{e(k)}$ is the main reason for this
lack of influence of the emission height on the simulated deposition fields. Similar results
have been obtained in studies of CO plumes, where the impact of emission height on atmospheric composition is significant locally, and only has a minor influence on regions distant from the plume source.\textsuperscript{39–41}

3.4.3 Sensitivity to oxidation mechanism

As mentioned in Section 2.2 the precise mechanism by which Hg$^0_{(g)}$ is oxidized in the atmosphere is not yet certain.\textsuperscript{25–27} Most models opt for a combination of O$_3$/OH, or alternatively a Br based oxidation mechanism. In either case Hg$^0_{(g)}$ has an atmospheric lifetime of approximately 8 to 12 months, which is consistent with the observed difference in the hemispherical background concentrations of Hg$^0_{(g)}$ (roughly 1.7 ng m$^{-3}$ in the Northern Hemisphere and 1.2 in the southern). For 2010 simulations were performed utilizing each oxidation pathway with each BB emission inventory. Further simulations, using fixed atmospheric lifetimes against oxidation (e-folding time, see description in the SI) of 12 and 6 months were also performed. The agreement maps for the simulations are presented in figure S6.

Although the number of cells where all the inventories agree that Hg deposition is high does not differ greatly between the different simulations, the distribution of the ‘agreement’ does. This is particularly true of the tropical Atlantic; using the Br mechanism there is no ‘high’ deposition area to the west of Africa, however the ‘high’ deposition region in the North Atlantic reaches Iceland, which it does not in the O$_3$/OH simulation. Again, in the Br simulation the ‘high’ deposition area reaches into the Gulf of Alaska, whereas in the O$_3$/OH simulation the ‘high’ deposition regions are more closely confined to a relatively narrow latitude band between the tropics, reflecting the distribution of O$_3$ in the troposphere. The Br simulation does show noticeably more areas where only the GFED inventory predicts high deposition, particularly in the Southern Ocean. This is in part due to the higher southern hemisphere emissions in GFED, but also because the period of the year when biomass burning is most prevalent in South America, July to September, coincides with low tropospheric Br concentrations, so that the emissions are transported much further in this simulation.
than in the O$_3$/OH simulation.

The simulations using a fixed atmospheric lifetime for Hg give results that are more similar to the Br mechanism, particularly in the case of the 12 month lifetime. In neither of the two simulations is the high deposition distribution as closely confined to the area between the tropics as in the O$_3$/OH case. In all the simulations most Hg deposition from BB emissions is deposited to the oceans. Clearly more monitoring sites in the Tropics would help immensely to understand more fully the importance of BB Hg emissions on oceanic Hg deposition. Table S3 summarises the simulated Hg deposition to the world’s ocean basins. The table includes the simulated deposition totals calculated using full atmospheric emissions (natural, anthropogenic and BB) for the two oxidation mechanisms for 2010.

### 3.4.4 Uncertainty in the Deposition fields

The Hg deposition fields obtained in this study vary and it is not immediately clear where and to what extent the results agree. In order to examine the ‘ensemble’ of results, rather than just averaging the full set of simulations, the model output has been tested against the Base run (GFED, O$_3$/OH, global ER, year 2010) to ascertain the probability that the deposition fields belong to the same distribution. This then permits those results which differ the most to be identified. This form of ‘inspected’ ensemble was recently described by Solazzo and Galmarini$^{42}$ for a multi-model ensemble. The non-parametric Kolmogorov-Smirnov two-sample test has been used to examine the results of the sensitivity tests performed using the GFED inventory. The test was repeated with the model output obtained using the FINN and GFAS inventories with the O$_3$/OH and Br oxidation mechanisms and with the 12 month pseudo-oxidation approach. The results of the test are shown in Table 3. A value of $\text{Prob}_{KS-test} \leq 0.05$ indicates that it is improbable that the simulated Hg deposition fields belong to the same distribution. The height at which the emissions are introduced into the model, and the first Enhancement Ratio variation ($\text{ER}_{coarse}$) make very little difference to the output results. The most important factors influencing the output fields are the inventory
Table 3: Correlations and probabilities that the sensitivity run Hg deposition fields belong to the same distribution as the GFED 2010 simulation deposition field, and comparison with FINN and GFAS

<table>
<thead>
<tr>
<th>RUN</th>
<th>Sensitivity assessment</th>
<th>12 m</th>
<th>O$_3$+OH</th>
<th>Br</th>
</tr>
</thead>
<tbody>
<tr>
<td>Emissions</td>
<td>Vertically distributed</td>
<td>1.00</td>
<td>1.00</td>
<td>1.00</td>
</tr>
<tr>
<td>Emissions</td>
<td>Inj. into ind. levels</td>
<td>1.00</td>
<td>1.00</td>
<td>1.00</td>
</tr>
<tr>
<td>Oxidation mech.</td>
<td>O$_3$/OH</td>
<td>0.91</td>
<td>&lt;0.05</td>
<td>0.81</td>
</tr>
<tr>
<td>Oxidation mech.</td>
<td>Br</td>
<td>0.96</td>
<td>&lt;0.05</td>
<td>0.81</td>
</tr>
<tr>
<td>Lifetime Hg(0)</td>
<td>12 months</td>
<td>–</td>
<td>–</td>
<td>0.96</td>
</tr>
<tr>
<td>Lifetime Hg(0)</td>
<td>6 months</td>
<td>0.99</td>
<td>0.41</td>
<td>0.97</td>
</tr>
<tr>
<td>Enhancement Ratio</td>
<td>ER$_{coarse}$</td>
<td>1.00</td>
<td>0.99</td>
<td>1.00</td>
</tr>
<tr>
<td>Enhancement Ratio</td>
<td>ER$_{fine}$</td>
<td>0.99</td>
<td>0.52</td>
<td>1.00</td>
</tr>
<tr>
<td>FINN 2010 emissions</td>
<td></td>
<td>0.97</td>
<td>&lt;0.05</td>
<td>0.98</td>
</tr>
<tr>
<td>GFAS 2010 emissions</td>
<td></td>
<td>0.97</td>
<td>&lt;0.05</td>
<td>0.99</td>
</tr>
</tbody>
</table>

and the oxidation mechanism. The second variation of the Enhancement Ratio (ER$_{fine}$) described in Section 3.4.1 also results in noticeably different deposition fields even if the hypothesis that the model output belongs to the same distribution as the Base case cannot be rejected, $\text{Prob}_{KS-test} = 0.17$ and 0.10, with the O$_3$/OH and Br oxidation mechanisms respectively. This is also true for the GFAS inventory $\text{Prob}_{KS-test} = 0.08$ and 0.09, however these values indicate that the probability of belonging to the same distribution is low. The results from the three inventories, and also the ER2 sensitivity run, with both the O$_3$/OH and Br oxidation mechanisms have therefore been averaged to obtain an 'ensemble' deposition field, which is illustrated in figure 5. The figure makes it evident that however much the simulated deposition fields differ, the regions most influenced by Hg deposition from biomass burning are the tropical areas of the oceans, the North Atlantic and the North Pacific.

4 Discussion

Just over 75% of the Hg released by BB is deposited to the world’s oceans and seas. As is well known, human exposure to methylmercury (the most toxic form) occurs predominantly through fish consumption. Hg$^{II}$ deposited to the ocean may be reduced and re-emitted...
Figure 5: Geographical distribution of the probability density function of the total Hg deposition obtained from an inspected ensemble of simulations for the year 2010. Total deposition is illustrated in terms of the average ($\mu$) and standard deviation $\sigma$ of the ensemble.

Total Dep : 580.45 ± 130.01 Mg y$^{-1}$

Flux : 0.97 ± 0.47 g km$^{-2}$ y$^{-1}$
from the sea surface, but a part can be methylated in surface or subsurface waters, where it can enter the food web\textsuperscript{2,43,44}. The maximum deposition fluxes in the individual oceanic regions, are very similar for all the BB inventories. From the results obtained from the five year runs it was found that the North Atlantic has the highest peak deposition flux value at \(21 \text{ g km}^{-2}\), followed by the North Pacific and Indian Oceans at \(\approx 20 \text{ g km}^{-2}\). The maximum Hg deposition flux in the Arctic reaches \(7 \text{ g km}^{-2}\), higher than the Mediterranean (\(6 \text{ g km}^{-2}\)) and the Southern Ocean (\(3 \text{ g km}^{-2}\)). The total calculated emissions of Hg from BB are similar for all three inventories used in this study, although there are differences in their geographical distribution. GFED has a higher proportion of emissions in the southern hemisphere (Figure 1c) in comparison to the other two inventories and this is also visible in the deposition fields (Figure 1d). However the lifetime of Hg\textsubscript{0} is such that the differences in the spatial distribution of the emissions is far less evident in the simulated deposition fields. GFED is a slight exception as the distribution, relatively to the other two inventories has a higher proportion of emissions in the southern hemisphere (Figure 1c) this is visible also in the deposition fields (Figure 1d). One effect of BB is to emit Hg from lower latitudes for eventual deposition at higher latitudes, in both hemispheres. The presence of higher latitude boreal forests in the Northern Hemisphere does mean that the Arctic is more impacted than the Antarctic by Hg deposition resulting from BB. The highest Hg deposition fluxes are found in the North Atlantic, while the the greatest total Hg deposition is to the North Pacific.

The oxidation mechanism and the choice of emission inventory have the greatest influence on the spatial distribution of the Hg deposition fields. The factor which most influences the total calculated Hg emission from BB is the enhancement ratio. More biome specific Hg/CO enhancement ratios are needed to better constrain the magnitude of Hg emissions from BB.

In order to build a bottom-up inventory it would be necessary to perform measurements of Hg and CO released by BB and also ideally to distinguish between the same biomes on different continents. As the number of Hg monitoring sites around the world increases, intermittent information will become more abundant as stations will at times be downwind
of BB plumes, however a more targeted approach addressing, tropical, savanna and boreal ecosystems would be far better. Biomass burning will continue to play a role in the cycling of Hg, and legacy Hg particularly, for a long time to come. As the Minamata Convention comes into force and anthropogenic emissions begin to be curbed, the role of BB in cycling Hg from the tropics to higher latitudes, and particularly in transferring Hg from terrestrial reservoirs to the oceans will become more important. Understanding the recycling of legacy Hg is particularly important in the assessment of the response times of ecosystems to changes in anthropogenic emissions, especially should the frequency and scale of BB increase as the climate changes.

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Supporting Information Available

Tables summarizing the simulations performed with ECHMERIT, the spatial correlation (R) for different ERs, annual Hg deposition to the major oceans basins. Figures illustrating the Base, Br and fixed lifetime model results, and agreement maps of Hg deposition. Sections describing the pseudo-language algorithm used to generate Agreement Maps, the calculation of ERs, and the oxidation mechanisms implemented in the model. This material is available free of charge via the Internet at http://pubs.acs.org/.
References


(33) Ichoku, C.; Kahn, R.; Chin, M. Satellite contributions to the quantitative characteri-

(34) Lapina, K.; Honrath, R. E.; Owen, R. C.; Val Martín, M.; Hyer, E. J.; Fialho, P. Late
497 summer changes in burning conditions in the boreal regions and their implications for
498 NOx and CO emissions from boreal fires. *J. Geophys. Res. Atmos.* **2008**, *113*, D11304–.

500 Smoke injection heights from fires in North America: analysis of 5 years of satellite

(36) Amiridis, V.; Giannakaki, E.; Balis, D. S.; Gerasopoulos, E.; Pytharoulis, I.; Zanis, P.;
505 Kazadzis, S.; Melas, D.; Zerefos, C. Smoke injection heights from agricultural burning

(37) Williams, J. E.; van Weele, M.; van Velthoven, P. F. J.; Scheele, M. P.; Liou, C.;
510 van der Werf, G. R. The Impact of Uncertainties in African Biomass Burning emission
511 estimates on modeling global air quality, long range transport and tropospheric chemical

(38) Dentener, F. et al. Emissions of primary aerosol and precursor gases in the years 2000

519 Yurganov, L. Impacts of enhanced biomass burning in the boreal forests in 1998 on
520 tropospheric chemistry and the sensitivity of model results to the injection height of

(40) Guan, H.; Chatfield, R. B.; Freitas, S. R.; Bergstrom, R. W.; Longo, K. M. Modeling
525 the effect of plume-rise on the transport of carbon monoxide over Africa with NCAR
(41) Jian, Y.; Fu, T.-M. Injection heights of springtime biomass-burning plumes over penin-

(42) Solazzo, E.; Galmarini, S. A science-based use of ensembles of opportunities for as-


(44) Blum, J. D.; Popp, B. N.; Drazen, J. C.; Anela Choy, C.; Johnson, M. W. Methylmer-
curry production below the mixed layer in the North Pacific Ocean. *Nat. Geosci.* 2013, 6, 879–884.