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Global budget of methanol: Constraints from atmospheric observations

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[1] We use a global three-dimensional model simulation of atmospheric methanol to examine the consistency between observed atmospheric concentrations and current understanding of sources and sinks. Global sources in the model include 128 Tg yr\(^{-1}\) from plant growth, 38 Tg yr\(^{-1}\) from atmospheric reactions of CH\(_3\)O\(_2\) with itself and other organic peroxy radicals, 23 Tg yr\(^{-1}\) from plant decay, 13 Tg yr\(^{-1}\) from biomass burning and biofuels, and 4 Tg yr\(^{-1}\) from vehicles and industry. The plant growth source is a factor of 3 higher for young than from mature leaves. The atmospheric lifetime of methanol in the model is 7 days; gas-phase oxidation by OH accounts for 63% of the global sink, dry deposition to land 26%, wet deposition 6%, uptake by the ocean 5%, and aqueous-phase oxidation in clouds less than 1%. The resulting simulation of atmospheric concentrations is generally unbiased in the Northern Hemisphere and reproduces the observed correlations of methanol with acetone, HCN, and CO in Asian outflow. Accounting for decreasing emission from leaves as they age is necessary to reproduce the observed seasonal variation of methanol concentrations at northern midlatitudes. The main model discrepancy is over the South Pacific, where simulated concentrations are a factor of 2 too low. Atmospheric production from the CH\(_3\)O\(_2\) self-reaction is the dominant model source in this region. A factor of 2 increase in this source (to 50–100 Tg yr\(^{-1}\)) would largely correct the discrepancy and appears consistent with independent constraints on CH\(_3\)O\(_2\) concentrations. Our resulting best estimate of the global source of methanol is 240 Tg yr\(^{-1}\). More observations of methanol concentrations and fluxes are needed over tropical continents. Better knowledge is needed of CH\(_3\)O\(_2\) concentrations in the remote troposphere and of the underlying organic chemistry.


1. Introduction

[2] Methanol is the second most abundant organic gas in the atmosphere after methane. It is present at typical concentrations of 1–10 ppbv in the continental boundary layer and 0.1–1 ppbv in the remote troposphere [Singh et al., 1995; Heikes et al., 2002]. It is a significant atmospheric source of formaldehyde [Riemer et al., 1998; Palmer et al., 2003a] and CO (B. N. Duncan et al., Global model study of the interannual variability and trends of carbon monoxide (1988–1997): 1. Model formulation, evaluation, and sensitivity, submitted to Journal of Geophysical Research, 2004, hereinafter referred to as Duncan et al., submitted manuscript, 2004), as well as a minor term in the carbon cycle [Heikes et al., 2002] and in the global budgets of tropospheric ozone and OH [Tie et al., 2003]. Most of the observations of atmospheric methanol concentrations consist of short-term records in surface air [Heikes et al., 2002]. Recent aircraft missions have added a new dimension to our knowledge of methanol concentrations in the global troposphere [Singh et al., 2000, 2001, 2003a, 2004; Lelieveld et al., 2002]. We use here a global 3-D chemical transport model (CTM) to examine the constraints that these aircraft observations provide on current understanding of methanol sources and sinks.

[3] Global budgets of atmospheric methanol have been presented previously by Singh et al. [2000], Galbally and Kristine [2002], Heikes et al. [2002], Tie et al. [2003], and von Kuhlmann et al. [2003a, 2003b]. They are summarized in Table 1. Plant growth is the principal source. Additional
sources include plant decay, biomass burning, atmospheric oxidation of methane and other volatile organic compounds (VOCs), vehicles, and industrial activities. Chemical loss by oxidation by OH results in an atmospheric lifetime of about 10 days. Deposition, exchange with the ocean, and heterogeneous reactions in aerosols and clouds are additional sinks but are poorly constrained.

Large discrepancies are apparent between the different budgets of Table 1. Singh et al. [2000] found a factor of three imbalance between their independently estimated sources and sinks, implying a missing sink. Heikes et al. [2002] achieved closure between sources and sinks in their budget but with large uncertainties. Their estimated plant growth source is four times larger than that of Singh et al. [2000] and is partly compensated by an assumed large deposition sink. Galbally and Kirstine [2002] enforced closure by simulating methanol in a six-box global model of the atmosphere-ocean system. Tie et al. [2003] conducted global 3-D model simulations of methanol with two estimates for the plant growth source, a high value of 312 Tg yr\(^{-1}\) from their bottom-up source inventory and a low value reduced by a factor of 3 to match that of Galbally and Kirstine [2002]. They presented order-of-magnitude comparisons to observations but did not conclude as to the implied constraints on the methanol budget. Von Kuhlmann et al. [2003b] evaluated a global 3-D model simulation of methanol with observations from two aircraft campaigns (SONEX and PEM-Tropics B). We present here a more detailed evaluation with a large ensemble of measurements from aircraft and ships (Figure 1).

2. Model Description

2.1. General Description

We conducted a 1-year simulation of atmospheric methanol for 2001 with the GEOS-CHEM CTM v4.32 (http://www-as.harvard.edu/chemistry/trop/geos) [Bey et al., 2001]. The model is driven by GEOS-3 assimilated meteorological observations (including convective mass fluxes) from the NASA Global Modeling and Assimilation Office. The GEOS-3 data have a temporal resolution of 6 hours (3 hours for mixing depths and surface quantities), a horizontal resolution of 1\(^\circ\) × 1\(^\circ\), and 48 layers in the vertical. We average them here over a 4\(^\circ\) latitude × 5\(^\circ\) longitude grid for input to GEOS-CHEM.
from 2001 but from other years as well, assuming that interannual variability is a relatively small source of error. Correlations of the methanol simulation with consistent simulations of CO, HCN, and acetone for 2001 are also presented for comparison with ship and aircraft observations. The CO and HCN simulations are as described by Li et al. (2003). The acetone simulation is as described by Jacob et al. (2002) but without an ocean source (for reasons to be discussed in section 4.2).

2.2. Sources of Methanol

2.2.1. Plant Growth

We use the plant physiology model of Galbally and Kirstine (2002) in which methanol emission scales as net primary productivity (NPP) with emission factors per unit carbon of 0.020% for grasses and 0.011% for other plants. This offers a process-based parameterization for global mapping of the methanol source from plant growth. We apply the emission factors to a monthly NPP database from the CASA 2 biosphere model with 1/17° spatial resolution (Potter et al., 1993; Randerson et al., 1997). The global NPP in that database is 17 Pg C yr⁻¹ for grasslands and 41 Pg C yr⁻¹ for other plants. The resulting methanol emission is 128 Tg yr⁻¹. Comparison to previous studies in Table 1 would suggest at least a factor of 2 uncertainty on this global source but our simulation of atmospheric observations implies a narrower range, as discussed later.

The Galbally and Kirstine (2002) relationship of methanol emission to NPP has yet to be tested with field observations, and NPP estimates are themselves subject to substantial uncertainty (Karl et al., 2004). Our initial simulations used monthly mean NPP values to distribute methanol emissions seasonally, but there resulted large model overestimates of observed methanol concentrations at northern midlatitudes in late summer and fall. Laboratory and field data indicate in fact that methanol emissions from young leaves are a factor of 2–3 higher than from mature leaves (McDonald and Fall, 1993; Nemecek-Marshall et al., 1995; Karl et al., 2003). We fit these results by scaling our monthly mean NPP-based emissions with the following parameterization adapted from A. Guenther:

\[ \alpha_i = \beta \left( 1 + 2 \max \left( \frac{L_i - L_{i-1}}{L_i}, 0 \right) \right) \]

where \( L_i \) is the local leaf area index for month \( i \), \( \alpha_i \) is the monthly scaling factor, and \( \beta \) is a normalizing factor such that \( \sum_{i=1}^{12} \alpha_i = 12 \) for each grid square. The normalization ensures consistency with the Galbally and Kirstine (2002) NPP-based algorithm on a yearly basis. Leaf area indices in the model are computed monthly as a function of ecosystem type, NPP, and global vegetation index (GVI), following the algorithm of Guenther et al. (1995) as implemented by Wang et al. (1998). The resulting methanol emissions at midlatitudes peak in spring, when they may be as much as a factor of three larger than in summer. Within a given month we distribute the methanol source evenly over the daytime hours, assuming zero emission from green plants at night (Nemecek-Marshall et al., 1995; Schade and Goldstein, 2001; Warneke et al., 2002).
2.2.2. Plant Decay

Wanneke et al. [1999] reported the abiotic emission of methanol from decaying plant matter with an emission factor of $3 \times 10^{-4}$ g per g of C oxidized. We apply this emission factor to monthly mean heterotrophic respiration rates with $1^o \times 1^o$ resolution from the CASA 2 model. The global heterotrophic respiration rate is 58 Pg C yr$^{-1}$ and the resulting methanol source is 17–29 Tg yr$^{-1}$ (best estimate 23 Tg yr$^{-1}$). Galbally and Kirstine [2002] point out that part of the methanol produced by plant decay may be consumed within the litter, but they also point out that additional biotic processes contribute to the methanol source from plant decay. Their best estimate for the total source from plant decay is 13 Tg yr$^{-1}$ (range 5–31).

2.2.3. Biomass Burning and Biofuels

We use a methanol/CO emission factor of 0.018 mol mol$^{-1}$ for combustion of different types of biomass, based on compilations of literature data [Yokelson et al., 1999; Andreea and Merlet, 2001]. Yokelson et al. [1999] find little variability in the emission factor between different types of fires (range 0.006–0.031 mol mol$^{-1}$). Singh et al. [2004] find a mean emission factor of 0.016 ± 0.002 mol mol$^{-1}$ for fire plumes from Southeast Asia sampled over the NW Pacific. Christian et al. [2003] find a mean value of 0.024 mol mol$^{-1}$ for Indonesian fuels. Holzinger et al. [2004] find a mean value of 0.038 mol mol$^{-1}$ for aged biomass burning plumes sampled over the Mediterranean Sea, which is relatively high and which they attribute to secondary production; but the fire plumes sampled by Singh et al. [2004] were also aged.

We apply the 0.018 mol mol$^{-1}$ emission factor to gridded CO emission inventories for biomass burning (climatological, monthly) [Duncan et al., 2003] and biofuels (aseasonal) [Yevich and Logan, 2003]. Global emissions in these inventories are 440 Tg CO yr$^{-1}$ for biomass burning and 161 Tg CO yr$^{-1}$ for biofuels, and the resulting methanol sources are 9 and 4 Tg yr$^{-1}$ respectively. Inverse modeling estimates of the global biomass burning source of CO constrained with surface air observations fall in the range 600–740 Tg CO yr$^{-1}$ [Bergamaschi et al., 2000; Pétron et al., 2002], about 50% higher than used here.

2.2.4. Urban

We refer to “urban” as the ensemble of methanol sources from fossil fuel combustion, other vehicular emissions, solvents, and industrial activity [Galbally and Kirstine, 2002]. We use the aseasonal gridded ($1^o \times 1^o$) EDGAR V2.0 global anthropogenic emission inventory for 1990, which gives a total urban alkane emission of 8.2 Tg yr$^{-1}$ [Olivier et al., 1994], and assume that methanol accounts for half of this total or 4.1 Tg yr$^{-1}$. Goldan et al. [1995] reported a concentration ratio of methanol to nitrogen oxides (NOx) of 0.17 mol mol$^{-1}$ in urban air in Colorado in winter; scaling of this source to a global fossil fuel combustion NOx source of 23 Tg N yr$^{-1}$ would yield an anthropogenic source of methanol of 9.1 Tg yr$^{-1}$. Aircraft measurements by J. deGouw (unpublished) indicate methanol/CO enhancement ratios of 0.050 mol mol$^{-1}$ in Denver but 0.011–0.014 mol mol$^{-1}$ in other U.S. cities. An emission ratio of 0.013 mol mol$^{-1}$, combined with a global fossil fuel source of CO of 480 Tg yr$^{-1}$ (Duncan et al., submitted manuscript, 2004), would imply a global methanol source of 7.1 Tg yr$^{-1}$. Holzinger et al. [2001] report a methanol:benzene enhancement ratio of 0.8 mol mol$^{-1}$ for urban air in Innsbruck, Austria. The EDGAR V2.0 inventory gives a global benzene emission of 1.2 Tg yr$^{-1}$ from vehicles and industry, which would imply a methanol source of only 0.4 Tg yr$^{-1}$. The urban source of methanol is thus highly uncertain but is clearly small on a global scale.

2.2.5. Atmospheric Production

Tyndall et al. [1990; Tyndall et al., 2001]:

\[(R1)\] \[\text{CH}_3\text{O}_2 + \text{CH}_3\text{O}_2 \rightarrow \text{CH}_3\text{OH} + \text{CH}_2\text{O} + \text{O}_2\]

\[(R2)\] \[\text{CH}_3\text{O}_2 + \text{RO}_2 \rightarrow \text{CH}_3\text{OH} + \text{R}^0\text{CHO} + \text{O}_2\]

Alternate branches for these reactions, not producing methanol, are:

\[(R1')\] \[\text{CH}_3\text{O}_2 + \text{CH}_3\text{O}_2 \rightarrow \text{CH}_3\text{O} + \text{CH}_3\text{O} + \text{O}_2\]

\[(R2')\] \[\text{CH}_3\text{O}_2 + \text{RO}_2 \rightarrow \text{CH}_3\text{O} + \text{RO} + \text{O}_2\]

The \text{CH}_3\text{O}_2 and \text{RO}_2 radicals are produced in the atmosphere by oxidation of VOCs. The sum of reactions (R1), (R1'), (R2), and (R2') typically accounts for less than 10% of the \text{CH}_3\text{O}_2 sink in current chemical mechanisms. The dominant atmospheric sinks are the reactions with \text{HO}_2 and \text{NO}, which do not produce methanol:

\[(R3)\] \[\text{CH}_3\text{O}_2 + \text{HO}_2 \rightarrow \text{CH}_3\text{OH} + \text{O}_2\]

\[(R4)\] \[\text{CH}_3\text{O}_2 + \text{NO} \rightarrow \text{CH}_3\text{O} + \text{NO}_2\]

We calculate the atmospheric source of methanol from (R1) and (R2) with a GEOS-CHEM simulation of tropospheric ozone-NOx-VOC chemistry [Fiore et al., 2003]. Primary VOCs in that simulation include methane, ethane, propane, higher alkanes, \text{C}_2 alkenes, isoprene, acetone, and methanol. The simulation uses recommended data from Tyndall et al. [2001] for the kinetics and methanol yields of (R1), and also for (R3) involving the acetonylperoxy radical (\text{CH}_3\text{C}(\text{O})\text{CH}_2\text{O}_2) produced by oxidation of acetone. The methanol yields at 298 K for these two reactions are 0.63 and 0.5, respectively. We assume a 0.5 yield for all other \text{CH}_3\text{O}_2 + \text{RO}_2 reactions, following Madronich and Calvert [1990]. The resulting global source of methanol is 38 Tg yr$^{-1}$.

Reaction (R1) contributes 85% of that global total. The remaining 15%, contributed by (R2), mainly involves \text{RO}_2 radicals produced from biogenic isoprene and is largely confined to the continental boundary layer, where it is much smaller than the primary emission from plant growth. In the remote atmosphere, where the atmospheric production of methanol is of most interest, the \text{CH}_3\text{O}_2 radicals driving (R1) originate mainly from the oxidation of methane.

Previous literature estimates of the atmospheric source of methanol from (R1) and (R2), similarly obtained with global tropospheric chemistry models, are in the range...
18–31 Tg yr$^{-1}$ (Table 1), lower than our estimate. These include 30 Tg yr$^{-1}$ from Heikes et al. [2002] computed with an earlier version of GEOS-CHEM. Discrepancies between estimates reflect differences in the abundances of NO and HO$_2$, the CH$_2$O$_2$ reaction rate constants, the yield of methanol from (R1) and (R2), and the importance of (R2). Uncertainties in the (R1) rate constant and in the corresponding methanol yield are each about 30% at 298K [Tyn dall et al., 2001] and higher at colder temperatures. As we will see, observations from the PEM-Tropics B aircraft mission over the South Pacific suggest an even larger atmospheric source of methanol than is used here.

2.3. Sinks of Methanol

2.3.1. Gas-Phase Oxidation by OH

[17] We use the rate constant $k = 3.6 \times 10^{-12} \exp[-415/T]$ cm$^3$ molecule$^{-1}$ s$^{-1}$ recommended by Jimenez et al. [2003] with an uncertainty of 20%. We apply this rate constant to a 3-D archive of monthly mean OH concentrations from the Fiore et al. [2003] GEOS-CHEM simulation of tropospheric ozone-NO$_x$-VOC chemistry. The lifetime of methylchloroform against tropospheric oxidation by OH (proxy for the global mean tropospheric OH concentration) is 5.6 years in that simulation [Martin et al., 2003]. This is within the range constrained by the methylchloroform observations, which imply a 25% uncertainty in global mean OH concentrations [Intergovernmental Panel on Climate Change (IPCC), 2001]. Stratospheric loss of methanol is computed using OH concentrations archived from a global 2-D stratospheric chemistry model [Schneider et al., 2000] and amounts to only 2% of the loss in the troposphere. Our computed lifetime of methanol against oxidation by OH is 11 days. Addition of errors in quadrature implies an uncertainty of 30% on this value.

2.3.2. Aqueous-Phase Oxidation by OH(aq)

[18] Methanol dissolves in aqueous aerosols (Henry’s law constant $H = 1.6 \times 10^{-5}$ exp [4900/T] M atm$^{-1}$) and can then be oxidized by OH(aq) ($k = 1.0 \times 10^{8}$ exp $[-590/T]$ M$^{-1}$ s$^{-1}$; Elliot and McCracken [1989]). The corresponding first-order loss rate constant (s$^{-1}$) for atmospheric methanol is $k = HLRT/[OH(aq)]$ where $L$ is the dimensionless liquid water content, $R$ is the gas constant, and [OH(aq)] is the OH(aq) concentration in units of M (mole per liter of water). The reaction takes place mainly in clouds, where $L$ is larger than ours. Heikes et al. [2002] and Galbally and Kirstine [2002] found a larger role for in-cloud oxidation by OH(aq) in their global methanol budgets, amounting to 5–10% of the gas-phase loss (Table 1). Their assumed OH(aq) concentrations and aqueous-phase CH$_3$OH + OH rate constants are higher than ours.

[19] A few studies have raised the possibility of a missing heterogeneous sink for methanol in aerosols or clouds. Singh et al. [2000] suggested that such a sink might explain their observed decrease of methanol concentrations from the middle to upper troposphere at northern midlatitudes (SONEX aircraft mission). Jaeglé et al. [2000] found that unexpectedly high formaldehyde concentrations measured in the upper troposphere during SONEX correlated with methanol, and speculated that reactive uptake of methanol in cirrus clouds with a reaction probability $\gamma = 0.01$ could provide an explanation. Yokelson et al. [2003] observed the depletion of methanol in a cloud polluted by biomass burning smoke, and Tabazadeh et al. [2004] proposed that surface reactions of methanol on cloud droplets could be responsible. However, statistical comparison of in-cloud versus clear-sky methanol concentrations in the large data set from the TRACE-P aircraft mission indicates no significant differences at any altitude [Singh et al., 2004]. Upper tropospheric observations from the SONEX mission indicate no depletion in air processed by deep convection or cirrus clouds [Jaeglé et al., 2000]. Laboratory studies indicate no significant reactive uptake of methanol by sulfuric acid aerosols [Iraci et al., 2002], ice [Hudson et al., 2002; Winkler et al., 2002], or mineral oxide surfaces [Carlos-Cuellar et al., 2003]. Kane and Leu [2001] report fast reaction of methanol with sulfuric acid in concentrated solutions but Iraci et al. [2002] attribute this result to an experimental artifact. At present, the weight of evidence does not support a fast heterogeneous sink for methanol.

2.3.3. Deposition to Land

[20] Microbial and foliar uptake of methanol by vegetation and soils is difficult to separate from the plant growth and decay sources, either observationally or for model purposes. Investigators have placed soil and leaf litter [Schade and Goldstein, 2001] and foliage [Nemecek-Marshall et al., 1995] in enclosures and observed a net emission of methanol. However, these studies used enclosures that were flushed with methanol-free air. Recent measurements by A. Guenther (unpublished) indicate that there can be a net uptake of methanol when enclosures are flushed with air containing ambient levels of methanol.

[21] Measurement of the diurnal cycle of methanol concentrations at continental surface sites provides some separation of microbial uptake from plant growth emission, which is restricted to daytime. Kesselmeier et al. [2002] observed no significant diurnal variation at a site in the Amazon forest, but other observations at sites in the tropics and northern midlatitudes indicate decreases over the course of the day ranging from about 30% [Goldan et al., 1995] to several-fold [Riemer et al., 1998; Holzinger et al., 2001; Karl et al., 2004], suggesting surface uptake. A 30% decrease over the course of an 8-hour summer night, and for a typical nighttime mixing depth of 100 m, would imply a methanol deposition velocity of 0.12 cm s$^{-1}$. Karl et al. [2004] measured methanol deposition to a tropical forest ecosystem by using a combination of eddy covariance and vertical gradients, and found a mean methanol deposition
velocity of $0.27 \pm 0.14 \text{ cm s}^{-1}$. On the basis of the above information, and acknowledging the large uncertainty, we assume here a constant dry deposition velocity of $0.2 \text{ cm s}^{-1}$ to land. We assume that this deposition velocity also applies to ice-covered surfaces since Boudries et al. [2002] found that Arctic snow is a net sink for methanol. The resulting atmospheric lifetime of methanol against dry deposition is 26 days. Including a methanol sink from dry deposition improves significantly the ability of the model to reproduce observations at northern midlatitudes in fall and winter.

### 2.3.4. Wet Deposition

We simulate wet deposition of methanol with the GEOS-CHEM wet deposition scheme described by Liu et al. [2001]. This scheme accounts for scavenging of water-soluble species by convective updrafts, convective anvils, and large-scale precipitation. The scavenging coefficients used by Liu et al. [2001] are scaled here to the dissolved fraction of methanol inferred from the local liquid water content and Henry’s law constant. Scavenging is assumed to take place in warm clouds only ($T > 268 \text{ K}$). The retention efficiency of methanol upon cloud droplet freezing is taken to be 0.02, by analogy with previous assumptions for CH$_3$OOH and CH$_2$O [Mari et al., 2000], so that scavenging is inefficient in mixed liquid-ice clouds where the precipitation involves riming. Laboratory studies indicate that methanol coverage of ice surfaces is low, less than $10^{-4}$ of a monolayer at equilibrium [Hudson et al., 2002; Winkler et al., 2002], supporting the assumption of a low retention efficiency. The resulting lifetime of methanol against wet deposition is 120 days. With such a long lifetime, wet deposition does not significantly affect the vertical profile of methanol in the troposphere [Crutzen and Lawrence, 2000].

### 2.3.5. Uptake by the Ocean

Simple solubility considerations imply that the ocean mixed layer must be a large reservoir for methanol [Galbally and Kirstine, 2002; Singh et al., 2003a]. Heikes et al. [2002] point out that methanol is both produced and consumed in the ocean. Singh et al. [2003a] observed a gradient of increasing methanol concentrations from the marine boundary layer to the free troposphere over the remote North Pacific during TRACE-P, implying an ocean sink with a mean deposition velocity of 0.08 cm s$^{-1}$. Measurements taken at the Mace Head coastal site in Ireland also show evidence of methanol uptake by the ocean, with a similar deposition velocity [Carpenter et al., 2004]. Our GEOS-CHEM results presented in the work of Singh et al. [2003a] show that an ocean saturation ratio of 90% for methanol, combined with a standard two-layer parameterization of ocean-atmosphere exchange, gives a good simulation of the observed TRACE-P vertical gradients. In the absence of better information we assume that the 90% saturation ratio holds globally. The resulting lifetime of methanol against net uptake by the ocean is 130 days.

### 3. Global Model Budget and Atmospheric Distribution of Methanol

Table 1 summarizes our global model budget of methanol. The global source of 206 Tg yr$^{-1}$ includes contributions from plant growth (62%), atmospheric oxidation of VOCs (18%), plant decay (11%), biomass burning and biofuels (6%), and vehicles and industrial activities (2%). The global sink balancing that source is dominated by gas-phase oxidation by OH (63%) with minor contributions from dry deposition to land (26%), wet deposition (6%), net uptake by the ocean (5%), and aqueous-phase oxidation in clouds (<1%). The resulting atmospheric lifetime of methanol is 7 days and the global atmospheric burden is 4.0 Tg.

A preliminary version of our model budget was presented by Singh et al. [2004] in a study comparing GEOS-CHEM results to their TRACE-P aircraft observations (reference to our budget is given there as B. D. Field et al., manuscript in preparation, 2003). That preliminary version did not include deposition of methanol to land or the dependence of methanol emissions on leaf age.

Our global source of methanol is in the range of previous literature cited in Table 1 (122–350 Tg yr$^{-1}$). Differences reflect principally the magnitude of the plant growth source. Our value for that source (128 Tg yr$^{-1}$) is close to that of Galbally and Kirstine [2002] (100 Tg yr$^{-1}$), as would be expected since we followed their algorithm. Our atmospheric source of methanol from CH$_3$O$_2$ reactions (38 Tg yr$^{-1}$) is larger than previous estimates (18–31 Tg yr$^{-1}$).

Gas-phase oxidation by OH is a major methanol sink in all the inventories of Table 1. However, the principal methanol sink in the Heikes et al. [2002] budget is dry deposition to land and oceans, with an assumed deposition velocity of 0.4 cm s$^{-1}$. As mentioned above, the TRACE-P aircraft data of Singh et al. [2003a] imply a much lower deposition velocity to the ocean. Fast deposition to land in the Heikes et al. [2002] budget (90 Tg yr$^{-1}$) partly compensates for their high terrestrial biogenic source (280 Tg yr$^{-1}$). The relative contributions to the methanol sink from gas-phase oxidation by OH, dry deposition, and wet deposition in our model are essentially the same as in the previous global 3-D model study by von Kuhlmann et al. [2003a] (63%, 30%, and 7% respectively). That study did not report separate deposition terms to land and ocean.

Figure 2 shows the mean concentrations of methanol simulated by the model in surface air and at 500 hPa, for January and July. Surface air concentrations over land exceed 5–10 ppbv in the tropics and at northern midlatitudes in summer. The particularly high concentrations over Siberia in July are due to the long continental fetch and the late emergence of leaves. Surface air concentrations at northern midlatitudes in winter drop to 0.5–2 ppbv. Concentrations over land decrease typically by a factor of 2–5 from the surface to 500 hPa, reflecting the surface source and the 7-day lifetime. There is by contrast little vertical gradient over the oceans. Concentrations over the northern hemisphere oceans (0.5–1 ppbv) show little seasonal variation, because faster chemical loss in summer balances the effect of the larger continental source. Concentrations over the southern hemisphere oceans are higher in winter (0.5–1 ppbv) than in summer (0.2–0.5 ppbv) since the continental source in the southern hemisphere is mainly in the tropics and has little seasonal variation. Concentrations are lowest (below 0.2 ppbv) in surface air over Antarctica in summer.

### 4. Evaluation With Observations

We examine in this section how the above estimates of methanol sources and sinks, when implemented in a
global CTM, succeed in reproducing the observed atmospheric concentrations of methanol. Our primary focus is on observations from aircraft and ships, which integrate source information over large regions. Observations from surface continental sites, reviewed by Heikes et al. [2002], are few and show high spatial variability that we cannot expect to reproduce given our crude model representation of sources. We use these surface observations mainly to examine large-scale features, including seasonal variations and tropical concentrations, that are of particular interest for testing the model.

4.1. Surface Concentrations

[29] The review of Heikes et al. [2002] gives representative surface air concentrations of 20 (range 0.03–47) ppbv in urban air, 10 (1–37) ppbv over forests, 6 (4–9) ppbv over grasslands, 2 (1–4) ppbv for continental background, and 0.9 (0.3–1.4) ppbv over the northern hemispheric oceans. Our model values are roughly consistent with these ranges, as shown in Figure 2. Urban air is not resolved on the scale of the model.

[30] The Heikes et al. [2002] compilation includes no surface data for tropical land ecosystems, where the model predicts high concentrations year-round. Measurements by Kesselmeier et al. [2002] in the Rondônia tropical forest of Brazil (10S, 63W) for a 7-day period in October 1999 (end of dry season) indicate a range of 1 to 6 ppbv. Our mean simulated concentration for that site and month is much higher, 10 ppbv, with dominant contributions from plant growth (6 ppbv), plant decay (2 ppbv), and biomass burning (2 ppbv). Measurements by Karl et al. [2004] at a tropical forest site in Costa Rica (10N, 84W) for a 20-day period in April–May 2003 (peak of dry season) indicate a mean concentration of 2.2 ppbv. Our mean simulated concentration for that site and 2-month period is 2.1 ppbv, in agreement with observations, with a major contribution from plant growth (1.2 ppbv) and minor contributions of about 0.3 ppbv each from plant decay, biomass burning, and atmospheric production. The lower model value in Costa Rica than in Rondônia reflects the shorter continental fetch. The model shows little seasonal variation at the Rondônia site (monthly means range from 8 to 10 ppbv) but more at the Costa Rica site (1.2 to 2.7 ppbv). At the latter site, the seasonal maximum is in the wet season (June–October) when the plant growth source is high, and the seasonal minimum is in the early dry season (January–February) before biomass burning. Karl et al. [2004] also made methanol flux measurements at the Costa Rica site, which indicate a value 50% higher than the parameterization of Galbally and Kristine [2002] when normalized to the estimated tropical forest NPP. On the other hand, the data from Kesselmeier et al. [2002] would suggest that the NPP-based parameterization of Galbally and Kristine [2002] is too high for the Amazon forest. More observations of
methanol concentrations and fluxes are clearly needed for tropical land ecosystems.

To our knowledge, the only methanol observations in the literature extending over a full yearly cycle are those of Holzinger et al. [2001] taken in the outskirts of Innsbruck, Austria (47N, 11E). These authors used concurrent measurements of benzene (emitted by vehicles) to separate the local urban from the regional nonurban components of methanol. Figure 3 shows the observed ranges for the nonurban component. There is a spring-summer maximum and fall-winter minimum, reflecting biogenic emission. Concentrations drop by more than a factor of 2 from June to September. Also shown in Figure 3 are the simulated methanol concentrations and the contributions from the individual model sources. Plant growth is the main model source except in winter, when plant decay and urban sources become relatively important. The model reproduces the seasonal variation in the observations, including the rise from winter to spring and the decrease from spring to fall. The latter reflects in the model the weaker emission from mature leaves.

Ship measurements from Wisthaler et al. [2002] over the Indian Ocean during INDOEX 1999 (March 1999) provide another important data set for methanol in surface air. The cruise track extended from 20N to 12 S, starting from the west coast of India and extending south of the intertropical convergence zone (Figure 1). Observed methanol concentrations exceed 1 ppbv in Indian outflow and drop to 0.5–0.6 ppbv in southern hemispheric air. Wisthaler et al. [2002] reported a strong correlation of methanol with CO, which we compare in Figure 4 to our mean monthly model results for 2001 sampled along the cruise track. The agreement is remarkably good. The model captures the

![Figure 3](image1.png)

**Figure 3.** Seasonal variation of methanol concentrations at Innsbruck, Austria. The ranges of observations reported by Holzinger et al. [2001] for different times of year in 1996–1997 are shown as vertical lines with symbols. These observations are for the non-urban component of methanol, after subtraction of the urban component based on correlation with benzene. Model results are shown as the solid line, with additional lines identifying contributions from individual sources in the model: plant growth (short dashes), plant decay (dots), urban (thin solid), biomass burning and biofuels (long dashes), and atmospheric production (dash-dot).

![Figure 4](image2.png)

**Figure 4.** Methanol-CO correlations along the INDOEX 1999 cruise track over the Indian Ocean (Figure 1). Observations from Wisthaler et al. [2002] are shown as small dots, with linear regression as solid line. Monthly mean GEOS-CHEM results along the cruise track are shown as large open circles, with linear regression as dashed line. Coefficients of determination ($r^2$) and slopes of the linear regressions ($S$) are shown inset.
clean 0.5–0.6 ppbv background and the observed relationship with CO, although it does not capture the population of observations with relatively high CO but low methanol (hence the stronger methanol-CO correlation in the model, $r^2 = 0.76$ versus $r^2 = 0.47$). Over India in the model the methanol is mostly from plant growth, while CO is mostly from combustion; the methanol-CO correlation in Indian outflow reflects varying degrees of dilution with the marine background rather than a commonality of methanol and CO source processes.

[33] Additional unpublished methanol measurements have been made by A. Wisthaler and A. Hansel on a ship cruise in the Arctic in July August 2001 (AOE-2001), and at Zugspitze in southern Germany (47N, 11E, 2650 m altitude) for four months in October 2002 to January 2003. These are shown in Figure 5 together with the corresponding model results. The model reproduces the low Arctic cruise observations north of 85N (0.30–0.35 ppbv), reflecting deposition to the ocean, but not the even lower concentrations (0.1–0.2 ppbv) observed further south; these low concentrations are from only a few measurements and suggest a strong local ocean sink. Observations at Zugspitze show a decrease from 0.7 ppbv in October to 0.3 ppbv in January. The model also shows a decrease, mostly from the deposition sink, but not as strong as observed. This would suggest a nonphotochemical sink of methanol missing from the model. However, the TOPSE aircraft observations over the North American Arctic in winter, to be discussed in section 4.3, show higher concentrations than at Zugspitze and do not suggest such a missing sink.

4.2. Asian Outflow (TRACE-P Mission)

[34] The TRACE-P aircraft mission in March–April 2001 provided extensive data for methanol and other species in Asian outflow over the North Pacific [Jacob et al., 2003]. The outflow included a major contribution from seasonal biomass burning in Southeast Asia [Heald et al., 2003a]. Flight tracks are shown in Figure 1. Our model simulation is for the same meteorological year as TRACE-P. We sample the model along the flight tracks and for the flight days for comparison to observations. GEOS-CHEM simulations for the TRACE-P period have been evaluated previously with observations for a number of species including in particular CO [Kiley et al., 2003; Palmer et al., 2003b] and HCN [Li et al., 2003], which we discuss below in the context of their correlations with methanol. The model gives a good simulation of Asian outflow pathways with no evident transport bias [Liu et al., 2003; I. Bey et al., Characterization of transport errors in chemical forecasts from a global tropospheric chemical transport model, submitted to Journal of Geophysical Research, 2005]. Simulation of the CO$_2$ observations indicates a 45% underestimate of the net biospheric carbon emission for China in the CASA 2 model during the TRACE-P period [Sunderalingam et al., 2004]. However, as we will see below, there is no evident bias in our methanol simulation.

4.2.1. Vertical Profiles

[35] Figure 6 compares the mean simulated and observed vertical profiles of methanol concentrations for the four TRACE-P quadrants separated at 30N, 150E. The four quadrants contrast fresh Asian outflow west of 150E with more background North Pacific air to the east.

[36] The observed methanol concentrations in the NW quadrant decrease from 1.5–2 ppbv in the boundary layer to 0.9–1.3 ppbv in the free troposphere, rise again to a maximum in excess of 2 ppbv in the upper troposphere, and then decline to 0.1 ppbv above the tropopause at 12 km.
The model reproduces these features although it is up to a factor 2 too low in the middle troposphere. Stratospheric concentrations cannot be compared because of excessive stratosphere-troposphere exchange in the GEOS assimilated meteorological data used to drive the model [Liu et al., 2001; Tan et al., 2004]. The plant growth source in the model contributes a relatively featureless background of 0.3 ppbv throughout the troposphere in this NW quadrant. The boundary layer enhancement reflects a mix of Chinese emissions from plant decay, biofuels, and urban sources. The upper tropospheric enhancement is due to outflow from deep convection in Southeast Asia and is contributed mainly by the plant growth and biomass burning components.

The observed mean profile for the SW quadrant in Figure 6 shows greater influence from Southeast Asia than the NW quadrant. Concentrations decrease gradually with altitude, from 1.5 ppbv in the boundary layer to 0.5–1 ppbv in the middle and upper troposphere but with a convective enhancement apparent at 8–9 km. The model reproduces this structure with no evident bias.

Observations in the NE quadrant show a marked increase with altitude, in contrast to the western quadrants. Asian influence is principally in the middle and upper troposphere. The model underestimates this Asian influence, by about the same factor as for the free tropospheric Asian outflow in the NW quadrant. The relatively low methanol concentrations in the lower troposphere, both in the model and in the observations, reflect gas-phase oxidation and ocean uptake of methanol as the Asian air masses subside [Singh et al., 2003a].

Asian influence during TRACE-P was weakest in the SE quadrant, which lies south of the dominant outflow track. The observed layer of high methanol concentrations at 2–4 km altitude is from one single Hawaii-Guam flight where the aircraft sampled repeatedly an Asian outflow plume that had traveled southward and subsided [Crawford et al., 2004]. High CO concentrations (200 ppbv) were observed in that layer. Previous analysis of the GEOSCHEM CO simulation shows that this layer is displaced upward and diluted in the model relative to the observations [Heald et al., 2003b]. A similar bias is found for methanol, as the model enhancement is at 4–5 km altitude and peaks at only 0.7 ppbv. The model is also low in the marine boundary layer (0.4 ppbv, versus 0.6 ppbv in the observations); atmospheric production is the most important model source there and is probably too weak, as discussed later in the context of the PEM-Tropics B observations over the South Pacific.

Singh et al. [2004] previously compared results from our preliminary model version (not including dry deposition to land or the decrease in the plant growth source as leaves age) to their mean observed background vertical profile of methanol concentrations for the ensemble of the mission. This showed agreement within 10% up to 6 km altitude, but a growing model overestimate at higher altitudes (up to a factor of 3 at 11 km) that they speculated could reflect heterogeneous chemical loss not captured by the model. In
the Singh et al. [2004] comparison, observed methanol concentrations decrease from 0.8 ppbv at 6 km to 0.35 ppbv at 11 km (filtered against stratospheric influence); whereas model values increase from 0.9 ppbv at 6 km to 1.1 ppbv at 11 km. Our comparisons in Figure 6 do not show such a model bias in the upper troposphere, but instead considerable regional difference in simulated and observed vertical profiles for the different quadrants, as discussed above. The decreasing trend from 6 to 11 km in the observations is found only for the southern quadrants. In addition, the model values presented here are lower than in the preliminary simulation reported by Singh et al. [2004] due to our subsequent inclusion of a land deposition sink in the model. This affects in particular the simulation of upper tropospheric concentrations, which include a major contribution from convective outflow of tropical continental air.

4.2.2. Correlations With Other Species

Further insight can be gained by examining the observed correlations of methanol with other species. In the ensemble of TRACE-P observations we find that methanol correlates most strongly with acetone ($r^2 = 0.83$, slope = 0.38 mol mol$^{-1}$), HCN ($r^2 = 0.78$, slope = 0.12 mol mol$^{-1}$), and CO ($r^2 = 0.63$, slope = 64 mol mol$^{-1}$). These correlations are shown in Figure 7. The slopes are given with methanol as denominator. We generated corresponding correlations from the model results sampled along the flight tracks. These are also shown in Figure 7. The CO and HCN simulations are as described in the work of Li et al. [2003]. The acetone simulation presented here is that of Jacob et al. [2002] applied to the TRACE-P period, but without an ocean source since the TRACE-P observations indicate that the ocean was in fact a net sink for acetone [Singh et al., 2003a]. We find that an ocean source for acetone in the model would destroy the correlation with methanol since the ocean is a sink for methanol.

The model reproduces the correlation between methanol and acetone found in the observations ($r^2 = 0.49$, slope = 0.48 mol mol$^{-1}$ in the model). The biogenic acetone source in the model is scaled to isoprene emission [Jacob et al., 2002], while that of methanol is scaled to NPP. The stronger correlation in the observations suggests that the sources of acetone and methanol are governed by more similar processes than is assumed in the model. Measurements at rural U.S. sites in summer have previously shown a strong correlation between acetone and methanol with an acetone/methanol slope of 0.21–0.27 mol mol$^{-1}$ [Goldan et al., 1995; Riemer et al., 1998]. The higher slope observed here is due to anthropogenic sources of acetone.

Figure 7. Correlations of methanol with CO, acetone, and HCN concentrations in the ensemble of TRACE-P data. Observations (top panels) are compared to model results (bottom panels). Coefficients of determination ($r^2$), regression lines, and corresponding slopes ($S$) are indicated.
The observed correlation between methanol and HCN is well simulated by the model ($r^2 = 0.43$, slope = 0.10 mol mol$^{-1}$/C$0$). The dominant source of HCN variability in the TRACE-P data is biomass burning in Southeast Asia [Li et al., 2003; Singh et al., 2003b]. The observed HCN/CO molar emission ratio from biomass burning varies in the literature over a large range from 0.03% to 1.1%, with a best fit for the TRACE-P conditions of 0.27% [Li et al., 2003]. Combined with our estimated methanol/CO molar emission ratio from biomass burning of 1.8% mol mol$^{-1}$/C$0$ this yields an HCN/methanol emission ratio of 0.15 mol mol$^{-1}$/C$0$. The weaker slope in the model and in the observations likely reflects biogenic methanol emissions in Southeast Asia that are collocated with the biomass burning emissions.

Good agreement between model and observations is also found in the correlation of methanol with CO ($r^2 = 0.44$, slope = 69 mol mol$^{-1}$/C$0$ in the model). CO here serves as a general tracer of Asian outflow and the correlation mainly provides support for the overall magnitude of methanol export from the Asian continent. Although the model underestimates the TRACE-P methanol observations in the middle troposphere (Figure 6), the methanol-CO correlation is principally driven by strong outflow events in the boundary layer [Liu et al., 2003].

### 4.3. Other Aircraft Observations

We now compare model results to methanol observations from other aircraft missions including TOPSE at North American high latitudes in February–May 2000 (D.R. Blake, unpublished data), SONEX over the North Atlantic in October–November 1997 [Singh et al., 2000], MINOS over the eastern Mediterranean in August 2001 [Lelieveld et al., 2002], ITCT 2K2 over the Northeast Pacific in April–May 2002 [Nowak et al., 2004], and PEM-Tropics B over the South Pacific in February–March 1999 [Singh et al., 2001]. Aside from MINOS, these missions were conducted for years other than the 2001 model year. We use monthly mean vertical profiles in the model over the flight regions (Figure 1) to compare to the mean observations. The measurements in TRACE-P, SONEX, and PEM-Tropics B were made by real-time gas chromatography (GC). The measurements in ITCT 2K2 and MINOS were made by proton transfer mass spectrometry (PTR-MS). A ship-based intercomparison of online PTR-MS and real-time GC-MS methods indicates high correlation between the two and agreement within a few percent [de Gouw et al., 2003]. Calibration differences of up to 20% may be expected between the GC measurements of Singh et al. from different missions. The TOPSE measurements were made by GC analysis from collected air canisters, all with the same relative humidity and the same lapse of time between collection and analysis. The accuracy is estimated to be 30% and the precision is much better.

Additional aircraft methanol data are available from the PEM-West B mission over the NW Pacific in February–March 1994 [Singh et al., 1995] and from the LBA-CLAIRE mission over the rain forest in Surinam in March 1998 [Williams et al., 2001]. The PEM-West B data are from the same region and season as TRACE-P, and show similar concentrations [Singh et al., 2004], but represent a much sparser data set. The LBA-CLAIRE data indicate low methanol concentrations, averaging 1.1 ppbv in the boundary layer and 0.6 ppbv in the free troposphere. These would suggest, consistent with Kesselmeier et al. [2002], that methanol emission from tropical forests of South America is much lower than predicted from the Galbally and Kirstine [2002] parameterization. However, quantitative comparison to the model is difficult because the observations were taken
under conditions of marine inflow and only about 12 hours downwind of the coast.

The TOPSE measurements are of particular interest as they characterize the latitudinal gradient at high northern latitudes during the transition from winter to spring [Atlas et al., 2003]. We show in Figure 8 the latitudinal profiles measured at 0–2 and 4–6 km altitude in February and April (only a few observations are available in May). Also shown are the corresponding model profiles, sampled as monthly means along the TOPSE flight tracks. Boundary layer concentrations in February show a latitudinal decrease from 1 ppbv at northern midlatitudes to 0.4 ppbv in the Arctic, both in the model and the observations, due to the shutting down of the biogenic source and the effect of surface deposition. The same latitudinal gradient is found in April but concentrations are about 0.2 ppbv higher, similarly in the model and in the observations, reflecting the springtime source. Concentrations at 4–6 km altitude show by contrast little latitudinal gradient, which the model explains as reflecting a weaker influence of the dry deposition sink. The February observations at 4–6 km (0.5 ppbv) are higher than the wintertime Zugspitze observations discussed in section 4.1 and are more consistent with the model results.

The model misses the observed April enhancement at 4–6 km south of 60N although it captures it in the 0–2 km data; this could be due to model error in the springtime onset of the source. We have no explanation for the high concentrations (above 1 ppbv) observed north of 80N at 4–6 km in April.

Aircraft observations at northern midlatitudes are available from the SONEX, MINOS, and ITCT 2K2 missions. These are shown in Figure 9 together with the corresponding model results. Observed concentrations in the middle and upper troposphere in SONEX are low, 0.4 ppbv on average, and this is reproduced in the model with only a slight positive bias. Reduced methanol emission from mature leaves is critical for simulation of the SONEX observations; without this reduction the simulated methanol concentrations would be much higher in fall because of the weak photochemical sink. The global 3-D model study of von Kuhlmann et al. [2003b] also found good agreement with the SONEX observations using a global methanol source 40% lower than ours (Table 1). They did not account for the decrease of methanol emission with leaf age.

The model reproduces the free tropospheric concentrations observed in MINOS (0.6–1 ppbv) over the Medi-
terrestrial in July, and also the high values in surface air. It is too low in the lower free troposphere (2–4 km), possibly reflecting difficulties in simulating vertical mixing and transport over this highly heterogeneous region. Salisbury et al. [2003] observed a mean concentration of 3.3 ppbv at a site in Crete during MINOS, and as shown in Figure 9 this is consistent with model results. Comparisons to the von Kuhlmann et al. [2003a, 2003b] model presented in that paper found it to be too low by a factor of four on average, possibly reflecting its low plant growth source.

[50] Observations from the ITCT 2K2 campaign off the California coast in April–May average 0.9–1.1 ppbv in the 1–7 km column with lower values near the surface, consistent with an ocean sink. The model is 0.2–0.6 ppbv too low. The comparison near the surface is subject to uncertainty because of land-ocean contrast (the model has more continental influence than the observations, but this may be due to numerical diffusion). The free tropospheric observations in ITCT 2K2 are similar in magnitude to the TRAC-E observations for the northeast quadrant (Figure 6), but the model shows lower values in ITCT 2K2 because of the longer distance from the Asian continent. Atmospheric production makes a relatively large contribution (0.2 ppbv) to the simulated concentrations in ITCT 2K2, reflecting the strong radiation and low NOx concentrations over the subsiding northeast Pacific. A possibility, discussed further below, is that this atmospheric source in the model is too weak.

[51] Observations from the PEM-Tropics B campaign over the South Pacific are typically 0.6–1.2 ppbv with little horizontal or vertical structure (Figure 9). These values are remarkably high considering the remoteness from land. The model is about a factor of 2 too low, averaging about 0.4–0.5 ppbv with little structure. A unique feature of model results for this region is that atmospheric production is the dominant source, contributing 0.2–0.4 ppbv. This source is favored by the high UV radiation (stimulating CH3O2 production), low NOx concentrations (suppressing competition from the CH3O2 + NO reaction), and low CO/CH4 ratio (resulting in a high CH3O2/HO2 ratio). The model underestimate in this region suggests that atmospheric production in the model is too low. A doubling of this source would largely correct the discrepancy and would be consistent with the featureless character of the observations.

[52] We previously discussed in section 2 the uncertainties in computing the atmospheric source of methanol from CH3O2 reactions. Examination of model results for the PEM-Tropics (B) region indicates that self-reaction (R1) accounts for 5–10% of the CH3O2 sink, reaction (R4) with NO for 20%, and reaction (R3) with HO2 for the rest; reaction (R2) is negligible. CH3OOH produced by (R3) has an atmospheric lifetime of about a day against losses by reaction with OH and photolysis:

\[ \text{CH}_3\text{OOH} + \text{OH} \rightarrow \text{CH}_3\text{O}_2 + \text{H}_2\text{O} \]  
(R5a)

\[ \text{CH}_3\text{OOH} + \text{OH} \rightarrow \text{CH}_2\text{O} + \text{OH} + \text{H}_2\text{O} \]  
(R5b)

\[ \text{CH}_3\text{OOH} + \text{hv} \rightarrow \text{CH}_3\text{O}_2 + \text{OH} \]  
(R6)

[53] Reaction (R5a) recycles CH3O2 while reactions (R5b) and (R6) do not. For typical atmospheric conditions, (R5a) accounts for about 50% of CH3OOH loss and (R5b) and (R6) each for about 25%. Assuming chemical steady state for CH3OOH, the effective sink of CH3O2 from (R3) is determined by \( k_3(1 - f) \), where the CH3O2 recycling efficiency \( f = k_{5a} / (k_{5a} + k_{5b} + k_5) \) is about 50%. Jet Propulsion Laboratory (JPL) [2003] gives uncertainty estimates of 30% for \( k_3 \), 40% for \( k_5 \), and 50% for \( k_{5a} \) (it gives no uncertainty estimate for the recommended branching ratio 70/30 for \( k_{5a}/k_{5b} \)). Thus the overall uncertainty on the effective CH3O2 sink from (R3) could be a factor of two. In addition, Martin et al. [2002] showed that the NO concentrations simulated by GEOS-CHEM over the South Pacific are about 50% higher than the PEM-Tropics (B) observations. Combination of these two factors could easily allow a doubling of the source of methanol from (R1), considering that this source is quadratic in the CH3O2 concentration. Photochemical model calculations by Olson et al. [2001] along the PEM-Tropics (B) flight tracks indicated a 50% overestimate of observed CH3OOH, which would be consistent with an overestimate of (R3). They also indicated a factor of 2 underestimate of CH2O (common product of CH3O2 degradation pathways) [Heikes et al., 2001]. As discussed by Heikes et al. [2001], the latter discrepancy suggests a major contribution from VOCs other than methane to the supply of CH3O2. This would provide an alternate explanation for increasing the methanol source from the CH3O2 self-reaction.

5. Discussion: Constraints on the Methanol Budget From Atmospheric Observations

[54] Observed atmospheric concentrations of methanol are consistent with the view that plant growth provides the principal global source. Laboratory and field studies indicate a factor of 2–3 decrease in this source from young to mature leaves [McDonald and Fall, 1993; Nemecek-Marshall et al., 1995; Karl et al., 2003]. We find that this is consistent with the observed seasonal variation of methanol concentrations at northern midlatitudes. A global plant growth source of 128 Tg yr\(^{-1}\), computed in our model using the NPP-based algorithm of Gulbally and Kirstine [2002], provides an overall unbiased simulation at northern midlatitudes. The tropics are the largest global contributor to this source but observations there are few. Data for tropical South America [Williams et al., 2001; Kesselmeier et al., 2002] suggest that our model source is too high, while data for Costa Rica [Karl et al., 2004] are consistent with the model. More work is clearly needed to improve understanding of methanol emission from tropical land ecosystems. The model also predicts high surface air concentrations (>10 ppbv) over Siberia in July, reflecting a large seasonal plant growth source as well as a long continental fetch. Measurements in this region would be of great value.

[55] The net flux of methanol from the terrestrial biosphere in the model is 96 Tg yr\(^{-1}\) including 128 Tg yr\(^{-1}\) from plant growth and 23 Tg yr\(^{-1}\) from plant decay, balanced by 55 Tg yr\(^{-1}\) dry deposition to land (Table 1). The atmospheric observations offer limited constraints for separating these three different terms. The plant growth source appears to be responsible for the observed decline of
concentrations from spring to fall, while the deposition sink appears to be responsible for the observed diurnal variation of concentrations as well as the decline at northern latitudes over the course of the winter.

[55] Observed concentrations of methanol over the South Pacific during the PEM-Tropics B aircraft mission (0.6–1.2 ppbv) are much higher than would be expected from the terrestrial vegetation source and show little vertical or longitudinal structure. Diffuse atmospheric production of methanol from the CH₂O₂ self-reaction (R1), where CH₂O₂ originates mainly from methane oxidation, is the principal model source in the region and contributes a 0.2–0.4 ppbv model background. Our computed global magnitude of this source (38 Tg yr⁻¹) is larger than previous estimates (18–31 Tg yr⁻¹) but still would need to be doubled to approach the South Pacific observations. Methanol observations over the remote north tropical and subtropical Pacific (TRACE-P and ITCT 2K2 aircraft missions) also suggest that atmospheric production of methanol from (R1) in the model is too low.

[57] Computation of the methanol source from (R1) in global tropospheric chemistry models is affected by uncertainties in the concentration of CH₂O₂ and in the branching ratio of CH₂O₂ reactions. A doubling to 50–100 Tg yr⁻¹ does not appear to be inconsistent with independent constraints, and is specifically within the constraints offered by ancillary PEM-Tropics B observations of CH₃OOH and CH₃O₂. It could reflect errors in kinetic rate constants, or the presence of VOCs other than methane in the remote marine atmosphere to provide sources of CH₂O₂. Some observations of aldehydes in remote marine air [Heikes et al., 2001; Singh et al., 2001, 2004] suggest the latter, and imply a more active organic photochemistry than is currently described in models, while other observations are more consistent with current understanding [Fried et al., 2003]. Measurements of HO₂ and total peroxy (HO₂ + CH₂O₂ + RO₂) radical concentrations by the chemical amplifier method during TRACE-P indicated concentration ratios of organic peroxy radicals to HO₂ that are consistent with current models [Cantrell et al., 2003], but there is substantial uncertainty in these measurements. Direct measurements of CH₂O₂ concentrations in the remote atmosphere are needed.

[58] Our model yields a global mean atmospheric lifetime for methanol of 7 days, with losses from gas-phase reaction with OH (63% of total loss), dry deposition to land (26%), wet deposition (6%), and uptake by the ocean (5%). Aqueous-phase oxidation by OH(aq) in clouds is negligible. The general ability of the model to reproduce the observed variability of methanol concentrations (cf. Figure 7) suggests a relatively narrow range of uncertainty in the lifetime, 5–10 days. Chemical loss of methanol by reaction with OH is uncertain by only about 30% [Jimenez et al., 2003] and most likely represents the dominant global sink for methanol. The low methanol concentrations observed at high northern latitudes in winter imply an additional nonphotocatalytic sink, which we attribute in the model to dry deposition to land. This attribution is supported by observations of methanol fluxes and of the diurnal variation of methanol concentrations at vegetated sites, although there is considerable variability from site to site. Heterogeneous loss in aerosols and clouds could provide an alternate explanation for the observed wintertime depletion of methanol but is not supported by independent evidence. Aircraft observations of vertical profiles over the oceans suggest that ocean uptake is only a weak sink, although ship data from the AOE-2001 summertime Arctic campaign suggest the possibility of rapid local uptake. Wet deposition, constrained by the Henry’s law solubility for methanol, is only a weak sink.

[59] Our best estimate of the global source of methanol (after allowing for doubling of atmospheric production to 76 Tg yr⁻¹) is 240 Tg yr⁻¹. Singh et al. [2004] estimated a much lower global source, 110 Tg yr⁻¹, by scaling the global acetone source of 95 Tg yr⁻¹ from Jacob et al. [2002] to the mean acetone/methanol concentration ratio of 2.1 g g⁻¹ that they measured in TRACE-P and to the ratio of lifetimes of acetone (15 days, from Jacob et al. [2002]) and methanol (9 days, from Heikes et al. [2002]). However, the concentration ratio measured in TRACE-P is lower than the expected global mean because of the large regional anthropogenic contribution to acetone [Jacob et al., 2002] and the relatively weak biogenic methanol emission from East Asia at that time of year. A methanol source of 110 Tg yr⁻¹ in our model would result in a low bias relative to observations. Singh et al. [2004] acknowledge that their emission estimate is subject to large uncertainty.

6. Implications for Atmospheric Chemistry

[60] Our budget implies a minor but nonnegligible role for methanol in global tropospheric chemistry. Oxidation of methanol by OH produces CH₃OH (major pathway) and CH₂O (minor), which both go on to produce CH₂O and HO₂ [Jimenez et al., 2003]. The global tropospheric rate of methanol oxidation calculated in our model on a per-carbon basis, 47 Tg C yr⁻¹, amounts to 12% of the corresponding value of 380 Tg C yr⁻¹ for methane oxidation [IPCC, 2001]. Methanol provides a significant source of CH₂O both in the continental boundary layer [Riemer et al., 1998; Palmer et al., 2003a] and in the remote troposphere [Heikes et al., 2001]. The production of CO from methanol oxidation amounts to 4–6% of the global CO source of 1800–2700 Tg yr⁻¹ (Duncan et al., submitted manuscript, 2004). Inclusion of methanol in GEOS-CHEM at the levels simulated here decreases the global tropospheric OH concentration by 2%, an effect similar to that reported previously by Tie et al. [2003]. Regional variations in this effect on OH are discussed by Tie et al. [2003] and are relatively small.

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