Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





1

1

The high-resolution version of TM5-MP for optimised satellite retrievals: Description and Validation.

3

2

J. E. Williams<sup>1</sup>, K. F. Boersma<sup>1,2</sup>, P. Le Sager<sup>1</sup>, W. W. Verstraeten<sup>1,2,\*</sup>

[1] {KNMI, De Bilt, The Netherlands}

[2] {Meteorology and Air Quality Group, Wageningen University, Wageningen, The Netherlands}

[\*] now at: KMI, Ukkel, Brussels, Belgium

12

13

14

15

16 17

18 19

20

21

22

23

24

25

26

27

28

29

30

31

32

33

34

35

36

37

38

39

40

41

We provide a comprehensive description of the high-resolution version of the TM5-MP global Chemistry-Transport Model which is employed to provide highly resolved vertical profiles of nitrogen dioxide (NO<sub>2</sub>), formaldehyde (CH<sub>2</sub>O), and sulphur dioxide (SO<sub>2</sub>) for use in satellite retrievals from platforms such as the Ozone Monitoring Instrument (OMI) and the Sentinel-5 Precursor, the TROPOspheric Monitoring Instrument (TROPOMI). Comparing seasonal differences in the global distributions of <sup>222</sup>Rn we show that there are generally differences of ±20%, with larger increases occurring near specific coastal and decreases over specific tropical ocean regions. Analyzing vertical profiles of <sup>222</sup>Rn above strong nitrogen oxide (NO<sub>x</sub>) source regions, we show that there are differences in the strength of the convective activity of around ~2-10% (~10-20%) at 1° x 1º constrained below 700hPa (200hPa) in the NH (tropics). We analyze the global distribution and chemical budget terms for tropospheric ozone (O<sub>3</sub>), the reactive NO<sub>x</sub> and N-reservoir species from simulations performed at 1° x 1° horizontal resolution. Compared to simulations at 3° x 2°, we show that although the impact on photolysis rates may be important regionally, changes in the seasonal means representative of the boundary layer are of the order of a few percent, in spite of the higher spatial variability of meteorological data fields from ERA-Interim. Surface concentrations of O<sub>3</sub> in high-NO<sub>x</sub> regions decrease by between ~5-10% at 1° x 1° as a result of a reduction in NO<sub>x</sub> recycling terms and an increase in the titration term of O<sub>3</sub> involving NO. At 1° x 1° the net global stratosphere-troposphere exchange of ozone decreases by ~7%, dominated by substantial reductions in the Northern Hemisphere somewhat offset by increases in down-welling which occurs in the Southern Hemisphere. By comparing NO, NO2, HNO3 and PAN profiles against a host of measurements made across large regional domains, we show that TM5-MP captures the vertical distribution of NO<sub>x</sub> and long-lived NO<sub>x</sub> reservoirs at background locations, but exhibits a too high NO/NO<sub>2</sub> ratio in the Free Troposphere, with changes at 1° x 1° being limited to a few percent. Surface mixing ratios in both NO and NO2 are generally underestimated in both low and high NO<sub>x</sub> scenarios. For Europe, we show that there is a negative bias in NO concentrations at the surface across the whole domain, with lower biases at 1° x 1° at only ~20% of sites. For NO<sub>2</sub>, biases are more variable, with lower (higher) biases occurring at ~35% (~20%) of sites. For nitric acid (HNO<sub>3</sub>) there is a seasonal cycle present at the surface in TM5-MP which is not observed, leading to a low bias during wintertime and a high bias during summertime. For the shorter-lived NO<sub>x</sub> reservoirs, no significant changes occur with the exception of  $N_2O_5$  due to ~10% perturbations in the  $NO_3$  radical. For  $CH_2O$ , the impact of higher resolution on the chemical budget terms is rather modest, with changes less than 5%. The simulated vertical distribution of CH<sub>2</sub>O agrees reasonably well with measurements in pristine locations, but CH<sub>2</sub>O vertical column densities are generally underestimated relative to satellite measurements in polluted regions. For SO<sub>2</sub>, the performance at 1° x 1° is principally governed by the quality of the emission inventory, with limited

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





2

improvements in the site specific biases with most showing no significant improvement. For the vertical column, improvements near strong source regions occur which reduce the biases in the integrated column.

44 45

### 1. Introduction

46

77

78

79

80

81

47 One application of Chemistry Transport Models (CTM) is to provide accurate vertical and horizontal global 48 distributions of trace gases such as ozone (O<sub>3</sub>), nitrogen dioxide (NO<sub>2</sub>), sulphur dioxide (SO<sub>2</sub>) and formaldehyde 49 (CH<sub>2</sub>O) that are used as a-priori best-guesses in the retrievals of tropospheric abundances from earth-orbiting 50 satellites including the Tropospheric Emission Sounder (TES, Worden et al., 2007) Global Ozone Monitoring 51 Experiment (GOME), SCanning Imaging Absorption spectroMeter for Atmospheric CHartographY 52 (SCIAMACHY, De Smedt et al., 2008), the Ozone Monitoring Instrument (OMI, Boersma et al., 2011), and 53 GOME-2 (Valks et al., 2011)). To date, although high-resolution regional models have been employed for 54 selected regions such as the US and Europe (e.g. Russell et al., 2011; Zhou et al., 2012; Vinken et al., 2014), at 55 global scale the CTMs resolutions employed are still rather coarse (between 2-4° latitude and 2-6° longitude), 56 resulting in 'footprints' of hundreds of kilometers in area. This has limitations as the resulting total columns are 57 sensitive to topography, surface albedo and the shape of the a-priori vertical profiles themselves. This sampling 58 is rather coarse and leads to substantial errors in the retrievals (e.g. Boersma et al., 2007; Heckel et al., 2011; 59 Russell et al., 2011) and imposes constraints on capturing the regional scale variability in short-lived trace gas 60 abundances observed from high-resolution satellite instruments such as OMI. 61 This lack of spatial detail is particularly relevant for situations where strong spatio-temporal variability in the 62 vertical distribution of NO2, SO2, and CH2O can be expected. Examples include shipping lanes in the relatively 63 unpolluted Marine Boundary Layer (e.g. Vinken et al., 2014) and coal-fired power plant SO2 pollution (e.g. 64 Fioletov et al., 2015). Moreover, during the day the local lifetime and mixing ratios of trace gases such as NO<sub>2</sub> 65 are critically dependent on a host of variables e.g. temperature, surface albedo, cloud cover (via photolysis), 66 chemical conversion (i.e. NO/NO2 ratio) and the extent of mixing by convective upwelling (i.e. land type) and 67 advective transport. Thus, the information provided for the retrievals are all affected by the coarsening procedure 68 that is necessary in the CTM. Recently, Heckel et al. (2011) demonstrated that there is an associated uncertainty 69 of ~2 using a-priori data from a global CTM rather than a regional CTM, principally due to loss of spatial 70 information. Two other studies focusing on the impact of horizontal resolution on the retrieval of vertical 71 column densities (VCD) of NO<sub>2</sub> suggested that errors of up to ~50% exist (Yamaji et al., 2014; Lin et al., 2014). 72 This problem becomes accentuated for the next generation of earth orbiting satellites such as the Tropospheric 73 Monitoring Instrument (tropOMI), which has a smaller footprint compared to its predecessors (Veefkind et al., 74 2012). Applications of TM5 include the retrieval of NO2, CH2O, and SO2 column densities from OMI and 75 tropOMI (e.g. van Geffen et al., 2016), where studies related to the influence of horizontal resolution have been 76 limited principally to NO<sub>2</sub>.

The dominant tropospheric loss terms for CH<sub>2</sub>O are photolysis and scavenging into cloud droplets (wet deposition; Jacob, 2000). Thus the atmospheric lifetime of CH<sub>2</sub>O is highly sensitive to the extent of cloud cover and the vertical profiles of the photolysis rates. A dominant application of CH<sub>2</sub>O retrievals is for placing constraints in tropical and sub-tropical isoprene emission fluxes (e.g. Palmer et al., 2006; Stavrakou et al., 2009; Marais et al., 2012). The resulting emission estimates are highly sensitive to the stoichiometric yield of CH<sub>2</sub>O

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





from isoprene oxidation, the chemical lifetime of CH<sub>2</sub>O and spatial differences in land cover. Other applications include estimating yields released during Biomass Burning (BB) episodes (Gonzi et al., 2011), whose spatial location is also smeared via the coarsening procedure. For SO<sub>2</sub>, which predominantly originates from point sources, an adequate spatial distribution of such sources is crucial for deriving accurate biases in existing emission inventories.

In this paper we provide a comprehensive description of the global, high-resolution  $1^{\circ}$  x  $1^{\circ}$  version of the TM5 CTM tailored for the application of satellite retrievals (hereafter referred to as TM5-MP). In Sect. 2 we give details related to the modifications which have been made to the TM5 model compared to previous versions, the emission inventories employed, updates that have been made to the modified CB05 chemical mechanism, stratospheric boundary conditions, photolysis scheme, heterogeneous conversion and model structure. In Sect. 3 we analyse the impact on convective and advective transport of trace species from the boundary layer as derived using Radon distributions. In Sect. 4 we investigate the effects on regional photolysis frequencies and in Sect. 5 we examine the differences introduced for both tropospheric  $O_3$ ,  $NO_x$  and N-containing species at higher resolution and make comparisons against both surface and aircraft measurements to investigate effects. In Sect 6 we examine impact on the global  $CH_2O$  budget and integrated columns, and in Sect. 7 we show the subsequent improvement in the distribution of  $SO_2$  at  $1^{\circ}$  x  $1^{\circ}$ . In Sect. 8, we present our conclusions.

## 2 Description of TM5-MP

Previous versions of TM5 (TM5-chem-v3.0, Huijnen et al., 2010) included a two-way nested zooming option as described by Krol et al. (2005). This option allowed high-resolution simulations to be performed over any predefined regional, with boundary conditions being determined by the global simulation at coarser resolution. Typically, global simulations at  $3^{\circ}$  x  $2^{\circ}$  with zoom regions at  $1^{\circ}$  x  $1^{\circ}$  were performed to alleviate the long runtime of a global  $1^{\circ}$  x  $1^{\circ}$  run. In the new version of TM5 (hereafter referred to as TM5-MP; the massivelyntersta parallel version), the usage of MPI has been totally rewritten. Zoom regions are no longer available, but data sets are distributed along longitudes and latitudes, instead of model levels and tracers. The advantages of that overhaul towards domain decomposition are a smaller memory requirement and the possibility to use more processors making global  $1^{\circ}$  x  $1^{\circ}$  simulations feasible in terms of runtime and affordable in terms of computing resources. A TM5-MP global  $3^{\circ}$  x  $2^{\circ}$  ( $1^{\circ}$  x  $1^{\circ}$ ) run is  $\sim$ 6 ( $\sim$  20) times faster than the previous version of TM5 (Huijnen et al., 2010) for similar resources.

Here we provide a comprehensive description of the modifications and updates introduced into TM5-MP compared to TM5 v3.0 (Huijnen et al., 2010). The model is driven using the ERA-interim meteorological reanalysis (Dee et al., 2011) and updated every 3 hours, with interpolation of fields for the intermediate time periods. The convective mass fluxes are taken from ERA-interim dataset and describe the updraft velocities from the boundary layer into the free troposphere (FT), which replaces the parameterization of Tiedtke (1989). An iterative time-step is employed to prevent too much mass being transported out of any particular grid cell for any particular time-step according to the Courant-Friedrichs-Lewy (CFL) criteria (Bregman et al., 2003), which is especially relevant when reducing the size of grid-cells as done here.

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

120

121

122

123

124

125

126

127

128

129

130

131

132

133

134

135

136

137

138

139

140

141

142

143

144

145

146

147

148

149

150

151

152

153

154

155

156

157

158

159

© Author(s) 2016. CC-BY 3.0 License.





4

The gas phase chemistry in TM5-MP is described by an expanded version of the modified CB05 chemical mechanism (hereafter mCB05; Williams et al., 2013). We have placed emphasis of updating and expanding the fast NO<sub>x</sub> chemistry to account for an accurate partitioning of nitrogen for higher NO<sub>x</sub> regimes than those occurring at coarser horizontal resolutions. All reaction rate data is now taken from the latest IUPAC recommendations (sited at http://iupac.pole-ether.fr/; last access June 2016) using updated formulations for third-body collisions, where the rate data for fast NO<sub>x</sub> and CH<sub>2</sub>O chemistry is given in Table 1. This includes the recent update to the formation rate of HNO<sub>3</sub> determined by Möllner et al. (2010). The most relevant modifications are: (i) The yield of CH<sub>2</sub>O, methanol (CH<sub>3</sub>OH) and the hydro-peroxy radical (HO<sub>2</sub>) from the selftermination of the methyl-peroxy radical (CH<sub>3</sub>O<sub>2</sub>) is increased according to Yarwood et al. (2005), (ii) the direct formation of CH<sub>2</sub>O from the reaction of CH<sub>3</sub>O<sub>2</sub> + HO<sub>2</sub> is added using the temperature dependent branching ratio defined in Atkinson et al. (2004), (iii) the production of HNO<sub>3</sub> during the oxidation of di-methyl sulphide (DMS) by the NO<sub>3</sub> is now included, (iv) explicit organic peroxy radicals have been introduced as products from the oxidation of propene  $(C_3H_6)$  and propane  $(C_3H_8)$  by OH, which are lost by either the reaction with nitric oxide (NO) or HO<sub>2</sub> allowing the in-situ chemical formation of acetone (CH<sub>3</sub>COCH<sub>3</sub>) and higher aldehydes (ALD2), respectively, following the stoichiometry given in Emmons et al. (2010), (v) a second product channel for N<sub>2</sub>O<sub>5</sub> photolysis is added producing NO, (vi) the formation and photo-dissociation of HONO has been included, (vi) the formation and transport of methyl peroxy nitrate (CH<sub>3</sub>O<sub>2</sub>NO<sub>2</sub>) is also included (Browne et al., 2011), and (vii) modifications to the gas-phase chemistry involving NH<sub>3</sub> have been introduced following the stoichiometry given in Hauglestaine et al. (2014). This version of the modified CB05 chemical mechanism is hereafter referred to as mCB05v2. The calculation of height resolved photolysis rates (J values) is performed using a tailored version of the Modified Band Approach (MBA). The implementation and performance of this parameterization in TM5 has been fully described in Williams et al. (2012). For the calculation of the height-resolved Actinic Fluxes at the seven specific wavelengths used for calculating the J values (these being 205.1nm, 287.9nm, 302.0nm, 311.0nm, 326.5nm, 385.0nm and 610.0nm), the 2-stream radiative transfer solver of Zdunkowski et al. (1980) is embedded into TM5-MP. Details regarding the parameterizations used to account for the scattering and absorption introduced by gaseous molecules, aerosols and clouds the reader is referred to Williams et al. (2012). For aerosols, the climatology of Shettle and Fenn (1979) is included. The calculation of the effective radius  $(r_{eff})$ of cloud droplets is now performed using the approach of Martin et al. (1994), where different parameter values are used for over the land and ocean using cloud condensation nuclei concentrations of 40 and 900, respectively. Due to potentially erroneous values at low horizontal resolution, we weight the final  $r_{eff}$  value using the land fraction in each grid-cell. We apply limits between 4-16um on the resulting reff values. This improves the representation of the scattering component due to cloud droplets used for the calculation of the actinic flux in the lower troposphere (not shown). For the scattering effects from cloud droplets, we subsequently downsize the physical  $r_{eff}$  by ~0.5-2 $\mu$ m to account for the relationship between the optical and physical  $r_{eff}$  values. For aerosols, an aerosol scheme is available for use within TM5-MP (aan den Brugh et al., 2010), but we choose not to use it for the purpose of satellite retrievals due to the extra computational expense needed when performing high resolution simulations that would potentially hinder operational use. We acknowledge that the description of aerosols in this study is rather crude and increasing scattering could have an impact under

instances of low cloud coverage. For the application of TM5-MP towards satellite retrieval, it is preferable to use

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

196

197

198

© Author(s) 2016. CC-BY 3.0 License.





5

160 any advancements in computational performance on further increases in the horizontal resolution employed. 161 Therefore it is not currently envisaged that a full description of aerosol processes will be included during 162 operational satellite retrievals. 163 However, heterogeneous conversion processes still need the description of the total reactive Surface Area 164 Density (SAD) from aerosols. In TM5-MP this is assumed as the cumulative value of contributions from 165 sulphate, nitrate, ammonium and methane sulphonic acid as calculated by the EQuilibrium Simplified Aerosol 166 Model (EQSAM) approach (Metzer et al., 2002), thus the secondary organic aerosol component is not included. 167 The distribution of these aerosol species is calculated online and coupled to the respective gaseous precursors. 168 The density of each aerosol type  $(1.7 \text{ g/cm}^3)$  and  $r_{eff}$  (of between  $0.18-0.2\mu\text{m}$ ) is prescribed as in Huijnen et al. 169 (2014). Swelling at higher relative humidities (> 70%) is crudely accounted for by increasing  $r_{eff}$  to between 170 0.25-0.27µm. The contributions due to sea-salt, black carbon and organic carbon towards heterogeneous loss are 171 not accounted for. Temperature dependent gas phase diffusion co-efficients (Dg) are used in the derivation of the 172 pseudo first-order heterogeneous rate constants based on the theory of Schwartz (1986). 173 For N<sub>2</sub>O<sub>5</sub>, the uptake coefficient (γ) is calculated using the parameterization of Evans and Jacob (2005), 174 therefore dependent on both temperature and relative humidity. Once a surface reaction with H<sub>2</sub>O occurs two 175 molecules of nitric acid (HNO<sub>3</sub>) are formed. No uptake on cirrus particles is included for HNO<sub>3</sub>, which can lead 176 to de-nitrification of the upper troposphere (Lawrence and Crutzen, 1998; von Kuhlmann and Lawrence, 2006). 177 For  $HO_2$  we adopt a fixed  $\gamma_{HO2}$ = 0.06 across all aerosol types as taken from Abbatt et al. (2012) and for  $NO_3$  we 178 adopt a fixed  $\gamma_{NO3} = 10^{-3}$  as recommended by Jacob (2000). For HO<sub>2</sub>, heterogeneous conversion forms 0.5 179 molecules of Hydrogen Peroxide (H<sub>2</sub>O<sub>2</sub>), whereas for NO<sub>3</sub> it forms one molecule of HNO<sub>3</sub> is formed, following 180 Emmons et al. (2010). For the SAD associated with cloud droplets we use the  $r_{eff}$  values that are calculated by 181 Martin et al. (1994) thus maintaining consistency between the size of the cloud droplets used for the scattering 182 component in the calculation of J values and heterogeneous loss rates on the clouds. By using the ECMWF 183 cloud fraction for each respective grid cell, we assume that instantaneous mixing throughout the grid cell does 184 not occur in order to avoid exaggerated conversion rates on cloud surfaces. 185 As TM5-MP contains no explicit stratospheric chemistry, we apply constraints above the tropopause to ensure 186 realistic Stratosphere-Troposphere Exchange (STE) of O<sub>3</sub> and for constraining the incoming radiation reaching 187 the troposphere needed for the MBA (Williams et al., 2012). For stratospheric O<sub>3</sub>, we use total column values 188 derived from the assimilation of satellite observations as provided in the improved version of the Multi-Sensor 189 Re-analysis (MSR, van der A., 2010), which is vertically distributed according to the climatology of Fortuin and 190 Kelder (1998). Three distinct zonal bands are used for nudging the stratospheric O<sub>3</sub> fields, these being 30°S-191 30°N, 30-66°S/N and > 66°S/N, where nudging occurs at pressure levels <45hPa, <95hPa and <120hPa, with 192 relaxation times of 2.5 days, 3 days and 4 days, respectively. 193 For stratospheric CH<sub>4</sub> we use the monthly 2D climatological fields provided by Grooß and Russell (2005), with 194 the nudging heights and relaxation times being identical to those used for stratospheric O<sub>3</sub>. For stratospheric CO 195 and HNO<sub>3</sub> we constrain mixing ratios by using monthly mean ratios between CO/O<sub>3</sub> (Dupuy et al., 2004) and

HNO<sub>3</sub>/O<sub>3</sub> (Jégou et al., 2008; Urban et al., 2009) based on the latitudinal climatologies derived from ODIN

observations between 2003/2004 and 2001-2009. In order to avoid jumps in the nudging constraints between

months, we gradually change between ratios using the total monthly difference/number of days in the month.

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





6

These ratios are applied using the monthly mean stratospheric O<sub>3</sub> distribution in TM5-MP, which is constrained by the MSR dataset (van der A et al., 2010). For both species, model fields are nudged at 5.5hPa, 10hPa and 28hPa using relaxation times of 5, 10 and 60 days, respectively. Previous versions of TM5 used a HNO<sub>3</sub> climatology from the UARS MLS instrument and applied nudging constraints at 10hPa only (Huijnen et al., 2010).

204205

206

207

208

209

210

211

212

213

214

215

216

217

218

219

220

221

222

223

224

225

226

227

228

229

230

## 2.2 Emission inventories

All emission inventories applied in TM5-MP are yearly specific meaning that the year-to-year variability in emission fluxes due to changes in anthropogenic activity, biogenic activity and burning extent are taken into account. For the anthropogenic emission of NOx, CO, SO2, NH3 and Non-Methane Hydrocarbons (NMVOC) we adopt the MACCity emission estimates described in Granier et al. (2011). The lack of sector-specific information complicates the use of daily cycles for e.g. the road transport component. For volcanic SO2 emissions, the estimated emission flux has been scaled up to 10 Tg S/year based on Halmer et al. (2002). For the biogenic component, where available we use the CLM-MEGANv2.1 emission inventories produced for the Southern Hemispheric Multi-model Intercomparison Project (SHMIP) as described in Zeng et al. (2015), with the missing trace species (e.g. ethane, propane, higher organics) coming from alternative MEGAN simulations as outlined in Sindelarova et al. (2014). A diurnal cycle is imposed on the isoprene emissions and introduced into the first ~50m between 20°S-20°N. The BB emissions are taken from the monthly estimates provided by the GFEDv3 inventory (van der Werf et al., 2010) and latitude dependent injection heights and a tropical burning cycle are implemented following Huijnen et al. (2010). All emission inventories are provided on a 0.5° x 0.5° resolution and subsequently coarsened onto the horizontal resolution used during any particular simulation. For lightning NO<sub>x</sub> we use the parameterization which uses convective precipitation of Meijer et al. (2001) and constrain to a total global emission of ~6Tg/N yr<sup>-1</sup>. This uses the convective flux values meaning that re-scaling of the nudging term was necessary in order to achieve similar total lightning NO<sub>x</sub> across simulations. In TM5-MP all NO<sub>x</sub> emissions are introduced as NO, rather than speciating a fraction which is emitted directly as NO<sub>2</sub> (Carslaw and Beevers, 2005). Global NO<sub>x</sub> emissions for the year 2006 total 49 Tg N yr<sup>-1</sup> (including lightning). Other notable species include CO (1081 Tg CO yr<sup>-1</sup>), SO<sub>2</sub> (117 Tg S yr<sup>-1</sup>), CH<sub>2</sub>O (13.5 Tg C yr<sup>-1</sup>) and isoprene (510 Tg C yr<sup>-1</sup>). An overview of the global and zonal emissions terms used in the simulations analysed here are given in Table 3. Latitudinal constraints on CH<sub>4</sub> global distributions are applied using the methodology outlined in Banda et al. (2015). We also introduce similar constraints based on the appropriate surface measurements for  $H_2$  in order to

231232233

# 2.3 Observations

Although the performance of mCB05 in TM5 v3.0 has been validated for selected Volatile Organic Compounds (VOC), O<sub>3</sub>, CH<sub>2</sub>O, CO and NOy in both hemispheres (Williams et al., 2013; 2014; Fisher et al., 2015; Zeng et al., 2015), the significant changes made to both the chemical scheme and the rate parameters in mCB05v2 necessitate independent validation at both 3° x 2° and 1° x 1°. We choose a range of ground-based and airborne

account for the latitudinal gradient and variability across seasons, which replaces the fixed global value of

550ppb used in previous versions. Finally, for Radon (Rn<sup>222</sup>) emissions we apply the estimates of Schery (2004).

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





7

238 measurements taken at diverse locations representing a diverse range of chemical regimes. Here we briefly 239 describe the observations utilised for this purpose. 240 For validation of simulated surface concentrations we use measurements of gaseous O3, NO, NO2, HNO3 and

241 SO<sub>2</sub> available from the European Monitoring and Evaluation program (EMEP, www.emep.int), where we

242 exploit measurements taken at various background sites in Norway, Finland, The Netherlands, Belgium, Poland,

243 Germany, Spain, Italy and Portugal. For the model composites we extract data from 3 hourly instantaneous 244 output in order to assemble both the weekly and monthly mean values from the simulations. For the weekly

245 comparisons of NO<sub>2</sub> and SO<sub>2</sub> we use values extracted at 13:00 local time, close to the overpass time of the OMI

246 instrument (e.g. Boersma et al, 2008). The selected stations allow validation of the seasonality for both rural

247 regions (FI37) and urban regions (NL09), where we include identical stations where possible for both species.

248 For HNO<sub>3</sub> we assemble the weekly values from the daily averages.

249 Measured O<sub>3</sub> surface concentrations in the EMEP network are obtained using UV monitors (Aas et al. 2001).

250 For all species, spatial interpolation of model data is performed accounting for the height of the measurement

251 station and by weighting using the distance of the station from the surrounding grid cells. The wide range of

252 measurement sites chosen insures that both background and polluted cases are assessed.

253 For validating the vertical distribution of relevant trace species such as O<sub>3</sub>, SO<sub>2</sub> and CH<sub>2</sub>O, we use

254 measurements by the DC-8 aircraft during the Intercontinental Chemical Transport Experiment B (INTEX-B;

255 Singh et al., 2009) that took place between March and May 2006. Observations of a host of co-located nitrogen-

256 containing species are available (namely NO, NO2, PAN and HNO3). These flights were conducted over a wide

257 region, and we use all three months of measurements. Each month sampled a different region representing

258 different meteorological conditions and local emission sources, namely: the Gulf of Mexico (90-100°W, 15-

259 30°N), the remote Pacific (176-140°W, 20-45°N) and around Alaska (160-135°W, 20-60°N) with flights to the

260 south and west. Measurements cover altitudes up to 10.5km, and bin the values with respect to pressure using 50 261

hPa bins or less in the LT. We interpolated three-hourly output against measurements for each respective day,

262 similar to the comparisons performed in previous evaluations of TM5 (e.g. Huijnen et al., 2010), but we

263 segregate our comparisons into the three distinct regions. For details relating to the location of each flight the

264 reader is referred to the campaign overview of Singh et al. (2009).

265 For tropospheric O<sub>3</sub>, we supplement the INTEX-B comparisons with those taken over more polluted regions as

266 part of the Measurement of Ozone, water vapour, carbon monoxide and nitrogen oxides by Airbus In-service

267 aircraft (MOZAIC; Thouret et al., 1998). We aggregate the measurements as seasonal means for December-

268 January-February (DJF) and June-July-August (JJA) in order to provide a robust number of samples for each

269 location. Here we chose to use profiles representative of the Northern mid-latitudes, namely: London (0.2°W,

270 51.2°N), Vienna (16.5°E, 48.1°N), Washington (77.5°W, 38.9°N), Portland (122.6°W, 45.6°N), Shanghai

271 (121.8°E, 31.2°N) and Tokyo (140.4°E, 35.8°N).

272 We also make comparisons of O<sub>3</sub>, NO, NO<sub>2</sub>, selected N reservoir species, SO<sub>2</sub> and CH<sub>2</sub>O profiles using

273 measurements made aboard the NOAA WP-3D aircraft as part of the Second Texas Air Quality Study (TexAQS

274 II; Parrish et al, 2009), which was conducted over the Texas sea-board during September and October 2006.

275 This allows the assessment of TM5-MP over a region with higher VOC emissions and industrial activity. These

measurements were typically sampled at altitudes below 500 hPa, therefore no UTLS measurements are

277 available from this campaign.

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





### 3 The Effect on Atmospheric Transport

Here we analyse the differences in convective transport out of the boundary layer by analysing the vertical and horizontal distribution of <sup>222</sup>Rn, which is a diagnostic typically used for assessing the differences in transport in CTMs (e.g. Jacob et al., 1997). <sup>222</sup>Rn is emitted at a steady rate and exhibits a half-life of ~3.8 days, which is long enough to be transported from the boundary layer into the FT due to chemical passivity, with loss via wet scavenging and deposition being negligible. Therefore, it acts as an ideal tracer to assess differences in convective transport from the surface out of the boundary layer (BL).

Figure 1 shows seasonal mean horizontal global distributions of  $^{222}$ Rn for DJF and JJA in the 1° x 1° simulation averaged between 800-900hPa (i.e.) at the top of the BL including the lower FT highlighting the spatial variability in convective upwelling near the top of the convective boundary layer. Also shown are the associated percentage differences against the re-binned 3° x 2°  $^{222}$ Rn distribution, allowing a direct comparison. Resolution dependent differences result from the cumulative effects of the use of higher resolution mass fluxes from the ERA-interim meteorological data for describing convective activity and the more accurate temporal distribution of regional  $^{222}$ Rn emissions at 1° x 1°. In general it can be seen that seasonal differences of  $\pm 20\%$  exist, typically with increases over continents and decreases over oceans. Maximal differences of >60% occur near selected coasted regions (California, West Africa, Madagaskar) or in outflow regions such as off South America and Africa, where differences exhibit a strong seasonal dependency. This is due to the large differences in convective strength due to the variability in heating rates, and thus temperatures, between land and ocean (e.g. Sutton et al., 2007).

A comparison of the ratio of the monthly mean <sup>222</sup>Rn profiles (1° x 1° /3° x 2°) extracted above selected European cities for January (black) and July (blue) 2006 are shown in Figure S1 in the supplementary material. The typical tropospheric profile of <sup>222</sup>Rn exhibits an exponential decay from the LT to the FT (not shown). In order to homogenise the emission flux in the comparison, we coarsen the 1° x 1° data onto the 3° x 2° grid by averaging the six individual values into a representative mean column. The extent of the changes in the vertical distribution of <sup>222</sup>Rn is somewhat site specific meaning an in depth analysis is beyond the scope of this paper. In summary, the 1° x 1° simulation generally provides stronger convective activity for January, with the main impact occurring below 700hpa (e.g. London and Paris). The changes in 222Rn in the lower troposphere range between ~2-10% (i.e. ratios of 0.9-1.1), implying both weaker and stronger convective transport depending on changes in location (i.e.) orography and land type. In that the impact at Berlin is larger than at e.g. Barcelona also shows that, surprisingly, the inclusion of a large ocean fraction (with weaker convective mixing) in the 3° x 2° cell does not seem to introduce dominating effects. For July the changes in the vertical distribution extend into the FT up to 500hPa, although changes in the upper FT have a significant component due to changes in long-range transport. The magnitude of the changes are similar to those exhibited during January, although maybe of the opposite sign (e.g. Rome). Thus the influence on e.g. NO2, CH2O and SO2 a-priori vertical profiles will be non-negligible and diverse.

For the tropical cities located in regions where convective mixing is stronger, the corresponding differences can reach  $\pm$  20%, especially near the surface (e.g. Caracas and Karachi). There is a site-specific seasonal dependency in the magnitude of the changes related to the regional land characteristics (e.g. Lagos versus Kuala Lumpar).

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





Thus, potential differences in *a-priori* vertical profiles can be considerable compared to those provided at a 3° x 2° resolution.

We also show comparisons of profiles from 1° x 1° simulations using the convective scheme of Tiedke (1989) against those using the convective mass-fluxes from the ERA-interim meteorological dataset for Europe (Figure S3). For this comparison no averaging is employed, where the selected grid cells are near the centre of each urban conurbation. The residuals show that the significant differences exist, with the convective mass-fluxes from ERA-interim being somewhat weaker than those calculated online using Tiedke (1989) (i.e) the residual is typically less than 1, especially during July.

### 4 The Impact on tropospheric photolysis frequencies

The changes in the spatio-temporal distribution of cloud cover and surface albedo has the potential to alter the penetration depth and upwelling of photolysing light, and thus photochemical production and destruction terms. The similarity in the monthly mean photolysis frequencies for  $O_3$  and  $NO_2$  across resolutions (hereafter denoted  $J_{O3}$  and  $J_{NO2}$ , respectively) are shown in Figure S4 of the supplementary material. Comparisons of the monthly mean  $J_{O3}$  and  $J_{NO2}$  values are shown at five different locations identical to those shown in Williams et al. (2012). For  $J_{O3}$  the impact of increasing resolution is limited to a few percent in the monthly mean values. At global scale this leads to a reduction of ~2% in the total mass of  $O_3$  photolysed (not shown). For  $J_{NO2}$ , the corresponding differences become more appreciable, with  $1^{\circ}$  x  $1^{\circ}$  exhibiting ~5-10% higher values at high Northern latitudes (associated with the high- $NO_x$  scenario).

Focusing on  $J_{NO2}$  and comparing seasonal mean values near the surface shows that very similar spatial patterns occur for both simulations at global scale (c.f. Figure S5). The highest  $J_{NO2}$  values occur over the tropical oceans and high altitude regions (e.g. Nepal). Although more regional fine-structure can be seen at 1° x 1° (e.g. South-Western US for DJF and around Iceland for JJA), these seasonal averages show that the small perturbations shown in Fig. S3 extend to the global scale leading to a reduction of ~5% at 1° x 1°.

Comparisons of monthly mean profiles of  $J_{O3}$  and  $J_{NO2}$  extracted over the location of selected tropical cities are shown in the Figs. S6a and b, respectively, in the Supplementary Material. Here no averaging is performed towards an identical horizontal resolution, therefore values are representative of the J values directly above the selected urban centres. The  $J_{O3}$  profiles are affected to a larger extent than the  $J_{NO2}$  profiles due to the characteristic absorption spectra of each species, which makes  $J_{O3}$  more sensitive to the additional scattering introduced due to clouds. Profiles over Dubai act as a proxy for clear-sky conditions, where values of unity exist in the residual of  $J_{O3}$  and  $J_{NO2}$  calculated through most of the column. The small difference at the surface is due to changes in the surface albedo between resolutions, with Dubai being situated on the coast meaning that a sharp horizontal gradient exists in surface albedo. For other cities, the largest perturbations occur away from the surface (e.g. Jakarta, Nairobi and Lagos) around the altitude where tropospheric clouds are most abundant. There are typically changes of between  $\pm 5$ -10% in the monthly mean profiles. The changes in  $J_{NO2}$  reflect those simulated for  $J_{O3}$ , with somewhat smaller perturbations.

# 5 Implications for oxidative capacity and tropospheric O<sub>3</sub>

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





10

358 The partitioning of reactive N between the short- and long-lived chemical N-reservoirs included in TM5-MP depends on the oxidative capacity simulated for the troposphere via competition between the various different 359 360 radicals i.e. OH, CH<sub>3</sub>C(O)O<sub>2</sub>, NO<sub>3</sub> and CH<sub>3</sub>O<sub>2</sub>. Therefore, changes to the distribution and resident mixing ratios 361 of tropospheric O<sub>3</sub> subsequently impose changes on the fractional composition of the NO<sub>y</sub> budget (e.g. Olszyna 362 et al., 1994) and the also the efficiency of the NO<sub>x</sub> re-cycling terms by altering the chain length (Lelieveld et al., 363 2004). In this section we analyse the global and zonal chemical budget terms for tropospheric O<sub>3</sub> to highlight the 364 inter-hemispheric differences which occur (i.e.) under low and high-NO<sub>x</sub> environments. 365 Table 4 provides the zonally segregated chemical budget terms for tropospheric O<sub>3</sub>, from which the global 366 component due to STE can be determined by closing the budget terms following the methodology given in 367 Stevenson et al. (2006). The chemical tropopause calculated for 3° x 2° is applied for the analysis of 1° x 1° 368 budget terms to ensure that a valid comparison is performed, i.e. the same mass of air is accounted for. For 369 computational efficiency the budget terms are aggregated in 10° latitudinal bins and summed across all 370 longitudes providing the cumulative terms. 371 The most significant change with resolution concerns STE. By using a dedicated tagged stratospheric O<sub>3</sub> tracer 372 (which only undergoes photo-chemical destruction and deposition in the troposphere; O<sub>3</sub>S) changes in the zonal 373 STE can be determined. The stratospheric burden of O<sub>3</sub> (BO<sub>3</sub>S) exhibits a strong hemispheric gradient with 374 much more down-welling occurring in the NH peaking during boreal springtime. At global scale the STE 375 exchange is 579 TgO3 yr<sup>-1</sup>, which agrees well with the multi-model STE mean of 556±154 TgO3 yr<sup>-1</sup> in 376 Stevenson et al. (2006), with observational estimates being ~550±140 TgO<sub>3</sub> yr<sup>-1</sup> (Olsen et al., 2001). This ~7% 377 reduction of STE at 1° x 1° is encouraging considering that previous studies using TM5 have concluded that STE 378 in TM5 at 3° x 2° was biased high compared to STE inferred from TES and MLS satellite observations 379 (Verstraeten et al. (2015)). The increase in STE in the SH, with an associated decrease in the NH (see below), 380 implies that there is a shift in circulation patterns at 1° x 1° even though the stratospheric BO3 remains essentially 381 unchanged. Previous studies have shown that in order to resolve the correct spatial and temporal stratosphere-382 troposphere flux, high resolution is required both in the horizontal and the vertical (e.g. Meloen et al., 2002). 383 The NH STE diagnosed with TM5-MP is an order of magnitude smaller than estimates derived in a recent study 384 also conducted at a 1° x 1° resolution (Tang et al, 2011; ~200 TgO<sub>3</sub> yr<sup>-1</sup>), which identified deep convection as 385 important for STE. Here we use a different vertical grid and meteorological dataset to drive TM5-MP, both of 386 which affect the ability towards capturing an accurate STE flux (Meloen et al., 2002). 387 The zonal seasonal means of the fraction of O<sub>3</sub>S to O<sub>3</sub> (O<sub>3</sub>S/O<sub>3</sub>) for both simulations are shown in Figure 2 for 388 DJF and JJA. There is a clear seasonal zonal shift in the fractional contribution due to the O3 transported 389 downwards from the Stratosphere exhibiting a longer lifetime in the winter hemisphere reflecting a lower 390 photochemical destruction rate. At 1° x 1° the largest increase in STE occurs in the SH during JJA. Here ~20-391 25% of tropospheric  $O_3$  is transported down from the stratosphere. Comparing the 0.2 contour for the NH mid-392 troposphere shows significant changes, extending further down towards the surface during boreal wintertime 393 leading to the higher total mass of O<sub>3</sub>S in the troposphere. The extent of nudging towards the MSR climatology 394 is essentially constant across simulations (c.f. Table 4). Interestingly, less O<sub>3</sub>S reaches the surface in the tropics 395 at 1° x 1° due to the enhanced chemical destruction term in the Free Troposphere. Approximately 10% of the 396 global deposition term for O<sub>3</sub> is associated with O<sub>3</sub> that originates from the Stratospheric at 1° x 1° (c.f. ~5% at 397 3° x 2°). For the NH, this contributes to the simulated increase in deposition of ~9%.

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

398

© Author(s) 2016. CC-BY 3.0 License.





11

For tropospheric O<sub>3</sub> there are similarities that occur between the NH, tropics and SH (i.e.) high and low-NO<sub>x</sub>

399 scenarios, resulting in a cumulative decrease in O<sub>3</sub> production of ~2-4% across zones. For the chemical loss 400 terms there is a decrease of ~3% (~1%) in the NH (SH) reflective of the changes discussed for J<sub>O3</sub>, which acts as 401 the primary destruction term. There is a zonal gradient in the tropospheric BO<sub>3</sub> following the zonal gradient in 402 NO<sub>x</sub> emissions. Comparing terms shows that BO<sub>3</sub> decreases at 1° x 1° by a few percent at global scale (~7 Tg O<sub>3</sub>) 403 making a rather small impact on oxidative capacity. This is of the same order of magnitude as that found in 404 previous studies concerned with horizontal resolution (e.g. Wild and Prather, 2006). Interestingly, changes in the 405 deposition flux of  $O_3$  are rather small, even though there is a larger amount of variability in the land surfaces and 406 better-resolved land-sea contrast at 1° x 1°, although differences in regional deposition fluxes can be more 407 significant. Multi-model inter-comparisons of surface deposition terms across models have shown previous 408 versions of TM5 to be a the low end of the model spread in terms of O<sub>3</sub> (Hardacre et al., 2015), suggesting that 409 the surface deposition flux to e.g. should be increased by ~10% in TM5-MP towards the multi-model mean 410 value. This can be partly attributed to the large uncertainty which exists related to the loss of O<sub>3</sub> to the ocean 411 (Hardacre et al., 2015). 412 Figure 3 shows comparisons of simulated and observed mass mixing ratios of surface O<sub>3</sub> at EMEP sites across 413 Europe (www.emep.int; Aas et al. 2001). Previous comparisons using mCB05 have revealed high biases in 414 surface O<sub>3</sub>, especially during boreal summertime (Williams et al., 2013). These high biases originate from 415 cumulative effects associated with the accuracy of the emission inventories, the convective mixing component, 416 the underestimation of the scattering and absorption of photolysing light due to aerosols and the chemical 417 mechanism that is employed. For the emission component it should be noted that even at 1° x 1° coarsening is 418 performed, where emission inventories are typically supplied at 0.5° x 0.5° resolution. The seasonal cycle in 419 surface  $O_3$  is captured to a large degree, and the high bias exhibited by the model is generally reduced by ~2-5 420 ppb (or ~20%) at 1° x 1°. This is associated with perturbations in the NO<sub>x</sub> recycling terms, chemical titration by 421 NO and convective mixing out of the boundary layer. In that the improvement in biases is largest during boreal 422 summertime is associated with the shorter chain length of the NO<sub>x</sub> recycling term during boreal wintertime (c.f. 423 Fig 2). However, there is still a significant monthly-mean bias in both simulations when compared against 424 observations throughout the year, especially for locations impacted by a large anthropogenic NO<sub>x</sub> source. This is 425 partly due to the low NO/NO2 ratio as discussed in Sect. 4 below. 426 Comparing vertical gradients from composites assembled from the MOZAIC measurements for DJF and JJA 427 (Figures S7a and S7b, respectively), INTEX B (Singh et al, 2009; Figure S8) and TexAQS II (Parrish et al, 428 2009; Figure S9) shows that differences are small and typically mimic those which occur at the surface. There is 429 a general positive bias of 20-40% in mixing ratios exhibited across all comparisons, although the variability in 430 the vertical gradients across regions is capture rather well. Such positive biases have consequences for both the 431 NO<sub>x</sub> recycling terms and HNO<sub>3</sub> formation discussed in the sections below.

4 Implications for the distribution of NO and NO<sub>2</sub>

433434435

436

437

432

Table 5 provides the zonally segregated annual  $NO_x$  re-cycling terms involving the main peroxy-radicals and the direct titration term involving NO for the 1° x 1° simulation. The conversion rate of NO back into  $NO_2$  decreases by ~2-3% across zones as a consequence of an associated increase in the titration term and re-partitioning of N

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

476

477

© Author(s) 2016. CC-BY 3.0 License.





12

438 into long-lived reservoir species (see below). For the titration term involving NO, although the globally 439 integrated flux remains relatively constant, there is contrasting behaviour for the two most important zones (TR, 440 NH), which exhibit a lower and higher titration term, respectively. It has been shown that for regions such as 441 Europe the increased titration results in lower surface O<sub>3</sub> mixing ratios (c.f. Fig. 3), improving the boreal 442 summertime high bias at the surface. 443 Important model uncertainties include the quality of the MACCity NO<sub>x</sub> emission inventory, the lifetime of NO<sub>2</sub> 444 simulated in TM5 and the re-cycling term via the chemical titration of O<sub>3</sub>. Figures 4 and 5 shows comparisons of 445 weekly [NO] and [NO<sub>2</sub>] surface measurements against the corresponding composites from both of the 446 simulations, sampled at 13:00 local time close to the time of overpass for OMI and tropOMI. To supplement 447 these comparisons we provide the seasonal mean biases for DJF and JJA from both simulations in Tables 6 and 448 7, respectively, calculated using weekly binned data from all EMEP sites, which measure hourly [NO] and 449 [NO<sub>2</sub>] levels. Here we perform an analysis across sites rather than focusing on the behaviour at selected 450 individual locations. 451 For the determination of [NO<sub>2</sub>], the reduction of NO on a Molybdenum convertor takes place with subsequent 452 detection by chemi-luminescense, with an associated detection limit of ~0.4ppb. Previous studies have shown 453 that some bias can result due the oxidation of nitrogen reservoirs such as PAN (Dunlea et al., 2007; Steinbacher 454 et al., 2007). In TM5-MP all  $NO_x$  emissions are introduced as NO, although a fraction for road transport is 455 known to be emitted directly as NO2 (e.g. Carslaw and Beevers, 2005). Many studies have been performed 456 comparing satellite NO2 columns with model values, implying that inadequacies in emission inventories are 457 somewhat region specific (e.g. Zyrichidou et al., 2015; Pope et al., 2015). 458 Table 6 shows a negative bias of a few µg m<sup>-3</sup> in TM5-MP in seasonal surface [NO] in Europe. This is a 459 cumulative effect of the accuracy of the MACC NO<sub>x</sub> emission estimates and, to a lesser extent, too high surface 460 [O<sub>3</sub>] (enhancing the oxidation rate of NO to NO<sub>2</sub>). As anthropogenic emissions are the principle source of NO, 461 no significant seasonal cycle exists in the weekly measurements, although differences in convective mixing do 462 cause somewhat higher surface [NO] during DJF, which is often captured in TM5-MP. For ~80% of the EMEP 463 sites we do not observe any significant change in the quality of the comparisons. For a 20% of sites, simulations 464 of [NO] at 1° x 1° introduce significant improvements over those at 3° x 2° and seasonal variability generally 465 improves (Fig. 4). 466 Table 7 shows that for [NO<sub>2</sub>] the biases are more variable being typically in the range of  $\pm 0$ -6  $\mu g$  m<sup>-3</sup>, with both 467 positive and negative biases occurring across sites. Both the conversion efficiency from NO, loss to reservoir 468 compounds i.e. HNO3, photo-dissociation rate and emission estimates contribute to these biases. The seasonal 469 biases show improvements at 1° x 1° for ~35% of the EMEP sites, accompanied with degradations at ~20% of 470 the sites. The maximal biases in [NO2] at 1° x 1° can be approximately double those for [NO]. For the 471 corresponding NO/NO<sub>2</sub> ratio, there will generally be an under-prediction in the model due to the negative biases 472 shown for the [NO] comparisons. 473 Beyond Europe, we compared monthly mean TM5-MP vertical distributions of NO and NO2 between March 474 and May 2006 against measurements taken during the INTEX-B campaign in Figure 6. In general differences 475 between 1° x 1° and 3° x 2° simulations are the order of a few percent, with NO2 biased low in the lower

troposphere by ~70-80%. This is partially associated with the take-off and landing of the aircraft from polluted

airfields, where point sources of high anthropogenic emissions cannot be resolved at 1° x 1°. For the FT, TM5-

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





13

478 MP captures the observed gradient to reasonable degree. In the UT there is a consistent high bias for NO and an 479 associated low bias for NO<sub>2</sub> suggesting that the conversion term is too low and the NO<sub>3</sub> cycle is out of synch at 480 these cold temperatures despite the addition of new reservoir species i.e. CH<sub>3</sub>O<sub>2</sub>NO<sub>2</sub>. 481 One important gauge as to whether the chemical mechanism can capture the correct re-cycling efficiency of NO 482 into NO<sub>2</sub> is to examine their ratio. These are presented in the third column of Fig. 6. In the lower troposphere (< 483 900 hPa) NO/NO<sub>2</sub> ratios of 0.1-0.2 exist which TM5-MP represents quite well, with negligible differences 484 between 3° x 2° and 1° x 1° simulations. For the FT, TM5-MP consistently over estimates the ratio in spite of a 485 high-bias in O<sub>3</sub> (c.f. Fig. 3) suggesting an exaggerated photo-dissociation rate or low bias in HO<sub>2</sub>. 486 Finally in Figure 7 we show the corresponding comparisons against measurements taken during the TexasAQS-487 II campaign (Parrish et al, 2009) during September 2006. There is a significant underestimation in NO and NO<sub>2</sub>, 488 mixing ratios, with both model profiles outside the 1-σ variability of the measured mixing ratios. This is clearly 489 related to the emission estimates for this region being underestimated in the emission inventories (e.g. Kim et al., 490 2011). For the resulting NO/NO2 ratio TM5-MP captures the correct ratio in the lowest few hundred meters of 491 the boundary layer, but overestimates the ratio at higher altitudes. In this case, the high NO and NO2 mixing 492 ratios at the top of the boundary layer imply that TM5-MP under-represents the NO2 fraction, regardless of the

494 495

493

# 5 Changes in the NOy budget

increased recycling term related to the titration of O<sub>3</sub> (c.f. Table 2).

496 497

## 5.1 Long-lived reservoirs

498

499 The resolution dependent changes in the temporal distribution of [NO<sub>2</sub>], and associated differences in VOC 500 chemical pre-cursor emissions have the potential to alter the partitioning of reactive NO<sub>x</sub> between the three main 501 chemical reservoirs included in mCB05v2 (i.e. HNO3, PAN and ORGNTR). Considering the differences in both 502 the deposition efficiency and tropospheric lifetimes between species means the fraction of NO<sub>x</sub> transported out 503 of source regions could change significantly. Here we briefly examine the zonally integrated nitrogen budget 504 terms between simulations to quantify the effect on higher spatial resolution. The seasonal distribution of these 505 three dominant reservoir species at 1° x 1° and their individual contributions to total NOy are shown in Figures 506 S10 to S13 for DJF and JJA, respectively. Here we define NO<sub>v</sub> as the cumulative total of NO, NO<sub>2</sub>, NO<sub>3</sub>, HNO<sub>3</sub>, 507 PAN, CH<sub>3</sub>O<sub>2</sub>NO<sub>2</sub>, HONO, 2\*N<sub>2</sub>O<sub>5</sub>, lumped organic nitrates and HNO<sub>4</sub>. These are provided as reference for the 508 reader to aid understanding of the discussion below. 509 Table S1 in the supplementary material provides a zonal decomposition of the tropospheric chemical budget 510 terms for HNO3, PAN and ORGNTR. For HNO3, even though the recent kinetic rate parameters increase 511 (decrease) the chemical production term at the surface (UTLS) compared to older rate data (e.g. Seltzer et al., 512 2015), changes in the integrated column term are small. The changes at 1° x 1° are somewhat latitude dependant 513 (low and high NO<sub>x</sub> regimes), with only small increases occurring in the NH and associated decreases in the 514 tropics related to lower [OH] (i.e.) chemical production. 515 For PAN, both the production and destruction terms decrease marginally by ~2-3% across all zones, meaning 516 the transport of NO<sub>x</sub> out of the main source regions remains relatively robust. The total mass of N cycled 517 through PAN is ~four times that of HNO<sub>3</sub>. The changes in the production term due to temporal increases in NO<sub>2</sub>

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

557

© Author(s) 2016. CC-BY 3.0 License.





14

518 near high NO<sub>x</sub> source regions (c.f. Fig. 5) are partially offset by a reduction in the mixing ratios of the acetyl-519 peroxy radical (C<sub>2</sub>O<sub>3</sub> in Table 1) due to e.g. increased dry deposition of organic precursors at 1° x 1°. Although 520 the chemical budget terms only exhibit small changes, it can be expected that the global distribution is somewhat 521 different due the changes in the convective and advective mixing terms. 522 For ORGNTR, there is a 5% reduction in the production term at 1° x 1°, with an associated decrease in the loss 523 by deposition. Both the largest production term and decrease occur in the tropics related to the strongest source 524 being biogenic pre-cursors. Thus at 1° x 1°, this intermediate become less important as a NO<sub>x</sub> reservoir. 525 Finally, the one additional intermediate not shown is CH<sub>3</sub>O<sub>2</sub>NO<sub>2</sub>, which is primary a stable vehicle for 526 transporting NO<sub>x</sub> from the surface up to the UTLS, where at cold temperatures it accounts for a significant 527 fraction of NO<sub>2</sub> speciation along with HNO<sub>4</sub> (Browne et al., 2011). At global scale three times as much nitrogen 528 cycles through CH<sub>3</sub>O<sub>2</sub>NO<sub>2</sub> compared to PAN, although the thermal stability is low at temperatures > 255°K thus 529 resident mixing ratios are typically small. This results in maximal mixing ratios occurring in the cold upper 530 troposphere (up to ~0.2 ppb) and subsequently dissociates primarily by thermal decomposition (photolytic 531 destruction accounting for <0.1% of all destruction). At 1° x 1° there is a few percent decrease in the chemical 532 production term as a result of lower CH<sub>3</sub>O<sub>2</sub> mixing ratios and more variability in the temporal temperature 533 distribution. 534 Comparisons of weekly [HNO<sub>3</sub>] at the surface in Europe are shown in Figure 8 against measurements from the 535 EMEP network. It has recently been determined that HNO3 measurements are also sensitive to ambient night-536 time  $N_2O_5$  concentrations, which could result in a positive bias in the observations (Phillips et al., 2013). In 537 general, the modelled seasonal cycle is not evident in the measurements, which exhibit a rather homogeneous 538 variation in mixing ratios throughout the year typically leading to an underestimation in TM5-MP during March 539 and an overestimation during JJA. No such seasonal pattern is observed for [NO<sub>2</sub>] (c.f. Fig. 5), thus seasonal 540 [OH] variability due to variations in photo-chemical activity and [H<sub>2</sub>O<sub>(g)</sub>] is a likely cause. The impact of 541 resolution on [HNO<sub>3</sub>] is rather muted for most weeks resulting in no significant changes to the seasonal biases 542 (not given), as constrained by the improvements in surface [NO<sub>2</sub>] (c.f. Fig. 5). The heterogeneous scavenging of 543 HNO<sub>3</sub> into ammonium nitrate can act as a moderator toward gaseous HNO<sub>3</sub> and, although included in TM5-MP, 544 generally produces low concentrations of e.g. ammonium nitrate (not shown). Thus, gaseous [HNO<sub>3</sub>] remains 545 too high due to too little conversion into particles. 546 For other regions, we make comparisons of vertical profiles of HNO<sub>3</sub> and PAN between March and September 547 2006 against those measured during INTEX-B (Figure 9) and Texas (Figure S14). PAN is a good marker for 548 transport in the free-troposphere due to the relatively long-lifetime at colder temperatures. For all regions the 549 vertical gradients for both species are captured quite well, although some fine-structure is lost due to the vertical 550 resolution of TM5-MP. This implies that the under-estimation in NO2 (Fig. 6) in not due to lack of transport 551 away from source regions and therefore should be more attributed to local under-estimations in emission fluxes. 552 Finally, we present the corresponding comparisons for September 2006 for HNO<sub>3</sub> and PAN measured during the 553 TexAQS II campaign (Parrish et al., 2009) as Figure S14 in the supplementary material. For HNO<sub>3</sub>, although the 554 vertical gradient is captured quite well, there is a significant low bias related to the low bias in NO<sub>2</sub> shown in 555 Fig. 7, with the 1° x 1° showing a marginal improvement in the LT for HNO<sub>3</sub>. For PAN the vertical profile in 556 TM5-MP is somewhat anti-correlated around 900hPa in both simulations, with the rapid decrease at the surface

not being captured by either simulation and the bias being larger for 1° x 1°.

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





15

558

#### 5.2 Short-lived reservoirs

559560561

562

563

564

565

566

567

568

569

570

571

572

573

574

575

Here we briefly discuss the perturbations introduced for the short-lived N-reservoirs, namely HONO, HNO4 and N<sub>2</sub>O<sub>5</sub>, where the chemical budget terms for all three species are provided in Table S2 in the supplementary material. For HONO it should be noted that many tropospheric CTMs have difficulty in simulating observed mixing ratios (e.g. Goncalves et al, 2012) suggesting missing (heterogeneous) source terms. The global production for HONO is an order of magnitude less than that for the other short-lived N reservoirs. At 1° x 1° there is ~10% more chemical production of HONO in high NO<sub>x</sub> regions and no appreciable effect in the low NO<sub>x</sub> regions. Thus the impact of increased resolution on HONO production is rather small, which is surprising considering the higher NO mixing ratios that occur in high NO<sub>x</sub> regions (c.f. Fig 4). The muted response is due to competing oxidative processes which effectively lower the OH available for forming HONO. For HNO<sub>4</sub>, approximately the same mass of N cycles through this species as for PAN, although the shorter lifetime means that is it more important at regional scale. Again, the impact of resolution on this species is small, where decreases in [HO<sub>2</sub>] result in no significant net change in production for the NH. The most significant changes occur for the global production and heterogeneous conversion of N<sub>2</sub>O<sub>5</sub>, with enhanced chemical production of ~12% at global scale, increasing the heterogeneous sink term by ~6%, although the changes in the total mass of N converted are small. In general, this is due to an increase in the production of the NO<sub>3</sub> radical by ~10% at 1° x 1° (not shown) resulting in enhanced N<sub>2</sub>O<sub>5</sub> mixing ratios.

576577

# 6 Implications for tropospheric CH<sub>2</sub>O retrieval

578579580

581

582

583

584

585

586

587

588

589

590

591

592

593

594

595

596

597

The implications of higher resolution for the global distribution of  $CH_2O$  are rather modest. In Table 8, we show zonally integrated chemical production and destruction terms for  $CH_2O$ , which suggests changes of the order of a few percent at global scale. The most notable difference is the increase in the cumulative deposition term of ~4% at 1° x 1°, thus reducing the atmospheric lifetime of  $CH_2O$  in TM5-MP. Again this low impact shows that the increase in the temporal variability of the meteorological data at 1° x 1°, and thus the local variability of cloud Surface Area Data, only changes the net deposition term by a few percent. Even though the temporal distribution of the surface mixing ratios shows more variability at 1° x 1° due to the better representation of regional pre-cursor sources terms (e.g.) isoprene and terpene, only moderate improvements occur to the simulated profiles and total columns due to changes in transport. For instance, when analysing individual production terms (not given) for the tropics, decreases are related to small changes in the dominating chemical source terms (e.g. oxidation of  $CH_3OOH$ ; ~3-5 Tg less  $CH_2O$   $yr^{-1}$ ). For the chemical destruction term, the relative insensitivity of the photolysis of  $CH_2O$  towards resolution (similar to  $J_{O3}$ ; c.f. Fig S4) results in small net decreases in line with changes in the chemical production term.

Figure 10 compares monthly mean tropospheric profiles of CH<sub>2</sub>O measured during INTEX-B (Singh et al., 2009) with those from both TM5-MP simulations for March to May 2006. In general, there is a fair representation of the vertical gradient of CH<sub>2</sub>O by TM5-MP for all months shown, although surface mixing ratios are typically too high suggesting loss by deposition is under-estimated or that the chemical production term is too. Moreover, there appears to be a missing (chemical) source term in the UTLS in TM5-MP leading to

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





16

a  $\sim$  30-50% ( $\sim$ 0.05 ppb) low bias above 600hPa, therefore no significant improvement to the under-estimation in the SH CH<sub>2</sub>O column in TM5-MP occurs compared to previous versions (Zeng et al, 2015). Comparing profiles shows that the changes in the vertical distribution of CH<sub>2</sub>O at 1° x 1° are minimal in the chemical background compared to 3° x 2°, with the main differences originating from more efficient transport out of source regions (c.f. March). These findings are further confirmed by the comparisons of TM5-MP against TexAQS II measurements for September 2006 (Figure S13).

603 604 605

598

599

600

601

602

## 7 Implications for tropospheric SO<sub>2</sub> retrieval

606 607

608 609

610

611

612

613

614

615

616

617

618

619

620

621

622

623

624

625

626

627

628

629

630

631

632

633

634

635

636

637

In Figure 11, we compare weekly [SO<sub>2</sub>] for 2006 at number of EMEP sites in Austria (ATO2, Forest), the Netherlands (NL09, Rural), Great Britain (GB43, Rural) and Spain (ES10, Rural), with most being positioned away from strong point sources. For SO<sub>2</sub> in Europe, the main source is primarily anthropogenic (e.g. Energy Sector) followed by oxidation to sulphate, with the tropospheric lifetime varying from ~2 days during winter and ~19 hours during summer (Lee et al, 2011). High [SO<sub>2</sub>] has been observed throughout the EMEP network in e.g. The Netherlands and Spain, which is significantly higher than that measured in Central Europe (Tørseth et al., 2012). Although the measurement uncertainty is somewhat site specific due to the different methodology employed, it is typically around ~1.3 ug/m<sup>3</sup> (e.g. Hamad et al., 2010). Comparing weekly averages shows that for most sites shown there is a significant low bias at 3° x 2°, indicating inaccuracies in the MACC emission inventory and the effect of coarsening to the model resolution. At 1° x 1° significant improvements occur as a result of the better temporal resolution of the emission sources. Table 9 provides an overview of the changes in the seasonal biases for all of the EMEP sites that measure hourly [SO<sub>2</sub>], with the biases calculated for the overpass time of tropOMI aggregated on a weekly basis. Improvements occur at 1° x 1° for ~20% of the sites during both seasons, with the majority (~50%) of sites showing no significant improvement (< 5%). In such instances the local [SO<sub>2</sub>] is determined more by long-range transport (thus sensitive to wash-out) rather than a local emission source, where strong mitigation practises have been implemented in Europe over the last few decades reducing resident [SO<sub>2</sub>] significantly (Tørseth et al., 2012). For some sites there is a notable increase in biases at 1° x 1° (20% DJF, 25% JJA) indicating that too strong local emission sources occur in the MACC inventories (e.g. ES13 and GR01). For others (e.g. ES08 and NL07) significantly low biases occur suggesting the opposite problem. Finally, for the vertical profiles we make comparisons against monthly mean composites assembled from measurements taken during INTEX-B (Figure S14) and TexAQS II (Figure 12) as for the other trace species. For the more pristine locations there are typically low biases at 3° x 2° for all months, especially at the surface during March indicating a significant under-estimation in the emission fluxes of SO2. Increasing to 1° x 1° only provides an improved correlation for March, due to the transport in the FT being described better as for that shown for NO<sub>2</sub> in Fig. 6. For April, the comparison shows a significant underestimation in the column for both simulations, where corresponding comparisons of the vertical profiles of Di-Methyl Sulphide, which acts as a key source of SO<sub>2</sub> in the Equatorial Pacific (Alonza Gray et al., 2011), also show significant low biases (not shown). For May again no significant improvement occurs, although both simulations capture the peak in SO<sub>2</sub> mixing ratios at the top of the boundary layer. More relevant for satellite based retrievals is the observed column near strong anthropogenic source regions as shown in Figure 13 over Texas during September 2006. Here a clear

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





17

improvement occurs at 1° x 1°, with the low bias in the boundary layer being reduced significantly although the integrated column is still too low. Again this is due to the under-estimation in the source emission fluxes in the anthropogenic emission inventory employed.

640 641

638

639

674

675

676

642 **8 Conclusions** 643 644 In this paper we have provided a comprehensive description of the high-resolution 1° x 1° version of TM5, 645 which is to be used for the purpose of providing a-priori columns for the satellite retrieval of trace gas columns 646 of NO<sub>2</sub>, CH<sub>2</sub>O and SO<sub>2</sub>. By performing identical simulations at both 3° x 2° and 1° x 1° horizontal resolution and 647 comparing the resulting global distributions of trace gas species, photolysis frequencies and chemical budget 648 terms, we quantify and validate both the near-surface and vertical distributions for the evaluation year of 2006. Comparing the seasonal distribution in 222Rn shows that differences of ±20% occur at global scale, with 649 650 significantly larger differences for specific coastal regions and tropical oceans. In order to assess the changes in 651 convective activity above strong NO<sub>x</sub> sources, we show that differences of between ~2-10% (~10-20%) exist for 652 the Northern mid-latitudes (tropics) at higher resolution, with both weaker and stronger upwelling occurring 653 depending on the region and the season. The differences are site specific being affected by location orography. 654 The impact on global monthly mean  $J_{O3}$  and  $J_{NO2}$  surface values over a range of conditions is limited to ~2% and 655 ~5-10%, respectively. This is surprising considering the larger variability in cloud cover and surface albedo at 1° 656 x 1°. Examining changes in J<sub>O3</sub> and J<sub>NO2</sub> which occur throughout the tropospheric column reveals that significant 657 differences of >10% can occur at the top of the boundary layer at tropical locations. Such modest changes 658 associated with this dominant loss term result in the change in the integrated chemical budget terms to be rather 659 low. 660 Analysing the chemical budget terms for tropospheric O<sub>3</sub> shows (i) a reduction in the Stratosphere-Troposphere 661 exchange flux of ~7% to 597 Tg O<sub>3</sub> yr<sup>-1</sup>, (ii) a repartitioning of the contribution from Stratospheric down welling 662 in both the Northern and Southern hemispheres, (iii) no significant change in the tropospheric burden of O<sub>3</sub> and 663 (iv) modest changes in the integrated chemical production and destruction terms. Comparing simulated mixing 664 ratios against surface measurements in Europe shows that the positive bias present in TM5 decreases by ~20% at 665 1° x 1° between 2-5 ppb month<sup>-1</sup>. This bias persists throughout the vertical column across diverse global regions, 666 although the vertical gradient in tropospheric O<sub>3</sub> is captured quite well. 667 For NO and NO<sub>2</sub> increasing horizontal resolution results in only modest differences in the zonal mean recycling 668 terms and the loss of O<sub>3</sub> by chemical titration. Comparisons against surface measurements in Europe shows that 669 there is a consistent negative bias in NO weekly concentrations of a few  $\mu g \ m^{-3}$  associated with both too high 670 surface O<sub>3</sub> (enhanced NO titration) and the accuracy of the emission inventories. For NO<sub>2</sub>, the biases in the 671 weekly concentrations are larger and can be both positive and negative. Increasing horizontal resolution has little 672 effect on improving the NO biases, but results in improvements for NO2 at ~35% of the available sites, with 673 ~45% of sites showing limited changes. For the tropospheric column the improvement in the comparisons in

only a of the order of few percent, with a significant under-estimation in both NO and NO<sub>2</sub> throughout the

tropospheric column. Analysing the NO/NO2 ratio and comparing against observations shows that although

partitioning is captured in the boundary layer there is a significant overestimation in the upper troposphere.

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





Finally for  $CH_2O$  and  $SO_2$ , which can also be retrieved from satellite measurements, the effect of increased resolution is rather modest due to compensating changes towards the chemical budget terms. When compared against observations there is a persistent low bias for tropospheric  $CH_2O$  due to missing production terms especially on the Free Troposphere. For  $SO_2$  comparison with surface observations in Europe show lower biases at ~20% of sites due to more accurate local emission fluxes, whereas for the majority of cases (~50%) there is no significant change. Comparing vertical profiles shows a significant under-estimation in the tropospheric column likely associated with either precursors or direct emission terms.

## **Code Availability**

The TM5-MP code can be downloaded from the SVN server hosted at KNMI, The Netherlands. A request to generate a new user account for access can be made by e-mailing <a href="mailto:sager@knmi.nl">sager@knmi.nl</a>. Any new user groups need to agree to the protocol set out for use, where it is expected that any developments are accessible to all users after publication of results. Attendance at 9-monthly TM5 international meetings is encouraged to avoid duplicity and conflict of interests.

## Acknowledgements

We thank M. van Weele for processing the MSR2 stratospheric ozone data record used for constraining the overhead O<sub>3</sub> field and T. P. C. van Noije for updating the SO<sub>x</sub> emission estimates. We thank V. Huijnen for providing estimates on the heterogeneous uptake co-efficients.

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





19

Table 1: Details of the reaction rate data applied for  $NO_x$  radical-radical reactions and nitrogen reservoirs. The  $k_0$  terms are multiplied by the relevant air density to calculate the correct forward and backward rate constants. The reaction data and stiochiometery is taken from Atkinson et al. (2004) accommodating the latest evaluation at <a href="http://iupac.pole-ether.fr">http://iupac.pole-ether.fr</a>.

Reactants	Products	Rate parameters
$NO + O_3$	NO <sub>2</sub>	3.0 x 10 <sup>-12</sup> *exp(-1500/T)
$NO_2 + O_3$	$NO_3$	1.4 x 10 <sup>-13</sup> *exp(-2470/T)
$NO + HO_2$	$NO_2 + OH$	$3.3 \times 10^{-12} * \exp(270/T)$
$NO + CH_3O_2$	$CH_2O + HO_2 + NO_2$	2.8 x 10 <sup>-12</sup> *exp(300/T)
$OH + NO_2$	$HNO_3$	$k_0=3.2 \times 10^{-30}*(300/T)^{4.5}$
$NO + NO_3$	$NO_2 + NO_2$	$k_{\infty} = 3.0 \times 10^{-11}$ 1.8 x $10^{-11} *exp(110/T)$
$NO_2 + NO_3$	$N_2O_5$	$k_0=8.0 \times 10^{-27}*(300/T)^{3.5}$
		$k_{\infty}=3.0 \times 10^{-11}*(300/T)^{1.0}$
$N_2O_5 + M$	$NO_2 + NO_3$	$k_0=1.3 \times 10^{-3}*(300/T)^{3.5}*$
		exp(-11000/T)
		$k_{\infty} = 9.7 \times 10^{-14} * (300/T)^{-0.1} *$
		exp(-11080/T)
$HO_2 + NO_2$	$\mathrm{HNO}_4$	$k_0 = 1.4 \times 10^{-31} * (300/T)^{3.1}$
		$k_{\infty} = 4.0 \text{ x } 10^{-12}$
$HNO_4 + M$	$HO_2 + NO_2$	$k_0 = 4.1 \times 10^{-5} * exp(-10650/T)$
		$k_{\infty} = 6.0 \times 10^{15} * \exp(-11170/T)$
OH + HNO <sub>4</sub>	$NO_2$	$1.3 \times 10^{-12} * \exp(380/T)$
OH + NO + M	HONO	$k_0 = 7.0 \times 10^{-31} * (300/T)^{4.4}$
		$k_{\infty}=3.6 \times 10^{-11}*(300/T)^{0.1}$
HONO + hυ	OH + NO	
OH + HONO	$NO_2$	2.5 x 10 <sup>-12</sup> *exp(260/T)
$NO_2 + CH_3C(O)O_2$	PAN	$k_0 = 3.28 \times 10^{-28} * (300/T)^{6.87}$
		$k_{\infty} = 1.125 \text{ x } 10^{-11} * (300/\text{T})^{1.105}$
PAN	$NO_2 + CH_3C(O)O_2$	$k_0 = 1.1 \times 10^{-5} * exp(-10100/T)$
		$k_{\infty} = 1.9 \times 10^{17} * \exp(-14100/T)$

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





20

PAN + hv	$CH_3C(O)O_2 + NO_2$	
	$CH_3O_2 + NO_3$	
$CH_3O_2 + NO_2$	CH <sub>3</sub> O <sub>2</sub> NO <sub>2</sub>	$k_0 = 2.5 \times 10^{-30} * (300/T)$
$CH_3O_2NO_2$	$CH_3O_2 + NO_2$	$k_{\infty} = 1.8 \times 10^{-11}$ $k_0 = 9.0 \times 10^{-5} * exp(-9690/T)$
$NO_3 + HO_2$	HNO <sub>3</sub>	$k_{\infty} = 1.1 \times 10^{16} * \exp(-10560/T)$ $4.0 \times 10^{-12}$
701	222,103	

Published: 11 July 2016

707 708

709

© Author(s) 2016. CC-BY 3.0 License.





703 Table 2: Details of updates made to the reaction data and stoichiometry of the modified CB05 chemical mechanism for other reactions. Data is taken from the following: [1] Atkinson et al. (2004) accommodating the 705 latest evaluation at <a href="http://iupac.pole-ether.fr">http://iupac.pole-ether.fr</a>, [2] Branching ratio (R) equal to 1/(1+498.\*exp(-1160./T), [3] Yarwood et al. (2005), [4] Sander et al. (2011), [5], Atkinson et al. (2006), [6] Emmons et al. (2010), [7] Hauglustaine et al. (2014), [8] Rate assumed equal to NH<sub>2</sub> analogue, [9] assumed to be equal to HNO<sub>4</sub> after Browne et al. (2011).

Reactants	Products	Rate expression	Ref.
$CH_3O_2 + HO_2$	CH₃OOH	3.8 x 10 <sup>-13</sup> *exp(750/T)*R	[1],[2]
$CH_3O_2 + HO_2$	$CH_2O$	3.8 x 10 <sup>-13</sup> *exp(750/T)	[1],[2]
		*(1-R)	
$CH_3O_2 + CH_3O_2$	$1.37\text{CH}_2\text{O} + 0.74\text{HO}_2 + 0.63\text{CH}_3\text{OH}$	9.5 x 10 <sup>-14</sup> *exp(390/T)	[3],[4]
$OH + C_3H_8$	$IC_3H_7O_2$	7.6 x 10 <sup>-12</sup> *exp(-585/T)	[5],[6]
$NO + IC_3H_7O_2$	$0.82\text{CH}_3\text{COCH}_3 + \text{HO}_2 + 0.27\text{ALD2} \\ + \text{NO}_2$	4.2 x 10 <sup>-12</sup> *exp(180/T)	[6]
$HO_2 + IC_3H_7O_2$	ROOH	7.5 x 10 <sup>-13</sup> *exp(700/T)	[6]
$OH + C_3H_6$	$\mathrm{C_3H_6O_2}$	$k_0 = 8.0 \times 10^{-27} * (-300/T)^{3.5}$	[5],[6]
		$k_{\infty}$ =3.0 x 10 <sup>-11</sup> * $(-300/T)^{1.0}$	
$NO_3 + C_5H_8$	$0.2ISPD + XO_2 + 0.8HO_2 +$	2.95 x 10 <sup>-12</sup> *exp(465/T)	[5]
	0.8ORGNTR + 0.8ALD2 +		
	$2.4 \text{ PAR} + 0.2 \text{ NO}_2$		
$NO + C_3H_6O_2$	$ALD2 + CH_2O +$	4.2 x 10 <sup>-12</sup> *exp(180/T)	[6]
	$HO_2 + NO_2$		
$HO_2 + C_3H_6O_2$	ROOH	$7.5 \times 10^{-13} * \exp(700/T)$	[6]
$NO_3 + DMS$	$SO_2 + HNO_3$	1.9 x 10 <sup>-13</sup> *exp(520/T)	[1]
$NH_2 + OH$		$3.4 \times 10^{-11}$	[E]
$NH_2 + HO_2$	$NH_3$	$3.4 \times 10^{-11}$	[4],[7]
$NH_2 + O_3$	$\mathrm{NH_2O_2}$	4.3 x 10 <sup>-12</sup> *exp(-930/T)	[4],[7]
$NH_2 + O_2$	NO	6.0 x 10 <sup>-21</sup>	[1],[7]
$NH_2O_2 + NO$	$NH_2 + NO_2$	4.0 x 10 <sup>-12</sup> *exp(450/T)	[7],[8]
$NH_2O_2 + O_3$	$\mathrm{NH}_2$	4.3 x 10 <sup>-12</sup> *exp(-930/T)	[7],[8]
$NH_2O_2 + HO_2$	$\mathrm{NH}_2$	$3.4 \times 10^{-11}$	[8]
$CH_3O_2NO_2 + h\upsilon$	$CH_3O_2 + NO_2$		[9]

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





22

	$CH_3O_2NO_2 + hv$	$CH_2O + HO_2 + NO_3$	[9]
	$HO_2$ + aero $NO_3$ + aero	$0.5\mathrm{H}_2\mathrm{O}_2$ $\mathrm{HNO}_3$	[6]
710	1103 + acro	111103	[o]

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





23

712 Table 3: The zonally segregated emission totals introduced into TM5-MP for the year 2006. All organic 113 hydrocarbons are given in Tg C yr $^{-1}$ , except for CO, CH $_2$ O and CH $_3$ OH and all NO $_x$  emissions are 114 introduced as NO. No direct emissions occur for HNO $_3$ , PAN, ORGNTR, HONO, N $_2$ O $_5$ , NO $_2$ , 115 introduced as NO. 715  $CH_3O_2NO_2$  or  $O_3$ .

Species (Tg/Year)	Global	30-90°S	30S-30°N	30-90°N
CO	1081.0	24.4	755.1	301.27
$NO_x$ (as N)	49.0	1.5	24.0	23.6
$SO_2$	117.0	3.0	49.2	64.3
DMS (as S)	19.2	6.7	9.3	3.2
$NH_3$	56.6	3.1	27.9	25.6
CH <sub>2</sub> O	13.5	0.3	10.5	2.7
PAR	34.1	0.7	18.5	14.9
OLE	22.4	0.9	16.6	4.9
ALD2	13.4	0.4	11.2	1.8
CH₃CHCHO	2.2	0.0	1.2	1.0
CH <sub>3</sub> OH	100.7	3.3	82.5	14.9
CH <sub>3</sub> CH <sub>2</sub> OH	70.4	2.8	52.6	15.1
$C_2H_4$	25.9	1.0	19.0	5.9
$C_2H_6$	6.1	0.3	5.3	1.5
$C_3H_8$	5.6	0.4	3.6	1.6
$C_3H_6$	19.6	0.9	14.8	3.9
CH <sub>3</sub> COCH <sub>3</sub>	27.4	0.8	22.0	4.6
НСООН	1.8	0.0	1.5	0.3
CH₃COOH	7.1	0.1	6.0	1.0
$C_5H_8$	510.0	23.2	441.9	45.0
$C_{10}H_{16}$	85.4	2.3	70.2	12.9

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





Table 4: The tropospheric chemical budget terms for  $O_3$  during 2006 for the  $1^\circ$  x  $1^\circ$  simulation given in Tg  $O_3$  yr $^{-1}$ . The associated percentage changes are given when comparing against the  $3^\circ$  x  $2^\circ$  simulation ( $1^\circ$  x  $1^\circ$ / $3^\circ$  x  $2^\circ$ ). The definition of the chemical tropopause and the calculation of the STE are defined using the methodology outlined in Stevenson et al. (2006). The stratospheric nudging term refers to total change in the mass of  $O_3$  in the stratospheric column when nudging towards observational constraints from the MSR (Huijnen et al., 2010). The contribution to each term from the SH extratropics/tropics/NH extra-tropics (defined as  $90\text{-}30^\circ\text{S}/30^\circ\text{S}-30^\circ\text{N}/30\text{-}90^\circ\text{N})$  are provided. The fraction of the tropospheric burden originating from the stratosphere is also given.

Term	Global	%	SH	%	Tropics	%	NH	%
Net STE	579	-6.7	166		396		16	
Strat. Nudging	1440	-0.7	-224	2.8	1615	-	49	5.8
Trop.Chem.Prod	5532	-1.9	389	-2.2	3938	-3.5	1206	-2.2
Trop.Chem.Loss	5162	-2.4	440	-1.0	3869	-2.5	853	-2.8
$BO_3$	378	-2.0	72	1.7	203	-2.3	104	-3.4
Strat BO <sub>3</sub>	80	-2.0	23	9.1	38	-6.5	24	-2.0
Deposition	949	0.8	115	0.6	465	-	369	1.9
O <sub>3</sub> S Deposition	97	5.0	19	7.5	37	-1.2	42	10.0

Table 5: The annual NO to NO $_2$  re-cycling terms involving peroxy-radicals given in Tg N yr $^1$  for 2006 at 1° x 1° resolution. In mCB05v2 XO $_2$  represents lumped alkyl-peroxy radicals (Yarwood et al, 2005). The RO $_2$  term is an aggregate of numerous specific peroxy-radical conversion terms in the modified CB05 mechanism (Williams et al., 2013; Tables 1 and 2). Also provided are the approximate percentage differences when comparing with 3° x 2° (1° x 1°/3° x 2°). The chemical tropopause is defined using the methodology outline in Stevenson et al. (2006).

Reaction	Global	%	SH	%	Tropics	%	NH	%
$NO + HO_2$	1058	-1.2	79	-1.2	740	-1.9	239	0.8
$NO + CH_3O_2$	407	-2.2	31	-2.6	294	-2.8	82	-2.0
$NO + XO_2$	147	-2.1	7	-3.6	111	-2.6	29	-
$NO + RO_2$	9.4	-4.4	0.4	-2.6	6.3	-4.4	2.7	-4.2
$NO + O_3$	5403	0.1	518	7.5	2933	-3.9	1953	4.9

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





25

740

Table 6: The seasonal mean absolute biases of weekly [NO] ( $\mu$ g m<sup>-3</sup>) composed from daily measurements at 13:00 for DJF and JJA (measurements-model). Values are shown for both the 3° x 2° and 1° x 1° simulations. Those with differences < 5% are considered to exhibit no discernible change in the bias.

EMEP	Lat	Lon	DJF	DJF	JJA	JJA
Station			3° x 2°	1° x 1°	3° x 2°	1° x 1°
CH01	46.32	7.59	-0.01	-0.01	0.00	-0.01
CZ03	49.35	15.50	-4.05	-3.30	-1.61	-1.35
DE43	47.48	11.10	-2.37	-2.36	-0.47	-0.48
DK05	54.44	10.44	-2.51	-2.61	-1.29	-1.51
ES07	58.23	21.49	-3.80	-3.84	-1.45	-1.48
ES08	43.26	-4.51	-2.08	-2.09	-1.00	-1.01
ES09	41.16	-3.80	-0.93	-0.93	-1.07	-1.07
ES10	38.28	3.19	-1.14	-1.24	-0.75	-0.88
ES11	39.50	-6.55	-1.07	-1.07	-0.44	-0.45
ES12	41.17	-1.60	-1.34	-1.34	-0.95	-0.95
ES13	41.24	-5.52	-2.50	-1.90	-0.74	-0.62
ES14	39.31	0.43	-2.21	-2.20	-1.27	-1.27
ES15	43.13	-4.21	-1.62	-1.61	-0.99	-1.00
ES16	43.37	-7.41	-2.39	-2.39	-1.11	-1.11
FR13	46.39	0.11	-1.90	-1.94	-0.52	-0.52
FR15	55.18	0.45	-3.09	-3.04	-1.51	-1.58
GB02	50.35	-3.12	-1.23	-1.23	-0.93	-0.92
GB13	54.20	-3.42	-1.28	-1.32	-0.58	-0.55
GB14	52.30	-0.48	-3.03	-3.04	-0.98	-0.98
GB31	53.23	-3.11	-1.74	-1.74	-0.90	-0.91
GB37	50.47	-1.45	-3.09	-3.08	-1.22	-1.21
GB38	51.13	0.10	-2.87	-2.78	-1.92	-1.72
GB44	51.17	-3.20	-1.65	-1.45	-0.27	-0.67
GB45	52.17	0.17	-1.80	0.11	0.20	0.19
GB51	52.33	0.46	-3.68	-3.42	-1.29	-1.17
NL91	52.18	4.30	-4.47	-3.51	-1.98	-1.86

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





26

750 Table 7: As for Table 5 except for  $NO_2$ .

			D. 17	***	** 1
Lat	Lon	DJF	DJF	JJA	JJA
					1° x 1°
					-2.13
					-0.01
.,			0.00		0.34
47.48	11.10	-2.37	-2.36	-0.49	-0.48
54.44	10.44	6.07	5.56	1.04	-0.02
58.23	21.49	-1.13	-1.79	-0.96	-1.15
43.26	-4.51	-2.01	-2.10	-0.93	-1.01
41.16	-3.80	-0.95	-0.94	-1.07	-1.07
42.19	3.19	2.18	1.46	0.92	-0.01
38.28	-6.55	-1.08	-1.08	-0.44	-0.44
39.50	-1.60	-1.36	-1.35	-0.95	-0.95
41.17	-5.52	-2.51	-0.29	-0.74	0.19
41.24	0.43	-2.22	-2.21	-1.27	-1.26
39.31	-4.21	-1.64	-1.63	-0.99	-0.99
43.13	-7.41	-2.40	-2.39	-1.11	-1.11
59.46	21.22	0.79	-0.91	-0.10	-0.53
60.31	27.41	10.04	9.60	1.70	0.92
62.35	24.11	0.40	0.34	0.31	0.17
68.00	24.09	-1.96	-1.95	-0.54	-0.54
43.37	0.11	-3.84	-3.91	-1.78	-1.81
46.39	0.45	3.12	3.42	0.82	0.54
55.18	-3.12	-2.17	-2.13	-1.00	-0.98
50.35	-3.42	5.02	3.93	1.17	1.50
54.20	-0.48	-1.71	-1.71	-0.90	-0.90
52.30	-3.11	-3.10	-3.08	-1.25	-1.25
53.23	-1.45	-4.05	-4.03	-2.20	-2.19
50.47	0.10	6.21	6.69	0.61	1.42
51.13	-3.20	6.28	6.00	3.48	1.94
52.17	-0.17	12.56	16.60	4.11	4.50
41.45	42.49	0.50	2.07	0.40	1.05
53.2	6.16	-3.21	-1.63	-1.18	-0.19
51.32	5.51	3.48	3.52	0.92	-0.29
52.18	4.30	11.81	6.42	1.48	-0.81
	54.44 58.23 43.26 41.16 42.19 38.28 39.50 41.17 41.24 39.31 43.13 59.46 60.31 62.35 68.00 43.37 46.39 55.18 50.35 54.20 52.30 53.23 50.47 51.13 52.17 41.45 53.2 51.32	46.32 7.59   49.35 15.50   47.48 11.10   54.44 10.44   58.23 21.49   43.26 -4.51   41.16 -3.80   42.19 3.19   38.28 -6.55   39.50 -1.60   41.17 -5.52   41.24 0.43   39.31 -4.21   43.13 -7.41   59.46 21.22   60.31 27.41   62.35 24.11   68.00 24.09   43.37 0.11   46.39 0.45   55.18 -3.12   50.35 -3.42   54.20 -0.48   52.30 -3.11   53.23 -1.45   50.47 0.10   51.13 -3.20   52.17 -0.17   41.45 42.49   53.2 6.16   51.32 5.51	46.32 7.59 -0.04   49.35 15.50 -4.04   47.48 11.10 -2.37   54.44 10.44 6.07   58.23 21.49 -1.13   43.26 -4.51 -2.01   41.16 -3.80 -0.95   42.19 3.19 2.18   38.28 -6.55 -1.08   39.50 -1.60 -1.36   41.17 -5.52 -2.51   41.24 0.43 -2.22   39.31 -4.21 -1.64   43.13 -7.41 -2.40   59.46 21.22 0.79   60.31 27.41 10.04   62.35 24.11 0.40   68.00 24.09 -1.96   43.37 0.11 -3.84   46.39 0.45 3.12   55.18 -3.12 -2.17   50.35 -3.42 5.02   54.20 -0.48 -1.71   52.30	50.30 4.59 10.56 1.27   46.32 7.59 -0.04 -0.03   49.35 15.50 -4.04 0.03   47.48 11.10 -2.37 -2.36   54.44 10.44 6.07 5.56   58.23 21.49 -1.13 -1.79   43.26 -4.51 -2.01 -2.10   41.16 -3.80 -0.95 -0.94   42.19 3.19 2.18 1.46   38.28 -6.55 -1.08 -1.08   39.50 -1.60 -1.36 -1.35   41.17 -5.52 -2.51 -0.29   41.24 0.43 -2.22 -2.21   39.31 -4.21 -1.64 -1.63   43.13 -7.41 -2.40 -2.39   59.46 21.22 0.79 -0.91   60.31 27.41 10.04 9.60   62.35 24.11 0.40 0.34   68.00 24.09	50.30 4.59 10.56 1.27 1.69   46.32 7.59 -0.04 -0.03 -0.02   49.35 15.50 -4.04 0.03 -1.62   47.48 11.10 -2.37 -2.36 -0.49   54.44 10.44 6.07 5.56 1.04   58.23 21.49 -1.13 -1.79 -0.96   43.26 -4.51 -2.01 -2.10 -0.93   41.16 -3.80 -0.95 -0.94 -1.07   42.19 3.19 2.18 1.46 0.92   38.28 -6.55 -1.08 -1.08 -0.44   39.50 -1.60 -1.36 -1.35 -0.95   41.17 -5.52 -2.51 -0.29 -0.74   41.24 0.43 -2.22 -2.21 -1.27   39.31 -4.21 -1.64 -1.63 -0.99   43.13 -7.41 -2.40 -2.39 -1.11   59.46 21.

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





Table 8: The tropospheric chemical budget for the  $CH_2O$  given in Tg  $CH_2O$   $yr^1$  during 2006 for the  $1^{\circ}$  x  $1^{\circ}$  simulation. Percentage differences are shown against the corresponding  $3^{\circ}$  x  $2^{\circ}$  simulation.

Budget Term	Global	%	SH	%	Tropics	%	NH	%
CH <sub>2</sub> O CP	1919	-1.1	147	-0.3	1491	-1.0	281	-2.0
$CH_2O$ $CD$ $CH_2O$ $Dep$ .	1739 193	-1.6 3.1	134 15	-0.5 2.0	1349 149	-1.1 3.9	256 29	-2.3

Table 9: The seasonal mean biases of daily [SO<sub>2</sub>] ( $\mu g \ m^{-3}$ ) at 13:00 for DJF and JJA, when taking the difference between measurements-model values. Values are shown for both the 3° x 2° and 1° x 1° simulations. Those with differences < 5% are considered to exhibit no discernible change in the bias.

EMEP	Lat	Lon	DJF	DJF	JJA	JJA
Station			3° x 2°	1° x 1°	3° x 2°	1° x 1°
AT02	47.46	16.46	-3.34	-3.15	-0.89	-0.53
AT05	46.40	12.58	-0.42	-0.41	-0.14	-0.14
AT48	47.50	14.26	-0.64	-0.63	-0.14	-0.15
CZ03	49.35	15.50	-3.52	3.65	-0.69	0.64
ES07	58.23	21.49	1.22	0.73	0.38	0.31
ES08	43.26	-4.51	-2.98	-3.21	-1.19	-1.58
ES09	41.16	-3.80	-0.62	-0.61	-0.42	-0.42
ES10	42.19	3.19	2.37	2.45	1.93	1.53
ES11	38.28	-6.55	-0.63	-0.61	-0.70	-0.70
ES12	39.50	-1.60	-0.47	-0.45	-0.32	-0.32
ES13	41.17	-5.52	-0.81	2.71	-0.78	0.55
ES14	41.24	0.43	-0.70	-0.67	-0.47	-0.47
ES15	39.31	-4.21	-0.40	-0.37	-0.45	-0.46
ES16	43.13	-7.41	-3.84	-3.82	-1.66	-1.66
GB37	52.30	-3.11	-2.92	-2.91	-1.72	-1.72
GB38	53.23	-1.45	2.93	2.75	0.39	1.33
GB43	51.14	-4.42	-1.49	4.77	-2.03	-0.63
GB45	52.17	-0.17	3.87	7.20	1.01	1.77
GR01	38.22	23.50	1.70	2.77	0.74	1.50
NL07	52.50	6.34	2.58	-1.67	0.46	-1.03
NL08	52.70	5.12	1.77	1.56	-0.14	-0.30
NL09	53.2	6.16	2.53	2.16	0.47	0.27

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





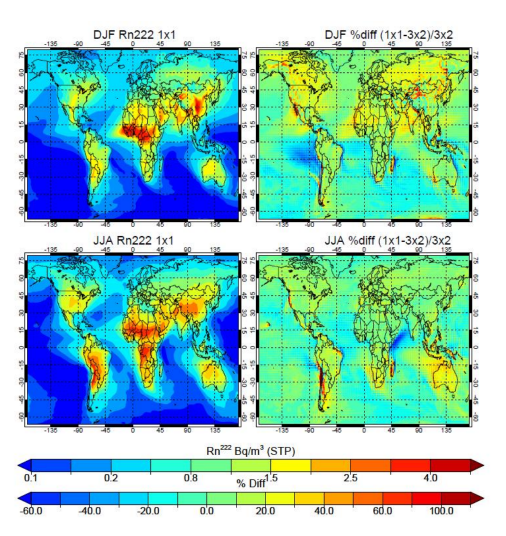


Figure 1: The seasonal distributions of  $Rn^{222}$  between 800-900hPa for DJF (top) and JJA (bottom) for the  $1^{\circ}$  x  $1^{\circ}$  (right) simulation, with the associated percentage differences when compared against the  $3^{\circ}$  x  $2^{\circ}$  simulation.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





29

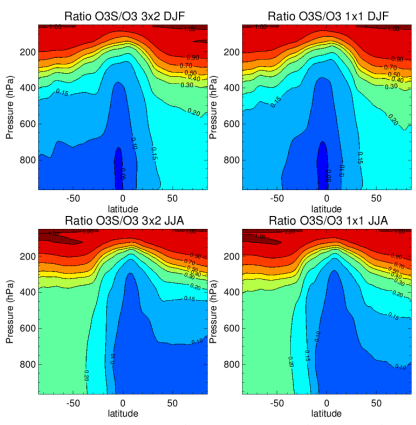


Figure 2: Zonal mean seasonal distribution of the TM5-MP  $O_3S/O_3$  ratio for the 3° x 2° (left) and 1° x1° (right) simulations.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





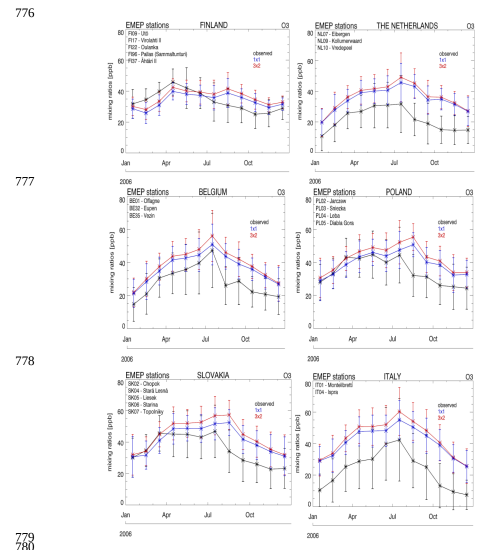


Figure 3: Comparisons of the seasonal variability in TM5-MP mass mixing ratios for surface  $O_3$  against composites of measurements taken across the EMEP monitoring network for 2006. Both the co-located TM5-MP 3° x 2° and 1° x 1° monthly mean values are shown, along with the 1- $\sigma$  variability for Finland, The Netherlands, Belgium, Poland, Slovakia and Italy. Individual stations that are aggregated are given in the panels.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





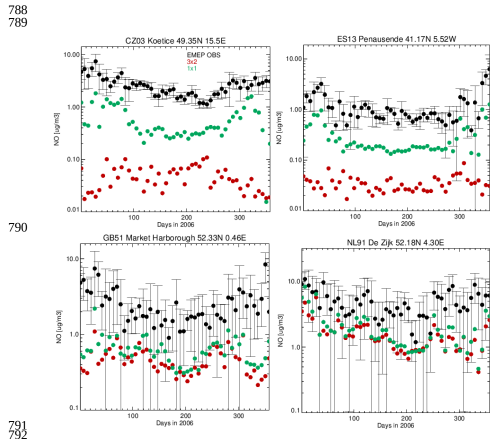


Figure 4: Comparison of TM5-MP weekly NO surface concentrations sampled at 13:00 UT each day during 2006 with observed NO surface concentrations ( $\mu g \ m^{-3}$ ). The selected sites shown are in the Czech Republic (top left), Spain (top right), Great Britain (bottom left) and The Netherlands (bottom right).

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





32

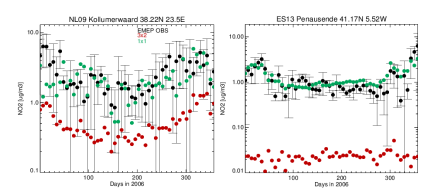


Figure 5: Comparison of weekly TM5-MP  $NO_2$  surface concentrations sampled at 13:00 UT each day during 2006 with observed  $NO_2$  surface concentrations ( $\mu g \ m^{-3}$ ). The selected sites shown are in The Netherlands (left) and Spain (right).

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





33

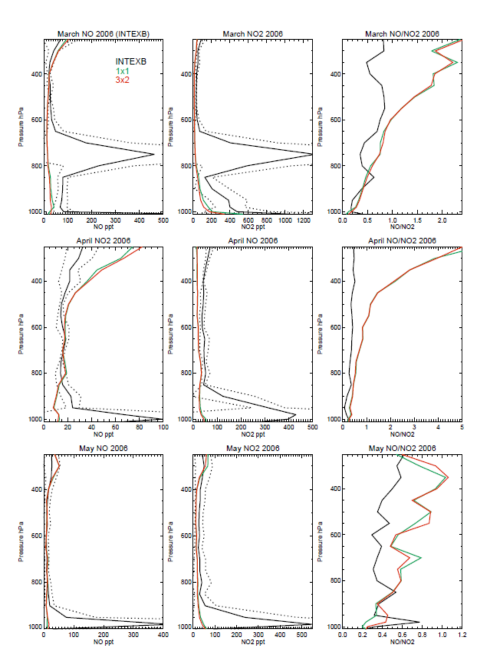


Figure 6: Monthly mean comparisons of NO (left), NO<sub>2</sub> (middle) and the resulting NO/NO<sub>2</sub> ratio from the INTEX-B measurements and TM5-MP simulations. The dotted line represents the  $1-\sigma$  deviation in the mean of the measurements. For details of the locations for each month the reader is referred to Singh et al. (2009).

812 813 814

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





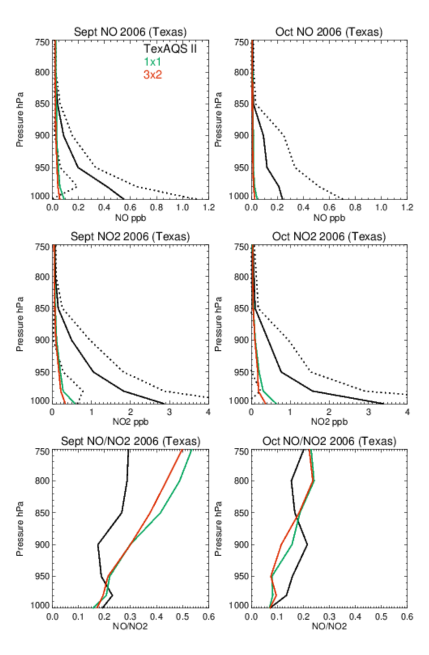


Figure 7: Monthly mean comparisons of NO (left), NO $_2$  (middle) and the resulting NO/NO $_2$  ratio from the TexAQSII campaign during September 2006 and TM5-MP simulations. The dotted line represents the 1- $\sigma$  deviation in the mean of the measurements. For details of the locations for each month the reader is referred to Parrish et al. (2009).

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





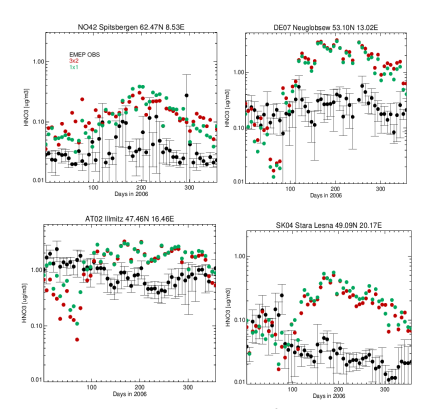


Figure 8: Comparison of weekly  $HNO_3$  concentrations ( $\mu g \ m^{-3}$ ) from both 3° x 2° and 1° x 1° simulations at 4 selected EMEP sites for 2006. The 1- $\sigma$  deviation in the weekly observations are shown as error bars. The selected sites shown are in Norway (top left), Germany (top right), Austria (bottom left) and Slovakia (bottom right).

Published: 11 July 2016

839 840

841 842

© Author(s) 2016. CC-BY 3.0 License.





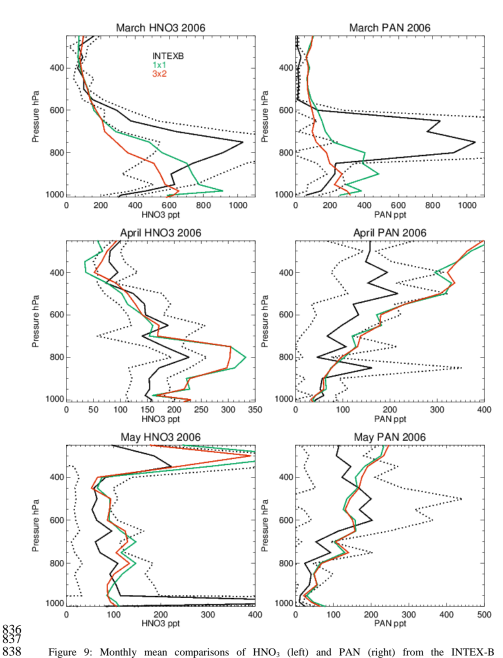


Figure 9: Monthly mean comparisons of  $HNO_3$  (left) and PAN (right) from the INTEX-B measurements and TM5-MP simulations. The dotted line represents the 1- $\sigma$  deviation in the mean of the measurements. For details of the locations for each month the reader is referred to Singh et al. (2009).

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





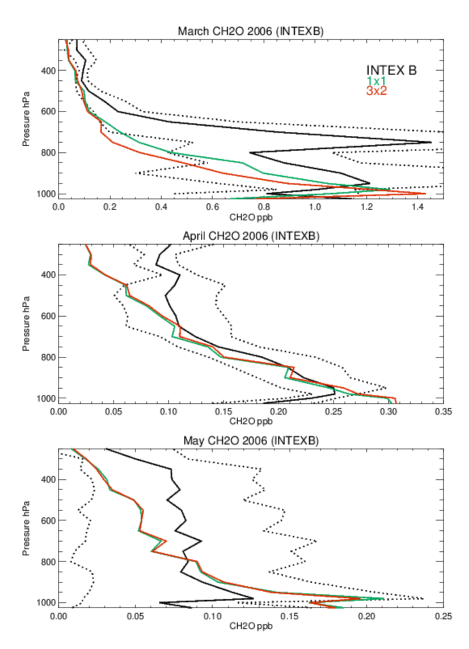


Figure 10: Comparisons of the vertical distribution of  $CH_2O$  from both 3° x 2° and 1° x 1° simulations against measurements made as part of the INTEX B during 2006. The dotted line represents the 1- $\sigma$  deviation in the mean of the measurements. For details on the exact location of the flights the reader is referred to Parrish et al. (2009).

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





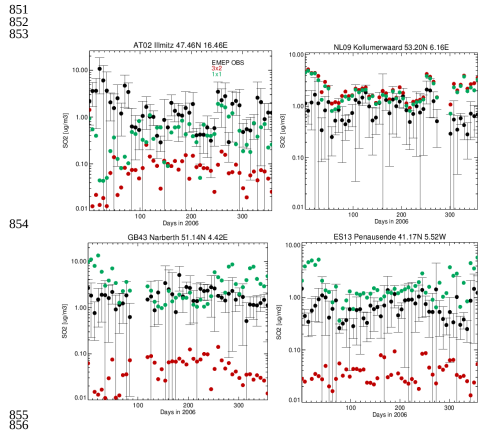


Figure 11: Comparison of hourly  $SO_2$  concentrations ( $\mu g \ m^{-3}$ ) at 13:00 from both the 3° x 2° and 1° x 1° simulations at 4 selected EMEP sites for 2006. The selected sites shown are in Austria (top left), the Netherlands (top right), Great Britain (bottom left) and Spain (bottom right).

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





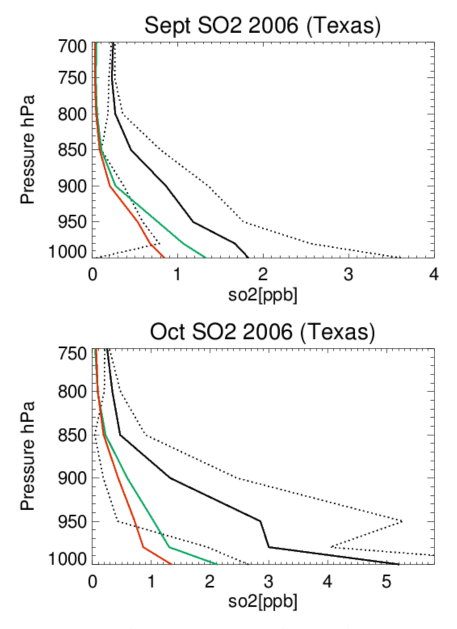


Figure 12: Comparisons of the monthly tropospheric  $SO_2$  profile assembled from data taken during September and October 2006 as part of the Texas-AQS II. The 1- $\sigma$  deviation from the measurements is shown as the dotted line for each of the days. For details of the flight paths the reader is referred to the details given in Parrish et al. (2009).

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





- 870 References
- 871 Aan de Brugh, J. M. J., Schaap, M., Vignati, E., Dentener, F., Kahnert, M., Sofiev, M., Huijnen, V., and Krol,
- 872 M. C.: The European aerosol budget in 2006, Atms. Phys. Chem., 11, 1117-1139, doi:10.5194/acp-11-1117-
- 873 2011, 2011.
- 874 Aas, W., Hjellbrekke, A.-G., Schaug, J., and Solberg, S.: Data quality 1999, quality assurance and field
- comparisons, Kjeller, Norwegian Institute for Air Research, EMEP/CCC Report 6/2001, 2001.
- 876 Abbatt, J. P. D., Lee, A. K. Y., and Thornton, J. A.: quantifying trace gas uptake to tropospheric aerosol: recent
- 877 advances and remaining challenges, Chem. Soc. Rev., 41, 6555–6581, doi:10.1039/c2cs35052a, 2012.
- 878 Alonza Gray, B., Wang, Y., Gu, D., Bandy, A., Mauldin, L., Clarke, A., Alexander, B., and Davis, D. D.:
- 879 Sources, transport, and sinks of SO<sub>2</sub> over the equatorial Pacific during the Pacific Atmospheric Sulfur
- 880 Experiment, J. Atmos. Chem., 68, 27-53, doi: 10.1007/s10874-010-9177-7, 2011.
- Atkinson, R., Baulch, D. L., Cox, R. A., Crowley, J. N., Hampson, R. F., Hynes, R. G., Jenkin, M. E., Rossi, M.
- 882 J., and Tore, J., Evaluated kinetic and photochemical data for atmospheric chemistry: Volume I -gas phase
- reactions of O<sub>x</sub>, HO<sub>x</sub>, NO<sub>x</sub> and SO<sub>x</sub> species, Atmos. Chem. Phys., 4, 1461-1738, 2004.
- Atkinson, R., Baulch, D. L., Cox, R. A., Crowley, J. N., Hampson, R. F., Hynes, R. G., Jenkin, M. E., Rossi, M.
- 885 J., and Tore, J., Evaluated kinetic and photochemical data for atmospheric chemistry: Volume II -gas phase
- reactions of organic species, Atmos. Chem. Phys., 6, 3625-4055, 2006.
- 887 Bândă, N., Krol., M., van Noije, T., van Weele, M., Williams J. E., Le Sager, P., Niemeier, U., Thomason, L.
- 888 and Röckmann, T.: The effect of stratospheric sulfur from Mount Pinatubo on tropospheric oxidizing capacity
- $889 \qquad \text{and methane, J. Geophys. Res. Atmos., } 120, \\ \text{doi:} 10.1002/2014 \\ \text{JDO22137, } 2015.$
- 890 Boersma, K. F., Jacob, D. J., Eskes, H. J., Pinder, R. W., Wang, J. and van der A, R. J.: Intercomparison of
- 891 SCIAMACHY and OMI tropospheric NO2 columns: observing the diurnal evolution of chemistry and emissions
- 892 from space, J. Geophys. Res., 2, 113, 1-14, doi:10.1029/2007JD008816, 2008.
- 893 Boersma, K. F., Eskes, H. J., Dirksen, R. J., van der A, R. J., Veefkind, J. P., Stammes, P., Huijnen, V.,
- 894 Kleipool, Q. L., Sneep, M., Claas, J., Leitao, J., Richter, A., Zhou, Y. and Brunner, D.: An improved
- 895 tropospheric NO<sub>2</sub> column retrieval algorithm for the Ozone Monitoring Instrument, Atmos. Meas. Tech., 4,
- 896 1905-1928, 2011.
- 897 Bregman, B., Segers, A., Krol, M., Meijer, E., and van Velthoven, P.: On the use of mass-conserving wind fields
- 898 in chemistry-transport models, Atmos. Chem. Phys., 3, 447–457, doi:10.5194/acp-3-447-2003, 2003.
- 899 Browne, E. C., Perring, A. E., Wooldridge, P. J., Apel, E., Hall, S. R., Huey, L. G., Mao, J., Spencer, K. M., St.
- 900 Clair, J. M., Weinheimer, A. J., Wisthaler, A., and Cohen, R. C.: Global and regional effects of the
- 901 photochemistry of CH<sub>3</sub>ONO<sub>2</sub>: evidence from ARCTAS, Atmos. Chem. Phys., 11, 4209-4219, 2011.
- 902 Carslaw, D. C. and Beevers, S. D.: Estimations of road vehicle primary NO<sub>2</sub> exhaust emission freations using
- 903 monitoring data in London, Atmos. Environ., 39(1), 167-177, 2005.

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





- 904 De Smedt, I., Muller, J.-F., Stavrakou, T., van der A., R., Eskes, H. and Van Roozendael, M.: Twelve years of
- 905 global observations of formaldehyde in the troposphere using GOME and SCIAMACHY sensors, Atmos. Chem.
- 906 Phys., 8, 4947-4963, 2008.
- 907 Dee, D. P., Uppala, S. M., Simmons, A. J., Berrisford, P., Poli, P., Kobayashi, S., Andrae, U., Balmaseda, M. A.,
- 908 Balsamo, G., Bauer, P., Bechtold, P., Beljaars, A. C. M., van de Berg, L., Bidlot, J., Bormann, N., Delsol, C.,
- 909 Dragani, R., Fuentes, M., Geer, A. J., Haimberger, L., Healy, S. B., Hersbach, H., Hólm, E. V., Isaksen, L.,
- 910 Kållberg, P., Köhler, M., Matricardi, M., McNally, A. P., Monge-Sanz, B. M., Morcrette, J.-J., Park, B.-K.,
- 911 Peubey, C., de Rosnay, P., Tavolato, C., Thépaut, J.-N. and Vitart, F.: The ERA-Interim reanalysis:
- 912 configuration and performance of the data assimilation system, Q. J. Royal Met. Soc, 137, 656, 553-597, 2011.
- 913 Dunlea, E. J., Herndon, S. C., Nelson, D. D., Volkamer, R. M., San Martini, F., Sheehy, P. M., Zahniser,
- 914 M. S., Shorter, J. H., Wormhoudt, J. C., Lamb, B. K., Allwine, E. J., Gaffney, J. S., Marley, N. A., Grutter, M.,
- 915 Marquez, C., Blanco, S., Cardenas, B., Retama, A., Ramos Villegas, C. R., Kolb, C. E., Molina, L. T., and
- 916 Molina, M. J.: Evaluation of nitrogen dioxide chemiluminescence monitors in a polluted urban environment,
- 917 Atmos. Chem. Phys., 7, 2691–2704, 2007, http://www.atmos-chem-phys.net/7/2691/2007/
- 918 Dupuy, E., Urban, J., Ricaud, P., Le Flochmoën, E., Lautié, N., Murtagh, D., De La Noë, J., El Amraoui, L.,
- 919 Eriksson, P., Forkman, P., Frisk, U., Jégou, F., Jiménez, C. and Olberg, M.: Starto-mesospheric measurements of
- 920 carbon monoxide with the Odin Sub-Millimetre Radiometer: Retrieval and first results, Geophys. Res. Letts., 31,
- 921 L20101, doi:10.1029/2004GL020558, 2004.
- 922 Emmons, L. K., Walters, S., Hess, P. G., Lamarque, J.-F., Pfister, G. G., Fillmore, D., Granier, C., Guenther, A.,
- 923 Kinnison, D., Laepple, T., Orlando, J., Tie, X., Tyndall, G., Wiedinmyer, C., Baughcum, S. L., and Kloster, S.:
- 924 Description and evaluation of the Model for Ozone and Related chemical Tracers, version 4 (MOZART-4),
- 925 Geosci. Model Dev., 3, 43–67, doi: 10.5194/gmd-3-43-2010, 2010.
- 926 Evans, M. J. and Jacob, D. J.: Impact of new laboratory studies of N<sub>2</sub>O<sub>5</sub> hydrolysis on global model budgets of
- 927 tropospheric nitrogen oxides, ozone and OH, Geophys. Res. Lett., 32, doi:10.1029/2005GL022469, 2005.
- 928 Fortuin, J. P. F. and Kelder, H.: An ozone climatology based on ozonesonde and satellite measurements. J.
- 929 Geophys. Res., 103, 31709–31734, 1998.
- 930 Gonçalves, M., Dabdub, D., Chang, W. L., Jorba, O. and Baldasanoa, J. M.: Impact of HONO sources on the
- 931 performance of mesoscale air quality models, Atmos. Environ., 54, Pages 168-176
- 932 doi:10.1016/j.atmosenv.2012.02.079, 2012.
- 933 Gonzi, S., Palmer, P. I., Barkley, M., De Smedt, I. and Roozendael, M. V.: Biomass burning emission estimates
- 934 inferred from satellite column measurements of HCHO: Sensitivity to co-emitted aerosol and injection height,
- 935 Geophys. Res. Letts., 38, L14807, 2011.
- 936 Granier, C., Bessagnet, B., Bond, T., D'Angiola, A., Denier van der Gon, H., Frost, G. J., Heil, A., Kaiser, J. W.,
- 937 Kinne, S., Klimont, Z., Kloster, S.-F., Lamarque, J., Liousse, C., Masui, T., Meleux, F., Mieville, A., Ohara, T.,
- 938 Raut, J.-C., Riahi, K., Schultz, M. G., Smith, S. J., Thompson, A., van Aardenne, J., van der Werf, G. R., and
- 939 van Vuuren, D. P.: Evolution of anthropogenic and biomass burning emissions of air pollutants at global and

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





- 940 regional scales during the 1980–2010 period, Climate Change, 109, 163–190, doi: 10.1007/s10584-011-0154-
- 941 115, 2011.
- 942 Grooß, J.-U. and Russell III, J. M.: Technical note: A stratospheric climatology for O<sub>3</sub>, H<sub>2</sub>O, CH<sub>4</sub>, NO<sub>x</sub>, HCl and
- 943 HF derived from HALOE measurements, Atmos. Chem. Phys., 5, 2797–2807, doi:10.5194/acp-5-2797-2005,
- 944 2005.
- 945 Hardacre, C., Wild, O. and Emberson, L.: An evaluation of ozone dry deposition in global scale chemistry
- 946 climate models, Atms. Chem. Phys., 15, 6419-6436, doi:10.5194/acp-15-6419-2015, 2015.
- 947 Hauglustaine, D. A., Balkanski, Y. and Schulz, M.: A global model simulation of present and future nitrate
- 948 aerosols and their direct radiative forcing of climate, Atms. Chem. Phys., 14, 11031-11063, doi:10.5194/acp-14-
- 949 11031-2014, 2014.
- 950 Heckel, A., Kim, S.-W., Frost, G. J., Richter, A., Trainer, M. and Burrows, J. P.: Influence of low spatial
- 951 resolution a priori data on tropospheric NO2 satellite retreivals, Atmos. Meas. Tech., 4, 1805-1820, 2011.
- 952 Huijnen, V., Williams, J., van Weele, M., van Noije, T., Krol, M., Dentener, F., Segers, A., Houweling, S.,
- 953 Peters, W., de Laat, J., Boersma, F., Bergamaschi, P., van Velthoven, P., Le Sager, P., Eskes, H., Alkemade, F.,
- 954 Scheele, R., Nédelec, P., and Pätz, H.-W., The global chemistry transport model TM5: description and
- evaluation of the tropospheric chemistry version 3.0, Geosci. Model Dev., 3, 445-473, 2010.
- 956 Huijnen, V., Williams, J. E., and Flemming, J.: Modeling global impacts of heterogeneous loss of HO<sub>2</sub> on cloud
- 957 droplets, ice particles and aerosols, Atmos. Chem. Phys. Discuss., 14, 8575-8632, doi: 10.5194/acpd-14-8575-
- 958 2014, 2014.
- 959 Jacob, D. J., Prather, M. J., Rasch, P. J., Shia, R. L., Balkanski, Y. J., Beagley, S. R., Bergmann, D. J.,
- 960 Blackshear, W. T., Brown, M., Chiba, M., Chipperfield, M. P., de Grandpré, J., Dignon, J. E., Feichter, J.,
- 961 Genthon, C., W. L. Grose, W. L., Kasibhatla, P. S., Köhler, I., Kritz, M. A., Law, K., Penner, J. E., Ramonet,
- 962 M., Reeves, C. E., Rotman, D. A., Stockwell, D. Z., Van Velthoven, P. F. J., Verver, G., Wild, O., Yang, H and
- 2008 Zimmermann, P.: Evaluation and intercomparison of global atmospheric transport models using 2222Rn and other
- 964 short-lived tracers, J. Geophys. Res., 102(D5), 5953-5970, doi:10.1029/96JD02955, 1997.
- 965 Jacob, J. D.: Heterogeneous chemistry and tropospheric ozone, Atmos. Environ., 34, 2131–2159, 2000.
- 966 Jégou, F., Urban, J., de La Noë, J., Ricaud, P., Le Flochmoën, E., Murtagh, D. P., Eriksson, P., Jones, A.,
- 967 Petelina, S., Llewellyn, E. J., Lloyd, N. D., Haley, C., Lumpe, J., Randall, C., Bevilacqua, R. M., Catoire, V.,
- 968 Huret, N., Berthet, G., Renard, J. B., Strong, K., Davies, J., Mc Elroy, C. T., Goutail, F., and Pommereau, J. P.:
- 969 Technical Note: Validation of Odin/SMR limb observations of ozone, comparisons with OSIRIS, POAM III,
- 970 ground-based and balloon-borne instruments, Atmos. Chem. Phys., 8, 3385-3409, doi:10.5194/acp-8-3385-
- 971 2008, 2008.
- 972 Kim, S.-W., McKeen, S. A., Frost, G. J., S.-H. Lee, S.-H., M. Trainer, M., Richter, A., Angevine, W. M., Atlas,
- 973 E., Bianco, L., Boersma, K. F., Brioude, J., Burrows, J. P., de Gouw, J., Fried, A., Gleason, J., Hilboll, A.,
- 974 Mellqvist, J., Peischl, J., Richter, D., Rivera, C., Ryerson, T., te Lintel Hekkert, S., Walega, J., Warneke,
- 975 C., Weibring, P., and Williams, E.: Evaluations of NOx and highly reactive VOC emission inventories in Texas

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





- 976 and their implications for ozone plume simulations during the Texas Air Quality Study 2006, Atmos. Chem.
- 977 Phys., 11, 11361–11386, doi:10.5194/acp-11-11361-2011, 2011.
- 978 Krol, M., Houweling, S., Bregman, B., van den Broek, M., Segers, A., van Velthoven, P., Peters, W., Dentener,
- 979 F., and Bergamaschi, P.: The two-way nested global chemistry-transport zoom model TM5: algorithm and
- 980 applications, Atmos. Chem. Phys., 5, 417–432, doi:10.5194/acp-5-417-2005, 2005.
- 981 Lawrence, M. G. and Crutzen, P. J.: The impact of cloud particle gravitational settling on soluble trace gas
- 982 distributions, Tellus, 50B, 263-289, 1998.
- 983 Lee, C., Martin, R. V., van Donkelaar, A., Lee, H., Dickerson, R. R., Hains, J. C., Krotkov, N., Richter, A.,
- 984 Vinnikov, K. and Schwab, J. J.: SO<sub>2</sub> emissions and lifetimes: Estimates from inverse modeling using in situ and
- 985 global, space-based (SCIAMACHY and OMI) observations, J. Geophys. Res., 116, D06304,
- 986 doi:10.1029/2010JD014758, 2011.
- 987 Lelieveld, J., Peters, W., Dentener, F. J., and Krol, M. C.: Stability of tropospheric hydroxyl chemistry, J.
- 988 Geophys. Res., 107(D23), 4715, doi:10.1029/2002JD002272, 2002.
- 989 Lin, J. T., Martin, R. V., Boersma, K. F., Sneep, M., Stammes, P., Spurr, R., Wang, P., Van Roozendael, M.,
- 990 Clémer, K., and Irie, H.: Retrieving tropospheric nitrogen dioxide from the Ozone Monitoring Instrument:
- 991 effects of aerosols, surface reflectance anisotropy, and vertical profile of nitrogen dioxide, Atmos. Chem. Phys.,
- 992 14, 1441–1461, doi:10.5194/acp- 14-1441-2014, 2014.
- 993 Lee, C., Martin, R. V., van Donkelaar, A., Lee, H., Dickerson, R. R., Hains, J. C., Krotkov, N., Richter, A.,
- 994 Vinnikov, K. and Schwab, J. J.: SO<sub>2</sub> emissions and lifetimes: Estimates from inverse modeling using in situ and
- 995 global, space-based (SCIAMACHY and OMI) observations, J. Geophys. Res., 116, D06304,
- 996 doi:10.1029/2010JD014758, 2011.
- 997 Marais, E. A., Jacob, D. J., Kurosu, T. P., Chance, K., Murphy, J. C., Reeves, C., Mills, G., Casadio, S., D. B.
- 998 Millet, D. B., Barkley, M. P., Paulot, F., and Mao, J.: Isoprene emissions in Africa inferred from OMI
- 999 observations of formaldehyde columns, Atms. Chem. Phys., 12, 6219-6235, doi:10.5194/acp-12p6219-2012,
- 1000 2012
- 1001 Martin, G. M., Johnson, D. W., and Spice, A.: The measurement and parameterization of effective radius of
- droplets in warm stratocumulus clouds, J. Atmos. Sci., 51, 1823–1842, 1994.
- 1003 Meijer, E. W., van Velthoven, P. F. J., Brunner, D. W., Huntrieser, H., and Kelder, H.: Improvement and
- evaluation of the parameterisation of nitrogen oxide production by lightning, Phys. Chem. Earth, 26, 557–583,
- 1005 2001
- 1006 Meloen, J., Siegmund, P., van Velthoven, P., Kelder, H., Sprenger, M., Wernli, H., Kentarchos, A., Roelofs, G.,
- 1007 Feichter, J., Land, C., Forster, C., James, P., Stohl, A., Collins, W., and Cristofanelli, P.: Stratosphere-
- 1008 troposphere exchange: A model and method intercomparison, J. Geophys. Res., 108(D12), 8256,
- 1009 doi:1029/2002JD002274, 2002.
- 1010 Möllner, A. K., Valluvadasan, S., Feng, L. Sprague, M. K., Okumura, M., Milligan, D. B., Bloss, W.J., Sander,
- 1011 S. P., Martien, P. T., Harley, R. A.,

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





- 1012 Olsen, S. C., McLinden, C. A., and Prather, M. J.: Stratospheric  $N_2O$ -NOy system: Testing uncertainties in a
- three-dimensional framework, J. Geophys. Res., 106, 28771–28784, 2001.
- 1014 Olszyna, K. J., Bailey, E. M., Simonaitis, R. and Meagher, J. F.: O<sub>3</sub> and NO<sub>3</sub> relationships at a rural site, 99(D7),
- 1015 14557–14563, doi:10.1029/94JD00739, 1994.
- 1016 Palmer, P. I., Abbot, D. S., Fu, T-M., Jacob, D. J., Chance, K., Kurosu, P., Guenther, A., Wiedinmyer, C.,
- 1017 Stanton, J. C., Pilling, M. J., Pressley, S. N., Lamb, B. and Sumner, A. L.: Quantifying the seasonal and
- 1018 interannual variability of North American isoprene emissions using satellite observations of the formaldehyde
- 1019 column, J. Geophys. Res., 111, D12315, doi:10.1029/2005JD006689, 2006.
- 1020 Parrish, D. D., Allen, D. T., Bates, T. S., Estes, M., Fehsenfeld, F. C., Feingold, G., Ferrare, R., Hardesty, R. M.,
- 1021 Meagher, J. F., Nielsen-Gammon, J. W., Pierce, R. B., Ryerson, T. B., Seinfeld, J. H. and Williams, E. J.:
- 1022 Overview of the Second Texas Air Quality Study (TexAQS II) and the Gulf of Mexico Atmospheric
- 1023 Composition and Climate study (GoMACCS), J. Geophys. Res., 114 (D00F13), doi: 10.1029/2009JD011842,
- 1024 2009.
- 1025 Phillips, G. J., Makkonen, U., Schuster, G., Sobanski, N., Hakola, H. and Crowley, J. N.: The detection of
- 1026 nocturnal N<sub>2</sub>O<sub>5</sub> as HNO<sub>3</sub> by alkali- and aqueous-denuder techniques, Atmos. Meas. Tech., 6, 231-237,
- 1027 doi:10.51094/amt-6-231-2013, 2013.
- 1028 Pope, R. J., Chipperfield, M. P., Savage, N. H., Ordóñez, C., Neal, L.S., Lee, L.A., Dhomsel, S. S., Richards, N.
- 1029 A. D. and Keslake, T. D.: Evaluation of a regional air quality model using satellite column NO2: treatment of
- 1030 observation errors and model boundary conditions and emissions, Atmos. Chem. Phys., 15, 5611-5626,
- 1031 doi:10.5194/acp-15-5611-2015, 2015
- 1032 Russell, A. R., Perring, A. E., Valin, L. C., Bucsela, E. J., Browne, E. C., Wooldridge, P. J. and Cohen, R. C.: A
- 1033 high spatial resolution retrieval of NO<sub>2</sub> column densities from OMI: method and evaluation, Atmos. Chem.
- 1034 Phys., 11, 8543-8554, doi:10.5194/acp-11-8543-2011, 2011.
- 1035 Schery, S. D.: Progress on Global Rn<sup>222</sup> Flux Maps and Recommendations for Future Research, in: 1st
- 1036 International Expert Meeting on Sources and Measurements of Natural Radionuclides Applied to Climate and
- 1037 Air Quality Studies (Gif-sur-Yvette, France, June 2003), edited by: Barrie, L. A. and Lee, H. N., WMO TD
- 1038 1201, Gif-sur-Yvette, France, 43–47, 2004.
- 1039 Schwartz, S. E.: Mass-transport considerations pertinent to aqueous-phase reactions of gases in liquid-water
- 1040 clouds, in: Chemistry of Multiphase Atmospheric Systems, edited by: Jaechske, W., Springer, Heidelberg, 415-
- 1041 471, 1986.
- 1042 Seltzer, K. M., Vizuete, W. and Henderson, B. H.: Evaluation of updated nitric acid chemistry on ozone
- precursors and radiative effects, Atmos. Chem. Phys. Discuss., 15, 3219–3255, 2015.
- Shettle, E. P. and Fenn, R. W.: Models for the aerosols of the lower atmosphere and the effects of the humidity
- variations on their optical properties, Environ. Res. Paper, 676, AFGL-TR-79-0114, 91 pp, 1979.

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





- 1046 Sindelarova, K., Granier, C., Bouarar, I., Guenther, A., Tilmes, S., Stavrakou, T., Muller, J.-F., Kuhn, U.,
- 1047 Stefani, P. and Knorr, W.: Global dataset of biogenic VOC emissions calculated by the MEGAN model over the
- 1048 last 30 years, Atmos. Phys. Chem., 14, 9317–9341, doi:10.5194/acp-14-9317-2014, 2014.
- 1049 Singh, H. B., Brune, W. H., Crawford, J. H., et al.: Chemistry and transport of pollution over the Gulf of Mexico
- and the Pacific: spring 2006 INTEX-B campaign overview and first results, Atmos. Chem. Phys., 9, 2301-2318,
- 1051 doi:10.5194/acp-9-2301-2009, 2009.
- 1052 Stavrakou, T., Müller, J.-F., De Smedt, I., Van Roozendael, M., van der Werf, G. R., Giglio, L., and Guenther,
- 1053 A.: Global emissions of non-methane hydrocarbons deduced from SCIAMACHY formaldehyde columns
- 1054 through 2003-2006, Atms. Chem. Phys., 9, 3663-3679, 2009.
- 1055 Steinbacher, M., Zellweger, C., Schwarzenbach, B., Bugmann, S., Buchmann, B., Ordónez, C., Prevot, A. S. H.,
- 1056 and Hueglin, C.: Nitrogen oxides measurements at rural sites in Switzerland: bias of conventional measurement
- 1057 techniques, J. Geophys. Res., 112, D11307, doi:10.1029/2006JD007971, 2007.
- 1058 Stevenson, D. S., Dentener, F. J., Schultz, M. G., Ellingsen, K., van Noije, T. P. C., Wild, O., Zeng, G., Amann,
- 1059 M., Atherton, M., Bell, N., Bergmann, D. J., Bey, I., Bulter, T., Cofala, J., Collins, W. J., Derwent, R. G.,
- 1060 Doherty, R. M., Drevet, J., Eskes, H. J., Fiore, A. M., Gauss, M., Hauglustaine, D. A., Horowitz, L. W., Isaksen,
- 1061 I. S. A., Krol, M. C., Lamarque, J.-F., Lawrence, M. G., Montanaro, V., M'uller, J. F., Pitari, G., Prather, M. J.,
- 1062 Pyle, J. A., Rast, S., Rodriguez, J. M., Sanderson, M. G., Savage, N. H., Shindell, D. T., Strahan, S. E., Sudo, K.,
- and Szopa, S.: Multimodel ensemble simulations of present-day and near future tropospheric ozone, J. Geophys.
- 1064 Res., 111, D08301, doi: 10.1029/2005JD006338, 2006.
- 1065 Sutton, R. T., Dong, B. and Gregory, J. M.: Land/sea warming ratio in response to climate change: IPCC AR4
- model results and comparison with observations, Geophys. Res. Letts., 34(2), 10.1029/2006GL028164, 2007.
- 1067 Tang, Q., Prather, M. J. and Hsu, J.: Stratosphere-troposphere exchange ozone flux related to deep convection,
- 1068 Geophys. Res. Letts., 38, L03806, doi:10.1029/2010GL046039, 2011.
- 1069 Thouret, V., Marenco, A., Logan, J. A., Nédélec, P., and Grouhel, C.: Comparisons of ozone measurements from
- 1070 the MOZAIC airborne program and the ozone sounding network at eight locations, J. Geophys. Res., 103,
- 1071 25695–25720, 1998.
- 1072 Tiedtke, M.: A comprehensive mass flux scheme for cumulus parameterization in large-scale models, Mon.
- 1073 Weather. Rev., 117(8), 1779–1800, 1989.
- 1074 Tørseth, K., Aas, W., Breivik, K., Fjaeraa, A. M., Fiebig, M., Hjellbrekke, A. G., Lund-Myrhe, C., Solberg, S.
- 1075 and Yttri, K. E.: Introduction to the European Monitoring and Evaluation Programme (EMEP) and observed
- atmospheric composition change during 1972-2009, Atmospheric Chemistry and Physics, 12, 5447-5481, 2012.
- 1077 Urban, J., Pommier, M., Murtagh, D. P., Santee, M. L., and Orsolini, Y. J.: Nitric acid in the stratosphere based
- 1078 on Odin observations from 2001 to 2009 Part 1: A global climatology, Atmos. Chem. Phys., 9, 7031-7044,
- 1079 doi:10.5194/acp-9-7031-2009, 2009.

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





- 1080 Valks, P., Pinardi, G., Richter, A., Lambert, J.-C., Hao, N., Loyola, D., Van Roozendael, M., and Emmadi, S.:
- 1081 Operational total and tropospheric NO2 column retrieval for GOME-2, Atmos. Meas. Tech., 4, 1491-1514,
- 1082 doi:10.5194/amt-4-1491-2011, 2011.
- 1083 van Geffen, J.H.G.M., Boersma, K. F., Eskes, H. J., Maasakkers J. D. and Veefkind, J. P., TROPOMI
- Algorithm Theoretical Basis Document (ATBD) tropospheric and total NO<sub>2</sub>, S5P-KNMI-L2-0005-RP, 56pp,
- 1085 2016.
- 1086 van der A, R. J., Allaart, M. A. F., and Eskes, H. J.: Multi sensor reanalysis of total ozone, Atmos. Chem. Phys.
- 1087 Discuss., 10, 11401-11448, doi: 10.5194/acpd-10-11401-2010, 2010.
- 1088 van der Werf, G. R., Randerson, J. T., Giglio, L., Collatz, G. J., Mu, M., Kasibhatla, P. S., Morton, D. C.,
- 1089 DeFries, R. S., Jin, Y., and van Leeuwen, T. T.: Global fire emissions and the contribution of deforestation,
- 1090 savanna, forest, agriculture, and peat fires (1997-2009) Atmos. Chem. Phys., 10, 11707-11735, 2010,
- 1091 <u>http://www.atmos-chem-phys.net/10/11707/2010/.</u>
- 1092 Veefkind, J.P., Aben, I., McMullan, K., Förster, H., de Vries, J., Otter, G., Claas, J., Eskes, H. J., de Haan, J. F.,
- 1093 Kleipool, Q., van Weele, M., Hasekamp, O., Hoogeveen, R., Landgraf, J., Snel, R., Tol, P., Ingmann, P., Voors,
- 1094 R., Kruizinga, B., Vink, R., Visser, H. and Levelt, P. F.: TROPOMI on the ESA Sentinel-5 Precursor: A GMES
- 1095 mission for global observations of the atmospheric composition for climate, air quality and ozone layer
- 1096 applications, Remote. Sens. Environ., 120, 70-83, doi:10.1016/j.rse.2011.09.027, 2012.
- 1097 Verstraeten, W. V., Neu, J. L., Williams, J. E., Bowman, K. W. and Worden, J. R.: Rapid increases in
- 1098 tropospheric ozone production and export from China, Nature Geosci., 8, 690-695, doi: 10.1038/ngeo2493,
- 1099 2015.
- 1100 Vinken, G. C. M., Boersma, K. F., Jacob, D.J. and Meijer, E. W.: Accounting for non-linear chemistry of ship
- 1101 plumes in the GEOS-Chem global chemistry transport model, Atms. Chem. Phys., 11, Atmos. Chem. Phys.,
- 1102 11707-11722, doi:10.5194/acp-11-11707-2011, 2011.
- 1103 Vinken, G. C. M., Boersma, K. F., Maasakkers, J. D., Adon, M. and Martin, R. V.: Worldwide biogenic soil
- NO<sub>x</sub> emissions inferred from OMI NO<sub>2</sub> observations, Atmos. Chem. Phys., 14, doi:10.5194/acp-14-10363-2014,
- 1105 10363–10381, 2014.
- 1106 Von Kuhlmann, R. and Lawrence, M. G.: The impact of ice uptake of nitric acid on atmospheric chemistry,
- 1107 Atms. Chem. Phys., 6, 225-235, 2006.
- 1108 Wild, O. and Prather, M. J.: Global tropospheric ozone modeling: Quantifying errors due to grid resolution, J.
- 1109 Geophys. Res., 111, D11305, doi:10.1029/2005JD006605, 2006.
- 1110 Williams, J. E., Strunk, A., Huijnen, V. and van Weele, M.: The application of the Modified Band Approach for
- 1111 the calculation of on-line photodissociation rate constants in TM5: implications for oxidative capacity, Geosci.
- 1112 Model Dev., 5, 15-35, doi:10.5194/gmd-5-15-2012,2012.
- 1113 Williams, J. E., van Velthoven, P. F. J., and Brenninkmeijer, C. A. M.: Quantifying the uncertainty in simulating
- 1114 global tropospheric composition due to the variability in global emission estimates of Biogenic Volatile Organic
- 1115 Compounds, Atmos. Chem. Phys., 13, 2857–2891, doi: 10.5194/acp-13-2857-2013, 2013.

Manuscript under review for journal Geosci. Model Dev.

Published: 11 July 2016

© Author(s) 2016. CC-BY 3.0 License.





- Williams, J. E., Le Bras, G., Kukai, A., Ziereis, H., and Brenninkmeijer, C. A. M.: The impact of the chemical
- 1117 production of methyl nitrate from the NO + CH<sub>3</sub>O<sub>2</sub> reaction on the global distribution of alkyl nitrates, nitrogen
- oxides and tropospheric ozone: a global modeling study, Atmos. Chem. Phys., 14, 2363-2382, 2014.
- 1119 Worden, J., Liu, X., Bowman, K., Chance, K., Beer, R., Eldering, A., Gunson, M. and Worden, H., Improved
- 1120 tropospheric ozone profile retrievals using OMI and TES radiances, Geophys. Res. Lett., 34, L01809,
- 1121 doi:10.1029/2006GL027806, 2007.
- 1122 Yamaji, K., Ikeda, K., Irie, H., Kurokawa, J. and Ohara, T.: Influence of model grid resolution on NO<sub>2</sub> vertical
- 1123 column densities over East Asia, J. Air. Waste Manage. Assoc., 64(4), 436-444, 2014.
- Yarwood, G., Rao, S., Yocke, M., and Whitten, G.: Updates to the carbon bond chemical mechanism: CB05,
- Final report to the US EPA, EPA Report Number: RT-0400675, available at: www.camx.com (last access: 15
- 1126 January 2014), 2005.
- Zeng. G., Williams, J. E., Fisher, J.A., Emmons, L. K., Jones, N. B., Morgenstern, O., Robinson, J., Smale, D.,
- 1128 Paton-Walsh, C. and Griffith, D. W. T.: Multi-model simulation of CO and HCHO in the Southern Hemisphere:
- 1129 comparison with observations and impact of biogenic emissions, Atmos. Chem. Phys., 15, 7217-7245,
- 1130 doi:10.5194/acp-15-7217-2015, 2015.
- Zdunkowski, W. G., Welsch, R. M., and Kord, G. J.: An investigation of the structure of typical 2-stream
- methods for the calculation of solar fluxes and heating rates in clouds, Contrib. Atms. Phys., 53, 215-238, 1980.
- 1133 Zhou, Y., Brunner, D., Hueglin, C., Henne, S., and Staehelin, J.: Changes in OMI tropospheric NO2 columns
- over Europe from 2004 to 2009 and the influence of meteorological variability. Atmos Environ, 46, 482-495,
- 1135 2012.
- 2136 Zyrichidou, I., Koukouli, M. E., Balis, D., Markakis, K., Poupkou, A., Katragkou, E., Kioutsioukis, I., Melas,
- 1137 D., Boersma, K. F. and van Roozendael, M.: Identification of surface NOx emission sources on a regional scale
- 1138 using OMI NO<sub>2</sub>. Atms. Environ., 101, 82-93, 2015.