Air quality and radiative impacts of Arctic shipping emissions in the summertime in northern Norway: from the local to the regional scale

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Abstract. In this study, we quantify the impacts of shipping pollution on air quality and shortwave radiative effect in northern Norway, using WRF-Chem (Weather Research and Forecasting with chemistry) simulations combined with high-resolution, real-time STEAM2 (Ship Traffic Emissions Assessment Model version 2) shipping emissions. STEAM2 emissions are evaluated using airborne measurements from the ACCESS (Arctic Climate Change, Economy and Society) aircraft campaign, which was conducted in the summer 2012, in two ways. First, emissions of nitrogen oxides (NOx) and sulfur dioxide (SO2) are derived for specific ships by combining in situ measurements in ship plumes and FLEXPART-WRF plume dispersion modeling, and these values are compared to STEAM2 emissions for the same ships. Second, regional WRF-Chem runs with and without STEAM2 ship emissions are performed at two different resolutions, 3 km × 3 km and 15 km × 15 km, and evaluated against measurements along flight tracks and average campaign profiles in the marine boundary layer and lower troposphere. These comparisons show that differences between STEAM2 emissions and calculated emissions can be quite large (−57 to +148 %) for individual ships, but that WRF-Chem simulations using STEAM2 emissions reproduce well the average NOx, SO2 and O3 measured during ACCESS flights. The same WRF-Chem simulations show that the magnitude of NOx and ozone (O3) production from ship emissions at the surface is not very sensitive (<5 %) to the horizontal grid resolution (15 or 3 km), while surface PM10 particulate matter enhancements due to ships are moderately sensitive (15 %) to resolution. The 15 km resolution WRF-Chem simulations are used to estimate the regional impacts of shipping pollution in northern Norway. Our results indicate that ship emissions are an important source of pollution along the Norwegian coast, enhancing 15-day-averaged surface concentrations of NOx (∼+80 %), SO2 (∼+80 %), O3 (∼+5 %), black carbon (∼+40 %), and PM2.5 (∼+10 %). The residence time of black carbon originating from shipping emissions is 1.4 days. Over the same 15-day period, ship emissions in northern Norway have a global shortwave (direct + semi-direct + indirect) radiative effect of −9.3 m W m−2.

1 Introduction

Shipping is an important source of air pollutants and their precursors, including carbon monoxide (CO), nitrogen oxides (NOx), sulfur dioxide (SO2), volatile organic compounds (VOCs) as well as organic carbon (OC) and black carbon (BC) aerosols (Corbett and Fischbeck, 1997; Corbett and Köhler, 2003). It is well known that shipping emissions have an important influence on air quality in coastal regions,
often enhancing ozone (O\textsubscript{3}) and increasing aerosol concentrations (e.g., Endresen et al., 2003). Corbett et al. (2007) and Winebrake et al. (2009) showed that aerosol pollution from ships might be linked to cardiopulmonary and lung diseases globally. Because of their negative impacts, shipping emissions are increasingly subjected to environmental regulations. The International Maritime Organization (IMO) has designated several regions as Sulfur Emission Control Areas (SECA; including the North Sea and Baltic Sea in Europe), where low sulfur fuels must be utilized to minimize the air quality impacts of shipping on particulate matter (PM) levels. The sulfur content in ship fuels in SECA was limited to 1% by mass in 2010, decreasing to 0.1% in 2015, while the global average is 2.4% (IMO, 2010). Less strict sulfur emission controls (0.5%) will also be implemented worldwide, at the latest in 2025, depending on current negotiations. Ships produced or heavily modified recently must also comply to lower NO\textsubscript{x} emissions factors limits, reducing emission factors (in g kWh\textsuperscript{-1}) by approximately −10% (after 2000) and another −15% (after 2011) compared to ships built before year 2000 (IMO, 2010). Jonson et al. (2015) showed that the creation of the North Sea and Baltic Sea SECA was effective in reducing current pollution levels in Europe, and that further NO\textsubscript{x} and sulfur emission controls in these regions could help to achieve strong health benefits by 2030 by reducing PM levels. In addition to its impacts on air quality, maritime traffic already contributes to climate change, by increasing the concentrations of greenhouse gases (CO\textsubscript{2}, O\textsubscript{3}) and aerosols (SO\textsubscript{4}, OC, BC) (Capaldo et al., 1999; Endresen et al., 2003). The current radiative forcing of shipping emissions is negative and is dominated by the cooling influence of sulfate aerosols formed from SO\textsubscript{2} emissions (Eyring et al., 2010). However, due to the long lifetime of CO\textsubscript{2} compared to sulfate, shipping emissions warm the climate in the long term (after 350 years; Fuglestvedt et al., 2009). In the future, global shipping emissions of SO\textsubscript{2} are expected to decrease due to IMO regulations, while global CO\textsubscript{2} emissions from shipping will continue to grow due to increased traffic. This combination is expected to cause warming relative to the present day (Fuglestvedt et al., 2009; Dalsøren et al., 2013). In addition to their global impacts, shipping emissions are of particular concern in the Arctic, where they are projected to increase in the future as sea ice declines (for details of future sea ice, e.g., Stroeve et al., 2011). Decreased summer sea ice, associated with warmer temperatures, is progressively opening the Arctic region to transit shipping, and projections indicate that new trans-Arctic shipping routes should be available by mid-century (Smith and Stephenson, 2013). Other shipping activities are also predicted to increase, including shipping associated with oil and gas extraction (Peters et al., 2011). Sightseeing cruises have increased significantly during the last decades (Eckhardt et al., 2013), although it is uncertain whether or not this trend will continue. Future Arctic shipping is expected to have important impacts on air quality in a now relatively pristine region (e.g., Granier et al., 2006), and will influence both Arctic and global climate (Dalsøren et al., 2013; Lund et al., 2012). In addition, it has recently been shown that routing international maritime traffic through the Arctic, as opposed to traditional routes through the Suez and Panama canals, will result in warming in the coming century and cooling on the long term (150 years). This is due to the opposite sign of impacts due to reduced SO\textsubscript{2} linked to IMO regulations and reduced CO\textsubscript{2} and O\textsubscript{3} associated with fuel savings from using these shorter Arctic routes (Fuglestvedt et al., 2014). In addition, sulfate is predicted to cause a weaker cooling effect for the northern routes (Fuglestvedt et al., 2014). Although maritime traffic is relatively minor at present in the Arctic compared to global shipping, even a small number of ships can significantly degrade air quality in regions where other anthropogenic emissions are low (Aliabadi et al., 2015; Eckhardt et al., 2013). Dalsøren et al. (2007) and Ødemark et al. (2012) have shown that shipping emissions also influence air quality and climate along the Norwegian and Russian coasts, where current Arctic ship traffic is the largest. Both studies (for years 2000 and 2004) were based on emission data sets constructed using ship activity data from the AMVER (Automated Mutual-Assistance VEssel Rescue system) and COADS (Comprehensive Ocean–Atmosphere Data Set) data sets. However, the AMVER data set is biased towards larger vessels (> 20 000 t) and cargo ships (Endresen et al., 2003), and both data sets have limited coverage in Europe (Miola and Ciufla, 2011). More recently, ship emissions using new approaches have been developed that use ship activity data more representative of European maritime traffic, based on the AIS (Automatic Identification System) ship positioning system. These include the STEAM2 (Ship Traffic Emissions Assessment Model version 2) shipping emissions, described in Jalkanen et al. (2012) and an Arctic-wide emission inventory described in Winther et al. (2014). To date, quantifying the impacts of Arctic shipping on air quality and climate has also been largely based on global model studies, which are limited in horizontal resolution. In addition, there have not been specific field measurements focused on Arctic shipping that could be used to study the local influence of shipping emissions in the European Arctic and to validate model predicted air quality impacts. In this study, we aim to quantify the impacts of shipping along the Norwegian coast in July 2012, using airborne measurements from the ACCESS (Arctic Climate Change, Economy and Society) aircraft campaign (Roiger et al., 2015). This campaign (Sect. 2) took place in summer 2012 in northern Norway, and was primarily dedicated to the study of local pollution sources in the Arctic, including pollution originating from shipping. ACCESS measurements are combined with two modeling approaches, described in Sect. 3. First, we use the Weather Research and Forecasting (WRF) model to drive the Lagrangian particle dispersion model FLEXPART-WRF run in forward mode to predict the dispersion of ship...
emissions. FLEXPART-WRF results are used in combination with ACCESS aircraft measurements in Sect. 4 to derive emissions of NO\textsubscript{x} and SO\textsubscript{2} for specific ships sampled during ACCESS. The derived emissions are compared to emissions from the STEAM2 model for the same ships. Then, we perform simulations with the WRF-Chem model, including STEAM2 ship emissions, in order to examine in Sect. 5 the local (i.e., at the plume scale) and regional impacts of shipping pollution on air quality and shortwave radiative effects along the coast of northern Norway.

2 The ACCESS aircraft campaign

The ACCESS aircraft campaign took place in July 2012 from Andenes, Norway (69.3° N, 16.1° W); it included characterization of pollution originating from shipping (four flights) as well as other local Arctic pollution sources (details are available in the ACCESS campaign overview paper; Roiger et al., 2015). The aircraft (DLR Falcon 20) payload included a wide range of instruments measuring meteorological variables and trace gases, described in detail by Roiger et al. (2015). Briefly, O\textsubscript{3} was measured by UV (ultraviolet) absorption (5% precision, 0.2 Hz), nitrogen oxide (NO), and nitrogen dioxide (NO\textsubscript{2}) by chemiluminescence and photolytic conversion (10% precision for NO, 15% for NO\textsubscript{2}; 1 Hz), and SO\textsubscript{2} by chemical ionization ion trap mass spectrometry (20% precision; 0.3 to 0.5 Hz). Aerosol size distributions between 60 nm and 1 μm were measured using a Ultra-High Sensitivity Aerosol Spectrometer Airborne.

The four flights focused on shipping pollution took place on 11, 12, 19, and 25 July 2012 and are shown in Fig. 1a (details on the 11 and 12 July 2012 flights shown in Fig. 1b). The three flights on 11, 12, and 25 July 2012 sampled pollution from specific ships (referred to as single-plume flights). During these flights, the research aircraft repeatedly sampled relatively fresh emissions from one or more ships during flight legs at constant altitudes, at several distances from the emission source, and in some cases at different altitudes. In this study, measurements from these single-plume flights are used in combination with ship plume dispersion simulations (described in Sects. 3.1 and 4.1) to estimate emissions from individual ships. This method relies on knowing the precise locations of the ships during sampling. Because those locations are not known for the ship emissions sampled on 25 July 2012 flight, emissions are only calculated for the three ships targeted during the 11 and 12 July 2012 flights (the Costa Deliziosa, Wilson Leer, and Wilson Nanjing), and for an additional ship (the Alaed) sampled during the 12 July flight, whose location could be retrieved from the STEAM2 shipping emission inventory (presented in Sect. 3.3). Table 1 gives more information about these four ships, one large cruise ship and three cargo ships. On 11 and 12 July 2012, the research aircraft sampled fresh ship emissions within the boundary layer, during flight legs at low altitudes (< 200 m).

Table 1. Description of the ships sampled during the ACCESS flights on 11 and 12 July 2012.

<table>
<thead>
<tr>
<th>Ship name</th>
<th>Vessel type</th>
<th>Gross tonnage (tons)</th>
<th>Fuel type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wilson Leer</td>
<td>Cargo ship</td>
<td>2446</td>
<td>Marine gas oil</td>
</tr>
<tr>
<td>Costa Deliziosa</td>
<td>Passenger ship</td>
<td>92 720</td>
<td>Heavy fuel oil</td>
</tr>
<tr>
<td>Wilson Nanjing</td>
<td>Cargo ship</td>
<td>6118</td>
<td>Heavy fuel oil</td>
</tr>
<tr>
<td>Alaed*</td>
<td>Cargo ship</td>
<td>7579</td>
<td>Heavy fuel oil</td>
</tr>
</tbody>
</table>

* Ship present in STEAM2, not targeted during the campaign.

Fresh ship emissions were sampled less than 4 h after emission. In addition to the single-plume flights, the 19 July 2012 ACCESS flight targeted aged ship emissions in the marine boundary layer near Trondheim. Data collected during the 11 and 12 July 2012 flights are used to derive emissions from operating ships (Sect. 4), and data from the four flights (11, 12, 19, and 25 July 2012) are used to evaluate regional chemical transport simulations investigating the impacts of shipping in northern Norway (Sect. 5). Other flights from the ACCESS campaign were not used in this study because their flight objectives biased the measurements towards other emissions sources (e.g., oil platforms in the Norwegian Sea) or because they included limited sampling in the boundary layer (flights north to Svalbard and into the Arctic free troposphere; Roiger et al., 2015).

3 Modeling tools

3.1 FLEXPART-WRF and WRF

Plume dispersion simulations are performed with FLEXPART-WRF for the four ships presented in Table 1, in order to estimate their emissions of NO\textsubscript{x} and SO\textsubscript{2}. FLEXPART-WRF (Brioude et al., 2013) is a version of the Lagrangian particle dispersion model FLEXPART (Stohl et al., 2005), driven by meteorological fields from the mesoscale weather forecasting model WRF (Skamarock et al., 2008). In order to drive FLEXPART-WRF, a meteorological simulation was performed with WRF version 3.5.1, from 4 to 25 July 2012, over the domain presented in Fig. 1a. The domain (15 km × 15 km horizontal resolution with 65 vertical eta levels between the surface and 50 hPa) covers most of northern Norway (~62 to 75° N) and includes the region of all ACCESS flights focused on ship emissions. The first week of the simulation (4 to 10 July included) is used for model spin-up. WRF options and parameterizations used in these simulations are shown in Table 2. Meteorological initial and boundary conditions are obtained from the FNL (abbreviation for “final”) analysis from NCEP (National Centers for Environmental Prediction). The simulation is also nudged to FNL winds, temperature, and humidity every

Figure 1. WRF and WRF-Chem domain (a) outer domains used for the MET, CTRL, and NOSHIP runs. ACCESS flight tracks during 11, 12, 19a (a – denotes that this was the first flight that occurred on this day, flight 19b – the second flight was dedicated to hydrocarbon extraction facilities) and 25 July 2012 flights are shown in color. (b) Inner domain used for the CTRL3 and NOSHIPS3 simulations, with the tracks of the four ships sampled during the 11 and 12 July 2012 flights (routes extracted from the STEAM2 inventory).

Table 2. Parameterizations and options used for the WRF and WRF-Chem simulations.

<table>
<thead>
<tr>
<th>Atmospheric process</th>
<th>WRF-Chem option</th>
</tr>
</thead>
<tbody>
<tr>
<td>Planetary boundary layer</td>
<td>MYNN (Nakanishi and Niino, 2006)</td>
</tr>
<tr>
<td>Surface layer</td>
<td>MM5 Similarity scheme, Carlson–Boland viscous sublayer (Zhang and Anthes, 1982; Carlson and Boland, 1978)</td>
</tr>
<tr>
<td>Land surface</td>
<td>Unified Noah land-surface model (Chen and Dudhia, 2001)</td>
</tr>
<tr>
<td>Microphysics</td>
<td>Morrison (Morrison et al., 2009)</td>
</tr>
<tr>
<td>Shortwave radiation</td>
<td>Goddard (Chou and Suarez, 1999)</td>
</tr>
<tr>
<td>Longwave radiation</td>
<td>RRTM (Mlawer et al., 1997)</td>
</tr>
<tr>
<td>Cumulus parameterization</td>
<td>Grell-3D (Grell and Devenyi, 2002)</td>
</tr>
<tr>
<td>Photolysis</td>
<td>Fast-J (Wild et al., 2000)</td>
</tr>
<tr>
<td>Gas phase chemistry</td>
<td>CBM-Z (Zaveri and Peters, 1999)</td>
</tr>
<tr>
<td>Aerosol model</td>
<td>MOSAIC 8 bins (Zaveri et al., 2008)</td>
</tr>
</tbody>
</table>

6 h. This WRF meteorological simulation is referred to as the MET simulation.

Ship emissions are represented in the FLEXPART-WRF plume dispersion simulations as moving 2 m × 2 m × 2 m box sources, whose locations are updated every 10 s along the ship trajectory (routes shown in Fig. 1b). In all, 1000 particles are released every 10 s into these volume sources, representing a constant emission flux with time of an inert tracer. During the ACCESS flights, targeted ships were moving at relatively constant speeds during the ~3 h of the flight, meaning that fuel consumption and emission fluxes are likely to be constant during the flights if environmental conditions (wind speed, waves, and currents) were not varying strongly. FLEXPART-WRF takes into account a simple exponential decay using a prescribed lifetime. In our case, the lifetime of NOx relative to reaction with OH was estimated using results from WRF-Chem simulations presented in Sect. 3.2. Specifically, we use OH concentrations, temperature, and air density from the CTRL3 simulation (Sects. 3.2 and 5.1). The NOx lifetime was estimated to be 12 h on 11 July and 5 h on 12 July. The SO2 lifetime was not taken into account, consistent with the findings of Lee et al. (2011), who reported a lifetime of ~20 h over the mid-Atlantic during summer, which is significantly longer than the ages of plumes measured during ACCESS. The FLEXPART-WRF output consists of particle positions, each associated with a pollutant mass; these particles are mapped onto a 3-D output grid (600 m × 600 m, with 18 vertical levels between 0 and 1500 m.a.s.l.) to derive fields of volume mixing ratios every minute. Since emissions are assumed to be constant with time, and since our simulations only take into account transport processes depending linearly on concentrations, the intensity of these mixing ratio fields also depend linearly on the emission strength chosen for the simulation. Therefore, the model results can be scaled a posteriori to represent any constant emission flux value.
Ship emissions can continue to rise after leaving the exhaust, due to their vertical momentum and buoyancy. This was taken into account in the FLEXPART-WRF simulations by calculating effective injection heights for each targeted ship, using a simple plume rise model (Briggs, 1965). This model takes into account ambient temperature and wind speed, as well as the volume flow rate and temperature at the ship exhaust, to calculate a plume injection height above the ship stack. Ambient temperature and wind speed values at each ship’s position are obtained from the WRF simulation. We use an average of measurements by Lyyranen et al. (1999) and Cooper (2001) for the exhaust temperature of the four targeted ships (350 °C). The volume flows at the exhaust are derived for each ship using CO₂ emissions from the STEAM2 ship emission model (STEAM2 emissions described in Sect. 3.3). Specifically, CO₂ emissions from STEAM2 for the four targeted ships are converted to an exhaust gas flow based on the average composition of ship exhaust gases measured by Cooper (2001) and Petzold et al. (2008). Average injection heights, including stack heights and plume rise, are found to be approximately 230 m for the Costa Deliziosa, 50 m for the Wilson Nanjing, 30 m for the Wilson Leer, and 65 m for the Alaed. In order to estimate the sensitivity of plume dispersion to these calculated injection heights, two other simulations are performed for each ship, where injection heights are decreased and increased by 50%. Details of the FLEXPART-WRF runs and how they are used to estimate emissions are presented in Sect. 4.

### 3.2 WRF-Chem

In order to estimate the impacts of shipping on air quality and radiative effects in northern Norway, simulations are performed using the 3-D chemical transport model WRF-Chem (Weather Research and Forecasting model, including chemistry, Grell et al., 2005; Fast et al., 2006). WRF-Chem has been used previously by Molders et al. (2010) to quantify the influence of ship emissions on air quality in southern Alaska. Table 2 summarizes all the WRF-Chem options and parameterizations used in the present study, detailed briefly below. The gas phase mechanism is the carbon bond mechanism, version Z (CBM-Z; Zaveri and Peters, 1999). The version of the mechanism used in this study includes dimethylsulfide (DMS) chemistry. Aerosols are represented by the 8 bin sectional MOSAIC (Model for Simulating Aerosol Interactions and Chemistry; Zaveri et al., 2008) mechanism. Aerosol optical properties are calculated by a Mie code within WRF-Chem, based on the simulated aerosol composition, concentrations, and size distributions. These optical properties are linked with the radiation modules (aerosol direct effect), and this interaction also modifies the modeled dynamics and can affect cloud formation (semi-direct effect). The simulations also include cloud-aerosol interactions, representing aerosol activation in clouds, aqueous chemistry for activated aerosols, and wet scavenging within and below clouds. Aerosol activation changes the cloud droplet number concentrations and cloud droplet radii in the Morrison microphysics scheme, thus influencing cloud optical properties (first indirect aerosol effect). Aerosol activation in MOSAIC also influences cloud lifetime by changing precipitation rates (second indirect aerosol effect).

Chemical initial and boundary conditions are taken from the global chemical-transport model MOZART-4 (model for ozone and related chemical tracers version 4; Emmons et al., 2010). In our simulations, the dry deposition routine for trace gases (Wesely, 1989) was modified to improve dry deposition on snow, following the recommendations of Ahmadov et al. (2015). The seasonal variation of dry deposition was also updated to include a more detailed dependence of dry deposition parameters on land use, latitude, and date, which was already in use in WRF-Chem for the MOZART-4 gas-phase mechanism. Anthropogenic emissions (except ships) are taken from the HTAPv2 (Hemispheric transport of air pollution version 2) inventory (0.1° × 0.1° resolution). Bulk VOCs are specified for both shipping and anthropogenic emissions, based on Murrells et al. (2010). Ship VOC emissions are specified using the “other transport” sector (transport emissions, excluding road transport) and anthropogenic VOC emissions are specified using the average speciation for the remaining sectors. DMS emissions are calculated following the methodology of Nightingale et al. (2000) and Saltzman et al. (1993). The oceanic concentration of DMS in the Norwegian Sea in July, taken from Lana et al. (2011), is 5.8 × 10⁻⁶ mol m⁻³. Other biogenic emissions are calculated online by the MEGAN (Model of Emissions of Gases and Aerosols from Nature; Guenther et al., 2006) model within WRF-Chem. Sea salt emissions are also calculated online within WRF-Chem.

The WRF-Chem simulations performed in this study are summarized in Table 3. The CTRL simulation uses the settings and emissions presented above, as well as ship emissions produced by the model STEAM2 (Sect. 3.3). The NO-SHIPS simulation is similar to CTRL, but does not include ship emissions. The NO-SHIPS and CTRL simulations are carried out from 4 to 25 July 2012, over the 15 km × 15 km simulation domain presented in Fig. 1a. The CTRL3 and NO-SHIPS3 simulations are similar to CTRL and NO-SHIPS, but are run on a smaller 3 km × 3 km resolution domain, shown in Fig. 1b, from 10 to 13 July 2012. The CTRL3 and NO-SHIPS3 simulations are not nudged to FNL and do not include a subgrid parameterization for cumulus due to their high resolution. Boundary conditions for CTRL3 and NO-SHIPS3 are taken from the CTRL and NO-SHIPS simulations (using one-way nesting within WRF-Chem) and are updated every hour.

The CTRL and CTRL3 simulations are not nudged to the reanalysis fields in the boundary layer, in order to obtain a more realistic boundary layer structure. However, comparison with ACCESS meteorological measurements
STEAM2 emissions were generated on a 5 km grid every 15 min for the CTRL3 simulation, and were re-gridded on the WRF-Chem simulation grids. Shipping emissions of NO\textsubscript{x}, SO\textsubscript{2}, black carbon, and organic carbon are presented in Fig. 2 for the 15 km × 15 km simulation domain (emissions totals during the simulation period are indicated within the figure panels). For comparison, the HTAPv2 emissions (without shipping emissions) are also shown. Ship emissions are, on average, located in main shipping lanes along the Norwegian coastline. However, they also include less traveled routes, which are apparent closer to shore. Other anthropogenic emissions are mainly located along the Norwegian coast (mostly in southern Norway) or farther inland and to the south in Sweden and Finland. Over the whole domain, NO\textsubscript{x} and OC emissions from shipping are approximately one-third of total anthropogenic NO\textsubscript{x} and OC emissions, but represent a lower proportion of anthropogenic SO\textsubscript{2} and BC emissions (5 and 10 %, respectively). However, other anthropogenic emissions are not co-located with shipping emissions, which represent an important source further north along the coast, as many ships are in transit between European ports and Murmansk in Russia. Very strong SO\textsubscript{2} emissions in Russia are included in the model domain (in the area highlighted in Fig. 2d), associated with smelting activities that occur on the Russian Kola Peninsula (Virkkula et al., 1997; Prank et al., 2010). The Kola Peninsula emissions represent 79 % of the total HTAPv2 SO\textsubscript{2} emissions in the domain.

STEAM2 emissions are based on AIS signals that are transmitted to base stations on shore that have a limited range of 50–90 km, which explains why the emissions presented in Fig. 2 only represent near-shore traffic. In addition, our study is focused on shipping emissions in northern Norway, therefore STEAM2 emissions were only generated along the Norwegian coast. As a result, ship emissions in the northern Baltic and along the northwestern Russian coast are not included in this study. However, these missing

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
<th>Period</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>MET</td>
<td>WRF meteorological simulation, 15 km × 15 km resolution (d01)</td>
<td>4–25 July 2012</td>
<td>Nudged to FNL</td>
</tr>
<tr>
<td>CTRL</td>
<td>WRF-Chem simulation, HTAPv2 anthropogenic emissions, STEAM2 ship emissions, online MEGAN biogenic emissions, online DMS and sea salt emissions, 15 km × 15 km horizontal resolution (d01)</td>
<td>4–25 July 2012</td>
<td>Nudged to FNL in the free troposphere only</td>
</tr>
<tr>
<td>NOSHIJPS</td>
<td>CTRL without STEAM2 emissions, 15 km × 15 km horizontal resolution (d01)</td>
<td>4–25 July 2012</td>
<td>Nudged to FNL in the free troposphere only</td>
</tr>
<tr>
<td>CTRL3</td>
<td>CTRL setup and emissions, 3 km × 3 km horizontal resolution (d02)</td>
<td>10–12 July 2012</td>
<td>Boundary conditions from CTRL</td>
</tr>
<tr>
<td>NOSHIJPS3</td>
<td>NOSHIJPS setup and emissions, 3 km × 3 km horizontal resolution (d02)</td>
<td>10–12 July 2012</td>
<td>Boundary conditions from NOSHIJPS</td>
</tr>
</tbody>
</table>

shows that on 11 July 2012 this leads to an overestimation of marine boundary layer wind speeds (normalized mean bias = +38 %). Since wind speed is one of the most critical parameters in the FLEXPART-WRF simulations, we decided to drive FLEXPART-WRF with the MET simulation instead of using CTRL or CTRL3. In the MET simulation, results are also nudged to FNL in the boundary layer in order to reproduce wind speeds (normalized mean bias of +14 % on 11 July 2012). All CTRL, NOSHIJPS, CTRL3, NOSHIJPS3 and MET simulations agree well with meteorological measurements during the other ACCESS ship flights.

### 3.3 High-resolution ship emissions from STEAM2

STEAM2 is a high-resolution, real-time bottom-up shipping emissions model based on AIS positioning data (Jalkanen et al., 2012). STEAM2 calculates fuel consumption for each ship based on its speed, engine type, fuel type, vessel length, and propeller type. The model can also take into account the effect of waves, and distinguishes ships at berth, maneuvering ships, and cruising ships. Contributions from weather effects were not included in this study, however. The presence of AIS transmitters is mandatory for large ships (gross tonnage ≥ 300 t) and voluntary for smaller ships.

Emissions from STEAM2 are compared with emissions derived from measurements for individual ships in Sect. 4. STEAM2 emissions of CO, NO\textsubscript{x}, OC, BC (technically elemental carbon in STEAM2), sulfur oxides (SO\textsubscript{2}), SO\textsubscript{4}, and exhaust ashes are also used in the WRF-Chem CTRL and CTRL3 simulations. SO\textsubscript{2} emissions are emitted as SO\textsubscript{2} in WRF-Chem, and NO\textsubscript{x} are emitted as 94 % NO, and 6 % NO\textsubscript{2} (EPA, 2000). VOC emissions are estimated from STEAM2 CO emissions using a bulk VOC / CO mass ratio of 53.15 %, the ratio used in the Arctic ship inventory from Corbett et al. (2010). STEAM2 emissions were generated on a 5 km × 5 km grid every 30 min for the CTRL simulation, and on a 1 km × 1 km
Figure 2. (a, c, e, g) STEAM2 ship emissions and (b, d, f, h) HTAPv2 anthropogenic emissions (without ships) of (a, b) NO\textsubscript{x}, (c, d) SO\textsubscript{2}, (e, f) BC, and (g, h) OC in kg km\textsuperscript{-2} over the CTRL and NOSHIPS WRF-Chem domain, during the simulation period (00:00 UTC 4 July 2012 to 00:00 UTC 26 July 2012). On panel (d), the location of the intense Kola Peninsula SO\textsubscript{2} emissions is highlighted by a gray box. The emissions totals for the simulation period are noted in each panel.

Shipping emissions are much lower than other anthropogenic sources inside the model domain. In the CTRL and CTRL3 simulations, ship emissions are injected in altitude using the plume rise model presented in Sect. 3.1. Stack height and exhaust fluxes are unknown for most of the ships present in the STEAM2 emissions, which were not specifically targeted during ACCESS. For these ships, exhaust parameters for the Wilson Leer (∼6000 gross tonnage) are used as a compromise between the smaller fishing ships (∼40% of Arctic shipping emissions; Winther et al., 2014), and larger ships like the ones targeted during ACCESS. In the CTRL3 simulation, the four ships targeted during ACCESS are usually alone in a 3 km × 3 km grid cell, which enabled us to treat these ships separately and to inject their emissions in altitude using individual exhaust parameters (Sect. 3.1). In the CTRL simulation, there are usually several ships in the same 15 km × 15 km grid cell, and the four targeted ships were treated in the same way together with all unidentified ships, using the exhaust parameters of the Wilson Leer and local meteorological conditions to estimate injection heights. This means that, for the Costa Deliziosa, Alaed and Wilson Nanjing, the plume rise model is used in CTRL with exhaust parameters from a smaller ship (the Wilson Leer) than in CTRL3. Because of this, emission injection heights for these ships are lower in CTRL (0 to 30 m) than in CTRL3 (230 m for the Costa Deliziosa, 50 m for the Wilson Nanjing, 30 m for the Wilson Leer, and 65 m for the Alaed).

Primary aerosol emissions from STEAM2 (BC, OC, SO\textsubscript{4}, and ash) are distributed into the eight MOSAIC aerosol bins in WRF-Chem, according to the mass size distribution measured in the exhaust of ships equipped with medium-speed diesel engines by Lyyranen et al. (1999). The submicron mode of this measured distribution is used to distribute primary BC, OC, and SO\textsubscript{4}, while the coarse mode is used to distribute exhaust ash particles (represented as “other inorganics” in MOSAIC).

4 Ship emission evaluation

In this section, emissions of NO\textsubscript{x} and SO\textsubscript{2} are determined for the four ships sampled during ACCESS flights (shown in Table 1). We compare airborne measurements in ship plumes and concentrations predicted by FLEXPART-WRF plume dispersion simulations. In order to derive emission fluxes, good agreement between measured and modeled plume locations is required (discussed in Sect. 4.1). The methods, derived emissions values for the four ships, and comparison with STEAM2 emissions are presented in Sect. 4.2.

4.1 Ship plume representation in FLEXPART-WRF and comparison with airborne measurements

FLEXPART-WRF plume dispersion simulations driven by the MET simulation are performed for the four ships sampled during ACCESS (Sect. 3.1). The MET simulation agrees well with airborne meteorological measurements on both days (shown in the Supplement, Fig. S1) in terms of wind direction (mean bias of −16° on 11 July, +6° on 12 July) and wind speed (normalized mean bias of +14% on 11 July, −17% on 12 July). Figure 3 shows the comparison between maps of the measured NO\textsubscript{x} and plume locations predicted by FLEXPART-WRF. This figure also shows the typical meandering pattern of the plane during ACCESS, measuring the same ship plumes several times as they age, while moving further away from the ship (Roiger et al., 2015). Wilson Leer and Costa Deliziosa plumes were sampled during two different runs at two altitudes on 11 July 2012, and presented in Fig. 3a and b (z = 49 m) and Fig. 3c and d (z = 165 m). During the second altitude level on 11 July (Fig. 3c and d) the Wilson Leer was farther south and the Costa Deliziosa had moved further north. Therefore, the plumes are farther
Modeled air tracer mixing ratios are interpolated in space and time to the aircraft location, and compared with airborne NO\textsubscript{x} and SO\textsubscript{2} measurements (Fig. 4). Each peak in Fig. 4 corresponds to the aircraft crossing the ship plume once during the meandering pattern before turning around for an additional plume crossing. Figure 4a and b only show measurements for the first altitude level at \( z = 49 \text{ m} \) on 11 July 2012 (results for the second altitude level are shown in the Supplement in Fig. S2). As expected from the comparison shown in Fig. 3, modeled peaks are co-located with measured peaks in Fig. 4. The model is also able to reproduce the gradual decrease of concentrations measured in the plume of the Wilson Nanjing on Fig. 4c–e, as the plane flies further away from the ship and the plume gets more dispersed. These peak concentrations vary less for the measured and modeled plume of the Costa Deliziosa (Fig. 4a and b). Measured plumes are less concentrated for the Wilson Leer since it is a smaller vessel, and for the Alaed because its emissions were sampled further away from their source.

### 4.2 Ship emission derivation and comparison with STEAM2

In this section, we describe the method for deriving ship emissions of NO\textsubscript{x} and SO\textsubscript{2} using FLEXPART-WRF and measurements. This method relies on the fact that in the FLEXPART-WRF simulations presented in Sect. 3.1, there is a linear relationship between the constant emission flux of the tracer chosen for the simulation and the tracer concentrations in the modeled plume. The only source of non-linearity that cannot be taken into account is changes in the emission source strength, which is assumed to be constant in time for the plumes sampled. Given that the ship and meteorological conditions were consistent during sampling (shown in the Supplement, Fig. S1), we expect that these effects would be very small. In our simulations, this constant emission flux is picked at \( E = 0.1 \text{ kg s}^{-1} \) and is identical for all ships. This initial value \( E \) is scaled for each ship by the ratio of the measured and modeled areas of the peaks in concentration corresponding to plume crossings, as shown in Fig. 4. Equation (1) shows how SO\textsubscript{2} emissions are derived by this method.

\[
E_i = E \times \frac{\int_{t_i}^{\text{end}} (\text{SO}_2(t) - \text{SO}_2\text{background}) \, dt}{\int_{t_i}^{\text{end}} \text{Tracer}(t) \, dt} \times \frac{M_{\text{SO}_2}}{M_{\text{air}}} \tag{1}
\]

In Eq. (1), \( \text{SO}_2(t) \) is the measured \( \text{SO}_2 \) mixing ratio (pptv), \( \text{SO}_2\text{background} \) is the background \( \text{SO}_2 \) mixing ratio for each peak, Tracer\((t)\) is the modeled tracer mixing ratio interpolated along the ACCESS flight track (pptv), \( t_i^{\text{begin}} \) and \( t_i^{\text{end}} \) are the beginning and end time of peak \( i \) (modeled or measured, in s) and \( M_{\text{SO}_2} \) and \( M_{\text{air}} \) are the molar masses of \( \text{SO}_2 \) and air (g mol\(^{-1}\)). This method produces a different \( \text{SO}_2 \) emission flux value \( E_i \) (kg s\(^{-1}\)) for each of the \( i = 1 \) to \( N \) peaks corresponding to all the crossings of a single ship plume.
by the aircraft. These \( N \) different estimates are averaged together to reduce the uncertainty in the estimated \( \text{SO}_2 \) emissions. A similar approach is used to estimate \( \text{NO}_x \) emissions. The background mixing ratios were determined by applying a 30 s running average to the \( \text{SO}_2 \) and \( \text{NO}_x \) measurements. Background values were then determined manually from the filtered time series. For each \( \text{NO}_x \) peak, an individual background value was identified and used to determine the \( \text{NO}_x \) enhancement for the same plume. For \( \text{SO}_2 \), a single background value was used for each flight leg (constant altitude).

In order to reduce sensitivity to the calculated emission injection heights, FLEXPART-WRF peaks that are sensitive to a \( \pm50\% \) change in injection height are excluded from the analysis. Results are considered sensitive to injection heights if the peak area in tracer concentration changes by more than \( 50\% \) in the injection height sensitivity runs. Using a lower threshold of \( 25\% \) alters the final emission estimates by less than \( 6\% \). Peaks sensitive to the calculated injection height typically correspond to samplings close to the ship, where the plumes are narrow. An intense \( \text{SO}_2 \) peak most likely associated with the \textit{Costa Deliziosa} and sampled around 17:25 UTC on 11 July 2012 is also excluded from the calculations, because this large increase in \( \text{SO}_2 \) in an older, diluted part of the ship plume suggests contamination from another source. \( \text{SO}_2 \) emissions are not determined for the \textit{Wilson Leer} and the \textit{Alaed}, since \( \text{SO}_2 \) measurements in their plumes are too low to be distinguished from the background variability. For the same reason, only the higher \( \text{SO}_2 \) peaks (four peaks \( > 1 \) ppbv) were used to derive emissions for the \textit{Wilson Nanjing}. The number of peaks used to derive emissions for each ship is \( N = 13 \) for the \textit{Costa Deliziosa}, \( N = 4 \) for the \textit{Wilson Leer}, \( N = 8 \) for the \textit{Wilson Nanjing} (\( N = 4 \) for \( \text{SO}_2 \)) and \( N = 5 \) for the \textit{Alaed}.

The derived emissions of \( \text{NO}_x \) (equivalent \( \text{NO}_2 \) mass flux in kg day\(^{-1}\)) and \( \text{SO}_2 \) are given in Table 4. The emissions extracted from the STEAM2 inventory for the same ships and the derived emissions are higher than the value derived for the \textit{Costa Deliziosa}, and lower than the value derived for the \textit{Wilson Nanjing}. \( \text{NO}_x \) emissions from STEAM2 are higher than our calculations for all ships. In STEAM2, the \( \text{NO}_x \) emission factor is assigned according to IMO MARPOL (marine pollution) Annex VI requirements (IMO, 2008) and engine revolutions per minute (RPM), but all engines subject to these limits must emit less \( \text{NO}_x \) than this required value. For the \textit{Wilson Leer}, two calculated values are reported: one calculated by averaging the estimates from the four measured peaks, and one value where an outlier value was removed before calculating the average. During the 11 July flight, the \textit{Wilson Leer} was traveling south at an average speed of 4.5 m s\(^{-1}\), with relatively slow tailwinds of 5.5 m s\(^{-1}\). Because of this, the dispersion of this ship’s plume on this day could be sensitive to small changes in modeled wind speeds, and calculated emissions are more uncertain.

The most important difference between the inventory \( \text{NO}_x \) and our estimates is \( \sim 150\% \) for the \textit{Costa Deliziosa}. Reasons for large discrepancy in predicted and measured \( \text{NO}_x \) emissions of \textit{Costa Deliziosa} were investigated in more detail. A complete technical description of \textit{Costa Deliziosa} was not available, but her sister vessel \textit{Costa Luminosa} was described at length recently (RINA, 2010). The details of \textit{Costa Luminosa} and \textit{Costa Deliziosa} are practically identical and allow for in-depth analysis of emission modeling. With complete technical data, the STEAM2 \( \text{SO}_2 \) and \( \text{NO}_x \) emissions of \textit{Costa Deliziosa} were estimated to be 2684 and 5243 kg day\(^{-1}\), respectively, whereas our derived estimates indicate 2399 and 2728 kg day\(^{-1}\) (difference of \(+12\% \) for \( \text{SO}_2 \) and \(+92\% \) for \( \text{NO}_x \)). The good agreement for \( \text{SO}_2 \) indicates that the power prediction at vessel speed reported in

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Figure 4. (a, c, d) \( \text{NO}_x \) and (b, e) \( \text{SO}_2 \) aircraft measurements (black) compared to FLEXPART-WRF air tracer mixing ratios interpolated along flight tracks, for the plumes of the (a, b) \textit{Costa Deliziosa} and \textit{Wilson Leer} on 11 July 2012 (first constant altitude level \( Z \sim 49 \) m), also shown in Fig. 3a) and (c, d, e) \textit{Wilson Nanjing} and \textit{Alaed} on 12 July 2012. Panel (d) shows the same results as panel (e) in detail. Since model results depend linearly on the emission flux chosen a priori for each ship, model results have been scaled so that peak heights are comparable to the measurements.
AIS and associated fuel flow is well predicted by STEAM2, but emissions of NO\textsubscript{x} are twice as high as the value derived from measurements. In case of \textit{Costa Deliziosa}, the NO\textsubscript{x} emission factor of 10.5 g kWh\textsuperscript{-1} for a tier II compliant vessel with 500 RPM engine is assumed by STEAM2. Based on the measurement-derived value, a NO\textsubscript{x} emission factor of 5.5 g kWh\textsuperscript{-1} would be necessary, which is well below the tier II requirements. It was reported recently (IPCO, 2015) that NO\textsubscript{x} emission reduction technology was installed on \textit{Costa Deliziosa}, but it is unclear whether this technology was in place during the airborne measurement campaign in 2012.

The case of \textit{Costa Deliziosa} underlines the need for accurate and up-to-date technical data for ships when bottom-up emission inventories are constructed. It also necessitates the inclusion of the effect of emission abatement technologies in ship emission inventories. Furthermore, model predictions for individual vessels are complicated by external contributions, like weather and sea currents, affecting vessel performance. However, the STEAM2 emission model is based on AIS real-time positioning data, which has a much better coverage than activity data sets used to generate older shipping emission inventories (e.g., COADS and AMVER). These earlier data sets also have known biases for ships of specific sizes or types. In addition, components of the STEAM2 inventory, such as fuel consumption, engine loads, and emission factors have already been studied in detail in the Baltic Sea by Jalkanen et al. (2009, 2012) and Beecken et al. (2015). Beecken et al. (2015) compared STEAM2 emission factors to measurements for ~300 ships in the Baltic Sea. Their results showed that, while important biases were possible for individual ships, STEAM2 performed much better on average for a large fleet. In the Baltic Sea, STEAM2 NO\textsubscript{x} emission factors were found to be biased by +4\% for passenger ships, based on 29 ships, and −11\% for cargo ships, based on 118 ships. For SO\textsubscript{2}, the biases were respectively +1 and +14\% for the same ships. Therefore, we expect that the large discrepancy in NO\textsubscript{x} for one individual ship (the \textit{Costa Deliziosa}) has only a small impact on the total regional emissions generated by STEAM2. The results presented later in Sect. 5.1 also indicate that STEAM2 likely performs better on average in the Norwegian Sea during ACCESS than for individual ships.

### 4.3 Comparison of STEAM2 to other shipping emission inventories for northern Norway

We compare in Table 5 the July emission totals for NO\textsubscript{x}, SO\textsubscript{2}, BC, OC and SO\textsubscript{2}\textsubscript{y} in northern Norway (latitudes 60.6 to 73° N, longitudes 0 to 31° W) for STEAM2 and four other shipping emission inventories used in previous studies investigating shipping impacts in the Arctic. We include emissions from the Winther et al. (2014), Dalsøren et al. (2009, 2007), and Corbett et al. (2010) inventories. The highest shipping emissions in the region of northern Norway are found in the STEAM2 and Winther et al. (2014) inventories, which are both based on 2012 AIS ship activity data (Sect. 3.3 for a description of the methodology used for STEAM2). We note that, except for OC, the emissions are higher in the Winther et al. (2014) inventory because of the larger geographical coverage: Winther et al. (2014) used both ground-based and satellite retrieved AIS signals, whereas the current study is restricted to data received by ground based AIS stations (capturing ships within 50 to 90 km of the Norwegian coastline). Despite lower coverage, the horizontal and temporal resolutions are better described in land-based AIS networks than satellite AIS data. The terrestrial AIS data used in this study is thus more comparable to the spatial extent and temporal resolution of the measurements collected close to the Norwegian coast. STEAM2 is the only inventory including sulfate emissions, which account for SO\textsubscript{2} to SO\textsubscript{2}\textsubscript{y} conversion in the ship exhaust. Ship emissions from Dalsøren et al. (2009) and Corbett et al. (2010) are based on ship activity data from 2004, when marine traffic was lower than in 2012. Furthermore, the gridded inventory from Corbett et al. (2010) does not include emissions from fishing ships, which represent close to 40\% of Arctic shipping emissions (Winther et al., 2014). These emissions could not be precisely distributed geospatially using earlier methodologies, since fishing ships do not typically follow a simple course (Corbett et al., 2010). Dalsøren et al. (2007) emissions for coastal shipping in Nor-
Inventories are estimated based on Norwegian shipping statistics for the year 2000, and contain higher NO\textsubscript{x}, BC, and OC emissions, but less SO\textsubscript{2}, than the 2004 inventories. This comparison indicates that earlier ship emission inventories usually contain lower emissions in this region, which can be explained by the current growth in shipping traffic in northern Norway. This means that up-to-date emissions are required in order to assess the current impacts of shipping in this region.

5 Modeling the impacts of ship emissions along the Norwegian coast

In this section, WRF-Chem, using STEAM2 ship emissions, is employed to study the influence of ship pollution on atmospheric composition along the Norwegian coast, at both the local (i.e., at the plume scale) and regional scale. As shown in Fig. 4, shipping pollution measured during ACCESS is inhomogeneous, with sharp NO\textsubscript{x} and SO\textsubscript{2} peaks in thin ship plumes, emitted into relatively clean background concentrations. The measured concentrations are on spatial scales that can only be reproduced using very high-resolution WRF-Chem simulations (a few kilometers of horizontal resolution), but such simulations can only be performed for short periods and over small domains. Therefore, high-resolution simulations cannot be used to estimate the regional impacts of shipping emissions. In order to bridge the scale between measurements and model runs that can be used to make conclusions about the regional impacts of shipping pollution, we compare in Sect. 5.1 WRF-Chem simulations using STEAM2 ship emissions, at 3 km × 3 km resolution (CTRL3) and at 15 km × 15 km resolution (CTRL). Specifically, we show in Sect. 5.1 that both the CTRL3 and CTRL simulations reproduce the average regional influence of ships on NO\textsubscript{x}, O\textsubscript{3}, and SO\textsubscript{2}, compared to ACCESS measurements. In Sect. 5.2 we use the CTRL simulation to quantify the regional contribution of ships to surface pollution and shortwave radiative fluxes in northern Norway.

5.1 Model evaluation from the plume scale to the regional scale

It is well known that ship plumes contain fine-scale features that cannot be captured by most regional or global chemical transport models. This fine plume structure influences the processing of ship emissions, including O\textsubscript{3} and aerosol formation, which are non-linear processes that largely depend on the concentration of species inside the plume. Some models take into account the influence of the instantaneous mixing of ship emissions in the model grid box by including corrections to the O\textsubscript{3} production and destruction rates (Huszar et al., 2010) or take into account plume ageing before dilution by using corrections based on plume chemistry models (Vinken et al., 2011). Here, we take an alternative approach by running the model at a sufficient resolution to distinguish individual ships in the Norwegian Sea (CTRL3 run at 3 km × 3 km resolution), and at a lower resolution (CTRL run at 15 km × 15 km resolution). It is clear that a 3 km × 3 km horizontal resolution is not sufficiently small to capture all small-scale plume processes. However, by comparing the CTRL3 simulation to ACCESS measurements, we show in this section that this resolution is sufficient to resolve individual ship plumes and to reproduce some of the plume macroscopic properties. The CTRL and CTRL3 simulations (presented in Table 3) are then compared to evaluate if non-linear effects are important for this study period and region.

WRF-Chem results from CTRL and CTRL3 for surface (∼ 0 to 30 m) NO\textsubscript{x} and O\textsubscript{3} are shown in Fig. 5. On 11 and 12 July, the aircraft specifically targeted plumes from the Wilson Leer, Costa Deliziaosa, Wilson Nanjing and, in addition, sampled emissions from the Alaed, identified later during the post-campaign analysis (Fig. 3). All these ships are individually present in the STEAM2 emissions inventory (Sect. 4 and Table 4). Emissions from these ships, as well as from other vessels traveling in that area, are clearly resolved in the CTRL3 model results for NO\textsubscript{x} (Fig. 5a and e). Ship NO\textsubscript{x} emissions are smoothed out in the CTRL run, seen in Fig. 5b and f, and the individual ship plumes cannot be clearly distinguished in the NO\textsubscript{x} surface concentrations. The predicted surface O\textsubscript{3} concentrations are shown in Fig. 5c, d, g, and h. On the 11 and 12 July 2012, titration of O\textsubscript{3} by NO\textsubscript{x} from fresh ship emissions can be identified in Fig. 5c and g for the 3 km run (areas indicated by black arrows on Fig. 5c.

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Table 5. July emission totals in northern Norway (60.6–73° N, 0 to 31° W) of NO\textsubscript{x}, SO\textsubscript{2}, BC, OC, and SO\textsubscript{2} in different ship emission inventories.

<table>
<thead>
<tr>
<th>Inventory</th>
<th>Year</th>
<th>NO\textsubscript{x} (kt)</th>
<th>SO\textsubscript{2} (kt)</th>
<th>BC (t)</th>
<th>OC (t)</th>
<th>SO\textsubscript{2} (t)</th>
</tr>
</thead>
<tbody>
<tr>
<td>STEAM2</td>
<td>2012</td>
<td>7.1</td>
<td>2.4</td>
<td>48.1</td>
<td>123.4</td>
<td>197.3</td>
</tr>
<tr>
<td>Winther et al. (2014)</td>
<td>2012</td>
<td>9.3</td>
<td>3.4</td>
<td>47.7</td>
<td>82.9</td>
<td>–</td>
</tr>
<tr>
<td>Dalsøren et al. (2009)</td>
<td>2004</td>
<td>3.1</td>
<td>1.9</td>
<td>7.3</td>
<td>24.5</td>
<td>–</td>
</tr>
<tr>
<td>Corbett et al. (2010)</td>
<td>2004</td>
<td>2.4</td>
<td>1.6</td>
<td>10.6</td>
<td>32.5</td>
<td>–</td>
</tr>
<tr>
<td>Dalsøren et al. (2007)</td>
<td>2000</td>
<td>5.5</td>
<td>1.1</td>
<td>24.</td>
<td>479.3</td>
<td>–</td>
</tr>
</tbody>
</table>
Figure 5. Snapshots of model predicted surface NO\textsubscript{x} and O\textsubscript{3} from the CTRL3 (3 km) simulation (a, c, e, g) and the CTRL (15 km) simulation (b, d, f, h) during the flights on 11 and 12 July 2012. Model results for the CTRL3 simulation are shown over the full model domain. CTRL run results are shown over the same region for comparison. The aircraft flight tracks are indicated in blue. On panels (e) and (g), black arrows indicate several areas of O\textsubscript{3} titration due to high NO\textsubscript{x} from ships. (i, j) NO\textsubscript{x} and (k, l) O\textsubscript{3} 2-day average surface enhancements (00:00 UTC 11 July 2012 to 00:00 UTC 13 July 2012) due to shipping emissions, (i, k) CTRL3 simulation, (j, l) CTRL simulation. The 2-day average enhancements of NO\textsubscript{x} and O\textsubscript{3} over the whole area are given below each respective panel.

However, evidence for O\textsubscript{3} titration quickly disappears away from the fresh emissions sources. In contrast, O\textsubscript{3} titration is not apparent in the CTRL run. However, NO\textsubscript{x} and O\textsubscript{3} patterns and average surface concentrations are very similar. This is illustrated in the lower panels, showing 2-day-averaged NO\textsubscript{x} and O\textsubscript{3} enhancements due to ships in the CTRL3 (CTRL3 – NOSHIPS3) and CTRL (CTRL – NOSHIPS) simulations. The results show that changing the horizontal resolution from 3 km \times 3 km (1 km \times 1 km emissions, 15 min emissions injection) to 15 km \times 15 km (5 km \times 5 km emissions, 1 h emissions injection) does not have a large influence on the domain-wide average NO\textsubscript{x} (−3.2 \%) or O\textsubscript{3} (+0.08 ppbv, +4.2 \%) enhancements due to ships. This is in agreement with earlier results by Cohan et al. (2006), who showed that regional model simulations at similar resolutions (12 km) were sufficient to reproduce the average O\textsubscript{3} response. Results by Vinken et al. (2011) suggest that simulations at a lower resolution more typical of global models (2° \times 2.5°) would lead to an overestimation of O\textsubscript{3} production from ships in this region by 1 to 2 ppbv. The influence of model resolution on surface aerosol concentrations is also moderate, and PM\textsubscript{10} due to ships are 15 \% lower on average in CTRL than in CTRL3 (not shown here).

To further investigate the ability of these different model runs to represent single ship plumes, we compare measured NO\textsubscript{x}, SO\textsubscript{2}, and O\textsubscript{3} along the flight track on 11 July 2012 with WRF-Chem predictions (Fig. 6). Corresponding results for 12 July 2012 are shown in the Supplement (Fig. S3). Large enhancements of NO\textsubscript{x} and SO\textsubscript{2} are seen during plume crossings in measurements, as already noted in Sect. 4. For comparison with WRF-Chem, we have averaged the measured data using a 56 s running average, equivalent to the aircraft crossing 6 km (two model grid cells) at its average speed during this flight (107 m s\textsuperscript{−1}). Using a running average takes into account plume dilution in grid cells, as well as additional smoothing introduced when modeled results are spatially interpolated onto the flight track. The CTRL3 simulation captures both the width and magnitude of NO\textsubscript{x} and SO\textsubscript{2} peaks, suggesting that the individual plumes are correctly represented in space and time. During the second part of the flight (17:20 UTC), the model does not reproduce two intense measured SO\textsubscript{2} peaks. We already noted in Sect. 4.2 that measurements in this part of the flight might be contaminated by another source. In contrast, the CTRL run has wider NO\textsubscript{x} and SO\textsubscript{2} peaks and lower peak concentrations, because of dilution in larger grids. Another difference between the simulations is the treatment of plume rise (Sect. 3.3), such
that the Costa Deliziosa plume is located at lower altitudes in CTRL than in CTRL3. The CTRL3 simulation tends to overestimate NO\textsubscript{x} in ship plumes, which is in agreement with the results shown in Table 4, indicating that STEAM2 NO\textsubscript{x} emissions are overestimated for the ships targeted during ACCESS. This overestimation is unlikely to be caused by chemistry issues, since an overestimated NO\textsubscript{x} lifetime would lead to comparatively larger biases at the end of the constant altitude runs, when older parts of the plume were sampled. Figure 6b shows O\textsubscript{3} during the same flight. The CTRL3 simulation reproduces the ozone variability better than the CTRL run, but both runs perform relatively well on average (mean bias = −3 ppbv during the constant altitude legs). This negative bias is due to a small underestimation in the background ozone, which could be caused by a number of reasons, including the boundary chemical conditions from the MOZART4 model, photolysis rates, cloud properties and locations, ozone deposition, and/or emissions. Both measurements and CTRL3 results show evidence of O\textsubscript{3} titration in the most concentrated NO\textsubscript{x} plumes, where ozone is 1.5 to 3 ppbv lower than out of the plumes. However, precise quantification of this titration is difficult because these values are the same order of magnitude as the spatial variability of O\textsubscript{3} outside of the plumes. O\textsubscript{3} titration is not apparent in the CTRL run. Results are similar for the 12 July 2012 flight (shown in the Supplement, Fig. S3), with lower model biases for O\textsubscript{3} but a stronger overestimation of NO\textsubscript{x}.

In order to evaluate modeled aerosols in ship plumes, modeled aerosols are evaluated using size distributions measured during the 11 July 2012 flight. Size distributions are integrated to estimate submicron aerosol mass (PM\textsubscript{1}), assuming a density of 1700 kg m\textsuperscript{-3} and spherical particles. This indicates that observed PM\textsubscript{1} enhancements in plumes (≈0.1 to 0.5 µg m\textsuperscript{-3}) are relatively low compared to background PM\textsubscript{1} (≈0.7 to 1.1 µg m\textsuperscript{-3}), because of the presence of high sea salt concentrations in the marine boundary layer (54% of the modeled background PM\textsubscript{1} during ship plume sampling is sea salt in NOSHIPS3). Because of this, comparing modeled and observed in-plume PM\textsubscript{1} directly would be mostly representative of background aerosols, especially sea salt, which is not the focus of this paper. Figure 7 shows the comparison between modeled and measured enhancements in PM\textsubscript{1} in the plume of the Costa Deliziosa (11 July 2012), removing from the model and measurements the contribution from sea salt and other aerosols not associated with shipping. Similarly to Fig. 6, a 56 s moving average was applied to the measurement (representing plume dilution in the model grid). This comparison indicates a generally good agreement between modeled and measured enhancements in PM\textsubscript{1} in ship plumes. There is a discrepancy between the model and the measurements for the first two PM\textsubscript{1} plumes measured close to the ships (around 16:05 UTC), which could be an artifact of the limited resolution of this simulation (3 km). If these peaks are excluded, the model slightly overestimates peak PM\textsubscript{1} enhancements in ship plumes (+26%). Since this enhancement is modeled as 80% SO\textsubscript{2}, this overestimation can be linked to the +37% overestimation of SO\textsubscript{2} emissions for the Costa Deliziosa in STEAM2 (Table 4).

Analysis of O\textsubscript{3} maps, average surface enhancements due to ships (Fig. 5) and analysis of model results along flight tracks (Fig. 6) show that both runs capture the NO\textsubscript{x} and O\textsubscript{3} concentrations in this region reasonably well. Further-
more, Fig. 7 shows that PM$_1$ enhancements in ship plumes are well reproduced in the CTRL3 simulation, and we found that PM$_{10}$ production from ships over the simulation domain was not very sensitive to resolution. This suggests that the CTRL simulation is sufficient to assess the impacts of ship emissions at a larger scale during July 2012. This is investigated further by comparing modeled NO$_x$, SO$_2$, and O$_3$ in the CTRL and NOSHIPS simulations with the average vertical profiles (200–1500 m) measured during four ACCESS flights from 11 to 25 July 2012 (flights shown in Fig. 1a); this comparison is shown in Fig. 8. Modeled vertical profiles of PM$_{2.5}$ are also shown in Fig. 8. This comparison allows us to estimate how well CTRL represents the average impact of shipping over a larger area and a longer period.

Figure 8 shows that the NOSHIPS simulation significantly underestimates NO$_x$ and SO$_2$, and moderately underestimates O$_3$ along the ACCESS flights, indicating that ship emissions are needed to improve the agreement between the model and observations. In the CTRL simulation, NO$_x$, SO$_2$, and O$_3$ vertical structure and concentrations are generally well reproduced, with normalized mean biases of $+14.2$, $-6.8$, and $-7.0\%$, respectively. Correlations between modeled (CTRL) and measured profiles are significant for NO$_x$ and O$_3$ ($r^2 = 0.82$ and 0.90). However, the correlation is very low between measured and modeled SO$_2$ ($r^2 = 0.02$), and it is not improved compared to the NOSHIPS simulation. Ships have the largest influence on NO$_x$ and SO$_2$ profiles, a moderate influence on O$_3$ and do not strongly influence PM$_{2.5}$ profiles along the ACCESS flights. However, this small increase in PM$_{2.5}$ corresponds to a larger relative increase in sulfate concentrations and in particle numbers in the size ranges typically activated as cloud condensation nuclei (shown in Fig. S4 in the Supplement).

NO$_x$ concentrations are overestimated in the parts of the profile strongly influenced by shipping emissions. This is in agreement with the findings of Sect. 4.2, showing that STEAM2 NO$_x$ emissions were overestimated for the ships sampled during ACCESS. However, the CTRL simulation performs well on average, suggesting that the STEAM2 inventory is able to represent the average NO$_x$ emissions from ships along the northern Norwegian coast during the study period. The bias for SO$_2$ is very low compared to results from Eyring et al. (2007), which showed that global models significantly underestimated SO$_2$ in the polluted marine boundary layer in July. Since aerosols from ships contain mostly secondary sulfate formed from SO$_2$ oxidation, the validation of modeled SO$_2$ presented in Fig. 8 also gives some confidence in our aerosol results compared to earlier studies investigating the air quality and radiative impacts of shipping aerosols. We therefore use the 15 km $\times$ 15 km CTRL run for further analysis of the regional influence of ships on pollution and the shortwave radiative effect in this region in Sect. 5.2.

5.2 Regional influence of ship emissions in July 2012

5.2.1 Surface air pollution from ship emissions in northern Norway

The regional-scale impacts of ships on surface atmospheric composition in northern Norway are estimated by calculating the 15-day (00:00 UTC, 11 July 2012 to 00:00 UTC, 26 July 2012) average difference between the CTRL and NOSHIPS simulations. Figure 9 shows maps of these anom-
lies at the surface, for SO$_2$, NO$_x$, and O$_3$. Ship emissions have the largest influence on surface NO$_x$ and SO$_2$ concentrations, with 75 to 100 % increases along the coast. Average O$_3$ increases from shipping are $\sim$ 6 % ($\sim$ 1.5 ppbv) in the coastal regions, with slightly lower enhancements ($\sim$ 1 ppbv, $\sim$ 4 %,) further inland over Sweden.

Dalsøren et al. (2007) studied the impact of maritime traffic in northern Norway in the summer using ship emission estimates for the year 2000. They found, for July 2000, a 1 to 1.5 % increase in surface O$_3$ from coastal shipping in Norwegian waters. However, unlike the present study, the estimate of Dalsøren et al. (2007) did not include the impact of international transit shipping along the Norwegian coast. Our estimated impact on O$_3$ in this region (6 % and 1.5 ppbv increase) is about half of the one determined by Ødemark et al. (2012) (12 % and 3 ppbv), for the total Arctic fleet in the summer (June–Aug–Sept) 2004, using ship emissions for the year 2004 from Dalsøren et al. (2009). It is important to note that we expect lower impacts of shipping in studies based on earlier years, because of the continued growth of shipping emissions along the Norwegian coast (as discussed in Sect. 4.3 and illustrated in Table 5). However, stronger or lower emissions do not seem to completely explain the different modeled impacts. Ødemark et al. (2012) found that Arctic ships had a strong influence on surface O$_3$ in northern Norway for relatively low 2004 shipping emissions. This could be explained by the different processes included in both models, or by different meteorological situations in the two studies based on two different meteorological years (2004 and 2012). However, it is also likely that the higher O$_3$ in the Ødemark et al. (2012) study could be caused, in part, by nonlinear effects associated with global models run at low resolutions. For example Vinken et al. (2011) estimated that instant dilution of shipping NO$_x$ emissions in $2^\circ \times 2.5^\circ$ model grids leads to a 1 to 2 ppbv overestimation in ozone in the Norwegian and Barents seas during July 2005. This effect could explain a large part of the difference in O$_3$ enhancements from shipping between the simulations of Ødemark et al. (2012) ($2.8^\circ \times 2.8^\circ$ resolution) and the simulations presented in this paper ($15 \text{ km} \times 15 \text{ km}$ resolution).

The impact of ships in northern Norway on surface PM$_{2.5}$, BC, and SO$_2$ during the same period is shown in Fig. 10. The impact on PM$_{2.5}$ is relatively modest, less than 0.5 µg m$^{-3}$. However, these values correspond to an important relative increase of $\sim$ 10 % over inland Norway and Sweden because of the low background PM$_{2.5}$ in this region. Over the sea surface, the relative effect of ship emissions is quite low because of higher sea salt aerosol background. Aliabadi et al. (2014) have observed similar increases in PM$_{2.5}$ (0.5 to 1.9 µg m$^{-3}$) in air masses influenced by shipping pollution in the remote Canadian Arctic. In spite of the higher traffic in northern Norway, we find lower values than Aliabadi et al. (2014) because results in Fig. 10 are smoothed by the 15-day average. Impacts on surface sulfate and BC concentrations are quite large, reaching up to 20 and 50 %, respectively. We note that Eckhardt et al. (2013) found enhancements in summertime equivalent BC of 11 % in Svalbard from cruise ships alone. As expected, absolute SO$_2$ and BC enhancements in our simulations are higher in the southern part of the domain, where ship emissions are the strongest. We estimated the lifetime (residence time) of BC originating from ship emissions using the method presented in Samset et al. (2014). This residence time is defined as the ratio of the average BC burden from ships divided by the average BC emissions in STEAM2.

Figure 9. 15-day average (00:00 UTC 11 July 2012 to 00:00 UTC 26 July 2012) of (top) absolute and (bottom) relative surface enhancements (CTRL – NOSHIPS) in (a, d) SO$_2$, (b, e) NO$_x$, and (c, f) O$_3$ due to ship emissions in northern Norway from STEAM2.
during the simulation. Using this method, we find a BC lifetime of 1.4 days. This short lifetime can be explained by the negative sea level pressure anomalies over northern Norway during the ACCESS campaign (Roiger et al., 2015), which indicates more rain and clouds than normal during summer. Given this short lifetime, BC is not efficiently transported away from the source region.

5.2.2 Shortwave radiative effect of ship emissions in northern Norway

The present-day climate effect of ship emissions is mostly due to aerosols, especially sulfate, which cool the climate through their direct and indirect effects (Capaldo et al., 1999). However, large uncertainties still exist concerning the magnitude of the aerosol indirect effects (Boucher et al., 2013). In this section, we determine the total shortwave radiative effect of ships by calculating the difference between the top-of-atmosphere (TOA) upwards shortwave (0.125 to 10 µm wavelengths) radiative flux in the CTRL and the NOSHIPS simulations. Since the CTRL and NOSHIPS simulations take into account aerosol–radiation interactions and their feedbacks (the so-called direct and semi-direct effects) as well as cloud–aerosol interactions (indirect effects), this quantity represents the sum of modeled direct, semi-direct and indirect effects from aerosols associated with ship emissions. Yang et al. (2011) and Saide et al. (2012) showed that including cloud aerosol couplings in WRF-Chem improved significantly the representation of simulated clouds, indicating that the indirect effect was relatively well simulated using CBM-Z/MOSAIC chemistry within WRF-Chem. Our calculations do not include the effect of BC on snow, since this effect is not currently included in WRF-Chem.

The shortwave radiative effect at TOA of in-domain ship emissions is $-1.77 \text{ W m}^{-2}$ (15-day average). We multiply this value by the area of our simulation domain to obtain a forcing value in watts (W), and divide it by the surface area of the Earth in order to obtain an equivalent global radiative effect in m W m$^{-2}$ that can be compared to results from global studies. This equivalent global radiative effect at TOA is $-9.3 \text{ m W m}^{-2}$. This value is strongly negative, indicating that ship emissions cause a net cooling effect in this region (likely due to sulfate) despite the strong relative increase in BC concentrations from shipping emissions (up to $+50\%$, Fig. 10). This can be explained by the fact that these strong relative enhancements in BC correspond to low absolute values (at most 20 ng m$^{-3}$) above very low background concentrations.

The radiative effect calculated in this study, $-9.3 \text{ m W m}^{-2}$, is similar to the estimate by Ødemark et al. (2012), who found a direct and indirect shortwave effect of aerosols from Arctic-wide shipping in July 2004 of $-10.4 \text{ m W m}^{-2}$. However, since the present study only represents the effect of shipping along the Norwegian coast, this implies that current ship emissions in northern Norway have a stronger effect in this study than in Ødemark et al. (2012), which was based on ship emissions from Dalsøren et al. (2009) corresponding to 24 % less SO$_2$ emissions than STEAM2. Higher emissions in our simulations could explain the stronger local shortwave effect of Arctic ships, since this effect is mostly associated with the direct and indirect effect of sulfate aerosols. However, the total sulfate column due to ship emissions in our study is 100 to
200 μg m⁻² along the Norwegian coast, about half of the value (250 to 300 μg m⁻²) found by Ødemark et al. (2012). This means that the stronger effect found here is not due to increased sulfate concentrations from larger emissions, but is likely due to the way aerosol–cloud interactions are treated in both models: the indirect aerosol effect was calculated by Ødemark et al. (2012) based on parameterizations of the relationship between clouds droplet numbers and aerosol mass, whereas the MOSAIC aerosol module used in this study explicitly treats aerosol activation within clouds and their impacts on cloud properties (Yang et al., 2011). It is also important to note here that the indirect radiative effect of shipping emissions is uncertain and that the difference between the estimate of Ødemark et al. (2012) and the one in this work can also be explained by these uncertainties. Based on the work of Eyring et al. (2007), Lauer et al. (2007), and Fuglestvedt et al. (2008), Eyring et al. (2010) estimated that the radiative forcing of global shipping emissions was −0.408 W m⁻², but found an uncertainty range of ±0.425 W m⁻². Ødemark et al. (2012) considered that the uncertainty in the indirect effect in their simulations was the same as the uncertainty in the global indirect forcing of aerosols as estimated by the IPCC (Forster et al., 2007, Table 2.12). Using this method, Ødemark et al. (2012) estimated a range of [−3.9 m W m⁻², −1.3 m W m⁻²] for the global and annual indirect effect of Arctic shipping emissions. It is important to better understand and constrain this effect, which would require more aerosol measurements in shipping lanes (including number concentrations and aerosol compositions in ship plumes) and more model case studies.

6 Conclusions

The focus of this work, linking modeling and measurements, is to better quantify regional atmospheric impacts of ships in northern Norway in July 2012. The study relies on measurements from the ACCESS aircraft campaign, emissions evaluation, and regional modeling in order to evaluate both individual ship plumes and their regional-scale effects. STEAM2 emissions, which represent individual ships based on high-resolution AIS ship positioning data, are compared with emissions for specific ships derived from measurements and plume dispersion modeling using FLEXPART-WRF. Regional WRF-Chem simulations run with and without ship emissions are performed at two different resolutions to quantify the surface air quality changes and radiative effects from ship emissions in northern Norway in July 2012. The most important conclusions from our study are

1. Validation of the STEAM2 emissions – emissions of NOₓ and SO₂ are determined for individual ships, by comparing airborne measurements with plume dispersion modeling results. These calculated emissions are compared with bottom-up emissions determined for the same ships by the STEAM2 emission model. Results show that STEAM2 overestimates NOₓ emissions for the four ships sampled during ACCESS. SO₂ emissions are also determined for two ships. Large biases are possible for individual ships in STEAM2, especially for ships for which there is incomplete technical data or where emission reduction techniques have been employed. Nevertheless, combining WRF-Chem simulations and STEAM2 emissions leads to reasonable predictions of NOₓ, SO₂, and O₃ compared to ACCESS profiles in the lower troposphere (normalized mean biases of +14.2, −6.8, and −7.0 %, respectively). These results also indicate that shipping emissions comprise a significant source of NOₓ and SO₂ at low altitudes during the ACCESS flights, even though specific ship plume sampling near the surface was excluded from these profiles. Pollution sampled during these flights thus represents shipping pollution that had time to mix vertically in the marine boundary layer and is more representative of the regional pollution from shipping in northern Norway. These results are in agreement with the recent evaluation of STEAM2 in the Baltic Sea by Beecken et al. (2015), which showed that STEAM2 performed well for an average fleet (~200 ships), despite biases for individual ships.

2. Regional model representation of ship plumes and their local-scale influence – WRF-Chem runs including shipping emissions from STEAM2 are performed at 15 km × 15 km and 3 km × 3 km horizontal resolutions, and compared with airborne measurements of NOₓ and ozone. The high-resolution simulation is better at reproducing measured NOₓ peaks and suggests some ozone titration in ship plumes, but the NOₓ and ozone enhancements due to ships in both simulations are within less than 5 % of each other when averaged over the whole domain and simulation period. The 3 km × 3 km simulation also reproduces observed PM₁ enhancement in ship plumes. Surface PM₁₀ enhancements due to ships are 15 % higher in the 3 km × 3 km resolution simulation.

3. Average influence of ship pollution in July 2012 – the difference between runs with and without ship emissions are compared with campaign average profiles (excluding flights focused on oil platforms, smelters, and biomass burning emissions from outside the simulation domain). Including STEAM2 emissions reduces the mean bias between measured and modeled trace gases NOₓ, SO₂, and O₃. At the surface, ship emissions enhance 15-day-averaged concentrations along the Norwegian coast by approximately 80 % for NOₓ, 80 % for SO₂, 5 % for O₃, 40 % for BC, and 10 % for PM₂.₅, suggesting that these emissions are already having an impact on atmospheric composition in this region. Regional model results presented in this study predict
lower ozone production from ships compared to certain earlier studies using global models. However, it is known that global models run at low resolution tend to overestimate ozone production (underestimate ozone titration) from fresh ship emissions because of nonlinearities introduced when diluting concentrated emissions from ships into coarse model grid cells.

4. Influence on the radiative budget – northern Norwegian ship emissions contribute $-9.3 \text{ mW m}^{-2}$ to the global shortwave radiative budget of ship emissions, including semi-direct and indirect effects. These results are more significant than found previously in a study using a global model that did not explicitly resolve aerosol activation in clouds. This suggests that global models may be underestimating the radiative impacts of shipping in this region.

Our study shows that local shipping emissions along the northern Norwegian coast already have a significant influence on regional air quality and aerosol shortwave radiative effects. As Arctic shipping continues to grow and new regulations are implemented, the magnitude of these impacts is expected to change. Due to the limited region (northern Norway) and the short timescale (15 days) considered here, it is not possible to assess the radiative effect of other climate forcers associated with shipping in northern Norway, including O$_3$ which global model studies have suggested are also significant (Dalsøren et al., 2013; Ødemark et al., 2012). However, since shipping emissions are highly variable and localized, quantifying impacts using global models can be challenging. Our approach used a regional chemical transport model at different scales, with high-resolution ship emissions, to evaluate model results against observations and estimate the regional impact of shipping emissions. In the future, additional work is needed in other regions and at different spatial scales (measurements and modeling) in order to investigate the impacts of shipping over the wider Arctic area.

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