Ozone Impacts of Gas-Aerosol Uptake in Global Chemistry Transport Models

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Abstract.

The impact of six heterogeneous gas-aerosol uptake reactions on tropospheric ozone and nitrogen species was studied using two chemical transport models, EMEP MSC-W and ECHAM-HAMMOZ. Species undergoing heterogeneous reactions in both models include N_2O_5 , NO_3 , NO_2 , O_3 , HNO_3 and HO_2 . Since heterogeneous reactions take place at the aerosol surface area,

5 the modeled surface area density S_a of both models was compared to a satellite product retrieving the surface area. This comparison shows a good agreement in global pattern and especially the capability of both models to capture the extreme aerosol loadings in East Asia.

The impact of the heterogeneous reactions was evaluated by the simulation of a reference run containing all heterogeneous reactions and several sensitivity runs. One reaction was turned off in each sensitivity run to compare it with the reference

- 10 run. The analysis of the sensitivity runs confirms that the globally most important heterogeneous reaction is the one of N_2O_5 . Nevertheless, NO_2 , HNO_3 and HO_2 heterogeneous reaction gain relevance particularly in East Asia due to the presence of high NO_x concentrations and high S_a in the same region. although ECHAM-HAMMOZ showed much stronger responses than EMEP in this respect. The heterogeneous reaction of O_3 itself on dust is of minor relevance compared to the other heterogeneous reactions. The impacts of the N_2O_5 reactions show strong seasonal variations, with biggest impacts on O_3
- 15 in spring time when photochemical reactions are active and N_2O_5 levels still high. Evaluation of the models with northern hemispheric ozone surface observations yields a better agreement of the models with observations in terms of concentration levels, variability, and temporal correlations at most sites when the heterogeneous reactions are incorporated. Our results are loosely consistent with results from earlier studies, although the magnitude of changes induced by N_2O_5 reaction is at the low end of estimates, which seems to fit a trend whereby the more recent the study the lower the impacts of these reactions.

20 1 Introduction

Nitrogen species, ozone and atmospheric aerosols are major pollutants in the atmosphere, having strong impacts on ecosystems and human health, and also interacting with climate (Ainsworth et al., 2012; Harrison and Yin, 2000; Simpson et al., 2014;

IPCC, 2013). In regions, where gas phase and aerosol pollutants meet, heterogeneous chemistry can play a significant role (Jacob, 2000). The first heterogeneous process to become prominent in atmospheric chemistry was the heterogeneous destruction of stratospheric ozone on polar stratospheric clouds (Solomon et al., 1986; Solomon, 1999). However, heterogeneous processes are also relevant in the lower atmosphere, influencing tropospheric ozone and therefore oxidation capacity of the atmosphere

- 5 (Ravishankara, 1997; Pöschl, 2005; Seinfeld and Pandis, 2012). An important example is the heterogeneous reaction of N_2O_5 on aerosols, which is known to impact the NO_x - O_3 cycle while mainly removing NO_x from the troposphere (Mozurkewich and Calvert, 1988; Dentener and Crutzen, 1993; Evans and Jacob, 2005; Chang et al., 2011; Brown and Stutz, 2012), which can lead to ozone reduction (Macintyre and Evans, 2010). Other oxidised nitrogen species also undergo heterogeneous reactions on different aerosol types. NO_2 , HNO_3 and NO_3 react on wet surfaces of different aerosol types and increase aerosol nitrate
- 10 content (Rudich et al., 1998; Goodman et al., 1999). HNO₃ reacts also with dust and sea salt particles which is again a sink for NO_x and a source for particulate nitrate (Davies and Cox, 1998; Hanisch and Crowley, 2001). Moreover, heterogeneous reaction of NO₂ produces HONO which plays the role of a reservoir specie for NO and OH production (Platt et al., 1980). Other species also undergo heterogeneous reactions. O₃ reacts on dust particles and this has been estimated to lead to an ozone loss of about 20% in dusty regions (Usher et al., 2003). HO₂ reacts on wet particles leading to H₂O₂ production (Thornton)
- 15 and Abbatt, 2005). Furthermore, heterogeneous reactions lead to halogen release from sea salt aerosols (Frenzel et al., 1998; Yang et al., 2008; Lowe et al., 2011). Many modeling studies have been conducted over the years on these processes, but usually heterogeneous reactions were studied individually, and typically considering annual global budgets rather than detailed temporal or spatial resolution of the impacts (Dentener and Crutzen, 1993; Rudich et al., 1998; Saathoff et al., 2001; Bauer et al., 2004; Hodzic et al., 2006; Thornton et al., 2008; Chang et al., 2011).
- This paper presents estimates of the global impact of heterogeneous reactions of N_2O_5 , NO_3 , NO_2 , HNO_3 , HO_2 and O_3 and evaluates each reaction in a systematic way. The influence of each reaction on the magnitude and spatial and temporal variation in surface ozone is illustrated. The greatest impacts are seen in northern hemispheric regions of North America, Europe, South and East Asia. The N_2O_5 reaction is shown to significantly affect the spring-peak of surface O_3 at sites in all these regions. Although the impact of N_2O_5 reaction on O_3 is analysed, due to technical limitations in both models no ClNO₂ chemistry is
- 25 included, which could decrease the impact of N_2O_5 on O_3 , since it is a competing NO_x loss process. Different limitations in both model formulations are imposed here, as the lack of halogen chemistry and no possibility to transfer chlorine from the particle phase in the gas phase.

Sect. 2 presents the two global scale chemical transport models, EMEP MSC-W and ECHAM-HAMMOZ, as well as details of the reaction parameterisations and sensitivity tests. In Sect. 3 a short review of the range of reaction probabilities for each

- 30 heterogeneous reaction is given. Model setups and sensitivity runs are described in Sect. 4. Sect. 5 first presents a comparison of the simulated surface area from the models with satellite derived product, since the surface area of aerosols is crucial for heterogeneous chemistry. Especially in polluted regions where high trace gas concentrations meet large surface areas provided by aerosols heterogeneous chemistry might be of significant importance explaining aerosol composition and trace gas mixing ratios (Jacob, 2000; Pathak et al., 2009). Sect. 5 also presents the results of the sensitivity tests, and comparisons of daily
- 35 maximum ozone time-series for 2012 with surface station observations for selected sites. In Sect. 5.4 we discuss these results

compared to previous studies, as well as commenting on a number of open questions concerning heterogeneous reactions. Finally, Sect. 6 summarizes the results and implications for atmospheric chemistry.

2 Model description

Two models, the chemical transport model EMEP MSC-W (v4.16) (Simpson et al., 2012, 2017) and the global chemistry
aerosol climate model ECHAM6.3-HAM2.3MOZ1.0 (Schultz et al., in preparation) were used to study the heterogeneous chemistry of various compounds in the atmosphere.

2.1 EMEP

The basis of the EMEP MSC-W chemical transport model has been described in detail by Simpson et al. (2012), but substantial updates have been made in the treatment of aerosols, biogenic emissions and chemistry in recent years. Simpson et al. (2015,

2017) have documented the main changes in aerosol surface area and biogenic VOC emissions as discussed below, up to version rv4.15. The model version used in this report, rv4.16, is the latest version of the model. The main changes in rv4.16 have been the inclusion of the dry and wet deposition for N₂O₅, using the same rates as for HNO₃, and the implementation of an improved radiation scheme, based upon Weiss and Norman (1985). These changes have not affected basic model performance very much compared to previous publications, but of course the concentrations of N₂O₅ and its impact on ozone are reduced
 somewhat compared to earlier model versions.

The default model setup includes 20 vertical layers up to 100 hPa, using terrain-following coordinates, and the lowest layer has a thickness of about 90 m. Although originally designed for European applications (previously using a grid of resolution 50 km, more recently 28 km), the model is very flexible and is now applied on scales ranging from global (Jonson et al., 2010) to local (1-7 km grids), e.g. Vieno et al. (2010, 2014), Schaap et al. (2015). Anthropogenic emissions from land-based sources

20 are here taken from the so-called PANHAM database from the EU PANDA project (http://panda-project.eu), which combined emissions from the global HTAP data base (http://edgar.jrc.ec.europa.eu/htap_v2/index.php?SECURE=123) with the MEIC database for China (http://www.meicmodel.org/).

Emissions of VOC from biogenic sources are calculated in the model based upon land-cover and meteorological conditions. Emission factors for earlier versions of the EMEP model were mainly intended for European simulations (Simpson et al., 1999,

- 25 2012), but during 2016-2017 the factors used in non-European areas were substantially revised see Simpson et al. (2017) for details. For details of other emissions (soil-NO, lightning, aircraft, biomass-burning), see Simpson et al. (2012). For the present study meteorological data from the European Centre for Medium Range Weather Forecasting Integrated Forecasting System (ECMWF-IFS) model (http://www.ecmwf.int/research/ifsdocs/) were used, and the model runs with 1 × 1° latitude-longitude resolution.
- The chemical scheme in the EMEP MSC-W model, denoted 'EmChem16', consists of a standard gas-phase mechanism (132 species, 183 reactions, a recent update of the earlier EmChem03 evaluated by Andersson-Sköld and Simpson 1999), extended with organic aerosols using a volatility-basis-set scheme (Bergström et al., 2012; Simpson et al., 2012), plus sea-salt

(Tsyro et al., 2011) and dust aerosol. Unlike ECHAM-HAMMOZ, the EMEP model includes NH_3 and handles the resulting interactions with sulphate, HNO_3 and ammonium-nitrate through the use of the MARS equilibrium solver (Binkowski and Shankar, 1995). Unfortunately, interactions with sea-salt have not yet been implemented in EMEP-MARS. The chemical equations are solved using the TWOSTEP algorithm (Verwer and Simpson, 1995; Verwer et al., 1996).

- The EMEP MSC-W model has been extensively compared with measurements of many different compounds with generally good performance (e.g. Simpson et al., 2006a, b; Fagerli and Aas, 2008; Aas et al., 2012; Gauss et al., 2011), although most of these studies have focused on Europe. Still, in comparisons with global data and other models, the EMEP MSC-W model seems to perform well, especially more recent versions (Jonson et al., 2010, 2015; Angelbratt et al., 2011; Bian et al., 2017).
- As of EMEP MSC-W model version rv4.7 (Simpson et al., 2015), aerosol surface area (S_a) is estimated using the em-10 pirical relations of Gerber (1985), which simply requires aerosol mass concentrations and assumed aerosol density and sizeparameters. Values of S_a are calculated for fine and coarse particulate matter (PM_f, PM_c) both as totals (including all components, for reaction R1-3,5 in Table 1), and separately for coarse sea-salt and dust particles - which we denote as S_{SS} and S_{du} respectively. The distinction between total area S_a and S_{SS} and S_{du} was made to allow the use of Gerber's specific param-

eterizations for sea-salt and dust for reactions R5 and R6 (Table 1), with the assumption that where concentrations are large

15 (eg over oceans, deserts) these give a better estimate of S_a than the rural parameterisation would give. Further, for S_{du} the aerosol is assumed to be dry; which is not always true but is intended to reflect the nature of desert dust dominated aerosol. The EMEP model does not include fine-mode formation of NO_3^- through reaction R4, since the relationship between HNO₃ and fine-mode nitrate is given by the thermodynamic equilibrium solver MARS.

2.2 ECHAM-HAMMOZ

- 20 ECHAM-HAMMOZ is an aerosol chemistry climate model capable of performing interactive aerosol chemistry simulations. For this study simulations were done using version ECHAM6.3-HAM2.3MOZ1.0 (https://redmine.hammoz.ethz.ch/projects/ hammoz/wiki/Echam630-ham23-moz10). The model system ECHAM-HAMMOZ consists of the general circulation model ECHAM6.3 (Stevens et al., 2013), the aerosol model HAM2.3 (Neubauer et al., in preparation; Zhang et al., 2012) and the chemistry model MOZ1.0 (Schultz et al., in preparation). ECHAM calculates meteorological variables, cloud processes and
- 25 radiative transfer considering greenhouse gases and aerosols. The simulations in this study use hybrid sigma coordinates with 47 vertical layers, while the surface layer thickness is about 50 m. The horizontal resolution T63 leads to an associated 1.875°× 1.875 °Gaussian grid.

HAM simulates the evolution of aerosols considering aerosol and aerosol precursor emissions, microphysical processes as nucleation, coagulation, accumulation, sedimentation, dry and wet deposition. Via direct and indirect aerosol effects a feedback to climate system is simulated (Neubauer et al., in preparation). The aerosols in HAM are assumed to be internally mixed and

30 to climate system is simulated (Neubauer et al., in preparation). The aerosols in HAM are assumed to be internally mixed and consist of up to 5 components: sulphate, sea salt, dust, organic carbon and black carbon. To describe the aerosol number the microphysical driver M7 uses distribution seven log normal functions describing four wet aerosol modes and three dry aerosol modes. Hence, the wet functions cover nucleation, Aitken, accumulation, and coarse modes and the dry functions do not cover the nucleation mode. The height and median radius of the distribution are calculated, just its width is fixed. Due to aerosol

aging it is possible for insoluble particles to become soluble (Vignati et al., 2004). Dust and sea salt emissions are interactively calculated considering the wind speed at 10 m. Dimethylsulphate emissions are parametrized and emissions of sulphate dioxide, sulphate aerosol, black carbon and organic carbon are taken from the Representative Concentration Pathway (RCP) 8.5 emissions (Van Vuuren et al., 2011). Finally, optical properties of the aerosol are calculated and impact the atmospheric

5 circulation in ECHAM (Zhang et al., 2012).

Atmospheric chemistry is simulated by MOZ which is based on MOZART3.5 (Model for Ozone and Related chemical Tracers version 3.5) (Stein et al., 2012) connecting tropospheric chemistry of MOZART4 (Emmons et al., 2010) and stratospheric chemistry of MOZART3 (Kinnison et al., 2007). Further development since Stein et al. (2012) lead to MOZ being a chemical mechanism resembling to CAM-chem (Community Atmosphere Model with Chemistry) (Lamarque et al., 2012) with sev-

- 10 eral revisions, extended chemistry of aromatic compounds and a more detailed isoprene chemistry based on Taraborrelli et al. (2009). The version MOZ1.0 used here consists of 242 tracers, 733 chemical reactions which contain 142 photolysis reactions, 6 heterogeneous tropospheric reactions and 16 stratospheric heterogeneous reactions. Further, MOZ calculates dry and wet deposition of gases. Anthropogenic emissions are taken from the emission inventory RCP 8.5 (Van Vuuren et al., 2011). Biogenic emissions of VOC and NO₂ are calculated interactively by MEGAN (Model of Emissions of Gases and Aerosols from Nature)
- 15 (Guenther et al., 2006; Henrot et al., 2017). NO lightning emission are parametrized as described by Grewe et al. (2001). HAM and MOZ interact via two physical processes. First, assuming spherical aerosols, the surface area density for heterogeneous reactions is calculated using aerosol distribution and median radius. Second, MOZ provides fields of oxidants for aerosol formation from gas-phase precursors. The HAMMOZ coupling does not include ammonium nitrate formation due to the lack of nitrate aerosol in the current HAM version. Therefore, reactive uptake of nitric acid leads to a total loss, based on the assumption of a quick loss of gas phase HNO₃ and particulate nitrate. To underline, heterogeneous reactions in ECHAM-HAMMOZ do not form HNO₃ in the gas phase, but introduce a direct loss to the products HNO₃ and NO₃⁻.

3 Heterogeneous reactive uptake

Experimental studies show that oxidised nitrogen species, ozone and the hydroperoxy radical undergo heterogeneous reactions on wet and dry aerosols. Heterogeneous reactions can be modeled as a pseudo-first order process (Ammann et al., 2013).

$$25 \quad \frac{d[\mathbf{X}]_g}{dt} = -k_{\mathbf{X}}[\mathbf{X}]_g \tag{1}$$

The change in gas phase concentration of the species $X = N_2O_5$, NO_3 , NO_2 , HNO_3 , HO_2 , O_3 is proportional to its gas phase concentration $[X]_g$ and a reaction rate coefficient k_X (Schwartz, 1986)

$$k_{\rm X} = \left(\frac{r_p}{D_g} + \frac{4}{c_{\rm X} \cdot \gamma_{\rm X}}\right)^{-1} S_a \tag{2}$$

where D_g represents the gas phase diffusion coefficient, r_p is the particle radius, c_X is the mean molecular velocity of the 30 species X, γ_X represents the reaction probability and S_a the surface area density. The γ_X values are generally determined from laboratory measurements. The first term in Eqn. 2 is very small for particles of accumulation mode and larger, and is neglected in the EMEP model. The main challenges for chemistry transport models are the calculation of a proper surface area density S_a and the parametrization of the reaction probability γ_X .

First, Table 1 summarizes the heterogeneous reactions investigated in this study. Second, sections 3.1-3.6 discuss literature values of γ associated with each reaction. An overview of the parametrization and values used for the different reaction probabilities is given in Table 2. ECHAM-HAMMOZ and EMEP MSC-W use the same reaction probabilities or functions for many reaction, with the most important difference being the lack of ammonium nitrate aerosol in ECHAM-HAMMOZ. Lastly, to check if the surface area density is realistic, simulated S_a is compared to a satellite-model-product in section 5.1.

3.1 N₂O₅

- 10 N_2O_5 reaction probability depends on aerosol water content and aerosol composition. Therefore, several laboratory studies measured γ values on different aerosol types leading to the possibility to derive detailed parametrizations (Riemer et al., 2003b, 2009; Evans and Jacob, 2005; Liao and Seinfeld, 2005; Davis et al., 2008; Bertram and Thornton, 2009; Griffiths et al., 2009; Brown and Stutz, 2012). For dry sulphate aerosol, reaction probabilities range between 10^{-4} and 10^{-3} ; for wet aerosol γ ranges between 10^{-3} and 8.6×10^{-2} depending on relative humidity. The N_2O_5 heterogeneous reaction humidity dependence
- 15 also explains the range of reaction probabilities of sea salt aerosol of 6×10^{-3} to 4×10^{-2} . On nitrate containing aerosol lower reaction probabilities were found due to nitrate effect (Wahner et al., 1998), between 3×10^{-4} and 3×10^{-3} (Chang et al. 2011, and references therein). Moreover, N₂O₅ can react on organic aerosol under dry conditions with low reaction probabilities in the order of 10^{-6} and 10^{-5} (Gross et al., 2009). This value increases to $10^{-4} - 10^{-3}$ under wet conditions, because the higher water content allows N₂O₅ to hydrolyze (Thornton et al., 2003). Even dust aerosols can be covered by a layer of water
- 20 leading to a reaction probability between 3×10^{-3} at 30 % and 2×10^{-2} at 70% relative humidity (Bauer et al., 2004). For N₂O₅ reaction on black carbon, Sander et al. (2006) reported a wide range of reaction probabilities, between $2 \times 10^{-2} 10^{-6}$.

Most studies have used laboratory data to estimate γ values, but some have made use of ambient data. Brown et al. (2009) used aircraft measurements over Texas, and found observation-based γ values of ca. $5 \times 10^{-4} - 6 \times 10^{-3}$, usually substantially lower (often a factor of 10) than values calculated using laboratory-based values. Using aircraft measurements around the

- 25 United Kingdom, Morgan et al. (2015) found rather high γ values for N₂O₅, from ca. $1 \times 10^{-2} 3 \times 10^{-2}$, with strong dependencies on sulphate, and a clear suppression of γ due to nitrate. They concluded that including the suppressive effect of organic aerosol in the uptake parameterisation leads to significant underprediction of the γ values. Further, direct N₂O₅ measurements retrieved a highly daily variation of $\gamma_{N_2O_5}$ also explained by the nitrate effect leading to a mean value of 5.4×10^{-3} ranging from $3 \cdot 10^{-5}$ to $2.9 \cdot 10^{-2}$ (Riedel et al., 2012). In Stone et al. (2014) and Wagner et al. (2013) in situ measurements of N₂O₅
- 30 were used to retrieve the reaction probability within the framework of a box model. In Stone et al. (2014) $\gamma_{N_2O_5}$ is varied over a range of values between 0 and 1, and found that values of $2 \cdot 10^{-1} - 2 \cdot 10^{-2}$ agreed best with observations. Wagner et al. (2013) retrieved the reaction probability of N₂O₅ using a box model driven by ambient wintertime observations. The reaction probability distribution ranges between $2 \cdot 10^{-3}$ and $1 \cdot 10^{-1}$, displaying a maximum at $2 \cdot 10^{-2}$.

It is clear from the studies mentioned above that great uncertainties surround both the magnitude and the chemical dependence of γ values for N₂O₅. Even thorough evaluations such as those of Davis et al. (2008) or Chang et al. (2011) have little consideration of important components of the aerosol such as organic matter, and even such schemes seem to be inconsistent with the aircraft-observations discussed above. For our modelling studies, we have not tried to develop or use yet another

5 scheme, but rather to make use of the γ schemes already implemented in each model, with some small efforts at harmonisation to build similar reference schemes.

The equations used for EMEP MSC-W and ECHAM-HAMMOZ can be found in Table 2. Both models make extensive use of the parameterizations developed by Evans and Jacob (2005), with the largest difference being that EMEP includes ammonium nitrate (in fine particles) among the nitrate species. For N_2O_5 the uptake coefficients for sulphate, sea salt and

10 organic aerosol are identical in the two models. For the reaction on dust, both models rely on Bauer et al. (2004) which was interpreted differently by Evans and Jacob (2005) and Liao and Seinfeld (2005). This small difference in the uptake coefficient formulation on dust does not lead to large differences in the resulting uptake coefficient.

EMEP MSC-W also modifies the γ value for secondary inorganic aerosol to account for a nitrate inhibition effect (Wahner et al., 1998; Riemer et al., 2003a). This makes use of the γ_{AN} factor presented in Davis et al. (2008) for ammonium nitrate, and merged with the sulphate factor in a manner reminiscent of Riemer et al. (2003b). First a sulphate mass fraction within the

secondary inorganic aerosol (SIA) is calculated, $f_{SU} = m_{SO_4}/(m_{SO_4} + m_{NO_3})$, then γ_{SIA} is calculated as given in Table 2.

Figure 1 illustrates the γ values for sulphate aerosol from Evans and Jacob (2005) as a function of relative humidity (RH) and temperature for sulphate, and the RH dependency of γ for nitrate from the Davis et al. (2008) formulation. The negative temperature dependence after 280K can be explained by increasing volatility with increasing temperature leading to less uptake on the aerosol. As described before, reaction probability increases with increasing water content in the aerosol due to enhanced

 N_2O_5 hydrolysis. Even at high RH, reaction probability on nitrate containing aerosol is not as high as in sulphate aerosol. Nevertheless, the very high γ values found at high RH seem questionable, because the aerosol itself becomes saturated at high RH and these small water content changes should not have such a huge impact on the heterogeneous reaction.

No further parameterization considering organic coatings is used in either EMEP MSC-W or ECHAM-HAMMOZ due to 25 the large uncertainties in this effect (e.g. Brown et al., 2009; Morgan et al., 2015), and the fact that ambient OM and its 25 thermodynamic properties are so poorly understood (Hallquist et al., 2009). Further, sensitivity runs done with ECHAM-HAMMOZ have shown minor global impact of organic coatings (Stadtler, 2015). Figure S1 in the Supplement illustrates the $\gamma_{N_2O_5}$ values from the two models using the setups described in Sect. 4, showing values of around 0.01 - 0.04 over much of the globe.

30 3.2 NO₃

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Hydrolysis of the nitrate radical NO₃ happens on various aerosol types depending on the water content. NO₃ heterogeneous reaction produces HNO₃ and OH in the aqueous particle phase and can be counted as a NO_x sink (Rudich et al., 1998). Several laboratory studies shown γ ranging between 10^{-4} and 10^{-3} (Rudich et al., 1996; Rudich et al., 2002). Jacob (2000)

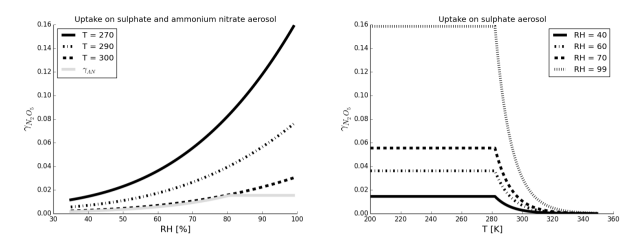


Figure 1. Upper plot: N_2O_5 reaction probability dependence on relative humidity for sulphate (black) and nitrate (grey) aerosol. For sulfate aerosol three temperatures are shown. Lower plot: N_2O_5 reaction probability temperature dependence on sulphate aerosol for four relative humidity. Parameterizations from Evans and Jacob (2005) and Davis et al. (2008), see text.

recommended to use $\gamma = 10^{-3}$ for atmospheric chemistry model simulations, and this value was adopted for EMEP and ECHAM-HAMMOZ.

The nitrate radical NO_3 undergoes hydrolysis in wet aerosols, but was also observed to react with organic compounds on the aerosol surface. Hydrolysis of nitrate radicals NO_3 happens on various aerossol types depending on the water content. NO_3 heterogeneous reaction produces HNO_3 and OH in the aqueous particle phase and can be counted as a NO_x sink (Rudich et al.,

5 heterogeneous reaction produces HNO₃ and OH in the aqueous particle phase and can be counted as a NO_x sink (Rudich et al., 1998). Several laboratory studies shown γ ranging between 10^{-4} and 10^{-3} (Rudich et al., 1996; Moise et al., 2002). Jacob (2000) recommended to use $\gamma = 10^{-3}$ for atmospheric chemistry model simulations.

Reactions with different organic compounds were explored in laboratory experiments. Gross and Bertram (2008) measured the reaction probabilities between 0.059 and 0.79 of NO₃ with different polycyclic aromatic hydrocarbons leading to NO₂
and HNO₃ formation. Two following studies also found high reaction probabilities of NO₃ with alkenoic acid (>0.07) (Gross et al., 2009) and alkene monolayers (0.034) (Gross and Bertram, 2009). Organic coatings could enhance NO₃ reactive uptake, nevertheless knowledge of explicit organic compounds in the organic fraction of aerosol is unknown in both mode systems, therefore the recommended of γ = 10⁻³ for NO₃ hydrolysis value was adopted for EMEP and ECHAM-HAMMOZ.

3.3 NO₂

15 NO₂ heterogeneous reaction leads to the production of HNO₃ and HONO. Especially in humid environments, the heterogeneous reaction may account for up to 95% of HONO production (Goodman et al., 1999). During nighttime HONO can accumulate in the atmosphere and therefore be an efficient OH radical source during the morning when sun rise starts photolysis (Goodman et al., 1999). Estimates of γ for NO₂ vary widely, however, with several laboratory studies giving a range between 10^{-8} and 10^{-3} (Harrison and Collins, 1998; Kleffmann et al., 1998; Arens et al., 2001; Underwood et al., 2001). Jacob (2000) recommended $\gamma = 10^{-4}$ and this value is used for this study.

3.4 HNO₃

Nitric acid reacts on sea salt and dust aerosol surfaces, producing nitrate which stays in the aerosol phase (Davies and Cox,

- 5 1998; Hodzic et al., 2006). Experimentally derived γ values for HNO₃ on sea salt range between 10^{-4} and 10^{-2} (Davies and Cox 1998, and references therein). A relative humidity dependent uptake coefficient was proposed in Hauglustaine et al. (2014) increasing γ from 10^{-3} and 10^{-1} to cover low and high relative humidity. No such relative humidity dependence was used in this study, because for the conditions in the marine boundary layer, the value of 10^{-2} fits well and is used here.
- Heterogeneous reaction of HNO₃ on dust was studied on different types of minerals, atmospheric dust types and for a range of relative humidities giving γ in the range of 10^{-6} and 10^{-1} (Hanisch and Crowley, 2001; Usher et al., 2003; Liu et al., 2008; Hauglustaine et al., 2014). Although in Fairlie et al. (2010) a relative humidity dependence of varying γ between 10^{-5} and 10^{-3} is described, the value is used here is based on Hodzic et al. (2006), who tested γ values between 10^{-6} and 0.3, deriving 0.1 as the best γ value minimizing the model error compared to observations. Compared to the other referenced studies, this is an upper limit.

15 3.5 O₃

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Studies of the heterogeneous reaction of ozone on dust give a wide range for possible reaction probabilities, from 10^{-10} to 10^{-4} (Reus et al., 2000; Usher et al., 2003; Mogili et al., 2006; George et al., 2015). Reus et al. (2000) gives 10^{-4} as an upper limit for this reaction probability, but George et al. (2015) suggests a more conservative upper limit value of 10^{-5} . Nevertheless, Nicolas et al. (2009) conclude that a reaction probability of 10^{-6} is a realistic number in terms of atmospheric environmental conditions, and this value was adopted here.

HO₂ reaction probability is highly variable and strongly depends on transition metal ions contained in the aerosol (Tilgner

3.6 HO₂

et al., 2005; Mao et al., 2013; George et al., 2013; Huijnen et al., 2014). Furthermore, this reaction can also take place on cloud droplets. Estimates for γ range between 0.02 and 1 (Jacob, 2000; Remorov et al., 2002; Thornton and Abbatt, 2005; Taketani

droplets. Estimates for γ range between 0.02 and 1 (Jacob, 2000; Remorov et al., 2002; Thornton and Abbatt, 2005; Taketani et al., 2008; George et al., 2013; Mao et al., 2013). Whalley et al. (2015) measured HO₂ in clouds and found a decrease in HO₂ concentrations up to 90%. Depending on the compounds in the particle aqueous phase heterogeneous reaction of HO₂ produces either H₂O₂ or H₂O. Consequently, this heterogeneous reaction can be a terminal radical sink or not (Mao et al., 2013; Whalley et al., 2015). Here we do not account for a terminal sink, but let the heterogeneous reaction of HO₂ produce H₂O₂ and use the γ recommended by Jacob (2000) of 0.2.

4 Setup of sensitivity runs

To test the six heterogeneous reactions (see Table 1), six sensitivity runs were designed and performed with both models. For EMEP MSC-W a spin up of six months and for ECHAM-HAMMOZ one of twelve months is used. Afterwards the results for the whole year 2012 are evaluated. The reference run REF contains all heterogeneous reactions with the parameterizations

5 given in Table 2. Each sensitivity run is done with five out of six heterogeneous reactions; the names of the runs show which compound does not undergo heterogeneous reaction. For example, in the noN2O5 run, only the N_2O_5 heterogeneous reaction is turned off. An overview of the simulations is given in Table 3.

5 Results and Discussion

5.1 Surface area density

- 10 Aerosols consist of a variety of compounds in the gas, liquid or solid phase, and the shapes of aerosols vary greatly (Pöschl, 2005). Large scale models can not explicitly treat the morphology of aerosols. In EMEP MSC-W and ECHAM-HAMMOZ distribution functions and median radii are used to simulate the aerosol population. Based on this approach surface area density S_a is calculated considering the aerosol distribution, the median radius and assuming spherical particles. This assumption is good for liquid aerosols behaving as small water droplets. For dry particles this assumption can lead to an underestimation of S_a due to folded or porous structures (Buseck and Posfai, 1999).
 - In van Donkelaar et al. (2015) satellite retrievals and the GEOS-Chem chemical transport model are used to derive global surface $PM_{2.5}$ estimates with a resolution of 10 km × 10 km in the time period between 1998 and 2012. The physical relation between AOD and surface area is described in the supplementary material of van Donkelaar et al. (2015).

Figure 2 shows the estimated $PM_{2.5}$ surface area by van Donkelaar et al. (2015) and the modeled surface area density S_a

- from EMEP MSC-W and ECHAM-HAMMOZ as ground-level annual mean 2012 over land. Although these data-sets are not strictly comparable, since the van Donkelaar et al. 2015 estimate in itself relies partly on various assumptions of a third chemical transport model, GEOS-Chem (van Donkelaar et al., 2015), the general patterns of the models agree well with the surface area density estimation. Both models capture the east west gradient in S_a over North America even if the total S_a value is comparably lower in both models. Similarly, Europe in the satellite GEOS-Chem product has slightly higher S_a values than
- 25 the models produce. In contrast, the S_a values over India are captured very well, and the peak values in East Asia are also produced by both models, while ECHAM-HAMMOZ simulates highest S_a values among the three data sets in East Asia. An overestimation of both models compared to satellite GEOS-Chem happens over North Africa. In South America EMEP MSC-W performs better than ECHAM-HAMMOZ due to larger contributions from secondary organic aerosol (SOA) formation. EMEP uses a more complex SOA scheme (Bergström et al., 2012; Simpson et al., 2012) which allows for oxidation ('aging')
- 30 of semivolatile organic vapours. In ECHAM-HAMMOZ an adjusted amount of organic material covering also SOA is emitted, but the amount does not close the gap leading to a lower S_a compared to EMEP MSC-W and satellite GEOS-Chem.

5.2 Impacts of sensitivity tests

reactions.

To evaluate the impact of our heterogeneous reactions, the six sensitivity runs were compared to the reference run containing all heterogeneous reactions. By turning off one heterogeneous reaction in each sensitivity run, the impact of each reaction can be estimated. Tables 4-5 show the differences in percent between the sensitivity runs and the reference run for EMEP MSC-

- 5 W and ECHAM-HAMMOZ, as averaged over regions of North America (NA), Europe (EUR), East Asia (EA) and South Asia (SA) (regions defined as in Fiore et al. 2009). Tables S1-S2 in Supplementary give absolute differences, in ppb or ppt. The main focus of this evaluation is on the effect of the heterogeneous reactions on ozone mixing ratios. (Also, in order to test the importance of year to year variability, the EMEP model was additionally run for the year 2011. The results, given in Supplementary Table S3, are almost identical to those shown for 2012 in Table 4 and so not discussed further here.)
- For the reference runs, EMEP MSC-W and ECHAM-HAMMOZ simulate very similar values for ozone, ECHAM-HAMMOZ giving somewhat lower mixing ratios. Also NO_x values are similar in Asia, but differ by ca. 60–100% in North America and Europe. In terms of other reactive nitrogen species, EMEP MSC-W has overall higher NO_y levels and especially PAN. This difference in NO_y availability is expected given the impact of EMEP's NH_3 emissions in the formation of ammonium nitrate, thus extending the lifetime of reactive nitrogen species, on top of general differences in emissions and chemical mechanisms.
- Tables 4–5 clearly show increases in O_3 from all the sensitivity runs when the heterogeneous reaction is turned off, except for the sensitivity run without heterogeneous NO_2 reaction in East Asia. In this region the special case of ozone titration (Wild and Akimoto, 2001) leads to an ozone loss due to NO_2 instead of production: lowering NO_2 in this region of very high NO_x regions means reducing a loss process. Even if the models agree on the direction of the impact of heterogeneous reaction on O_3 , they do not agree on the strength of the reactions.
- For both models the N₂O₅ reactions have generally (ECHAM-HAMMOZ) or always (EMEP) the biggest effect on O₃, with changes of ca. 2–3 ppb (5–9%). Some other heterogeneous reactions (especially NO₂, HO₂ and HNO₃) gain some significance in highly polluted areas where aerosol surface areas are high, but the two models show quite different responses though in their response to these other gas-aerosol reactions. The EMEP model actually shows rather small impacts of all reactions on O₃ except N₂O₅, except in East and South Asia where some impacts can approach 10-20% of that of N₂O₅.
 ECHAM-HAMMOZ, on the other hand, shows quite marked responses to especially the HNO₃ reactions, but also the HO₂

The strong response of O_3 in ECHAM-HAMMOZ to the HNO₃ reaction compared to EMEP seems to be the result of a number of factors. The simplest is that EMEP allows this reaction only on coarse aerosol, and so has a smaller surface area for this reaction, especially on dust. Another explanation is that the model sensitivities to NO_x changes may be different, possibly

30 caused by chemical differences or the different horizontal resolutions of the models. Ozone chemistry (and even the switch from production to loss) can be very sensitive to NO_x concentration levels, especially in unpolluted areas (Crutzen et al., 1999; Sillman et al., 1990). NO_x plumes from ships or power plants emitted into large model grid cells might well produce more O_3 in one model than the other, leading to different sensitivities to NO_x emissions (von Glasow et al., 2003; Vinken et al., 2011). The EMEP model has in fact a psuedo-species 'SHIPNOx' by which 50% of NO_x from ship plumes are given a pathway to HNO_3 production, skipping the intermediate NO_2 production associated with overestimating O_3 production from NO in pristine environments (Simpson et al., 2015). A further factor is the lack of nitrate aerosol in ECHAM-HAMMOZ. In the EMEP model HNO_3 can take part in ammonium nitrate aerosol (AN) formation, thus extending the lifetime of NO_y . Due to the AN, some HNO_3 can be recycled back into the atmosphere stabilizing the HNO_3 and NO_3 mixing ratios.

- Table S2 (and 5) shows that for ECHAM-HAMMOZ omitting the HNO₃ reaction on dust and sea salt aerosol increases NO_x by ca. 10-20 ppt (1 %), whereas in EMEP the change is tiny (\leq 1 ppt). The impact in ECHAM-HAMMOZ can be found over the whole globe, but especially over the oceans, where NO_x is low, but still much higher than NO_z (=NO₃+N₂O₅). Changes in NO_z with this noHNO3 scenario are far higher in ECHAM-HAMMOZ than in EMEP. Even if heterogeneous HNO₃ loss does not hugely impact NO_x, a small NO_x increase, even if really small, is ubiquitous and can shift the equilibrium between
- 10 ozone production and loss towards more production, reaching a higher steady state O_3 concentration. Also, this reaction has a significant effect on NO₃, reducing it in the northern oceans by about 10 % (not shown). NO₃ rapidly photolyses, and resulting NO₂ likewise, so has a high ozone-formation potential. NO₃ rapidly photolyses and produces NO₂ and atomic oxygen $O_3(^{3}P)$. NO₂ subsequently photolyses and results in NO and a second $O_3(^{3}P)$. From these two reactions two ozone molecules can be formed, therefore NO₃ has a high ozone-formation potential. Reducing HNO₃ and therefore NO₃ drastically by the
- 15 surface reaction in this highly sensitive region leads to a nonlinear response of the model changing the gross ozone production in ECHAM-HAMMOZ by 350 Tg, which is a reduction of 7 %. This leads to a global more or less uniformly distributed difference of 1 - 4 ppbv in ozone mixing ratios.

Analyzing all the possible differences in these two different models is beyond of the scope of this study, but it may well be that ECHAM-HAMMOZ overestimates the impact of HNO_3 due to missing nitrate aerosol formation and EMEP underestimates the impact, due to the use of only coarse sea salt and dust aerosol for the HNO_3 and HO_2 reactions.

As the N_2O_5 reactions have the greatest impact on tracer concentrations among our sensitivity tests, the spatial and temporal differences between the reference run and the sensitivity run noN2O5 have been investigated in more detail. Figs. 3 and 4 show the difference between the mixing ratios of O_3 and NO_x in the reference run and in the sensitivity run without the N_2O_5 reactions. Both models show the largest changes in regions where high aerosol loadings and high NO_x emissions can be found, such as Northeast America. Europe, South and East Asia

25 such as Northeast America, Europe, South and East Asia.

20

Converting N_2O_5 to HNO_3 on aerosol surfaces introduces an additional sink for NO_x , because HNO_3 is rapidly (in EMEP) or immediately (in ECHAM-HAMMOZ) lost via dry and wet deposition, and reactive uptake on aerosols, after it is produced. Therefore, NO_x mixing ratios are lowered in the reference run REF compared to the simulation without the heterogeneous reaction noN2O5, as can be seen in Figure 4.

30 For ozone, the differences propagate through the whole northern hemisphere due to the longer lifetime of O_3 compared to NO_x (Fig. 3). Again both models simulate similar patterns with regard to the spatial distribution of changes due to N_2O_5 . As described before, EMEP MSC-W shows a bigger impact on O_3 concentrations than ECHAM-HAMMOZ because of O_3 is already lowered due to the direct heterogeneous loss of HNO₃.

Especially for East Asia the impact of heterogeneous reactions cannot be neglected. High nitrate loadings in ammonium 35 poor regions verify the importance shown by other models (Pathak et al., 2008). In the southern hemisphere, N_2O_5 and the

other heterogeneous reactions evaluated in this study have much smaller impacts on ozone and NO_x than seen in the northern hemisphere (Figures 3, 4).

To explore the seasonal impact of N_2O_5 reactions, Fig. 5 shows monthly values for tracer mixing ratios and surface area density from both models for the different northern hemispheric regions. In general, the models produce comparable seasonal

- 5 cycles for the gas tracers and surface area density. Strongest changes in seasonal cycles are found in the noN2O5 run. and in ozone for both runs. In the noN2O5 run, N₂O₅ builds up during winter time, because it is thermally unstable and photolabile. Including the heterogeneous uptake leads to a strong N₂O₅ reduction in both models, yielding a flatter seasonal curve. The loss of N₂O₅ in going from noN2O5 to REF leads to a decrease in NO₂, NO₃ and PAN. Here models slightly differ. EMEP displays a stronger reduction in NO₃ and PAN, since it has in both runs higher mixing ratios compared to ECHAM-HAMMOZ.
- 10 Removing NO₂ from the system leads in both models to a reduction of ozone. Although, the impact of N_2O_5 heterogeneous reaction on NO₂ is higher in winter and lowest during summer, the greatest change in O₃ can be found during spring. This can be explained by the availability of N_2O_5 and ozone production strength. As stated before, N_2O_5 is formed during night time, therefore less sun is favorable. In contrast, to form ozone light is needed. Still high N_2O_5 concentrations, enough surface area and a sufficiently high ozone production can be found during spring, leading to the biggest change in O₃ production during this
- 15 season. During winter, nights are longer, the sun is less active, therefore heterogeneous chemistry is efficient, but less ozone production reduction occurs. During winter, nights are longer leading to inactive photochemistry. Therefore, heterogeneous chemistry is efficient. Nevertheless, a rather inactive photochemistry also leads to less ozone production. Comparing to spring, the impact seen here is lower because of already low ozone formation rate.

Table 1. Heterogeneous reactions in the EMEP MSC-W and ECHAM-HAMMOZ models. The second column specifies the aerosol type on which the reaction proceeds in the models:. SS: seasalt, DU: dust, PM: particulate matter

No.	Reaction		Aerosol type	Notes
R1	N_2O_5	$\rightarrow 2HNO_3$	РМ	1
R2	NO_3	$\rightarrow \mathrm{HNO}_3$	PM	2
R3	NO_2	$\rightarrow 1/2HNO_3 + 1/2HONO$	PM	2
R4	HNO_3	$ ightarrow \mathrm{NO}_3^-$	SS, DU	2,3
R5	HO_2	$ ightarrow 1/2H_2O_2$	PM	2
R6	O_3	$ ightarrow HO_2$	DU	

¹ Just for RH>40% in EMEP.

² Just on wet aerosol in ECHAM-HAMMOZ.

 3 Just on coarse mode dust and sea-salt in EMEP, using $S_{\rm SS}$ and/or $S_{
m du}$, see Sect. 2.1.

Table 2. Reaction probabilities for the different species. Unless explicitly labelled (in parentheses afer the equation), both models use the same formulation. Here, RH denotes relative humidity in range, $RH \in (0, 100)$, and fRH denotes fractional relative humidity in range, $fRH \in (0, 1)$.

Specie	γ	Reference		
N ₂ O ₅	$\gamma_{SS} = \begin{cases} 0.005, \text{RH} \le 62 \ \% \\ 0.03, \text{RH} \ge 62 \ \% \end{cases}$			
	$\gamma_{SU} = \alpha \cdot 10^{-\beta}$	EVA05		
	$\alpha = 2.79 \cdot 10^{-4} \text{fRH} + 1.3 \cdot 10^{-4} \text{fRH} - 3.43 \cdot 10^{-6} \text{fRH}^2 + 7.52 \cdot 10^{-8} \text{fRH}^3$			
	$\int 4 \cdot 10^{-2} (T - 294), T > 282 \text{ K}$			
	$\beta = \begin{cases} 4 \cdot 10^{-2} (\mathrm{T} - 294), T > 282 \mathrm{K} \\ -0.48, T \le 282 \mathrm{K} \end{cases}$			
	$\gamma_{DU} = 0.01 \text{ (EMEP)}$	EVA05		
	$\gamma_{DU} = 4.25 \cdot 10^{-4} \text{RH} - 9.75 \cdot 10^{-3} (30\% \le \text{RH} \le 70\%) \text{ (ECHAM)}$	LIA05		
	$\int 0.03, \text{RH} > 57\%$			
	$\gamma_{OC} = \begin{cases} 0.03, \text{RH} > 57\% \\ 5.2 \cdot 10^{-2}, \text{RH} \le 57\% \end{cases}$			
	$\gamma_{BC} = 0.005$	EVA05		
	$\gamma_{AN} = \min(0.0154, 1/(1 + \exp(8.10774 - 0.04902 \cdot \text{RH}))) \text{ (EMEP)}$	DAV08		
	$\gamma_{SIA} = f_{SU}\gamma_{SU} + (1 - f_{SU})\gamma_{AN} \text{ (EMEP)}$	SIM15		
	(where f_{SU} = mass fraction of sulfate in inorganic aerosol, see Sect. 3.1)			
NO_3	$\gamma = 0.001$	JAC00		
NO_2	$\gamma = 10^{-4}$	JAC00		
HNO ₃	$\gamma_{SS} = 0.01$	DAV98		
	$\gamma_{DU}=0.1$	HOD06		
HO_2	$\gamma = 0.2$	JAC00		
O ₃	$\gamma_{DU} = 10^{-6}$	NIC09		

The subscripts refer to the aerosol compounds as given in Table 1, plus OC:Organic carbon/Organic matter, SU: Sulphate,

SIA:secondary inorganic aerosol, BC black carbon.

Refs: DAV98 Davies and Cox (1998), DAV08 Davis et al. (2008), EVA05 Evans and Jacob (2005), JAC00 Jacob (2000), LIA05 Liao and Seinfeld (2005), THO08 Thornton et al. (2008), HOD06 Hodzic et al. (2006), NIC09 Nicolas et al. (2009) SIM15 Simpson et al. (2015)

Run	Description
REF	All heterogeneous reactions
noN2O5	All except N_2O_5 reaction
noNO3	All except NO ₃ reaction
noNO2	All except NO ₂ reaction
noHNO3	All except HNO3 reaction
noHO2	All except HO ₂ reaction
noO3	All except O_3 reaction

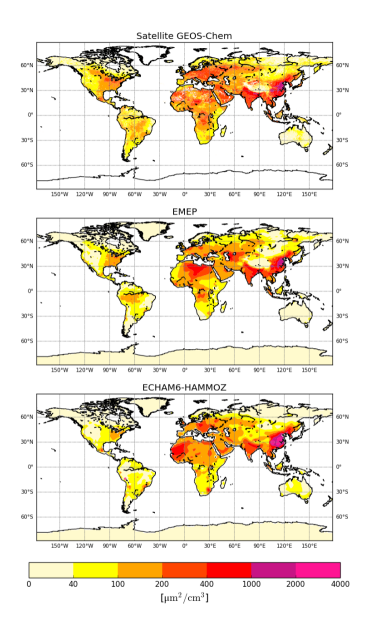


Figure 2. Satellite estimated (top) and simulated surface area densities by EMEP (middle) and ECHAM-HAMMOZ (bottom) at ground level. The Satellite data is an average value for the time period 2010 - 2012, from van Donkelaar et al. (2015). The model data is for 2012 and the lowest model level.

Table 4. Impacts of gas-aerosol reactions on regional ground level average mixing ratios of O_3 and key NO_y compounds: EMEP model, year 2012.

Region	Run	Unit	O ₃	NO _x	NOy	HNO ₃	PAN	N_2O_5	NO ₃
NA	REF	Conc*:	40.33	0.82	1.81	0.21	0.55	5.08	4.54
NA	noN2O5	%:	5	9	4	-10	8	160	59
NA	noHO2	%:	0	-1	0	1	1	-2	-10
NA	noHNO3	%:	0	0	-2	18	0	0	0
NA	noNO2	%:	0	2	1	-1	0	4	2
NA	noNO3	%:	0	0	0	0	0	1	3
NA	noO3	%:	0	0	0	0	0	0	0
EUR	REF	Conc*:	40.89	1.01	2.43	0.25	0.54	7.73	6.48
EUR	noN2O5	%:	7	16	3	-16	10	280	72
EUR	noHO2	%:	1	-3	0	1	4	-4	-14
EUR	noHNO3	%:	1	0	-6	58	0	0	3
EUR	noNO2	%:	0	5	1	-1	-1	6	4
EUR	noNO3	%:	0	0	0	0	0	2	10
EUR	noO3	%:	0	0	0	0	0	0	2
EA	REF	Conc*:	43.96	2.23	4.63	0.54	0.89	12.59	5.52
EA	noN2O5	%:	8	14	4	-19	13	278	106
EA	noHO2	%:	2	-4	0	2	7	0	-7
EA	noHNO3	%:	0	0	-2	13	0	0	0
EA	noNO2	%:	-1	30	9	-11	-8	13	4
EA	noNO3	%:	0	0	0	0	0	1	3
EA	noO3	%:	0	0	0	0	0	0	0
SA	REF	Conc*:	47.33	1.12	2.90	0.42	0.33	10.37	12.0
SA	noN2O5	%:	6	11	1	-4	15	139	63
SA	noHO2	%:	1	-3	0	1	5	-5	-12
SA	noHNO3	%:	1	0	-8	61	0	1	4
SA	noNO2	%:	1	4	1	0	1	10	5
SA	noNO3	%:	1	0	0	0	1	5	11
SA	noO3	%:	0	0	0	0	0	0	0

Notes: Base-case concentrations from the surface-level of the model are given in ppt for NO_3 and N_2O_5 , otherwise ppb (Conc^{*} flags this difference in units). Results for the sensitivity tests are given as (test-base)/base in %. The first column refers to the region over which the annual mean is spatially averaged, and the second column refers to the corresponding run. Regions are defined as follows: NA ($15^\circ N-55^\circ N$; $60^\circ W-125^\circ W$), EU ($25^\circ N-65^\circ N$; $10^\circ W-50^\circ E$), EA ($15^\circ N-50^\circ N$; $95^\circ E-160^\circ E$), and SA ($5^\circ N-35^\circ N$; $50^\circ E-95^\circ E$).

Region	Run	Unit	O ₃	NO _x	NOy	HNO ₃	PAN	N_2O_5	NO ₃
NA	REF	Conc*	38.94	1.29	1.59	0.15	0.14	14.85	2.71
NA	noN2O5	%	6	7	8	8	8	94	56
NA	noHO2	%	0	-1	-1	2	1	0	-3
NA	noHNO3	%	6	-1	11	127	2	6	11
NA	noNO2	%	0	2	2	-1	-1	3	2
NA	noNO3	%	0	0	0	0	0	0	1
NA	noO3	%	0	0	0	0	0	0	0
EUR	REF	Conc*	39.57	2.03	2.38	0.15	0.16	21.51	4.7
EUR	noN2O5	%	7	11	13	12	14	177	61
EUR	noHO2	%	1	-2	-1	3	6	-1	-4
EUR	noHNO3	%	5	-1	14	227	2	6	14
EUR	noNO2	%	0	6	4	-3	-5	5	3
EUR	noNO3	%	0	0	0	0	0	1	3
EUR	noO3	%	0	0	0	0	0	0	0
EA	REF	Conc*	38.51	2.1	2.54	0.17	0.26	10.05	2.64
EA	noN2O5	%	9	11	13	15	15	311	114
EA	noHO2	%	2	-5	-2	5	13	1	-2
EA	noHNO3	%	5	-1	10	148	1	5	9
EA	noNO2	%	0	29	21	-8	-21	10	5
EA	noNO3	%	0	0	0	0	0	0	2
EA	noO3	%	0	0	0	0	0	0	0
SA	REF	Conc*	44.26	1.29	1.53	0.1	0.12	15.5	6.15
SA	noN2O5	%	5	6	8	13	14	96	35
SA	noHO2	%	1	-3	-2	5	6	-1	-4
SA	noHNO3	%	8	-1	39	612	4	8	17
SA	noNO2	%	1	4	3	-1	0	8	5
SA	noNO3	%	0	0	0	0	0	2	3
SA	noO3	%	0	0	0	0	0	0	0

Table 5. Impacts of gas-aerosol reactions on regional ground level average mixing ratios of O_3 and key NO_y compounds. As Table 4, but for the ECHAM-HAMMOZ model.

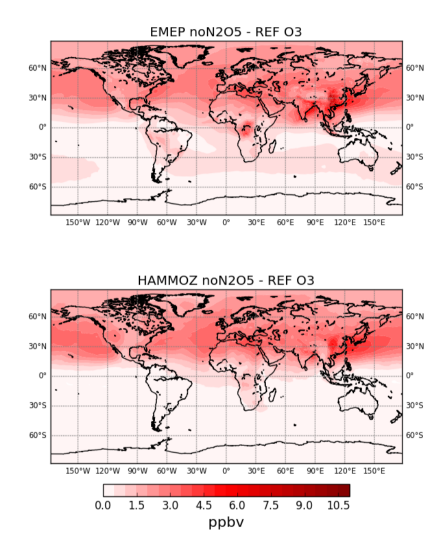


Figure 3. Differences in annual mean ground level ozone mixing ratio between the reference run REF and the sensitivity run noN2O5 for 2012. Since the Reference run was subtracted from the noN2O5 run, positive values show higher values in noN2O5 than in REF.

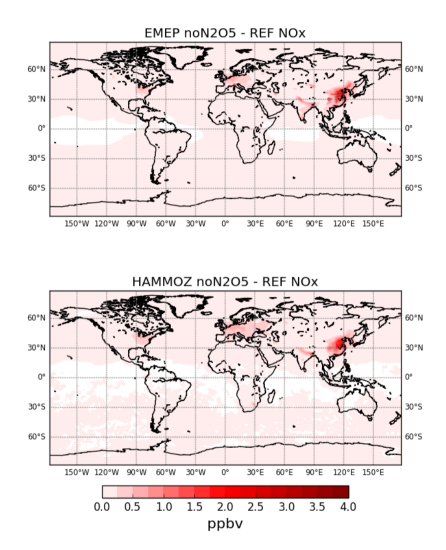


Figure 4. As Fig. 3, but for annual mean ground level NO_{x} mixing ratios.

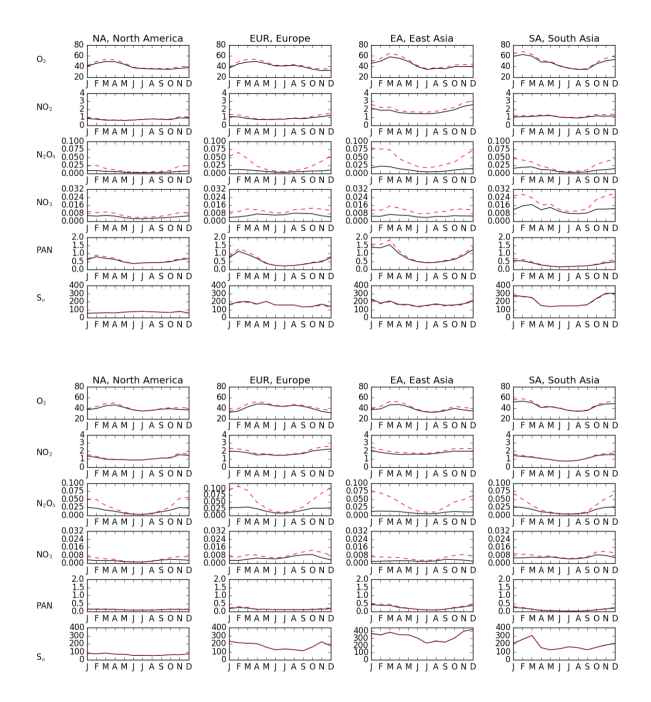


Figure 5. Changes in near-surface O_3 , NO_2 , N_2O_5 , NO_3 , PAN and S_a for the base-case (solid black line) and noN2O5 case (dashed red line) for EMEP (top) and ECHAM-HAMMOZ (bottom). Plots show monthly gas phase mixing ratios in ppbv and surface area density in $\mu m^2 cm^{-3}$ for different regions as defined above.

An important question is how sensitive the results are to the particular values chosen for the γ values. This is a complicated question, since these reactions also change the composition of NO_y in the atmosphere, the lifetime of NO₂ and hence the photoxidation processes leading to O₃. In order to address this, additional runs with the EMEP model in four new configurations were performed:

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- 1. $\gamma = 0.01$ for N₂O₅, a value lower than typical values, and at the low end of estimates (see Section 3.1).
 - 2. $\gamma = 0.1$ for N₂O₅, equivalent to values used by e.g. Dentener and Crutzen (1993); Tie et al. (2001, 2003), which is substantially higher than values obtained for $\gamma_{N_2O_5}$ used here (Table 2, Figure S1).
 - 3. $\gamma = 1.0 \times 10^{-3}$ for NO₂, at the top end of estimates (Section 3.3).
 - 4. $\gamma = 0.0$ for NO₂, since the lowest estimates are extremely low.
- 10 The model has been run for new base-cases including γ as listed above, and for the noN2O5, noHNO3 and (except for test (4)) noNO2 cases. Results for the regional averages (equivalent to manuscript Tables 4-5) are shown in Tables S4 and S5 in the supplement. Considering the N₂O₅ tests first, the changes in ozone over for example North America range from 3% (γ = 0.01) to 8% (γ = 0.1), compared to the original estimate in REF of 5% (Table 4). Changes for NO_x follow a similar pattern (e.g. 6-13% for NA, versus original 9%), but changes for N₂O₅ itself are much more significant (80% versus 354%, compared 15 to the original 160%).

Considering the γ tests for NO₂, the test results for the noN2O5 tests generally span those of the original runs, e.g. changes of 4–6% for ozone in North America versus 5% in the original run, or 113–170% for N₂O₅ versus 160% for the original case. Test (3), with the high $\gamma = 1.0 \times 10^{-3}$ for NO₂ does have significant impacts on the NO_x levels though, from e.g. 2% in the original run to 16% in test (3) for NA, or from 30% to 109% in East Asia. In these runs the impacts of noNO2 on ozone become comparable to those of noN2O5, and in South Asia the ozone changes from noNO2 actually exceed those from noN2O5.

Test (4), using zero γ actually gives results which are very similar to our default $\gamma = 1.0 \times 10^{-4}$, suggesting that this reaction only becomes important if higher values than $\times 10^{-4}$ can be justified.

Thus, we find that the exact changes in ozone and N-compounds do depend on the assumed γ values, but the relative importance of the different heterogeneous reactions generally remains. The N₂O₅ reactions are in nearly all cases the most important driver of ozone changes, but the use of a very high values for γ for NO₂ changes the picture somewhat. We can note

though that use of the high 0.1 values for $\gamma(NO_2)$ leads to quite significant reductions in annual NO₂ concentrations, resulting in degraded performance of the EMEP model compared to measurements (not shown), at least across the EMEP observational network in Europe (Tørseth et al., 2012).

5.3 Comparison with observations

30 Surface observations from 20 sites of the GAW and TOAR networks (Global Atmospheric Watch, Schultz et al. 2015, 2017), with stations distributed over the world, were used to evaluate ozone concentrations in the reference and N_2O_5 sensitivity runs of both models. The GAW data set consists of many sites in North America and Europe, but unfortunately few in Asia (e.g. none in China for 2012). Still, sites exist in Japan and these should provide a good indication of ozone formation downwind of mainland China. Mountain sites were excluded from this comparison in order to avoid problems with the interpretation of which model level is most appropriate for comparison. Trinidad Head on the west coast of USA and Mace Head on the west coast of Ireland are also good background stations which capture trends in hemispheric air masses arriving from the Pacific

5 and Atlantic respectively (Parrish et al., 2009, 2014). To capture the seasonal dependence of N_2O_5 uptake on aerosol, daily maximum ozone values were compared with the corresponding interpolated model data. Since the stations were selected to be relatively remote and low-elevation ground stations, the comparison with the coarse grids of the models might be representative.

Six out of the twenty stations are shown in Figs. 6, 7 and 8. Both models generally capture the seasonal variation well, fine structures and fluctuations are often reproduced, but not equally well by both models and depending on the station. For exam-

10 ple in Tsukuba, Japan both models simulate the increasing variability during summer time, nevertheless peak concentrations are still underestimated. EMEP calculates higher peak values, than ECHAM-HAMMOZ, in contrast in Waldhof, Germany, ECHAM-HAMMOZ simulates higher peak values, partially overestimating them compared to the observations.

A closer look at the dashed line compared to the solid line reveals the seasonal highest impact of N_2O_5 during spring time. The high impact in spring pattern can be found in both model simulations , but is stronger in EMEP, since ECHAM-HAMMOZ

- 15 includes the year through ozone reduction due to direct HNO_3 loss explained in the previous section 5.2. For example in Mace Head, Ireland the springtime ozone formation is clearly decreased by N_2O_5 reaction, while during summer the impact is marginal and increases again during winter. Both models start with a spin-up from the reference run, therefore the winter impact can not be seen in January. If the models would run for another month, this would show too, indicated by the gap between reference run and noN2O5 sensitivity run at the very end of the year.
- 20 Concluding, the impact of N_2O_5 heterogeneous reactions on chemical ozone production leads to a better agreement of EMEP and ECHAM-HAMMOZ and daily maximum ozone station observations in remote stations. Both models show improvements in model bias, which is expected with due to prior slight ozone overestimation; heterogeneous chemistry removes ozone, hence in models which tend to overestimate it, logically inclusion of such reactions tends to improve the model performance with regard to bias. Such improvements could also arise if introducing other nitrogen species loss processes to reduce ozone
- 25 production, having less NO_x emissions or dynamically inhibiting downward transport of stratospheric ozone. Especially the stratospheric ozone intrusion is assumed to strongly happen during spring time, which would cause the same pattern as we see here in Figures 6, 7 and 8.

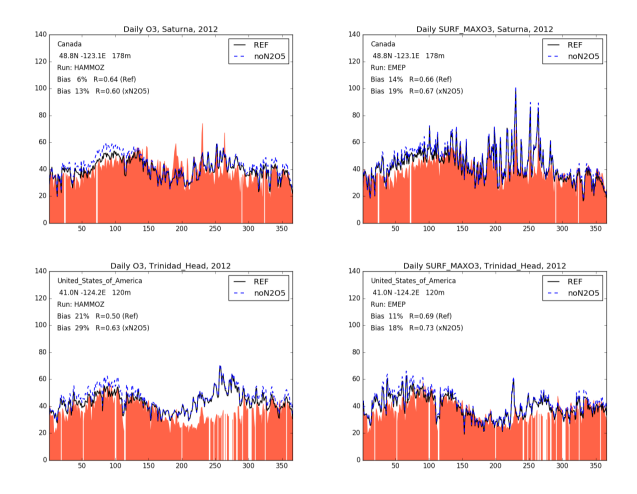


Figure 6. Modelled versus observed daily maximum ozone (ppbv) for two North American sites (Saturna, Canada, Trinidad Head, USA). The shaded area refers to surface station observations, the solid line is the reference run of the model and the dashed line the sensitivity run noN2O5 excluding heterogenous N_2O_5 reaction. On the upper left corner the station location, model, bias and correlation R are specified.

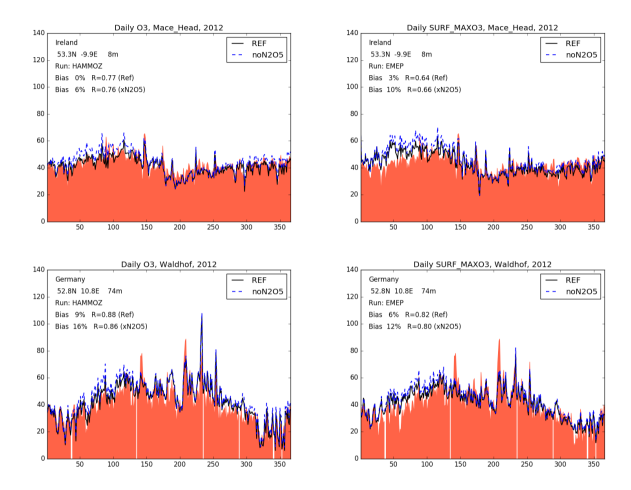


Figure 7. As Fig. 6 but for two European sites, Mace Head (Ireland) and Waldhof (Germany).

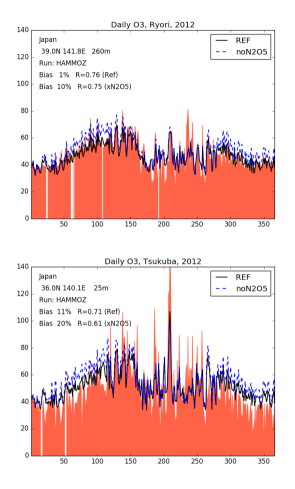
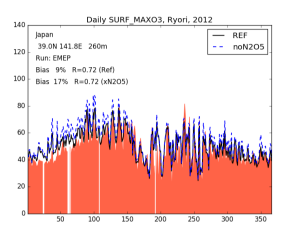
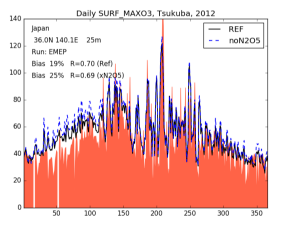


Figure 8. As Fig. 6 but for two Japanese sites, Ryori and Tsukuba.





5.4 Discussion

The influence of heterogeneous chemistry is known to be important in global chemical transport models, but it is also generally difficult to parameterize for many reasons (Jacob, 2000; Chang et al., 2011; Brown and Stutz, 2012). These include the difficulty of accurately simulating aerosol surface area density available for heterogeneous reactions and the large uncertainty in uptake

- 5 coefficients. In this section we briefly compare our results with some previous studies, and also comment on some of the remaining difficulties which will need to be tackled in future studies. Concerning modelling, many studies have been published on especially the importance of the N₂O₅ reactions, (e.g. Dentener and Crutzen, 1993; Tie et al., 2001, 2003; Evans and Jacob, 2005; Alexander et al., 2009; Macintyre and Evans, 2010; Chang et al., 2011).
- Table 6 presents a comparison of some reductions in O_3 and NO_x due to N_2O_5 aerosol uptake. Starting with the annual values, the classic study of Dentener and Crutzen (1993) produced the most dramatic changes in O_3 and especially NO_x (40-49%, depending on assumed γ value), with subsequent studies including ours producing smaller changes. Macintyre and Evans (2010) explored runs with a variety of γ coefficients, showing how O_3 and NO_x sensitivities change with different values. Our global results can be see to lie not too far from the Macintyre and Evans (2010) results obtained with $\gamma = 0.01$.
- As seen in Sect. 5.2, the impact of N_2O_5 hydrolysis is higher during winter in our study, and Table 6 confirms this for 15 other studies. Dentener and Crutzen (1993) report about a 75 % NO_x and 20 % O_3 reduction in their winter period with $\gamma = 0.1$. Although Tie et al. (2001) found such dramatic NO_x changes at 45°N, reductions were much smaller elsewhere (e.g. 3% at the equator). A follow-up study of Tie et al. (2003) gave global average NO_x and O_3 reductions of 38% and 6%, respectively, significantly lower than that found by Dentener and Crutzen (1993). Our models produce smaller changes again, for example ECHAM-HAMMOZ simulates a reduction in tropospheric NO_x due to N_2O_5 hydrolysis of 9% in winter,
- 20 (reductions in surface-level concentrations are greater, at 16 %). Also, O₃ reductions with our models are somewhat lower compared to these other models. EMEP shows the lowest changes in wintertime NO_x, though O₃ changes are closer to those of ECHAM-HAMMOZ.

Summertime results from Table 6 will not be discussed in detail, but again we see the same trend of more recent models producing smaller changes.

- There are many possible reasons for these differences. Firstly, the $\gamma_{N_2O_5}$ values used by Dentener and Crutzen 1993 and Tie et al. (2001, 2003) (= 0.1) are significantly larger than the typical values of around 0.01-0.04 as calculated in this study (c.f. Fig. S1) and seen in atmospheric observations (Brown et al., 2009; Brown and Stutz, 2012). Macintyre and Evans (2010) tested the model sensitivity to uniform $\gamma_{N_2O_5}$ values and report the highest sensitivity between 0.001 and 0.02. This is exactly the range of values given frequently by the $\gamma_{N_2O_5}$ parametrization used here. The impact of the hydrolysis reaction on ozone
- 30 is indeed stronger with higher γ , but our main results are relatively insensitive to these necessarily very uncertain choices (Sect. 5.2, Table S4). The ECHAM-HAMMOZ and EMEP models also have a set of other heterogeneous reactions competing with N₂O₅ hydrolysis, which again lowers the possible impact of this hydrolysis reaction.

There have been many changes in models, emissions, and indeed the atmosphere since these early studies. For example, the pioneering study of Dentener and Crutzen (1993) used had a model with a horizontal resolution of $10 \times 10^{\circ}$, giving grid cells

Table 6. Comparison of impacts found by turning off N_2O_5 hydrolysis in global model studies. Reductions in NO_x and O_3 found in the different global model studies are given in %.

Model	domain	$\gamma_{\rm N_2O_5}$	NO_x^a	O_3	Comments
Annual					
Den93	Trop.	0.1	49^a	9	globe
Den93	Trop.	0.01	40^a	4	globe
ME10	Trop	0.01	~ 12	~ 2.5	globe
ME10	Trop	0.01	~ 30	~ 6	N. Extra Trop.
ME10	Trop	0.1	~ 20	~ 7	globe
ME10	Trop	0.1	~ 38	~ 12	N. Extra Trop.
HAMMOZ	Trop.	Table 2	9.1	2.0	globe, this study
HAMMOZ	Trop.	Table 2	15	3.1	NH, this study
EMEP	Trop.	Table 2	16	2.4	globe, this study
Winter					
Den93	Trop.	0.1	75^a	20	NH, Nov-Apr.
Den93	Trop.	0.01	66^a	12	NH, Nov-Apr.
Tie01	Trop.	0.1^{b}	73	11	45°N, Dec.
Tie01	Trop.	0.1^{b}	3	3	Equator, Dec.
Tie03	Trop.	0.1^{b}	38	6	globe, Dec.
Tie03	Trop.	0.1^{b}	47	7	NH, Dec.
HAMMOZ	Trop.	Table 2	9.1	2.0	globe, Dec-Feb., this study
HAMMOZ	Trop.	Table 2	24.	3.8	NH, Dec-Feb., this study
HAMMOZ	surface	Table 2	16.	8	globe, Dec-Feb., this study
EMEP	surface	Table 2	5.1	4.9	globe, Dec-Feb., this study
Summer					
Den93	Trop.	0.1	45^a	13	NH, May-Oct.
Den93	Trop.	0.01	30^a	5	NH, May-Oct.
Tie01	Trop.	0.1^b	7	7	45°N, June
Tie01	Trop.	0.1^b	2	2	Equator, June
Tie03	Trop.	0.1^{b}	6	4	globe, Dec.
Tie03	Trop.	0.1^{b}	7	~ 5.5	NH, Dec.
EMEP	surface	Table 2	2.3	3.0	globe, Jun-Aug, this study
HAMMOZ	surface	Table 2	0.4	2	NH, Jun-Aug, this study
HAMMOZ	Trop.	Table 2	2.4	1.3	NH, Jun-Aug, this study

Refs: Den93: Dentener and Crutzen (1993) ,ME10: Macintyre and Evans (2010) ,Tie01: Tie et al. (2001) ;

Notes: a.Dentener and Crutzen (1993) reported changes in NOX=NO+NO2+NO3+2N2O5+HNO4, not NOx; b. Tie et al. (2001) used surface area of sulfate aerosols only; Trop. denotes full model domain, e.g. 0–100 hPa for EMEP, 0–4 hPa for Tie et al. (2001); NH denotes northen hemisphere. Data extracted from figures by eye indicated with ~. approximate.

with 100 times the area of the $1 \times 1^{\circ}$ grid used in EMEP or almost 30 times that of ECHAM-HAMMOZ's $1.85 \times 1.85^{\circ}$ grid. This alone will lead to different regimes of ozone productivity. It can also be noted that global CTMs (including changes due to emissions and chemical mechanisms) have improved over the years, so recent models should be expected to have different sensitivities to earlier studies (Wu et al., 2007). Emissions have also changed enormously over this period, especially in Asia (Granier et al., 2011); again with implications for the atmospheric oxidation capacity.

There are many other aspects of heterogeneous chemistry which are potentially important, but extremely complex and beyond the scope and abilities of our models. This includes for example the strong interactions of NO₂ with aerosol water and sulphate formation seen in wintertime haze events in Beijing (Cheng et al., 2016). However, Cheng et al. (2016) were concerned with extreme aerosol pollution events with concentrations exceeding 100 μ g m⁻³. These cannot be modeled at

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10 present in global scale models because of the dilution effect of the coarse grid resolution, and such extreme pollution events are likely to only have a local importance. In any case, it is not certain that the mechanism suggested by Cheng et al. (2016), is sufficient to explain some other extreme smog events (e.g. Guo et al., 2017).

Another important aspect for N_2O_5 heterogeneous chemistry is the formation of $ClNO_2$. In this case, N_2O_5 reacts with particulate chlorine to form gas-phase $ClNO_2$, which can photolyse and recycle NO_2 (Wang et al., 2016) and alter NO_y

- 15 composition (Sarwar et al., 2012). Especially in the planetary boundary layer of southern China, high mixing ratios of ClNO₂ (> 400 pptv) and N₂O₅ (> 1 ppbv) have been observed (Wang et al., 2016). The formation of ClNO₂ lowers the impact of N₂O₅ reaction on ozone, because it recycles NO₂ and was observed to enhance the ozone peak in southern China up to 16% (Wang et al., 2016). Unfortunately, our models (and indeed most global models) lack chlorine chemistry and treatment of chlorine in the aerosol thermodynamics, so cannot tackle these issues.
- Heterogeneous reactions on cloud surfaces, which can be important especially for HO₂ uptake depending of the presence of transition metal ions, were also excluded from our study. However, Dentener and Crutzen (1993) included the reaction of N₂O₅ on cloud droplets, but just found minor changes in NO_x and O₃. Jacob (2000) argue that for O₃, HO_x and NO_x, life times are not significantly reduced in clouds and current knowledge is insufficient to include cloud chemistry in O₃ models. In fact, most global model studies exclude the heterogeneous reactions of nitrogen species on clouds. Therefore, further development of CTM cloud-chemical systems will be needed before this question can be properly addressed.

In summary, our study finds a lower but still important impact of N_2O_5 hydrolysis on ozone and nitrogen oxides compared to previous model studies. However, earlier studies used rather high γ values for N_2O_5 , and neglected the other heterogeneous reactions. Further, chemical transport models have developed in many ways over the last 20-30 years, and indeed emissions across the globe have dramatically changed over this time period. In this paper we have illustrated that ECHAM-HAMMOZ

30 and EMEP, two up-to-date models systems, are rather consistent in the importance of N_2O_5 reactions, and that such reactions seem to be the most important among the six reactions we tested. Although one can never know if models produce good results for the right reasons, we have shown that both ECHAM-HAMMOZ and EMEP can reproduce even daily ozone variations remarkably well at sites across the globe (one can contrast results for Mace Head between Fig. 7 and the wide range of data from earlier models presented in Wild et al. 2012). We have also demonstrated that both models do a fair job of reproducing surface area density, so we believe our new estimates provide a valuable revision of calculations concerning the impact of heterogeneous reactions in such CTMs.

6 Conclusions

Two global transport models were used to investigate the implications of six heterogeneous (gas-aerosol uptake) reactions on
ground-level ozone concentrations. Both models were harmonized to use similar parameterizations for most of these reactions, enabling us to compare the impacts of N₂O₅, NO₃, NO₂, O₃, HNO₃, and HO₂ on ozone mixing ratios. Each reaction was evaluated systematically comparing the reference run to sensitivity simulations excluding one reaction at a time. Since heterogeneous reactions take place at the aerosol surface area, the modeled surface area density S_a of both models was compared to a satellite product retrieving the surface area. This comparison shows a good agreement in global pattern and especially the
capability of both models to capture the extreme aerosol loadings in East Asia.

The analysis of the sensitivity runs confirms that the globally most important heterogeneous reaction is the one of N_2O_5 . This impact was expected from previous studies, with the surface reactions of N_2O_5 having an impact on ozone mixing ratios through removal of reactive NO_x species. This result is loosely consistent with results from earlier studies (e.g. Dentener and Crutzen, 1993; Tie et al., 2001, 2003; Alexander et al., 2009; Macintyre and Evans, 2010), although here the magnitude of

- 15 changes induced by N_2O_5 reaction is at the low end of estimates, which seems to fit a trend whereby the more recent the study the lower the impacts of these reactions. Some other heterogeneous reactions (especially the ones of NO_2 , HO_2 and HNO_3) gain some significance in highly polluted areas where aerosol surface areas are high, but the two models show quite different response in their response to these other gas-aerosol reactions. The EMEP model actually shows rather small impacts of these reactions, except in East and South Asia where some impacts can approach 10-20% of that of N_2O_5 . ECHAM-HAMMOZ, on
- 20 the other hand, shows quite marked responses to especially the HNO_3 reactions. The reasons for this are related to differences in nitrate chemistry and surface area assumptions in the models, and to the differing spatial resolutions. It may well be that ECHAM-HAMMOZ overestimates the impact of HNO_3 due to missing nitrate aerosol formation and EMEP underestimates the impact, due to the use of only coarse sea salt and dust aerosol for the HNO_3 and HO_2 reactions.

The reactions of O_3 on dust and NO_3 on aerosols were found to have only minor effects on ozone in comparison to the other reactions in both models. In terms of global spatial impact, all reactions related to nitrogen species alter atmospheric chemistry downwind of source areas to some extent, with changes being much larger in the polluted northern hemisphere than in the southern hemisphere.

Evaluation of the models with northern hemispheric ozone surface observations from the GAW/TOAR networks yields a better agreement of the models with observations in terms of daily maximum concentrations, variability and temporal correla-

30 tions at most sites when the heterogeneous reactions are incorporated. The impacts of the N_2O_5 reactions show strong seasonal variations, with biggest impacts in spring time when photochemical reactions are active and N_2O_5 levels still high.

Due to lack of direct observations substantial uncertainties remain regarding the impact of heterogeneous reactions on tropospheric reactive gases. It should be noted, that neither model had an implementation of the particle-liquid-water/nitrate/chloride effects suggested by Bertram and Thornton (2009) and tested by e.g. Lowe et al. (2015). Further, neither model includes halogen chemistry, which is also known to impact O_3 in polluted regions (e.g. Sarwar et al., 2014; Li et al., 2016). The large impact of N_2O_5 seen in our work might be somewhat overestimated compared to that we would obtain if the chemistry of ClNO₂ (which would recycle NO_x) and other halogens could be included. Such improvements should result in better particle phase chemistry, and will be the subject of future work.

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Competing interests. No competing interests are present

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References

- Aas, W., Tsyro, S., Bieber, E., Bergström, R., Ceburnis, D., Ellermann, T., Fagerli, H., Frölich, M., Gehrig, R., Makkonen, U., Nemitz, E., Otjes, R., Perez, N., Perrino, C., Prévôt, A. S. H., Putaud, J.-P., Simpson, D., Spindler, G., Vana, M., and Yttri, K. E.: Lessons learnt from the first EMEP intensive measurement periods, Atmos. Chem. Physics, 12, 8073–8094, doi:10.5194/acp-12-8073-2012, http:
- 5 //www.atmos-chem-phys.net/12/8073/2012/, 2012.
- Ainsworth, E. A., Yendrek, C. R., Sitch, S., Collins, W. J., and Emberson, L. D.: The Effects of Tropospheric Ozone on Net Primary Productivity and Implications for Climate Change, Ann. Rev. Plant Biol, 63, 637–661, doi:10.1146/annurev-arplant-042110-103829, 2012.
 - Alexander, B., Hastings, M., Allman, D., Dachs, J., Thornton, J., and Kunasek, S.: Quantifying atmospheric nitrate formation pathways
- 10 based on a global model of the oxygen isotopic composition (Δ 17 O) of atmospheric nitrate, Atmospheric Chemistry and Physics, 9, 5043–5056, 2009.
 - Ammann, M., Cox, R., Crowley, J., Jenkin, M., Mellouki, A., Rossi, M., Troe, J., and Wallington, T.: Evaluated kinetic and photochemical data for atmospheric chemistry: Volume VI–heterogeneous reactions with liquid substrates, Atmospheric Chemistry and Physics, 13, 8045–8228, 2013.
- 15 Andersson-Sköld, Y. and Simpson, D.: Comparison of the chemical schemes of the EMEP MSC-W and the IVL photochemical trajectory models, Atmos. Environ., 33, 111–1129, 1999.
 - Angelbratt, J., Mellqvist, J., Simpson, D., Jonson, J. E., Blumenstock, T., Borsdorff, T., Duchatelet, P., Forster, F., Hase, F., Mahieu, E., Notholt, J., Petersen, A. K., Raffalski, U., Servais, C., Sussman, R., and Warneke, T.: Carbon monoxide (CO) and ethane (C₂H₆) trends from ground-based solar FTIR measurements at six European stations, comparison and sensitivity analysis with the EMEP model, Atmos.
- 20 Chem. Physics, 11, 9253–9269, doi:10.5194/acpd-11-13723-2011, http://www.atmos-chem-phys-discuss.net/11/13723/2011/, 2011. Arens, F., Gutzwiller, L., Baltensperger, U., Gäggeler, H. W., and Ammann, M.: Heterogeneous reaction of NO₂ on diesel soot particles, Environmental science & technology, 35, 2191–2199, 2001.
 - Bauer, S., Balkanski, Y., Schulz, M., Hauglustaine, D., and Dentener, F.: Global modeling of heterogeneous chemistry on mineral aerosol surfaces: Influence on tropospheric ozone chemistry and comparison to observations, Journal of Geophysical Research: Atmospheres

- Bergström, R., Denier van der Gon, H. A. C., Prévôt, A. S. H., Yttri, K. E., and Simpson, D.: Modelling of organic aerosols over Europe (2002–2007) using a volatility basis set (VBS) framework: application of different assumptions regarding the formation of secondary organic aerosol, Atmos. Chem. Physics, 12, 8499–8527, doi:10.5194/acp-12-8499-2012, http://www.atmos-chem-phys.net/12/8499/2012/, 2012.
- 30 Bertram, T. and Thornton, J.: Toward a general parameterization of N₂O₅ reactivity on aqueous particles: the competing effects of particle liquid water, nitrate and chloride, Atmospheric Chemistry and Physics, 9, 8351–8363, 2009.
 - Bian, H., Chin, M., Hauglustaine, D. A., Schulz, M., Myhre, G., Bauer, S. E., Lund, M. T., Karydis, V. A., Kucsera, T. L., Pan, X., Pozzer, A., Skeie, R. B., Steenrod, S. D., Sudo, K., Tsigaridis, K., Tsimpidi, A. P., and Tsyro, S. G.: Investigation of global particulate nitrate from the AeroCom phase III experiment, Atmospheric Chemistry and Physics, 17, 12911–12940, doi:10.5194/acp-17-12911-2017, https://www.com/doi/10.5194/acp-17-12911-2017, https://www.com/doi/10.5194/acp-17-12911-2017.
- 35 //www.atmos-chem-phys.net/17/12911/2017/, 2017.
 - Binkowski, F. and Shankar, U.: The Regional Particulate Matter Model .1. Model description and preliminary results, J. Geophys. Res., 100, 26 191–26 209, 1995.

^{25 (1984–2012), 109, 2004.}

- Brown, S. S. and Stutz, J.: Nighttime radical observations and chemistry, Chem. Soc. Rev., 41, 6405–6447, doi:10.1039/C2CS35181A, http://dx.doi.org/10.1039/C2CS35181A, 2012.
- Brown, S. S., Dube, W. P., Fuchs, H., Ryerson, T. B., Wollny, A. G., Brock, C. A., Bahreini, R., Middlebrook, A. M., Neuman, J. A., Atlas, E., Roberts, J. M., Osthoff, H. D., Trainer, M., Fehsenfeld, F. C., and Ravishankara, A. R.: Reactive uptake coefficients for N₂O₅ determined
- 5 from aircraft measurements during the Second Texas Air Quality Study: Comparison to current model parameterizations, J. Geophys. Res., 114, D00F10, doi:10.1029/2008JD011679, 2009.
 - Buseck, P. R. and Posfai, M.: Airborne minerals and related aerosol particles: Effects on climate and the environment, Proceedings of the National Academy of Sciences, 96, 3372–3379, 1999.
 - Chang, W. L., Bhave, P. V., Brown, S. S., Riemer, N., Stutz, J., and Dabdub, D.: Heterogeneous atmospheric chemistry, ambient measurements, and model calculations of N₂O₅: A review, Aerosol Science and Technology, 45, 665–695, 2011.
- Cheng, Y., Zheng, G., Wei, C., Mu, Q., Zheng, B., Wang, Z., Gao, M., Zhang, Q., He, K., Carmichael, G., et al.: Reactive nitrogen chemistry in aerosol water as a source of sulfate during haze events in China, Science advances, 2, e1601 530, 2016.
- Crutzen, P., Lawrence, M., and Poschl, U.: On the background photochemistry of tropospheric ozone, Tellus, 51, 123–146, international Symposium to Commemorate the 100th Anniversary of the Birth of Carl Gustaf Rossby (Rossby-100), STOCKHOLM, SWEDEN, JUN
- 15 08-12, 1998, 1999.

10

30

- Davies, J. A. and Cox, R. A.: Kinetics of the heterogeneous reaction of HNO₃ with NaCl: Effect of water vapor, The Journal of Physical Chemistry A, 102, 7631–7642, 1998.
- Davis, J., Bhave, P., and Foley, K.: Parameterization of N₂O₅ reaction probabilities on the surface of particles containing ammonium, sulfate, and nitrate, Atmospheric Chemistry and Physics, 8, 5295–5311, 2008.
- 20 Dentener, F. J. and Crutzen, P. J.: Reaction of N₂O₅ on tropospheric aerosols: Impact on the global distributions of NO_x, O₃, and OH, Journal of Geophysical Research: Atmospheres (1984–2012), 98, 7149–7163, 1993.
 - Emmons, L., Walters, S., Hess, P., Lamarque, J.-F., Pfister, G., Fillmore, D., Granier, C., Guenther, A., Kinnison, D., Laepple, T., et al.: Description and evaluation of the Model for Ozone and Related chemical Tracers, version 4 (MOZART-4), Geoscientific Model Development, 3, 43–67, 2010.
- 25 Evans, M. and Jacob, D. J.: Impact of new laboratory studies of N₂O₅ hydrolysis on global model budgets of tropospheric nitrogen oxides, ozone, and OH, Geophysical Research Letters, 32, 2005.
 - Fagerli, H. and Aas, W.: Trends of nitrogen in air and precipitation: Model results and observations at EMEP sites in Europe, 1980-2003, Environ. Poll., 154, 448–461, 2008.
 - Fairlie, T. D., Jacob, D. J., Dibb, J. E., Alexander, B., Avery, M. A., Donkelaar, A. v., and Zhang, L.: Impact of mineral dust on nitrate, sulfate, and ozone in transpacific Asian pollution plumes, Atmospheric Chemistry and Physics, 10, 3999–4012, 2010.
- Fiore, A., Dentener, F., Wild, O., Cuvelier, C., Schultz, M., Hess, P., Textor, C., Schulz, M., Doherty, R., Horowitz, L., et al.: Multimodel estimates of intercontinental source-receptor relationships for ozone pollution, Journal of Geophysical Research: Atmospheres (1984–2012), 114, 2009.
 - Frenzel, A., Scheer, V., Sikorski, R., George, C., Behnke, W., and Zetzsch, C.: Heterogeneous Interconversion Reactions of BrNO₂, ClNO₂,
- 35 Br₂, and Cl₂, The Journal of Physical Chemistry A, 102, 1329–1337, doi:10.1021/jp973044b, http://dx.doi.org/10.1021/jp973044b, 1998. Gauss, M., Benedictow, A., and Hjellbrekke, A.-G.: Photo-oxidants: validation and combined maps, Supplementary material to emep status report 1/2011, available online at www.emep.int, The Norwegian Meteorological Institute, Oslo, Norway, 2011.

- George, C., Ammann, M., D'Anna, B., Donaldson, D., and Nizkorodov, S. A.: Heterogeneous Photochemistry in the Atmosphere, Chemical reviews, 2015.
- George, I., Matthews, P., Whalley, L., Brooks, B., Goddard, A., Baeza-Romero, M., and Heard, D.: Measurements of uptake coefficients for heterogeneous loss of HO₂ onto submicron inorganic salt aerosols, Physical Chemistry Chemical Physics, 15, 12829–12845, 2013.
- 5 Gerber, H. E.: Relative-Humidity Parameterization of the Navy Aerosol Model (NAM), NRL Report 8956, Naval Research Laboratory, Washington, DC, 1985.
 - Goodman, A., Underwood, G., and Grassian, V.: Heterogeneous reaction of NO₂: Characterization of gas-phase and adsorbed products from the reaction, 2 NO₂ (g)+ H₂O (a) \rightarrow HONO (g)+ HNO₃ (a) on hydrated silica particles, The Journal of Physical Chemistry A, 103, 7217–7223, 1999.
- 10 Granier, C., Bessagnet, B., Bond, T., D'Angiola, A., van der Gon, H. D., Frost, G. J., Heil, A., Kaiser, J. W., Kinne, S., Klimont, Z., Kloster, S., Lamarque, J.-F., Liousse, C., Masui, T., Meleux, F., Mieville, A., Ohara, T., Raut, J.-C., Riahi, K., Schultz, M. G., Smith, S. J., Thompson, A., van Aardenne, J., van der Werf, G. R., and van Vuuren, D. P.: Evolution of anthropogenic and biomass burning emissions of air pollutants at global and regional scales during the 1980-2010 period, Climatic Change, 109, 163–190, doi:10.1007/s10584-011-0154-1, 2011.
- 15 Grewe, V., Brunner, D., Dameris, M., Grenfell, J., Hein, R., Shindell, D., and Staehelin, J.: Origin and variability of upper tropospheric nitrogen oxides and ozone at northern mid-latitudes, Atmospheric Environment, 35, 3421–3433, 2001.
 - Griffiths, P. T., Badger, C. L., Cox, R. A., Folkers, M., Henk, H. H., and Mentel, T. F.: Reactive uptake of N₂O₅ by aerosols containing dicarboxylic acids. Effect of particle phase, composition, and nitrate content, The Journal of Physical Chemistry A, 113, 5082–5090, 2009.
- 20 Gross, S. and Bertram, A. K.: Reactive uptake of NO₃, N₂O₅, NO₂, HNO₃, and O₃ on three types of polycyclic aromatic hydrocarbon surfaces, The Journal of Physical Chemistry A, 112, 3104–3113, 2008.
 - Gross, S. and Bertram, A. K.: Products and kinetics of the reactions of an alkane monolayer and a terminal alkene monolayer with NO₃ radicals, Journal of Geophysical Research: Atmospheres, 114, 2009.

Gross, S., Iannone, R., Xiao, S., and Bertram, A. K.: Reactive uptake studies of NO3 and N2O5 on alkenoic acid, alkanoate, and polyalcohol

- substrates to probe nighttime aerosol chemistry, Physical Chemistry Chemical Physics, 11, 7792–7803, 2009.
 - Guenther, A., Karl, T., Harley, P., Wiedinmyer, C., Palmer, P., and Geron, C.: Estimates of global terrestrial isoprene emissions using MEGAN (Model of Emissions of Gases and Aerosols from Nature), Atmospheric Chemistry and Physics Discussions, 6, 107–173, 2006.
 - Guo, H., Weber, R. J., and Nenes, A.: High levels of ammonia do not raise fine particle pH sufficiently to yield nitrogen oxide-dominated sulfate production, Scientific reports, 7, 12 109, 2017.
- 30 Hallquist, M., Wenger, J., Baltensperger, U., Rudich, Y., Simpson, D., Claeys, M., Dommen, J., Donahue, N., George, C., Goldstein, A., et al.: The formation, properties and impact of secondary organic aerosol: current and emerging issues, Atmospheric Chemistry and Physics, 9, 5155–5236, 2009.
 - Hanisch, F. and Crowley, J. N.: The heterogeneous reactivity of gaseous nitric acid on authentic mineral dust samples, and on individual mineral and clay mineral components, Physical Chemistry Chemical Physics, 3, 2474–2482, 2001.
- 35 Harrison, R. M. and Collins, G. M.: Measurements of reaction coefficients of NO₂ and HONO on aerosol particles, Journal of atmospheric chemistry, 30, 397–406, 1998.
 - Harrison, R. M. and Yin, J.: Particulate matter in the atmosphere: which particle properties are important for its effects on health?, Science of the total environment, 249, 85–101, 2000.

- Hauglustaine, D., Balkanski, Y., and Schulz, M.: A global model simulation of present and future nitrate aerosols and their direct radiative forcing of climate., Atmospheric Chemistry & Physics, 14, 2014.
- Henrot, A.-J., Stanelle, T., Schröder, S., Siegenthaler, C., Taraborrelli, D., and Schultz, M.: Implementation of the biogenic emission model MEGAN (v2. 1) into the ECHAM6-HAMMOZ chemistry climate model, Geoscientific Model Development [= GMD], 2017.
- 5 Hodzic, A., Bessagnet, B., and Vautard, R.: A model evaluation of coarse-mode nitrate heterogeneous formation on dust particles, Atmospheric Environment, 40, 4158–4171, 2006.
 - Huijnen, V., Williams, J., and Flemming, J.: Modeling global impacts of heterogeneous loss of HO₂ on cloud droplets, ice particles and aerosols, Atmospheric Chemistry and Physics Discussions, 14, 8575–8632, 2014.
 - IPCC: Annex I: Atlas of Global and Regional Climate Projections, in: Climate Change 2013: The Physical Science Basis. Contribution
- 10 of Working Group I to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change, edited by Stocker, T., Qin, D., Plattner, G.-K., Tignor, M., Allen, S., Boschung, J., Nauels, A., Xia, Y., Bex, V., and Midgley, P., book section AI, pp. 1311–1394, Cambridge University Press, Cambridge, United Kingdom and New York, NY, USA, doi:10.1017/CBO9781107415324.029, www.climatechange2013.org, 2013.

Jacob, D. J.: Heterogeneous chemistry and tropospheric ozone, Atmospheric Environment, 34, 2131-2159, 2000.

- 15 Jonson, J., Stohl, A., Fiore, A., Hess, P., Szopa, S., Wild, O., Zeng, G., Dentener, F., Lupu, A., Schultz, M., Duncan, B., Sudo, K., Wind, P., Schulz, M., Marmer, E., Cuvelier, C., Keating, T., Zuber, A., Valdebenito, A., Dorokhov, V., De Backer, H., Davies, J., Chen, G., Johnson, B., Tarasick, D., Stübi, R., Newchurch, M., von der Gathen, P., Steinbrecht, W., and Claude, H.: A multi-model analysis of vertical ozone profiles, Atmos. Chem. Physics, 10, 5759–5783, doi:10.5194/acp-10-5759-2010, 2010.
- Jonson, J. E., Semeena, V. S., and Simpson, D.: Global ozone bias, in: Transboundary particulate matter, photo-oxidants, acidifying and eutrophying components. Status Report 1/2015, pp. 115–128. The Norwegian Meteorological Institute, Oslo, Norway, 2015.
- Jülich Supercomputing Centre: JURECA: General-purpose supercomputer at Jülich Supercomputing Centre, Journal of large-scale research facilities, 2, doi:10.17815/jlsrf-2-121, http://dx.doi.org/10.17815/jlsrf-2-121, 2016.
 - Kinnison, D., Brasseur, G., Walters, S., Garcia, R., Marsh, D., Sassi, F., Harvey, V., Randall, C., Emmons, L., Lamarque, J., et al.: Sensitivity of chemical tracers to meteorological parameters in the MOZART-3 chemical transport model, Journal of Geophysical Research: Atmospheres (1984, 2012), 112, 2007
- 25 Atmospheres (1984–2012), 112, 2007.

30

- Kleffmann, J., Becker, K., and Wiesen, P.: Heterogeneous NO₂ conversion processes on acid surfaces: possible atmospheric implications, Atmospheric Environment, 32, 2721–2729, 1998.
- Lamarque, J., Emmons, L., Hess, P., Kinnison, D. E., Tilmes, S., Vitt, F., Heald, C., Holland, E. A., Lauritzen, P., Neu, J., et al.: CAM-chem: Description and evaluation of interactive atmospheric chemistry in the Community Earth System Model, Geosci. Model Dev, 5, 369–411, 2012.
- Li, Q., Zhang, L., Wang, T., Tham, Y. J., Ahmadov, R., Xue, L., Zhang, Q., and Zheng, J.: Impacts of heterogeneous uptake of dinitrogen pentoxide and chlorine activation on ozone and reactive nitrogen partitioning: improvement and application of the WRF-Chem model in southern China, Atmos. Chem. Physics, 16, 14875–14890, doi:10.5194/acp-16-14875-2016, 2016.
- Liao, H. and Seinfeld, J. H.: Global impacts of gas-phase chemistry-aerosol interactions on direct radiative forcing by anthropogenic aerosols
- and ozone, Journal of Geophysical Research: Atmospheres (1984–2012), 110, 2005.
 - Liu, Y., Gibson, E. R., Cain, J. P., Wang, H., Grassian, V. H., and Laskin, A.: Kinetics of heterogeneous reaction of CaCO₃ particles with gaseous HNO₃ over a wide range of humidity, The Journal of Physical Chemistry A, 112, 1561–1571, 2008.

Lowe, D., Archer-Nicholls, S., Morgan, W., Allan, J., Utembe, S., Ouyang, B., Aruffo, E., Le Breton, M., Zaveri, R. A., Di Carlo, P., Percival, C., Coe, H., Jones, R., and McFiggans, G.: WRF-Chem model predictions of the regional impacts of N2O5 heterogeneous processes on night-time chemistry over north-western Europe, Atmos. Chem. Physics, 15, 1385–1409, doi:10.5194/acp-15-1385-2015, 2015.

Lowe, D., Ryder, J., Leigh, R., Dorsey, J. R., and McFiggans, G.: Modelling multi-phase halogen chemistry in the coastal marine bound-

- 5 ary layer: investigation of the relative importance of local chemistry vs. long-range transport, Atmos. Chem. Physics, 11, 979–994, doi:10.5194/acp-11-979-2011, 2011.
 - Macintyre, H. and Evans, M.: Sensitivity of a global model to the uptake of N₂O₅ by tropospheric aerosol, Atmospheric Chemistry and Physics, 10, 7409–7414, 2010.

- Mogili, P. K., Kleiber, P. D., Young, M. A., and Grassian, V. H.: Heterogeneous uptake of ozone on reactive components of mineral dust aerosol: An environmental aerosol reaction chamber study, The Journal of Physical Chemistry A, 110, 13799–13807, 2006.
- Moise, T., Talukdar, R., Frost, G., Fox, R., and Rudich, Y.: Reactive uptake of NO₃ by liquid and frozen organics, Journal of Geophysical Research: Atmospheres (1984–2012), 107, AAC–6, 2002.
- 15 Morgan, W. T., Ouyang, B., Allan, J. D., Aruffo, E., Di Carlo, P., Kennedy, O. J., Lowe, D., Flynn, M. J., Rosenberg, P. D., Williams, P. I., Jones, R., McFiggans, G. B., and Coe, H.: Influence of aerosol chemical composition on N2O5 uptake: airborne regional measurements in northwestern Europe, Atmospheric Chemistry and Physics, 15, 973–990, doi:10.5194/acp-15-973-2015, 2015.
 - Mozurkewich, M. and Calvert, J. G.: Reaction possibility of N₂O₅ on aqueous aerosols, Journal of Geophysical Research: Atmospheres, 93, 15,889–15,896, 1988.
- 20 Neubauer, D., Siegenthaler-Le Drian, C., Ferrachat, S., Bey, I., Lohmann, U., Stanelle, T., Frontoso, G., Stier, P., Schutgens, N., Schmidt, H., Rast, S., Schultz, M. G., Schroeder, S., Tegen, I., and Heinold, B.: The new version of the global aerosol-climate model ECHAM6-HAM: aerosol evaluation against observations and previous versions, Special Issue Global Model Development and Atmospheric Chemistry and Physics, in preparation.

Nicolas, M., Ndour, M., Ka, O., D'Anna, B., and George, C.: Photochemistry of atmospheric dust: ozone decomposition on illuminated titanium dioxide, Environmental science & technology, 43, 7437–7442, 2009.

- Parrish, D., Millet, D., and Goldstein, A.: Increasing ozone in marine boundary layer inflow at the west coasts of North America and Europe, Atmospheric Chemistry and Physics, 9, 1303–1323, 2009.
- Parrish, D., Lamarque, J.-F., Naik, V., Horowitz, L., Shindell, D., Staehelin, J., Derwent, R., Cooper, O., Tanimoto, H., Volz-Thomas, A., et al.: Long-term changes in lower tropospheric baseline ozone concentrations: Comparing chemistry-climate models and observations at
- 30 northern midlatitudes, Journal of Geophysical Research: Atmospheres, 119, 5719–5736, 2014.

10

25

Pathak, R., Wu, W., and Wang, T.: Summertime PM_{2.5} ionic species in four major cities of China: nitrate formation in an ammonia-deficient atmosphere, Atmospheric Chemistry and Physics, 9, 1711–1722, 2009.

Pathak, R. K., Wu, W. S., and Wang, T.: Summertime PM_{2.5} ionic species in four major cities of China: nitrate formation in an ammoniadeficient atmosphere, Atmospheric Chemistry and Physics Discussions, 8, 11 487–11 517, 2008.

- 35 Platt, U., Perner, D., Harris, G., Winer, A., and Pitts, J. j.: Observations of nitrous acid in an urban atmosphere by differential optical absorption, Nature, 285, 312–314, 1980.
 - Pöschl, U.: Atmospheric aerosols: composition, transformation, climate and health effects, Angewandte Chemie International Edition, 44, 7520–7540, 2005.

Mao, J., Fan, S., Jacob, D. J., and Travis, K. R.: Radical loss in the atmosphere from Cu-Fe redox coupling in aerosols, Atmospheric Chemistry and Physics, 13, 509–519, 2013.

Ravishankara, A.: Heterogeneous and multiphase chemistry in the troposphere, Science, 276, 1058–1065, 1997.

Remorov, R., Gershenzon, Y. M., Molina, L., and Molina, M.: Kinetics and mechanism of HO₂ uptake on solid NaCl, The Journal of Physical Chemistry A, 106, 4558–4565, 2002.

Reus, M., Dentener, F., Thomas, A., Borrmann, S., Ström, J., and Lelieveld, J.: Airborne observations of dust aerosol over the North Atlantic

- 5 Ocean during ACE 2: Indications for heterogeneous ozone destruction, Journal of Geophysical Research: Atmospheres (1984–2012), 105, 15 263–15 275, 2000.
 - Riedel, T., Bertram, T., Ryder, O., Liu, S., Day, D., Russell, L., Gaston, C., Prather, K., and Thornton, J.: Direct N₂O₅ reactivity measurements at a polluted coastal site, Atmospheric Chemistry and Physics, 12, 2959–2968, 2012.
 - Riemer, N., Vogel, H., Vogel, B., Schell, B., Ackermann, I., Kessler, C., and Hass, H.: Impact of the heteorogenous hydrolysis
- 10 of N_2O_5 on chemistry and nitrate formation in the lower troposhere under photosmog conditions, J. Geophys. Res., 108, 4144, doi:10.1029/2002JD002436, 2003a.
 - Riemer, N., Vogel, H., Vogel, B., Schell, B., Ackermann, I., Kessler, C., and Hass, H.: Impact of the heterogeneous hydrolysis of N₂O₅ on chemistry and nitrate aerosol formation in the lower troposphere under photosmog conditions, Journal of Geophysical Research: Atmospheres (1984–2012), 108, 2003b.
- 15 Riemer, N., Vogel, H., Vogel, B., Anttila, T., Kiendler-Scharr, A., and Mentel, T.: Relative importance of organic coatings for the heterogeneous hydrolysis of N₂O₅ during summer in Europe, Journal of Geophysical Research: Atmospheres (1984–2012), 114, 2009.

Rudich, Y., Talukdar, R. K., Ravishankara, A., and Fox, R.: Reactive uptake of NO₃ on pure water and ionic solutions, Journal of Geophysical Research: Atmospheres (1984–2012), 101, 21 023–21 031, 1996.

- Rudich, Y., Talukdar, R., and Ravishankara, A.: Multiphase chemistry of NO₃ in the remote troposphere, Journal of Geophysical Research:
 Atmospheres (1984–2012), 103, 16 133–16 143, 1998.
 - Saathoff, H., Naumann, K.-H., Riemer, N., Kamm, S., Möhler, O., Schurath, U., Vogel, H., and Vogel, B.: The loss of NO₂, HNO₃, NO₃/N₂O₅, and HO₂/HOONO₂ on soot aerosol: A chamber and modeling study, Geophysical Research Letters, 28, 1957–1960, 2001.

Sander, S. P., Friedl, R., Golden, D., Kurylo, M., Moortgat, G., Keller-Rudek, H., Wine, P., Ravishankara, A., Kolb, C., Molina, M., et al.: Chemical kinetics and photochemical data for use in atmospheric studies: evaluation number 15, National Aeronautics and Space Admin-

- 25 istration, Jet Propulsion Laboratory, California Institute of Technology Pasadena, CA, 2006.
 - Sarwar, G., Simon, H., Bhave, P., and Yarwood, G.: Examining the impact of heterogeneous nitryl chloride production on air quality across the United States, Atmospheric Chemistry and Physics, 12, 6455–6473, 2012.
 - Sarwar, G., Simon, H., Xing, J., and Mathur, R.: Importance of tropospheric CINO₂ chemistry across the Northern Hemisphere, Geophys. Res. Lett., 41, 4050–4058, doi:10.1002/2014GL059962, 2014.
- 30 Schaap, M., Cuvelier, C., Hendriks, C., Bessagnet, B., Baldasano, J., Colette, A., Thunis, P., Karam, D., Fagerli, H., Graff, A., Kranenburg, R., Nyiri, A., Pay, M., RouÃ-I, L., Schulz, M., Simpson, D., Stern, R., Terrenoire, E., and Wind, P.: Performance of European chemistry transport models as function of horizontal resolution, Atmospheric Environment, 112, 90 105, doi:http://dx.doi.org/10.1016/j.atmosenv.2015.04.003, http://www.sciencedirect.com/science/article/pii/S1352231015300066, 2015.
- Schultz, M. G., Akimoto, H., Bottenheim, J., Buchmann, B., Galbally, I. E., Gilge, S., Helmig, D., Koide, H., Lewis, A. C., Novelli, P. C., Plass-Duelmer, C., Ryerson, T. B., Steinbacher, M., Steinbrecher, R., Tarasova, O., Torseth, K., Thouret, V., and
- Zellweger, C.: The Global Atmosphere Watch reactive gases measurement network, Elementa-Science of the Anthropocene, 3, doi:http://doi.org/10.12952/journal.elementa.000067, 2015.

38

- Schultz, M. G., Schroder, S., Lyapina, O., Cooper, O. R., Galbally, I., Petropavlovskikh, I., von Schneidemesser, E., Tanimoto, H., Elshorbany, Y., Naja, M., Seguel, R. J., Dauert, U., Eckhardt, P., Feigenspan, S., Fiebig, M., Hjellbrekke, A.-G., Hong, Y.-D., Kjeld, P. C., Koide, H., Lear, G., Tarasick, D., Ueno, M., Wallasch, M., Baumgardner, D., Chuang, M.-T., Gillett, R., Lee, M., Molloy, S., Moolla, R., Wang, T., Sharps, K., Adame, J. A., Ancellet, G., Apadula, F., Artaxo, P., Barlasina, M. E., Bogucka, M., Bonasoni, P., Chang, L., Colomb, A.,
- 5 Cuevas-Agullo, E., Cupeiro, M., Degorska, A., Ding, A., FrHlich, M., Frolova, M., Gadhavi, H., Gheusi, F., Gilge, S., Gonzalez, M. Y., Gros, V., Hamad, S. H., Helmig, D., Henriques, D., Hermansen, O., Holla, R., Hueber, J., Im, U., Jaffe, D. A., Komala, N., Kubistin, D., Lam, K.-S., Laurila, T., Lee, H., Levy, I., Mazzoleni, C., Mazzoleni, L. R., McClure-Begley, A., Mohamad, M., Murovec, M., Navarro-Comas, M., Nicodim, F., Parrish, D., Read, K. A., Reid, N., Ries, L., Saxena, P., Schwab, J. J., Scorgie, Y., Senik, I., Simmonds, P., Sinha, V., Skorokhod, A. I., Spain, G., Spangl, W., Spoor, R., Springston, S. R., Steer, K., Steinbacher, M., Suharguniyawan, E., Torre, P., Trickl,
- 10 T., Weili, L., Weller, R., Xiaobin, X., Xue, L., and Zhiqiang, M.: Tropospheric Ozone Assessment Report: Database and metrics data of global surface ozone observations, Elementa-Science of the Anthropocene, 5, 58–Article, doi:10.1525/elementa.244, 2017.
 - Schultz, M. G., Stadtler, S., Schroeder, S., Taraborrelli, D., Henrot, A., Kaffashzadeh, N., Franco, B., Ferrachat, S., Siegenthaler-Le Drian, C., Lohmann, U., Neubauer, D., Wahl, S., Kokkola, H., Kuehn, T., Stier, P., Kinnison, D., Tyndall, G., and Orlando, J.: The Chemistry Climate Model ECHAM-HAMMOZ, Golbal Model Development, in preparation.
- 15 Schwartz, S. E.: Mass-transport considerations pertinent to aqueous phase reactions of gases in liquid-water clouds, in: Chemistry of multiphase atmospheric systems, pp. 415–471, Springer, 1986.

Seinfeld, J. H. and Pandis, S. N.: Atmospheric chemistry and physics: from air pollution to climate change, John Wiley & Sons, 2012.

- Sillman, S., Logan, J., and Wofsy, S.: The sensitivity of ozone to nitrogen oxides and hydrocarbons in regional ozone episodes, J. Geophys. Res., 95, 1837–1851, 1990.
- 20 Simpson, D., Winiwarter, W., Börjesson, G., Cinderby, S., Ferreiro, A., Guenther, A., Hewitt, C. N., Janson, R., Khalil, M. A. K., Owen, S., Pierce, T. E., Puxbaum, H., Shearer, M., Skiba, U., Steinbrecher, R., Tarrasón, L., and Öquist, M. G.: Inventorying emissions from Nature in Europe, J. Geophys. Res., 104, 8113–8152, 1999.
 - Simpson, D., Butterbach-Bahl, K., Fagerli, H., Kesik, M., Skiba, U., and Tang, S.: Deposition and Emissions of Reactive Nitrogen over European Forests: A Modelling Study, Atmos. Environ., 40, 5712–5726, doi:10.1016/j.atmosenv.2006.04.063, 2006a.
- 25 Simpson, D., Fagerli, H., Hellsten, S., Knulst, J., and Westling, O.: Comparison of modelled and monitored deposition fluxes of sulphur and nitrogen to ICP-forest sites in Europe, Biogeosciences, 3, 337–355, 2006b.
 - Simpson, D., Benedictow, A., Berge, H., Bergström, R., Emberson, L. D., Fagerli, H., Flechard, C. R., Hayman, G. D., Gauss, M., Jonson, J. E., Jenkin, M. E., Nyíri, A., Richter, C., Semeena, V. S., Tsyro, S., Tuovinen, J.-P., Valdebenito, A., and Wind, P.: The EMEP MSC-W chemical transport model technical description, Atmos. Chem. Physics, 12, 7825–7865, doi:10.5194/acp-12-7825-2012, http://www.
- 30 atmos-chem-phys.net/12/7825/2012/acp-12-7825-2012.html, 2012.
 - Simpson, D., Arneth, A., Mills, G., Solberg, S., and Uddling, J.: Ozone the persistent menace; interactions with the N cycle and climate change, Current Op. Environ. Sust., 9-10, 9–19, doi:http://dx.doi.org/10.1016/j.cosust.2014.07.008, sI: System dynamics and sustainability, 2014.
- Simpson, D., Tsyro, S., and Wind, P.: Updates to the EMEP/MSC-W model, in: Transboundary particulate matter, photo-oxidants, acidifying
 and eutrophying components. Status Report 1/2015, pp. 129–138, The Norwegian Meteorological Institute, Oslo, Norway, 2015.
 - Simpson, D., Bergström, R., Denby, B., Wind, P., and Others, X.: Updates to the EMEP/MSC-W model, 2016–2017, in: Transboundary particulate matter, photo-oxidants, acidifying and eutrophying components. EMEP Status Report 1/2016, p. xx, The Norwegian Meteorological Institute, Oslo, Norway, 2017.

Solomon, S.: Stratospheric ozone depletion: A review of concepts and history, Reviews of Geophysics-Richmond Virginia Washington, 37, 275-316. 1999.

Solomon, S., Garcia, R. R., Rowland, F. S., and Wuebbles, D. J.: On the depletion of Antarctic ozone, Nature, 321, 755–758, 1986.

Stadtler, S.: Heterogeneous N₂O₅ Chemistry in the Aerosol-Chemistry-Climate Model ECHAM6-HAMMOZ, Master's thesis, University of Bonn, 2015.

- 5
 - Stein, O., Flemming, J., Inness, A., Kaiser, J. W., and Schultz, M. G.: Global reactive gases forecasts and reanalysis in the MACC project, Journal of Integrative Environmental Sciences, 9, 57-70, 2012.
 - Stevens, B., Giorgetta, M., Esch, M., Mauritsen, T., Crueger, T., Rast, S., Salzmann, M., Schmidt, H., Bader, J., Block, K., et al.: Atmospheric component of the MPI-M Earth System Model: ECHAM6, Journal of Advances in Modeling Earth Systems, 5, 146-172, 2013.
- 10 Stone, D., Evans, M. J., Walker, H., Ingham, T., Vaughan, S., Ouyang, B., Kennedy, O. J., McLeod, M. W., Jones, R. L., Hopkins, J., Punjabi, S., Lidster, R., Hamilton, J. F., Lee, J. D., Lewis, A. C., Carpenter, L. J., Forster, G., Oram, D. E., Reeves, C. E., Bauguitte, S., Morgan, W., Coe, H., Aruffo, E., Dari-Salisburgo, C., Giammaria, F., Di Carlo, P., and Heard, D. E.: Radical chemistry at night: comparisons between observed and modelled HOx, NO₃ and N₂O₅ during the RONOCO project, Atmospheric Chemistry and Physics, 14, 1299–1321, doi:10.5194/acp-14-1299-2014, 2014.
- 15 Taketani, F., Kanaya, Y., and Akimoto, H.: Kinetics of heterogeneous reactions of HO2 radical at ambient concentration levels with (NH₄)₂SO₄ and NaCl aerosol particles, The Journal of Physical Chemistry A, 112, 2370–2377, 2008.
 - Taraborrelli, D., Lawrence, M., Butler, T., Sander, R., and Lelieveld, J.: Mainz Isoprene Mechanism 2 (MIM2): an isoprene oxidation mechanism for regional and global atmospheric modelling, Atmospheric Chemistry and Physics, 9, 2751-2777, 2009.
- Thornton, J. and Abbatt, J. P.: Measurements of HO_2 uptake to aqueous aerosol: Mass accommodation coefficients and net reactive loss, 20 Journal of Geophysical Research: Atmospheres (1984–2012), 110, 2005.
 - Thornton, J. A., Braban, C. F., and Abbatt, J. P.: N₂O₅ hydrolysis on sub-micron organic aerosols: the effect of relative humidity, particle phase, and particle size, Physical Chemistry Chemical Physics, 5, 4593–4603, 2003.
 - Thornton, J. A., Jaeglé, L., and McNeill, V. F.: Assessing known pathways for HO₂ loss in aqueous atmospheric aerosols: Regional and global impacts on tropospheric oxidants, Journal of Geophysical Research: Atmospheres (1984–2012), 113, 2008.
- 25 Tie, X., Brasseur, G., Emmons, L., Horowitz, L., and Kinnison, D.: Effects of aerosols on tropospheric oxidants: A global model study, Journal of Geophysical Research: Atmospheres, 106, 22931-22964, 2001.
 - Tie, X., Emmons, L., Horowitz, L., Brasseur, G., Ridley, B., Atlas, E., Stround, C., Hess, P., Klonecki, A., Madronich, S., et al.: Effect of sulfate aerosol on tropospheric NOx and ozone budgets: Model simulations and TOPSE evidence, Journal of Geophysical Research: Atmospheres, 108, 2003.
- 30 Tilgner, A., Majdik, Z., Sehili, A., Simmel, M., Wolke, R., and Herrmann, H.: SPACCIM: Simulations of the multiphase chemistry occurring in the FEBUKO hill cap cloud experiments, Atmospheric Environment, 39, 4389-4401, 2005.
 - Tørseth, K., Aas, W., Breivik, K., Fjæraa, A. M., Fiebig, M., Hjellbrekke, A. G., Lund Myhre, C., Solberg, S., and Yttri, K. E.: Introduction to the European Monitoring and Evaluation Programme (EMEP) and observed atmospheric composition change during 1972-2009, Atmos. Chem. Phys., 12, 5447–5481, doi:10.5194/acp-12-5447-2012, http://www.atmos-chem-phys.net/12/5447/2012/, 2012.
- Tsyro, S., Aas, W., Soares, J., Sofiev, M., Berge, H., and Spindler, G.: Modelling of sea salt concentrations over Europe: key uncer-35 tainties and comparison with observations, Atmos. Chem. Physics, 11, 10367-10388, doi:10.5194/acp-11-10367-2011, http://www. atmos-chem-phys.net/11/10367/2011/, 2011.

Underwood, G., Song, C., Phadnis, M., Carmichael, G., and Grassian, V.: Heterogeneous reactions of NO₂ and HNO₃ on oxides and mineral dust: A combined laboratory and modeling study, Journal of Geophysical Research: Atmospheres (1984–2012), 106, 18 055–18 066, 2001.
Usher, C. R., Michel, A. E., and Grassian, V. H.: Reactions on mineral dust, Chemical reviews, 103, 4883–4940, 2003.
van Donkelaar, A., Martin, R. V., Brauer, M., and Boys, B. L.: Use of satellite observations for long-term exposure assessment of global

- 5 concentrations of fine particulate matter, Environmental health perspectives, 123, 135, 2015.
- Van Vuuren, D. P., Edmonds, J., Kainuma, M., Riahi, K., Thomson, A., Hibbard, K., Hurtt, G. C., Kram, T., Krey, V., Lamarque, J.-F., et al.: The representative concentration pathways: an overview, Climatic change, 109, 5, 2011.
 - Verwer, J. and Simpson, D.: Explicit methods for stiff ODEs from atmospheric chemistry, Applied Numerical Mathematics, 18, 413–430, 1995.
- 10 Verwer, J. G., Blom, J. G., Van Loon, M., and Spee, E. J.: A comparison of stiff ODE solvers for atmospheric chemistry problems, Atmos. Environ., 30, 49–58, 1996.
 - Vieno, M., Dore, A. J., Bealey, W. J., Stevenson, D. S., and Sutton, M. A.: The importance of source configuration in quantifying footprints of regional atmospheric sulphur deposition, Science of the Total Environment, 408, 985–995, doi:10.1016/j.scitotenv.2009.10.048, 2010.
 Vieno, M., Heal, M. R., Hallsworth, S., Famulari, D., Doherty, R. M., Dore, A. J., Tang, Y. S., Braban, C. F., Leaver, D., Sutton, M. A., and
- 15 Reis, S.: The role of long-range transport and domestic emissions in determining atmospheric secondary inorganic particle concentrations across the UK, Atmos. Chem. Physics, 14, 8435–8447, doi:10.5194/acp-14-8435-2014, http://www.atmos-chem-phys.net/14/8435/2014/, 2014.

Vignati, E., Wilson, J., and Stier, P.: M7: An efficient size-resolved aerosol microphysics module for large-scale aerosol transport models, Journal of Geophysical Research: Atmospheres (1984–2012), 109, 2004.

- 20 Vinken, G. C. M., Boersma, K. F., Jacob, D. J., and Meijer, E. W.: Accounting for non-linear chemistry of ship plumes in the GEOS-Chem global chemistry transport model, Atmos. Chem. Physics, 11, 11707–11722, doi:10.5194/acp-11-11707-2011, http://www. atmos-chem-phys.net/11/11707/2011/, 2011.
 - von Glasow, R., Lawrence, M. G., Sander, R., and Crutzen, P. J.: Modeling the chemical effects of ship exhaust in the cloud-free marine boundary layer, Atmos. Chem. Physics, 3, 233–250, http://www.atmos-chem-phys.net/3/233/2003/, 2003.
- 25 Wagner, N., Riedel, T., Young, C., Bahreini, R., Brock, C., Dubé, W., Kim, S., Middlebrook, A., Öztürk, F., Roberts, J., et al.: N₂O₅ uptake coefficients and nocturnal NO₂ removal rates determined from ambient wintertime measurements, Journal of Geophysical Research: Atmospheres, 118, 9331–9350, 2013.
 - Wahner, A., Mentel, T. F., Sohn, M., and Stier, J.: Heterogeneous reaction of N₂O₅ on sodium nitrate aerosol, Journal of Geophysical Research: Atmospheres (1984–2012), 103, 31 103–31 112, 1998.
- 30 Wang, T., Tham, Y. J., Xue, L., Li, Q., Zha, Q., Wang, Z., Poon, S. C., Dubé, W. P., Blake, D. R., Louie, P. K., et al.: Observations of nitryl chloride and modeling its source and effect on ozone in the planetary boundary layer of southern China, Journal of Geophysical Research: Atmospheres, 121, 2476–2489, 2016.

Weiss, A. and Norman, J. M.: Partitioning Solar-radiation into Direct and Diffuse, Visible and Near-infrared Components, Agricultural and Forest Meteorology, 34, 205–213, doi:10.1016/0168-1923(85)90020-6, 1985.

35 Whalley, L., Stone, D., George, I., Mertes, S., van Pinxteren, D., Tilgner, A., Herrmann, H., Evans, M., and Heard, D.: The influence of clouds on radical concentrations: observations and modelling studies of HO_x during the Hill Cap Cloud Thuringia (HCCT) campaign in 2010, Atmospheric Chemistry and Physics, 15, 3289–3301, 2015.

- Wild, O. and Akimoto, H.: Intercontinental transport of ozone and its precursors in a three-dimensional global CTM, Journal of Geophysical Research: Atmospheres, 106, 27729–27744, 2001.
- Wild, O., Fiore, A. M., Shindell, D. T., Doherty, R. M., Collins, W. J., Dentener, F. J., Schultz, M. G., Gong, S., MacKenzie, I. A., Zeng, G., Hess, P., Duncan, B. N., Bergmann, D. J., Szopa, S., Jonson, J. E., Keating, T. J., and Zuber, A.: Modelling future changes in surface ozone: a parameterized approach, Atmos. Chem. Phys., 12, 2037–2054, doi:10.5194/acp-12-2037-2012, 2012.
- Wu, S., Mickley, L. J., Jacob, D. J., Logan, J. A., Yantosca, R. M., and Rind, D.: Why are there large differences between models in global budgets of tropospheric ozone?, J. Geophys. Res.-Atm., 112, doi:10.1029/2006JD007801, 2007.

5

- Yang, X., Pyle, J. A., and Cox, R. A.: Sea salt aerosol production and bromine release: Role of snow on sea ice, Geophysical Research Letters, 35, 2008.
- 10 Zhang, K., O'donnell, D., Kazil, J., Stier, P., Kinne, S., Lohmann, U., Ferrachat, S., Croft, B., Quaas, J., Wan, H., et al.: The global aerosolclimate model ECHAM-HAM, version 2: sensitivity to improvements in process representations, Atmospheric Chemistry and Physics, 12, 8911–8949, 2012.