The high-resolution version of TM5-MP for optimized satellite retrievals: description and validation

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Received: 18 May 2016 – Discussion started: 11 July 2016
Revised: 13 December 2016 – Accepted: 4 January 2017 – Published: 15 February 2017

Abstract. We provide a comprehensive description of the high-resolution version of the TM5-MP global chemistry transport model, which is to be employed for deriving highly resolved vertical profiles of nitrogen dioxide (NO₂), formaldehyde (CH₂O), and sulfur dioxide (SO₂) for use in satellite retrievals from platforms such as the Ozone Monitoring Instrument (OMI) and the Sentinel-5 Precursor, and the TROPOspheric Monitoring Instrument (tropOMI). Comparing simulations conducted at horizontal resolutions of 3° × 2° and 1° × 1° reveals differences of ±20 % exist in the global seasonal distribution of ²²²Rn, being larger near specific coastal locations and tropical oceans. For tropospheric ozone (O₃), analysis of the chemical budget terms shows that the impact on globally integrated photolysis rates is rather low, in spite of the higher spatial variability of meteorological data fields from ERA-Interim at 1° × 1°. Surface concentrations of O₃ in high-NOₓ regions decrease between 5 and 10 % at 1° × 1° due to a reduction in NOₓ recycling terms and an increase in the associated titration term of O₃ by NO. At 1° × 1°, the net global stratosphere–troposphere exchange of O₃ decreases by ∼ 7 %, with an associated shift in the hemispheric gradient. By comparing NO, NO₂, HNO₃ and peroxy-acetyl-nitrate (PAN) profiles against measurement composites, we show that TM5-MP captures the vertical distribution of NOₓ and long-lived NOₓ reservoirs at background locations, again with modest changes at 1° × 1°. Comparing monthly mean distributions in lightning NOₓ and applying ERA-Interim convective mass fluxes, we show that the vertical re-distribution of lightning NOₓ changes with enhanced release of NOₓ in the upper troposphere. We show that surface mixing ratios in both NO and NO₂ are generally underestimated in both low- and high-NOₓ scenarios. For Europe, a negative bias exists for [NO] at the surface across the whole domain, with lower biases at 1° × 1° at only ∼ 20 % of sites. For NO₂, biases are more variable, with lower (higher) biases at 1° × 1° occurring at ∼ 35 % (∼ 20 %) of sites, with the remainder showing little change. For CH₂O, the impact of higher resolution on the chemical budget terms is rather modest, with changes of less than 5 %. The simulated vertical distribution of CH₂O agrees reasonably well with measurements in pristine locations, although column-integrated values are generally underestimated relative to satellite measurements in polluted regions. For SO₂, the performance at 1° × 1° is principally governed by the quality of the emission inventory, with limited 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. For remote regions missing biogenic source terms are inferred.

1 Introduction

One application of chemistry transport models (CTMs) is to provide accurate vertical and horizontal global distributions of trace gases such as ozone (O₃), nitrogen dioxide (NO₂), sulfur dioxide (SO₂) and formaldehyde (CH₂O) that are used as a priori best guesses in the retrievals of tropospheric abundances from instruments mounted on Earth-orbiting satellites such as the Tropospheric Emission Sounder (TES; Worden et al., 2007) Global Ozone Monitoring Experiment (GOME),
the SCanning Imaging Absorption spectroMeter for Atmospheric CHartographY (SCIAMACHY; De Smedt et al., 2008), the Ozone Monitoring Instrument (OMI; Boersma et al., 2011), and GOME-2 (Valks et al., 2011). To date, although high-resolution regional models have been employed for selected regions such as the US and Europe (e.g. Russell et al., 2011; Zhou et al., 2012; Vinken et al., 2014), at the global scale the CTM resolutions employed are still rather coarse (between 1.1 and 4.0° in latitude and between 1.1 and 6° in longitude), resulting in “footprints” which aggregate hundreds of kilometres in area. This has limitations as the resulting total columns are sensitive to topography, surface albedo and the shape of the a priori vertical profiles themselves. Using rather coarse resolution leads to substantial errors in the retrievals (e.g. Boersma et al., 2008; Heckel et al., 2011; Russell et al., 2011) and imposes limitations on capturing the regional-scale variability in short-lived trace gas abundances observed from high-resolution satellite instruments such as the OMI.

This lack of spatial detail is particularly relevant for situations where strong spatio-temporal variability in the vertical distribution of NO\textsubscript{2}, SO\textsubscript{2}, and CH\textsubscript{2}O can be expected. Examples include shipping lanes in the relatively unpolluted marine boundary layer (e.g. Vinken et al., 2014) and coal-fired power plant SO\textsubscript{2} pollution (e.g. Fioletov et al., 2015). Moreover, during the day the local lifetime and mixing ratios of trace gases such as nitric oxide (NO) and NO\textsubscript{2} are critically dependent on a host of variables, e.g. temperature, surface albedo, cloud cover (via photolysis), chemical conversion (i.e. the NO / NO\textsubscript{2} ratio) and the extent of mixing by convective upwelling (i.e. land type) and advective transport.

Thus, the information provided for the retrievals is affected by the coarsening of the high-resolution meteorological data used to drive the CTM. Recently, Heckel et al. (2011) demonstrated that there is an associated uncertainty of ∼ 2 using a priori data from a global CTM rather than a regional CTM, principally due to loss of spatial information. Two other studies focusing on the impact of horizontal resolution on the retrieval of vertical column densities of NO\textsubscript{2} suggested that errors of up to ∼ 50% exist (Yamaji et al., 2014; Lin et al., 2014). This problem becomes accentuated for the next generation of Earth-orbiting satellites such as the Tropospheric Monitoring Instrument (tropOMI), which has a smaller footprint compared to its predecessors (Veenkamp et al., 2012). Applications of TM5 include the retrieval of NO\textsubscript{2}, CH\textsubscript{2}O, and SO\textsubscript{2} column densities from the OMI and tropOMI (e.g. van Geffen et al., 2016), where studies related to the influence of horizontal resolution have been limited principally to NO\textsubscript{2}.

The dominant tropospheric loss terms for CH\textsubscript{2}O are photolysis and scavenging into cloud droplets (wet deposition; Jacob, 2000). Thus the atmospheric lifetime of CH\textsubscript{2}O is highly sensitive to the extent of cloud cover and the vertical profiles of the photolysis rates. A dominant application of CH\textsubscript{2}O retrievals is to provide constraints on tropical and sub-
tropical isoprene emission fluxes (e.g. Palmer et al., 2006; Stavrovkou et al., 2009; Marais et al., 2012). The resulting emission estimates are highly sensitive to the stoichiometric yield of CH\textsubscript{2}O from isoprene oxidation, the chemical lifetime of CH\textsubscript{2}O and spatial differences in land cover. Other applications include estimating emissions released during biomass burning (BB) episodes (Gonzi et al., 2011), whose spatial location is also smeared via coarsening in TM5-MP. For SO\textsubscript{2}, which predominantly originates from point sources, an adequate spatial distribution of such sources is crucial for estimating accurate biases in existing emission inventories.

In this paper we provide a comprehensive description of the global, high-resolution 1° × 1° 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, the stratospheric boundary conditions, the photolysis scheme, the heterogeneous conversion and the overall model structure. In Sect. 3 we analyse the impact on convective and advective transport of trace species from the boundary layer (BL) of both increased horizontal resolution and use of ERA-Interim convective mass fluxes, as derived using radon (222\textsuperscript{Rn}) distributions. In Sect. 4 we investigate the effects on regional and global photolysis frequencies. In Sects. 5–9 we examine the differences in the vertical and horizontal distributions of tropospheric O\textsubscript{3}, NO\textsubscript{2}, lightning-induced NO\textsubscript{x}, N-containing species (i.e. nitric acid (HNO\textsubscript{3}), PAN and lumped organic nitrates (ORGNTR)), CH\textsubscript{2}O and SO\textsubscript{2}, where we make comparisons against both surface and aircraft measurements to validate mixing ratios. Finally, in Sect. 10, 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 pre-defined regional domain, with boundary conditions being determined by the global simulation at coarser resolution. Typically, global simulations at 3° × 2° with zoom regions at 1° × 1° were performed to alleviate the long runtime of a global 1° × 1° run. In the new version of TM5 (hereafter referred to as TM5-MP; the massively parallel version), the usage of the Message Passing Interface (MPI) has been totally rewritten. Zoom regions are no longer available, but datasets are distributed along longitudes and latitudes, instead of model levels and tracers. The advantages of that overhaul of domain decomposition are a smaller memory requirement and the possibility of using more processors making global 1° × 1° simulations feasible in terms of runtime and affordable in terms of computing resources. A TM5-MP global 3° × 2°
(1° × 1°) run is ∼6 (∼20) times faster than the previous version of TM5 (Huijnen et al., 2010) for similar resources. The following model description pertains to both 3° × 2° and 1° × 1° simulations discussed in this paper.

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 re-analysis (Dee et al., 2011) and updated every 3 h, with interpolation of fields for the intermediate time periods. Although TM5-MP can adopt all 60 vertical levels provided by the ECMWF ERA-Interim reanalysis, we employ 34 vertical levels for this study, with higher resolution in the troposphere and upper troposphere–lower stratosphere (UTLS). Convective mass fluxes and detrainment rates are taken from the ERA-Interim dataset to describe the updraft velocities from the BL into the free troposphere, which replaces the parameterization of Tiedtke (1989) used in previous versions. The vertical diffusion in the free troposphere is calculated according to Louis (1979), and in the BL by the approach of Holtslag and Boville (1993). Diurnal variability in the BL height is determined using the parameterization of Vogelezang and Holtslag (1996). We use the first-order moments scheme with an iterative time step to prevent too much mass being transported out of any particular grid cell according to the preservation of the Courant–Friedrichs–Lewy (CFL) criterion (Bregman et al., 2003), which is especially relevant when reducing the size of grid cells as done here.

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 on updating and expanding the fast NOx chemistry to account for an accurate partitioning of nitrogen for higher NOx regimes than those occurring at coarser horizontal resolutions. All reaction rate data are 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 NOx and CH2O chemistry are given in Table 1. This includes the recent update to the formation rate of HNO3 determined by Möllner et al. (2010). The most relevant modifications are that (i) the yield of CH2O, methanol (CH3OH) and the hydro-peroxy radical (HO2) from the self-termination of the methyl-peroxy radical (CH3O2) is increased according to Yarwood et al. (2005), (ii) the direct formation of CH2O from the reaction of CH3O2+ HO2 is added using the temperature-dependent branching ratio defined in Atkinson et al. (2004), (iii) the production of HNO3 during the oxidation of dimethyl sulfide (DMS) by the NO3 radical is now included, (iv) explicit organic peroxy radicals have been introduced as products from the oxidation of propane (C3H8) and propane (C3H6) by OH, which are lost either by the reaction with NO or HO2 allowing the in situ chemical formation of acetone (CH3COCH3) and higher aldehydes (ALD2), respectively, following the stoichiometry given in Emmons et al. (2010), (v) a second product channel for N2O5 photolysis is added producing NO, (vi) the formation and photo-dissociation of HONO have been included, (vii) the formation and transport of methyl-peroxy-nitrate (CH3O2NO2) are also included (Browne et al., 2011), and (viii) modifications to the gas-phase chemistry involving NH3 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 have been fully described in Williams et al. (2012). For the calculation of the height-resolved actinic fluxes at the seven specific wavelength used for calculating the J values (these being 205.1, 287.9, 302.0, 311.0, 326.5, 385.0 and 610.0 nm), the two-stream radiative transfer solver of Zdunkowski et al. (1980) is embedded in TM5-MP. For 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 (reff) 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 reff value using the land fraction in each grid cell. We apply limits between 4 and 16 µm to 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 (LT; not shown). For the scattering effects from cloud droplets, we subsequently downsize the physical reff by ∼0.5–2 µm to account for the relationship between the optical and physical reff values.

For aerosols, an aerosol scheme is available for use within TM5-MP (Aan den Brugh et al., 2011), 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 to satellite retrieval, it is preferable to use any advancements in computational performance on further increases in the horizontal resolution employed. Therefore it is not currently envisaged that a full description of aerosol processes will be included during operational satellite retrievals.

However, heterogeneous conversion processes still need the description of the total reactive surface area density (SAD) from aerosols. In TM5-MP this is assumed as the cumulative value of contributions from sulfate, nitrate, ammo-
the parameterization of Evans and Jacob (2005), and is there-
some account for. Temperature-dependent gas-phase diffu-
sion of Schwartz (1986).

<table>
<thead>
<tr>
<th>Reactants</th>
<th>Products</th>
<th>Rate parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>NO + O₃</td>
<td>NO₂</td>
<td>$3.0 \times 10^{-12} \times \exp(-1500/T)$</td>
</tr>
<tr>
<td>NO₂ + O₃</td>
<td>NO₃</td>
<td>$1.4 \times 10^{-13} \times \exp(-2470/T)$</td>
</tr>
<tr>
<td>NO + HO₂</td>
<td>NO₂ + OH</td>
<td>$3.3 \times 10^{-12} \times \exp(270/T)$</td>
</tr>
<tr>
<td>NO + CH₃O₂</td>
<td>CH₂O + HO₂ + NO₂</td>
<td>$2.8 \times 10^{-12} \times \exp(300/T)$</td>
</tr>
<tr>
<td>OH + NO₂</td>
<td>HNO₃</td>
<td>$k_0 = 3.2 \times 10^{-30} \times (300/T)^{4.5}$ $k_\infty = 3.0 \times 10^{-11}$</td>
</tr>
<tr>
<td>NO + NO₃</td>
<td>NO₂ + NO₂</td>
<td>$1.8 \times 10^{-11} \times \exp(110/T)$</td>
</tr>
<tr>
<td>NO₂ + NO₃</td>
<td>N₂O₅</td>
<td>$k_0 = 8.0 \times 10^{-27} \times (300/T)^{3.5}$ $k_\infty = 3.0 \times 10^{-11} \times (300/T)^{1.0}$</td>
</tr>
<tr>
<td>N₂O₅ + M</td>
<td>NO₂ + NO₃</td>
<td>$k_0 = 1.3 \times 10^{-3} \times (300/T)^{3.5} \times \exp(-11000/T)$ $k_\infty = 9.7 \times 10^{14} \times (300/T)^{-0.1} \times \exp(-11080/T)$</td>
</tr>
<tr>
<td>HO₂ + NO₂</td>
<td>HNO₄</td>
<td>$k_0 = 1.4 \times 10^{-31} \times (300/T)^{3.1}$ $k_\infty = 4.0 \times 10^{-12}$</td>
</tr>
<tr>
<td>HNO₄ + M</td>
<td>HO₂ + NO₂</td>
<td>$k_0 = 4.1 \times 10^{-5} \times \exp(-10650/T)$ $k_\infty = 6.0 \times 10^{15} \times \exp(-11170/T)$</td>
</tr>
<tr>
<td>OH + NO₂O₄</td>
<td>NO₂</td>
<td>$1.3 \times 10^{-12} \times \exp(380/T)$</td>
</tr>
<tr>
<td>OH + NO + M</td>
<td>HONO</td>
<td>$k_0 = 7.0 \times 10^{-31} \times (300/T)^{4.4}$ $k_\infty = 3.6 \times 10^{-11} \times (300/T)^{0.1}$</td>
</tr>
<tr>
<td>HONO + hν</td>
<td>OH + NO</td>
<td>$2.5 \times 10^{-12} \times \exp(260/T)$</td>
</tr>
<tr>
<td>OH + HONO</td>
<td>NO₂</td>
<td>$k_0 = 3.28 \times 10^{-28} \times (300/T)^{6.87}$ $k_\infty = 1.125 \times 10^{-11} \times (300/T)^{1.105}$</td>
</tr>
<tr>
<td>NO₂ + CH₃C(O)O₂</td>
<td>PAN</td>
<td>$k_0 = 1.1 \times 10^{-5} \times \exp(-10100/T)$ $k_\infty = 1.9 \times 10^{17} \times \exp(-14100/T)$</td>
</tr>
<tr>
<td>PAN + hν</td>
<td>CH₃C(O)O₂ + NO₂CH₃O₂ + NO₃</td>
<td>$k_0 = 2.5 \times 10^{-30} \times (300/T)$ $k_\infty = 1.8 \times 10^{-11}$</td>
</tr>
<tr>
<td>CH₃O₂ + NO₂</td>
<td>CH₃O₂NO₂</td>
<td>$k_0 = 9.0 \times 10^{-5} \times \exp(-9690/T)$ $k_\infty = 1.1 \times 10^{16} \times \exp(-10560/T)$ $4.0 \times 10^{-12}$</td>
</tr>
<tr>
<td>NO₃ + HO₂</td>
<td>HNO₃</td>
<td></td>
</tr>
</tbody>
</table>

Table 1. Details of the reaction rate data applied for NO₄ 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 stoichiometry are taken from Atkinson et al. (2004) accommodating the latest evaluation at http://iapac.pole-ether.fr.

Once a surface reaction with H₂O occurs, 2 molecules of HNO₃ are formed. No uptake on cirrus particles is included for HNO₃, which can lead to de-nitrification of the upper troposphere (Lawrence and Crutzen, 1998; von Kuhlmann and Lawrence, 2006). For HO₂ we adopt a fixed $γ_{HO_2} = 0.06$ across all aerosol types as taken from Abbatt et al. (2012), and for NO₃ we adopt a fixed $γ_{NO_3} = 10^{-3}$ as recommended by Jacob (2000). For HO₂, heterogeneous conversion forms 0.5 molecules of hydrogen peroxide (H₂O₂), whereas for NO₃ it forms 1 molecule of HNO₃ following Emmons et al. (2010). For the SAD associated with cloud droplets we use the $r_{eff}$ values that are calculated by Martin et al. (1994), thus maintaining consistency between the size of the cloud droplets used for the scattering component in the calculation of $J$ values and heterogeneous loss rates on the clouds. By using the ECMWF cloud fraction for each respective grid cell, we assume that instantaneous mixing throughout the...
grid cell does not occur, in order to avoid exaggerated con-
version rates on cloud surfaces.

As TM5-MP contains no explicit stratospheric chemistry,
we apply constraints above the tropopause to ensure realistic stratosphere–troposphere exchange (STE) of O₃ and
for constraining the incoming radiation reaching the troposphere
needed for the MBA (Williams et al., 2012). For strato-
spheric O₃, we use total column values derived from the
assimilation of satellite observations as provided in the
improved version of the Multi-Sensor Re-analysis (MSR, van
der A et al., 2010), which is vertically distributed according
to the climatology of Fortuin and Kelder (1998). Three dis-
tinct zonal bands are used for nudging the stratospheric O₃
fields, these being 30° S−30° N, 30−66° S/N and > 66° S/N,
where nudging occurs at pressure levels < 45 hPa, < 95 hPa
and < 120 hPa, with relaxation times of 2.5, 3 and 4 days,
respectively.

For stratospheric CH₄ we use the monthly 2-D climato-
logical fields provided by Grooß and Russell (2005), with
the nudging heights and relaxation times being identical to
those used for stratospheric O₃. For stratospheric CO and
HNO₃ we constrain mixing ratios by using monthly mean
ratios of CO / O₃ (Dupuy et al., 2004) and HNO₃ / O₃ (Jé-
gou et al., 2008; Urban et al., 2009) based on the latitu-
dinal climatologies derived from ODIN observations using
data for 2003/2004 (CO) and 2001–2009 (HNO₃). 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. These ratios are
applied using the monthly mean stratospheric O₃ distribu-
tion in TM5-MP, which is constrained by the MSR dataset
(van der A et al., 2010). For both species, model fields are
nudged at 5.5, 10 and 28 hPa using relaxation times of 5,
10 and 60 days, respectively. Previous versions of TM5 used
a HNO₃ climatology from the UARS MLS instrument and
applied nudging constraints at 10 hPa only (Huijnen et al.,
2010).

For our study on the impact of horizontal resolution on
the performance of TM5-MP, we present simulations for the
year 2006, which has been used for previous benchmark-
ing studies (Huijnen et al., 2010; Williams et al., 2012). We
use a 1-year spin-up from the same initial conditions, where
the initial conditions are representative of the state-of-the-
atmosphere for January 2005 taken from a previous simula-
tion (see Zeng et al., 2015). The model is run using 34 lev-
els, as it will be used operationally for satellite retrievals,
with details of the pressure levels being given in Huijnen et al.
(2010).

2.1 Emission inventories

All emission inventories applied in TM5-MP are year-
specific, meaning that the year-to-year variability in emis-
sion fluxes due to changes in anthropogenic activity, bio-
genic activity and burning extent is taken into account. For
the anthropogenic emission of NOₓ, CO, SO₂, NH₃ and non-
methane volatile organic compounds (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, where a bi-sinusoidal distribution could be ap-
plied, peaking in the morning and late afternoon to repre-
sent variability in traffic volume. Aircraft emissions are in-
cluded only for NOₓ using a homogenous hourly flux esti-
mate related to regional flight times. For volcanic SO₂
emissions, the estimated emission flux has been scaled up to
10 Tg S yr⁻¹ 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 ∼ 50 m between 20° S and 20° N,
whereas for other latitudes a continuous daily flux is applied.
The BB emissions are taken from the monthly estimates pro-
vided by the GFEDv3 inventory (van der Werf et al., 2010)
and latitude-dependent injection heights and a tropical burn-
ing cycle are implemented following Huijnen et al. (2010).
All emission inventories are provided on a 0.5° × 0.5° reso-
lution and subsequently coarsened onto the horizontal res-
olution employed in any simulation. In TM5-MP all NOₓ
emissions are introduced as NOₓ rather than specifying a
fraction that is emitted directly as NO (Carslaw and Beever,
2005). Global NOₓ emissions for the year 2006 total
49 Tg N yr⁻¹ (including lightning). Other notable species in-
clude CO (1081 Tg CO yr⁻¹), SO₂ (117 Tg S yr⁻¹), CH₃O
(13.5 Tg C yr⁻¹) and isoprene (510 Tg C yr⁻¹). An overview of
the global and zonal emission terms used in the simula-
tions analysed here is given in Table 3.

For lightning NOₓ we use the parameterization which uses
convective precipitation fields (Meijer et al., 2001) and con-
strain the annual global emission term at ∼ 6 Tg N yr⁻¹. This
uses the convective flux values, meaning that re-scaling of
the nudging term was necessary in order to achieve simi-
lar total lightning NOₓ emissions across simulations. An ex-
ample of the resulting horizontal distributions in lightning
NOₓ at ∼ 400 hPa for the tropics for both horizontal reso-
lutions is shown in the top panel of Fig. S1a in the Supple-
ment. Although the spatial variability increases at 1° × 1°,
the global distribution remains essentially the same, where
the constraints on annual lightning NOₓ emissions homoge-
nize the total emission flux between resolutions. One other
factor affecting the vertical distribution of lightning NOₓ
emissions is the convective parameterization which is used.
The lower panels of Fig. S1a show that, at this altitude, the
Tiedke (1989) approach increases the NOₓ emissions at this
level by ∼ 14 %, accompanied by a significant re-distribution
between regions (cf. the Southern Hemisphere (SH) below.
30° S and an significant increase in the tropical component in the Tiedke simulation). However, this is altitude-dependent, where the absolute differences in the vertical distribution in the monthly NOx emissions for a selection of latitudes are shown in Fig. S1b. Here comparisons are shown both over the continents (e.g. 50.5° N) and the oceans (e.g. 59.5° S). Although the differences in the integrated monthly global emission NOx flux are only around ~1–2 %, the temporal and vertical distribution can be quite different between convective schemes. Profiles show that in the upper troposphere ERA-Interim consistently results in higher NOx emissions around 300 hPa, especially for July.

Latitudinal constraints on CH4 global distributions are applied using the methodology given in Bänd et al. (2015) with a 3-day relaxation time. We also introduce similar constraints based on the appropriate surface measurements for H2 in order to account for the latitudinal gradient and variability across seasons, which replaces the fixed global value of 550 ppb used in previous versions. Finally, for radon (Rn$^{222}$) emissions we apply the estimates of Schery (2004), whose global distribution is given in Zhang et al. (2011).

2.2 Observations

Although the performance of mCB05 in TM5 v3.0 has been validated for selected NMVOC, O3, CH$_2$O, CO and NO$_x$ 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° × 2° and 1° × 1°. We choose a range of ground-based and airborne measurements taken at diverse locations during the year 2006, representing different chemical regimes. Here we briefly describe the observations utilized for this purpose.

For validation of simulated surface concentrations we use measurements of gaseous O$_3$, NO, NO$_2$, HNO$_3$ and SO$_2$ available from the European Monitoring and Evaluation program (EMEP, www.emep.int), where we exploit measurements taken at various background sites in Norway, Finland, the Netherlands, Belgium, Poland, the Czech Republic, Ger-

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<th>Rate expression</th>
<th>Ref.</th>
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<tbody>
<tr>
<td>CH$_3$O$_2$ + HO$_2$</td>
<td>CH$_3$OOH</td>
<td>$3.8 \times 10^{-13} \times \exp(750/T) \times R$</td>
<td>(1), (2)</td>
</tr>
<tr>
<td>CH$_2$O + HO$_2$</td>
<td>CH$_2$O</td>
<td>$3.8 \times 10^{-13} \times \exp(750/T) \times (1 - R)$</td>
<td>(1), (2)</td>
</tr>
<tr>
<td>CH$_2$O + CH$_3$O$_2$</td>
<td>1.37CH$_2$O + 0.74HO$_2$ + 0.63CH$_3$OH</td>
<td>$9.5 \times 10^{-14} \times \exp(390/T)$</td>
<td>(3), (4)</td>
</tr>
<tr>
<td>OH + C$_3$H$_8$</td>
<td>i-C$_3$H$_7$O$_2$</td>
<td>$7.6 \times 10^{-12} \times \exp(-585/T)$</td>
<td>(5), (6)</td>
</tr>
<tr>
<td>NO + IC$_3$H$_2$O$_2$</td>
<td>0.82CH$_3$COCH$_3$ + HO$_2$ + 0.27ALD2 + NO$_2$</td>
<td>$4.2 \times 10^{-12} \times \exp(180/T)$</td>
<td>(6)</td>
</tr>
<tr>
<td>HO$_2$ + IC$_3$H$_7$O$_2$</td>
<td>ROOH</td>
<td>$7.5 \times 10^{-13} \times \exp(700/T)$</td>
<td>(6)</td>
</tr>
<tr>
<td>OH + C$_3$H$_6$</td>
<td>C$_3$H$_4$O$_2$</td>
<td>$k_0 = 8.0 \times 10^{-27} \times (-300/T)^{3.5}$</td>
<td>(5), (6)</td>
</tr>
<tr>
<td>NO$_3$ + C$_4$H$_8$</td>
<td>0.21SPD + XO$_2$ + 0.8HO$_2$ + 0.8ORGNTR + 0.8ALD2 + 2.4 PAR + 0.2 NO$_2$</td>
<td>$2.95 \times 10^{-12} \times \exp(465/T)$</td>
<td>(5)</td>
</tr>
<tr>
<td>NO + C$_3$H$_6$O$_2$</td>
<td>ALD2 + CH$_2$O + HO$_2$ + NO$_2$</td>
<td>$4.2 \times 10^{-12} \times \exp(180/T)$</td>
<td>(6)</td>
</tr>
<tr>
<td>HO$_2$ + C$_3$H$_6$O$_2$</td>
<td>ROOH</td>
<td>$7.5 \times 10^{-13} \times \exp(700/T)$</td>
<td>(6)</td>
</tr>
<tr>
<td>NO$_3$ + DMS</td>
<td>SO$_2$ + HNO$_3$</td>
<td>$1.9 \times 10^{-13} \times \exp(520/T)$</td>
<td>(1)</td>
</tr>
<tr>
<td>NH$_2$ + OH</td>
<td></td>
<td>$3.4 \times 10^{-11}$</td>
<td>(E)</td>
</tr>
<tr>
<td>NH$_2$ + HO$_2$</td>
<td>NH$_3$</td>
<td>$3.4 \times 10^{-11}$</td>
<td>(4), (7)</td>
</tr>
<tr>
<td>NH$_2$ + O$_3$</td>
<td>NH$_2$O$_2$</td>
<td>$4.3 \times 10^{-12} \times \exp(-930/T)$</td>
<td>(4), (7)</td>
</tr>
<tr>
<td>NH$_2$ + O$_2$</td>
<td>NO</td>
<td>$6.0 \times 10^{-21}$</td>
<td>(1), (7)</td>
</tr>
<tr>
<td>NH$_2$O$_2$ + NO</td>
<td>NH$_2$ + NO$_2$</td>
<td>$4.0 \times 10^{-12} \times \exp(450/T)$</td>
<td>(7), (8)</td>
</tr>
<tr>
<td>NH$_2$O$_2$ + O$_3$</td>
<td>NH$_2$</td>
<td>$4.3 \times 10^{-12} \times \exp(-930/T)$</td>
<td>(7), (8)</td>
</tr>
<tr>
<td>NH$_2$O$_2$ + HO$_2$</td>
<td>NH$_2$</td>
<td>$3.4 \times 10^{-11}$</td>
<td>(8)</td>
</tr>
<tr>
<td>CH$_3$O$_2$NO$_2$ + hv</td>
<td>CH$_3$O$_2$ + NO$_2$</td>
<td></td>
<td>(9)</td>
</tr>
<tr>
<td>CH$_2$O$_2$NO$_2$ + hv</td>
<td>CH$_2$O + HO$_2$ + NO$_3$</td>
<td></td>
<td>(9)</td>
</tr>
<tr>
<td>HO$_2$ + aero</td>
<td>0.5H$_2$O$_2$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NO$_3$ + aero</td>
<td>HNO$_3$</td>
<td></td>
<td>(6)</td>
</tr>
</tbody>
</table>
Table 3. The zonally segregated emission totals introduced into TM5-MP for the year 2006. All organic hydrocarbons are given in Tg C yr\(^{-1}\), except for CO (Tg CO yr\(^{-1}\)), CH\(_2\)O (Tg CH\(_2\)O yr\(^{-1}\)) and CH\(_3\)OH (Tg CH\(_3\)OH yr\(^{-1}\)). All NO\(_x\) emissions are introduced as NO (Tg N yr\(^{-1}\)). For SO\(_2\) emission totals are given as Tg SO\(_2\) yr\(^{-1}\) and for NH\(_3\) as Tg NH\(_3\) yr\(^{-1}\). No direct emissions occur for HNO\(_3\), PAN, ORGNTR, HONO, N\(_2\)O\(_5\), NO\(_2\), CH\(_3\)O\(_2\)NO\(_2\) or O\(_3\).

<table>
<thead>
<tr>
<th>Species</th>
<th>Tg yr(^{-1})</th>
<th>Global</th>
<th>30–90° S</th>
<th>30° S–30° N</th>
<th>30–90° N</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO</td>
<td>1081.0</td>
<td>24.4</td>
<td>755.1</td>
<td>301.27</td>
<td></td>
</tr>
<tr>
<td>NO(_x) (as N)</td>
<td>49.0</td>
<td>1.5</td>
<td>24.0</td>
<td>23.6</td>
<td></td>
</tr>
<tr>
<td>SO(_2)</td>
<td>117.0</td>
<td>3.0</td>
<td>49.2</td>
<td>64.3</td>
<td></td>
</tr>
<tr>
<td>DMS (as S)</td>
<td>19.2</td>
<td>6.7</td>
<td>9.3</td>
<td>3.2</td>
<td></td>
</tr>
<tr>
<td>NH(_3)</td>
<td>56.6</td>
<td>3.1</td>
<td>27.9</td>
<td>25.6</td>
<td></td>
</tr>
<tr>
<td>CH(_3)O</td>
<td>13.5</td>
<td>0.3</td>
<td>10.5</td>
<td>2.7</td>
<td></td>
</tr>
<tr>
<td>PAR</td>
<td>34.1</td>
<td>0.7</td>
<td>18.5</td>
<td>14.9</td>
<td></td>
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<tr>
<td>OLE</td>
<td>22.4</td>
<td>0.9</td>
<td>16.6</td>
<td>4.9</td>
<td></td>
</tr>
<tr>
<td>ALD2</td>
<td>13.4</td>
<td>0.4</td>
<td>11.2</td>
<td>1.8</td>
<td></td>
</tr>
<tr>
<td>CH(_3)CHCHO</td>
<td>2.2</td>
<td></td>
<td>1.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CH(_3)OH</td>
<td>100.7</td>
<td>3.3</td>
<td>82.5</td>
<td>14.9</td>
<td></td>
</tr>
<tr>
<td>CH(_3)CH(_2)OH</td>
<td>70.4</td>
<td>2.8</td>
<td>52.6</td>
<td>15.1</td>
<td></td>
</tr>
<tr>
<td>C(_2)H(_4)</td>
<td>25.9</td>
<td>1.0</td>
<td>19.0</td>
<td>5.9</td>
<td></td>
</tr>
<tr>
<td>C(_2)H(_6)</td>
<td>6.1</td>
<td>0.3</td>
<td>5.3</td>
<td>1.5</td>
<td></td>
</tr>
<tr>
<td>C(_3)H(_8)</td>
<td>5.6</td>
<td>0.4</td>
<td>3.6</td>
<td>1.6</td>
<td></td>
</tr>
<tr>
<td>C(_3)H(_6)</td>
<td>19.6</td>
<td>0.9</td>
<td>14.8</td>
<td>3.9</td>
<td></td>
</tr>
<tr>
<td>CH(_3)CO(_3)CH(_3)</td>
<td>27.4</td>
<td>0.8</td>
<td>22.0</td>
<td>4.6</td>
<td></td>
</tr>
<tr>
<td>HCOOH</td>
<td>1.8</td>
<td>0.0</td>
<td>1.5</td>
<td>0.3</td>
<td></td>
</tr>
<tr>
<td>CH(_3)COOH</td>
<td>7.1</td>
<td>0.1</td>
<td>6.0</td>
<td>1.0</td>
<td></td>
</tr>
<tr>
<td>C(_4)H(_8)</td>
<td>510.0</td>
<td>23.2</td>
<td>441.9</td>
<td>45.0</td>
<td></td>
</tr>
<tr>
<td>C(_{10})H(_16)</td>
<td>85.4</td>
<td>2.3</td>
<td>70.2</td>
<td>12.9</td>
<td></td>
</tr>
</tbody>
</table>

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For validating the vertical distribution of relevant trace species such as O\(_3\), SO\(_2\) and CH\(_2\)O, we use measurements by the DC-8 aircraft during the Intercontinental Chemical Transport Experiment B (INTEX-B; Singh et al., 2009) that took place between March and May 2006. Observations of a host of co-located nitrogen-containing species are available (namely NO, NO\(_2\), PAN and HNO\(_3\)). These flights were conducted over a wide region, and we use all 3 months of measurements. Each month sampled a different region representing different meteorological conditions and local emission sources, namely, the Gulf of Mexico (90–100° W, 15–30° N), the remote Pacific (176–140° W, 20–45° N) and to the south and west of Alaska over the ocean (160–135° W, 20–60° N). Measurements cover altitudes up to 10.5 km, and we bin the values with respect to pressure using 50 hPa bins or less in the LT. We interpolated 3-hourly output against measurements for each respective day, similar to the comparisons performed in previous evaluations of TM5 (e.g. Huijnen et al., 2010), but we segregate our comparisons into the three distinct regions. For details relating to the location of each flight the reader is referred to the campaign overview of Singh et al. (2009).

For tropospheric O\(_3\), we supplement the INTEX-B comparisons with measurements taken over more polluted regions as part of the Measurement of Ozone, water vapour, carbon monoxide and nitrogen oxides by Airbus In-service aircraft initiative (MOZAIC; Thouret et al., 1998). We aggregate the measurements as seasonal means for December–January–February (DJF) and June–July–August (JJA) in order to provide a robust number of samples for each location. Here we choose to use profiles representative of the northern mid-latitudes, namely, London (0.2° W, 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 (121.8° E, 31.2° N) and Tokyo (140.4° E, 35.8° N).

We also make comparisons of O\(_3\), NO, NO\(_2\), selected N-reservoir species, SO\(_2\) and CH\(_2\)O profiles using measurements made aboard the NOAA WP-3D aircraft as part of the Second Texas Air Quality Study (TexAQS II; Parrish et al., 2009), which was conducted over the Texas seaboard during September and October 2006. This allows the assessment of TM5-MP over a region with higher NMVOC emissions and industrial activity. These measurements were typically sampled at altitudes below 500 hPa; therefore, no measurements in the UTLS are available from this campaign.

3 The effect on atmospheric transport

Here we analyse the differences in convective transport out of the BL by analysing the vertical and horizontal distribution of \(^{222}\)Rn, which is a diagnostic typically used for assessing the differences in transport in CTMs (e.g. Jacob et al., 1997). \(^{222}\)Rn is emitted at a steady rate and exhibits a half-life of \(\sim\) 3.8 days, which is long enough to be transported from the BL into the FT due to chemical passivity, with loss via wet scavenging and dry deposition being negligible. Therefore, it acts as an ideal tracer to assess differences in convective transport from the surface out of the BL. The representation of BL dynamics for TM5-MP has recently been assessed at 1° × 1° using \(^{222}\)Rn distributions for both the Tiedtke (1989)
scheme and when adopting convective mass transport values from the ERA-Interim meteorological data (Koffi et al., 2016).

Figure 1 shows seasonal mean horizontal global distributions of $^{222}$Rn for DJF and JJA in the $1^\circ \times 1^\circ$ simulation averaged between 800 and 900 hPa (i.e. sampling the LT). Also shown are the associated percentage differences against the re-binned $3^\circ \times 2^\circ$ $^{222}$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 $1^\circ \times 1^\circ$. In general it can be seen that seasonal differences of $\pm 20\%$ exist, typically with increases over continents and decreases over oceans in the $1^\circ \times 1^\circ$ simulations. Maximum differences of $> 60\%$ occur near selected coastal regions (California, western Africa, Madagascar) 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 $^{222}$Rn profiles ($1^\circ \times 1^\circ/3^\circ \times 2^\circ$) extracted above for selected European cities for January (black) and July (blue) 2006 is shown in Fig. S3 in the Supplement. The typical tropospheric profile of $^{222}$Rn exhibits an exponential decay from the LT to the FT (not shown). In order to homogenize the emission flux in the comparison, we coarsen the $1^\circ \times 1^\circ$ data onto the $3^\circ \times 2^\circ$ grid by averaging the six individual values into a representative mean column. The extent of the changes in the vertical distribution of $^{222}$Rn is somewhat site-specific, meaning an in-depth analysis is beyond the scope of this paper. In summary, the $1^\circ \times 1^\circ$ simulation generally provides stronger convective activity for January, with the main impact occurring below 700 hPa (e.g. London and Paris). The changes in $^{222}$Rn in the LT range between 2 and 10\% (i.e. ratios of 0.9 to 1.1), implying both weaker and stronger convective transport, depending on changes in location (e.g. orography and land type). The impact at Berlin is larger than e.g. Barcelona, which shows that, surprisingly, the inclusion of a large ocean fraction (with weaker convective mixing) in the $3^\circ \times 2^\circ$ cell does not seem to introduce dominating effects. Recently Koffi et al. (2016) have shown that comparisons of $^{222}$Rn at coastal sites in Europe at $1^\circ \times 1^\circ$ exhibit significant discrepancies compared to more continental stations. For July the changes in the vertical distribution extend into the FT up to 500 hPa, although changes in the upper FT have a significant component due to changes in long-range transport. The magnitudes 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. NO$_2$, CH$_2$O and SO$_2$ 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 between resolutions 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 Lumpur) and the extent of the ocean within any particular grid cell. Thus, differences in a priori vertical profiles of trace gases using a resolution of $1^\circ \times 1^\circ$ can be considerable compared to those provided at a $3^\circ \times 2^\circ$ resolution.

We also show ratios of profiles from $1^\circ \times 1^\circ$ simulations using the convective scheme of Tiedtke (1989) against those using the convective mass fluxes from the ERA-Interim meteorological dataset (Fig. S4), defined as ERA($1^\circ \times 1^\circ$)/Tiedtke ($1^\circ \times 1^\circ$). For this comparison no daily averaging is employed, with $^{222}$Rn profiles extracted from 3-hourly instantaneous sampling, with the profiles shown being interpolated directly above urban conurbations (with high trace gas emissions). The ratios show that the

<table>
<thead>
<tr>
<th>Term</th>
<th>Global</th>
<th>%</th>
<th>SH</th>
<th>%</th>
<th>Tropics</th>
<th>%</th>
<th>NH</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Net STE</td>
<td>579</td>
<td>-6.7</td>
<td></td>
<td></td>
<td>1615</td>
<td>-</td>
<td>49</td>
<td>5.8</td>
</tr>
<tr>
<td>Strat. nudging</td>
<td>1440</td>
<td>-0.7</td>
<td>-224</td>
<td>2.8</td>
<td>3938</td>
<td>-3.5</td>
<td>1206</td>
<td>-2.2</td>
</tr>
<tr>
<td>Trop. chem. pro.</td>
<td>5532</td>
<td>-1.9</td>
<td>389</td>
<td>-2.2</td>
<td>369</td>
<td>-2.5</td>
<td>853</td>
<td>-2.8</td>
</tr>
<tr>
<td>Trop. chem. loss</td>
<td>5162</td>
<td>-2.4</td>
<td>440</td>
<td>-1.0</td>
<td>3869</td>
<td>-2.5</td>
<td>853</td>
<td>-2.8</td>
</tr>
<tr>
<td>BO$_3$</td>
<td>378</td>
<td>-2.0</td>
<td>72</td>
<td>1.7</td>
<td>203</td>
<td>-2.3</td>
<td>104</td>
<td>-3.4</td>
</tr>
<tr>
<td>Strat. BO$_3$</td>
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<td>-2.0</td>
<td>23</td>
<td>9.1</td>
<td>38</td>
<td>-6.5</td>
<td>24</td>
<td>-2.0</td>
</tr>
<tr>
<td>Deposition</td>
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<td>0.8</td>
<td>115</td>
<td>0.6</td>
<td>465</td>
<td>-</td>
<td>369</td>
<td>1.9</td>
</tr>
<tr>
<td>O$_3$S deposition</td>
<td>97</td>
<td>5.0</td>
<td>19</td>
<td>7.5</td>
<td>37</td>
<td>-1.2</td>
<td>42</td>
<td>10.0</td>
</tr>
</tbody>
</table>
significant differences exist, with the convective mass fluxes from ERA-Interim being somewhat weaker than those calculated online using Tiedtke (1989), i.e. the ratio is typically less than 1, especially during July. In the recent study by Koffi et al. (2016) performed at 1° × 1° for the European domain, there was no appreciable improvement in the correlation coefficients when distributions of 222Rn were compared against measurements, resulting in no strong conclusion about which of the parameterizations results in better atmospheric transport.

4 The impact on tropospheric photolysis frequencies

The changes in the spatio-temporal distribution of tropospheric clouds and surface albedo have the potential to alter the incident flux of photolysing light reaching the surface, and thus photochemical production and destruction terms. When present, clouds dominate the integrated optical density in the tropospheric column. TM5-MP uses a random overlap method for determining the impact of clouds on actinic flux, which is weighted by cloud cover (Williams et al., 2012). Comparing seasonal mean cloud coverage for DJF and JJA (Fig. S5), we show that there are significant increases in the fractional cloud cover (fcc) at 1° × 1°, resulting in fcc values ranging between 0.1 and 0.8 (cf. 0.1–0.5 for 3° × 2°). Moreover, the definition of tropical equatorial cloud systems becomes much more defined and there are significant differences in the cloud distributions around the western coast of South America. For DJF, the largest changes occur at high latitudes over the tundra and oceans, but correspond to low-intensity incident radiation due to the polar winter. For the SH, the seasonal fcc increases significantly, which will potentially impose effects on Antarctic oxidative capacity (see Sect. 5). For JJA, most increases in fcc do not occur directly above high-NOx sources, but rather over the oceans. This limits the impact on the lifetime of chemical precursors (e.g. NO2) as discussed in Sect. 6. Examining similar plots for surface albedo (not shown) reveals that the maximum differ-

![Figure 1. The seasonal distributions of Rn²²² averaged between 800 and 900 hPa for DJF (top) and JJA (bottom) for the 3° × 2° (left) and 1° × 1° (right) simulations, with the associated percentage differences when compared against the 3° × 2° simulation.](image-url)
ences (increases at $1^\circ \times 1^\circ$) again occur in the polar regions under low temperatures related to sea ice and snow coverage, typically during polar winters. For mid-latitudes and tropics, although differences in the absolute albedo value can be significant ($\pm 50\%$), values are typically below 0.1, which will contribute to the perturbations in the final $J$ value tropospheric profiles as discussed below. The monthly mean comparisons in surface $J$ values provided in Fig. S6 show that any differences in instantaneous cloud cover are moderated to the order of a few percent when looking at longer periods.

The similarity in the monthly mean photolysis frequencies for $O_3$ and $NO_2$ across resolutions (hereafter denoted $J_{O_3}$ and $J_{NO_2}$, respectively) are shown in Fig. S6 of the Supplement. Comparisons of the monthly mean $J_{O_3}$ and $J_{NO_2}$ values are shown at five different locations identical to those shown in Williams et al. (2012). For $J_{O_3}$, the impact of increasing resolution is limited to a few percent in the monthly mean values, even for regions which have high surface albedo. At the global scale this leads to a reduction of $\sim 2\%$ in the total mass of $O_3$ photolysed (not shown). For $J_{NO_2}$, the corresponding differences become more appreciable, with $1^\circ \times 1^\circ$ exhibiting $\sim 5$–$10\%$ higher values at high northern latitudes (associated with locations with high-$NO_x$ regimes). Focusing on $J_{NO_2}$ and comparing seasonal mean values near the surface show that very similar large-scale spatial patterns occur for both simulations at the global scale (cf. Fig. S7). The highest $J_{NO_2}$ values occur over the tropical oceans and high-altitude regions (e.g. Nepal), with a latitudinal shift related to seasonal changes in daylight hours. Although a more regional fine structure can be seen at $1^\circ \times 1^\circ$ (e.g. the south-western US and south-western China for DJF), these seasonal averages show that the small perturbations in $J_{NO_2}$ shown in Fig. S6 extend to the global scale, leading to only modest changes in the tropospheric lifetime of $NO_2$ (see Sect. 6).

Comparisons of monthly mean vertical profiles of $J_{O_3}$ and $J_{NO_2}$ as sampled over selected tropical cities are shown in Fig. S8a and b, respectively, in the Supplement. 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 $J_{O_3}$ profiles are affected to a larger extent than the $J_{NO_2}$ profiles, due to the characteristic absorption spectra of each species which make $J_{O_3}$ 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 $J_{O_3}$ and $J_{NO_2}$ 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 $\pm 5$ and $10\%$ in the monthly mean profiles. The changes in $J_{NO_2}$ reflect those simulated for $J_{O_3}$, with somewhat smaller perturbations.

5 Implications for oxidative capacity and tropospheric $O_3$

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 radicals (i.e. OH, CH$_3$O(O)O$_2$, NO$_3$ and CH$_3$O$_2$). Therefore, changes to the distribution and resident mixing ratios of tropospheric $O_3$ subsequently impose changes in the fractional composition of the NO$_x$ budget (Olszyna et al., 1994) and also the efficiency of the NO$_x$ recycling terms by altering the chain length (Lelieveld et al., 2002). In this section we analyse the global and zonal chemical budget terms for tropospheric $O_3$ to highlight the inter-hemispheric differences which occur (i.e. under low- and high-NO$_x$ environments). An overview of the resulting near-surface global distribution of tropospheric $O_3$ for May 2006 is shown in Fig. 3, which also includes the location of the regional comparisons presented below in a larger global context. In general, the pattern of minimum and maximum mixing ratios in $O_3$ occurs in similar locations, with the long-range transport component being more clearly defined in the $1^\circ \times 1^\circ$ simulation. There is a distinct latitudinal gradient in $O_3$ mixing ratios imposed by the global distribution of NO$_x$ emissions.

Table 4 provides the zonally segregated chemical budget terms for tropospheric $O_3$, from which the global component due to STE can be determined by closing the budget terms following the methodology given in Stevenson et al. (2006). The chemical tropopause calculated for $3^\circ \times 2^\circ$ is applied for the analysis of $1^\circ \times 1^\circ$ budget terms to ensure that a valid comparison is performed, (i.e. the same mass of air is accounted for). For computational efficiency the budget terms are aggregated in $10^\circ$ latitudinal bins for each vertical level and summed across all longitudes, providing the cumulative latitudinal terms.

The most significant change with resolution concerns STE. By using a dedicated tagged stratospheric $O_3$ tracer (which only undergoes photo-chemical destruction and deposition in the troposphere; hereafter denoted as $O_3S$), changes in the zonal mean STE can be determined. The stratospheric burden of $O_3$ (BO$_3$ (strat)) exhibits a strong hemispheric gradient with much more downwelling occurring in the Northern Hemisphere (NH), peaking during boreal springtime. At the global scale the STE exchange is $579$ Tg $O_3$ yr$^{-1}$, which agrees well with the multi-model mean for STE of $556 \pm 154$ Tg $O_3$ yr$^{-1}$ in Stevenson et al. (2006), with observational estimates being $\sim 550 \pm 140$ Tg $O_3$ yr$^{-1}$ (Olsen et al., 2001). The $\sim 7\%$ reduction of STE at $1^\circ \times 1^\circ$ is encouraging considering that previous studies using TM5 have concluded that STE in TM5 at $3^\circ \times 2^\circ$ was biased high com-
Figure 2. Zonal mean seasonal distribution of the TM5-MP O$_3$/O$_3$ ratio for the 3$^\circ$ × 2$^\circ$ (left) and 1$^\circ$ × 1$^\circ$ (right) simulations.

pared to STE inferred from TES and MLS satellite observations (Verstraeten et al., 2015). The increase in STE in the SH, with an associated decrease in the NH (see below), implies that there is a shift in circulation patterns at 1$^\circ$ × 1$^\circ$ even though BO$_3$ (strat) remains essentially unchanged. Previous studies have shown that in order to resolve the correct spatial and temporal stratosphere–troposphere flux, high resolution is required both in the horizontal and the vertical gridding (e.g., Meloen et al., 2002). The NH STE diagnosed with TM5-MP is an order of magnitude smaller than estimates derived in a CTM study also conducted at a 1$^\circ$ × 1$^\circ$ resolution (Tang et al., 2011; 382 70–20 Tg O$_3$ yr$^{-1}$), which identified deep convection as important for STE. Here we use a different vertical grid and meteorological dataset to drive TM5-MP, both of which affect the ability to capture an accurate STE flux (Meloen et al., 2002). For the 1$^\circ$ × 1$^\circ$ simulation using the Tiedtke scheme, there is a further reduction in the STE component of 21 Tg O$_3$ yr$^{-1}$, resulting in an STE component almost identical to the multi-model mean in Stevenson et al. (2006).

The zonal seasonal means of the fraction of O$_3$S to O$_3$ (O$_3$S/O$_3$) for both simulations are shown in Fig. 2 for DJF and JJA. There is a clear seasonal zonal shift in the fractional contribution due to the O$_3$ transported downwards from the stratosphere exhibiting a longer lifetime in the winter hemisphere, reflecting a lower photochemical destruction rate. At 1$^\circ$ × 1$^\circ$ the largest increase in STE occurs in the SH during JJA. Here 20–25 % of tropospheric O$_3$ is transported down from the stratosphere. Comparing the 0.2 contour for the NH mid-troposphere shows significant changes, extending further down towards the surface during boreal wintertime, leading to the higher total mass of O$_3$S in the troposphere. The extent of nudging towards the MSR climatology is essentially constant across simulations (cf. Table 4). Interestingly, less O$_3$S reaches the surface in the tropics at 1$^\circ$ × 1$^\circ$ due to the enhanced chemical destruction term in the free troposphere. Approximately 10 % of the global deposition term for O$_3$ is associated with O$_3$ that originates from the stratosphere at 1$^\circ$ × 1$^\circ$ (H. 5 % at 3$^\circ$ × 2$^\circ$). For the NH, this contributes to the simulated increase in deposition of 9 %.

For tropospheric O$_3$ there are similarities that occur between the NH, tropics and SH, i.e. high- and low-NO$_x$ scenarios, resulting in a cumulative decrease in O$_3$ production of 2–4 % across zones. For the chemical loss terms there is a modest decrease of 3 % (~ 2 %) in the NH (SH) reflective of the changes discussed for JJA, which acts as the primary destruction term. Therefore, in the SH the significant differences shown for fcc do not significantly impose a lower photochemical destruction term on the annual tropospheric O$_3$ budget. There is a zonal gradient in the tropospheric burden of O$_3$ (BO$_3$) following the zonal gradient in NO$_x$ emissions (cf. Fig. 3). Comparing terms shows that BO$_3$ decreases at 1$^\circ$ × 1$^\circ$ by a few percent at the global scale (~ 7 Tg O$_3$), making a rather small impact on oxidative capacity. This is of the same order of magnitude as that found in previous studies concerned with horizontal resolution (e.g., Wild and Prather, 2006). Interestingly, changes in the deposition flux of O$_3$ are rather small, even though there is a larger amount of variability in the land surfaces and a better-resolved land–sea contrast at 3$^\circ$ × 1$^\circ$, although differences in regional deposition fluxes can be more significant. Multi-model inter-comparisons of surface deposition terms across models have shown previous versions of TM5 to be at the low end of the model spread in terms of O$_3$ (Hardacre et al., 2015), suggesting that the surface deposition flux should be increased by 10 % in TM5-MP towards the multi-model mean value. This can be partly attributed to the large uncertainty which exists related to the loss of O$_3$ to the ocean (Hardacre et al., 2015).

Figure 4 shows comparisons of simulated and observed mass mixing ratios of surface O$_3$ at EMEP sites across Europe (www.emep.int; Aas et al., 2001), with stations chosen to cover a range of latitudes. Previous comparisons using mCB05 have revealed high biases in surface O$_3$, especially during boreal summertime (Williams et al., 2013). These high biases originate from cumulative effects associated with the accuracy of the emission inventories, the convective and turbulent mixing component, the underestimation of the scattering and absorption of photolysing light due to aerosols and the chemical mechanism that is employed. For the emission component it should be noted that even at 1$^\circ$ × 1$^\circ$ coarsening is performed, where emission inventories are typically supplied at 0.5$^\circ$ × 0.5$^\circ$ resolution. The seasonal cycle in surface O$_3$ is captured to a large degree, and the high bias exhibited by the model is generally reduced by 2–5 ppb (or 20 %).
at 1° × 1°. This is associated with perturbations in the NO\textsubscript{x} recycling terms, chemical titration by NO, changes to the turbulent diffusion and convective mixing out of the BL. The improvement in biases is largest during boreal summertime, associated with the shorter chain length of the NO\textsubscript{x} recycling term during boreal wintertime. However, there is still a significant monthly mean bias in both simulations when compared against observations throughout the year, especially for locations impacted by a large anthropogenic NO\textsubscript{x} source. This is partly due to the low NO / NO\textsubscript{2} ratio as discussed in Sect. 6 below.

Comparing vertical profiles from composites assembled from the MOZAIC measurements for DJF and JJA (Figs. S9a and S8b, respectively), INTEX-B (Singh et al., 2009; Fig. S10) and TexAQS II (Parrish et al., 2009; Fig. S11) show consistently that differences are small between simulations across regions, and typically mimic those which occur at the surface. There is a general positive bias of ~20–40 % in mixing ratios exhibited across all comparisons, although the variability in the vertical gradients across regions is captured rather well. Such positive biases have consequences for both the NO\textsubscript{x} recycling terms and HNO\textsubscript{3} formation discussed in the sections below.

6 Implications for the distribution of NO and NO\textsubscript{2}

Table 5 provides the zonally segregated annual NO\textsubscript{x} recycling terms involving the main peroxy radicals and the direct titration term involving NO for the 1° × 1° simulation. The conversion rate of NO back into NO\textsubscript{2} decreases by ~2–3 % across zones as a consequence of an associated increase in the titration term and re-partitioning of N into long-lived reservoir species (see below). For the titration term involving NO, although the globally integrated flux remains relatively constant, there is contrasting behaviour for the two most important zones (TR, NH), which exhibit a lower and higher titration term, respectively. It has been shown that for regions such as Europe the increased titration results in lower surface
O$_3$ mixing ratios (cf. Fig. 4), improving the boreal summertime high bias at the surface.

Important model uncertainties include the quality of the MACCity NO$_x$ emission inventory, the lifetime of NO$_2$ simulated in TM5, BL mixing and the NO$_x$ recycling term via the chemical titration of O$_3$. Figure 3 provides an illustration of the global distribution in surface NO$_2$ during May 2006 for both the 3° × 2° and 1° × 1° simulations, where the short lifetime means that the maximum mixing ratios occur directly near the strong source regions. Most NO$_x$ is anthropogenic in origin; therefore, there is a strong latitudinal gradient between the NH and SH, with ship tracks also visible. The regions where validation occurs are also superimposed.
in the figure, including the extent of the EMEP domain over which NO\textsubscript{2} weekly comparisons are made.

Figures 5 and 6 show comparisons of weekly [NO] and [NO\textsubscript{2}] surface measurements with the corresponding composites from both simulations, sampled at 13:00 local time, which is close to the local overpass time for both OMI and tropOMI. Although the number of EMEP sites conducting NO\textsubscript{2} measurements is smaller than those measuring O\textsubscript{3}, we choose stations located throughout Europe in both high- and low-NO\textsubscript{x} regimes. To supplement these comparisons we provide the seasonal mean biases for DJF and JJA from both simulations in Tables 6 and 7, respectively, calculated using weekly binned data from all EMEP sites that measure hourly [NO] and [NO\textsubscript{2}]. Here we perform an analysis across sites rather than focusing on the behaviour at selected individual locations.

For the determination of [NO\textsubscript{2}], the reduction of NO on a Molybdenum convertor takes place with subsequent detection by chemi-luminescence, with an associated detection limit of \( \sim 0.4 \text{ ppb} \). Previous studies have shown that some bias can result due to the oxidation of nitrogen reservoirs such as PAN (Dunlea et al., 2007; Steinbacher et al., 2007). In TM5-MP all NO\textsubscript{2} emissions are introduced as NO, although a fraction for road transport is known to be emitted directly as NO\textsubscript{2} (e.g. Carslaw and Beevers, 2005). Many studies have been performed comparing satellite NO\textsubscript{2} columns with model values, implying that inadequacies in emission inventories are somewhat region-specific (e.g. Zyrichidou et al., 2015; Pope et al., 2015).

Table 6 shows a negative bias of a few \( \mu \text{g m}^{-3} \) in TM5-MP in seasonal surface [NO] in Europe. This is a cumulative effect of the accuracy of the MACC NO\textsubscript{3} emission estimates, an overestimate in daytime vertical mixing (Koffi et al., 2016) (enhanced dilution) and, to a lesser extent, overly high surface [O\textsubscript{3}] (increasing the oxidation rate of NO to NO\textsubscript{2}). As anthropogenic emissions are the principle source of NO, there is no significant seasonal cycle in the monthly emission estimates in the NH. Seasonal differences in convective mixing (i.e. lower BL heights) do cause somewhat higher surface [NO] during DJF for approximately equal emission terms. This is captured by TM5-MP, although under night-time conditions TM5-MP has been shown to overestimate nocturnal BL heights (Koffi et al., 2016). For \( \sim 80 \% \) of the EMEP sites we do not observe any significant change in the quality of the comparisons. For \( \sim 20 \% \) of sites, simulations of [NO] at 1\(^\circ\) \( \times \) 1\(^\circ\) introduce significant improvements over those at 3\(^\circ\) \( \times \) 2\(^\circ\), and there is an improvement regarding the extent of seasonal variability (Fig. 4).

Table 7 shows that for [NO\textsubscript{2}] the biases are more variable, being typically in the range of \( \pm 0.6 \mu \text{g m}^{-3} \), with both positive and negative biases occurring across sites. The conversion efficiency from NO, loss to reservoir compounds (e.g. HNO\textsubscript{3}), photo-dissociation rate, convective mixing and emission estimates all contribute to these biases. The seasonal biases show improvements at 1\(^\circ\) \( \times \) 1\(^\circ\) for \( \sim 35 \% \) of the EMEP sites, accompanied by degradations at \( \sim 20 \% \) of the sites. The maximum biases in [NO\textsubscript{2}] at 1\(^\circ\) \( \times \) 1\(^\circ\) can be approximately double those for [NO]. For the corresponding NO / NO\textsubscript{2} ratio, there will generally be an underprediction in the model due to the negative biases shown for the [NO] comparisons. Analysing the corresponding seasonal correlation coefficients (not shown) shows that in \( \sim 25 \% \) of the cases there is little seasonal correlation between the weekly [NO\textsubscript{2}] in TM5-MP and the measurements regardless of resolution for both seasons (Pearson’s \( r \) in the range -0.3 to 0.3). In \( \sim 30 \% \) of cases there is actually a degradation in \( r \) between resolutions, the changes somewhat reflect those seen in the seasonal biases, i.e. simultaneous changes to both the meteorology and local emission fluxes do not necessarily improve the performance of the model. Comparing 1\(^\circ\) \( \times \) 1\(^\circ\) values both with and without the Tiedtke convection scheme shows that for the most convective regions (e.g. south of 45\(^\circ\)N) increases in \( r \) generally occur during JJA when employing the ERA-Interim mass fluxes. Conversely for e.g. Finland the correlation becomes worse.

Beyond Europe, we compared monthly mean TM5-MP vertical distributions of NO and NO\textsubscript{2} between March and May 2006 against measurements taken during the INTEX-B campaign in Fig. 7. In general differences between 1\(^\circ\) \( \times \) 1\(^\circ\) and 3\(^\circ\) \( \times \) 2\(^\circ\) simulations are of the order of a few percent,
with NO₂ biased low in the LT 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 even at 1° × 1°. For March, there is a strong signature from biomass burning plumes in the middle troposphere which is not captured using the monthly burning estimates. For the FT, TM5-MP captures the observed gradient to a reasonable degree. In the UT there is a consistent high bias for NO and an associated low bias for NO₂, suggesting that the conversion term is too low and that the NOₓ cycle is out of synch at these cold temperatures despite the addition of new reservoir species (i.e. CH₃O₂NO₂) and application of new rate data.
One important gauge as to whether the chemical mechanism can capture the correct recycling efficiency of NO into NO$_2$ is to examine their ratio, which is presented in the third column of Fig. 7. In the LT ($<900$ hPa) NO / NO$_2$ ratios of 0.1–0.2 exist that are captured quite well by TM5-MP, with negligible differences between $3^\circ \times 2^\circ$ and $1^\circ \times 1^\circ$ simulations. For the FT, TM5-MP consistently overestimates the ratio in spite of a high bias in O$_3$ (cf. Fig. 4) which is imposed by the overestimates in NO$_2$. This implies the chemical conversion is too slow and, assuming representative $J_{NO_2}$ values, indicates a low bias in HO$_2$ or an underestimation in the mixing ratios of other long-lived and short-lived NO$_x$ compounds (see Sect. 7).

Finally in Fig. 8 we show the corresponding comparisons against measurements taken during the TexAQS II campaign (Parrish et al., 2009) for both September and October 2006. As for the EMEP comparisons shown in Figs. 5 and 6, there is a significant underestimation in NO and NO$_2$ mixing ratios, with both model profiles being outside the 1σ variability in the observational mean. This is clearly related to the emission estimates for this region being underestimated in the current emission inventories (e.g. Kim et al., 2011). For the resulting NO / NO$_2$ ratio, TM5-MP captures the correct ratio in the lowest few hundred metres of the BL, but overestimates the ratio at higher altitudes as for more pristine environments, although there is marked improvement in the ratios simulated for October.

7 Changes in the NO$_x$ budget

7.1 Long-lived reservoirs

The resolution-dependent changes in the temporal distribution of [NO$_2$] and associated differences in NMVOC chemical precursor emissions have the potential to alter the partitioning of reactive NO$_x$ between the three main chemical reservoirs included in mCB05v2 (i.e. HNO$_3$, PAN and ORGNTR). The differences in both the deposition efficiency and tropospheric lifetimes between trace species at $1^\circ \times 1^\circ$ suggests that the fraction of NO$_x$ that can be transported out of source regions could change significantly. Here we briefly examine the zonally integrated nitrogen budget terms between simulations to quantify the effect of applying a higher spatial resolution. The seasonal distribution of these three
dominant reservoir species at 1° × 1° and their individual contributions to total NO\textsubscript{y} are shown in Figs. S12–S15 for DJF and JJA, respectively. Here we define NO\textsubscript{y} as the cumulative total of NO, NO\textsubscript{2}, NO\textsubscript{3}, HNO\textsubscript{3}, PAN, CH\textsubscript{3}O\textsubscript{2}NO\textsubscript{2}, HONO, 2-N\textsubscript{2}O\textsubscript{5}, lumped organic nitrates (ORGNTR) and HNO\textsubscript{4}. It should be noted that methyl-nitrate is not in this version of TM5-MP. These figures are provided as a reference for the reader to aid understanding of the discussion below.

Table S1 in the Supplement provides a zonal decomposition of the tropospheric chemical budget terms for HNO\textsubscript{3}, PAN and ORGNTR. For HNO\textsubscript{3}, even though the recent kinetic rate parameters increase (decrease) the chemical production term at the surface (UTLS) compared to older rate data (e.g. Seltzer et al., 2015), changes in the integrated column term are small. The changes at 1° × 1° are somewhat latitude-dependent (corresponding to low- and high-NO\textsubscript{y} regimes), with only small increases occurring in the NH and associated decreases in the tropics related to lower [OH] (i.e. chemical production). Loss by cumulative deposition terms only changes by a few percent, due to wet scavenging being so efficient for HNO\textsubscript{3} for any associated change in the SAD of cloud droplets (cm\textsuperscript{3}/cm\textsuperscript{3}) introduced by changes in the liquid water product.

For PAN, both the production and destruction terms decrease marginally by ∼1–3% across all zones, meaning the transport of NO\textsubscript{y} out of the main source regions remains relatively robust. The total mass of N cycled through PAN is ∼4 times that sequestered as HNO\textsubscript{3}, although the lifetime of PAN is shorter due to the efficient thermal decomposition. The changes in the production term due to temporal increases in NO\textsubscript{2} near high-NO\textsubscript{y} source regions (cf. Fig. 3) are partially offset by a reduction in the mixing ratios of the acetyl-peroxy radical (C\textsubscript{2}O\textsubscript{2}H) due to e.g. increased dry deposition of organic precursors at 1° × 1°. Although the chemical budget terms only exhibit small changes, it can be expected that the global distribution of PAN will be somewhat different due to the changes in the convective and advective mixing due to

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<td>−1.63</td>
<td>−1.18</td>
<td>−0.19</td>
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<tr>
<td>NL10</td>
<td>51.32</td>
<td>5.51</td>
<td>3.48</td>
<td>3.52</td>
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<td>−0.29</td>
</tr>
<tr>
<td>NL91</td>
<td>52.18</td>
<td>4.30</td>
<td>11.81</td>
<td>6.42</td>
<td>1.48</td>
<td>−0.81</td>
</tr>
</tbody>
</table>
Figure 7. Monthly mean comparisons of NO (left), NO$_2$ (middle) and the resulting NO / NO$_2$ ratio (right) from INTEX-B measurements and TM5-MP simulations. The dotted line represents the 1σ deviation in the mean of the measurements. For details of the locations for each month the reader is referred to Singh et al. (2009).

the application of higher-resolution meteorological data (cf. Sect. 3).

For ORGNTR, there is a 5% reduction in the annual production term at 1° × 1°, with an associated decrease in the loss by deposition. Both the largest production and, thus, destruction terms occur in the tropics related to the strongest source of ORGNTR being biogenic precursors in mCB05v2. Thus, at 1° × 1°, this intermediate trace species becomes less important as a NO$_x$ reservoir.

Finally, the one additional intermediate not shown is CH$_3$O$_2$NO$_2$, which is primarily a stable vehicle for transporting NO$_x$ from the surface up to the UTLS, where at cold temperatures it accounts for a significant fraction of NO$_2$ speciation along with HNO$_4$ (Browne et al., 2011). At the global scale 3 times as much nitrogen cycles through CH$_3$O$_2$NO$_2$ compared to PAN, although the thermal stability is low at temperatures > 255 K; thus, resident mixing ratios are typically small. This results in maximum mixing ratios occurring in the cold upper troposphere (up to ~0.2 ppb) and sub-
where the chemical budget terms for all three species are provided in Table S2 in the Supplement. For HONO it should be noted that many tropospheric CTMs have difficulty in simulating observed mixing ratios (e.g. Gonçalves et al., 2012), suggesting missing (heterogeneous) source terms. The global impact of increased resolution on HONO production is rather muted for most weeks, resulting in no significant changes to the seasonal biases (not given), as constrained by the improvements in surface [NO2] (cf. Fig. 6). The heterogeneous scavenging of HNO3 into ammonium nitrate can act as a moderator of gaseous HNO3. Although this heterogeneous conversion process is included in TM5-MP as described by the EQSAM approach, low concentrations of e.g. ammonium nitrate (not shown) typically result. Thus, gaseous [HNO3] remains too high due to too little conversion into particles and subsequent deposition.

For other regions outside Europe, we make comparisons of vertical profiles of HNO3 and PAN between March and September 2006 with those measured during INTEX-B (Fig. 10) and TexAQSII (Fig. S16). PAN is a good marker for transport in the free troposphere due to the relatively long lifetime at colder temperatures. For all regions the vertical gradients for both species are captured quite well, although some fine structure is lost due to the vertical resolution of TM5-MP and insufficient precursor emissions. This implies that the underestimation in NO2 simulated in the UTLS (Fig. 7) is not due to insufficient transport of NO2 away from source regions, but rather should be attributed to either missing chemistry or insufficient transport down from the stratosphere. Finally, for more polluted regions, the vertical gradient of HNO3 is rather less steep than that observed, with significant low biases in the lower troposphere related to the low bias in NO2 shown in Fig. 8. The impact of the 1° × 1° resolution only results in a marginal improvement in the LT for HNO3 (again similar to NO2). For PAN the vertical profile in TM5-MP agrees remarkably well, but is somewhat anti-correlated around 900 hPa in both simulations, and the rapid decrease in the lowest kilometre is captured sufficiently. This overestimation would likely be larger if the emission estimates were increased as required to consolidate the NO2 comparisons for the same campaign.

7.2 Short-lived reservoirs

Here we briefly discuss the perturbations introduced for the short-lived N-reservoirs, namely HONO, HNO4 and N2O5, where the chemical budget terms for all three species are provided in Table S2 in the Supplement. For HONO it should be noted that many tropospheric CTMs have difficulty in simulating observed mixing ratios (e.g. Gonçalves 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° × 1° there is ~10 % more chemical production of HONO in high-NOx regions and no appreciable effect in low-NOx 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$_x$ regions (cf. Fig. 5). The muted response is due to competing oxidative processes, which effectively lower the OH available. For HNO$_4$, approximately the same mass of N cycles through this species as for PAN, although the shorter lifetime means that it is more important at regional scale. Again, the impact of resolution on this species is small, where decreases in [HO$_2$] 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$_2$O$_5$, with enhanced chemical production of $\sim$12% at the global scale, increasing the heterogeneous sink term by $\sim$6%, although the changes in the total mass of N converted are small. In general, this is due to an increase in the production of the NO$_3$ radical by $\sim 10\%$ at 1° × 1° (not shown) resulting in enhanced N$_2$O$_5$ mixing ratios.

8 Implications for tropospheric CH$_2$O retrieval

The implications of applying a higher-resolution CTM for the global distribution of CH$_2$O are rather modest. Figure S17 shows the near-surface global distributions of CH$_2$O for May 2006, where maximum mixing ratios occur near forested regions due to the link with isoprene oxidation (e.g. Palmer et al., 2006). The tropospheric lifetime of CH$_2$O is of the order of a few days, meaning that transport has little impact between simulations, apart from in low-emission areas. Also shown are the locations where regional validation occurs. In Table 8, we show zonally integrated chemical production and destruction terms for CH$_2$O, which suggests changes of the order of a few percent at the global scale. The most notable difference is the increase in the cumulative deposition term of $\sim 4\%$ at 1° × 1°, thus reducing the atmospheric lifetime of CH$_2$O in TM5-MP. Again, this low impact shows that the increase in the temporal variability of the meteorological data at 1° × 1°, and thus the local variability of cloud SAD, 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° × 1° due to the better representation of regional precursor source 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$_3$OOH; a reduction of $\sim 3$–5 Tg CH$_2$O yr$^{-1}$).

For the chemical destruction term, the relative insensitivity of the photolysis of CH$_2$O to resolution (similar to $J_{O_3}$; cf. Fig. S4) results in small net decreases in line with changes in the chemical production term.

Figure 11 compares monthly mean tropospheric profiles of CH$_2$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$_2$O by TM5-MP for all months shown, although surface mixing ratios are typically too high, suggesting that loss by deposition to the ocean is underestimated (potentially related to underestimations in surface area due to lack of 3-D wave structure) or that the chemical production term is too efficient. Moreover, there appears to be a missing (chemical) source term in the UTLS in TM5-MP, leading to a $\sim 30$–$50$ % ($\sim 0.05$ ppb) low bias above 600 hPa, i.e. there is
Table 8. The tropospheric chemical budget for the CH$_2$O given in Tg CH$_2$O yr$^{-1}$ during 2006 for the 1$\degree$ × 1$\degree$ simulation. Percentage differences are shown against the corresponding 3$\degree$ × 2$\degree$ simulation.

<table>
<thead>
<tr>
<th>Budget term</th>
<th>Global</th>
<th>% SH</th>
<th>% Tropics</th>
<th>% NH</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH$_2$O CP</td>
<td>1919</td>
<td>−1.1</td>
<td>147</td>
<td>−0.3</td>
<td>1491</td>
</tr>
<tr>
<td>CH$_2$O CD</td>
<td>1739</td>
<td>−1.6</td>
<td>134</td>
<td>−0.5</td>
<td>1349</td>
</tr>
<tr>
<td>CH$_2$O dep.</td>
<td>193</td>
<td>3.1</td>
<td>15</td>
<td>2.0</td>
<td>149</td>
</tr>
</tbody>
</table>

9 Implications for tropospheric SO$_2$ retrieval

The global distributions of near-surface SO$_2$ mixing ratios for both the 3$\degree$ × 2$\degree$ and 1$\degree$ × 1$\degree$ simulations are shown in the bottom panels of Fig. S17, where the distribution shows the land-based point sources as applied from the MACCity emission inventory. The high mixing ratios of SO$_2$ correlate with the location of strong anthropogenic emission sources due to the relatively short atmospheric lifetime of SO$_2$ (varying between ~2 days during winter and ~19 h during summer; Lee et al., 2011), being rapidly oxidized to sulfate (SO$_4$$^2$-). Although the regional distributions are similar, the 1$\degree$ × 1$\degree$ simulation is able to differentiate point sources to a much better degree, which enhances the ability to derive more accurate emission fluxes. The in situ chemical production term of SO$_2$ from the oxidation of oceanic di-methyl sulfide (DMS) is low; thus, there are very low SO$_2$ mixing ratios in the chemical background. Also shown are the regions used for validating the SO$_2$ surface concentrations and vertical profiles (insets in Fig. S17; see below).

In Fig. 12, we compare weekly [SO$_2$] for 2006 at a number of EMEP sites in Austria (AT02, forested), the Netherlands (NL09, rural), Great Britain (GB43, rural) and Spain (ES10, rural), with most sites being positioned away from strong point sources. For SO$_2$ in Europe, the main emission source is anthropogenic (e.g. from the energy sector). High [SO$_2$] 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 methodologies employed, it is typically around ~1.3 µg m$^{-3}$ (e.g. Hamad et al., 2010). Comparing weekly averages shows that for most sites shown there is a significant low bias at 3$\degree$ × 2$\degree$, indicating inaccuracies in the MACC emission inventory and the effect of coarsening on the model resolution. At 1$\degree$ × 1$\degree$ significant improvements occur in the correlations as a result of the better temporal distribution of anthropogenic emission sources.

Table 9 provides an overview of the seasonal biases for all of the EMEP sites that measure hourly [SO$_2$], with the biases calculated for the overpass time of tropOMI aggregated on a weekly basis. Improvements occur at 1$\degree$ × 1$\degree$ 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$_2$] is determined more by long-range...
transport (and is thus sensitive to wash-out) than a local emission source, where strong mitigation practises have been implemented in Europe over the last few decades reducing resident [SO2] significantly (Tørseth et al., 2012). For some sites there is a notable increase in biases at 1° × 1° (20 % DJF, 25 % JJA), indicating that overly 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 (Fig. S19) and TexAQS II (Fig. 13) as for the other trace gas species. For the more pristine locations there are typically low biases at 3° × 2° for all months, especially at the surface during March, indicating a significant underestimation of the emission fluxes of SO2. Increasing to 1° × 1° only provides an improved correlation for March, due to the transport in the FT being described better, as shown for NO2 in Fig. 7. For April, the comparison shows a significant underestimation in the column for both simulations, where corresponding comparisons of the vertical profiles of DMS, which acts as a key source of SO2 in the equatorial Pacific (Alonza Gray et al., 2011), agree quite well (not shown). This points to an appreciable biogenic source term that is currently missing from the inventories as proposed for organic nitrates (Williams et al., 2014). For May, again no significant improvement occurs at 1° × 1°, although both simulations capture the peak in SO2 mixing ratios at the top of the BL. More relevant for satellite-based retrievals is the observed column near strong anthropogenic source regions as shown in Fig. 13 over Texas during September and October 2006. Here a clear improvement occurs at 1° × 1°, with the low bias in the BL being reduced significantly, although the integrated column is still too low. Again this is due to the underestimation in the source emission fluxes in the anthropogenic emission inventory employed.

### 10 Conclusions

In this paper we have provided a comprehensive description of the high-resolution 1° × 1° version of TM5, which is to be used for the purpose of providing a priori columns for the satellite retrieval of trace gas columns of NO2, CH2O and SO2. By performing identical simulations at a horizontal resolution of 3° × 2° and 1° × 1°, and comparing the resulting global distributions of trace gas species, photolysis frequencies and chemical budget terms, we quantify and validate both the near-surface and vertical distributions for the evaluation year of 2006.

Comparing the seasonal distribution in 222Rn between resolutions, we show differences in the vertical distribution of up to ±20 % at the global scale, with significantly larger impacts for specific coastal regions and tropical oceans. In order

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**Table 9.** The seasonal mean biases of daily [SO2] (µg m$^{-3}$) at 13:00 for DJF and JJA, when taking the difference between measurements–model values. Values are shown for both the 3° × 1° and 1° × 1° simulations. Those with differences < 5 % are considered to exhibit no discernible change in the bias.

<table>
<thead>
<tr>
<th>EMEP station</th>
<th>Lat</th>
<th>Lon</th>
<th>DJF 3° × 2°</th>
<th>DJF 1° × 1°</th>
<th>JJA 3° × 2°</th>
<th>JJA 1° × 1°</th>
</tr>
</thead>
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<td>AT02</td>
<td>47.46</td>
<td>16.46</td>
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<td>−3.15</td>
<td>−0.89</td>
<td>−0.53</td>
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<td>−0.41</td>
<td>−0.14</td>
<td>−0.14</td>
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<tr>
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<td>−0.63</td>
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<td>−0.15</td>
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<td>CZ03</td>
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<td>3.65</td>
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<tr>
<td>ES07</td>
<td>37.14</td>
<td>3.32</td>
<td>1.22</td>
<td>0.73</td>
<td>0.38</td>
<td>0.31</td>
</tr>
<tr>
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<td>43.26</td>
<td>16.46</td>
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<td>−3.21</td>
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<tr>
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</tr>
<tr>
<td>ES13</td>
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<td>−0.81</td>
<td>2.71</td>
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<td>41.24</td>
<td>0.43</td>
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<tr>
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<td>39.31</td>
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<td>52.17</td>
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<td>52.50</td>
<td>6.34</td>
<td>2.58</td>
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<td>0.46</td>
<td>−1.03</td>
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<tr>
<td>NL08</td>
<td>52.70</td>
<td>5.12</td>
<td>1.77</td>
<td>1.56</td>
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<td>−0.30</td>
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<tr>
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<td>6.16</td>
<td>2.53</td>
<td>2.16</td>
<td>0.47</td>
<td>0.27</td>
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</table>
to assess the changes in convective activity above strong NOx sources, we show that differences of between \( \sim 2 \) and 10\% (\( \sim 10–20\% \)) exist for the northern mid-latitudes (tropics) at higher resolution, with both weaker and stronger upwelling occurring depending on the region and the season. The magnitudes of the changes are site-specific, being affected by local orography. We have also made comparisons using a \( 1^\circ \times 1^\circ \) simulation applying the Tiedtke (1989) convection scheme, showing that ERA-Interim mass fluxes result in less transport of \(^{222}\text{Rn}\) out of the boundary layer, where it has been shown that the use of ERA-Interim mass fluxes introduces inconclusive improvements in surface \(^{222}\text{Rn}\) distributions (Koffi et al., 2016). For lightning NOx there is a vertical re-distribution in the emission flux, with enhanced NOx injected into the upper troposphere when using the ERA-Interim mass fluxes.

Although the impact of resolution on daily photolysis rates may be appreciable, analysing global monthly mean \( J_{\text{O}_3} \) and \( J_{\text{NO}_2} \) surface values over a range of conditions shows that effects are limited to \( \sim 2\% \) and \( \sim 5–10\% \), respectively. One contributing factor to this rather muted impact is that the changes in surface albedo that occur at \( 1^\circ \times 1^\circ \) are largest at the poles during winter, which has no impact on photo-chemistry due to the absence of photolysing light (not shown). For cloud cover, a dominant term for determining total optical depth, there are significant increases at \( 1^\circ \times 1^\circ \) over the oceans, although this is generally related to instances of low photochemical activity. Examining the resulting changes in \( J_{\text{O}_3} \) and \( J_{\text{NO}_2} \) which occur throughout the tropospheric column reveals that significant differences of > 10\% can occur at the top of the BL at tropical locations. Such modest changes associated with this dominant loss term result in the change in the integrated chemical budget terms for tropospheric O\(_3\) and NO\(_2\) being rather low.

Analysing the chemical budget terms for tropospheric O\(_3\) shows (i) a reduction in the stratosphere–troposphere exchange flux of \( \sim 7\% \) to 597 Tg O\(_3\) yr\(^{-1}\), (ii) a repartitioning of the contribution from stratospheric downwelling in both the Northern and Southern hemispheres, (iii) no significant change in the tropospheric burden of O\(_3\) and (iv) modest changes in the integrated chemical production and destruction terms. Comparing simulated mixing ratios against surface measurements in Europe shows that the positive bias present in TM5 decreases by \( \sim 20\% \) at \( 1^\circ \times 1^\circ \) between 2 and 5 ppb month\(^{-1}\). This positive bias persists throughout the vertical column across diverse global regions regardless of

**Figure 12.** Comparison of weekly [SO\(_2\)] (µg m\(^{-3}\)) at 13:00 from both the \( 3^\circ \times 2^\circ \) and \( 1^\circ \times 1^\circ \) simulations at four 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).
the local NO\textsubscript{3} emissions, although the vertical gradient in tropospheric O\textsubscript{3} through the tropospheric column is captured quite well.

For NO and NO\textsubscript{2} increasing horizontal resolution results in only modest differences in the zonal mean recycling terms and the loss of O\textsubscript{3} by chemical titration. Comparisons against surface measurements in Europe show that there is a consistent negative bias in weekly [NO] of a few µg m\textsuperscript{-3} associated with both overly high surface O\textsubscript{3} (enhanced NO titration) and the inaccuracy of the NO\textsubscript{3} emission inventories. For NO\textsubscript{2}, the biases in the weekly concentrations are larger and can be both positive and negative. Increasing horizontal resolution has little effect on reducing the NO biases, but results in improvements for NO\textsubscript{2} at ~35 % of the available sites, with ~45 % of sites showing limited changes. Examining correlation coefficients shows that although there is typically a higher correlation at 1° × 1°, many sites still exhibit very low correlation or anti-correlation for some seasons. For the tropospheric column the improvement in the comparisons is only by a few percent, with a significant underestimation in both NO and NO\textsubscript{2} throughout the tropospheric column. Analysing the NO / NO\textsubscript{2} ratio and comparing against observations show that although partitioning is captured in the BL, there is a significant overestimation in the upper troposphere.

Finally, for CH\textsubscript{2}O and SO\textsubscript{2}, which can also be retrieved from satellite measurements, the effect of increased resolution is rather modest due to compensating changes to the chemical budget terms. When compared against observations there is a persistent low bias for tropospheric CH\textsubscript{2}O due to missing production terms, especially in the free troposphere. SO\textsubscript{2} comparison with surface observations in Europe shows 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 underestimation in the tropospheric column likely associated with either missing precursors or an underestimation in the direct emission terms.

Future updates to TM5-MP will most likely focus on developing an online secondary organic aerosol scheme, tropospheric halogen chemistry and incorporating an updated isoprene oxidation scheme involving more intermediate species. It will also be applied in the context of an Earth system model (EC-EARTH) for allowing future studies concerning chemistry–climate feedbacks. When computing resources allow, more expensive simulations can be performed using the 60 vertical levels as defined in the ERA-Interim meteorological dataset, approximately doubling the resolution of the simulations presented here. An additional update to improve the STE would be to apply the second-order moments scheme (Prather, 1986), whose application has been shown to capture the seasonality and magnitude of STE exchange to a better degree (Bönisch et al., 2008). In terms of oxidative capacity, one means of reducing the tropospheric near-surface O\textsubscript{3} mixing ratios would be to improve loss to land surfaces (Hardacre et al., 2010), although mixing ratios have been shown to be insensitive to the additional loss term to oceans, which is currently missing from many CTMs (Ganzeveld et al., 2009). Our comparisons of CH\textsubscript{2}O and SO\textsubscript{2} show that there is a significant uncertainty of chemical processes that affect distributions in the pristine marine environment. For instance, the physical process of deposition seems to be under-represented, possibly due to an overly low surface area of the surface, i.e. the lack of a flat surface. The significant underestimates in SO\textsubscript{2} suggest missing biogenic source terms; therefore, more understanding of biogenic emission terms is necessary.

11 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 sager@knmi.nl. 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 re-
The Supplement related to this article is available online at doi:10.5194/gmd-10-721-2017-supplement.

Competing interests. The authors declare that they have no conflict of interest.

Acknowledgements. This research has been supported by FP7 project Quality Assurance for Essential Climate Variables (QA4ECV), no. 607405. We thank M. van Weele for processing the overhead $O_3$ field and T. P. C. van Noije for updating the $SO_x$ emission estimates. We thank V. Huijnen for providing estimates on the heterogeneous uptake coefficients.

Edited by: F. O’Connor
Reviewed by: two anonymous referees

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