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Representation of the Community Earth System Model (CESM1) CAM4-chem within the Chemistry-Climate Model Initiative (CCMI)

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Abstract. The Community Earth System Model (CESM1) CAM4-chem has been used to perform the Chemistry Climate Model Initiative (CCMI) reference and sensitivity simulations. In this model, the Community Atmospheric Model version 4 (CAM4) is fully coupled to tropospheric and stratospheric chemistry. Details and specifics of each configuration, including new developments and improvements are described. CESM1 CAM4-chem is a low-top model that reaches up to approximately 40 km and uses a horizontal resolution of 1.9° latitude and 2.5° longitude. For the specified dynamics experiments, the model is nudged to Modern-Era Retrospective Analysis for Research and Applications (MERRA) reanalysis. We summarize the performance of the three reference simulations suggested by CCMI, with a focus on the last 15 years of the simulation when most observations are available. Comparisons with selected data sets are employed to demonstrate the general performance of the model. We highlight new data sets that are suited for multimodel evaluation studies. Most important improvements of the model are the treatment of stratospheric aerosols and the corresponding adjustments for radiation and optics, the updated chemistry scheme including improved polar chemistry and stratospheric dynamics and improved dry deposition rates. These updates lead to a very good representation of tropospheric ozone within 20 % of values from available observations for most regions. In particular, the trend and magnitude of surface ozone is much improved compared to earlier versions of the model. Furthermore, stratospheric column ozone of the Southern Hemisphere in winter and spring is reasonably well represented. All experiments still underestimate CO most significantly in Northern Hemisphere spring and show a significant underestimation of hydrocarbons based on surface observations.

1 Introduction

The Chemistry Climate Model Initiative (CCMI) coordinates evaluation and modeling activities for both tropospheric and stratospheric global chemistry-climate models. The CCMI-1 model experiments include three reference and several sensitivity experiments to evaluate the performance of chemistry-climate models in the troposphere and stratosphere for past and present conditions between 1960 and 2010 (REFC1 and REFC1SD), and to identify future climate trends between 1960 and 2100 (REFC2) (Eyring et al., 2013). The REFC1SD simulation differs from the REFC1 simulation in that the dynamics are specified from reanalysis. Comprehensive tropospheric and stratospheric chemistry has been integrated into the Community Atmospheric Model version 4 (CAM4-chem) of the Community Earth System Model (CESM1) and shows a reasonable representation of present-day atmospheric composition in the troposphere (Lamarque et al., 2012; Tilmes et al., 2015) and stratosphere

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(Lamarque et al., 2010). This model is therefore well suited to participate in the CCMI model intercomparison project.

The purpose of this paper is to summarize the CESM1 CAM4-chem model configurations that were used to perform the reference CCMI model experiments (Sect. 2) including physics, dynamics, the chemical mechanism and aerosol description, as well as a summary of newly integrated diagnostics. We also describe issues that have been identified after the simulations were performed and their likely impacts. In addition, we summarize the global performance of the model in Sect. 3, and evaluate selected diagnostics based on observational data sets in Sect. 4. We employ existing and new data sets to evaluate the general performance of the model. More in-depth analysis and evaluations will follow in multi-model comparison studies. Improvements in comparison to earlier versions of the model are discussed in the Conclusions.

2 Model description

CESM is a fully coupled Earth System model, which includes atmosphere, land, ocean and sea-ice components. All CCMI simulations are carried out with the same model code that is based on CESM version 1.1.1 (CESM1) (Neale et al., 2013), with modifications discussed below. The configuration of the model used here fully couples the Community Atmosphere Model version 4 (CAM4), the Community Land model version 4.0 (CLM4.0), the Parallel Ocean Program version 2 (POP2) and the Los Alamos sea ice model (CICE version 4). The land model was run without an interactive carbon or nitrogen cycle and only the atmospheric and land components are coupled to the chemistry. The climatological present-day land cover is used for all simulations.

2.1 The atmosphere model

Detailed information about the physics of the atmosphere model used here are described in Neale et al. (2013) and Richter and Rasch (2008), and also summarized in Lamarque et al. (2012, and references therein). In summary, deep convection is treated by Zhang and McFarlanle (1995) with improvements in the convective momentum transport (Richter and Rasch, 2008), which improved surface winds, stresses and tropical convection. At the same time, an entraining plume was added to the convection parameterization, which together with the momentum transport improved the representation of El Niño-Southern Oscillation (ENSO) significantly (Neale et al., 2008). The photolysis calculation uses a look-up table between 200 and 750 nm and online calculations for wavelengths < 200 nm. Only changes in the ozone column, but not in the aerosol burden, impact photolysis rates. Attenuation of the spectral irradiance above the model top is calculated using the approach of Kinnison et al. (2007) and Lamarque et al. (2012).

Processes in the planetary boundary layer are represented using the Holtslag and Boville (1993) parameterization. Wet deposition of gas and aerosol compounds is based on Neu and Prather (2012), as described in Lamarque et al. (2012). In this version of CAM4-chem all aerosols in the cloudy fraction of the grid cell are assumed to reside within cloud droplets and are removed in proportion to the cloud water removal rate. Aerosols directly impact the radiation and chemistry, but do not change the radiative properties of clouds (i.e., no representation of the aerosol indirect effects is included).

Lightning NO_x is parameterized following Price and Vaughan (1993) and Price et al. (1997). The global amount of produced lightning NO_x is scaled differently for the specified dynamics (SD) and the free-running (FR) experiments due to differences in the meteorology to ensure values of approximately 3–5 Tg N year⁻¹ for present-day conditions.

2.1.1 Model grid

For all CCMI reference simulations, CESM1 CAM4-chem uses a horizontal grid with a resolution of $1.9^{\circ} \times 2.5^{\circ}$ (latitude by longitude), and uses the finite volume dynamical core. The top of the model is located at 3 hPa (about 40 km). The vertical coordinate is sigma (hybrid terrainfollowing pressure) in the troposphere, switching over to isobaric at pressure levels less than 100 hPa; the vertical resolution of the model depends on the configuration of the experiment. The atmosphere model, CAM4, makes use of two different configurations, the FR (with 26 vertical levels) and the SD(with 56 vertical levels adopted from the analysis fields); see Lamarque et al. (2012). For the SD configuration, internally derived meteorological fields are nudged every time step of 30 min by 1 % towards reanalysis fields (equivalent to a 50h Newtonian relaxation timescale for nudging) from Modern-Era Retrospective Analysis for Research and Applications (MERRA) reanalysis (http://gmao. gsfc.nasa.gov/merra/) (Rienecker et al., 2011). Nudged meteorological fields include wind components, temperatures, surface pressure, surface stress, latent and sensible heat flux. The MERRA reanalysis fields are interpolated to the horizontal resolution of the model prior to running the simulation. The MERRA surface geopotential height is used for the SD simulations to be consistent with the reanalysis fields.

2.1.2 Quasi-Biennial Oscillation

The SD configuration of the model incorporates the observed quasi-biennial oscillation (QBO), which is present in the meteorological analysis fields. The limited vertical resolution of the FR model configurations does not allow for the generation of an internal QBO in CAM4-chem. Therefore, for the FR CCMI experiments, REFC1 and REFC2, the QBO is imposed in the model by relaxing equatorial zonal winds between 90 to 3 hPa to the observed interannual variability, following the approach by Matthes et al. (2010). Here, we

vary the QBO phase between eastward and westward phase using an approximate 28-month period, similar to what was done by Marsh et al. (2013).

2.1.3 Improved gravity wave representation

The representation of sub-grid-scale gravity waves (GW) in CAM was formerly limited to orographic gravity waves using the parameterization adapted from McFarlane (1987). In the present simulations, the parameterizations of non-orographic gravity waves generated by convection (Beres et al., 2005) and fronts (Richter et al., 2010), which were developed for the Whole Atmosphere Community Climate Model (WACCM), are also included.

In addition, we have added another gravity wave module to represent the waves with large horizontal wavelengths that are often observed in the stratosphere (e.g., Zink and Vincent, 2001). The new GW module is adopted from the inertia-gravity wave (IGW) parameterization developed by Xue et al. (2012) for an interactive QBO. The formulation includes the impact of the Coriolis force on gravity wave propagation and breaking. Rather than applying it in the equatorial region, as done by Xue et al. (2012), we use a more general mechanism for determining sources; gravity waves are triggered by the same frontal threshold used for the mesoscale gravity waves (Richter et al., 2010). This has the impact of shifting the bulk of the waves from the tropics to middle and high latitudes. In the current implementation, gravity waves have a narrow phase speed spectrum (-20 to) $20\,\mathrm{m\,s^{-1}}$) and long horizontal wavelength (1000 km). The momentum forcing associated with this module particularly impacts the winter stratosphere. In the Southern Hemisphere (SH), it enhances downwelling and increases the winter stratospheric temperature, which in previous simulations was substantially colder than observed.

However, it was found, that the version of the IGW parameterization used for the performed experiments has a narrow IGW spectrum centered on zero phase velocity instead of being centered on the speed of the background wind at the GW launch level, as in the standard GW parameterization. Even with this shortcoming, the model produces a much improved temperature evolution in the stratosphere, in particular in the SH high latitudes compared to earlier versions. This results in a well-resolved ozone hole in winter and spring over Antarctica. No significant changes are expected from a corrected IGW parameterization for the troposphere.

2.1.4 Tropospheric aerosols

CAM4-chem runs with the bulk aerosol model (BAM), which simulates the distribution of externally mixed sulfate, black carbon (BC), primary organic carbon (OC), sea-salt and dust, as described in Lamarque et al. (2012). The dust emissions are calibrated so that the global dust aerosol optical depth (AOD) is about 0.025 to 0.030 (Mahowald et al.,

2006). The distribution of sea-salt and dust are described using four size bins. In CAM4-chem, the formation of secondary organic aerosols (SOA) is coupled to the chemistry and biogenic emissions. SOA are derived using the twoproduct model approach using laboratory determined yields for SOA formation from monoterpene, isoprene and aromatic photooxidation, as described in Heald et al. (2008). The aging process of BC and OC from hydrophobic to hydrophilic is included through a specified conversion timescale. For all aerosol species, the size distributions are specified as in Lamarque et al. (2012). Aerosols interact with the gas-phase chemistry through heterogeneous reactions that depend on the available surface area density (SAD), as discussed below. For the tropospheric SAD calculation, sulfate, hydrophilic black carbon, primary organic carbon and nitrates are included, where SOA has not been included. This may lead to a significant underestimation of tropospheric SAD in the experiments.

2.1.5 Representation of aerosols in the stratosphere

Aerosol mass, heating rates and SAD are revised in this version compared to earlier configurations. Most significantly, the model uses a new stratospheric aerosol and SAD data set, derived based on observations, to force models participating in CCMI (Eyring et al., 2013). In addition, in order to fully utilize the aerosol size information provided by the new model input file, the optics in the radiative transfer code associated with CAM4 (i.e., CAMRT) (Neale et al., 2010) have been modified to include a lookup table for aerosol effective radius in the shortwave radiation scheme. The new description leads to an updated representation of volcanic heating for REFC1 and REFC2, whereas in REFC1SD volcanic heating is included through the nudged temperature fields. See Neely et al. (2015) for a full description of changes to the stratospheric aerosol scheme. Tropospheric aerosols that enter the stratosphere are promptly removed (as listed in Table A2) since the aerosol burden in the stratosphere is prescribed.

2.1.6 Coupling to the land model

Dry deposition velocity for tracers in the atmosphere are calculated online in CLM4.0. An updated calculation is used, where leaf and stomatal resistances are coupled to the leaf area index (LAI) and are also linked to the photosynthesis provided by the land model, as described in Val Martin et al. (2014).

Biogenic emissions are calculated online in CLM using the Model of Emissions of Gases and Aerosols from Nature (MEGAN), version 2.1 (Guenther et al., 2012). An erroneous implementation of MEGAN in this version differs from the description of Guenther et al. (2012) by using the LAI from the previous model time step (30 min) instead of the average of the previous 10 days. In addition, in this version we are using a fixed CO₂ mixing ratio, instead of the simulated

atmospheric value, in the calculation of the CO_2 inhibition effect on isoprene emissions. The corrected implementation is closer to the algorithm of Guenther et al. (2012).

2.2 Chemical mechanism

The chemical mechanism of CAM4-chem includes 169 species, listed in Table A1 in the Appendix. Depending on the chemical lifetime of each species, an explicit or semi-implicit solver is used. Emissions of gas-phase and aerosol species, as indicated in Table A1, are in general distributed at the surface. Only aircraft emissions of BC and nitrogen dioxide, and volcanic emissions of sulfur and sulfate, are vertically distributed. Species with lower boundary conditions, as indicated in Table A1, as discussed in Sect. 2.3.2. Different species experience wet and/or dry deposition, as also listed in Table A1. Furthermore, 14 artificial tracers are implemented as recommended by CCMI (Eyring et al., 2013, Sect. 4.2.1): NH₅, NH₅₀, NH₅₀W, AOA_{NH}, ST80₂₅, CO₂₅, CO₅₀, SO_{2t}, SF6em, O₃S, E90, E90_{NH}, E90_{SH}. O₃S is a stratospheric ozone tracer that represents the amount of ozone in the troposphere with its source in the stratosphere. O₃S is set to stratospheric values at the tropopause, and experiences the same loss rates as ozone in the troposphere, as defined by CCMI. As interpreted from the CCMI recommendation, dry deposition is not included, which will lead to an overestimation of O₃S in the lower boundary layer when compared to ozone (which is dry deposited).

The chemical mechanism, is based on the Model for Ozone and Related chemical Tracers (MOZART), version 4 mechanism for the troposphere (Emmons et al., 2010). It further includes extended stratospheric chemistry (Kinnison et al., 2007) and updates, as described in Lamarque et al. (2012) and Tilmes et al. (2015). The reactions include photolysis, gas-phase chemistry and heterogeneous chemistry, in both troposphere and stratosphere. The complete chemical mechanism is listed in Table A2 and incorporates all the latest updates. All aerosols and some gas-phase species, including H₂O, O₂, CO₂, O₃, N₂O, CH₄, CFC11, CFC12, are radiatively active.

Reaction rates are updated following JPL2010 recommendations (Sander et al., 2011). Bromoform (CHBr₃) and dibromomethane (CH₂Br₂) were added to the model to represent the stratospheric bromine loading from very short-lived (VSL) species. The surface volume mixing ratio for these two VSL species was set globally to 1.2 ppt (i.e., 6 ppt total bromine). This approach adds an additional ≈ 5 ppt of inorganic bromine to the stratosphere. The resulting stratospheric total inorganic bromine abundance (for present-day conditions) from both long-lived and VSL species is ≈ 21.5 ppt. Besides the current lower boundary condition (LBC) approach for VSL species, CAM4-chem can be also configured with a full-VSL mechanism, including detailed gasphase halogen chemistry mechanism, geographically and

time-dependent distributed sources of nine halocarbons and improved representation of heterogeneous recycling and removal rates in the troposphere (Fernandez et al., 2014; Saiz-Lopez et al., 2014).

Details on updated reactions and processes for chemistry in the polar stratosphere are described in Wegner et al. (2013) and Solomon et al. (2015).

Diagnostics of the tropospheric ozone production and loss rates are explicitly calculated; see Table A3, in adding the listed reaction rates r of two species A and B, r(A-B), as well as the photolysis reaction of ONITR (defined as lumped organic nitrate species that includes nitrates derived from the OH- and NO₃-initiated oxidation of isoprene and terpenes, and related species), called jonitr:

$$\begin{aligned} \text{O3-Prod} &= r(\text{NO} - \text{HO2}) + r(\text{CH3O2} - \text{NO}) \\ &+ r(\text{PO2} - \text{NO}) + r(\text{CH3CO3} - \text{NO}) \\ &+ r(\text{C2H5O2} - \text{NO}) + 0.92 \cdot r(\text{ISOPO2} - \text{NO}) \\ &+ r(\text{MACRO2} - \text{NOa}) + r(\text{MCO3} - \text{NO}) \\ &+ r(\text{MACRO2} - \text{NO}) + r(\text{RO2} - \text{NO}) \\ &+ r(\text{C3H7O2} - \text{NO}) + 0.9 \cdot r(\text{TOLO2} - \text{NO}) \\ &+ r(\text{XO2} - \text{NO}) + 0.9 \cdot r(\text{ALKO2} - \text{NO}) \\ &+ r(\text{TERPO2} - \text{NO}) + 0.9 \cdot r(\text{ALKO2} - \text{NO}) \\ &+ r(\text{ENEO2} - \text{NO}) + r(\text{EO2} - \text{NO}) \\ &+ r(\text{MEKO2} - \text{NO}) + 0.4 \cdot r(\text{ONITR} - \text{OH}) \\ &+ \text{jonitr} \end{aligned}$$

$$\begin{aligned} \text{O3-Loss} &= r(\text{O1D} - \text{H2O}) + r(\text{OH} - \text{O3}) \\ &+ r(\text{HO2} - \text{O3}) + r(\text{C3H6} - \text{O3}) \\ &+ 0.9 \cdot r(\text{ISOP} - \text{O3}) + r(\text{C2H4} - \text{O3}) \\ &+ 0.8 \cdot r(\text{MVK} - \text{O3}) + 0.8 \cdot r(\text{MACR} - \text{O3}) \\ &+ r(\text{C10H16} - \text{O3}). \end{aligned}$$

These are defined based on the rate-limiting terms for the gas-phase reactions of the O_x family (O_3, O, O_1D, NO_2) , not including $O_2 + hv \rightarrow 2O$ production, O_x , ClO_x , and BrO_x losses, and are therefore not valid for the stratosphere. The sum of those rates are very similar to the explicit calculation of the net chemical change of ozone (as listed in Table A2).

2.3 Experimental Setup

The reference experiments are set up according to the CCMI recommendation, including surface and altitude-dependent emissions, and lower boundary conditions. The three reference experiments are performed with the recommended emissions. REFC1 and REFC1SD (years 1960–2010) use the same emissions, excluding biogenic emissions. Anthropogenic and biomass burning emissions are from the Monitoring Atmospheric Composition and Climate/CityZen inventory (MAC City) emission data set and change every year (Granier et al., 2011). For REFC2 (years 1960–2100), anthropogenic and biomass burning emissions are taken from AR5 (Eyring et al., 2013) (see Fig. A1), which only vary ev-

ery 5–10 years. All emissions include a seasonal cycle. Biogenic emissions are calculated every time step by MEGAN, as described in Sect. 2.1.6.

The REFC1SD experiment is nudged to analyzed air temperatures, winds, surface fluxes and surface pressure, and uses the Hadley Centre Global Sea Ice and Sea Surface Temperature version 2 (HadISST2) observed time-dependent data set for sea surface temperatures (SSTs) and sea ice. The REFC1 experiment also uses prescribed SSTs and sea ice, while the REFC2 simulation calculates temperatures in the ocean and atmosphere. We have carried out one simulation for REFC1SD, and an ensemble of three members for each REFC1 and REFC2.

The solar cycle is prescribed using observed daily fields for the years until 2010. For the future period in REFC2, we follow the CCMI recommendation and repeat a sequence of the last four solar cycles (20–23), as defined in http://solarisheppa.geomar.de/ccmi.

2.3.1 Initial conditions and spin-up

CAM4-chem initial conditions for the three REFC1 and REFC2 ensemble members are taken from 3 realizations of CESM1-WACCM 20th Century ensemble for CMIP5 (Marsh et al., 2013). The spin-up period started in 1950 and ran through 1959. The experiments simulated the years 1960 to 2010 (REFC1) and 1960 to 2100 (REFC2). Initial conditions for the REFC1SD simulation are taken from the first REFC1 ensemble member in 1975. The spin-up of this experiment covered the years 1975 to 1979, repeating 1979 meteorological analysis for each year. The experiment was performed between 1980 and 2010.

2.3.2 Lower boundary conditions

For all of the three reference experiments the same monthly and annually varying lower boundary conditions are used based on the Representation Concentration Pathway 6.0 (RCP6.0) Coupled Model Intercomparison Project Phase 5 (CMIP5) future projection (Taylor et al., 2012). We prescribe CO₂, N₂O, CH₄, as well as the following halogen species based on the CCMI recommendations: CCl₄, CF₂ClBr, CF₃Br, CFC11, CFC113, CFC12, CH₃Br, CH₃CCl₃, CH₃Cl, H₂, HCFC22, CFC114, CFC115, HCFC141b, HCFC142b, CH₂Br₂, CHBr₃, H1202, H2402, SF₆. A north–south gradient was added for CH₃Br, HCFC22, HCFC141b, HCFC142b, based on the HIAPER (High-Performance Instrumented Airborne Platform for Environmental Research) Pole-to-Pole Observations (HIPPO) (Wofsy et al., 2011; Mijeong Park, personal communication, 2015).

3 Model performance

3.1 Global diagnostics

The general state of the model is investigated by comparing diagnostics of globally averaged values between different model experiments that are averaged between 1995 and 2010 (Table 1). The global surface temperatures (TS) of all three experiments are in agreement within 0.15 K for the observed period (Table 1). REFC1SD land temperature (TS land) is on average 0.25 K higher than for REFC1 and 0.15 K higher than for the REFC2 experiments (Table 1). The largest deviations occur over high latitudes (not shown). In the REFC1SD experiment, low cloud fraction is significantly larger than in the other experiments, which results in a much smaller shortwave cloud forcing (SWCF) of $-83 \, \mathrm{W \, m^{-2}}$ compared the other experiments that are with 54–56 W m⁻² more in line with observations.

Differences in clouds and land surface temperatures between the reference experiments result in different biogenic emissions of volatile organic components (VOCs) (Fig. 1). REFC1SD biogenic emissions are about 10 % lower than in the REFC1 experiment and about 15% lower than in the REFC2 experiment. The emissions differ the most in summer during their peak (Fig. 1, bottom row). Despite the fact that surface temperatures in REFC1SD are warmer than in REFC1, more low cloud clouds and reduced solar radiation (as evident in photolysis rates) near the surface may be the important driver for the reduced biogenic emissions in REFC1SD, which has to be further investigated. Other differences in the REFC1 and REFC2 VOC emissions arise from different anthropogenic and biomass burning emissions, while biogenic emissions differ by less than 10 % (Table 1). Despite the variation in the reference experiments, biogenic emissions are in agreement with earlier estimates (e.g., Young et al., 2013).

The performance of the model in simulating tropospheric chemical variables (Table 1) is similar to earlier studies (e.g., Tilmes et al., 2015). Methane lifetime is low compared to observational estimates of 11.2 years (Prather et al., 2012). Ozone budgets, including ozone burden, stratosphere—troposphere exchange, and budgets of carbon monoxide (CO), are in agreement with earlier model studies (Young et al., 2013). Aerosol burdens of primary organic matter (POM) and SOA are low, but within the spread of other model results (Tsigaridis et al., 2014). The SO₄ burden with 0.45 to 0.51 TgS and the lifetime of 3.0 to 3.5 days is somewhat low compared to the Aerocom multi-model mean of 0.66 TgS and 4.12 days, respectively (e.g., Liu et al., 2012). The dust optical depth around 0.04 is somewhat higher than suggested by Mahowald et al. (2006).

Table 1. Overview of global diagnostics for different experiments, averaged between 1995 and 2010. Lifetimes and burdens are calculated for the troposphere defined for regions where ozone is below 150 ppb.

CESM1 CAM4chem	REFC1SD	REFC1.1	REFC1.2	REFC1.3	REFC2.1	REFC2.2	REFC2.3
Meteorology	MERRA	CAM4	CAM4	CAM4	CAM4	CAM4	CAM4
Vert. Res.	56L	26L	26L	26L	26L	26L	26L
TS Global	288.43	288.27	288.27	288.28	288.35	288.40	288.41
TS Land	282.37	282.10	282.12	282.17	282.23	282.20	282.23
SWCF	-82.47	-55.96	-56.01	-55.97	-54.66	-54.65	-54.78
CH ₄ Burden (Tg)	3991.3	4100.5	4103.8	4099.3	4101.4	4105.0	4103.1
CH ₄ Lifet. (year)	7.6	8.0	8.1	8.1	8.2	8.2	8.2
CH ₃ CCl ₃ Lifet. (year)	4.5	4.8	4.8	4.8	4.9	4.9	4.9
CO Burden (Tg)	289.6	303.6	305.3	305.7	315.4	316.7	315.3
CO Emis. (Tg year ⁻¹)	1114.8	1119.3	1126.5	1126.8	1170.1	1171.1	1169.9
CO Dep. (Tg year ⁻¹)	125.8	120.7	122.0	122.1	122.7	123.0	122.9
CO Chem. Loss (Tg year $^{-1}$)	2264.1	2294.2	2295.3	2298.0	2348.4	2353.3	2345.5
CO Lifet. (days)	44.2	45.9	46.1	46.1	46.6	46.7	46.6
O ₃ Burden (Tg)	332.5	326.9	326.5	326.4	327.8	327.2	327.0
O_3 Dep. (Tg year ⁻¹)	871.7	894.4	893.9	894.2	895.0	892.8	894.7
O ₃ Chem. Loss (Tg year ⁻¹)	4256.0	4268.3	4250.6	4259.0	4287.6	4293.5	4278.9
O_3 Chem. Prod. (Tg year ⁻¹)	4693.8	4710.0	4706.5	4708.3	4747.2	4756.9	4744.1
O_3 Net Chem.Change (Tg year ⁻¹)	392.9	420.9	430.5	426.0	432.5	436.5	438.2
O_3 STE (Tg year ⁻¹)	478.8	473.4	463.4	468.2	462.5	456.4	456.5
Isop. Emis. (Tg year ⁻¹)	454.2	512.6	511.8	515.0	546.6	551.6	545.6
Monoterp. Emis. (Tg year ⁻¹)	138.9	150.0	150.0	150.3	155.4	156.4	155.0
Methanol Emis. (Tg year ^{−1})	100.4	114.6	114.8	114.9	113.7	114.9	113.4
Aceton Emis. (Tg year ⁻¹)	41.6	44.3	44.3	44.3	47.8	48.1	47.7
Lightning Prod. (TgN year ⁻¹)	4.5	4.8	4.7	4.8	4.7	4.7	4.7
Total optical depth	0.107	0.119	0.119	0.119	0.118	0.118	0.118
Dust optical depth	0.041	0.043	0.043	0.043	0.040	0.041	0.040
POM Burden (TgC)	0.75	0.73	0.73	0.74	0.77	0.77	0.77
POM Emis. (TgC year ⁻¹)	48.38	47.99	48.38	48.38	51.23	51.23	51.23
POM Lifet. (days)	7.23	7.18	7.15	7.19	7.05	7.06	7.01
SOA Burden (TgC)	0.54	0.49	0.49	0.49	0.51	0.51	0.50
SOA Chem. Prod. (TgC year ⁻¹)	32.79	34.45	34.43	34.79	35.86	36.32	35.54
SOA Lifet. (days)	0.54	0.49	0.49	0.49	0.51	0.51	0.50
BC Burden (TgC)	0.12	0.12	0.12	0.12	0.12	0.12	0.12
BC Emis. $(TgC year^{-1})$	7.71	7.68	7.71	7.71	7.95	7.95	7.95
BC Lifet. (days)	7.44	7.48	7.46	7.49	5.88	5.89	5.86
DUST Burden (TgC)	43.87	45.04	45.03	45.20	42.60	42.75	42.31
SALT Burden (TgC)	6.02	10.88	10.88	10.87	11.14	11.10	11.11
SO ₄ Burden (TgS)	0.45	0.49	0.49	0.49	0.51	0.51	0.51
SO ₄ Emis. (TgS year ⁻¹)	0.25	0.25	0.25	0.25	0.25	0.25	0.25
SO ₄ Dry Dep. (TgS year ⁻¹)	5.29	5.76	5.78	5.77	5.94	6.00	5.99
SO ₄ Wet Dep. (TgS year ⁻¹)	-49.93	-46.36	-46.28	-46.30	-46.36	-46.42	-46.49
SO ₄ Chem. Prod. (TgS year ⁻¹)	10.35	10.81	10.83	10.82	10.98	11.02	11.02
SO ₄ AQ. Prod. (TgS year ⁻¹)	44.95	41.41	41.34	41.35	41.44	41.53	41.58
SO ₄ Total Prod. (TgS year ⁻¹)	55.30	52.23	52.17	52.18	52.42	52.55	52.60
SO ₄ Lifet. (days)	2.97	3.41	3.42	3.41	3.52	3.54	3.53

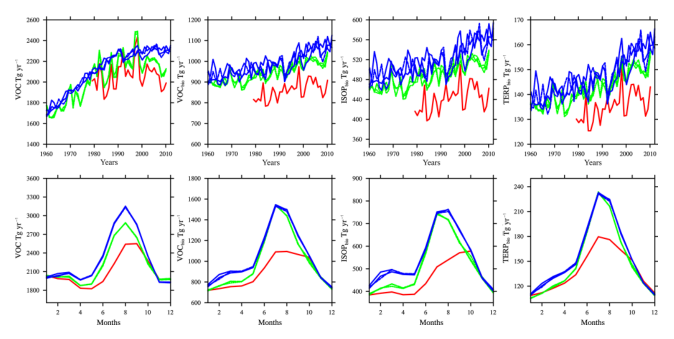


Figure 1. Global-averaged surface emissions of total volatile organic compounds (VOCs) (first column), biogenic VOCs (second column), biogenic isoprene (third column), and biogenic terpenes (fourth column), for different experiments, REFC1SD (red), REFC1 (green), REFC2 (blue). The seasonal cycle of zonal averages between 1960 and 2010 are shown at the bottom row.

3.2 Trends of tropospheric components

Time-varying emissions of ozone precursors and aerosols impact the oxidation capacity of the atmosphere. In the following, we discuss the evolution of different chemical species and surface area density in the tropical troposphere between 30° S and 30° N, tropospheric methane lifetime and stratospheric column ozone (Fig. 2), since methane is mostly controlled by processes in the tropics. Increasing nitrogen dioxide (NO₂), CO and VOC burdens between 1960 and 1990 result in increasing tropospheric ozone with the strongest trend between 1960 and 1990. Increasing aerosols between 1960 and 1990 result in an increase in SAD, with little change after 1990. Together with the increase in CO burden, this results in a decrease of OH. Other factors result in an increase in tropospheric OH, including decreasing stratospheric column ozone between 1960 and 2010, increasing tropospheric column ozone, increasing nitrogen dioxides (NO_x) burden, and decreasing methane emissions (e.g., Murray et al., 2014). Both counteracting effects on OH result in little change in methane lifetime between 1960 and 1990. After 1990, SAD, as well as CO and VOC, trends are leveling off, but nitrogen dioxide and ozone burdens are still increasing, partly due to increasing lightning NO_x production (not shown). This results in a decreasing trend in methane lifetime after 1990 for all reference experiments.

The burden of chemical tracers differs between REFC1SD and REFC1/REFC2 (Fig. 2). Variations in emissions and atmospheric dynamics, including surface temperature, clouds and convection, influence the chemical composition of the

atmosphere. Exchange processes between the upper troposphere and lower stratosphere are also different in the model experiments and impact ozone. The shorter lifetime of methane in REFC1SD compared to the other experiments may be a result of a reduction in high clouds, and, to some amount, the larger ozone mixing ratios in the tropical troposphere, which would increase the oxidation capacity in the tropics. This has to be investigated in more detail in future studies.

Besides a continuous decrease, the stratospheric ozone column shows a significant drop after major volcanic eruptions (e.g., WMO, 2006). This is expected due to an increase in stratospheric SAD after the eruption, which causes enhanced halogen activation, resulting in ozone depletion (see Fig. 2).

4 Evaluation against selected diagnostics

The purpose of this section is to give an overview of selected variables and diagnostics that summarize the performance of the model, including some of its shortcomings, in comparison to observations. Additional and more detailed investigations are expected in future multi-model comparison studies. We only discuss the performance of the reference experiments for past and present day. Model results from other sensitivity studies are not analyzed and will be discussed in future studies.

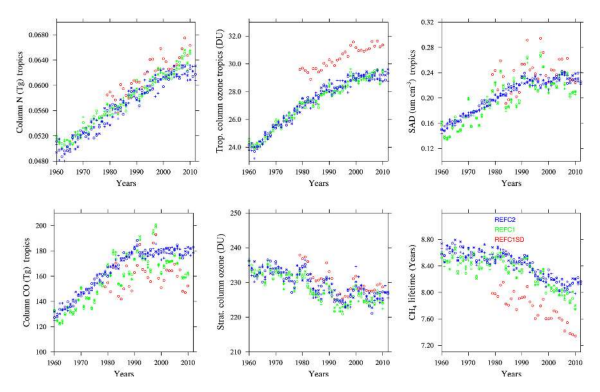


Figure 2. Time series of annually averaged column integrated tropospheric and tropical nitrogen dioxide (in Tg N), tropospheric ozone burden, and CO, in (30° S–30° N), tropical average of tropospheric surface area density, global stratospheric column ozone and tropospheric methane lifetime.

4.1 Ozone

Ozone is an important atmospheric tracer in both the troposphere and the stratosphere. In the troposphere and at the surface, ozone is an air pollutant and is impacted by various precursors, most importantly CO and NO_x. A reasonable performance of tropospheric ozone is required for air quality studies. In the stratosphere, ozone is strongly influenced by dynamics, photo-chemistry and catalytic reactions (e.g., WMO, 2011). The strength of the transport of stratospheric ozone into the troposphere follows a seasonal cycle controlled by the Brewer-Dobson circulation (BDC). Shortcomings in the representation of the strength of the BDC and mixing processes between stratosphere and troposphere influence the performance of tropospheric ozone, as discussed below. In addition, ozone is an important greenhouse gas in the upper troposphere and lower stratosphere (UTLS) and influences tropospheric climate (e.g., WMO, 2014).

4.1.1 Trends and seasonality of ozone

Ozone trends and seasonality in the reference experiments are compared to ozonesonde observations (Tilmes et al., 2012) in the free troposphere (at 500 hPa) and the boundary layer (at 900 hPa). For Japan, we employ an additional climatology derived by Tanimoto et al. (2015), which is based on surface observations at five marine boundary layer

sites from the Acid Deposition Monitoring Network in East Asia (EANET) for pressure levels larger than 900 hPa, and a combination of the historical Measurements of OZone, water vapor, carbon monoxide and nitrogen oxides by inservice AIrbus airCraft (MOZAIC; URL: http://www.iagos. fr/mozaic) data (over Narita airport) and ozonesonde observations (at Tateno/Tsukuba) for pressure levels between 472 and 616 hPa. We use an artificial stratospheric ozone tracer (O₃S) to identify differences in stratosphere–troposphere exchange (STE) between different model experiments for four selected regions (see Figs. 3 and 4).

In high northern latitudes, REFC1SD reproduces the magnitude and trend of ozone very well, including variability within the standard deviation of the observations for all seasons, as shown in the example of the Northern Hemisphere (NH) polar west region (Fig. 3, first and second row). A very good agreement between the model experiment and ozonesondes also exists for western Europe, with the exception of the high bias between October and February at 500 hPa of 5–10 ppb (Fig. 3, third and fourth row).

Results from REFC1 and REFC2 show larger deviations from the observations than REFC1SD over these two regions. These are in part due to not only differences in the amount of stratospheric ozone entering the troposphere for the different experiments (see Fig. 3, right column, dashed lines), but also changes in ozone loss and production. Discrepancies in ozone between the experiments can be ex-

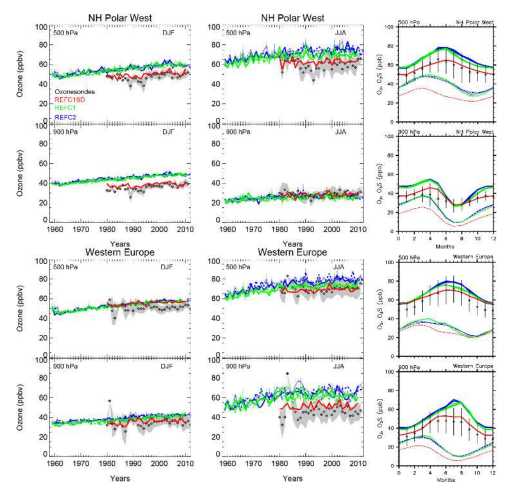


Figure 3. Left and middle column: time evolution of seasonal averaged and regionally aggregated ozone mixing ratios derived from ozone soundings (black diamonds) and model results (colored lines) at two different pressure levels, two different seasons (DJF: left, JJA: right) and regions (NH polar west, and western Europe). Grey shading indicates the standard deviation of the observations that include at least 12 observed profiles per season in a year. Colored error bars indicate the standard deviation based on monthly averaged model output. Right column: Regionally aggregated seasonal cycle comparisons of ozone soundings (black lines) and model simulations (colored lines), averaged between 1995 and 2010. Dashed lines indicate mixing ratios of the stratospheric ozone tracer (see text for more details).

plained by differences in O₃S for the whole year at 500 hPa and for winter months at 900 hPa. During summer months, differences in chemical production at the surface for the different experiments seem to play an additional role and explain about 5–10 ppb of the deviations for western Europe.

Selected ozonesondes over eastern USA and Japan are located further south and are more strongly influenced by tropical air masses and tropospheric intrusion in the lowermost stratosphere in particular in winter, as discussed in Tilmes et al. (2012). Each of the regions covers only two stations and so uses fewer observations for the different years than other regions, which increases the uncertainty of trends (Saunois et al., 2012).

Comparisons for eastern USA and Japan are illustrated in Fig. 4. For Japan, we are using two data sets to compare to model results. Ozone mixing ratios and trends at 900 hPa over Japan using ozonesondes, as compiled by Tilmes et al.

(2012), Fig. 4 (black diamonds), largely differ from the climatology by Tanimoto et al. (2015), which is based on surface observations (black triangles). This is due to uncertainties in the ozonesonde observations at these altitudes, which should be treated with caution. On the other hand, the two climatologies agree well in the free troposphere at 500 hPa.

For eastern USA and Japan the REFC1SD model experiment nicely reproduces the observed trend and magnitude of ozone within the variability of the observations at 900 hPa. The seasonal cycle for both regions are well reproduced. This significant improvement compared to earlier versions of the model is in part a result of the improved calculation of dry deposition rates, as discussed in Val Martin et al. (2014) over the USA. REFC1 and REFC2 experiments show slightly larger values at 900 hPa in comparison to the REFC1SD experiment particularly in winter, aligned with a larger O₃ contribution from the stratosphere, as determined by the O₃S

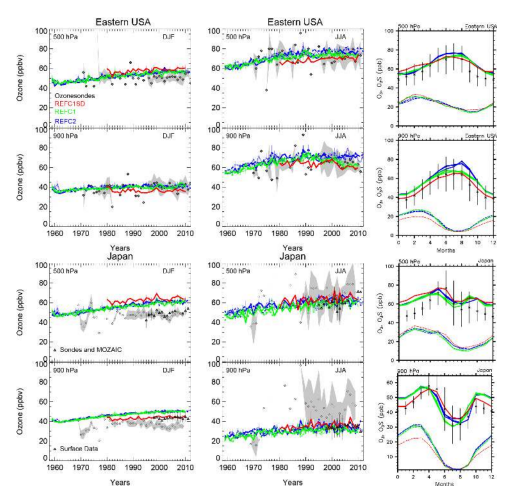


Figure 4. As Fig. 3, but for Eastern USA and Japan instead. For Japan, ozone time series compiled by Tanimoto et al. (2015) are added (black triangles) (see text for more details) and used to compare with the seasonal cycle of the model for Japan.

tracer (see Fig. 4). At 500 hPa, ozone mixing ratios and trends are well reproduced for all experiments in summer. However, the model overestimates winter ozone mixing ratios in the upper troposphere.

4.1.2 Present-day ozone

A comparison with ozonesonde observations over different regions for simulated years between 1995 and 2010 is presented in Fig. 5. Besides some differences in ozone compared to observations, as discussed above, all model experiments reproduce observed tropospheric ozone within 25 % for most of the regions. At 250 hPa, which is the UTLS at mid- and high latitudes, REFC1SD overestimates ozone by up to 50 %, particularly at mid-latitudes in both hemispheres. This could be the result of strong mixing in the UTLS associated with the use of the small nudging amount of 1 % in this study; however, this needs to be investigated in more detail in future studies. The other experiments show smaller deviations from the observations of about 20 % or less. Tropical values at 50 hPa are overestimated by no more than 20 % compared

to observations for all the experiments, while ozone in the mid- and high latitudes in the stratosphere agrees within $10\,\%$ with observations.

Model results further agree well with HIPPO aircraft observations for profiles sampled from 85° N to 65° S over the Pacific Ocean between 2009 and 2011 (Fig. A2). In REFC1SD, lower troposphere values (1–2 km) are within the range of the observations, while for REFC1 and REFC2 ozone is overestimated by about 5 ppb in high northern latitudes, in particular in winter and spring, which points to a transport problem as discussed above. Some differences, especially at higher altitudes (7–8 km) are likely caused by the specific meteorological situation for the flight conditions compared to the climatological model results.

The regional performance of tropospheric ozone in the model is further illustrated in Fig. 6, comparing simulated ozone mixing ratios with ozone sondes and various aircraft observations at 3–7 km, as compiled in Tilmes et al. (2015). Observed features, for example the summertime maximum of ozone over eastern Mediterranean/Middle East

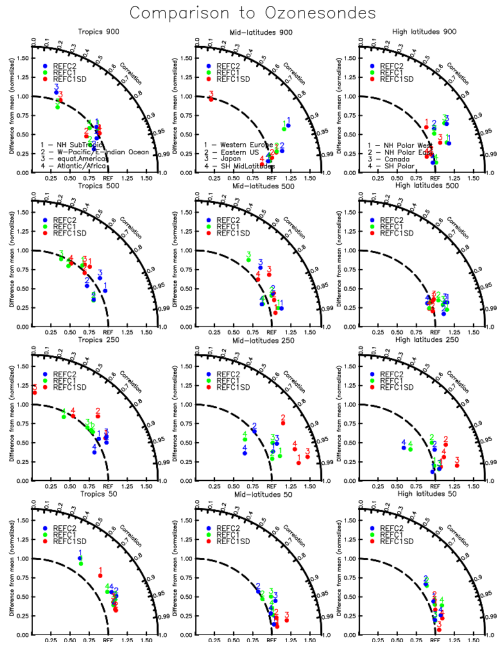


Figure 5. Taylor-like diagram comparing the mean and correlation of the seasonal cycle between observations using a present-day ozonesonde climatology between 1995 and 2011 and model results between 1995 and 2010, interpolated to the same locations as sampled by the observations and for different pressure levels, 900 hPa (top panel), 500 hPa (second panel), 250 hPa (third panel) and 50 hPa (bottom panel). Different numbers correspond to a specific region, as defined in Tilmes et al. (2012). Left panels: 1 – NH-subtropics; 2 – W-Pacific/eastern Indian Ocean; 3 – equat. Americas; 4 – Atlantic/Africa. Middle panels: 1 – western Europe; 2 – eastern USA; 3 – Japan; 4 – SH mid-latitudes. Right panels: 1 – NH polar west; 2 – NH polar east; 3 – Canada; 4 – SH polar.

(Kalabokas et al., 2013; Zanis et al., 2014), are reproduced by the REFC1 and REFC1SD experiments. The ozone gradient between mid-latitudes and tropics is for the most part well captured, for example over Japan in summer. The pole to mid-latitude ozone gradient in the SD simulation showed a larger southward ozone gradient than the REFC1 simulation, which is more consistent with the measurements. Regional differences in tropospheric ozone between the different model experiments have to be investigated in future studies.

We further perform comparisons of model results to a present-day ozone climatology based on Ozone Monitor-

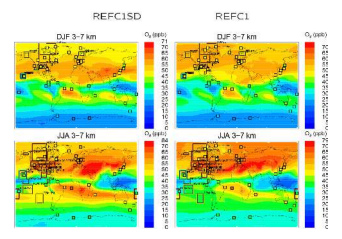


Figure 6. Comparison between model results in contours (REFC1SD left and REFC1.1 right) and observations of ozone mixing ratios, averaged over 3–7 km for December–January–February (DJF), top, and June–July–August (JJA), bottom. The color of each square represents the value of the observed ozonesonde measurement for the same period and altitude interval, and the color of framed regions corresponds to values derived from aircraft observations averaged over the particular region for each experiment (Tilmes et al., 2015).

ing Instrument (OMI) and Microwave Limb Sounder (MLS) satellite observations between 2004 and 2010, compiled by Ziemke et al. (2011), in the troposphere (Fig. 7) and stratosphere (Fig. 8). The model tropopause for this diagnostic is defined as the 150 ppb ozone level, which may lead to small differences between observations and model simulations, but not between model experiments themselves. The comparisons reveal additional characteristics of the model performance compared to observations. Tropospheric column ozone is reproduced within ± 10 DU of the observations, with a close agreement to the satellite climatology within less than ± 5 DU in low and mid-latitudes in spring and summer (Fig. 7). All model experiments show a low bias in midlatitudes in the SH and high bias by 10-15 DU in the NH mid-latitudes in winter and fall. NH tropospheric ozone is in general large in the REFC1 and REFC2 simulations compared to the REFC1SD experiments, as discussed above.

Stratospheric ozone in all model experiments agree within ±30 DU in mid- and low latitudes compared to the satellite climatology (Fig. 8). Larger deviations from the observations occur in the NH mid- and high latitudes in winter and spring with a high bias of up to 60 DU. Ozone in the SH is within about 25 DU from the observations and is reasonably well reproduced by all model experiments, especially for the freerunning experiments.

4.2 Carbon monoxide

Carbon monoxide, non-methane hydrocarbons and nitrogen dioxides are the most important precursors to the formation

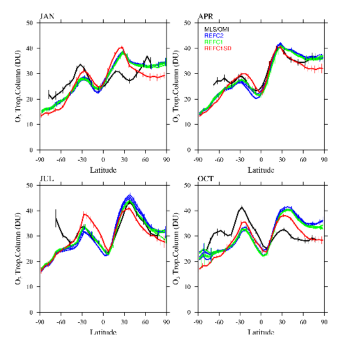


Figure 7. Monthly and zonally averaged tropospheric ozone column (in DU) comparison between OMI/MLS observations (black) and different model experiments; see legend (for ozone < 150 ppb in the model) for 4 months. Error bars describe the zonally averaged 2 sigma 6-year root mean square standard error of the mean at a giving grid point, derived from the 10° N to 10° S gridded product (Ziemke et al., 2011). Model results are interpolated to the same grid and error bars indicate the standard deviation of the interannual variability per latitude interval.

of tropospheric ozone. Carbon monoxide also impacts the oxidation capacity of the atmosphere and therefore methane lifetime. We compare the CO burden from different experiments to monthly and zonally averaged tropospheric column carbon monoxide derived from Measurements of Pollution in the Troposphere (MOPITT) version 6 level 3 satellite observations, as described in Tilmes et al. (2015) (see Fig. 9). The climatological averaging kernel and a priori is applied to both observations and model experiments in the same way.

The most obvious difference between observations and model results occurs in NH winter and spring. All model experiments are biased low by about one-third relative to observations, similar to results from the Atmospheric Chemistry and Climate Model Intercomparison Project (ACCMIP) (Naik et al., 2013; Lamarque et al., 2012). In summer and fall, the CO representation differs between different experiments, in agreement with differences in biogenic emissions. The lowest CO burden is simulated for the REFC1SD experiment, which also shows the lowest emissions of VOCs in summer (see Fig. 1). This may translate into lower CO values in fall. Furthermore, the tropospheric OH burden is significantly larger in REFC1SD compared to the other ex-

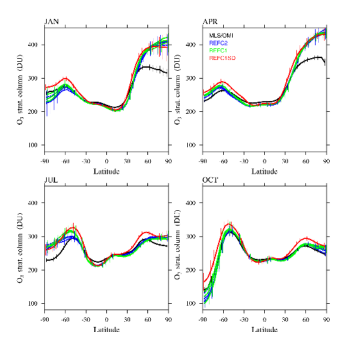


Figure 8. As Fig. 7, but showing monthly and zonally averaged stratospheric ozone column comparison between OMI/MLS observations (black) and different model experiments; see legend (for ozone > 150 ppb in the model) for 4 months.

periments (not shown), which is consistent with more ozone in the tropical troposphere (see Fig. 2).

The simulated CO column in the tropics agrees with the satellite climatology within the interannual variability. However, the model underestimates CO column in the SH for all the experiments, in particular in summer. In contrast, comparisons to HIPPO CO in situ observations indicate very good agreement between CO mixing ratios in the SH over the remote region of the Pacific Ocean for most of the seasons (see Fig. A3). Furthermore, CO mixing ratios are largely underestimated in March and April in comparison to the aircraft observations, consistent with the satellite comparison. Differences in CO will be investigated in more detail in future studies.

4.3 Hydrocarbons

Hydrocarbons are important tropospheric compounds that are emitted from vegetation, biomass burning and anthropogenic sources, including oil and gas extraction activities. They are important ozone precursors, influence the oxidation capacity of the atmosphere, and eventually form CO.

Ethane and other hydrocarbons have been measured using canister samples along coastal and island sites in the Pacific Ocean since 1984 typically every 3 months, December, March, June and September (Simpson et al., 2012); data are available at http://cdiac.ornl.gov/trends/otheratg/blake/blake. html. We have compiled a climatology using ethane mixing ratios between 1995 and 2010 that covers latitudes between

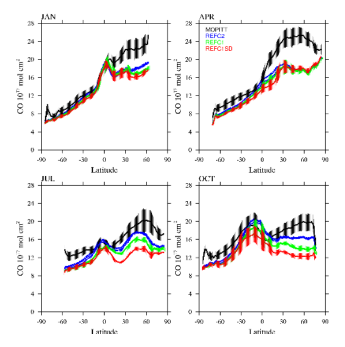


Figure 9. Monthly and zonally averaged tropospheric CO column comparison (in molec. cm⁻²) between MOPITT satellite observations (black) and different model experiments; see legend (for ozone < 150 ppb in the model) for 4 months. Error bars for observations and model experiments show the standard deviation of the interannual variability per latitude interval.

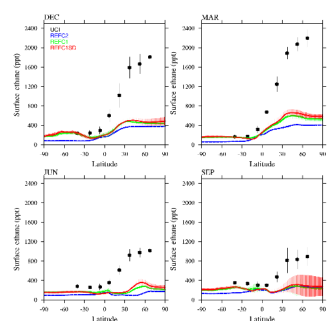


Figure 10. Comparison of observed and modeled surface ethane (C_2H_6) mixing ratios in each season averaged over 1995–2010 along the length of the Pacific Ocean. Monthly mean CAM4-chem ethane mixing ratios at 190° E are shown for the three model experiments.

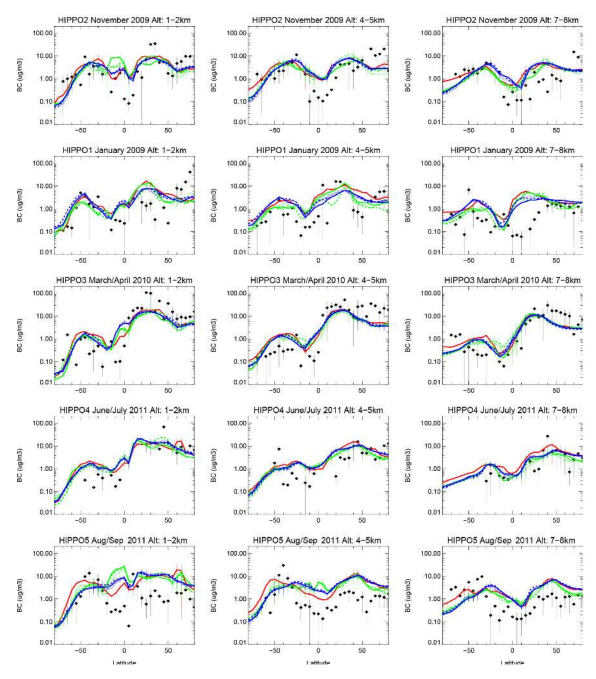


Figure 11. Black carbon comparison between different HIPPO aircraft campaigns taken over the Pacific Ocean (black symbols) and results from the reference simulations REFC1SD (red), REFC1 (green), REFC2 (blue), averaged over different altitude intervals. The sampled aircraft profiles during different HIPPO campaigns were averaged over 5° latitude intervals along the flight path over the Pacific Ocean and compared to model output averaged over the same grid points, as done in Tilmes et al. (2015). The average profiles are averaged over three altitudes regions, 1-2, 4-5 and 7-8 km.

50° S and 75° N (shown in Fig. 10). Comparisons to the three model experiments reveal a very large underestimation of ethane mixing ratios by up to 80% in spring. The smallest deviations occur in NH fall. These deviations are likely contributing to the underestimation of CO and overestimation of OH.

While there is significant uncertainty in the speciation of VOC emissions (e.g., Li et al., 2014), which could lead to this discrepancy, it is likely there is an underestimation of all VOC emissions. Globally, ethane concentrations have been declining since long-term global record keeping began. Simpson et al. (2012) reported a 21 % decline in global ethane concentrations from 1984 to 2010, which is much smaller than the discrepancy between the model and observations.

4.4 Aerosols

A reasonable description of aerosols in climate models, including interactions with chemistry and clouds, is important for the representation of radiative processes. The aerosol optical depth, global aerosol burden of organic matter, black carbon and sulfate aerosol are global diagnostics to evaluate the general performance of aerosol processes (Table 1). This version of CAM4-chem produces values for these diagnostics very similar to earlier model studies using CAM4-chem (e.g., Tilmes et al., 2015). Here, we focus on the evaluation of background black carbon in comparison to HIPPO observations. The HIPPO campaign between 2009 and 2011 provided a comprehensive data set of black carbon over the remote region of the Pacific. Black carbon results from the model are averaged over the same locations, and altitude levels and compared to the observations, as described above.

All model simulations show a very similar distribution (Fig. 11), with only a few deviations from each other mostly in the SH. The model reproduces BC values in the SH and NH mid-latitudes for most seasons within the range of uncertainty. A significant high bias in BC occurs in the tropics for all altitude levels and most seasons. Otherwise, in spring and summer, the hemispheric gradient of BC is represented well, following the observed larger burden in the NH compared to the SH, with some overestimation in the SH. The largest BC values in the NH spring are however underestimated. On the other hand, BC values in August/September, and partly November, are overestimated in the NH and in March/April and June/July in the SH.

5 Conclusions

The CESM1 CAM4-chem model has been used to perform the CCMI reference and sensitivity simulations. This paper provides an overview of the model setup of the reference experiments, including a detailed description of new developments. The most important improvements of the model beyond what has been discussed in earlier studies (Lamarque et al., 2012; Tilmes et al., 2015) are the treatment of stratospheric aerosols and the corresponding radiation and optics, which is important for the free-running experiments (Neely et al., 2015). Further, the chemistry scheme has been updated to reaction rates of JPL 2010, and improved polar chemistry has been implemented (Wegner et al., 2013; Solomon et al., 2015). A new gravity wave description, while implemented incorrectly in the code, led to an improved representation of the evolution of polar stratospheric ozone in the SH. The updated dry deposition scheme by Val Martin et al. (2014) resulted in a much improved ozone near the surface, as also shown in Tilmes et al. (2015), and leads to a very good representation of ozone mixing ratios and trends in the REFC1SD simulation.

Global model diagnostics are investigated and a selected evaluation of key chemical species, including ozone, carbon monoxide, hydrocarbons and black carbon is performed. We limit our evaluation to present-day results of the REFC1SD, REFC1 and REFC2 experiments. Comparisons to observations are focused mostly on the troposphere. Nevertheless, stratospheric column ozone reproduces observed values, in particular in SH winter and spring, but overestimates values in the NH high latitudes.

For the troposphere, near-surface ozone mixing ratios and trends are very well reproduced and within 25 % of the values from ozonesonde and satellite observations throughout the troposphere. A high bias in mid- and high northern latitudes for the REFC1 and REFC2 experiments can be explained by a stronger influence of stratospheric air masses compared to the REFC1SD simulation. This points to shortcomings in the stratosphere—troposphere exchange in the free-running simulations. On the other hand, the specified dynamics model experiment shows an overestimation of ozone in mid-latitude UTLS, as well as enhanced ozone in the upper tropical troposphere compared to the free-running experiments. The impact of shortcomings in the dynamical description of the model needs to be investigated in multi-model comparison studies.

Some biases in the model have not been resolved compared to earlier versions of the model (e.g., Tilmes et al., 2015). CO is still biased low in all model experiments in the NH, especially in spring. Some differences between the experiments may be attributed to differences in biogenic emissions. Correspondingly, methane lifetime is low compared to observational estimates, which is likely related not only to shortcomings in emissions, but also to too large an oxidation capacity of the atmosphere. Significant shortcomings of hydrocarbons (shown for ethane) are identified in particular in the NH. The hemispheric gradient of BC in the model is reproduced well in most seasons, while the fall and winter values in mid-latitudes are often overestimated in mid-latitudes. BC in the tropics is largely overestimated for most seasons. This points to potential shortcomings in emissions, but also loss processes in the model.

Code and data availability

The model code of the documented simulations is based on the Community Earth System Model, CESM version 1.1.1 (CESM1); http://www.cesm.ucar.edu/models/cesm1.1/index.html. Modifications to the model code will be documented at http://www2.cesm.ucar.edu/models/scientifically-supported. The data of the simulations are available for download at the NCAR Earth System Grid (ESG) (https://www.earthsystemgrid.org/home.html) and are submitted to the British Atmospheric Data Center (BADC) database for the CCMI project.

Appendix A

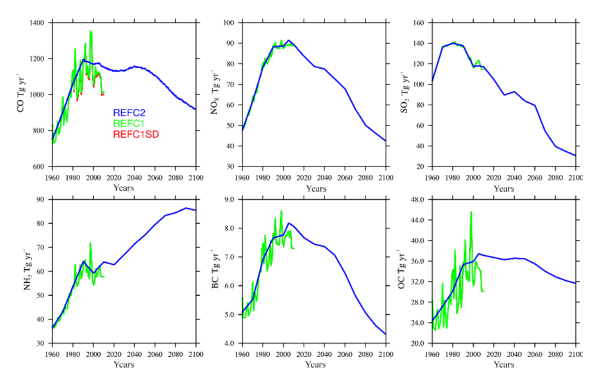


Figure A1. Selected surface emissions used for the different reference experiments.

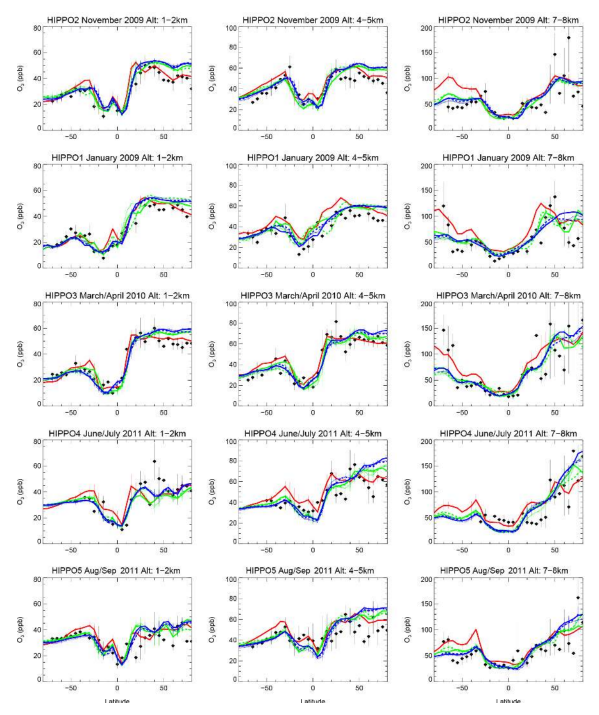


Figure A2. O₃ comparison between different HIPPO aircraft campaigns taken over the Pacific Ocean (black symbols) and results from the reference simulations REFC1SD (red), REFC1 (green), REFC2 (blue), averaged over different altitude intervals. The sampled aircraft profiles during different HIPPO campaigns were averaged over 5° latitude intervals along the flight path over the Pacific Ocean and compared to model output averaged over the same grid points, as done in Tilmes et al. (2015). The average profiles are averaged over three altitudes regions, 1-2, 4-5 and 7-8 km.

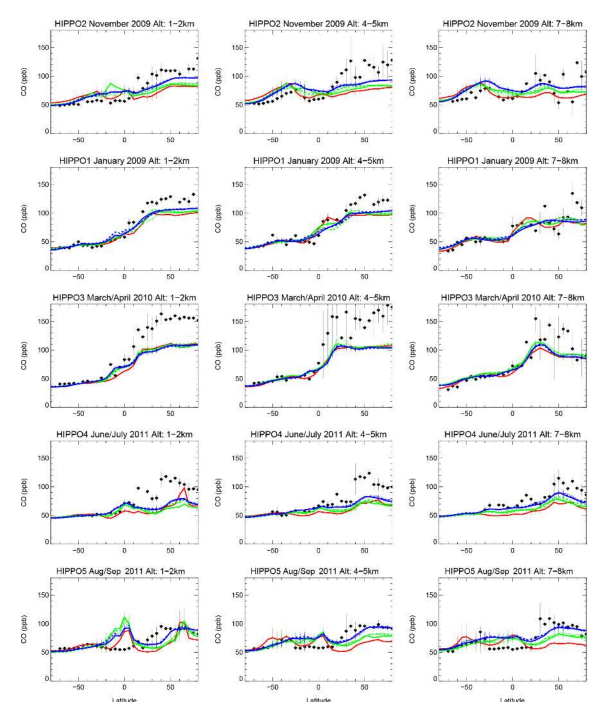


Figure A3. As Fig. A2, but for carbon monoxide.

Table A1. Chemical species in CAM4-chem, chemical formula, solver (either explicit (E) or semi-implicit (I)), lower boundary conditions (LBC), and wet and dry deposition of species.

No.	Species	Formula	Solver	Emis.	LBC	Wet dep.	Dry dep.
1	ALKO2	(C5H11O2)	I				
2	ALKOOH	(C5H12O2)	I			X	X
3	BENO2	(C6H7O3)	I				
4	BENOOH	(C6H8O3)	I				
5	BENZENE	(C6H6)	I	X			
6	BIGALD	(C5H6O2)	I	X			
7	BIGALK	(C5H12)	I	X			
8	BIGENE	(C4H8)	I	X			
9	BR	(Br)	I				
10	BRCL	(BrCl)	I				
11	BRO	(BrO)	I				
12	BRONO2	(BrONO2)	I			X	
13	BRY		E				
14	C10H16		I	X			
15	C2H2		I	X			
16	C2H4		I	X			
17	C2H5O2		I				
18	C2H5OH		I	X		X	X
19	C2H5OOH		I			X	X
20	C2H6		I	X			
21	C3H6		I	X			
22	C3H7O2		I				
23	СЗН7ООН		I			X	X
24	СЗН8		I	X			
25	CCL4	(CCl4)	E		X		
26	CF2CLBR	(CF2ClBr)	E		X		
27	CF3BR	(CF3Br)	E		X		
28	CFC11	(CFCl3)	E		X		
29	CFC113	(CCl2FCClF2)	E		X		
30	CFC114	(CCIF2CCIF2)	E		X		
31	CFC115	(CCIF2CF3)	E		X		
32	CFC12	(CF2Cl2)	E		X		
33	CH2BR2	(CH2Br2)	E		X		
34	CH2O	(CH2B12)	I	X	21	X	X
35	CH3BR	(CH3Br)	Ē	21	X	21	21
36	CH3CCL3	(CH3CCl3)	E		X		
37	СНЗСНО	(CHSCCIS)	I	X	21	X	X
38	CH3CL	(CH3Cl)	E	71	X	Λ	Α
39	CH3CN	(CH3CI)	I	X	71	X	X
40	CH3CO3		I	Λ		Λ	Λ
41	CH3COCH3		I	X		X	X
42	СНЗСОСНО		I	Λ		X	X
42	СНЗСОСНО		I	X		Λ	X
43 44	СНЗСООН		I	Λ		X	X X
44 45						Λ	Λ
	CH3O2		I	v		\mathbf{v}	v
46 47	CH3OH		I	X		X	X
47 49	СН3ООН		I		v	X	X
48	CHPB2	(CHD.,2)	E		X		
49 50	CHBR3	(CHBr3)	E		X		
50	CL	(Cl)	I				

Table A1. Continued.

No.	Species	Formula	Solver	Emis.	LBC	Wet dep.	Dry dep.
51	CL2	(Cl2)	I				
52	CL2O2	(Cl2O2)	I				
53	CLO	(ClO)	I				
54	CLONO2	(ClONO2)	I			X	
55	CLY		E				
56	CO		I	X			X
57	CO2		E		X		
58	CRESOL	(C7H8O)	I				
59	DMS	(CH3SCH3)	I	X			
60	ENEO2	(C4H9O3)	I				
61	EO	(HOCH2CH2O)	I				
62	EO2	(HOCH2CH2O2)	I				
63	EOOH	(HOCH2CH2OOH)	I			X	X
64	GLYALD	(HOCH2CHO)	I			X	X
65	GLYOXAL	(C2H2O2)	I				
66	Н		I				
67	H1202	(CBr2F2)	Е		X		
68	H2		I		X		
69	H2402	(CBrF2CBrF2)	Е		X		
70	H2O	,	I				
71	H2O2		I			X	X
72	HBR	(HBr)	I			X	
73	HCFC141B	(CH3CCl2F)	Е		X		
74	HCFC142B	(CH3CClF2)	Е		X		
75	HCFC22	(CHF2Cl)	E		X		
76	HCL	(HCl)	I			X	
77	HCN	(====)	Ī	X		X	X
78	НСООН		Ī	X		X	X
79	HNO3		I			X	X
80	HO2		Ī				
81	HO2NO2		Ī			X	X
82	HOBR	(HOBr)	I			X	
83	HOCH2OO	(11021)	I				
84	HOCL	(HOCl)	Ī			X	
85	HYAC	(CH3COCH2OH)	I			X	X
86	HYDRALD	(HOCH2CCH3CHCHO)	I			X	X
87	ISOP	(C5H8)	I	X		∡ 1.	21
88	ISOPNO3	(CH2CHCCH3OOCH2ONO2)	I	71		X	
89	ISOPO2	(HOCH2COOCH3CHCH2)	I			Λ	
90	ISOPOOH	(HOCH2COOHCH3CHCH2)	I			X	X
91	MACR	(CH2CCH3CHO)	I			X	Λ
92	MACRO2	(CH3COCHO2CH2OH)	I			Λ	
93	MACRO2 MACROOH	(CH3COCHO2CH2OH) (CH3COCHOOHCH2OH)	I			X	X
93 94	MCO3	(CH2CCH3CO3)	I			Λ	Λ
9 4 95	MEK	(C4H8O)	I	X			
				Λ			
96 07	MEKO2	(C4H7O3)	I			\mathbf{v}	v
97	MEKOOH	(C4H8O3)	I			X	X
98	MPAN MVK	(CH2CCH3CO3NO2)	I			\mathbf{v}	X
99	MVK	(CH2CHCOCH3)	I			X	
100	N		I				

Table A1. Continued.

No.	Species	Formula	Solver	Emis.	LBC	Wet dep.	Dry dep.
101	N2O		Е		X		
102	N2O5		I				
103	NH3		I	X		X	X
104	NO		I	X			X
105	NO2		I				X
106	NO3		I				
107	O		I				
108	O1D	(O)	I				
109	O3		I				X
110	OCLO	(OClO)	I				
111	OH		I				
112	ONIT	(CH3COCH2ONO2)	I			X	X
113	ONITR	(CH2CCH3CHONO2CH2OH)	I			X	X
114	PAN	(CH3CO3NO2)	I				X
115	PO2	(C3H6OHO2)	I				
116	POOH	(СЗН6ОНООН)	I			X	X
117	RO2	(CH3COCH2O2)	I				
118	ROOH	(CH3COCH2OOH)	I			X	X
119	SF6		E		X		
120	SO2		I	X		X	X
121	SOGB	(C6H7O3)	I			X	X
122	SOGI	(CH3C4H9O4)	I			X	X
123	SOGM	(C10H16O4)	I			X	X
124	SOGT	(C7H9O3)	I			X	X
125	SOGX	(C8H11O3)	I			X	X
126	TERPO2	(C10H17O3)	I				
127	TERPOOH	(C10H18O3)	I			X	X
128	TOLO2	(C7H9O5)	I				
129	TOLOOH	(C7H10O5)	I			X	X
130	TOLUENE	(C7H8)	I	X			
131	XO2	(HOCH2COOCH3CHOHCHO)	I				
132	XOH	(C7H10O6)	I				
133	XOOH	(НОСН2СООНСН3СНОНСНО)	I			X	X
134	XYLENE	(C8H10)	I				
135	XYLO2	(C8H11O3)	I				
136	XYLOOH	(C8H12O3)	I				

Table A1. Continued.

No.	Aerosols	Formula	Solver	Emis.	LBC	Wet dep.	Dry dep.
1	CB1	(C), hydrophobic BC	I	X		X	
2	CB2	(C) hydrophilic BC	I	X		X	
3	NH4		I				NH4
4	NH4NO3		I				X
5	OC1	(C), hydrophobic OC	I	X			X
6	OC2	(C) hydrophilic OC	I	X			X
7	DST01	(AlSiO5)	I				
8	DST02	(AlSiO5)	I				
9	DST03	(AlSiO5)	I				
10	DST04	(AlSiO5)	I				
11	SO4		I				X
12	SOAB	(C6H7O3)	I				X
13	SOAI	(CH3C4H9O4)	I				X
14	SOAM	(C10H16O4)	I				X
15	SOAT	(C7H9O3)	I				X
16	SOAX	(C8H11O3)	I				X
17	SSLT01	(NaCl)	I				
18	SSLT02	(NaCl)	I				
19	SSLT03	(NaCl)	I				
20	SSLT04	(NaCl)	I				
No.	Artificial tracers	Formula	Solver	Emis.	LBC	Wet dep.	Dry dep.
1	AOA_{NH}	(H)	E				
2	CO_{25}	(CO)	E	X			
3	CO_{50}	(CO)	E	X			
4	E90	(CO)	E	X			
5	E90 _{NH}	(CO)	E	X			
6	E90 _{SH}	(CO)	E	X			
7	NH ₅	(H)	E				
8	NH_{50}	(H)	E				
9	$NH_{50}W$	(H)	E			X	
10	O3S	(O3)	E				
11	SF6em	(SF6)	E	X			
12	SO2t	(SO2)	E			X	
13	ST80 ₂ 5	(H)	E				

Table A2. Chemical reactions in CAM4-chem.

Photolysis
$O2 + hv \rightarrow 2*O$
$O3 + hv \rightarrow O1D + O2$
$O3 + hv \rightarrow O + O2$
$N2O + hv \rightarrow O1D + N2$
$NO + hv \rightarrow N + O$
$NO2 + hv \rightarrow NO + O$
$N2O5 + hv \rightarrow NO2 + NO3$
$N2O5 + hv \rightarrow NO + O + NO3$
$HNO3 + hv \rightarrow NO2 + OH$
$NO3 + hv \rightarrow NO2 + O$
$NO3 + hv \rightarrow NO + O2$
$HO2NO2 + hv \rightarrow OH + NO3$
$HO2NO2 + hv \rightarrow NO2 + HO2$
$CH3OOH + hv \rightarrow CH2O + H + OH$
$CH2O + hv \rightarrow CO + 2*H$
$CH2O + hv \rightarrow CO + H2$
$H2O + hv \rightarrow OH + H$
$H2O + hv \rightarrow H2 + O1D$
$H2O + hv \rightarrow 2*H + O$
$H2O2 + hv \rightarrow 2*OH$
$CL2 + hv \rightarrow 2*CL$
$CLO + hv \rightarrow CL + O$
$OCLO + hv \rightarrow O + CLO$
$CL2O2 + hv \rightarrow 2*CL$
$HOCL + hv \rightarrow OH + CL$
$HCL + hv \rightarrow H + CL$
$CLONO2 + hv \rightarrow CL + NO3$
$CLONO2 + hv \rightarrow CLO + NO2$
$BRCL + hv \rightarrow BR + CL$
$BRO + hv \rightarrow BR + O$
$HOBR + hv \rightarrow BR + OH$
$HBR + hv \rightarrow BR + H$
$BRONO2 + hv \rightarrow BR + NO3$
$BRONO2 + hv \rightarrow BRO + NO2$
$CH3CL + hv \rightarrow CL + CH3O2$
$CCL4 + hv \rightarrow 4*CL$
$CH3CCL3 + hv \rightarrow 3*CL$
$CFC11 + hv \rightarrow 3*CL$
$CFC12 + hv \rightarrow 2*CL$
$CFC113 + hv \rightarrow 3*CL$
$HCFC22 + hv \rightarrow CL$
$CFC114 + hv \rightarrow 2*CL$
$CFC115 + hv \rightarrow CL$

Table A2. Continued.

```
Photolysis
HCFC141B + hv \rightarrow 2*CL
HCFC142B + hv \rightarrow CL
CH3BR + hv \rightarrow BR + CH3O2
CF3BR + hv \rightarrow BR
H1202 + hv \rightarrow 2*BR
H2402 + hv \rightarrow 2*BR
CF2CLBR + hv \rightarrow BR + CL
CHBR3 + hv \rightarrow 3*BR
CH2BR2 + hv \rightarrow 2*BR
CO2 + hv \rightarrow CO + O
CH4 + hv \rightarrow H + CH3O2
CH4 + hv \rightarrow 1.44*H2 + 0.18*CH2O + 0.18*O + 0.33*OH + 0.33*H
+0.44*CO2 + 0.38*CO + 0.05*H2O
CH3CHO + hv \rightarrow CH3O2 + CO + HO2
POOH + hv \rightarrow CH3CHO + CH2O + HO2 + OH
CH3COOOH + hv \rightarrow CH3O2 + OH + CO2
PAN + hv \rightarrow .6*CH3CO3 + .6*NO2 + .4*CH3O2 + .4*NO3 + .4*CO2
MPAN + hv \rightarrow MCO3 + NO2
MACR + hv \rightarrow 1.34*HO2 + .66*MCO3 + 1.34*CH2O + 1.34*CH3CO3
MACR + hv \rightarrow .66*HO2 + 1.34*CO
MVK + hv \rightarrow .7*C3H6 + .7*CO + .3*CH3O2 + .3*CH3CO3
C2H5OOH + hv \rightarrow CH3CHO + HO2 + OH
EOOH + hv \rightarrow EO + OH
C3H7OOH + hv \rightarrow 0.82*CH3COCH3 + OH + HO2
ROOH + hv \rightarrow CH3CO3 + CH2O + OH
CH3COCH3 + hv \rightarrow CH3CO3 + CH3O2
\text{CH3COCHO} + hv \rightarrow \text{CH3CO3} + \text{CO} + \text{HO2}
XOOH + hv \rightarrow OH
ONITR + hv \rightarrow HO2 + CO + NO2 + CH2O
ISOPOOH + hv \rightarrow .402*MVK + .288*MACR + .69*CH2O + HO2
HYAC + hv \rightarrow CH3CO3 + HO2 + CH2O
GLYALD + hv \rightarrow 2*HO2 + CO + CH2O
MEK + hv \rightarrow CH3CO3 + C2H5O2
BIGALD + hv \rightarrow .45*CO + .13*GLYOXAL + .56*HO2 + .13*CH3CO3
+ .18*CH3COCHO
GLYOXAL + hv \rightarrow 2*CO + 2*HO2
ALKOOH + hv \rightarrow .4*CH3CHO + .1*CH2O + .25*CH3COCH3 + .9*HO2 + .8*MEK
MEKOOH + hv \rightarrow OH + CH3CO3 + CH3CHO
TOLOOH + hv \rightarrow OH + .45*GLYOXAL + .45*CH3COCHO + .9*BIGALD
TERPOOH + hv \rightarrow OH + .1*CH3COCH3 + HO2 + MVK + MACR
SF6 + hv \rightarrow sink
SF6em + hv \rightarrow sink
```

Table A2. Continued.

Odd oxygen reactions	Rate
$O + O2 + M \rightarrow O3 + M$	6.E-34*(300/T)**2.4
$O + O3 \rightarrow 2*O2$	8.00E-12*exp(-2060./t)
$O + O + M \rightarrow O2 + M$	2.76E-34*exp(720./t)
Odd oxygen reactions (O1D only)	
$O1D + N2 \rightarrow O + N2$	2.15E-11*exp(110./t)
$O1D + O2 \rightarrow O + O2$	3.30E-11*exp(55./t)
$O1D + H2O \rightarrow 2*OH$	1.63E-10*exp(60./t)
$O1D + N2O \rightarrow 2*NO$	7.25E-11*exp(20./t)
$O1D + N2O \rightarrow N2 + O2$	4.63E-11*exp(20./t)
$O1D + O3 \rightarrow O2 + O2$	1.20E-10
O1D + CFC11 \rightarrow 3*CL	2.02E-10
O1D + CFC12 \rightarrow 2*CL	1.20E-10
O1D + CFC113 \rightarrow 3*CL	1.50E-10
O1D + CFC114 \rightarrow 2*CL	9.75E-11
$O1D + CFC115 \rightarrow CL$	1.50E-11
O1D + HCFC22 \rightarrow CL	7.20E-11
O1D + HCFC141B \rightarrow 2*CL	1.79E-10
$O1D + HCFC142B \rightarrow CL$	1.63E-10
$O1D + CCL4 \rightarrow 4*CL$	2.84E-10
$O1D + CH3BR \rightarrow BR$	1.67E-10
$O1D + CF2CLBR \rightarrow CL + BR$	9.60E-11
$O1D + CF3BR \rightarrow BR$	4.10E-11
$O1D + H1202 \rightarrow 2*BR$	1.01E-10
$O1D + H2402 \rightarrow 2*BR$	1.20E-10
O1D + CHBR3 \rightarrow 3*BR	4.49E-10
$O1D + CH2BR2 \rightarrow 2*BR$	2.57E-10
$O1D + CH4 \rightarrow CH3O2 + OH$	1.31E-10
$O1D + CH4 \rightarrow CH2O + H + HO2$	3.50E-11
$O1D + CH4 \rightarrow CH2O + H2$	9.00E-12
$O1D + H2 \rightarrow H + OH$	1.20E-10
$O1D + HCL \rightarrow CL + OH$	1.50E-10
$O1D + HBR \rightarrow BR + OH$	1.20E-10
$O1D + HCN \rightarrow OH$	7.70E-11*exp(100./t)
Odd hydrogen reactions	
$H + O2 + M \rightarrow HO2 + M$	ko=4.40E-32*(300/t)**1.30
32 · 22 · 22 32 · 22	ki=7.50E-11*(300/t)**-0.20
	f=0.60
$H + O3 \rightarrow OH + O2$	1.40E-10*exp(-470./t)
$H + HO2 \rightarrow 2*OH$	7.20E-11
$H + HO2 \rightarrow E O11$ $H + HO2 \rightarrow H2 + O2$	6.90E-12
$H + HO2 \rightarrow H2O + O$	1.60E-12
$OH + O \rightarrow H + O2$	1.80E-11*exp(180./t)
$OH + O3 \rightarrow HO2 + O2$	1.70E-12*exp(-940./t)
$OH + HO2 \rightarrow H2O + O2$	4.80E-11*exp(250./t)
$OH + OH \rightarrow H2O + O$	1.80E-12
$OH + OH + M \rightarrow H2O2 + M$	ko=6.90E-31*(300/t)**1.00
	ki=2.60E-11

Table A2. Continued.

OH + H2 → H2O + H	2.80E-12*exp(-1800./t)
$OH + H2O2 \rightarrow H2O + HO2$ $OH + H2O2 \rightarrow H2O + HO2$	1.80E-12
$H2 + O \rightarrow OH + H$	1.60E-12 1.60E-11*exp(-4570./t)
$HO2 + O \rightarrow OH + O2$	3.00E-11*exp(200./t)
$HO2 + O3 \rightarrow OH + O2$ $HO2 + O3 \rightarrow OH + 2*O2$	1.00E-14*exp(-490./t)
$HO2 + O3 \rightarrow OH + 2 \cdot O2$ $HO2 + HO2 \rightarrow H2O2 + O2$	$3.0\text{E}-13*\exp(460/t)$
$HO2 + HO2 \rightarrow H2O2 + O2$	
	+ 2.1E-33 * [M] * exp (920/t)) * (1 + 1.4E-21 * [H2O] exp (2200/t)
11202 - 0 - 011 - 1102	
$H2O2 + O \rightarrow OH + HO2$ $HCN + OH + M \rightarrow HO2 + M$	1.40E-12*exp(-2000./t) ko=4.28E-33
$HCN + OH + M \rightarrow HOZ + M$	
	ki=9.30E-15*(300/t)**-4.42
$CH3CN + OH \rightarrow HO2$	f=0.80 7.80E-13*exp(-1050./t)
Odd nitrogen reactions	THOSE TO CAP(TOCOMO)
	1.505 11* (2000 4)
$N + O2 \rightarrow NO + O$	1.50E-11*exp(-3600./t)
$N + NO \rightarrow N2 + O$	2.10E-11*exp(100./t)
$N + NO2 \rightarrow N2O + O$	2.90E-12*exp(220./t)
$N + NO2 \rightarrow 2*NO$	1.45E-12*exp(220./t)
$N + NO2 \rightarrow N2 + O2$	1.45E-12*exp(220./t)
$NO + O + M \rightarrow NO2 + M$	ko=9.00E-32*(300/t)**1.50
	ki=3.00E-11
	f=0.60
$NO + HO2 \rightarrow NO2 + OH$	3.30E-12*exp(270./t)
$NO + O3 \rightarrow NO2 + O2$	3.00E-12*exp(-1500./t)
$NO2 + O \rightarrow NO + O2$	5.10E-12*exp(210./t)
$NO2 + O + M \rightarrow NO3 + M$	ko=2.50E-31*(300/t)**1.80
	ki=2.20E-11*(300/t)**0.70
	f=0.60
$NO2 + O3 \rightarrow NO3 + O2$	1.20E-13*exp(-2450./t)
$NO2 + NO3 + M \rightarrow N2O5 + M$	ko=2.00E-30*(300/t)**4.40
	ki=1.40E-12*(300/t)**0.70
	f=0.60
$N2O5 + M \rightarrow NO2 + NO3 + M$	k(NO2 + NO3 + M)
	* 3.704E26 * exp(-11000./t)
$NO2 + OH + M \rightarrow HNO3 + M$	ko=1.80E-30*(300/t)**3.00
	ki=2.80E-11
	f=0.60
$HNO3 + OH \rightarrow NO3 + H2O$	k0 + k3[M]/(1 + k3[M]/k2)
	k0 = 2.4E-14*exp(460/t)
	k2 = 2.7E-17*exp(2199/t)
	k3 = 6.5E-34*exp(1335/t)
$NO3 + NO \rightarrow 2*NO2$	1.50E-11*exp(170./t)
$NO3 + O \rightarrow NO2 + O2$	1.00E-11
$NO3 + OH \rightarrow HO2 + NO2$	2.20E-11
$NO3 + HO2 \rightarrow OH + NO2 + O2$	3.50E-12
$NO2 + HO2 + M \rightarrow HO2NO2 + M$	ko=2.00E-31*(300/t)**3.40
	ki=2.90E-12*(300/t)**1.10
	f=0.60
$HO2NO2 + OH \rightarrow H2O + NO2 + O2$	1.30E-12*exp(380./t)
$HO2NO2 + M \rightarrow HO2 + NO2 + M$	k(NO2+HO2+M)
	* exp(-10900/t)/2.1E-27

Table A2. Continued.

Odd chlorine reactions	
$CL + O3 \rightarrow CLO + O2$	2.30E-11*exp(-200./t)
$CL + H2 \rightarrow HCL + H$	3.05E-11*exp(-2270./t)
$CL + H2O2 \rightarrow HCL + HO2$	1.10E-11*exp(-980./t)
$CL + HO2 \rightarrow HCL + O2$	1.40E-11*exp(270./t)
$CL + HO2 \rightarrow OH + CLO$	3.60E-11*exp(-375./t)
$CL + CH2O \rightarrow HCL + HO2 + CO$	8.10E-11*exp(-30./t)
$CL + CH4 \rightarrow CH3O2 + HCL$	7.30E-12*exp(-1280./t)
$CLO + O \rightarrow CL + O2$	2.80E-11*exp(85./t)
$CLO + OH \rightarrow CL + HO2$	7.40E-12*exp(270./t)
$CLO + OH \rightarrow HCL + O2$	6.00E-13*exp(230./t)
$CLO + HO2 \rightarrow O2 + HOCL$	2.60E-12*exp(290./t)
$CLO + CH3O2 \rightarrow CL + HO2 + CH2O$	3.30E-12*exp(-115./t)
$CLO + NO \rightarrow NO2 + CL$	6.40E-12*exp(290./t)
$CLO + NO2 + M \rightarrow CLONO2 + M$	ko=1.80E-31*(300/t)**3.40
	ki=1.50E-11*(300/t)**1.90
	f=0.60
$CLO + CLO \rightarrow 2*CL + O2$	3.00E-11*exp(-2450./t)
$CLO + CLO \rightarrow CL2 + O2$	1.00E-12*exp(-1590./t)
$CLO + CLO \rightarrow CL + OCLO$	3.50E-13*exp(-1370./t)
$CLO + CLO + M \rightarrow CL2O2 + M$	ko=1.60E-32*(300/t)**4.50
	ki=3.00E-12*(300/t)**2.00
	f=0.60
$CL2O2 + M \rightarrow CLO + CLO + M$	k(CLO+CLO+M) / (1.72E-27*exp(8649./t))
$HCL + OH \rightarrow H2O + CL$	1.80E-12*exp(-250./t)
$HCL + O \rightarrow CL + OH$	1.00E-11*exp(-3300./t)
$HOCL + O \rightarrow CLO + OH$	1.70E-13
$HOCL + CL \rightarrow HCL + CLO$	3.40E-12*exp(-130./t)
$HOCL + OH \rightarrow H2O + CLO$	3.00E-12*exp(-500./t)
$CLONO2 + O \rightarrow CLO + NO3$	3.60E-12*exp(-840./t)
$CLONO2 + OH \rightarrow HOCL + NO3$	1.20E-12*exp(-330./t)
$CLONO2 + CL \rightarrow CL2 + NO3$	6.50E-12*exp(135./t)
Odd bromine reactions	
$BR + O3 \rightarrow BRO + O2$	1.60E-11*exp(-780./t)
$BR + HO2 \rightarrow HBR + O2$	4.80E-12*exp(-310./t)
$BR + CH2O \rightarrow HBR + HO2 + CO$	1.70E-11*exp(-800./t)
$BRO + O \rightarrow BR + O2$	1.90E-11*exp(230./t)
$BRO + OH \rightarrow BR + HO2$	1.70E-11*exp(250./t)
$BRO + HO2 \rightarrow HOBR + O2$	4.50E-12*exp(460./t)
$BRO + NO \rightarrow BR + NO2$	8.80E-12*exp(260./t)
$BRO + NO2 + M \rightarrow BRONO2 + M$	ko=5.20E-31*(300/t)**3.20
	ki=6.90E-12*(300/t)**2.90
	f=0.60
$BRO + CLO \to BR + OCLO$	9.50E-13*exp(550./t)
$BRO + CLO \rightarrow BR + CL + O2$	2.30E-12*exp(260./t)
$BRO + CLO \rightarrow BRCL + O2$	4.10E-13*exp(290./t)
BRO + BRO \rightarrow 2*BR + O2	1.50E-12*exp(230./t)
$HBR + OH \rightarrow BR + H2O$	5.50E-12*exp(200./t)
$HBR + O \rightarrow BR + OH$	5.80E-12*exp(-1500./t)
$HOBR + O \rightarrow BRO + OH$	1.20E-10*exp(-430./t)
$BRONO2 + O \rightarrow BRO + NO3$	1.90E-11*exp(215./t)

Table A2. Continued.

Organic halogens reactions with Cl, OH	Rate
$CH3CL + CL \rightarrow HO2 + CO + 2*HCL$	2.17E-11*exp(-1130./t)
$CH3CL + OH \rightarrow CL + H2O + HO2$	2.40E-12*exp(-1250./t)
$CH3CCL3 + OH \rightarrow H2O + 3*CL$	1.64E-12*exp(-1520./t)
$HCFC22 + OH \rightarrow H2O + CL$	1.05E-12*exp(-1600./t)
$CH3BR + OH \rightarrow BR + H2O + HO2$	2.35E-12*exp(-1300./t)
$CH3BR + CL \rightarrow HCL + HO2 + BR$	1.40E-11*exp(-1030./t)
$HCFC141B + OH \rightarrow 2*CL$	1.25E-12*exp(-1600./t)
$HCFC142B + OH \rightarrow CL$	1.30E-12*exp(-1770./t)
$CH2BR2 + OH \rightarrow 2*BR + H2O$	2.00E-12*exp(-840./t)
$CHBR3 + OH \rightarrow 3*BR$	1.35E-12*exp(-600./t)
$CH2BR2 + CL \rightarrow 2*BR + HCL$	6.30E-12*exp(-800./t)
CHBR3 + CL \rightarrow 3*BR + HCL	4.85E-12*exp(-850./t)
C-1 degradation (Methane, CO, CH2O and derivatives)	
$CH4 + OH \rightarrow CH3O2 + H2O$	2.45E-12*exp(-1775./t)
$CO + OH \rightarrow CO2 + H$	ki = 2.1E09 * (t/300)**6.1
	ko = 1.5E-13 * (t/300)**0.6
	rate=ko/(1+ko/(ki/M))
	*0.6**(1/(1+log10(ko/(ki/M)**2))
$CO + OH + M \rightarrow CO2 + HO2 + M$	ko=5.90E-33*(300/t)**1.40
	ki=1.10E-12*(300/t)**-1.30
	f=0.60
$CH2O + NO3 \rightarrow CO + HO2 + HNO3$	6.00E-13*exp(-2058./t)
$CH2O + OH \rightarrow CO + H2O + H$	5.50E-12*exp(125./t)
$CH2O + O \rightarrow HO2 + OH + CO$	3.40E-11*exp(-1600./t)
$CH2O + HO2 \rightarrow HOCH2OO$	9.70E-15*exp(625./t)
$CH3O2 + NO \rightarrow CH2O + NO2 + HO2$	2.80E-12*exp(300./t)
$CH3O2 + HO2 \rightarrow CH3OOH + O2$ $CH3O2 + GH3O2 - 2*GH3O - 2*HO2$	4.10E-13*exp(750./t)
$CH3O2 + CH3O2 \rightarrow 2*CH2O + 2*HO2$	5.00E-13*exp(-424./t)
$CH3O2 + CH3O2 \rightarrow CH2O + CH3OH$	1.90E-14*exp(706./t)
$CH3OH + OH \rightarrow HO2 + CH2O$ $CH3OOH + OH \rightarrow 77CH2OO + 37CH2O + 142O$	2.90E-12*exp(-345./t)
CH3OOH + OH \rightarrow .7*CH3O2 + .3*OH + .3*CH2O + H2O	3.80E-12*exp(200./t)
$HCOOH + OH \rightarrow HO2 + CO2 + H2O$	4.50E-13
HOCH2OO → CH2O + HO2	2.40E+12*exp(-7000./t)
$HOCH2OO + NO \rightarrow HCOOH + NO2 + HO2$	2.60E-12*exp(265./t)
HOCH2OO + HO2 → HCOOH	7.50E-13*exp(700./t)
C-2 degradation	
$C2H2 + CL + M \rightarrow CL + M$	ko=5.20E-30*(300/t)**2.40
	ki=2.20E-10*(300/t)**0.70
COLLA . CL . M CL . M	f=0.60
$C2H4 + CL + M \rightarrow CL + M$	ko=1.60E-29*(300/t)**3.30
	ki=3.10E-10*(300/t)
COLIC - CL	f=0.60
$C2H6 + CL \rightarrow HCL + C2H5O2$	7.20E-11*exp(-70./t)
$C2H2 + OH + M \rightarrow .65*GLYOXAL + .65*OH + .35*HCOOH + .35*HO2$	ko=5.50E-30
+ .35*CO + M	ki=8.30E-13*(300/t)**-2.00
GALLY OLL GALLEGA 1720	f=0.60
$C2H6 + OH \rightarrow C2H5O2 + H2O$	7.66E-12*exp(-1020./t)
COLIA + CII + M + FOO + M	ko=8.60E-29*(300/t)**3.10
$C2H4 + OH + M \rightarrow EO2 + M$	ki=9.00E-12*(300/t)**0.85

Table A2. Continued.

C-2 degradation	
$EO2 + NO \rightarrow 0.5*CH2O + 0.25*HO2 + 0.75*EO + NO2$	4.20E-12*exp(180./t)
$EO2 + HO2 \rightarrow EOOH$	7.50E-13*exp(700./t)
$EO + O2 \rightarrow GLYALD + HO2$	1.00E-14
$EO \rightarrow 2*CH2O + HO2$	1.60E+11*exp(-4150./t)
$C2H4 + O3 \rightarrow CH2O + .12*HO2 + .5*CO + .12*OH + .5*HCOOH$	1.20E-14*exp(-2630./t)
$CH3COOH + OH \rightarrow CH3O2 + CO2 + H2O$	7.00E-13
$C2H5O2 + NO \rightarrow CH3CHO + HO2 + NO2$	2.60E-12*exp(365./t)
$C2H5O2 + HO2 \rightarrow C2H5OOH + O2$	7.50E-13*exp(700./t)
C2H5O2 + CH3O2 \rightarrow .7*CH2O + .8*CH3CHO + HO2 + .3*CH3OH + .2*C2H5OH	2.00E-13
$C2H5O2 + C2H5O2 \rightarrow 1.6*CH3CHO + 1.2*HO2 + .4*C2H5OH$	6.80E-14
$C2H5OOH + OH \rightarrow .5*C2H5O2 + .5*CH3CHO + .5*OH$	3.80E-12*exp(200./t)
$CH3CHO + OH \rightarrow CH3CO3 + H2O$	4.63E-12*exp(350./t)
$CH3CHO + NO3 \rightarrow CH3CO3 + HNO3$	1.40E-12*exp(-1900./t)
$CH3CO3 + NO \rightarrow CH3O2 + CO2 + NO2$	8.10E-12*exp(270./t)
$CH3CO3 + NO2 + M \rightarrow PAN + M$	ko=9.70E-29*(300/t)**5.6
	ki=9.30E-12*(300/t)**1.50
	f=0.60
$CH3CO3 + HO2 \rightarrow .75*CH3COOOH + .25*CH3COOH + .25*O3$	4.30E-13*exp(1040./t)
$CH3CO3 + CH3O2 \rightarrow .9*CH3O2 + CH2O + .9*HO2 + .9*CO2$	2.00E-12*exp(500./t)
+ .1*CH3COOH	•
$CH3CO3 + CH3CO3 \rightarrow 2*CH3O2 + 2*CO2$	2.50E-12*exp(500./t)
CH3COOOH + OH \rightarrow .5*CH3CO3 + .5*CH2O + .5*CO2 + H2O	1.00E-12
$GLYALD + OH \rightarrow HO2 + .2*GLYOXAL + .8*CH2O + .8*CO2$	1.00E-11
$GLYOXAL + OH \rightarrow HO2 + CO + CO2$	1.15E-11
$C2H5OH + OH \rightarrow HO2 + CH3CHO$	6.90E-12*exp(-230./t)
$PAN + M \rightarrow CH3CO3 + NO2 + M$	k(CH3CO3+NO2+M)
	*1.111E28 * exp(-14000/t)
$PAN + OH \rightarrow CH2O + NO3$	4.00E-14
C-3 degradation	Rate
$C3H6 + OH + M \rightarrow PO2 + M$	ko=8.00E-27*(300/t)**3.50
0010 1 011 1 11 1 1 02 1 112	ki=3.00E-11
	f=0.50
C3H6 + O3 → .54*CH2O + .19*HO2 + .33*OH + .08*CH4 + .56*CO + .5*CH3CHO + .31*CH3O2 + .25*CH3COOH	6.50E-15*exp(-1900./t)
$C3H6 + NO3 \rightarrow ONIT$	4.60E-13*exp(-1156./t)
C3H7O2 + NO \rightarrow .82*CH3COCH3 + NO2 + HO2 + .27*CH3CHO	4.20E-12*exp(180./t)
$C3H7O2 + HO2 \rightarrow C3H7OOH + O2$	7.50E-13*exp(700./t)
$C3H7O2 + CH3O2 \rightarrow CH2O + HO2 + .82*CH3COCH3$	3.75E-13*exp(-40./t)
$C3H7OOH + OH \rightarrow H2O + C3H7O2$	3.80E-12*exp(200./t)
$C3H8 + OH \rightarrow C3H7O2 + H2O$	8.70E-12*exp(-615./t)
$PO2 + NO \rightarrow CH3CHO + CH2O + HO2 + NO2$	4.20E-12*exp(180./t)
$PO2 + HO2 \rightarrow POOH + O2$	7.50E-13*exp(700./t)
$POOH + OH \rightarrow .5*PO2 + .5*OH + .5*HYAC + H2O$	•
$POOH + OH \rightarrow .5^{\circ}PO2 + .5^{\circ}OH + .5^{\circ}H \text{ YAC} + H2O$ $CH3COCH3 + OH \rightarrow RO2 + H2O$	3.80E-12*exp(200./t)
CHISCOCHS \dagger Uff \rightarrow RU2 \dagger fl2U	3.82E-11*exp(-2000/t)
DO2 + NO > CU2CO2 + CU2O + NO2	+ .33E-13
$RO2 + NO \rightarrow CH3CO3 + CH2O + NO2$ $RO2 + HO2 \rightarrow ROOH + O2$	2.90E-12*exp(300./t)
$K \cap V \to B \cap V \longrightarrow K \cap V \cap B \to V \cap V$	8.60E-13*exp(700./t)
$RO2 + RO2 \rightarrow ROO1 + O2$ $RO2 + CH3O2 \rightarrow .3*CH3CO3 + .8*CH2O + .3*HO2 + .2*HYAC$	7.10E-13*exp(500./t)

Table A2. Continued.

C-3 degradation	Rate
$ROOH + OH \rightarrow RO2 + H2O$	3.80E-12*exp(200./t)
$HYAC + OH \rightarrow CH3COCHO + HO2$	3.00E-12
CH3COCHO + OH \rightarrow CH3CO3 + CO + H2O	8.40E-13*exp(830./t)
$CH3COCHO + NO3 \rightarrow HNO3 + CO + CH3CO3$	1.40E-12*exp(-1860./t)
ONIT + OH \rightarrow NO2 + CH3COCHO	6.80E-13
C-4 degradation	
$BIGENE + OH \rightarrow ENEO2$	5.40E-11
ENEO2 + NO \rightarrow CH3CHO + .5*CH2O + .5*CH3COCH3 + HO2 + NO2	4.20E-12*exp(180./t)
$MVK + OH \rightarrow MACRO2$	4.13E-12*exp(452./t)
$MVK + O3 \rightarrow .8*CH2O + .95*CH3COCHO + .08*OH + .2*O3 + .06*HO2$	7.52E-16*exp(-1521./t)
+ .05*CO + .04*CH3CHO	
$MEK + OH \rightarrow MEKO2$	2.30E-12*exp(-170./t)
$MEKO2 + NO \rightarrow CH3CO3 + CH3CHO + NO2$	4.20E-12*exp(180./t)
$MEKO2 + HO2 \rightarrow MEKOOH$	7.50E-13*exp(700./t)
$MEKOOH + OH \rightarrow MEKO2$	3.80E-12*exp(200./t)
$MACR + OH \rightarrow .5*MACRO2 + .5*H2O + .5*MCO3$	1.86E-11*exp(175./t)
$MACR + O3 \rightarrow .8*CH3COCHO + .275*HO2 + .2*CO + .2*O3 + .7*CH2O$	4.40E-15*exp(-2500./t)
+ .215*OH	
$MACRO2 + NO \rightarrow NO2 + .47*HO2 + .25*CH2O + .53*GLYALD$	2.70E-12*exp(360./t)
+ .25*CH3COCHO + .53*CH3CO3 + .22*HYAC + .22*CO	
$MACRO2 + NO \rightarrow 0.8*ONITR$	1.30E-13*exp(360./t)
$MACRO2 + NO3 \rightarrow NO2 + .47*HO2 + .25*CH2O + .25*CH3COCHO$	2.40E-12
+ .22*CO + .53*GLYALD + .22*HYAC + .53*CH3CO3	
$MACRO2 + HO2 \rightarrow MACROOH$	8.00E-13*exp(700./t)
$MACRO2 + CH3O2 \rightarrow .73*HO2 + .88*CH2O + .11*CO + .24*CH3COCHO$	5.00E-13*exp(400./t)
+ .26*GLYALD + .26*CH3CO3 + .25*CH3OH + .23*HYAC	
$MACRO2 + CH3CO3 \rightarrow .25*CH3COCHO + CH3O2 + .22*CO + .47*HO2$	1.40E-11
+ .53*GLYALD + .22*HYAC + .25*CH2O + .53*CH3CO3	
$MACROOH + OH \rightarrow .5*MCO3 + .2*MACRO2 + .1*OH + .2*HO2$	2.30E-11*exp(200./t)
$MCO3 + NO \rightarrow NO2 + CH2O + CH3CO3$	5.30E-12*exp(360./t)
$MCO3 + NO3 \rightarrow NO2 + CH2O + CH3CO3$	5.00E-12
$MCO3 + HO2 \rightarrow .25*O3 + .25*CH3COOH + .75*CH3COOOH + .75*O2$	4.30E-13*exp(1040./t)
$MCO3 + CH3O2 \rightarrow 2*CH2O + HO2 + CO2 + CH3CO3$	2.00E-12*exp(500./t)
$MCO3 + CH3CO3 \rightarrow 2*CO2 + CH3O2 + CH2O + CH3CO3$	4.60E-12*exp(530./t)
$MCO3 + MCO3 \rightarrow 2*CO2 + 2*CH2O + 2*CH3CO3$	2.30E-12*exp(530./t)
$MCO3 + NO2 + M \rightarrow MPAN + M$	1.1E-11*300./t/[M]
$MPAN + M \rightarrow MCO3 + NO2 + M$	k(MCO3 + NO2 + M)
	* 1.111E28 * exp(-14000/t)
MPAN + OH + M \rightarrow .5*HYAC + .5*NO3 + .5*CH2O + .5*HO2	ko=8.00E-27*(300/t)**3.50
+0.5*CO2 + M	ki=3.00E-11
	f=0.50

Table A2. Continued.

C-5 degradation	
$\overline{\text{ISOP} + \text{OH} \rightarrow \text{ISOPO2}}$	2.54E-11*exp(410./t)
ISOP + O3 \rightarrow .4*MACR + .2*MVK + .07*C3H6 + .27*OH + .06*HO2	1.05E-14*exp(-2000./t)
+ .6*CH2O + .3*CO + .1*O3 + .2*MCO3 + .2*CH3COOH	**
$ISOP + NO3 \rightarrow ISOPNO3$	3.03E-12*exp(-446./t)
ISOPO2 + NO \rightarrow .08*ONITR + .92*NO2 + .23*MACR + .32*MVK	4.40E-12*exp(180./t)
+ .33*HYDRALD + .02*GLYOXAL + .02*GLYALD	
+ .02*CH3COCHO + .02*HYAC + .55*CH2O + .92*HO2	
$ISOPO2 + NO3 \rightarrow HO2 + NO2 + .6*CH2O + .25*MACR + .35*MVK$	2.40E-12
+ .4*HYDRALD	
$ISOPO2 + HO2 \rightarrow ISOPOOH$	8.00E-13*exp(700./t)
$ISOPOOH + OH \rightarrow .8*XO2 + .2*ISOPO2$	1.52E-11*exp(200./t)
$ISOPO2 + CH3O2 \rightarrow .25*CH3OH + HO2 + 1.2*CH2O + .19*MACR$	5.00E-13*exp(400./t)
+ .26*MVK + .3*HYDRALD	
$ISOPO2 + CH3CO3 \rightarrow CH3O2 + HO2 + .6*CH2O + .25*MACR$	1.40E-11
+ .35*MVK + .4*HYDRALD	
ISOPNO3 + NO \rightarrow 1.206*NO2 + .794*HO2 + .072*CH2O + .167*MACR	2.70E-12*exp(360./t)
+ .039*MVK + .794*ONITR	
ISOPNO3 + NO3 \rightarrow 1.206*NO2 + .072*CH2O + .167*MACR	2.40E-12
+ .039*MVK + .794*ONITR + .794*HO2	
ISOPNO3 + HO2 \rightarrow .206*NO2 + .206*CH2O + .206*OH + .167*MACR	8.00E-13*exp(700./t)
+ .039*MVK + .794*ONITR	
$BIGALK + OH \rightarrow ALKO2$	3.50E-12
$ONITR + OH \rightarrow HYDRALD + .4*NO2 + HO2$	4.50E-11
$ONITR + NO3 \rightarrow HO2 + NO2 + HYDRALD$	1.40E-12*exp(-1860./t)
$HYDRALD + OH \rightarrow XO2$	1.86E-11*exp(175./t)
$ALKO2 + NO \rightarrow .4*CH3CHO + .1*CH2O + .25*CH3COCH3 + .9*HO2$	4.20E-12*exp(180./t)
+ .8*MEK + .9*NO2 + .1*ONIT	
$ALKO2 + HO2 \rightarrow ALKOOH$	7.50E-13*exp(700./t)
$ALKOOH + OH \rightarrow ALKO2$	3.80E-12*exp(200./t)
$XO2 + NO \rightarrow NO2 + HO2 + .25*CO + .25*CH2O$	2.70E-12*exp(360./t)
+ .25*GLYOXAL + .25*CH3COCHO + .25*HYAC + .25*GLYALD	
$XO2 + NO3 \rightarrow NO2 + HO2 + 0.5*CO + .25*HYAC + 0.25*GLYOXAL$	2.40E-12
+ .25*CH3COCHO + .25*GLYALD	0.007.401
$XO2 + HO2 \rightarrow XOOH$	8.00E-13*exp(700./t)
$XO2 + CH3O2 \rightarrow .3*CH3OH + .8*HO2 + .8*CH2O + .2*CO$	5.00E-13*exp(400./t)
+ .1*GLYOXAL + .1*CH3COCHO + .1*HYAC + .1*GLYALD	1.20E 12*(
$XO2 + CH3CO3 \rightarrow .25*CO + .25*CH2O + .25*GLYOXAL + CH3O2$	1.30E-12*exp(640./t)
+ HO2 + .25*CH3COCHO + .25*HYAC + .25*GLYALD + CO2	1.005 12* (100 //)
$XOOH + OH \rightarrow H2O + XO2$	1.90E-12*exp(190./t)
$XOOH + OH \rightarrow H2O + OH$	T**2 * 7.69E-17 * exp(253./t)

Table A2. Continued.

C-7 degradation	Rate
TOLUENE + OH \rightarrow .25*CRESOL + .25*HO2 + .7*TOLO2	1.70E-12*exp(352./t)
TOLO2 + NO → .45*GLYOXAL + .45*CH3COCHO + .9*BIGALD	4.20E-12*exp(180./t)
+ .9*NO2 + .9*HO2	
$TOLO2 + HO2 \rightarrow TOLOOH$	7.50E-13*exp(700./t)
$TOLOOH + OH \rightarrow TOLO2$	3.80E-12*exp(200./t)
$CRESOL + OH \rightarrow XOH$	3.00E-12
$XOH + NO2 \rightarrow .7*NO2 + .7*BIGALD + .7*HO2$	1.00E-11
BENZENE + OH \rightarrow BENO2	2.30E-12*exp(-193./t)
$BENO2 + HO2 \rightarrow BENOOH$	1.40E-12*exp(700./t)
$BENO2 + NO \rightarrow 0.9*GLYOXAL + 0.9*BIGALD + 0.9*NO2 + 0.9*HO2$	2.60E-12*exp(350./t)
$XYLENE + OH \rightarrow XYLO2$	2.30E-11
$XYLO2 + HO2 \rightarrow XYLOOH$	1.40E-12*exp(700./t)
XYLO2 + NO \rightarrow 0.62*BIGALD + 0.34*GLYOXAL + 0.54*CH3COCHO + 0.9*NO2 + 0.9*HO2	2.60E-12*exp(350./t)
C-10 degradation	
$C10H16 + OH \rightarrow TERPO2$	1.20E-11*exp(444./t)
$C10H16 + O3 \rightarrow .7*OH + MVK + MACR + HO2$	1.00E-15*exp(-732./t)
$C10H16 + NO3 \rightarrow TERPO2 + NO2$	1.20E-12*exp(490./t)
TERPO2 + NO \rightarrow .1*CH3COCH3 + HO2 + MVK + MACR + NO2	4.20E-12*exp(180./t)
$TERPO2 + HO2 \rightarrow TERPOOH$	7.50E-13*exp(700./t)
TERPOOH + OH \rightarrow TERPO2	3.80E-12*exp(200./t)
Tropospheric heterogeneous reactions	
$N2O5 \rightarrow 2*HNO3$	
$NO3 \rightarrow HNO3$	
$NO2 \rightarrow 0.5*OH + 0.5*NO + 0.5*HNO3$	
$CB1 \rightarrow CB2$	7.10E-06
$SO2 + OH \rightarrow SO4$	
$DMS + OH \rightarrow SO2$	9.60E-12*exp(-234./t)
DMS + OH \rightarrow .5*SO2 + .5*HO2	
$DMS + NO3 \rightarrow SO2 + HNO3$	1.90E-13*exp(520./t)
$NH3 + OH \rightarrow H2O$	1.70E-12*exp(-710./t)
$OC1 \rightarrow OC2$	7.10E-06
$HO2 \rightarrow 0.5*H2O2$	
Stratospheric removal rates for BAM aerosols	
$CB1 \rightarrow (No products)$	6.34E-08
$CB2 \rightarrow (No products)$	6.34E-08
$OC1 \rightarrow (No products)$	6.34E-08
$OC2 \rightarrow (No \text{ products})$	6.34E-08
$SO4 \rightarrow (No products)$	6.34E-08

Table A2. Continued.

Stratospheric removal rates for BAM aero	sols
$SOAM \rightarrow (No products)$	6.34E-08
$SOAI \rightarrow (No products)$	6.34E-08
$SOAB \rightarrow (No products)$	6.34E-08
$SOAT \rightarrow (No products)$	6.34E-08
$SOAX \rightarrow (No products)$	6.34E-08
$NH4 \rightarrow (No products)$	6.34E-08
$NH4NO3 \rightarrow (No products)$	6.34E-08
$SSLT01 \rightarrow (No products)$	6.34E-08
$SSLT02 \rightarrow (No products)$	6.34E-08
$SSLT03 \rightarrow (No products)$	6.34E-08
$SSLT04 \rightarrow (No products)$	6.34E-08
$DST01 \rightarrow (No products)$	6.34E-08
$DST02 \rightarrow (No products)$	6.34E-08
$DST03 \rightarrow (No \text{ products})$	6.34E-08
$DST04 \rightarrow (No products)$	6.34E-08
$SO2t \rightarrow (No \text{ products})$	6.34E-08
Sulfate aerosol reactions	
N2O5 → 2*HNO3	f (sulfuric acid wt%)
$CLONO2 \rightarrow HOCL + HNO3$	f (T,P,HCl,H2O,r)
$BRONO2 \rightarrow HOBR + HNO3$	f (T,P,H2O,r)
$CLONO2 + HCL \rightarrow CL2 + HNO3$	f (T,P,HCl,H2O,r)
$HOCL + HCL \rightarrow CL2 + H2O$	f (T,P,HCl,HOCl,H2O,r)
$HOBR + HCL \rightarrow BRCL + H2O$	f (T,P,HCl,HOBr,H2O,r)
Nitric acid Di-hydrate reactions	
N2O5 → 2*HNO3	$\gamma = 0.0004$
$CLONO2 \rightarrow HOCL + HNO3$	$\gamma = 0.004$
$CLONO2 + HCL \rightarrow CL2 + HNO3$	$\gamma = 0.2$
$HOCL + HCL \rightarrow CL2 + H2O$	$\gamma = 0.1$
$BRONO2 \rightarrow HOBR + HNO3$	$\gamma = 0.3$
Ice aerosol reactions	
N2O5 → 2*HNO3	$\gamma = 0.02$
$CLONO2 \rightarrow HOCL + HNO3$	$\gamma = 0.3$
$BRONO2 \rightarrow HOBR + HNO3$	$\gamma = 0.3$
$CLONO2 + HCL \rightarrow CL2 + HNO3$	$\gamma = 0.3$
$HOCL + HCL \rightarrow CL2 + H2O$	$\gamma = 0.2$
$HOBR + HCL \rightarrow BRCL + H2O$	$\gamma = 0.3$
Synthetic tracer reactions	
$NH_5 \rightarrow (No \text{ products})$	2.31E-06
$NH_{50} \rightarrow (No \text{ products})$	2.31E-07
$NH_{50W} \rightarrow (No \text{ products})$	2.31E-07
$ST80_{25} \rightarrow (No \text{ products})$	4.63E-07
$CO_{25} \rightarrow (No \text{ products})$	4.63E-07
$CO_{50} \rightarrow (No \text{ products})$	2.31E-07
$E90 \rightarrow (No \text{ products})$	1.29E-07
$E90_{NH} \rightarrow (No \text{ products})$	1.29E-07
$E90_{SH} \rightarrow (No \text{ products})$	1.29E-07

Table A3. Tropospheric ozone production and loss rates calculated for explicit reaction rates, O3-Prod and O3-Loss are the sum of the specific reaction rates.

Production/loss (Tg year ⁻¹)	REFC1SD	REFC1	REFC2
O3-Prod	4701.1	4716.5	4758.0
NO-HO2	3032.2	3017.3	3051.7
CH3O2-NO	1102.1	1078.6	1072.2
PO2-NO	19.8	20.9	21.1
CH3CO3-NO	159.6	168.8	172.3
C2H5O2-NO	8.2	8.1	7.5
0.92*ISOPO2-NO	113.0	131.8	136.1
MACRO2-NOa	60.9	68.3	69.9
MCO3-NO	25.6	28.9	29.8
C3H7O2-NO	n.a.	n.a.	n.a.
RO2-NO	10.6	11.2	11.6
XO2-NO	53.6	62.4	64.1
0.9*TOLO2-NO	2.7	2.8	3.8
TERPO2-NO	15.2	16.7	16.8
0.9*ALKO2-NO	21.6	21.3	21.7
ENEO2-NO	12.0	12.4	12.5
EO2-NO	34.9	37.2	37.0
MEKO2-NO	16.4	16.1	16.7
0.4*ONITR-OH	6.0	6.8	7.1
jonitr	1.1	1.2	1.3
O3-Loss	4118.0	4128.9	4157.6
O1D-H2O	2217.8	2295.8	2290.2
OH-O3	582.2	537.6	536.7
HO2-O3	1203.5	1179.0	1202.4
C3H6-O3	11.9	11.0	12.0
0.9*ISOP-O3	51.3	51.9	59.4
C2H4-O3	7.9	8.0	8.1
0.8*MVK-O3	12.9	13.5	14.8
0.8*MACR-O3	2.4	2.4	2.7
C10H16-O3	28.2	29.6	31.4

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