A global perspective on aerosol from low-volatility organic compounds

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Abstract. Global production of organic aerosol from primary emissions of semivolatile (SVOCs) and intermediate (IVOCs) volatility organic compounds is estimated using the global chemical transport model, GEOS-Chem. SVOC oxidation is predicted to be a larger global source of net aerosol production than oxidation of traditional parent hydrocarbons (terpenes, isoprene, and aromatics). Using a prescribed rate constant and reduction in volatility for atmospheric oxidation, the yield of aerosol from SVOCs is predicted to be about 75% on a global, annually-averaged basis. For IVOCs, the use of a naphthalene-like surrogate with different high-NO_x and low-NO_x parameterizations produces a global aerosol yield of about 30%, or roughly 5 Tg/yr of aerosol. Estimates of the total global organic aerosol source presented here range between 60 and 100 Tg/yr. This range reflects uncertainty in the parameters for SVOC volatility, SVOC emissions, and IVOC emissions, as well as wet deposition. The highest estimates result if SVOC emissions are significantly underestimated (by more than a factor of 2) or if wet deposition of the gas-phase semivolatile species is less effective than previous estimates. A significant increase in SVOC emissions, a reduction of the volatility of the SVOC emissions, or an increase in the enthalpy of vaporization of the organic aerosol all lead to an appreciable reduction of prediction/measurement discrepancy. In addition, if current primary organic aerosol (POA) inventories capture only about one-half of the SVOC emission and the Henrys Law coefficient for oxidized semivolatiles is on the order of 10^3 M/atm, a global estimate of OA production is not inconsistent with the top-down estimate of 140 Tg/yr by (Goldstein and Galbally, 2007). Additional information is needed to constrain the emissions and treatment of SVOCs and IVOCs, which have traditionally not been included in models.

1 Introduction

Organics represent a significant fraction of the aerosol mass in the atmosphere (Zhang et al., 2007). Bottom-up estimates of particulate matter concentrations, such as those that would be used to estimate aerosol radiative forcings, visibility, or implications for public health, must be able to represent the processes critical for organic aerosol formation. Work by Lipsky and Robinson (2006) and Robinson et al. (2007) as well as others (Huffman et al., 2009a,b) indicates that what has traditionally been considered non-volatile primary organic aerosol (POA) is actually a dynamic system of semivolatile species that partition between the gas and aerosol phases as well as undergo gas-phase oxidation to form species of different volatilities that can condense to form secondary organic aerosol (SOA). We present the first estimates of the contribution of primary semivolatile organic compounds and other low-volatility organic compounds to global aerosol production.

Emissions of low-volatility organic compounds can be subdivided somewhat arbitrarily into two classes based on volatility (Donahue et al., 2006): semivolatile organic compounds (SVOCs) and intermediate volatility organic compounds (IVOCs). Semivolatile organic compounds are those that partition directly between the gas and aerosol phases under ambient conditions and include compounds with saturation concentrations roughly between 0.1 and 10^4 µg/m^3. SVOC emissions include traditional POA plus any vapor phase species that are in direct equilibrium with the particle...
Aerosol formation from SVOCs, IVOCs, and their oxidation products has been implemented in regional models such as PMCAMx and CHIMERE (Robinson et al., 2007; Shrivastava et al., 2008; Murphy and Pandis, 2009; Hodzic et al., 2010) as well as box models (Dzepina et al., 2009; Grieshop et al., 2009a). The SVOCs, and IVOCs if applicable, are typically represented using the volatility basis set framework (Donahue et al., 2006) and allow for oxidation in the gas phase and formation of lower volatility products. Replacing the traditional non-volatile POA with semivolatile POA in regional models has lead to improvements in the urban to regional organic aerosol (OA) concentration ratio (Robinson et al., 2007; Shrivastava et al., 2008) and predicts an ambient aerosol more dominated by oxygenated species (Shrivastava et al., 2008; Murphy and Pandis, 2009), consistent with observations.

In this work, organic aerosol formation from SVOCs and IVOCs is studied using the global chemical transport model, GEOS-Chem. Section 2 describes the global model framework as well as the emissions and atmospheric transformation of SVOCs and IVOCs. Results are presented in Sect. 3 in terms of predicted aerosol levels, global budgets, and modern vs. fossil carbon. The paper finishes by addressing model uncertainties (Sect. 4) and placing the results in the larger context of top-down vs. bottom-up global aerosol budgets (Sect. 5).

2 Model description

2.1 Global model

The global chemical transport model, GEOS-Chem (version 8-01-04, http://acmg.seas.harvard.edu/geos/), is used to simulate year 2000 organic aerosol concentrations. Simulations are conducted at 2° latitude by 2.5° longitude horizontal resolution using GEOS-4 assimilated meteorology with 30 vertical layers going up to 0.01 hPa for baseline simulations. For computational speed, and in preparation for climate change-organic aerosol interaction studies, sensitivity tests are performed at 4° latitude by 5° longitude horizontal resolution with 23 vertical levels up to 0.002 hPa using GISS GCM Model III meteorology (Rind et al., 2007; Wu et al., 2007, 2008). All simulations are conducted for year 2000 with a minimum of 11 months of initialization. Simulations include fully coupled ozone-NOx-hydrocarbon chemistry (Bey et al., 2001), and formation of inorganic (Park et al., 2004; Pye et al., 2009) and organic aerosol (Park et al., 2003, 2006; Henze and Seinfeld, 2006; Liao et al., 2007; Henze et al., 2008).

2.2 Absorptive partitioning

Formation of organic aerosols occurs by absorptive partitioning (Odum et al., 1996), and the partitioning framework used here is based on the work of Chung and Seinfeld (2002).
An equilibrium partitioning coefficient, \( K_{OM,i} \), describing the partitioning between the gas and aerosol phases of a semivolatile species can be calculated assuming a pseudo-ideal solution and absorptive partitioning theory (Pankow, 1994):

\[
K_{OM,i} = \frac{RT}{\tilde{M}_i \gamma_i P_{i_{vap}}} \tag{1}
\]

where \( R \) is the gas constant, \( T \) is temperature, \( \tilde{M}_i \) is the molecular weight, \( \gamma_i \) is the activity coefficient of compound \( i \) in the aerosol phase, and \( P_{i_{vap}} \) is the vapor pressure of pure species \( i \) at temperature \( T \). Absorptive partitioning can also be described using a saturation concentration, \( C^*_i \), which is the inverse of the equilibrium partitioning coefficient (for a discussion of the difference between \( C^*_i \) and \( K_{OM,i} \) see the supporting material from Donahue et al., 2006):

\[
K_{OM,i} = \frac{1}{C^*_i}. \tag{2}
\]

In terms of concentrations of the semivolatile \( i \), \([G_i]\) and \([A_i]\), in the gas and aerosol phases, respectively,

\[
K_{OM,i} = \frac{[A_i]}{[G_i]\tilde{M}_i} \tag{3}
\]

where \([M_o]\) is the concentration of particle-phase absorptive material into which the semivolatile compound can partition. In this study, the partitioning medium includes only the particle-phase organics (inorganic constituents and water included in some studies are not considered here). Primary and secondary organic aerosol is assumed to form a single absorbing phase. When non-volatile POA is present,

\[
[M_o] = [POA] + \sum_i [A_i]. \tag{4}
\]

and if POA is semivolatile, this reduces to,

\[
[M_o] = \sum_i [A_i]. \tag{5}
\]

Combining Eqs. (3) and (4) or (5) as appropriate, along with a mass balance, yields an implicit equation for \( M_o \). Organic aerosol will form only when (i.e. the implicit equation for \( M_o \) will only have a solution when) (Chung and Seinfeld, 2002),

\[
\sum_i K_{OM,i} ([A_i] + [G_i]) > 1. \tag{6}
\]

Simulations indicate that this condition is generally satisfied with a few exceptions. Formation of organic aerosol through other means, such as cloud processing or reactive uptake, are not considered in the present study.

### 2.3 SOA from traditional precursors

SOA from traditional precursors follows earlier studies for terpenes (Chung and Seinfeld, 2002), isoprene (Henze and Seinfeld, 2006), and aromatics (Henze et al., 2008). Parent hydrocarbons are oxidized in the gas-phase to form a series of semivolatile species,

\[
HC + O_x \rightarrow \alpha_1 P_1 + \alpha_2 P_2 + \ldots \tag{R1}
\]

where \( \alpha_1, \alpha_2, \ldots \) are mass-based stoichiometric coefficients for products \( P_1, P_2, \ldots \) Sometimes, only one semivolatile or nonvolatile product is necessary to represent chamber data (like for low-NOx aromatic oxidation (Ng et al., 2007; Chan et al., 2009) or nitrate radical oxidation of terpenes (Chung and Seinfeld, 2002)), but typically two semivolatile products have been used (Odum et al., 1996) with partitioning parameters, \( \alpha_i \) and \( K_{OM,i} \), determined by fits to laboratory data. Models such as PMCAMx have been used to investigate the possibility of continued aging of the SOA formed from traditional parent hydrocarbons due to continued OH oxidation of the gas-phase semivolatile species (Lane et al., 2008). In the present work, traditional SOA species are not considered subject to continued oxidation beyond that captured in chamber studies.

### 2.4 Aerosol from SVOCs

POA is defined to be any SVOC that partitions directly to the particle phase after emission without undergoing oxidation. In source regions, there is likely to be net POA production, but as air masses move to more remote regions, POA will be driven out of the particle phase due to dilution and oxidation of the gas-phase species. Oxidation of the primary gas-phase SVOCs can lead to lower volatility products that partition to make SOA.

#### 2.4.1 SVOC emissions

SVOCs from all sources are assumed to be emitted as two semivolatile surrogate species, SVOC\(_1\) and SVOC\(_2\), in roughly equal fractions of 0.49 and 0.51 based on the work by Shrivastava et al. (2006) and Lipsky and Robinson (2006). Partitioning coefficients for SVOC\(_1\) and SVOC\(_2\) are given in Table 1 and correspond to saturation concentrations of roughly 1600 and 20 \( \mu g/m^3 \). Under most atmospherically relevant conditions, only the lower volatility component is expected to partition appreciably to the aerosol phase.

The SVOC emissions are based on the traditional nonvolatile POA emission inventory used in GEOS-Chem (Park et al., 2003, 2006). The non-volatile POA inventory includes contributions from biomass burning, biofuel burning, and anthropogenic sources (Table 2 and Fig. 1). Monthly biomass burning emissions are based on the Global Fire and Emission Database version 2 (GFEDv2) for year 2000 (van der Werf et al., 2006). Global biofuel and anthropogenic organic...
Table 1. Low-volatility organic compounds.

<table>
<thead>
<tr>
<th>Species</th>
<th>Description</th>
<th>$\alpha$</th>
<th>$K_{DM}^b$ [m$^3$/µg]</th>
<th>$C^*$ [µg/m$^3$]</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>SVOC$_1$</td>
<td>primary SVOC emission</td>
<td>0.49$^c$</td>
<td>0.0006</td>
<td>1646</td>
<td>Shrivastava et al. (2006)</td>
</tr>
<tr>
<td>SVOC$_2$</td>
<td>primary SVOC emission</td>
<td>0.51$^c$</td>
<td>0.05</td>
<td>20</td>
<td>Shrivastava et al. (2006)</td>
</tr>
<tr>
<td>O−SVOC$_1$</td>
<td>oxidized SVOC$_1$</td>
<td>1.5</td>
<td>0.06</td>
<td>16.46</td>
<td>Grieshop et al. (2009a)</td>
</tr>
<tr>
<td>O−SVOC$_2$</td>
<td>oxidized SVOC$_2$</td>
<td>1.5</td>
<td>5.0</td>
<td>0.20</td>
<td>Grieshop et al. (2009a)</td>
</tr>
<tr>
<td>IVOC</td>
<td>primary IVOC emission$^d$</td>
<td>NA</td>
<td>$1 \times 10^{-5}$</td>
<td>$10^3$</td>
<td></td>
</tr>
<tr>
<td>O−IVOC$_{N,1}$</td>
<td>high NO$_x$ IVOC oxidation product$^d$</td>
<td>0.21</td>
<td>0.59</td>
<td>1.69</td>
<td>Chan et al. (2009)</td>
</tr>
<tr>
<td>O−IVOC$_{N,2}$</td>
<td>high NO$_x$ IVOC oxidation product$^d$</td>
<td>1.07</td>
<td>0.0037</td>
<td>270</td>
<td>Chan et al. (2009)</td>
</tr>
<tr>
<td>O−IVOC$_{H,1}$</td>
<td>low NO$_x$ IVOC oxidation product$^d$</td>
<td>0.73</td>
<td>10,000</td>
<td>0.0001</td>
<td>Chan et al. (2009)</td>
</tr>
</tbody>
</table>

$^a$ See Reaction (R1).
$^b$ Reference temperature for SVOC parameters is 300 K. IVOC SOA reference temperature is 299 K.
$^c$ Fraction of total SVOC emissions.
$^d$ IVOC behavior based on naphthalene.

Table 2. Emissions of primary low-volatility organic compounds (year 2000).

<table>
<thead>
<tr>
<th>Species</th>
<th>Biomass burning [Tg C/yr]</th>
<th>Biofuel burning [Tg C/yr]</th>
<th>Anthropogenic sources$^a$ [Tg C/yr]</th>
<th>Total [Tg C/yr]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Traditional non-volatile POA</td>
<td>19</td>
<td>7.1</td>
<td>2.7</td>
<td>29</td>
</tr>
<tr>
<td>SVOCs</td>
<td>24</td>
<td>9.0</td>
<td>3.4</td>
<td>37</td>
</tr>
<tr>
<td>Naphthalene$^b$</td>
<td>0.09</td>
<td>0.05</td>
<td>0.09</td>
<td>0.22</td>
</tr>
<tr>
<td>I VOC surrogate</td>
<td>5.7</td>
<td>3.2</td>
<td>5.8</td>
<td>15</td>
</tr>
</tbody>
</table>

$^a$ Excluding biomass and biofuel burning.
$^b$ The baseline naphthalene emission inventory is used only to obtain the spatial distribution of IVOC emissions.

Carbon emissions are from the technology-based inventory by Bond et al. (2004). Over North America, anthropogenic emissions are superseded with those based on work by Cooke et al. (1999). US biofuel emissions are constructed based on residential and industrial wood fuel consumption, as implemented in the work of Park et al. (2003). The scaling and seasonal distribution of US anthropogenic and biofuel organic carbon (OC) emissions are also detailed in the work of Park et al. (2003).

Providing a global estimate of SVOC emissions is difficult due to the fact that it is unclear what portion of the SVOCs traditional POA represents, and traditional POA emissions, themselves are uncertain. Estimates of the traditional POA emission rate in global models has ranged from 17 to 142 Tg/yr as summarized in the work of Farina et al. (2010). Note that since biomass burning is a significant POA contributor, emissions of SVOCs likely vary from year to year as different areas become more or less prone to burning. Part of the uncertainty in global OC emissions, estimated at a factor of two in the work by Bond et al. (2004), is likely due to the fact that traditional POA inventories try to capture organic aerosol emissions over all atmospherically relevant conditions using measurements or observations of emission factors under a single or limited set of conditions in terms of temperature and organic aerosol loadings. For a species that is semivolatile, like POA, one set of conditions will not characterize the emissions well.

Debate continues on whether traditional POA inventories represent most SVOC emissions or a very limited subset. POA emission factors are often obtained under organic concentrations higher than atmospherically relevant, which may force organic vapors that would be in the gas phase under ambient conditions into the particle phase during sampling. Under such conditions, the POA inventory may be a good representation of SVOC emissions. However, an examination of the Schauer et al. (2001) inventory for wood burning, which gives OC emission factors consistent with those used in the GFEDv2 biomass burning inventory, shows individual PAH compounds with saturation concentrations of 12, 100, and $9 \times 10^3$ µg/m$^3$ being emitted 91, 56, and 4% in the particle phases. Thus, compounds over the entire SVOC volatility range exhibit appreciable mass in the gas and particle phase. Furthermore, models using non-volatile POA tend to underestimate, not overestimate, ambient organic aerosol concentrations (Heald et al., 2005; Liao et al., 2007). We proceed by assuming traditional non-volatile POA inventories likely represent a subset of SVOC emissions.
Table 3. Parameters for SVOC oxidation.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>This work</th>
<th>Robinson et al. (2007)</th>
<th>Grieshop et al. (2009a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas-phase OH rate constant [cm$^3$ molec$^{-1}$ s$^{-1}$]</td>
<td>$2 \times 10^{-11}$</td>
<td>$4 \times 10^{-11}$</td>
<td>$2 \times 10^{-11}$</td>
</tr>
<tr>
<td>Reduction in volatility per reaction</td>
<td>100×</td>
<td>10×</td>
<td>100×</td>
</tr>
<tr>
<td>Increase in mass per reaction</td>
<td>50%</td>
<td>7.5%</td>
<td>40%</td>
</tr>
<tr>
<td>Number of oxidation reactions per parent hydrocarbon</td>
<td>1</td>
<td>&gt; 1</td>
<td>&gt; 1</td>
</tr>
</tbody>
</table>

Fig. 1. Emissions of SVOCs and IVOCs. SVOC emissions shown here are the baseline POA emission inventory. Plots are annual averages for year 2000. Note that the color scale is not linear.

The SVOC emission rate in grid cell $I, J$ can be estimated using:

$$E_{SVOC}(I, J) = \left[1 + \frac{G_R}{POA_R}\right]E_{NVPOA}(I, J)$$  \hspace{1cm} (7)

where $E_{SVOC}$ is the emission rate of SVOCs (Tg C/yr), $G_R$ is the emission of the gas-phase SVOC surrogate in a representative study (mg/kg), $POA_R$ is the emission of organic aerosol in that same representative study (mg/kg), and $E_{NVPOA}$ is the traditional, non-volatile POA emission rate (Tg C/yr). For unit conversion purposes, the organic matter to organic carbon (OM/OC) ratio is assumed to be the same for $G_R$ and $POA_R$ in the above equation. Using the work of Schauer et al. (2001) as the representative study, the ratio of gas-phase SVOC surrogates to the particle-phase organics ($G_R/POA_R$) is about 0.27. The gas-phase SVOC surrogate estimate is based on adding up the gas-phase speciated emissions that were also found in the particle phase. Some of the SVOC emissions could be part of the gas-phase unresolved complex mixture in the work of Schauer et al. (2001) or were misclassified as IVOCs and are not reflected in the 0.27 estimate. Accounting for all the gas-phase species on the PUF sampling train/filter and the entire gas-phase UCM would increase the 0.27 estimate to 0.61. The possibility remains that POA inventories may represent a significantly
different fraction of SVOCs than in the study of Schauer et al.
(2001), and scaling POA emissions up 27% is likely a fairly
conservative estimate, given that the uncertainty in the base-
line POA emissions is easily a factor of 2 (Bond et al., 2004).
Global emissions of SVOCs are predicted to be 37 Tg C yr\(^{-1}\)
(Table 2 and Fig. 1).

Since traditional POA inventories are reported as the mass
of carbon emitted, the mass of organic matter that includes
any additional oxygen, nitrogen, or other species associated
with the carbon must be determined. The range of OM/OC
ratios for SVOCs likely ranges from just above 1 for carbon-
rich species to above 2 for highly oxygenated species such
as those found in wood smoke (Schauer et al., 2001). Simu-
lations here use a value of 1.4, which is only slightly higher
than the OM/OC ratio estimated for hydrocarbon like organic
aerosol, HOA (OM/OC: 1.2−1.3) (Aiken et al., 2008; Zhang
et al., 2005), slightly lower than the OM/OC ratio for pri-
mary biomass burning organic aerosol, P-BBOA (OM/OC:
1.6−1.7) (Aiken et al., 2008), and consistent with the wood
burning inventory of Schauer et al. (2001) (OM/OC: 1.4−
1.7). The OM/OC ratio affects the partitioning of organics
since it determines the total SVOC mass available for parti-
tioning.

2.4.2 SVOC oxidation

Primary SVOC emissions age in the gas phase by reaction
with OH, based on the wood burning chamber experi-
ments of Grieshop et al. (2009a). An OH rate constant
of \(2 \times 10^{-11} \text{ cm}^3/\text{molec/s}\) is assumed for the present work.
The mass of the parent SVOC is assumed to increase 50%
through functionalization, a value slightly larger than, but
still consistent with, Grieshop et al. (2009a) for oxidation
of wood smoke. The volatility of the SVOC is reduced by
a factor of 100 as a result of the OH reaction. This represents
a slightly slower rate of oxidation, but a more aggressive addi-
tion of oxygen and reduction in volatility, than that used
by Robinson et al. (2007) and was found to give better O:C
ratio agreement with experiments than the traditional Robin-
son et al. (2007) parameters in the work of Grieshop et al.
(2009a). Table 3 summarizes the SVOC oxidation parameter-
s and compares them with the optimized parameters in the
work of Grieshop et al. (2009a) and the parameters of Robin-
son et al. (2007).

The mechanism by which aging of low-volatility organic
compounds occurs is not well-constrained. Previous work
(Robinson et al., 2007) has suggested that aging occurs as
the result of sequential OH oxidation reactions in the gas-
phase. The SVOCs in the present study are assumed to un-
dergo only one generation of oxidation. This assumption
is made for three main reasons. First, chamber studies, which
are the source of oxidation data, tend to access only initial
reactions. Second, since the model assumes that oxidation
leads only to functionalization of the molecule and there-
fore a reduction in volatility, it is likely to be less valid for
later generations of oxidation in which molecular fragmen-
tation becomes more important (Kroll et al., 2009). Lastly,
for a 50% (or 40% as used by Grieshop et al., 2009a) in-
crease in mass per generation, a parent SVOC with an initial
OM/OC of 1.4 would require only 1 generation of oxidation
to reach a final OM/OC of about 2, consistent with the ob-
served OM/OC of aged aerosol in the experiment on which
the parameters are based (Grieshop et al., 2009a), with the
OM/OC of oxygenated organic aerosol, OOA (OM/OC: 1.8−
2.4, 2.2) (Aiken et al., 2008; Zhang et al., 2005), and with
other estimates of the OM/OC of oxidized regional aerosol
(OM/OC: 2.1) (Turpin and Lim, 2001).

2.5 Aerosol from IVOCs

IVOCs are emitted entirely in the gas phase and form aerosol
only upon oxidation. Since naphthalene is predicted to be
a major SOA precursor in the oxidation of wood burning
and diesel combustion exhaust (Chan et al., 2009), here,
IVOCs are represented as a naphthalene-like surrogate that
forms aerosol according to the chamber studies of Chan et al.
(2009).

In addition to naphthalene, other important classes of spe-
ciated IVOCs include alkanes and phenols (Schauer et al.,
2001, 2002). A significant portion of IVOCs likely belong
to the unresolved complex mixture and thus their identity
and aerosol yield is not known. Aerosol yield information
is available for many alkanes (Jordan et al., 2008) as well as
naphthalene and functionalized naphthalene species (Chan
et al., 2009). Gasoline combustion likely produces more
alkanes than aromatics (Schauer et al., 2002), but wood burn-
ing, globally the largest source of POA, likely emits more
gas-phase aromatics than alkanes (Schauer et al., 2001; Hays
et al., 2002). Approximately 75% of the speciated IVOC sur-
rogate in the wood burning inventory of Schauer et al. (2001)
is phenol (\(C_\alpha\) about \(10^6 \text{ µg/m}^3\)) or substituted phenol com-
ounds. Due to their ring structures, these phenol compounds
are likely to behave more like naphthalene than an alkane in
terms of yields under high-NOx vs. low-NOx conditions. So,
if all IVOCs are to be represented with one surrogate com-
pound, naphthalene is a good choice.

2.5.1 IVOC emissions

Since naphthalene is an important IVOC from many sources
including wood combustion and vehicle exhaust (Chan et al.,
2009; Schauer et al., 2001, 2002), IVOC emissions are as-
sumed to be spatially distributed like naphthalene. First,
a baseline emission inventory of naphthalene (NAP) is cre-
dated. For biomass and biofuel burning, this is done using an
emission ratio to CO. An emission factor of 0.025 g NAP/kg
dry matter burned is used along with CO emission factors
(Andreae and Merlet, 2001) to produce emission ratios of
0.0602 and 0.0701 mmol NAP/mol CO for biomass and bio-
fuel burning, respectively. The emission factor presented by
Andreae and Merlet (2001) is consistent with that of naphthalene presented by Hays et al. (2002) for the burning of fossil fuels, but about a factor of 10 lower than the value in the work of Schauer et al. (2001). Anthropogenic naphthalene emissions, from sources such as traffic oil combustion, consumer products, and industrial sources (Zhang and Tao, 2009), are spatially distributed like benzene from the EDGAR2 inventory and are aseasonal. Emissions given for 1990 are scaled to year 2000 based on CO2 emissions from liquid fossil fuel usage (Bey et al., 2001; Fu et al., 2008). The magnitude of non-biomass and biofuel burning emissions of naphthalene is estimated based on Zhang and Tao (2009) to be about 0.09 Tg C/yr. The baseline naphthalene emission inventory from all sources constructed here is approximately 0.22 Tg C/yr (Table 2).

As is the case for SVOCs, the magnitude of total IVOC emissions is uncertain. To estimate total IVOC emissions, the naphthalene emission inventory is scaled up. This assumes that the relative contributions of biomass burning vs. biofuel burning vs. anthropogenic sources to IVOC emissions are consistent with those of naphthalene. The emission rate of IVOCs in grid cell \( I, J \) \( (E_{IVOC}) \) is determined by scaling the naphthalene emission \( (E_{NAP}) \),

\[
E_{IVOC}(I, J) = \beta E_{NAP}(I, J)
\]

where, the scaling factor, \( \beta \), is estimated from,

\[
\beta = \frac{E_{NVPOA,BB+BF}}{E_{NAP,BB+BF}} \left( \frac{E_{IVOC,BB+BF}}{E_{NVPOA,BB+BF}} \right)_{R_1} \left( \frac{E_{IVOC,TOT}}{E_{IVOC,BB+BF}} \right)_{R_2} \left( \frac{1}{E_{NAP,TOT}} \right)
\]

\( E_{A,B} \) are the global emissions of species \( A \) (IVOC, NVPOA: traditional POA, or NAP: naphthalene) from source type \( B \) (BB: biomass burning, BF: biofuel burning, or TOT: total). Subscripts \( R_1 \) and \( R_2 \) are used to label the two ratios for further discussion.

The first ratio in Eq. (9), \( R_1 \), is estimated based on the Schauer et al. (2001) inventory for pine wood burning. \( E_{IVOC,BB+BF} \) is approximately as any species collected on the filter/polyurethane foam (PUF) sampling train with only a gas-phase emission reported plus the entire gas-phase unresolved complex mixture (UCM). The gas-phase UCM in the work of Schauer et al. (2001) for pine wood burning represents roughly 10% of the total non-methane organic carbon (gas+aerosol) mass emitted. The two contributions (PUF gases and UCM gases) to IVOC emissions are roughly equal. Using the organic aerosol emission rate \( (E_{NVPOA,BB+BF}) \) from the work of Schauer et al. (2001) as well, \( R_1 \) is estimated as,

\[
\frac{E_{IVOC,BB+BF}}{E_{NVPOA,BB+BF}} \approx 0.34.
\]

Since the IVOC is spatially distributed like naphthalene, \( R_2 \) can be replaced by:

\[
\frac{E_{IVOC,TOT}}{E_{IVOC,BB+BF}} = \frac{E_{NAP,TOT}}{E_{NAP,BB+BF}}
\]

and consequently, only \( E_{NAP,BB+BF} \) is needed to complete the scaling factor.

Thus, our scaling incorporates two ideas: IVOCs are spatially distributed like naphthalene and the ratio of IVOCs to traditional POA for wood burning sources is 0.34 as in the work of Schauer et al. (2001). As a result, the predicted scaling factor, \( \beta \), is 66 and yields a global IVOC emission rate of about 15 Tg C/yr (Table 2 and Fig. 1). The large value of the scaling factor, \( \beta \), means that naphthalene itself is actually a relatively small (<2%) contribution to global IVOC emissions.

The IVOC emissions are not linked to the POA emission inventory within each grid cell; rather, they are spatially distributed like naphthalene. On a global basis, the present work predicts IVOC emissions of roughly 0.5 x POA in magnitude. As a result of the separation of POA and IVOC emissions, the ratio of IVOC:traditional POA emissions is roughly 2.1 for anthropogenic sources and 0.34 for biomass and biofuel sources combined. These ratios are consistent with the discussion by Shrivastava et al. (2008) and box model studies by Grieshop et al. (2009a), indicating that diesel combustion (an anthropogenic source) may produce relatively more IVOCs than wood burning.

### 2.5.2 IVOC oxidation

IVOC aging behavior is based on the chamber studies of Chan et al. (2009) and Kautzman et al. (2010) examining the oxidation of naphthalene under high- and low-NOx conditions. Similar to light aromatic oxidation (Ng et al., 2007), naphthalene oxidation under high-NOx conditions was found to produce semivolatile SOA while oxidation under low-NOx conditions was found to produce essentially non-volatile SOA. The yield of aerosol ranged from 13 to 30% for the high-NOx oxidation and was constant at 73% for low-NOx oxidation. For naphthalene, the results indicate that the first oxidation step is rate-limiting for SOA formation. Calculations using diesel engine and wood burning emission profiles indicate that naphthalene and other PAHs are responsible for substantially more SOA than light aromatics on short timescales (about 12 h) (Chan et al., 2009), and ambient aerosol contains compounds indicative of naphthalene oxidation (Kautzman et al., 2010).

Formation of aerosol from IVOC oxidation is modeled similar to the aerosol from aromatic oxidation in the work of Henze et al. (2008). Reaction of the parent hydrocarbon with OH in the presence of oxygen results in the formation of a peroxy radical species,

\[
\text{Naphthalene} + \text{OH} \rightarrow \text{RO}_2
\]

Under high-NOx conditions, reaction of the \( \text{RO}_2 \) radical with NO likely dominates over reaction of \( \text{RO}_2 \) with \( \text{HO}_2 \) or \( \text{RO}_2 \) and thus SOA can be assumed to form from the following channel:

\[
\text{RO}_2 + \text{NO} \rightarrow \alpha_{N,1} \text{P}_{N,1} + \alpha_{N,2} \text{P}_{N,2}
\]
Table 4. Rate constants for IVOC oxidation. \( k = Ae^{b/T} \).

| Reaction          | \( A \) [cm\(^3\) molec\(^{-1}\) s\(^{-1}\)] | \( B \) [K] | \( k_{298} \) [cm\(^3\) molec\(^{-1}\) s\(^{-1}\)] |
|-------------------|-----------------------------|-----------|----------------|--------|
| NAP + OH          | 1.56 \times 10^{-11}       | 117       | 2.3 \times 10^{-11} |
| RO\(_2\) + HO\(_2\) | 1.4 \times 10^{-12}       | 700       | 1.5 \times 10^{-11} |
| RO\(_2\) + NO     | 2.6 \times 10^{-12}       | 350       | 8.5 \times 10^{-12} |

Sources: Atkinson and Arey (2003); Henze et al. (2008); Atkinson (1997)

where the two semivolatile products are described using parameters by Chan et al. (2009). Under low-NO\(_x\) conditions, the RO\(_2\) radical is expected to react predominantly with HO\(_2\) and lead to one essentially non-volatile product (Chan et al., 2009; Kautzman et al., 2010):

\[
RO_2 + HO_2 \rightarrow \alpha_{H,1}P_{H,1} \quad \text{(R4)}
\]

In practice, the non-volatile SOA product is represented in GEOS-Chem using a partitioning coefficient, \( K_{OM} \), of \( 10^4 \text{ m}^3/\text{mg} \). Values for the partitioning coefficients and mass-based stoichiometric coefficients of all the products are given in Table 1, and Table 4 contains the gas-phase rate constants.

2.6 Additional model parameters and specifications

In the current version of GEOS-Chem, the semivolatile products of parent hydrocarbon oxidation are lumped together into several gas-phase and aerosol-phase tracers (Henze and Seinfeld, 2006; Liao et al., 2007; Henze et al., 2008). Since the identity of each individual species is not preserved during transport, some artificial migration of mass between the volatilities may occur. In the present work, for the SVOC related species, separate tracers are used for each phase (gas and aerosol) as well as each volatility for a total of 4 tracers related to primary SVOC emissions and 4 tracers for SVOC oxidation products. The use of separate tracers prevents migration between the different volatilities during transport which was found to be significant at \( 4^{\circ} \times 5^{\circ} \) resolution using a lumped tracer. For IVOC oxidation, the effects of lumping were found to be less significant, but in order to obtain the best estimate of IVOC aerosol produced under high-NO\(_x\) conditions, separate tracers for each of the IVOC oxidation products are used. Each species with a saturation concentration below \( 10^3 \text{ mg} / \text{m}^3 \) in Table 1 has a separate gas and aerosol phase tracer. The IVOC has a gas-phase tracer only for a total of 15 additional tracers for the low-volatility organic aerosol simulation. A list of GEOS-Chem tracers for a standard full-chemistry SOA simulation can be found in Table 2 in the work of Liao et al. (2007).

Emitted SVOCs, aged SVOCs, and aged IVOCs in the gas and aerosol phases are subject to wet and dry deposition. The IVOC surrogate gas is not deposited. Dry deposition is represented by a resistance in series method (We- sely, 1989), with the surface resistances for aerosols following the work of Zhang et al. (2001). SVOCs that partition directly to the aerosol phase to form POA are treated as hydrophobic and are assumed insoluble. Gas-phase SVOC emissions are treated as relatively hydrophobic with an effective Henry’s law coefficient of 9.5 M/atm and a heat of dissolution, \( \Delta H/R \) of \( -4700 \text{ K} \) based on phenanthrene (Sander, 1999). Other SVOC species, such as long-chain alkanes, are more hydrophobic with Henry’s law coefficients of \( 10^{-4} \text{ M} \text{atm}^{-1} \) to 0.3 M/atm. However, SVOCs such as functionalized phenols or oxygenated species, will likely be more soluble than phenanthrene. The SVOC and IVOC oxidation products are treated like the traditional SOA species and are hydrophilic (Henry’s law coefficient of \( 10^{5} \text{ M} \text{atm}^{-1} \) for all gas-phase semivoltiles), and all SOA is scavenged with an 80% efficiency (Chung and Seinfeld, 2002). The effect of a lower Henry’s law coefficient will be examined in sensitivity studies.

The equilibrium partitioning coefficients of all species are adjusted for temperature based on the Clausius-Clapeyron equation. As in previous studies using GEOS-Chem, the enthalpy of vaporization is assumed to be 42 kJ/mol for all organic species (Chung and Seinfeld, 2002). However, a discrepancy exists in the predicted enthalpy of vaporization for semivolatile organic aerosol estimated for complex SOA systems (around 10–40 kJ/mol, Offenberg et al. (2006)) and based on theory or single component systems (around 100 kJ/mol, Epstein et al. (2010)). The enthalpy of vaporization is examined in Sect. 3.4.1.

2.7 Aerosol aging

Modeling efforts are still limited in their ability to represent aging that occurs on timescales longer than a few days, as these conditions are not readily accessed in chamber experiments (Jimenez et al., 2009). On long timescales, compounds may continue to functionalize and form more aerosol or may fragment and reduce aerosol formation. Chamber experiments tend to produce aerosol that resembles semivolatile oxygenated OA, SV-OOA, which is higher in volatility and lower in O:C than that observed in the atmosphere which tends to be dominated by low-volatility oxygenated OA, LV-OOA, with high O:C (Ng et al., 2009). Lane et al. (2008), Murphy and Pandis (2009), Jimenez et al. (2009), and Farina et al. (2010) have postulated that the semivolatile species from traditional precursors such as isoprene, terpenes, and aromatics continue to oxidize in the atmosphere in the gas phase. However, the analysis of Chhabra et al. (2010) indicates that the chamber oxidation of several parent hydrocarbons, including toluene, xylene, and naphthalene, does approach LV-OOA type aerosol with high O:C. In the work presented here, chemical reaction and aerosol formation from all parent hydrocarbons (traditional, SVOCs, and IVOCs) is limited to the behavior that is currently captured in chamber experiments.
Fig. 2. Predicted concentration of aerosol from SVOCs and IVOCs. Aerosol from SVOCs includes POA, (a and b), and SOA, (c and d), and is expressed in µg C/m$^3$. Aerosol from IVOCs is shown in panels (e and f) and is expressed in µg/m$^3$. Concentrations are shown at the surface for December-January-February (DJF) and June-July-August (JJA) for year 2000 (GEOS-4, 2°×2.5°). Note that the color scale is not linear.

3 Results and discussion

3.1 OA from SVOCs and IVOCs

Figure 2 shows the amount of organic aerosol predicted to form from direct partitioning of SVOC emissions (POA), oxidation of SVOCs, and oxidation of IVOCs. Concentrations are shown for the winter and summer at the surface. The highest POA concentrations (Fig. 2a, b) reflect biomass burning source regions, but anthropogenic source regions such as the US and East Asia also have high POA concentrations. Compared to POA, the SOA formed from SVOC oxidation is more regionally distributed (Fig. 2c, d). SOA from IVOC oxidation has a similar spatial distribution to SOA from SVOCs, but in general, concentrations are lower (Fig. 2e, f).

Compared to the traditional simulation with non-volatile POA and no IVOCs, generally less total organic aerosol is predicted at the surface in the revised simulation with semivolatile POA, primary SVOC aging in the gas phase, and SOA from IVOCs (see Fig. S1 in supplement (http://www.atmos-chem-phys.net/10/4377/2010/acp-10-4377-2010-supplement.pdf), note that both traditional and revised simulations form aerosol from traditional SOA precursors such as biogenic hydrocarbons and light aromatics). The largest decreases in total organic aerosol are over the biomass burning and isoprene source regions. Small increases of up to 0.2 µg/m$^3$ in organic aerosol are predicted at northern high latitudes in DJF (December-January-February). The largest surface level, seasonally averaged increase (0.4 µg/m$^3$) occurs during SON (September-October-November) over Eastern Russia near a
Fig. 3. Fraction of primary semivolatile material in aerosol phase as POA at the surface, seasonally averaged for year 2000. Fraction is simply the aerosol-phase SVOC concentrations (in ppb) divided by the total SVOC (gas+aerosol) concentration (in ppb). Simulations performed with GEOS-4 at 2° × 2.5°.

While the revised simulation generally predicts lower total OA concentrations near the surface, starting at roughly 4 km (model level 7) OA concentrations tend to be higher in the revised simulation (Fig. S1). Not all regional changes necessarily transition from decreases to increases at 4 km though. Concentrations over the US during JJA (June-July-August) tend to be lower than the traditional simulation predicts up until about 7 km. Biomass burning outflow (JJA Africa), South American outflow (JJA and SON), and Asian outflow regions are among the first locations other than high latitudes at which increases in total OA occur as altitude increases.

Several factors influence the change in organic aerosol concentration between the traditional and revised simulations. With semivolatile POA, concentrations will tend to decrease, as a large portion of the POA is predicted to evaporate. The SVOC emission rate is higher (27%) than the standard POA emission rate, but not high enough to compensate for POA evaporation. This shift to the gas phase is reflected in Fig. 3, which displays the fraction of primary SVOCs in the particle phase as POA in the revised simulation. During DJF over the eastern US, about 20% and as much as 27% of the primary SVOC is partitioned to the aerosol phase. (At 270 K, a typical wintertime temperature in the Northeast US, the saturation concentration of SVOC$_2$ is about 3 µg/m$^3$.) In JJA, 10 to 20% of the SVOC is in the aerosol phase. In the western US as much as 36% is in the particle phase due to high biomass burning emissions. In addition to the effect of evaporation, the OM/OC ratio for POA decreases from 2.1 in the traditional simulation to 1.4 in the revised simulation, which affects the amount of aerosol available for partitioning.

Concentrations of total OA could also increase for a number of reasons. Since SVOCs are oxidized in the gas phase, OA will tend to be shifted toward more remote or downwind areas relative to source regions. Also, changes in wet deposition (SVOCs and oxidized SVOCs are less aggressively wet deposited than hydrophilic traditional POA) allow further transport of some SVOC emissions. The introduction of IVOCs adds another source of OA. Oxidation of the IVOC through the RO$_2$+HO$_2$ route (which is more dominant than the RO$_2$+NO route in remote regions, Henze et al., 2008) produces non-volatile aerosol at a high yield. Still, even with the introduction of IVOCs, surface OA levels are lower than those in the traditional model (Fig. S1, http://www.atmos-chem-phys.net/10/4377/2010/acp-10-4377-2010-supplement.pdf).

Fig. 4. SVOC budget. Relative portions of pie indicate annual net production (32 Tg C/yr total). SVOC and O-SVOC (oxidized SVOC) are also wet and dry deposited (not shown). All numbers are for year 2000. Simulation performed at 2° × 2.5° with GEOS-4 meteorology.
Table 5. Global OA budget for traditional (non-volatile POA) and revised (semivolatile POA, SOA from SVOCs, and SOA from IVOCs) simulations. The OM/OC ratio for traditional, non-volatile POA is 2.1. The OM/OC ratio for semivolatile POA is 1.4. The OM/OC ratio for all SOA is 2.1. Simulations were performed at $2^\circ \times 2.5^\circ$ horizontal resolution with GEOS-4 meteorology.

<table>
<thead>
<tr>
<th>Tropospheric burden [Tg]</th>
<th>Net Sourcea [Tg/yr]</th>
<th>Wet Deposition [Tg/yr]</th>
<th>Dry Deposition [Tg/yr]</th>
<th>Lifetime [days]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Traditional Non-volatile POA Simulation</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Traditional POA</td>
<td>0.92</td>
<td>61</td>
<td>53</td>
<td>8.4</td>
</tr>
<tr>
<td>Traditional SOA</td>
<td>0.72</td>
<td>26</td>
<td>23</td>
<td>2.3</td>
</tr>
<tr>
<td>Total OA</td>
<td>1.64</td>
<td>87</td>
<td>76</td>
<td>11</td>
</tr>
<tr>
<td>Revised Simulation</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Semivolatile POA</td>
<td>0.03</td>
<td>0.70</td>
<td>0.22</td>
<td>0.49</td>
</tr>
<tr>
<td>SOA from SVOCs</td>
<td>0.81</td>
<td>38</td>
<td>34</td>
<td>4.4</td>
</tr>
<tr>
<td>SOA from IVOCs</td>
<td>0.09</td>
<td>5.2</td>
<td>4.6</td>
<td>0.6</td>
</tr>
<tr>
<td>Traditional SOA</td>
<td>0.71</td>
<td>23</td>
<td>21</td>
<td>2.0</td>
</tr>
<tr>
<td>Total OA</td>
<td>1.65</td>
<td>67</td>
<td>60</td>
<td>7.4</td>
</tr>
<tr>
<td>Percent Change in OA</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Traditional OA</td>
<td>−1%</td>
<td>−11%</td>
<td>−11%</td>
<td>−15%</td>
</tr>
<tr>
<td>Total OA</td>
<td>0%</td>
<td>−23%</td>
<td>−22%</td>
<td>−31%</td>
</tr>
</tbody>
</table>

a Net Source includes emission for POA in the traditional simulation.

3.2 Global budgets

Despite the fact that the revised simulation, with the new SVOC and IVOC inventories, has a larger pool of organics with the potential to form OA, global production of OA decreases 23% in the revised simulation compared to the traditional simulation. Table 5 shows the global OA budget for each simulation. In the traditional simulation, traditional SOA and non-volatile POA represent a global OA source of 87 Tg/yr. In the revised simulation, the net OA source decreases to 67 Tg/yr. The formation of traditional SOA decreases as well (although only 11%), likely as a result of a reduced partitioning medium into which the SOA may absorb. The tropospheric lifetime of OA against deposition (defined as the tropospheric burden divided by the sum of wet and dry deposition) increases in the revised simulation. In the traditional simulation, POA is assumed to be emitted as 50% hydrophobic and 50% hydrophilic and converted from hydrophobic to hydrophilic forms with an e-folding lifetime of 1.15 days. In the revised simulation, the hydrophobic nature of POA leads to a much longer POA lifetime (17 days against deposition). In addition, production of aerosol is shifted to higher altitudes where it is less subject to wet or dry deposition and thus has a longer lifetime. By coincidence, the global tropospheric burden of organic aerosol is roughly the same in the traditional and revised simulations.

As shown in Fig. 4, only about 50% of the carbon emitted as an SVOC leads to net aerosol production. The rest is wet or dry deposited in the gas phase. Of the 37 Tg C/yr emitted, only 0.5 Tg C/yr are predicted to lead to net POA formation, most of which (97%) is due to the lower volatility SVOC (SVOC$\_1$). 95% of the emitted SVOC carbon reacts in the gas phase with OH to form an oxidized SVOC (O-SVOC). The O-SVOC is fairly effectively wet and dry deposited (it is treated like traditional gas-phase SOA products with a Henry’s law coefficient of $10^3$ M/atm) and a significant fraction is lost to deposition in the gas phase. Only 18 Tg C/yr forms net SOA. Roughly 80% of the SOA from SVOCs is from the lower volatility product ($C^*=$0.2 µg/m$^3$) while 20% is from the higher volatility product ($C^*=$16 µg/m$^3$). Ultimately, the OA from SVOCs is lost to wet and dry deposition with wet deposition being dominant. The yield of OA from SVOCs is about 50% on a carbon basis (mass of carbon in aerosol/ mass of carbon emitted) and about 75% in terms of total mass of SVOC aerosol produced divided by total mass of SVOC emitted. Figure 4 also shows the relative rates of formation of aerosol from each source in the pie chart (in Tg C/yr). More than half of the global aerosol carbon is predicted to come from primary SVOC oxidation products. The second largest contribution is from traditional SOA followed by SOA from IVOCs and POA.

Figure 5 shows the emission, oxidation, and SOA formation predicted for light aromatics and IVOCs. Because of where they are emitted, most of the parent hydrocarbons react following the high-NOx RO$_2$+NO pathway. As shown by Henze et al. (2008), aromatics that react faster with OH will have a greater tendency to follow the RO$_2$+NO pathway since biofuel and fossil fuel burning emissions tend to be colocated with anthropogenic NOx sources, and the parent hydrocarbon is more likely to be oxidized in the source region. The IVOC surrogate, naphthalene, reacts faster with OH than benzene or toluene, but similar in rate to xylene. For both naphthalene and xylene, the amount of oxidation through the high-NOx pathway is about twice that through...
the low-NO\textsubscript{x} pathway. Benzene, with the slowest OH rate constant, reacts slightly more through the RO\textsubscript{2}+HO\textsubscript{2} than the RO\textsubscript{2}+NO pathway. The high-NO\textsubscript{x} IVOC oxidation pathway results in about 1 Tg/yr of SOA and the low-NO\textsubscript{x} pathway results in about 4 Tg/yr, leading to an overall yield of aerosol from IVOCs of about 30%.

### 3.3 United States organic aerosol

Figure 6 shows the predicted concentration of total OA over the United States for the winter and summer and the contributions of different OA types to that total. The contribution of light aromatics to total OA is not shown but is similar in magnitude to that from IVOCs. The highest concentrations in the winter are predicted to be located in the Southeast US as a result of a biomass burning event and high SVOC emissions. During the winter, POA is a significant contributor to total OA concentrations. POA contributions are highest in the Northeast where temperatures are lower and partitioning to the aerosol phase is favored. Anthropogenic and biofuel burning emissions also tend to be highest in the Northeast (Fig. 1). Despite significant POA contributions, SOA from SVOCs is the dominant wintertime OA component, generally contributing 50% or more to surface concentrations. In the summer, the highest OA also occurs in the Southeast, but is due primarily to biogenic SOA. The highest POA contributions in summer reflect biomass burning sources. Outside of the biomass burning locations, POA is generally lower in the summer than in the winter. Despite the significant contribution of biogenic SOA, SOA from SVOCs remains an important contributor to total OA, representing up to 50% of the OA in the Northeast and 50% or more in the Western US.

Since the winter organic aerosol is dominated by SVOC aerosol, further examination of wintertime concentrations allows us to assess the model performance while minimizing uncertainties in the biogenic aerosol parameterization as a source of discrepancy. Figure 7 shows winter (December-January-February 2000) simulated and observed aerosol OC concentrations over the US. Surface measurements of total OC from the Interagency Monitoring of Protected Visual Environments (IMPROVE) network (http://vista.cira.colostate.edu/improve/) are overlaid in circles on top of the simulated OC concentrations. Since the IMPROVE network observations are reported as mass of carbon rather than the mass of total organic aerosol, converting model values, which tend to be in total organic mass, to organic carbon mass is an additional source of potential discrepancy. Since the aerosol from SVOCs is tracked in GEOS-Chem as the mass of carbon and OM/OC ratios of 1.4 for POA and 2.1 for SOA are applied in the partitioning routines, focusing on the wintertime, where aerosol is dominated by SVOC sources with a model imposed OM/OC ratio, can reduce the effect of the OM/OC ratio which must be specified for traditional and IVOC SOA. To convert traditional and IVOC SOA to OC, an OM/OC ratio of 2.1 is used and is roughly consistent with chamber data with the notable exception of α-pinene SOA (Chhabra et al., 2010). Comparisons to the IMPROVE network by Liao et al. (2007) indicate that GEOS-Chem underpredicts annual OA levels by about 0.56 µg/m\textsuperscript{3} (mean bias) or 34% (normalized mean bias) with the bias being larger in magnitude in the western US and lower in magnitude in the eastern US. A comparison of simulated and observed OC levels for the US during DJF 2000 in this work, indicate that GEOS-Chem underestimates OC by 0.20 µgC/m\textsuperscript{3} (21%) for the 2°×2.5° simulations and 0.37 µgC/m\textsuperscript{3} (39%) for the 4°×5° simulation. The bias for the 4°×5° simulation is likely larger as a result of the coarser model resolution and the fact that a climatological meteorology has been used as opposed to the assimilated, year-specific meteorology used by 2°×2.5° simulations. Compared to the semivolatile simulation (Fig. 7 bottom), the traditional simulation (Fig. 7 top) is more consistent with the IMPROVE observations. Sensitivity tests will be used to determine the factors that allow model performance to be brought closer to observations.

### 3.4 Sensitivity tests

A number of key inputs and parameters involved in the global simulation of organic aerosol are uncertain; this fact suggests that sensitivity simulations will be useful in understanding the extent to which these uncertainties influence the predictions. The quantities examined in the sensitivity tests are listed in Table 6. For computational efficiency, the sensitivity tests are performed at 4°×5° resolution with GISS GCM meteorology. The effect of changes in meteorology and grid resolution on the global OA budget are examined in Appendix A. Although both the resolution and meteorology differ from those in the simulations discussed in the previous section, since the sensitivity tests address relative changes,
the conclusions should be robust and relatively independent of the meteorology used (a discussion of interannual variability and resolution appears in Sect. 4.3). A Traditional (non-volatile POA and traditional SOA only) simulation is performed for comparison purposes. Even though the Revised simulation better represents the current scientific understanding of low-volatility organic compounds, the predicted concentration of OC over the US is low compared to observations (see Fig. 7). Due to the sparse nature of the IMPROVE network data for year 2000 DJF and the fact that the revised simulation predicts such low OC values, sensitivity simulations are compared to the Traditional simulation as well as observations. A traditional GEOS-Chem simulation has also been extensively evaluated against observations in the work of Liao et al. (2007).

Table 6 lists the additional simulations performed. The Revised simulation with semivolatile POA, oxidation of SVOCs, IVOC aerosol, and traditional SOA is described in previous sections. Two sensitivity tests are performed to examine the effects of changes in emissions. IVOC emissions

Fig. 6. Total OA and contribution of each component to winter and summer concentrations over the US. Not shown is the contribution of aromatic SOA to total OA. Fraction is fraction of total OA. Simulations are $2^\circ \times 2.5^\circ$ with GEOS-4 meteorology.
Table 6. Sensitivity tests with GCM meteorology at 4° × 5°.

<table>
<thead>
<tr>
<th>Label</th>
<th>IVOC Emissions [Tg C/yr]</th>
<th>SVOC emissions [Tg C/yr]</th>
<th>(K_{OM}^{SOVC}) [m^3/µg]</th>
<th>(K_{OM}^{O-SVOC}) [m^3/µg]</th>
<th>Henry’s Law Coefficient^c</th>
<th>(\Delta H_{vap})/100</th>
<th>(\Delta H_{vap})/100</th>
<th>(\Delta H_{vap})/100</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Traditional(^a)</td>
<td>NA</td>
<td>29</td>
<td>large</td>
<td>NA</td>
<td>10(^5)</td>
<td>42(^d)</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Revised(^b)</td>
<td>15</td>
<td>37</td>
<td>0.0006, 0.05</td>
<td>0.06, 5.0</td>
<td>Traditional</td>
<td>Traditional</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2*IVOC</td>
<td>2 × Revised</td>
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</tr>
<tr>
<td>2*SVOC</td>
<td>Revised</td>
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<td>2 × Traditional</td>
<td>Revised</td>
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<tr>
<td>SVOC K*10</td>
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<tr>
<td>O-SVOC K*10</td>
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</tr>
</tbody>
</table>

\(^a\) Traditional simulation with non-volatile POA and SOA from traditional precursors.

\(^b\) Revised simulation uses semivolatile POA, oxidation of SVOCs, SOA from IVOCs, and SOA from traditional precursors.

\(^c\) For wet removal of gas-phase semivolatiles.

\(^d\) Implemented using a value of 41.6 kJ/mol.

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Fig. 7. Winter (December-January-February 2000) surface total OC concentration with IMPROVE Observations (circles).

Winter (DJF) Simulated Total OC Concentration with IMPROVE Observations (circles)

As mentioned previously, wintertime OA concentrations are dominated by contributions from POA and SOA from traditional precursors, IVOCs, and SVOCs. IMPROVE observations (averaged over DJF for year 2000) are overlaid in circles. An outlier value of >27 µg C/m\(^3\) for year 2000 has been removed from the IMPROVE observations. Only sites with valid data for at least half of the DJF 2000 season are shown.

3.4.1 Winter US concentrations

As mentioned previously, wintertime OA concentrations are dominated by contributions from POA and SOA from SVOCs and offer an opportunity to examine model performance while minimizing the effect of errors in the parameterization of biogenic SOA. Figures 8 and 9 show the response of wintertime surface concentrations to the sensitivity tests in Table 6 relative to the Traditional simulation. Table 7 shows the mean bias and normalized mean bias for the sensitivity simulations compared to the IMPROVE.
network. Panel (a) of Fig. 8 represents the difference in surface concentrations between the Revised and Traditional simulations and highlights the fact that the Revised simulation results in a large decrease in surface concentrations as a result of POA evaporation. By doubling the IVOC emissions, 2*IVOC, surface concentrations increase slightly, but a significant underestimate still exists as IVOCs contribute generally a small fraction of the wintertime OA. Increasing the SVOC emissions within the uncertainty of traditional POA inventories, 2*SVOC, brings surface concentrations much closer to the traditional simulation and thus observations, although a slight underestimate persists. The SVOC K*10 simulation is also effective in reducing the discrepancy between the new and traditional model formulations, since it is essentially making SVOCs more similar to traditional POA. In this simulation, roughly a factor of 6 to 10 more SVOCs with saturation concentrations of 1–2 µg/m^3 are emitted than recommended by Grieshop et al. (2009b). A comparison to the O-SVOC K*10 simulation shows that the improvement in the SVOC K*10 simulation must be primarily a result of greater POA formation, not greater formation of SOA from SVOCs. The O-SVOC K*10 indicates that reducing the volatility of the SVOC oxidation products alone is not especially effective for bringing the model closer to observations. The H-Law/100 simulation has only a small effect on winter surface concentrations, but can have a significant effect on the global budget of OA, which will be addressed in the next section. Figure 9 shows the change in OC surface concentration for DJF relative to the traditional simulation for the ΔH^2 simulation. The new enthalpy produces a result similar to that of doubling the POA inventory (2*SVOC simulation) in terms of reducing the discrepancy with the traditional simulation and observations. In conclusion, increasing the SVOC emissions, decreasing the SVOC emission volatility, or increasing the enthalpy of vaporization effectively reduce measurement/model discrepancy, although significant underestimates of the seasonal mean OC concentration persist (mean bias of −0.51 µgC/m^3 for 2*SVOC, −0.44 µgC/m^3 for SVOC K*10, and −0.50 µgC/m^3 for ΔH^2 compared to the IMPROVE observations). The reduction in volatility in SVOC K*10 may be too extreme, and the higher enthalpy of vaporization may not be appropriate for lumped organics that span a relatively large range of volatility (Donahue et al., 2006). However, the increase in SVOC emissions, 2*SVOC, lies within the uncertainty of the OA emission inventory in the work of Bond et al. (2004). Tripling the POA emission inventory to obtain SVOC emissions (not shown) results in DJF OC concentrations about 1 µg C/m^3 higher than in the traditional simulation. The possibility remains that IVOCs could be underestimated by a substantially larger amount than examined here, or that the doubling of IVOC emissions in combination with doubling the POA emission inventory may provide a good present-day simulation. Additional constraints as well as a correct treatment of SOA temperature dependence are needed to assess the optimal model parameters.

For reference, Fig. S2 (see supplement, http://www.atmos-chem-phys.net/10/4377/2010/acp-10-4377-2010-supplement.pdf) shows the effects of the first six sensitivity simulations on surface level June-July-August OC concentrations. The summertime simulations also indicate that increasing the SVOC emissions or decreasing the SVOC emission volatility are effective ways of reducing the discrepancy with the traditional simulation.
However, since the summer OA is dominated by contributions from biogenic SOA, future updates to this source of SOA will likely have significant effects on the summertime predictions.

### 3.4.2 Global budget sensitivity

Figure 10 shows the effect of the different sensitivity simulations on the global OA production rate compared to the *Traditional* simulation (all simulations use GISS meteorology at 4°×5° resolution). The *Traditional* simulation (at 4°×5°) predicts 80 Tg/yr of net OA production using an OM/OC ratio of 2.1 for POA. The net OA source for the *Traditional* simulation includes the emission rate of non-volatile POA. As mentioned previously, the *Revised* simulation leads to a decrease of the net global OA source (−26%, see Table 5 for 2°×2.5° simulation results), but an increase of the OA lifetime due to a shift of production away from the surface. Doubling the IVOC emissions (2*IVOC) increases the global OA production rate compared to the *Revised* simulation, but not compared to the *Traditional* simulation (−17% compared to *Traditional*). Doubling the POA emission inventory to obtain SVOC emissions (2*SVOC), however, leads to an OA source that exceeds the *Traditional* simulation by 12%. The SVOC K*10 and O-SVOC K*10 simulations lead to a global production rate slightly larger than in the *Traditional* simulation (+3% and +2% respectively). The decrease in effective Henry’s Law coefficient significantly increases the OA source (+12%) and lifetime due to less effective wet removal.

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**Table 7.** Mean bias (MB) in µgC/m³ and normalized mean bias (NMB) in percent (%) for the sensitivity simulations for DJF 2000 (MB = \( \frac{1}{N} \sum_{i=1}^{N} (P_i - O_i) \), NMB = 100% * \( \frac{1}{N} \sum_{i=1}^{N} (P_i - O_i)/\sum_{i=1}^{N} (O_i) \), where \( P_i \) are the model predictions and \( O_i \) are the IMPROVE observations at N locations). IMPROVE observations are shown in Fig. 7. Sensitivity simulations are outlined in Table 6. 2°×2.5° simulations use GEOS4 meteorology. 4°×5° simulations use GISS GCM meteorology.

<table>
<thead>
<tr>
<th>Simulation</th>
<th>MB [µgC/m³]</th>
<th>NMB [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>2°×2.5° Traditional</td>
<td>−0.20</td>
<td>−22</td>
</tr>
<tr>
<td>2°×2.5° Revised</td>
<td>−0.59</td>
<td>−63</td>
</tr>
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<td>4°×5° Revised</td>
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<td>−75</td>
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<tr>
<td>4°×5° 2*IVOC</td>
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<td>−73</td>
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<tr>
<td>4°×5° 2*SVOC</td>
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<td>−55</td>
</tr>
<tr>
<td>4°×5° SVOC K*10</td>
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<td>−47</td>
</tr>
<tr>
<td>4°×5° O-SVOC K*10</td>
<td>−0.60</td>
<td>−64</td>
</tr>
<tr>
<td>4°×5° H-Law/100</td>
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<td>−71</td>
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<tr>
<td>4°×5° ΔH*2</td>
<td>−0.50</td>
<td>−54</td>
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</tbody>
</table>

**Fig. 10.** Change in global OA net source (from all types of organic aerosol) for sensitivity tests. Values are relative to traditional simulation at 4°×5° using GISS meteorology. Sensitivity tests are given in Table 6.

**Fig. 11.** Fraction of SVOC emissions from modern carbon: (Biomass+Biofuel)/(Biomass+Biofuel+Anthropogenic).
Production of organic aerosol in the Revised framework at 2°×2.5° horizontal resolution with GEOS-4 is estimated to be 67 Tg/yr compared to 87 Tg/yr in the Traditional framework. The lowest global net production rate out of all simulations (at all resolutions and meteorologies) performed occurs with the Revised framework at 4°×5° horizontal resolution (GISS) and is about 60 Tg/yr. The sensitivity tests at 4°×5° horizontal resolution indicate that the global production rate could be 12% higher than that predicted in the traditional simulation if SVOC emissions are significantly underestimated or wet deposition of the gas-phase semivolatiles is less effective. The 2°*SVOC test was repeated using the 2°×2.5° resolution (with GEOS-4 meteorology, see supplement) and the increase in production over the Traditional simulation was confirmed to be roughly the same as in the coarser resolution simulations (13%). Thus, the range of estimates for global OA production examined here is approximately 60 to 100 Tg/yr.

One additional sensitivity test was performed to examine the effect of the biogenic emission inventory on the OA budget. Using the GEIA biogenic inventory (Guenther et al., 1995) instead of MEGAN (Guenther et al., 2006) results in a modest (about 7%) increase in the global OA production rate. For the 4°×5° resolution with GISS GCM III meteorology, GEIA estimates global isoprene sources to be 490 Tg/yr which is 24% higher than the MEGAN estimate. Terpene emissions increase by 14% compared to the MEGAN inventory. Isoprene SOA increases to 8 Tg/yr, and terpene SOA increases to 5 Tg/yr. However, the global OA budget (burdens, sources, and losses) remains within 10% of the estimate using the MEGAN inventory. Current global chemical transport models and chemical mechanisms are known to have significant issues simulating isoprene and OH levels under low-NOx conditions like over the Amazon (Lelieveld et al., 2008; Butler et al., 2008; Archibald et al., 2010). Improved isoprene chemistry and/or improved isoprene SOA formation parameterizations could lead to substantial changes in the global estimate of SOA from isoprene.

3.5 Modern vs. fossil carbon

The fraction of modern vs. fossil aerosol carbon can provide an additional model constraint. The major contributors to US surface OA are predicted to be biogenic hydrocarbons (mainly in summer) and SVOCs followed by IVOCs and aromatics. Slightly more than half of the global aromatic emissions are from fossil fuel sources (Henze et al., 2008). The fraction of modern carbon in POA and SOA formed from SVOCs is not separately tracked but depends on the composition of the SVOC emissions. Figure 11 shows the fraction of modern carbon in the SVOC emissions globally and for the US. Biomass and biofuel burning produce modern C while anthropogenic sources are assumed to be 100% fossil. Figure 11 indicates that aerosol from SVOCs over the US has a significant and often dominant modern component. The IVOC emissions over the US are dominated by fossil carbon (Fig. 12).

The radioisotope, 14C, can be used to distinguish between modern (contemporary) and fossil carbon in ambient samples. 14C is produced in the upper atmosphere by cosmic rays and has a half-life of about 5700 years. Fossil carbon should be depleted in 14C, while modern carbon from sources such as biogenic VOC oxidation or wood burning should be enriched. Carbon isotope measurements have been performed in various locations to determine the contribution of modern and fossil carbon to total aerosol carbon. Carbon isotope measurements in the Southern California air basin in 1987 indicate that 30–40% of the fine particulate carbon was modern (Kaplan and Gordon, 1994). Szidat (2009) found OC was usually more than 50% modern for cities in Switzerland and Sweden, with biogenic OA likely responsible for modern carbon in summer and wood burning responsible in winter. Carbon isotope analysis of individual PAHs in Sweden indicates that residential wood burning is the dominant source of PAHs in the winter (Sheesley et al., 2009). During two field campaigns (2003 and 2006), Mexico City was found to have, on average, 70% modern carbon (Marley et al., 2009).
In the analysis of aerosol at 12 US sites by Schichtel et al. (2008), total carbon was found to be about 50% modern in urban, 70–97% modern in near-urban, and 82 to 100% modern in remote areas. In short, the carbonaceous aerosol in a variety of locations exhibits significant amounts of modern carbon, with the fraction of modern carbon increasing with distance from urban centers (Schichtel et al., 2008). These observations lend support to the hypothesis that SVOCs contribute significantly to ambient OA and that scaling up SVOC emissions may be more justified than scaling up IVOC emissions. However, when scaling up the POA emission inventory to obtain SVOC emissions in this work, all sources were scaled up equally. Scaling up the POA emission inventories from all sources equally is supported by the work of Shrivastava et al. (2006) and others that indicate that diesel and wood combustion exhaust have very similar volatility profiles in the SVOC range. Thus, traditional POA inventories may represent the same fraction of SVOC emissions in each case. But, work by Shrivastava et al. (2008) and Grieshop et al. (2009a) indicate that diesel combustion or other anthropogenic sources may produce relatively more IVOCs than wood burning. The volatility at which the wood burning and anthropogenic emission volatility profiles diverge is not known (this works assumes it is about 10^4 μg/m^3 based on sources that have been examined), and the fraction of SVOCs captured by diesel exhaust and wood burning POA inventories might differ leading to the need for source specific SVOC scaling factors. SVOCs from anthropogenic sources could be underestimated to a greater extent by the traditional POA inventory than SVOCs from biomass or biofuel burning sources which means the fossil carbon SVOC emissions may be underestimated.

Using alkanes to obtain the spatial distribution of IVOC emissions would likely lead to large estimates of anthropogenic sources and small estimates of wood burning sources since wood burning is not a large source of intermediate volatility alkanes (approximately C17) (Schauer et al., 2001). Thus, the IVOC composition would be shifted even more toward fossil fuel sources than predicted by the naphthalene spatial distribution. Since ambient data indicate a large modern C component to aerosol, the alkane distribution would further diminish the expected role of IVOCs in ambient organic aerosol formation.

Modern carbon, however, should be distinguished from non-anthropogenic carbon. Biofuel combustion from activities like residential wood burning is a significant anthropogenic source of modern carbon. The high fraction of modern carbon in urban areas, especially compared to rural areas, in the winter suggests substantial contributions of wood burning to aerosol carbon in the US (Bench et al., 2007; Schichtel et al., 2008).

4 Model uncertainties

Modeling organic aerosol production from SVOCs and IVOCs requires extrapolating experimental results obtained under idealized conditions or from limited set of ambient observations to global conditions. As a result, potential discrepancies between experiments, observations, and the atmosphere must be addressed.

4.1 IVOC behavior

Naphthalene is used as a surrogate to represent SOA formation from the entire set of IVOC emissions (see Sect. 2.5). Other important classes of IVOCs include alkanes and phenol type compounds. A significant portion of the IVOCs may also be part of the gas-phase UCM. Naphthalene is expected to be more representative of phenol-type compounds than alkanes would be, but does naphthalene exhibit behavior representative of IVOCs? High-NOx oxidation of an alkane IVOC surrogate in the work of Presto et al. (2009) resulted in less volatile SOA than low-NOx oxidation, opposite of the behavior observed for naphthalene (Chan et al., 2009). However, in terms of yields, the IVOC, heptadecane, is predicted to react slightly faster with OH than naphthalene and result in a slightly lower but similar (20% vs. 26%) yield of SOA under high-NOx conditions (Chan et al., 2009).

Assuming naphthalene is a good surrogate for IVOCs, is naphthalene behavior accurately captured by the current parameterization? Chamber experiments are likely to be most representative of the atmosphere when aerosol loadings are low and oxidation times are long. Low-NOx conditions in which the peroxy radical from IVOC oxidation is expected to react with HO2 is predicted to form a non-volatile aerosol product based on chamber experiments (Chan et al., 2009). However, chamber aerosol studies typically do not access very low organic concentrations, and thus what appears non-volatile in a chamber study may actually be semivolatile under atmospheric conditions. As a result, the dominance of aromatic and IVOC aerosol in remote regions near the surface could be a model artifact. In addition, aerosol from the RO2+RO2 pathway, which should be minor, is not accounted for in our model.

Another option for treating SOA from IVOCs is the volatility basis set approach with parameters by Robinson et al. (2007) or Grieshop et al. (2009a). IVOC compounds can be lumped into a series of volatility bins and oxidized with a prescribed reduction in volatility and increase in mass. With multiple generations of oxidation and no fragmentation reactions, this approach can give a very high yield of aerosol from IVOCs which could be on the order of 200% or more (J. Jimenez, personal communication, 2010). The yield is especially high using the parameters introduced by Grieshop et al. (2009a). Modeling for the Mexico City area indicates that the Grieshop et al. (2009a) parameters with multigenerational oxidation tend to overestimate organic aerosol
downwind of the city (Hodzic et al., 2010). While the Robin-son et al. (2007) parameters may give more realistic predic-tions of the total OA, the aerosol they produce predicts too low of an O:C ratio (Hodzic et al., 2010). Currently, Robin-son et al. (2007) or Grieshop et al. (2009a) frameworks likely cannot be consistent with ambient measurements indicating that OOA is relatively non-volatile (Cappa and Jimenez, 2010) without predicting too much of an increase in mass or too low of an increase in O:C.

Simulations presented in this work obtain a lower net yield of aerosol from IVOCs of about 30% indicating a more minor role of IVOC aerosol. Both the basis set and the naphthalene-like approach used here may be similar in that near-source (i.e. high-NOx) SOA tends to be semivolatile, while remote (i.e. low-NOx) SOA tends to be relatively non-volatile. However, the reason for this trend and overall yields are very different in each case. Naphthalene can continue to serve as an IVOC surrogate until more information on IVOC emissions and/or IVOC oxidation behavior is obtained.

4.2 SVOC volatility

The parameterization for SVOC volatility uses two surro-gate compounds to represent SVOC emissions, in contrast to the volatility basis set which uses 4 or more volatility classes to represent emissions. Very low-volatility com-pounds (those with $C^* < 0.1 \mu g/m^3$) will be found in the aerosol phase under almost all atmospherically relevant conditions. The lowest volatility SVOCs emitted in this work is assumed to have a $C^*$ of 200 $\mu g/m^3$ (at 300 K), and we likely do not represent SVOCs with saturation concentrations below 1 $\mu g/m^3$. The volatility basis set approaches of Robin-son et al. (2007) and Grieshop et al. (2009b) distribute low-volatility organic compounds with $C^*$s down to 0.01 $\mu g/m^3$. Grieshop et al. (2009b) indicate that up to 5% of SVOC emis-sions may be considered non-volatile under atmospheric conditions since the $C^*$ values are below 1 $\mu g/m^3$. However, emissions of these very low-volatility compounds are not well constrained. Fits of wood-smoke data tend to diverge at organic concentrations less than about 100 $\mu g/m^3$ (Shrivastava et al., 2006). Thermodynamic data can provide additional constraints but also show significant variability with somewhat between 80 and 40% of the wood-smoke aerosol evaporating at 50°C (Grieshop et al., 2009b). Grieshop et al. (2009a) included low-volatility compounds in their model of chamber aging and found that including a 9% contribution of $C^*$ 0.1 and 0.01 degraded model performance. The two volatility components used here should roughly capture the partitioning of species with $C^*$ of 1 to 104 $\mu g/m^3$ and thus capture 95% (for wood burning) to 98% (for diesel) of the SVOC mass. If a 5% low-volatility emission were to be included in our simulation, it could account for up to 2 Tg C/yr of net primary organic aerosol formation. This could easily more than quadruple the net POA source, but would have a small effect on the global aerosol production rate, which is predicted to exceed 60 Tg/yr.

4.3 Horizontal resolution and inter-annual variability

Sensitivity tests performed indicate that doubling SVOC emissions leads to more realistic surface-level OC concentra-tions for the US in winter. This conclusion was reached using sensitivity tests for DJF in year 2000 with GISS mete-oroology at 4°×5° horizontal grid resolution. The same tests were repeated for DJF in year 2001 with GISS meteorology at 4°×5° to confirm that this conclusion is not highly sensitive to the choice of meteorological year (see Supple-ment Fig. S3, http://www.atmos-chem-phys.net/10/4377/ 2010/acp-10-4377-2010-supplement.pdf). Three of the tests in Table 6 were also performed for year 2000 DJF using GEOS-4 meteorology at 2°×2.5° horizontal grid resolu-tion. Thus, the robustness of the conclusions with re-pect to meteorological year as well as horizontal grid res-o-lation and meteorology can be determined. GISS tests for year 2000 and 2001 both indicate that doubling SVOC emis-sions brings concentrations closer to those observed, but still leaves a slight underestimate that may be larger in 2001. Using the GEOS-4 meteorology at 2°×2.5° indicates that concen-trations might actually be higher compared to the tradi-tional simulation in the northeast when SVOC emissions are doubled (Fig. S4, http://www.atmos-chem-phys.net/10/ 4377/2010/acp-10-4377-2010-supplement.pdf). In conclu-sion, years 2000 and 2001 are roughly similar in terms of the effects of the sensitivity simulations on DJF surface OC. The GEOS-4 2°×2.5° simulation confirms that doubling SVOC emissions is reasonable. However, these tests and the fact that the 2°×2.5° and 4°×5° simulations have different mean biases (Table 7) illustrate that tuning a model based on one simulation may not produce the same result for other simu-lations (like the Northeast at 2°×2.5° vs. 4°×5°). Any tun-ing of the model emissions should be based on multiple con-straints and/or a large observational data set.

5 Conclusions

In this study, we present a global estimate, using GEOSChem, of organic aerosols from primary emissions of gases and aerosols with saturation concentrations of roughly less than 10^6 $\mu g/m^3$. Sources of these compounds include biomass burning, biofuel burning, and anthropogenic activ-ities. POA, which has traditionally been considered non-volatile, is replaced by a pool of semivolatile organic com-pounds, denoted here as SVOCs, that can partition between the gas and aerosol phases and can be oxidized in the gas phase to less volatile species that partition even more effec-tively to the aerosol phase. In addition, intermediate volatili-ty organic compounds, denoted as IVOCs, which ex-ist entirely in the vapor phase, can undergo oxidation in the
gas phase to form lower volatility species that partition to the aerosol phase (based on naphthalene-like behavior). Aerosol that results from the oxidation of any of these gas-phase species is termed secondary organic aerosol (SOA). Globally, biomass burning and anthropogenic sources contribute similar amounts of IVOCs. Over the US, the dominant fraction of IVOCs arises from anthropogenic sources.

Implementation of semivolatile POA generally leads to decreases in predicted surface-level concentrations of organic aerosol due to a portion of the POA evaporating upon emission. US winter organic carbon concentrations from the IMPROVE network are used to assess the accuracy of model predictions under conditions in which the effect of uncertainties in biogenic SOA formation are likely to be at a minimum. Sensitivity tests indicate that uncertainties in the IVOC emissions, the Henry’s Law coefficient for scavenging of gas-phase semivolatilites, or the assumed 100x (or 1000x) decrease in volatility upon oxidation of the primary SVOCs are not especially influential in reducing the discrepancy between predictions and observations. However, a significant increase in SVOC emissions, a reduction of the volatility of the SVOC emissions, or an increase in the enthalpy of vaporization to 83 kJ/mol all lead to an appreciable reduction of the prediction/measurement discrepancy. The reduction in SVOC volatility examined is likely too extreme, but scaling up the SVOC emissions by a factor of 2 seems reasonable considering that traditional inventories do not necessarily capture SVOCs emitted in the gas phase. The higher enthalpy of vaporization is also supported by a recent study (Epstein et al., 2010) but may not be appropriate for lumped organics (Donahue et al., 2006).

The range of estimates for global organic aerosol production is 60–100 Tg/yr. Virtually all of this production is SOA, since POA tends to evaporate and oxidize in the gas phase after emission. The sensitivity tests examine uncertainties one parameter at a time, so the range in organic aerosol production could be larger if uncertainties in multiple parameters were to be accounted for simultaneously. If current traditional POA inventories capture only about one-half of the SVOC emissions and gas-phase semivolatilites are much less aggressively wet removed, this would lead to an estimate of global OA production that is not inconsistent with top-down calculations, such as those by (Goldstein and Galbally, 2007) who estimated SOA production ranging from 140 Tg/yr and up. Much of the increase in SOA shown here compared to previous estimates, like those by Henze et al. (2008), results from reclassification of most of the POA as SOA due to evaporation and subsequent oxidation. Also note that the present estimate of OA production is net production, so a species that partitions to the aerosol, but later evaporates, is not counted in net aerosol. The effects of chemical aging of SOA beyond that reflected in current chamber experiments is not explicitly considered.

SVOC and IVOC emissions are predicted to have different fractions of modern and fossil carbon which can provide constraints on estimates of SVOC and IVOC aerosol. In the US, SVOC emissions have a significant biofuel component which, along with biomass burning emissions, results in a significant fraction of modern C. US IVOC emissions, however, are predicted to be predominantly fossil. The high fraction of modern carbon observed in organic aerosol in the US is consistent with an important contribution of aerosol from SVOCs.

While representing POA as semivolatile is clear progress in modeling of organic aerosol, additional constraints are needed to sharpen estimates. Information in the form of improved SVOC and IVOC emission estimates or data (such as 14C fractions, O:C ratios, AMS PMF components (OOA, HOA), correlation with gas-phase tracers (Weber et al., 2007; de Gouw et al., 2005), or identification of marker compounds (Bhave et al., 2007)) that allow for the determination of sources of OA can help to constrain models.

Appendix A

Effect of meteorology and grid resolution on global OA budget

The effect of changes in meteorology and horizontal grid resolution on the global OA budget are briefly examined. GEOS-Chem with GEOS-4 and GISS GCM III meteorology at 4°×5° has been compared previously with a focus on tropospheric ozone budgets (Wu et al., 2007). Wu et al. (2007) found that an important difference between the GEOS assimilated and GISS GCM meteorology is the treatment of wet convection. The GISS model allows for non-precipitating condensed water, and in GEOS-Chem with GISS meteorology, soluble species are not scavenged in shallow convective updrafts below 700 hPa. Differences in clouds can lead to differences in oxidant levels. Boundary layer turbulence is also treated slightly differently in GEOS-Chem with GEOS meteorology versus GEOS-Chem with GISS meteorology (Wu et al., 2007). Although these differences exist between the two meteorologies, organic aerosols have been successfully simulated and compared to observations using GEOS meteorology at 2°×2.5° resolution (Park et al., 2003) and 1°×1° nested grid resolution (Park et al., 2006) and using GISS meteorology at 4°×5° resolution (Liao et al., 2007). Note that the 2°×2.5° GEOS and 4°×5° GISS simulations should be viewed as 2 different realizations of year 2000 conditions and should not be the same, but show similar responses to changes in parameters.

Table A1 shows the effect of changes in meteorology and resolution on precursor emissions, OA net production, aerosol lifetime, and global OH. Natural emissions such as lightning NOx and biogenic hydrocarbons that are parameterized based on meteorology are predicted to change significantly in some cases. The change to 4°×5° GISS meteorology results in 7% higher terpene emissions, but 11%
Table A1. Effect of model resolution and meteorology on global OA budget. 2°×2.5° simulations are performed with GEOS-4 assimilated meteorology. 4°×5° simulations were performed with GISS GCM Model 3 meteorology. Both simulations use revised framework (semivolatile POA, SOA from SVOCs, and SOA from IVOCs). To convert POA to Tg C, divide by 2.1. To convert SOA to Tg C, divide by 2.1.

<table>
<thead>
<tr>
<th>Process</th>
<th>2°×2.5°</th>
<th>4°×5°</th>
<th>Difference (%)</th>
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<tbody>
<tr>
<td>Precursor Emissions [Tg/yr]</td>
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<td>5.1</td>
<td>4.1</td>
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<td>3.3</td>
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<td>1.1</td>
<td>5.1</td>
<td>0.3</td>
<td>0.26</td>
<td>0.30</td>
<td>56%</td>
<td>31%</td>
<td>12%</td>
<td>6%</td>
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</table>

To gain further insight into the effects of changes in meteorology and resolution and why POA production may have increased at 4°×5°, the effect of the changes on a traditional non-volatile POA simulation are shown in Table A2. Hydrophobic and hydrophilic forms of POA are separated. Hydrophobic POA is similar to the semivolatile POA, in that it is emitted but can be converted to another species (hydrophilic POA for the traditional simulation or SOA in the revised simulation). In the traditional simulation, for 4°×5° resolution compared to 2°×2.5°, hydrophobic POA is more effectively wet and dry deposited as reflected by the higher deposition rates and shorter tropospheric lifetime against deposition. For production of semivolatile POA to increase at 4°×5° for the same SVOC emissions, POA must be more quickly lost to wet and dry deposition before it evaporates and reacts with OH. The aerosol deposition velocity over land is generally higher for the 4°×5° GISS study than 2°×2.5° GEOS-4 study due to a higher friction velocity over land in the GISS meteorological fields. Thus, OA production can generally decrease as a result of lower isoprene emissions, SOA production could also decrease as a result of more effective deposition of the gas-phase precursors or semivolatiles. OH levels over the Amazon are generally slightly lower in the GCAP 4°×5° simulation compared to the 2°×2.5° simulation which supports lower isoprene SOA formation as well.

Table A2. Traditional (non-volatile POA) simulation for 2°×2.5° and 4°×5° grid resolution. 2°×2.5° simulations are performed with GEOS-4 assimilated meteorology. 4°×5° simulations were performed with GISS GCM Model 3 meteorology.

<table>
<thead>
<tr>
<th>Tropospheric burden [Tg/yr]</th>
<th>Emission [Tg C/yr]</th>
<th>Wet Deposition [Tg C/yr]</th>
<th>Dry Deposition [Tg C/yr]</th>
<th>Tropospheric Life time [days]</th>
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</thead>
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<tr>
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<tr>
<td>Hydrophobic POA</td>
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<td>0.7</td>
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<td>Hydrophilic POA</td>
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<td>Total POA</td>
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<td>4</td>
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<tr>
<td>4°×5° Traditional Non-volatile POA Simulation</td>
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<tr>
<td>Hydrophobic POA</td>
<td>0.039</td>
<td>15</td>
<td>0.7</td>
<td>1.2</td>
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<tr>
<td>Hydrophilic POA</td>
<td>0.26</td>
<td>15</td>
<td>23</td>
<td>3.9</td>
</tr>
<tr>
<td>Total POA</td>
<td>0.30</td>
<td>29</td>
<td>24</td>
<td>5</td>
</tr>
</tbody>
</table>

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References


H. O. T. Pye and J. H. Seinfeld: SVOC and IVOC aerosol


