Multi-model simulations of the impact of international shipping on Atmospheric Chemistry and Climate in 2000 and 2030


¹DLR, Institut für Physik der Atmosphäre, Oberpfaffenhofen, Germany
²University of Edinburgh, School of GeoSciences, Edinburgh, UK
³European Commission, Joint Research Centre, Institute for Environment and Sustainability, Ispra, Italy
⁴Max Planck Institute for Chemistry, Mainz, Germany
⁵Met Office, Exeter, UK
⁶University of Oslo, Department of Geosciences, Oslo, Norway
⁷Laboratoire des Sciences du Climat et de l’Environnement, Gif-sur-Yvette, France
⁸University of Bremen, Institute for Environmental Physics, Bremen, Germany
⁹Goddard Earth Science & Technology Center (GEST), Maryland, Washington, DC, USA
¹⁰Frontier Research Center for Global Change, JAMSTEC, Yokohama, Japan
¹¹Royal Netherlands Meteorological Institute (KNMI), Atmospheric Composition Research, De Bilt, the Netherlands

* now at: University of Cambridge, Centre for Atmospheric Science, Cambridge, UK

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Abstract. The global impact of shipping on atmospheric chemistry and radiative forcing, as well as the associated uncertainties, have been quantified using an ensemble of state-of-the-art atmospheric chemistry models and a predefined set of emission data. The analysis is performed for present-day conditions (year 2000) and for two future ship emission scenarios. In one scenario ship emissions stabilize at 2000 levels; in the other ship emissions increase with a constant annual growth rate of 2.2% up to 2030 (termed the “Constant Growth Scenario” (CGS)). Most other anthropogenic emissions follow the IPCC (Intergovernmental Panel on Climate Change) SRES (Special Report on Emission Scenarios) A2 scenario, while biomass burning and natural emissions remain at year 2000 levels. An intercomparison of the model results with observations over the Northern Hemisphere (25°–60° N) oceanic regions in the lower troposphere showed that the models are capable to reproduce ozone (O₃) and nitrogen oxides (NOₓ=NO+NO₂) reasonably well, whereas sulphur dioxide (SO₂) in the marine boundary layer is significantly underestimated. The most pronounced changes in annual mean tropospheric NO₂ and sulphate columns are simulated over the Baltic and North Seas. Other significant changes occur over the North Atlantic, the Gulf of Mexico and along the main shipping lane from Europe to Asia, across the Red and Arabian Seas. Maximum contributions from shipping to annual mean near-surface O₃ are found over the North Atlantic (5–6 ppbv in 2000; up to 8 ppbv in 2030). Ship contributions to tropospheric O₃ columns over the North Atlantic and Indian Oceans reach 1 DU in 2000 and up to 1.8 DU in 2030. Tropospheric O₃ forcings due to shipping are 9.8±2.0 mW/m² in 2000 and 13.6±2.3 mW/m² in 2030. Whilst increasing O₃, ship NOₓ simultaneously enhances hydroxyl radicals over the remote ocean, reducing the global methane lifetime by 0.13 yr in 2000, and by up to 0.17 yr in 2030, introducing a negative radiative forcing. The models show future increases in NOₓ and O₃ burden which scale almost linearly with increases in NOₓ emission totals. Increasing emissions from shipping would significantly counteract the benefits derived from reducing SO₂ emissions from all other anthropogenic sources under the A2 scenario over the continents, for example in Europe. Globally, shipping contributes 3% to increases in O₃ burden between 2000 and 2030, and 4.5% to increases in sulphate under A2/CGS. However, if future ground based emissions follow a more stringent scenario, the relative importance of ship emissions will increase. Inter-model differences in the simulated O₃ contributions from ships are significantly smaller than estimated uncertainties stemming from the ship emission inventory, mainly the ship emission totals, the distribution of the emissions over the globe, and the neglect of ship plume dispersion.

Correspondence to: V. Eyring
(Veronika.Eyring@dlr.de)

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1 Introduction

Seagoing ships emit exhaust gases and particles into the marine boundary layer contributing significantly to the total budget of anthropogenic emissions from the transportation sector (e.g. Olivier et al., 2001; Eyring et al., 2005a). Emissions of nitrogen oxides \((\text{NO}_x=\text{NO+NO}_2)\) from shipping lead to tropospheric ozone \((\text{O}_3)\) formation and perturb the hydroxyl radical \((\text{OH})\) field, and hence the lifetime of methane \((\text{CH}_4)\). These changes affect the Earth’s radiation budget as \(\text{O}_3\) and \(\text{CH}_4\) are greenhouse gases. Evidence for the importance of ship emissions comes from satellite observations from GOME (Beirle et al., 2004) and SCIAMACHY (Richter et al., 2004) that show enhanced tropospheric \(\text{NO}_2\) columns along the major international shipping routes in the Red Sea and over the Indian Ocean. A number of atmospheric model studies quantifying the impact of ship emissions on the chemical composition of the atmosphere and on climate have been published in recent years. All these studies used a global fuel consumption of about 160 million metric tons \((\text{Mt}, \text{or Tg})\) per year derived from energy statistics (Corbett and Fischbeck, 1997; Corbett et al., 1999; Olivier et al., 2001; Endresen et al., 2003). However, there is an ongoing discussion on the present-day value, as recent estimates of the fuel consumption calculated with an activity-based approach indicate higher fuel consumption of around 280 Mt (Corbett and Köhler, 2003; Eyring et al., 2005a), suggesting that previous model studies may have significantly underestimated emissions. Despite this, these models tend to overestimate oceanic \(\text{NO}_x\) over parts of the North Atlantic and the Pacific (Lawrence and Crutzen, 1999; Kasibhatla et al., 2000; Davis et al., 2001; Endresen et al., 2003). One possibility to reduce this discrepancy might be to account for ship plume dispersion in global models (Kasibhatla et al., 2000; Davis et al., 2001; Song et al., 2003; von Glasow et al., 2003; Chen et al., 2005). In ship plumes the lifetime of \(\text{NO}_x\) is significantly reduced compared to the background and \(\text{NO}_x\) in the plume is rapidly oxidised. Some \(\text{NO}_x\) is therefore lost at scales smaller than the typical grid size of global models \((100–500 \text{ km})\). These sub-grid-scale processes have to be parameterised in global models, e.g., by the use of effective emission indices depending on the meteorology and background conditions. However, there is no clear consensus on the effective global emissions from ships. In this study we use relatively low global emission estimates and neglect ship plume chemistry.

In addition to \(\text{NO}_x\), shipping contributes significantly to global sulphur dioxide \((\text{SO}_2)\) emissions as the average sulphur content of the fuel burned in marine diesel engines of 2.4% is high compared to other transport sectors (EPA, 2002). In large areas of the Northern Hemisphere \(\text{SO}_2\) emissions from ships are comparable to biogenic dimethyl sulphide \((\text{DMS})\) emissions, which are the main natural source of sulphur over the oceans (Corbett et al., 1999; Capaldo et al., 1999; Derwent et al., 2005). The main oxidation pro-

cesses for \(\text{SO}_2\) are believed to be either via \(\text{OH}\) in the gas phase or in the liquid phase via \(\text{O}_3\) or \(\text{H}_2\text{O}_2\) in cloud droplets (Langner and Rohde, 1991). As \(\text{SO}_2\) is primarily controlled by aqueous processes, it is expected that \(\text{SO}_2\) is largely independent of enhanced \(\text{OH}\) in the plume (Davis et al., 2001). In addition to sulphate particles resulting from \(\text{SO}_2\) emissions, ships also release black carbon \((\text{BC})\) and particulate organic matter \((\text{POM})\). The increase in sulphate, \(\text{BC}\) and \(\text{POM}\) concentrations has a direct and indirect effect on climate. The direct effect results from enhanced scattering of solar radiation (Haywood and Shine, 1995). The indirect effect of particles from shipping results from changes in the microphysical, optical and radiative properties of low marine clouds (Scorer, 1987; Capaldo et al., 1999; Durkee et al., 2000; Schreier et al., 2006).

The numbers of ships in the world merchant fleet has increased by 35% over the past 50 years, accompanied by a significant increase in emission totals (Eyring et al., 2005a). At the end of the year 2001 it consisted of around 90 000 oceangoing ships of 100 gross tons \((\text{GT})\) and above (Lloyd’s, 2002). Shipping is currently one of the less regulated sources of anthropogenic emissions with a high reduction potential through technological improvements, alternative fuels and ship modifications. Emission scenario calculations up to the year 2050 show that if no control measures are taken beyond existing International Maritime Organization \((\text{IMO})\) regulations \((\text{IMO}, 1998)\), \(\text{NO}_x\) emissions might increase with an annual growth rate of 1.7% between 2000 and 2030 and up to a value of present day global road transport by 2050 \((38.8 \text{Tg} \ (\text{NO}_2)/\text{yr})\) (Eyring et al., 2005b). If the sulphur content remains at present day levels a doubling of ship \(\text{SO}_2\) emissions can be expected. However, given the air quality issue of shipping emissions, further emission reductions of total \(\text{NO}_x\) and \(\text{SO}_2\) emissions are likely. Using aggressive \(\text{NO}_x\) emission reduction technologies, a significant decrease up to 85% of today’s \(\text{NO}_x\) emissions could be reached through technological improvements by 2050, in spite of a growing fleet. A summary of current national and international maritime regulations is given in Eyring et al. (2005b).

Currently there is a large uncertainty about the overall impact of emissions from international shipping which can be explored using global atmospheric models. The first key question addressed in this study is how \(\text{NO}_x\) and \(\text{SO}_2\) emissions from international shipping might influence atmospheric chemistry, in particular tropospheric \(\text{O}_3\) and sulphate, in the next three decades, if these emissions increase unabated. To address this, impacts of \(\text{NO}_x\) and \(\text{SO}_2\) emissions from international shipping are assessed with the help of an ensemble of state-of-the-art global atmospheric chemistry models. The second major issue is to examine the range of results given by the individual models compared to the ensemble mean to estimate the uncertainties introduced by different modelling approaches. The participating models have also been evaluated and used in accompanying studies (e.g. Stevenson et al., 2006; Dentener et al., 2002).
The models and model simulations, together with the method to analyse the results are described in Sect. 2. The models' ability to simulate O$_3$, NO$_x$ and SO$_2$ over the remote ocean is evaluated in Sect. 3.1. Large-scale chemistry effects on NO$_2$ and O$_3$ distributions due to NO$_x$ emissions from ships are discussed in Sect. 3.2 for present-day conditions and in Sect. 3.3 for 2030, while impacts of SO$_2$ emissions from ships on sulphate distributions are discussed in Sect. 3.4. In Sect. 3.5 radiative forcings (RFs) from tropospheric O$_3$ calculated with the help of an offline radiation scheme are summarised and RFs due to carbon dioxide (CO$_2$), CH$_4$, and sulphate are roughly estimated. Section 4 discusses the impact of volatile organic compounds (VOC) and carbon monoxide (CO) emissions from ships and possible uncertainties in the presented results mainly stemming from the emission inventory itself, the neglect of plume

2006a,b; Shindell et al., 2006, van Noije et al., 2006) as part of the European Union project ACCENT (Atmospheric Composition Change: the European NeTwork of excellence; http://www.accent-network.org).

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chemsitry and assumptions in the future scenarios for background as well as ship emissions. Section 5 closes with a summary and conclusions.

2 Models and model simulations

2.1 Participating Models

Ten global atmospheric chemistry models have participated in this model inter-comparison. Seven of the ten models are Chemistry-Transport Models (CTMs) driven by meteorological assimilation fields and three models are atmospheric General Circulation Models (GCMs). Two of the GCMs are driven with the dynamical fields calculated by the GCM in climatological mode, but the fully coupled mode (interaction between changes in radiatively active gases and radiation) has been switched off in the simulations of this study. The other GCM runs in nudged mode, where winds and temperature fields are assimilated towards meteorological analyses. Therefore, changes in the chemical fields do not influence the radiation and hence the meteorology in any of the model simulations used here, and for a given model each scenario is driven by identical meteorology. The main characteristics of the ten models are summarised in Table 1 and the models are described in detail in the cited literature.

The horizontal resolution ranges from 5.6°×5.6° (MATCH-MPIC) to 2.8°×2.8° (CHASER-CTM, FRSGC/UCI, UIO.CTM2) and 3°×2° (TM4). The vertical resolution varies in terms of number of vertical layers and upper boundary and ranges from 9 layers with a model top at 100 hPa (STOChem-HadAM3) to 52 layers with a model top at 0.006 hPa (GMI/CCM3).

All models use detailed tropospheric chemistry schemes even though notable differences exist between the models, e.g. in the hydrocarbon chemistry. Of the ten models, four included the tropospheric sulphur cycle (CHASER-CTM, STOChem-HadAM3, STOChem-HadGEM, and TM4). These models include anthropogenic and natural sulphur sources, and account for oxidation in the gas phase by OH, and in the aqueous phase (in cloud droplets) by H2O2 and O3. Sulphate aerosol is predominantly removed by wet-deposition processes. Models also differ in the parameterisation of sub-grid scale convection, the representation of cloud and hydrological processes, dry and wet deposition, and boundary layer mixing and a variety of different advection schemes are used.

Differences in the main characteristics as listed above will lead to differences in the modelled response to NOx and SO2 emissions from shipping, even if near-identical forcings (e.g., sea surface temperatures, natural and anthropogenic emissions, and meteorology) are used. A more detailed discussion of the sources of differences between the models is included in Stevenson et al. (2006).

2.2 Model Simulations

Two of the five simulations that have been defined as part of the wider PHOTOCOMP-ACCENT-IPCC study have been used in this work: a year 2000 base case (S1) and a year 2030 emissions case (S4) following the IPCC (Intergovernmental Panel on Climate Change) SRES (Special Report on Emission Scenarios) A2 scenario (IPCC, 2000). Full details on the emissions used in the S1 and S4 simulations are summarised in Stevenson et al. (2006) and only the key aspects for this study are given here. All models used the same anthropogenic and biomass burning emissions, but variable natural emissions. For example, for NOx, natural sources from soils and lightning differ between the models, but this has little impact over the oceans. For SO2, natural sources from volcanoes and oceanic DMS differ. The latter has some impact on absolute values of SO2 over the oceans, but little impact on changes due to ships. The differences in natural emissions are not thought to be a major source of inter-model differences in the results presented here. To retain consistency with all other emissions, ship emissions in the year 2000 (S1) are based on the EDGAR3.2 dataset (Olivier et al., 2001) at a spatial resolution of 1° latitude×1° longitude. The global distribution of ship emissions in EDGAR3.2 is based on the world’s main shipping routes and traffic intensities (Times

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Table 2. Specified global annual anthropogenic (not including biomass burning emissions) surface emission totals for each scenario.

<table>
<thead>
<tr>
<th>Name</th>
<th>Meteorology</th>
<th>Emissions</th>
<th>Total</th>
<th>From Shipping</th>
<th>Total</th>
<th>From Shipping</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>NOx (Tg N)</td>
<td></td>
<td></td>
<td>SOx (Tg S)</td>
<td></td>
</tr>
<tr>
<td>S1</td>
<td>2000</td>
<td>2000 (EDGAR3.2)</td>
<td>27.80</td>
<td>3.10</td>
<td>54.00</td>
<td>3.88</td>
</tr>
<tr>
<td>S1w</td>
<td>2000</td>
<td>2000 (EDGAR3.2), but without ship emissions</td>
<td>24.70</td>
<td>0.00</td>
<td>50.12</td>
<td>0.00</td>
</tr>
<tr>
<td>S4</td>
<td>2000</td>
<td>2030 SRES A2, but with ship emissions of the year 2000</td>
<td>54.60</td>
<td>3.10</td>
<td>100.00</td>
<td>3.88</td>
</tr>
<tr>
<td>S4s</td>
<td>2000</td>
<td>2030 SRES A2, Traffic A2s; Ship emissions increase with a flat increase of 2.2%/yr compared to the year 2000</td>
<td>57.45</td>
<td>5.95</td>
<td>103.48</td>
<td>7.36</td>
</tr>
<tr>
<td>S4w</td>
<td>2000</td>
<td>2030 SRES A2, but without ship emissions</td>
<td>51.50</td>
<td>0.00</td>
<td>96.12</td>
<td>0.00</td>
</tr>
</tbody>
</table>
Books, 1992; IMO, 1992). EDGAR3.2 includes data for 1995, which have been scaled to 2000 values assuming a growth rate of 1.5%/yr, resulting in annual NOx and SO2 emissions of 3.10 Tg(N) and 3.88 Tg(S), respectively, similar to the emission totals published by Corbett et al. (1999). Table 2 summarises the global annual anthropogenic surface emission and ship emission totals for NOx and SO2. In the 2000 simulation (S1), ship emissions account for about 11.2% of all anthropogenic surface nitrogen oxide emissions and for about 7.2% of all anthropogenic sulphur emissions. Other emissions from ships such as CO, particulate matter, CH4 and VOC were not included in this inventory and were therefore not considered in the reference simulations. The effect of CO and VOC emissions from ships is quantified with the help of sensitivity simulations carried out with a single model (MATCH-MPIC) in Sect. 4.1.2.

As noted in Stevenson et al. (2006) in the S4 simulation emissions from ships were included at year 2000 levels by mistake. All other anthropogenic sources (except biomass burning emissions, which remain fixed at year 2000 levels) vary according to A2 broadly representing a pessimistic future situation. The simulation S4 is used here to assess the impact of ship emissions under different background levels. An additional model simulation for 2030 (S4s) has been designed to assess the impact of shipping if emission growth remains unabated. Ship emissions in S4s are based on a “Constant Growth Scenario” (CGS) in which emission factors are unchanged and emissions increase with an annual growth rate of 2.2% between 2000 and 2030. In the S4s scenario emissions from shipping increase to 5.95 Tg(N) for NOx and 7.36 Tg(S) for SO2 in 2030. As an example, the global distributions of surface NOx emissions for the years 2000 (S1) and 2030 (S4s) are displayed in Fig. 1. Vessel traffic distributions are assumed to stay the same for all model simulations presented here.

To assess the impact of ship emissions on chemistry and climate in 2000 and 2030, two sensitivity simulations have been defined that use identical conditions to S1 or S4/S4s except that ship emissions are excluded. The year 2000 and 2030 experiments without ship emissions are denoted as S1w and S4w.

All 2030 model experiments (S4, S4s and S4w) are driven by the same meteorological data as the 2000 simulations (S1 and S1w). Global methane mixing ratios have been specified across the model domain (1760 ppbv in 2000 and 2163 ppbv in 2030) to save time spinning up the models and to help constrain the results (for details see Stevenson et al., 2006). Nine of the ten models performed single year simulations with spin-ups of at least 3 months and one model (STOCHEM-HadGEM) performed a four year simulation. For the multi-annual simulations the results have been averaged over all four years to reduce the effects of inter-annual variability.

### 2.3 Model Analyses

Each model provided output for 3-D monthly mean NOx, O3 and sulphate mixing ratios as well as the mass of each grid-box on the native model grid for all five simulations. As models used a variety of vertical co-ordinate systems and resolutions, the output has been converted to the same common vertical grid as used in Stevenson et al. (2006). Model results are interpolated to the 19 hybrid (sigma-pressure) levels of the Met Office HadAM3 model where up to 14 of these levels span the troposphere. Results were also interpolated to a common horizontal resolution of 5°×5°. To calculate the ensemble mean on the common grid the simulated fields have been masked at the chemical tropopause (O3 =150 ppbv) similar to the method applied in Stevenson et al. (2006). For the 2000 simulations (S1 and S1w) we apply a consistent mask by using the O3 field from the S1w simulation for each model and all species. For the 2030 simulations (S4, S4s, and S4w) the S4w O3 field is used to mask the tropopause. To calculate global tropospheric burdens,
The models have been evaluated in accompanying studies. For example, \( \text{O}_3 \) fields have been compared to ozonesonde measurements (Stevenson et al., 2006), \( \text{NO}_2 \) columns have been compared to three state-of-the-art retrievals from measurements of the Global Ozone Monitoring Experiment (GOME) (van Noije et al., 2006), \( \text{CO} \) has been compared to near-global observations from the MOPITT instrument and local surface measurements (Shindell et al., 2006), deposition budgets have been compared to nearly all information on wet and dry deposition available worldwide (Dentener et al., 2006).
et al., 2006b), and finally modelled surface O$_3$ fields have been compared to observations from various measurement sites (Dentener et al., 2006a). In all these studies, the ensemble mean was among the best when comparing to measurements. In this study we build on the model evaluation work listed above as well as on the individual evaluation of the models and make the assumption that the models produce reasonable simulations of the key chemical species. We use the ensemble mean to assess large-scale chemistry effects resulting from ship emissions for the present day and in the future.

In addition to the above mentioned studies, we evaluate here the models’ ability to simulate O$_3$, NO$_x$ and SO$_2$ in the lower troposphere (<3 km) as well as tropospheric NO$_2$ columns over the ocean by comparing the results from the S1 present-day reference simulations of the individual models to aircraft measurements (Emmons et al., 2000; Singh et al., 2006) and to satellite data from SCIAMACHY (Richter et al., 2004).

Tropospheric data from a number of aircraft campaigns have been gridded onto a 5° longitude by 5° latitude grid by Emmons et al. (2000), with additional data from more recent campaigns (see http://gctm.acd.ucar.edu/data), up to and including TRACE-P in 2001 (we subsequently refer to this data as “Emmons et al. (2000) plus updates”). We compare data over the Northern Hemisphere (25°–60° N) oceanic regions in Fig. 2. Most models reproduce the observed annual and seasonal O$_3$ means in the lower troposphere (<3 km). Two models (GMI/CCM3 and GMI/DAO) simulate O$_3$ values larger than observed in autumn and winter, and MATCH-MPIC overestimates observed O$_3$ in winter. The oceanic 25°–60° N means in observations and models shown in Fig. 2 are averages over only those grid boxes that include observations for the particular species (see Fig. 3). To compare model results with observations in the Pacific and Atlantic, we use data collected during the INTEX-NA campaign in July/August 2004 in addition (see http://www-air.larc.nasa.gov/; Singh et al., 2006). The good agreement between modelled and observed O$_3$ in July holds for both the Pacific and the Atlantic (Fig. 4).

In winter and autumn, the two STOCHEM models, both using the same chemistry scheme, simulate the highest NO$_x$ values among all models (Fig. 2). An additional analysis showed that in these models in these seasons, high NO$_x$ plumes from land-based anthropogenic sources were extending sufficiently far out to sea to affect the nearest oceanic data comparison points. In addition, the treatment of heterogeneous NO$_x$ loss in the STOCHEM models is very simple, and may be leading to unrealistically long NO$_x$ lifetimes in winter and autumn. For all other models the simulated NO$_x$ and NO$_2$ (not shown) lies within one standard deviation (1σ) of the observational mean in all seasons with values above the observational mean in autumn and below in all other seasons as well as in the annual mean (Fig. 2). Nearly all observations for NO$_x$ over the Northern Hemisphere oceanic regions below 3 km in the Emmons et al. (2000) plus updates data set are made over the Pacific (Fig. 3). Compared to the INTEX-NA campaign, the simulated NO$_2$ lies within 1σ of the observational mean over the Atlantic and the Pacific (Fig. 4). This overall reasonable agreement between models and observations for NO$_x$ and NO$_2$ differs from the findings of Kasibhatla et al. (2000) and Davis et al. (2001), who found that their models overpredicted NO$_x$ in the Atlantic and Pacific marine boundary layer. These authors used similar ship emissions, but compared with somewhat different observational data. Kasibhatla et al. (2000) used NARE campaign data from the North Atlantic (37°–50° N; 35°–50° W) in September 1997 and Davis et al. (2001) used data from five campaigns over the North Pacific.

SO$_2$ in the four models that included a sulphur cycle generally lie within 1σ of the observations in Fig. 2, but there is little data outside the winter season, and the standard deviations in the marine boundary layer are large, making it difficult to assess the models. One of the models (STOCHEM-HadAM3) is consistently high, particularly in the free troposphere – this is probably related to the relatively large magnitude and elevated source of volcanic SO$_2$ in this model. Standard deviations are smaller in the summertime INTEX-NA data set over the Pacific (Fig. 4). All models significantly underestimate observed SO$_2$ in July, in agreement with previous findings (Davis et al., 2001).

Kasibhatla et al. (2000) and Davis et al. (2001) suggested

![Fig. 3. Geographical distribution of the number of observations in the Emmons et al. (2000) plus updates data set for O$_3$, NO$_x$, and SO$_2$ in the Northern Hemisphere (25°–60° N) oceanic regions in the altitude range 0–3 km. N1 gives the total number of individual observations and N2 the total number of 5°×5° grid boxes that include at least one observation.](http://www.atmos-chem-phys.net/7/757/2007/)
that ship plume chemistry may partly explain the discrepancy between observed and simulated NO\textsubscript{x}. While the neglect of ship plume chemistry in global models might contribute to differences between observations and models, we show here that the selection of data the models are compared to is also very important: if we reduce the Emmons et al. (2000) plus updates and the INTEX-NA data set and only compare the model results for NO\textsubscript{x} with the data used in Davis et al. (2001) we can reproduce their results. However, if we use the entire observational data set (Fig. 2), which is more widespread and therefore more robust (up to 32 campaigns between 1983 and 2001, including the five Pacific datasets, but not including the Atlantic NARE data) we find good agreement with observed NO\textsubscript{x}. In addition, differences in the modelled and observed NO\textsubscript{x} and SO\textsubscript{2} fields cannot be fully attributed to ship emissions. In particular, the comparison with the Emmons et al. (2000) plus updates and the INTEX-NA data sets leads to only small differences between the S1 and the S1\textsubscript{w} ensemble mean (Figs. 2 and 4), so that the shipping signal cannot be properly evaluated. Therefore, it remains unclear whether the underestimation of SO\textsubscript{2} in the marine boundary layer in the models is due to the ship emission inventory or due to the underestimation of another source (e.g. DMS), overestimation of a sink (e.g. oxidation),
errors associated with simulating specific events (e.g. horizontal/vertical transport and mixing), or a combination of several factors. More measurements are needed before final conclusions on the quality of the models to simulate the ship emission response can be made (see also Sect. 5).

Another approach to evaluate the shipping response in models is the use of satellite data. Recently, enhanced tropospheric NO$_2$ columns have been observed over the Red Sea and along the main shipping lane to the southern tip of India, to Indonesia and north towards China and Japan (Beirle et al., 2004; Richter et al., 2004). Here we intercompare the tropospheric NO$_2$ columns derived from SCIAMACHY nadir measurements from August 2002 to April 2004 (Richter et al., 2004) to the models' ensemble mean in 2000 (S1 simulation). The ensemble mean in Fig. 5 consists of the eight models that provided tropospheric NO$_2$ columns at 10:30 a.m. local time, which is close to the overpass time of the ERS-2 satellite. The two STOCHEM models provided output only as 24 h mean, and are not included. To compare the model data to the satellite measurements, individual model results and SCIAMACHY data were interpolated to a common grid ($0.5^\circ \times 0.5^\circ$). The models simulate similar tropospheric columns at 10.30 local time for 2000. The ensemble mean comprises 8 out of the 10 models. Individual model results and SCIAMACHY data were interpolated to a common grid ($0.5^\circ \times 0.5^\circ$).

Fig. 5. NO$_x$ signature of shipping in the Indian Ocean (5° S to 40° N and 25° E to 125° E). Left: Tropospheric NO$_2$ columns derived from SCIAMACHY data from August 2002 to April 2004 (Richter et al., 2004). Right: Ensemble mean NO$_2$ tropospheric columns at 10.30 local time for 2000. The ensemble mean comprises 8 out of the 10 models. Individual model results and SCIAMACHY data were interpolated to a common grid ($0.5^\circ \times 0.5^\circ$).

stronger than emissions from shipping. Thus, the shipping signal cannot be identified in the coarse resolution model data in this region. From this point of view, an intercomparison over the remote ocean (e.g. over the Atlantic) with satellite data far away from any land source would be preferable. However, up to now, no statistically significant satellite data on ship emissions are available for remote oceans. This is mainly due to the distributed nature of the emissions which leads to dilution and makes it difficult to distinguish the shipping signal from the effect of long-range transport from the US towards Europe. Increasing ship emission in the future should make detection possible, in particular if data from an instrument with good spatial coverage such as OMI or GOME-2 is used in combination with model calculations of the contribution of long range transport. However, unambiguous identification and quantification will always be more difficult than in the special case between India and Indonesia with its unique emission pattern.

3.2 Large-scale chemistry effects of NO$_x$ ship emissions in 2000

To examine the range of results given by the individual models compared to the ensemble mean, Fig. 6 shows differences in annually averaged zonal mean O$_3$ versus height between the 2000 base case simulation (S1) and the model simulation without shipping (S1w) for each model and the ensemble mean. Standard deviations are shown in addition indicating regions of large intermodel variability. All models show the maximum contributions from ships in zonal annual mean near-surface O$_3$ in northern mid-latitudes between 10° N and 55° N, where most of the global ship emissions are released into the atmosphere (see Fig. 1). A rapid decrease of the impact on O$_3$ distributions with height is simulated. There are notable differences in the magnitude of the response to ship emissions between the individual models which can be attributed to differences in the main
The main conclusions in the subsequent sections are based on the models’ ensemble mean.

Due to the short lifetime of NO$_2$ in the boundary layer, near-surface changes in NO$_2$ due to ship emissions (S1-S1w) follow closely the main shipping routes, but the dispersion is a few hundred kilometres, which is partly due to coarse resolutions of the models, but also real transport (Fig. 7). Maximum changes of 2.3 ppbv are found over the Baltic Sea in both months. In the English Channel and along the west coast of Europe (from Ireland to Morocco) NO$_2$ changes are also significant (around 0.5 ppbv). Enhanced NO$_2$ levels up to 0.2 ppbv are also simulated over large areas of the Atlantic and e.g. in the Red Sea and along the main shipping lane to the southern tip of India, to Indonesia and north towards China and Japan as well as in the Gulf of Mexico. In July NO$_2$ changes over the Baltic and over the Atlantic are smaller than in January and cover a smaller area, as the lifetime of NO$_3$ in the summer months is shorter due to higher chemical activity. Relative changes in NO$_2$ strongly depend on the level of NO$_3$ from other sources. For example, even
Fig. 7. Modelled ensemble mean near-surface NO\(_2\) and O\(_3\) change between case S1 (year 2000) and S1w (year 2000 without ship emissions) in January (left) and July (right). (a–b): absolute changes in near-surface NO\(_2\); (c–d): relative changes in near-surface NO\(_2\); (e–f) absolute changes in near-surface O\(_3\); (g–h): relative changes in near-surface O\(_3\).

though absolute NO\(_2\) changes over the Baltic and the North Sea are large, relative changes are less pronounced because of high background NO\(_2\) levels. The largest relative changes of up to 80% in January and 96% in July are found over the remote Atlantic and along the major shipping lanes, where background levels are small and NO\(_x\) emissions from ships are the dominant source (see Fig. 1).

Substantial differences are simulated between the near-surface January and July ensemble mean O\(_3\) changes despite the fact that there are no seasonal variations in the ship emission inventory (Fig. 7, lower panels). The reason for the difference is that during winter additional NO\(_x\) emissions from shipping can lead to O\(_3\) titration in the highly polluted regions over the continents, whereas in summer the additional NO\(_x\) leads to O\(_3\) production. O\(_3\) concentrations in January show a decrease of around 1 ppbv with maximum decreases of up to 2.8 ppbv over the Baltic. The changes in July over this region are rather small but positive. The largest increases in January are found southward of 30°N, where the available solar radiation is sufficient for O\(_3\) production to outweigh direct removal of NO. Most pronounced increases of 3.3 ppbv are simulated over the Indian Ocean and along the main shipping lane west of the coast of Southern Africa. The simulated large decrease in O\(_3\) over a large area in Europe in winter, which has not been reported in e.g. the Endresen et al. (2003) study, might be due to high vessel traffic densities in particular over the Baltic Sea in the inventory used here. Vessel traffic densities derived from different sources such as AMVER (Automated Mutual-assistance Vessel Rescue system) or the Purple Finder (PF) data set (available at http://www.purplefinder.com) report fewer observations there (Endresen et al., 2003), see further discussion in Sect. 4.1.1. In July, the geographical pattern in O\(_3\) changes is similar to the pattern in Lawrence and Crutzen (1999) showing most pronounced changes over wide areas of the Atlantic up to 12 ppbv (approx. 35%) and over the
Fig. 8. Modelled ensemble mean O₃ change between (a–c) case S1 (year 2000) and S1w (year 2000 without ship emissions), (d–f) case S4 (year 2030) and S4w (year 2030 without ship emissions), and (g–i) case S4s (year 2030) and S4w. Figures 8a, 8d, and 8g are zonal mean changes (ppbv), Figs. 8b, 8e, and 8h are near-surface O₃ changes (ppbv) and Figs. 8c, 8f, and 8i are tropospheric O₃ column changes (DU).

Western and Northern Pacific up to 5 ppbv (approx. 25%). Changes of the order of 5 ppbv (approx. 10%) are also simulated over the Indian Ocean. The simulated changes over the Atlantic and the Indian Ocean are in good agreement with results reported in Endresen et al. (2003), but smaller over the Pacific. Differences between the two studies are likely related to the difference in vessel traffic densities (see Sect. 4.1.1). Due to the longer lifetime of O₃ compared to NOₓ in the boundary layer, O₃ changes are less strictly confined to the main shipping lanes, and thus affect larger areas. Non-linear effects of O₃ photochemistry are significant. For example, over the Baltic and the North Sea, where background NOₓ levels are relatively high, changes in O₃ are comparatively small, whereas they are substantial over more remote areas.

3.3 Large-scale chemistry effects of NOₓ ship emissions in 2030

3.3.1 Ozone distributions

Figure 8 shows modelled ensemble mean O₃ changes due to ship emissions for the year 2000 (S1-S1w) and for two different scenarios in 2030 (S4-S4w; S4s-S4w). As vessel traffic densities are the same in all model simulations the 2030 results mainly show a scaling of the 2000 results, but non-linearity effects also play a role. O₃ changes versus height in 2000 (Fig. 8a) have already been discussed in Sect. 3.1. The largest O₃ response is found near the surface between 10°–55° N, and rapidly decreases with altitude. Keeping emissions from shipping at 2000 levels but with all other emissions increasing, the pattern in zonal mean changes remains the same but the impact is slightly less than in 2000 due to lower O₃ production rates under the influence of higher background of NOₓ concentrations (Fig. 8d; S4-S4w). Under the S4s scenario near-surface O₃ changes of about 1.7 ppbv are simulated in the zonal mean between 10°–55° N, and even in the free troposphere changes in O₃ reach up to 1.3 ppbv (Fig. 8g). Changes in the annual mean near-surface level reach 5.5 ppbv over the Atlantic (Fig. 8h) and increase up to 7.4 ppbv in the S4s scenario in 2030 (Fig. 8h). Over Northern Europe where there are high levels of NOₓ, the increase in NOₓ from shipping decreases, rather than increases, O₃ levels. This is due to reduction in oxidant levels as OH is removed by the reaction NO₂ + OH -> HNO₃, and in winter to direct titration of O₃ by NO₂. The effect is stronger in the 2030 scenarios due to the increased background level of NOₓ in Northern Europe, with shipping decreasing O₃ by 3 ppbv in the S4s scenario.

The most pronounced changes in tropospheric O₃ columns are found over the Indian Ocean (1.16 DU in 2000 and 1.72 DU in 2030), related to the higher tropopause there and to more effective transport of O₃ from the boundary into the upper troposphere. A second peak is simulated over the Atlantic (Figs. 8c,f,i).
3.3.2 Linearities in NO\textsubscript{x} and O\textsubscript{3} burden response to ship emissions

Changes in annual mean NO\textsubscript{x} and O\textsubscript{3} burdens are calculated for the two 2030 scenarios (S4-S4w; S4s-S4w) as described in Sect. 2.3 for four different regions (global mean, Atlantic Ocean, Baltic Sea, and Indian Ocean). A linear regression is performed for the changes in NO\textsubscript{x} and O\textsubscript{3} burdens due to shipping over the scenarios S4-S4w, S4s-S4w and the origin (zero NO\textsubscript{x} ship emissions / zero changes in NO\textsubscript{x} and O\textsubscript{3} burdens). The changes in global tropospheric NO\textsubscript{x} burden associated with these two scenarios show a fairly linear relationship for all models (Fig. 9, left), and a doubling of NO\textsubscript{x} emissions approximately results into a doubling of the mean NO\textsubscript{x} burden.

For O\textsubscript{3} the correlation is also broadly linear (Fig. 9, right). Only small saturation effects are visible as the O\textsubscript{3} burden for the low emission scenario (3.10 Tg(N)) lies slightly above the multi-regression line whereas the one for the high emission scenario (5.95 Tg(N)) lies slightly below the multi-regression line for all models. This is to be expected due to non-linearities in the O\textsubscript{3} chemistry. In contrast to the relatively small degree of saturation computed here, Labrador et al. (2004) computed a substantial saturation effect for lightning NO\textsubscript{x} emissions as they were increased from 0 to 20 Tg(N)/yr, with the saturation already becoming clearly evident between 5 and 10 Tg(N)/yr. Although there are only three data points available here (0, 3.1 and 5.95 Tg(N)/yr), there is evidence in these results that the ship NO\textsubscript{x} effect is only weakly subject to saturation in its current magnitude range, and that saturation cannot be expected to help mitigate the effects of near-future increases. Overall, similar to NO\textsubscript{x} burdens, a doubling in NO\textsubscript{x} ship emissions results in approximately a doubling in O\textsubscript{3} burdens in the ensemble mean. Note that, whereas the majority of the models (eight out of ten) show similar response, the two STOCHEM models simulate NO\textsubscript{x} burden changes around a factor of 3 (STOCHEM-HadAM3) or 6 (STOCHEM-HadGEM) higher than the other models, which explains the large standard deviations of the ensemble mean NO\textsubscript{x} burden. As discussed in Sect. 3.1, this can mainly be attributed to high NO\textsubscript{x} plumes from land-based anthropogenic sources and to the long NO\textsubscript{x} lifetimes in winter and autumn in these two models. For the annual O\textsubscript{3} burdens the two STOCHEM models show similar results compared to all other models, and the inter-model standard deviations for O\textsubscript{3} are small (<15%).

Similar to the global annual burdens, eight out of the ten models show similar response in the seasonal cycle of NO\textsubscript{x} and O\textsubscript{3} burden changes for the S4s-S4w scenario over the Atlantic Ocean, the Baltic Sea, the Indian Ocean and globally (Fig. 10). As expected, the seasonal cycle in both NO\textsubscript{x} and O\textsubscript{3} is most pronounced in the Baltic Sea (northern hemispheric mid-latitudes), whereas the amplitude of the seasonal

Fig. 9. Global total change in annual mean tropospheric NO\textsubscript{x} burden (left) and O\textsubscript{3} burden (right) due to ship emissions (S4-S4w and S4s-S4w) in each individual model (coloured lines) and the ensemble mean (black line). Inter-model standard deviations are shown as black bars.
Fig. 10. Seasonal variation in tropospheric NO\textsubscript{x} (left column) and O\textsubscript{3} (middle column) burden due to shipping for the scenario S4s-S4w in different regions: Atlantic Ocean (85°W–5°W; 15°N–60°N), Baltic Sea (10°E–30°E; 54°N–66°N), Indian Ocean (50°E–100°E; 0°N–25°N) and global.

cycle over the Indian Ocean is relatively small. Again the two STOCHEM models show significantly higher changes in NO\textsubscript{x} burdens in winter and autumn in northern mid-latitude regions (Atlantic and Baltic Sea). The change in ensemble mean O\textsubscript{3} burdens reaches peak values of about 0.8 Tg over the Atlantic in August, 0.4 Tg over the Indian Ocean in October, and 0.04 Tg over the Baltic Sea in June. The global change in NO\textsubscript{x} burden due to ship emissions as simulated by the ensemble mean in the S4s scenario is enhanced by 25 Gg in summer. The global O\textsubscript{3} burden is enhanced by about 4 Tg with peak changes in October and smallest changes in January.

3.4 Large-scale chemistry effects of SO\textsubscript{2} ship emissions in 2000 and 2030

A subset of four models (CHASER, STOCHEM-HadAM3, STOCHEM-HadGEM, and TM4) included the tropospheric sulphur cycle. The ensemble mean of these four models has been applied to quantify changes due to shipping in sulphate distributions now and in the future (Fig. 11). Maximum changes due to shipping in the ensemble mean zonal mean sulphate (SO\textsubscript{4}) distribution are located in the boundary layer of the northern mid-latitudes around 40°N. These changes result in about 30 pptv in the year 2000 simulations.
and 50 pptv in the year 2030 (S4s-S4w). With increasing height, the changes in SO₄ decrease continuously to about 3–5 pptv (2000) and 6–9 pptv (2030) in the upper troposphere. In the lowermost boundary layer over the Atlantic Ocean at the west coast of Europe and over the Baltic Sea, maximum annual sulphate changes amount to about 200 pptv in 2000 and 300 pptv in 2030. The geographical pattern shows the main shipping routes over the Atlantic Ocean between Europe and North America and over the Red Sea and the Indian Ocean between the Arabian Peninsula and India. In all other parts of the world, changes in sulphate due to SO₂ emissions from shipping remain low in general. Globally, shipping contributes with 4.5% to sulphate increases until 2030 under the A2/CGS.

### Table 3. O₃, SO₄, CO₂, and CH₄ radiative forcings due to shipping in 2000 and 2030.

<table>
<thead>
<tr>
<th>Year</th>
<th>O₃ mW/m²</th>
<th>SO₄(direct) mW/m²</th>
<th>CH₄ mW/m²</th>
<th>CO₂ mW/m²</th>
</tr>
</thead>
<tbody>
<tr>
<td>2000</td>
<td>9.8±2.0</td>
<td>-14</td>
<td>-14</td>
<td>26</td>
</tr>
<tr>
<td>2030</td>
<td>7.9±1.4</td>
<td>-13</td>
<td>-13</td>
<td>24</td>
</tr>
<tr>
<td>2030</td>
<td>13.6±2.3</td>
<td>-26</td>
<td>-21</td>
<td>46</td>
</tr>
</tbody>
</table>

#### 3.5 Radiative Forcing

O₃ distributions from all scenarios and all models were used as input for an offline radiation scheme (Edwards and Slingo, 1996), with all other parameters held constant, broadly representing the present-day atmosphere. In the stratosphere, O₃ was overwritten by a climatology, so the changes discussed here are purely tropospheric. The calculations include the radiative effects of clouds. Comparing instantaneous short-wave and long-wave radiative fluxes at the tropopause between scenarios yields O₃ radiative forcings (see Stevenson et al. (1998) for more details on the method). In order to make the O₃ forcings directly comparable to other forcings, stratospheric temperatures, which respond on timescales of a month or so to radiative perturbations, need to be adjusted until there is no change in stratospheric heating rates, the so-called “fixed dynamical heating” approximation. Stevenson
et al. (1998) found that this relaxation of stratospheric temperatures resulted in a 22% reduction in the \( \text{O}_3 \) forcing compared to the instantaneous value and we apply this as a globally constant correction. Figure 12a shows maps of multimodel ensemble mean annual mean instantaneous \( \text{O}_3 \) radiative forcings for the 2030 high emissions case (S4s) relative to the scenario without ships (S4w). Similar distributions were found for the other scenarios. The peak forcing occurs over the Indian Ocean, the site of the largest column \( \text{O}_3 \) changes (Fig. 8), but also a region with relatively high surface temperatures, and a high, cold tropopause. A secondary peak occurs over the Caribbean for similar reasons. Further north over the Atlantic the forcing is less, despite a significant \( \text{O}_3 \) change, reflecting the smaller surface-to-tropopause temperature contrast and increasing cloudiness.

Figure 12b shows the inter-model standard deviation, which is typically 15–25%. The ensemble mean forcings and standard deviations for the three cases (S1-S1w, S4-S4w and S4s-S4w), applying a 22% reduction to account for stratospheric temperature adjustment, are 9.8±2.0, 7.9±1.4 and 13.6±2.3 mW/m², respectively. The influence of ship emissions on the \( \text{O}_3 \) forcing slightly reduces as the background \( \text{O}_3 \) levels rise, but the relationship between ship \( \text{NO}_x \) emissions and resultant \( \text{O}_3 \) forcing is close to linear. Comparing with the total \( \text{O}_3 \) forcing between 2000 and 2030, as discussed in Stevenson et al. (2006), the contribution from ships in the S4s case to the global projected tropospheric \( \text{O}_3 \) forcing is 4%.

Ship \( \text{NO}_x \) emissions also affect the radiatively active gas methane, by increasing OH and reducing the methane lifetime. Five models (CHASER-CTM, FRSGC/UCI, LMDz-INCA, STOCHEM-HadAM3 and TM4) provided methane destruction fluxes for each scenario, and these were used to calculate whole atmosphere CH₄ lifetimes, as described in Stevenson et al. (2006). For four of the five models, changes in lifetime between scenarios were very consistent (within 4% of each other), but STOCHEM-HadAM3 was nearly twice as sensitive, and is thus considered an outlier. Here we use ensemble mean results from the four models to assess ship impacts on CH₄ lifetime. For present-day, ship \( \text{NO}_x \) shortens the CH₄ lifetime by 0.13 yr (1.56±0.05%) (S1-S1w); in 2030 the same ship \( \text{NO}_x \) perturbation reduces the lifetime by 0.10 yr (1.14±0.02%) (S4-S4w), again illustrating the slightly lower sensitivity when background levels are higher. Following the A2/CGS the methane lifetime is reduced by an additional 0.07 yr (0.77±0.02%) in 2030 (S4s-S4). Ship \( \text{NO}_x \) therefore introduces a negative radiative forcing by reducing the build-up of methane. We make a first-order estimate of the methane radiative forcing by linearly scaling the methane lifetime changes, assuming a feedback factor of 1.4 (IPCC, 2001) to infer a change in methane mixing ratio, and then calculate global mean forcing using the value of 0.37 W/m² ppb⁻¹ for a globally uniform methane change (Schimel et al., 1996). Estimated methane forcings are given in Table 3.

The various contributions to the radiative forcing from shipping also include radiative forcing due to \( \text{CO}_2 \) and sulphate changes. The corresponding radiative forcing of \( \text{CO}_2 \) is estimated from the fraction of the ship emission totals in the year 2000 (136.7 Tg(C)/yr, Endresen et al. (2003)) to the total annual \( \text{CO}_2 \) emissions in 2000 (7970 Tg(C)/yr, IPCC (2001), scenario A2). This fraction (1.7%) is used to scale the RF resulting from all \( \text{CO}_2 \) sources (1.51 W/m², IPCC (2001), scenario A2, ISAM reference case) linearly, resulting in a RF of 26 mW/m² due to shipping. The same approach is used to estimate \( \text{CO}_2 \) RF for the year 2030, with annual emissions of 14,720 Tg(C)/yr for all sources (IPCC (2001), scenario A2) and a total \( \text{CO}_2 \) RF of 2.59 W/m² (IPCC (2001), scenario A2, ISAM reference case). The emissions from shipping are assumed to increase at an annual rate of 2.2% since 2000, resulting in 262.6 Tg(C)/yr in 2030. For the simulation S4, shipping contributes with 0.9% to the total annual \( \text{CO}_2 \) emissions, for S4s with 1.8%. This results in \( \text{CO}_2 \) RF of about 24 mW/m² (S4-S4w) and 46 mW/m² (S4s-S4w) due to shipping in 2030. However, since \( \text{CO}_2 \) has a long average lifetime, the time integral of ship emissions would be needed to estimate a more accurate number for RF.

Direct sulphate forcings are calculated from the change in total SO₄ burdens due to shipping. The relative change of the SO₄ burdens is used to scale the total direct radiative forcing of sulphate particles given by IPCC (2001) (scenario A2) of −0.4 W/m² for the year 2000 and −0.65 W/m² for
the year 2030. The relative contribution of shipping to the total sulphate burdens is about 3.6% (S1-S1w), 2.0% (S4-S4w), and 4.0% (S4s-S4w). This results in a RF of –14 mW/m² (2000, S1-S1w), -13 mW/m² (2030, S4-S4w), and –26 mW/m² (2030, S4S-S4w). It should be noted that the sulphate RFs are only rough estimates, as it is expected that SO₄ forcings show a significant variability with emission site and cloud cover, which is not accounted for in this estimate.

RFs from shipping O₃, CH₄ and CO₂ as well as the direct sulphate forcings for the different scenarios are summarized in Table 3. Contributions to the RF resulting from the indirect aerosol effect (Capaldo et al., 1999) are not included, but are expected to be negative and larger in magnitude than the direct sulphate forcings estimated here.

4 Discussion

In the previous section, impacts of NOₓ and SO₂ ship emissions on O₃ and sulphate distributions have been presented for present-day and 2030 conditions under the same meteorological conditions. In general, the results derived from the ensemble mean comprising ten atmospheric chemistry models agree with previous studies based on single models (Lawrence and Crutzen, 1999; Kasibhatla et al., 2000; Davis et al., 2001; Endresen et al. 2003; Derwent et al., 2005). In this section we discuss the main uncertainties that could impact on results presented in Sect. 3 (Sect. 4.1) and how the results might change in the context of other emission scenarios (Sect. 4.2).

4.1 Uncertainties

4.1.1 Uncertainties in ship emission inventories

Uncertainties in the emission inventory include uncertainties in the vessel traffic densities and the emission totals. Several emission inventories for shipping based on energy statistics have been published in recent years resulting in a total fuel consumption below or around 160 Mt per year (e.g. Corbett and Fischbeck, 1997; Corbett et al., 1999; Olivier et al., 2001; Endresen et al., 2003). However, recent studies which used an activity-based approach and statistical information of the total fleet greater than 100 GT including the larger military vessels and auxiliary engines (Lloyd’s, 2002) suggest a fuel consumption of 289 Mt/yr (Corbett and Köhler, 2003) or 280 Mt/yr (Eyring et al. 2005a) for the year 2001. Ideally, the fuel consumption calculated from energy statistics and with an activity-based approach would be the same, but there is an ongoing discussion on the baseline value for the 2000 fuel consumption (Endresen et al., 2004; Corbett and Köhler, 2004; Eyring et al., 2005a). The emission totals for NOₓ used in this study are around a factor of two lower than those calculated from activity-based approaches.

Vessel traffic densities in the inventory used here are based on EDGAR 3.2 (Olivier et al., 2001). Compared to other data sets such as the Comprehensive Ocean-Atmosphere Data Set (COADS, see http://www.wmo.ch/web/www/ois/ois-home.htm) or the Automated Mutual-assistance Vessel Rescue system (AMVER; Endresen et al., 2003), shipping routes are probably too narrow in this inventory. As pointed out by Lawrence and Crutzen (1999) this could lead to an underestimation of O₃ production, as photochemical production is more efficient at lower NOₓ concentrations (Li et al., 1987). However, the emissions are instantaneously diluted into the large model grid boxes (see Table 1) which partly compensates this effect. Over the Baltic emissions in the EDGAR3.2 inventory are higher than in the COADS and AMVER data sets, and over the Mediterranean they are smaller. Note that significant differences between the COADS and AMVER data sets have also been reported (Endresen et al., 2003), as only small subsets of the world-merchant fleet find input into these data bases.

Assuming that changes in O₃ burden scale linearly with increasing NOₓ emissions (see Sect. 3.3.2), we conclude that the uncertainty arising from NOₓ emission totals in the inventory itself could lead to an underestimation of up to 100% in the simulated O₃ response. We also conclude that due to the inventory used, the process of titration might be overestimated over the Baltic Sea and the impact of ship emissions over the Mediterranean might be underestimated. In addition, the use of a yearly average ship emission inventory is a simplification and e.g. overestimates the impact of ship emissions over the partly frozen Baltic Sea in winter time.

The uncertainties in emission inventories discussed above apply also for the future scenarios. In the future, the global distribution of vessel traffic might change. For example, while international trade in north-south direction and within the Southern Hemisphere is likely increasing, the east-west trade on the long-term might reach a level of saturation and hence reduced growth (Eyring et al., 2005b). However, changes in vessel traffic densities are likely to have only small effects on the global scale, though they could regionally be important. For example, with sea ice expected to recede in the Arctic during the 21st century as a result of projected climate warming, Granier et al. (2006) showed that the opening of new shipping routes in the Arctic could lead to an important increase in O₃ levels in this remote region.

4.1.2 The impact of VOC and CO emissions from ships

We applied one model (MATCH-MPIC) to assess the impact of VOC and CO emissions from ships on tropospheric O₃ and NOₓ burdens. These sensitivity simulations with VOC and CO emissions from ships included (1.2 Tg(CO)/yr and 2.03 Tg(VOCs)/yr in 2000 from Endresen et al. (2003); 2.15 Tg(CO)/yr and 3.89 Tg(VOCs)/yr in 2030 under CGS) are displayed in Figs. 9 and 10 (MATCH-MPIC/VOC) and show only small differences compared to the MATCH-MPIC reference simulations. The differences are most apparent over the Baltic where NOₓ are dominated by high emissions.
from land in the $5^\circ \times 5^\circ$ model grid boxes. On the global scale as well as in the Pacific and Indian Ocean the differences are small, and even in the Baltic the difference between the sensitivity and reference simulations in the MATCH-MPIC model are much smaller than the inter-model differences. We therefore conclude that the neglect of ship CO and VOC emissions in the reference simulations does not change the main conclusions of this paper. This can be understood, because total CO emissions from ships are small and VOC emissions are not expected to play a significant role over the remote ocean. (1) In general, CO emission is the result of incomplete combustion. Since large-bore diesel engines operate at high air excess ratios and high combustion temperatures, total CO emissions from the fleet above 100 GT are small and much lower than from other internal combustion engines (see Eyring et al., 2005a). (2) Past global modelling studies (e.g. Stevenson et al., 1998; Stevenson et al., 2006) have found that the troposphere is mainly NO$_x$-limited with respect to O$_3$ production, as is to be expected since the vast majority of it is quite “remote” from NO$_x$ emissions. This is particularly true for oceanic regions. VOC-limited O$_3$ production regions are typically limited to highly polluted, urban areas. With the exception of the Baltic Sea these rarely occur, even in the most congested shipping lanes, at least when they are represented in the global model grid-boxes of $5^\circ$ longitude $\times 5^\circ$ latitude. Therefore, even though the models describe atmospheric photochemistry in considerable detail, and include multiple non-linear interactions, this doesn’t mean one should necessarily expect non-linear relationships in the shipping response. Indeed, the response of the modelled NO$_x$ and O$_3$ burdens to varying the magnitude of ship NO$_x$ emissions is basically linear whether or not VOC emissions from ships are included (Fig. 9), but as discussed in Sect. 3.3.2 small saturation effects are simulated. These findings are in agreement with results from Endresen et al. (2003) who also showed that the effects of VOC emissions from ships on present-day O$_3$ are very small.

4.1.3 The role of plume dispersion

Many physical and chemical processes in the atmosphere occur on spatial and temporal scales that cannot be resolved in current global numerical models, which have typical resolutions of a few hundred kilometres. One major open issue is the evolution and chemical transformation of trace gases and particulate matter emitted from individual point sources such as e.g. a single ship during the dispersion into the large scales of a grid cell of a global model. Emission totals are based on specific fuel-oil consumption rates and emission indices measured at the manufacturers’ engine test beds (e.g. Eyring et al., 2005a). The emission totals are distributed over the globe with the help of vessel traffic densities derived in various ways and are instantaneously spread onto large inventory grid boxes, usually $1^\circ$ longitude $\times 1^\circ$ degree latitude, without accounting for chemical dispersion on the sub-grid scale. Those inventories are used as inputs in emission databases such as EDGAR (Olivier et al., 2001) and global modelling studies. The horizontal resolution in the global models used in this study all have even larger grid sizes of a few hundred kilometres (see Table 1). Therefore, emissions from ships are further instantaneously distributed within the large grid boxes. However, several studies have pointed to the importance of chemical conversion in the near field of ships (Kasibhatla et al., 2000; Davis et al., 2001; Song et al., 2003; von Glasow et al., 2003; Chen et al., 2005). Uncertainties in the modelled O$_3$ changes arising from the neglect of plume chemistry are hard to quantify, in particular because the studies cited above are all based on certain meteorological situations, amount of emissions released by the ship, and daytime. However, the studies all agree in this respect that the lifetime of NO$_x$ is significantly reduced in the plumes, corresponding to high in-plume NO$_x$ destruction. Chen et al. (2005) used a box model and found that more than 80% of the NO$_x$ loss is due to the reaction of NO$_2$ with OH and the rest due to PAN formation. As parameterisations that account for these sub-grid scale processes are not yet available, the model simulations have been run without accounting for sub-grid plume chemistry, but with a relatively low total emission estimate (Sect. 4.1.1).

4.1.4 Uncertainties due to different model approaches

Uncertainties in different model approaches presented here and discussed in Sect. 3.2 arise from differences in the main characteristics in the models (see Sect. 2.1). The advantage of this study compared to all previous assessments on ship emissions is that an ensemble of ten models has been used, which makes the results more robust, because the models have performed the same experiments with different treatments of chemical and dynamical processes. In summary, the differences in the simulated contributions from shipping to O$_3$, NO$_2$, sulphate and RF are smaller than 20%, as revealed by the intermodel standard deviations.

4.2 Results in the context of other emission scenarios

Up to now we have only quantified the impact of ship emissions in 2030 under the assumption that all other emissions vary as projected in the A2 scenario. The A2 scenario is a rather pessimistic scenario, which describes a very heterogeneous world with high population growth. Economic development is primarily regionally oriented and per capita economic growth and technological changes are more fragmented and slower than in other IPCC SRES storylines. NO$_x$ emissions in A2 increase to 179 Tg(NO$_x$)yr$^{-1}$ in 2030. However, if ground based emissions grow less rapidly than under A2, the relative contribution of ship emissions will become more important. The ‘Current Legislation Scenario (CLE)’ scenario takes into account the current perspectives of individual countries on future economic development and
anticipated effects of presently decided emission control legislation in the individual countries and the “Maximum technically Feasible Reduction” (MFR) scenario considers the scope for emission reductions offered by full implementation of the presently available emission control technologies, while maintaining the projected levels of anthropogenic activities. A detailed description of the two scenarios can be found in Dentener et al. (2005). Compared to today’s anthropogenic NO\textsubscript{x} emissions (91.3 Tg(NO\textsubscript{2}) yr\textsuperscript{-1} in S1; not including biomass burning), NO\textsubscript{x} emissions in the CLE scenario increase to 108 Tg(NO\textsubscript{2}) yr\textsuperscript{-1} and decrease to 43.0 Tg(NO\textsubscript{2}) yr\textsuperscript{-1} in the MFR scenario in 2030 (Stevenson et al., 2006, Table 3). Differences in tropospheric O\textsubscript{3} burden between 2030 and 2000 for the three scenarios A2, CLE, and MFR have been calculated from an ensemble of 26 models in Stevenson et al. (2006), their Table 6. The simulated inter-scenario ensemble mean changes in O\textsubscript{3} burdens between 2030 and 2000 resulted in 53±10 Tg(O\textsubscript{3}) in the A2 scenario with ship emissions remaining at 2000 levels (S4-S1), 20±4 Tg(O\textsubscript{3}) in the CLE scenario, and −16±4 Tg(O\textsubscript{3}) in the MFR scenario. If we add on the 1.75 Tg(O\textsubscript{3}) increase in O\textsubscript{3} burdens due to ship emission increase under the A2/CGS calculated in this study (see Fig. 9) to the difference in O\textsubscript{3} burden between the S4 and the S1 simulation from Stevenson et al. (2006), the global mean relative contribution from shipping is around 3% under the A2 scenario.
However, under the CLE scenario the relative contribution of ships to O₃ burden trends until 2030 increases to around 9%. If land-based emissions fall, as under the MFR scenario, but ship emissions continue to grow, they will significantly counteract the benefits derived from the land-based emissions reductions.

As an example how increases in ship emissions might impact on future trends, Fig. 13 shows differences in near-surface NO₂, O₃ and sulphate between the 2030 A2 scenario and 2000 over Europe if ship emissions are zero in 2030 (S₄w-S₁; left column), remain constant at 2000 levels (S₄-S₁; middle column), or increase with a constant growth rate of 2.2%/yr (S₄s-S₁; right column).

If NOₓ emissions from shipping fall to zero in 2030, a decrease in near-surface NO₂ concentrations between 2030 and 2000 of up to 2 ppbv over Scandinavia and 0.4 ppbv over the Atlantic can be reached (a reduction of over 50%), whereas central Europe is dominated by increases due to the rise in land NOₓ emissions (Fig. 13a). On the other hand, with constant or increasing emissions from shipping (Fig. 13b,c) near-surface NO₂ levels are enhanced over the continent by up to 2 ppbv (>50%), in particular over the coastal regions of the North Sea and the Baltic Sea as well as over the southern part of Great Britain. Only small differences in O₃ are simulated under the three different ship emission scenarios (Figs. 13d-f). In a world without ship NOₓ emissions in 2030, the positive trend in near-surface O₃ will be reduced over most of Europe (Fig. 13d), whereas it will be enhanced in the A2/CGS (Fig. 13f).

The strongest impact of ship emissions to the trends up to 2030 is simulated in the sulphate distributions. A significant decrease in sulphate is simulated in a world without ship emissions in 2030 (Fig. 13g). If ship emissions however remain at today’s levels, this negative trend over Central Europe and the United Kingdom is reduced and even changes sign over large areas of the Atlantic and over Scandinavia under the A2/CGS (Fig. 13i). Thus, increasing emissions from shipping would significantly counteract the benefits derived from reducing SO₂ emissions from all other anthropogenic sources under the A2 scenario over the continents in Europe.

5 Summary and Conclusions

In this study we have used an ensemble of ten state-of-the-art global atmospheric chemistry models to assess the impact of NOₓ atmospheric chemistry models to assess the impact of NOₓ emissions from international shipping on O₃ for present-day conditions (year 2000). This multi-model approach accounts for intermodel differences and therefore makes the results more robust compared to previous studies. In addition this study for the first time quantifies the potential impact of ship emissions in the future (year 2030). A subset of four models included the tropospheric sulphur cycle. The ensemble mean of these four models has been applied to investigate the changes in sulphate distributions due to SO₂ emissions from international shipping for present-day conditions and in 2030.

For present-day conditions we find the most pronounced changes in annual mean tropospheric NO₂ and SO₂ columns over the Baltic and the North Sea, and also though smaller over the Atlantic, Gulf of Mexico, and along the main shipping lane from Europe to Asia. Maximum near-surface O₃ changes due to NOₓ ship emissions are simulated over the North Atlantic in July (∼12 ppbv) in agreement with previously reported results (Lawrence and Crutzen, 1999; Endresen et al., 2003). However, in contrast to Endresen et al. (2003), a decrease in O₃ in winter is found over large areas in Europe (∼3 ppbv) due to titration, which could be related to differences in the emission inventories. Overall NOₓ emissions most effectively produce O₃ over the remote ocean, where background NOₓ levels are small.

The two 2030 scenarios both specify emissions following the IPCC SRES A2 scenario (IPCC, 2000). The first future scenario assumes that ship emissions remain constant at 2000 levels and under this scenario a slightly smaller response in O₃ and sulphate changes due to shipping is found compared to the present-day contribution from shipping. This indicates that higher background levels tend to slightly reduce the perturbation from ships. The second emission scenario addresses the question of how NOₓ and SO₂ emissions from international shipping might influence atmospheric chemistry in the next three decades if these emissions grow unabated and one assumes a constant annual growth rate of 2.2% between 2000 and 2030 (“Constant Growth Scenario” (CGS)). The models show future increases in NOₓ and O₃ burden which scale almost linearly with increases in NOₓ emission totals under the same background conditions. Therefore, there is evidence that the ship NOₓ effect is only weakly subject to saturation in its current magnitude range, and that saturation cannot be expected to help mitigate the effects of near-future increases. In other words a doubling of NOₓ emissions from ships in the future might lead to a doubling in atmospheric O₃ burdens due to ship emissions. In addition, increasing emissions from shipping would significantly counteract the benefits derived from reducing SO₂ emissions from all other anthropogenic sources under the A2 scenario over the continents, for example in Europe. Under A2/CGS shipping globally contributes with 3% to increases in O₃ burden until 2030 and with 4.5% to increases in sulphate. The results discussed above are calculated under the assumption that all other emissions follow the A2 scenario broadly representing a pessimistic future situation. However, if future ground based emissions follow a more stringent scenario, the relative importance of ship emissions becomes larger.

Tropospheric O₃ forcings due to ships of 9.8 mW/m² in 2000 and 13.6 mW/m² in 2030 are simulated by the ensemble mean, with standard deviations of 10–15%. Compared to aviation (∼20 mW/m²; Sausen et al., 2005) tropospheric O₃ forcings from shipping are of the same order in 2000, despite the much higher NOₓ emissions from ships (Eyring et al., 2003), a decrease in O₃ in winter is found over large areas in Europe (∼3 ppbv) due to titration, which could be related to differences in the emission inventories. Overall NOₓ emissions most effectively produce O₃ over the remote ocean, where background NOₓ levels are small.

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This can be understood because peak changes in O$_3$ due to shipping occur close to the surface, whereas changes in O$_3$ due to aviation peak in the upper troposphere (Grewe et al., 2002). The net radiative forcing is most sensitive to NO$_x$ emissions at altitudes of about 8–12 km because of longer NO$_x$ and O$_3$ lifetimes, and colder temperatures compared to the surface (Lacis et al., 1990; Brasseur et al., 1998). Ship NO$_x$ reduces the CH$_4$ lifetime by 0.13 yr in 2000 and by up to 0.17 yr in 2030, introducing a negative radiative forcing of about $-14 \text{ mW/m}^2$ in 2000 and $-21 \text{ mW/m}^2$ in 2030. A rough estimate of RF from shipping CO$_2$ suggests 26 mW/m$^2$ in 2000 compared to 23 mW/m$^2$ from aviation CO$_2$. The direct effect from SO$_2$ ship emissions is approximately $-14 \text{ mW/m}^2$ in 2000 and decreases to a more negative value of $-26 \text{ mW/m}^2$ in 2030 under A2/CGS.

The recent rapid rise in ship emissions may have generated O$_3$ trends that have hitherto been attributed to increases in hemispheric background, and related to North American or Asian anthropogenic emissions, or to changes in forest fire activities (e.g., observations at Mace Head, Simmonds et al., 2005). For example, Lelieveld et al. (2004) report significant surface O$_3$ trends over the Atlantic Ocean, although not particularly over the North Atlantic, where ships appear to have their largest impact in our study. The large rise in ship emissions and the associated increase in O$_3$ may be compromising measures to reduce O$_3$ in some regions, e.g., Europe and potentially elsewhere (India, western North America). Of the estimated 0.35 W/m$^2$ radiative forcing since 1750 due to increases in tropospheric O$_3$, our results indicate that about 4% of this is due to ship NO$_x$ emissions.

This study has also investigated the range of results given by the individual models compared to other uncertainties. Uncertainties in the simulated O$_3$ contributions from ships for the different model approaches revealed by the inter-model standard deviations are found to be significantly smaller than estimated uncertainties stemming from the ship emission inventory, mainly the ship emission totals, the distribution of the emissions over the globe, and the neglect of ship plume dispersion. The relatively small intermodel variance lends to confidence to our use of current models to study the effects of shipping, while highlighting the urgent need to quantify the uncertainties that may be present or to the plume nature of shipping emissions.

The results of the ship induced O$_3$, NO$_x$, and SO$_2$ changes discussed above are derived from one particular ship emission inventory. The main goal for future work is to reduce uncertainties in the emissions totals and to further improve vessel traffic densities as well as to develop sophisticated parameterisations of unresolved plume processes for the use in global models.

An intercomparison of the model results with observations over the Northern Hemisphere (25°–60° N) oceanic regions in the lower troposphere showed that the models are capable to reproduce O$_3$ and NO$_x$ reasonably well, whereas SO$_2$ in the marine boundary layer is significantly underestimated. However, evaluation of the models’ response to ship emissions is still at a preliminary stage and is currently limited by the coarse spatial resolution of the models, the uncertainty in the measurements, the lack of sufficient in situ measurements over the ocean, and the difficulty to separate ship emissions from other even stronger emission sources close to land. Additional in situ measurements inside single ship plumes, but also in the corridor of the shipping lanes are needed and the set up of a measurement network onboard ships similar to MOZAIC (Measurements of Ozone and water vapour by in-service Alrbus airCraft; Marenco et al., 1998) or CARIBIC (Civil Aircraft for Global Measurement of Trace Gases and Aerosols in the Tropopause Region; Brenninkmeijer et al., 1999) onboard civil aircrafts would be desirable. Unambiguous detection of ship emissions in satellite data is currently only available for the region of the Red Sea and the Indian Ocean (Beirle et al., 2004; Richter et al., 2004), where shipping routes are close to the coastal area. Reduction in measurement uncertainties through use of long-term averages and data from more instruments (e.g. OMI and GOME-2) combined with better constraints on land-based sources and higher spatial resolution in the models should facilitate such an intercomparison in the future.

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