
Reply to Reviewer 1:

The authors would like to thank to Reviewer 1 for a thorough review of the manuscript and for the constructive comments. Here are our responses:

1. **Line 35-36: Is this the summer mean or annual mean, please rephrase the sentence. Please provide relative contributions (%) along with absolute contributions**

Response:

It should be summer mean. The sentence has been rephrased as “The local shipping emissions of NOx led to a decrease of the summer mean O3 levels in the city by 0.5 ppb (~2%) in average.”

2. **Line 157: “exposure-response function”**

Response:

The typo has been corrected

3. **Line 296: What is spatial resolution of the SMED database that is the source of these “other” emissions and how are they regrded into 1x1 km resolution? In addition, how are there emissions regridded to the TAPM resolution of 250 m?**

Response:

The spatial resolution of the SMED database is 1x1 km. The SMED gridded emissions for the different source categories were applied directly as gridded sources in the model.

4. **Section 2.2.2: Does road traffic includes resuspension so that it is the largest PM10 source in the domain?**

Response:

The road-traffic emissions include the wear particles, however, not the re-suspension. This fact has been added to the methods part.

5. **Section 2.4. Why no model evaluation for PM2.5? Section 3.2.4 includes some discussion on modelled vs measured PM2.5, why not include these in the model evaluation section? It is very important as PM2.5 is the main health impact pollutant and errors in PM2.5 simulations lead to underestimations in the health impacts.**

Response:

We would like to thank the reviewer for pointing out this omission. Model evaluation for PM2.5 has been added to the text and Figure 5 extended with panel for summary statistics for PM2.5. In addition, section on comparing measured versus modeled daily concentrations of NO2, O3, PM10 and PM2.5 has been included in the supplement in response to a comment of Reviewer 2.
6. **Section 2.5: Are the age intervals taken into account? If not, please discuss potential shortcomings. How are the chronic vs acute impacts taken into account?**

Response:

We would like to thank the reviewer for pointing out these deficiencies in description of the HIA methodology. Both aspect of the age intervals and form of the ERF for PM$_{2.5}$ have been added to Section 2.5:

ARP uses linear ERFs, recognizing the limited range of pollutant exposures in Europe. The YOLLs are calculated per year, applying the relative risk within national life tables. This is done through relation between life years lost per 100 000 population per unit PM$_{2.5}$ concentration and life expectancy of the population developed by Miller et al. (2003) based on analysis of life tables. The premature deaths are calculated using the total national mortality rate. This methodology is justified for European countries with health status and proportion of natural mortality of population corresponding to population studied in the epidemiological studies which brought forward the CRFs for all-cause mortalities. For regions with high concentration levels of PM$_{2.5}$ the HIA studies need to use different form of ERFs and for populations with different health status comparing to the US and Western Europe, cause-specific rather than all-cause mortalities need to be used.

7. **Section 3.2. Please provide with relative contributions along with absolute contributions throughout the text.**

Response:

The relative contributions were added for SO$_2$ concentrations (3.2.1), NO$_2$ (3.2.2.) and ozone (in Conclusions). For PM (3.2.4) the relative contributions were already presented.

8. **Section 5. How about the linearity of the ERF? There are studies clearly showing that assuming a linear relationship can lead to significant under or over estimation of health impacts depending on the concentration range. This should be discussed, I think.**

Response: The discussion on ERF model as well as on use of all-cause or cause-specific mortality ERFs was added to Section 5:

In ARP a linear form of ERFs is applied which is justified by a rather narrow interval of PM exposure levels in Europe. In terms of impact of the total exposure to PM$_{2.5}$ on natural mortality, the linear and log-linear form of the functions give similar results within the concentration range of 10–30 µg m$^{-3}$, the linear model giving slightly lower relative risks in this range and higher relative risks below and above (Ostro et. al., 2004). The PM$_{2.5}$ levels found in our study fall below 10 µg m$^{-3}$. For regions with high PM$_{2.5}$ levels different ERF models need to be applied and for HIA global studies or studies...
in other regions bur Western Europe or North America also ERFs for cause-specific mortalities, rather than natural mortalities are usually used.

In terms of incremental effects, the impacts can differ substantially between the two models at different concentration levels. Sofiev et al. (2018) show difference in relative risks of cause-specific mortalities for different base concentrations, at 1 µg m$^{-3}$ the log-linear model gives higher incremental relative risks than the linear model while at 5 and 10 µg m$^{-3}$ levels the log-linear model gives lower incremental relative risks.

9. Brandt et al., 2013 is not cited in the text.

Response:
The reference has been deleted.

10. Figure 8 could be made similar to figure 11, showing the monthly means as it is a bit crowded as it is now. In addition, both figures could use stacked bars instead. Finally, it would be great to create a similar figure where it shows the contributions from other pollutants as “others”, that can be split into local and outside of Gothenburg and Sweden if possible, as in Im et al., ACP, 2019.

Response:
The Figure 8 as well as Figure 11 has been updated by showing the contributions to monthly mean NO$_2$ and PM$_{2.5}$ concentrations from local shipping, regional shipping, road traffic and others.

![Figure 8: Modelled monthly mean contributions of the local shipping, regional shipping, local road traffic and other anthropogenic emissions (including contribution from the boundary conditions) to the NO$_2$ concentrations (ppb) at Eriksberg in year 2012.](image-url)
Reply to Reviewer 2:

We would like to thank the reviewer for a thorough review of the manuscript and for many good points and suggestions for improvement. We have MET these valuable comments mainly by including additional information to the manuscript and believe that it gained more clarity, especially in terms of the modelling methodology employed and its verification. In the following text, the comments are answered and the changes in the manuscript indicated. The Response includes some new citations. Those which are not included in the manuscript are included at the end of the Response.

**Major comments:**

1. *About the model set-up I feel some information about the advection and diffusion needs to be described, so that people can understand how the air pollutants transport horizontally and vertically over the Eulerian grid. What is the model top pressure? How many layers are there in the TAPM?*

   Response:
   
   We would like to thank the reviewer for pointing out need of better description of the model. We have extended this part with more details and added information on vertical layers to the place where the model domain is described:

   TAPM consists of a meteorological and an air pollution components. The meteorological component of TAPM is an incompressible, non-hydrostatic, primitive equation model with a terrain-following vertical sigma coordinate for 3-D simulations. The model solves the momentum equations for horizontal wind components, the incompressible continuity equation for vertical velocity, and scalar equations for potential virtual temperature and specific humidity of water vapour, cloud water/ice, rain water and snow. The turbulence terms in these equations have been determined by solving equations for turbulence kinetic energy and eddy dissipation rate, and then using these values to represent vertical fluxes by a gradient diffusion approach (Hurley, 2008 b). Using predicted meteorology and turbulence from the meteorological component, TAPM applies Eulerian grid module in its air pollution component which consists of nested grid-based solutions of the Eulerian concentration mean equations representing advection, diffusion, chemical reactions and emissions. Dry and wet deposition processes are also included. (l. 227-235)

   In TAPM, an Exner pressure function is integrated from mean sea level to the model top (10 Pa in this study) to determine the top boundary condition. The Exner pressure function is determined from the sum of the hydrostatic component and non-hydrostatic component (Hurley, 2008). The number of vertical grid levels was 30 in this study. Twenty of these layers are below approximately 2 km; the lowest layer extends to ca. 10 m above ground. (l. 248-251)
2. *The authors mentioned that only simple formation of secondary inorganic and organic aerosol exist in TAMP. How does that affect the simulation of PM2.5 and PM10? Any underestimation? I would imagine the ability of TAMP in reproducing particulate matters may not be as well as CMAQ. Please refer to my second concern, in which I strongly suggest a comparison be made between CMAQ and TAMP.*

Response:
The secondary aerosol formation in TAPM is heavily parameterized, however, captures the important features of the secondary particle formation, i.e. formation of sulphate and nitrate following the SO$_2$ and NO$_2$ oxidation, as well as formation of SOA as a fixed part of the degraded smog reactivity representing VOC species in the reaction scheme of TAPM (Hurley, 2008b). We have replaced the description of aerosol formation in TAPM in the PAPER with this wording.

On urban scale, formation of secondary PM is usually suppressed as the radical pool is depleted by the primary emissions and many urban models do not consider the secondary PM at all. We recognise that this assumption is questionable for shipping emissions, which are often emitted into relatively clean air masses coming from the sea over the harbour area to the city, additionally, also chemistry involving sea-salt aerosol particles can be of importance in this case. We have investigated contribution of secondary PM to the total PM modelled with TAPM photochemistry scheme, Fig. S4 shows contributions of max. 2% of the PM related to the local shipping in Gothenburg in winter months and negligible contributions in summer. In an earlier study (Haeger-Eugensson et al., 2010) we have compared oxidation processes in a ship plume transported over Gothenburg area simulated with TAPM photochemistry scheme and with much more detailed scheme including explicit aerosol chemistry of the MOCCA model (Sander et al., 1996, Pszenny et al., 2004) and found that during the day time the 2 schemes gave similar results while in dark hours the NO2 oxidation was underestimated, mainly due to the missing night-time NO3 chemistry. The MOCCA scheme does not, however, involve any advanced SOA chemistry so the performance of the two schemes regarding the SOA formation was not investigated.

The main idea of our city-scale study utilizing the boundary conditions of the CMAQ model simulations is to assess the urban-scale features of the shipping emissions, including differentiation of the regional shipping and the local shipping contributions to air pollution in the city. The regional-scale secondary PM formation is captured by the CMAQ model and is transferred to the local scale through the boundary concentration fields. CMAQ includes both, secondary inorganic (SIA) and secondary organic aerosol (SOA) formation. SIA formation builds mainly upon the widely distributed ISORROPIA mechanism (Nenes et al., 1998) and considers sulphate, nitrate, ammonium and interactions with sea salt. SOA can be formed from biogenic precursors (isoprene, terpenes, sesquiterpenes) and/or through oxidation of anthropogenic VOCs. As most regional modeling systems do, CMAQ typically underestimates PM concentrations, in particular SOA, because of unknown oxidation pathways or underestimated emissions (e.g. Solazzo et al., 2012).

The PM components of CMAQ, as well as the gases and radical species are re-calculated into the compounds included in TAPM. We expect that TAPM underestimates the secondary PM formation as discussed above; however, we don’t expect that this effect is large on the urban scale.
3. **Model evaluation** Figure 4. only shows the wind rose plots, and it is hard to tell whether the meteorological conditions perform well by the model. The authors mentioned that temperature, relative humidity, total solar radiation, wind speed, wind direction and precipitation show high correlation and low bias. How low is the bias? Is it within certain criteria, i.e., temperature bias within half to one degree Celsius?

Response:
We would like to thank the Reviewer for pointing out the need to carry out a better evaluation of modelled meteorological parameters. Thus, we added a section on comparing measured versus modeled meteorological parameters (as shown below) in the supplement. Additionally, we have added a reference in the manuscript section 3.1, which directs the reader to the supplement.

Moreover, we would like to point to a study by Tang et al. (2009), which performed an evaluation and comparison (with MM5) of meteorological parameters on the urban-scale in Gothenburg. The results of that study showed that “(1) TAPM performs better than MM5 in simulating near-surface air temperature and wind in urban area, (2) both models are able to reproduce nighttime vertical temperature gradient reasonably well, but underestimate daytime temperature gradient, and (3) the two models significantly underestimate the occurrences of low wind speed situation at night. These results indicate that the performance of TAPM in simulating meteorological features over the urban area is generally comparable to that of MM5. TAPM can be used with some confidence to describe the local-scale meteorology needed for air quality applications.” (Tang et al. 2009). Moreover, we applied urban-scale meteorology, simulated with TAPM, successfully in other harbor city studies (Ramacher et al. 2019, Ramacher et al. 2020). Evaluations in these studies also showed good performance of meteorological fields derived with TAPM. Table S1 has been added to the manuscript supplement.
4. The same applies to the evaluation of air quality variables. I feel it is very hard to read Fig. 5. The authors mainly show the annual mean comparison. How about daily scale? Any statistical metrics such as mean bias, mean normalized bias, etc. were calculated? I think it is useful to construct either a time series comparison or scatter plot to give readers an overall impression how the model performs in terms of the daily scale, or even hourly scale, if possible.

Response:
We would like to thank the Reviewer for pointing out the need to carry out a better evaluation of modelled concentrations. Thus, we added a section on comparing measured versus modeled daily concentrations of NO$_2$, O$_3$, PM$_{10}$ and PM$_{2.5}$ in the supplement together with description of the indicators presented in the table (Supplement section S1). This section contains a table of relevant statistical parameters (Table S2), as well as scatter plots of modeled versus measured daily concentrations for all stations and pollutants (Figure S1). Additionally, we added a reference in the manuscript section 3.1, which directs the reader to the supplement and enhanced the manuscript text by adding values for underestimations, which are the main drawback of the modeled results in terms of their use in health-effect calculations.
Nevertheless, we decided to keep the summary statistics as calculated with FAIRMODE DeltaTool in the manuscript, due to the focus and aim of the DeltaTool to evaluate air quality modeling results for policy applications.

S2 Statistical indicators and model performance indicators

In the statistical analysis of the model performance, the following statistical indicators are used: normalized mean bias (NMB), standard deviation (STD), root mean square error (RMSE), correlation coefficient (r), index of agreement (IOA) and the fraction of predictions within a factor of two of observations (FAC2). The overall bias captures the average deviations between the model and observed data and the NMB is given by:

$$\text{NMB} = \frac{\overline{M} - \overline{O}}{\overline{O}}$$

where $\overline{M}$ and $\overline{O}$ stand for the averaged model and observation results, respectively. The RMSE combines the magnitudes of the errors in predictions for various times into a single measure and is defined as

$$\text{RMSE} = \sqrt{\frac{1}{N} \sum_{i=1}^{N} (M_i - O_i)^2}$$

where subscript $i$ indicates the time step and $N$ the number of observations. RMSE is a measure of accuracy, to compare prediction errors of different models for a particular data and not between datasets, as it is scale-dependent. The correlation coefficient (Pearson $r$) for the temporal correlation is defined as:

$$r = \frac{\sum_{i=1}^{N} (O_i - \overline{O})(M_i - \overline{M})}{\sqrt{\sum_{i=1}^{N} (O_i - \overline{O})^2 \cdot \sum_{i=1}^{N} (M_i - \overline{M})^2}}$$

The index of agreement is defined as:

$$\text{IOA} = 1 - \frac{\sum_{i=1}^{N} (O_i - M_i)^2}{\sum_{i=1}^{N} (|M_i - \overline{M}| + |O_i - \overline{O}|)^2}$$

An IOA value close to 1 indicates agreement between modelled and observed data. The fraction of modelled values within a factor of two (FAC2) of the observed values are the fraction of model predictions that satisfy is defined as:

$$0.5 \leq \frac{M_i}{O_i} \leq 2.0$$

(9)

For evaluation of modelled values in rural areas, the acceptance criteria is FAC2 ≥ 0.5, while in urban areas it is FAC2 ≥ 0.3.
### Table S2: Evaluation of modeled versus measured daily concentrations of NO2, O3, PM10 and PM2.5

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5. How about the performance of the 4km * 4km CMAQ results? I believe it is interesting to do a comparison between the CMAQ results and the urban-scale model results, of course, together with the observations. Based upon this comparison, people can easily judge the usefulness of the ultra-fine scale city-level model. Currently, the city scale model has a higher spatial resolution of 250-m, however, if the model performs worse than CMAQ, what is the major purpose of the ultra-fine resolution? The same applies to the meteorology. I don’t feel the science was advanced by simply focusing on the city-scale model without detailed clarification of the advances of the model.

Response:

We would like to thank the Reviewer for pointing out the need to clarify the advantages of applying a city-scale model for the purpose of this study. As described in response to question 2, the main idea of our city-scale study is to utilize the boundary conditions of the CMAQ model simulations in a city-scale air quality model to assess the urban-scale features of the shipping emissions. In general, regional air quality models can give a reliable representation of concentrations in the urban background, but due to their limitation in resolving the near-field dispersion of emission sources and photochemistry at the sub-kilometre scale, around industrial stacks and on the neighbourhood level, they cannot provide the information needed by urban policymakers for population exposure mapping, city planning and the assessment of abatement measures. City-scale air quality models overcome the limitation inherent in regional-scale models by taking into account details of the urban topography, wind flow field characteristics, land use information and the geometry of local pollution sources. Thus, it is necessary to move beyond a resolution of e.g. 4km x 4km (resolution of CMAQ simulations used for the regional background). The city-scale air quality model TAPM was successfully applied to investigate urban air quality and scenarios in coastal urban areas all over the world (e.g. Matthias et al., 2018, Ramacher et al., 2020, Gallego et al., 2016, Fridell et al., 2014). Especially the meteorological module has proven to be capable of reproducing measured parameters such as temperature, wind speed and wind direction, because of its capability to capture meteorological effects sea-land circulations and complex terrain. This tackles also the minor comment on the use of COSMO-CLM instead of TAPM. COSMO-CLM does not take into account such effects on the urban-scale and thus, we decided to simulate and apply meteorological fields with TAPM. Previous studies (e.g. Tang et al. 2009 and Ramacher et
al. 2018) prove good meteorological simulation capabilities for the urban-scale, based on synoptic reanalysis, which we can confirm with the results of the presented study on Gothenburg (see also response to question 3).

When it comes to a possible comparison with other models on the regional scale, such as CMAQ, we would like to refer to a study by Karl et al. 2019, who compared the newly developed urban-scale CTM EPISODE-CityChem with TAPM and CMAQ. Karl et al. 2019 have carried out a full-year run with the TAPM air quality model to compare it with the urban-scale CTM EPISODE-CityChem. In this study, the TAPM run has been performed with the same horizontal resolution (1 km) as the EPISODE-CityChem run, identical emissions, but 2-D boundary concentrations instead of 3-D boundary conditions from CMAQ. CMAQ was not further included in the evaluation published in the manuscript because CMAQ cannot give realistic concentrations at the traffic sites and the industrial sites. CMAQ is a regional CTM system which does not handle local scale dispersion, i.e. a traffic site and a background site located within the same 4 x 4 km² grid cell would have the same concentration values. If the traffic stations and industrial stations were included, it would be obvious that CMAQ fails to reproduce concentrations at urban stations that are impacted by the local pollution. A realistic representation of local emissions is complicated by their high the spatial and temporal variability in the urban area. Urban-scale CTM such as EPISODE-CityChem and TAPM use the local scale emissions to compute the pollutant concentrations in the urban background areas, which are in turn affected by the highly resolved emissions. Therefore, urban scale models are much more sensitive to an incorrect representation of the local emissions than a regional scale model with coarser resolution.

Finally, we decided not to take into account a comparison of concentrations simulated with TAPM and concentrations simulated with CMAQ, also because the focus of this study is not the comparison of regional and city-scale CTM performances but much more the local effects, trends and challenges that arise for the urban population and policy.

6. The authors evaluated the species of PM$_{10}$, O$_3$ and NO$_2$, however, the health impact assessment is based on PM$_{2.5}$, O$_3$ and NO$_2$. Why not evaluating PM$_{2.5}$ directly? Line 493 mentioned that “In the chemistry mode of TAPM, simplified chemical reactions for the secondary PM are included and the secondary particulate matter consists of organic carbon, reactive nitrogen and sulfate.” I am also worried about the performance of PM$_{2.5}$ in the TAPM since only simple secondary inorganic and organic aerosol scheme was applied. How about the aerosol modes? Is the model using bulk mode aerosol or sectional bin model in TAPM?

Response:

Model evaluation for PM$_{2.5}$ has been added to the text and Figure 5 extended with panel for summary statistics for PM$_{2.5}$. Additionally, we added a section on detailed concentration evaluation in the supplement (see response to comment 4).

The secondary aerosol formation in TAPM is already discussed above at Discussion point 2. We have concluded that the regional-scale secondary aerosol formation is covered in the boundary concentrations calculated by CMAQ and that we don’t expect that effect of simplified secondary aerosol formation is large on the urban scale. Regarding the aerosol scheme, TAPM is using refined bulk mode, having separate scheme for Fine PM corresponding to PM$_{2.5}$ and Ambient PM corresponding to PM$_{10}$, which both include secondary PM. Two additional modes for particles in size fractions 10-20 and 20-30 µm not
involving secondary aerosol formation are included, these have not been used in our simulations.

(c) $PM_{2.5}$ At Femman and Haga

7. *Fig.9 the effect of local emission on ozone* The summer mean impact implies the NO titration effect. How about daily scale? The summer mean ozone is indeed quite low. Is there any day with slightly higher concentration, which may reveal different role of local shipping? It is not very persuasive by only using seasonal mean.

Response:

We would like to thank the reviewer for a very good suggestion to show the ozone formation due to the different sources on daily scale. We have added Fig. S5 in Supplement showing the modelled daily mean ozone concentrations contributed from local shipping, regional shipping and VOC emissions from local shipping at Eriksberg under summer and added the following text to the paper:

Further details of impact of the shipping emissions on ozone formation are illustrated in Figure S5 in the Supplement, showing summer ozone formation from regional and local shipping as well as from the local shipping VOC emissions at Eriksberg. At this location the local shipping emissions lead almost always to ozone depletion. On contrary, VOC emissions from local shipping cause the increase of ozone concentrations, confirming that the location is in a VOC-limited photochemical regime. The regional shipping tends to increase the local ozone concentrations in most of days (78 days under June–August). Inspecting details of the diurnal variation of ozone contributions (Figure S5b-d), one can see that during the rare occasions without ozone depletion by the local shipping, there is a small ozone formation from the local shipping emissions and no ozone formation from the local shipping VOC emission, indicating presence of NOx-limited regime (Fig. S5b), while during most of the studied days the local shipping emissions have an ozone depletion effect at daytime while the local shipping VOC emission have ozone formation effect peaking in the morning and sometimes also in the afternoon (Fig. S5c). The regional shipping increases the ozone concentrations in all three depicted cases, showing maxima in the afternoon.
(a) Diagram showing the concentration of O3 (ppb) over time from June to August 2012, with lines representing local shipping, regional shipping, and VOC local shipping.

(b) Diagram for 20120602, illustrating the concentration of O3 over time from 1 to 24 CET, again with lines for local shipping, regional shipping, and VOC local shipping.

(c) Diagram for 20120707, showing the concentration of O3 over time from 1 to 24 CET, with lines for local shipping, regional shipping, and VOC local shipping.
8. **the section of 3.2 Impact of ship emissions on local air quality** Most of this section simply describes the figure by using domain average, which does not make too much sense and not too much useful. Some comparisons might be made with either other sources or other studies to reveal the advancement of this study. For instance, what do the contributions from the local and regional shipping emission tell us? Is it useful in future strategies in the control policy? Only simple descriptions greatly discount the value of the study.

Response:

We would like to thank the Reviewer for pointing out the need of a deeper analysis to improve the value of the study. The air quality discussion arises from the annual mean for the later discussion on health impact. But we agree that more interesting comparisons will reveal the advancement of the study. The section 3.2 has been updated from several aspects, discussing local relative and absolute contributions, seasonal differences and especially exemplifying more details of the impacts of shipping and other sources for location Eriksberg, as described in more detail bellow. Regarding the control policies we would like to refer to the Conclusions part of the paper where we discuss potential impact of different policies for mitigation of air pollution from shipping on exposure to air pollutants in Gothenburg and on the health effects.

Section 3.2.1 SO₂
The modelled SO₂ concentrations in Gothenburg are relatively low and Fig. 6 shows highest concentrations around the city ports as well as around industrial areas north of Göta älv. The dominated south-westerly winds transport emissions from the shipping routes and port areas farther inlands to the north of Göta älv. Eriksberg, located on the north of Göta älv, is today a modern residential and commercial center built in place of an old dockyard area. We have selected this place to study relative impact of shipping in more detail. The shipping-related monthly contributions to SO₂ concentrations at Eriksberg were
47 % on average and over 60 % on June-August. Figure S3 in the supplement shows the modelled monthly mean relative contributions at Eriksberg.

Figure S3: Modelled monthly mean relative contributions from local shipping, regional shipping and all other emission sources (road traffic, industry etc.) to SO$_2$ concentrations at Eriksberg in year 2012.

Section 3.2.2 NO$_2$

Nearly 90 % of NO$_x$ emissions in Gothenburg are from road traffic (47 %) and local shipping (41 %). But local shipping impact is concentrated in areas inside the harbor along the Göta älv and decreases with growing distance to the port areas. Fig. 8 presents the impacts of local, regional shipping, as well as road traffic and other local anthropogenic sources on monthly level at Eriksberg, located on the north of Göta älv. The modelled annual mean NO$_2$ concentration from all sources is 7.5 ppb at Eriksberg, in which 2.5 ppb (33 %) from local shipping, 1.0 ppb (13 %) from regional shipping and 2.1 ppb (28 %) from road traffic. The maximum relative contributions from local shipping and regional shipping to monthly mean concentrations of NO$_2$ reach to 43 % in July and 16 % in June respectively. The monthly average contributions from local and regional shipping together are larger than or comparable to the contributions from road traffic in all months. Even though road traffic is the major contributor to the NO$_2$ concentrations in urban environment, the local ship emissions should not be neglected, especially in areas close to the city ports.
Section 3.2.3 O₃
Major update already shown in question 7.

Section 3.2.4 Particulate matter
At the near-harbour residential area Eriksberg, the modelled annual mean PM$_{2.5}$ concentration from all sources is 4.5 µg m$^{-3}$. The calculated annual mean contributions from local shipping and regional shipping are 0.2 µg m$^{-3}$ (~4%) and 0.4 µg m$^{-3}$ (~9%) respectively. The maximum monthly relative contribution from the local and regional shipping was about 29 % in July, in which 21 % from regional shipping (Fig. 11). Road traffic, the largest local source of PM$_{10}$, contributed up to 5 % of monthly PM$_{2.5}$ mean concentrations. The large contribution of PM$_{2.5}$ from regional shipping is agree with the character of source apportionment in Gothenburg. An early study shows that the main sources types of PM$_{2.5}$ in Gothenburg were long-range transport (LRT) (about 50 %), followed by ship emissions (20 %) and local combustion (19 %) (Molnár et al., 2017).
Figure 11: Modelled monthly mean contributions from local shipping, regional shipping and other sources (including contribution from the boundary condition) to PM$_{2.5}$ concentrations ($\mu$g m$^{-3}$) at Eriksberg for year 2012.

**Minor comments:**

1. **Line 48** Our study show “show” changed to “shows”
2. **Line 49:** emphasising changed to emphasizing
3. **Line 157:** exposureresponse Please add a space between exposure and response
   
   Response: Thank you, corrected accordingly.

4. **Line 233:** In this study, the meteorological component of TAPM was driven by the recently published ECMWF ERA5 synoptic Since the COSMO-CLM model has higher meteorological model and TAPM was driven by CMAQ 4km * 4km, why not using COSMO-CLM drives TAPM?
   
   Response:

   We would like to thank the reviewer for this comment, we agree that use of the same meteorological driver for both models would make sense. Reason for using the two different meteorological drivers for the TAPM and CMAQ modelling has a practical background – the work was following two different tasks which were initiated by two different modelling teams. The use of common meteorological driver was not agreed, mainly as TAPM produces its own fields and the 2 modelling studies were connected through the boundary concentration fields. The model comparison is discussed in more detail in Point 5 of the Major comment section.
5. Line 235: five nested domains What are the five domains? It is better to show a figure of the five nested domains. The authors also need to clarify the spatial resolutions of the five domains. Fig. 3b: the domain should be inferred in Fig. 3a, so the readers can tell where the domain of the finer resolution is.

Response: The Fig. 3a has been updated accordingly.

![Figure 3: (a) Five nested meteorological model domains with their sizes and spatial resolutions. The fifth domain with air pollution grid (250 x 250 m²) is pointed out in the figure, showing location of the three air quality monitoring sites Femman, Haga and Mölndal, as well as Eriksberg, a residential area close to the harbour;](image)

6. Line 307: “NOX” should be replaced by “NOx”.
7. Line 512: “µg m⁻³” should be replaced by “µg m⁻³”, and the same applies to Line 523 and 549.
8. Line 488: “A3 in Appendix”, but the Appendix only have S3, not A3. The same issue applies to “Fig. A4 in the Appendix” on line 503 and 511.

Response: Thank you, corrected accordingly.
9. **Fig 8, the x-axis label needs to be changed. For instance, either all using the mid-day of the month, i.e., 15/01, or something else to make it easy to follow.**

Response: Yes, using the mid-day of the month would be much easier to follow. However, the figure has been changed to monthly mean according to the suggestion of the first reviewer.

![Graph showing NO2 concentrations (ppb) by month](image)

*Figure 8: Modelled monthly mean contributions of the local shipping, regional shipping, local road traffic and other anthropogenic emissions (including contribution from the boundary conditions) to the NO2 concentrations (ppb) at Eriksberg in year 2012.*

10. **Figure captions can be more succinct. A lot of repetitive words.**

Response: Corrected accordingly.

**References**

Haeger-Eugensson, M., Moldanova, J., Ferm, M., Jerksjö, M., Fridell, E.: On the increasing levels of NO2 in some cities. The role of primary emissions and shipping. IVL report B1886, [www.ivl.se](http://www.ivl.se), 2010.


Ramacher, M.O.P.: Performance and evaluation of local scale wind flow fields for urban air pollution modeling with the coupled prognostic model TAPM driven by ERA5 climate reanalysis data, abstract EGU2018-8112 accepted for poster presentation at the European Geosciences Union General Assembly, Vienna, 2018.


The impact of ship emissions on air quality and human health in the Gothenburg area - Part I: 2012 emissions

Lin Tang\textsuperscript{1,2}, Martin O.P. Ramacher\textsuperscript{3}, Jana Moldanová\textsuperscript{1}, Volker Matthias\textsuperscript{3}, Matthias Karl\textsuperscript{3}, Lasse Johansson\textsuperscript{4}, Jukka-Pekka Jalkanen\textsuperscript{4}, Katarina Yaramenka\textsuperscript{1}, Armin Aulinger\textsuperscript{3}, Malin Gustafsson\textsuperscript{1}

\textsuperscript{1}IVL, Swedish Environmental Research Institute, P.O. Box 530 21, 40014 Gothenburg, Sweden
\textsuperscript{2}WSP Environment Sweden, Box 13033, 402 51 Gothenburg, Sweden
\textsuperscript{3}Chemistry Transport Modelling, Helmholtz-Zentrum Geesthacht, 21502, Geesthacht, Germany
\textsuperscript{4}Finnish Meteorological Institute, P.O. Box 503, 00101 Helsinki, Finland

Correspondence to: Jana Moldanová (jana.modanova@ivl.se) and Lin Tang (lin.tang@wsp.com)

Abstract. Ship emissions in and around ports are of interest for urban air quality management in many harbour cities. We investigated the impact of regional and local ship emissions on urban air quality for 2012-year conditions in the city of Gothenburg, Sweden, the largest cargo port in Scandinavia. In order to assess the effects of ship emissions, a coupled regional and local-scale model system has been set up, using ship emissions in the Baltic Sea and the North Sea, as well as in and around the port of Gothenburg. Ship emissions were calculated with the Ship Traffic Emission Assessment Model (STEAM) model, taking into account individual vessel characteristics and vessel activity data. The calculated contributions from local and regional shipping to local air pollution in Gothenburg were found substantial, especially in areas around the city ports. The relative contribution from local shipping to annual mean NO\textsubscript{2} concentrations was up to 3.3 ppb, together with 14 \% as the model-domain average, meanwhile the relative contribution from regional shipping at the North Sea and the Baltic Sea, the contribution was up to 4.3 ppb.\textsuperscript{26} In an area close to the city terminals, the contribution of NO\textsubscript{2} from local shipping (33 \%) was higher than that of the road traffic, (28 \%), which indicates importance of controlling the local shipping emissions. The local shipping emissions of NO\textsubscript{2} decreased to decreasing of the summer mean O\textsubscript{3} levels in the city by 0.5 ppb on annual mean(\textasciitilde2\%) in average. The regional shipping lead to a slight increase in the O\textsubscript{3} concentrations, however, the overall effect of the regional and the local shipping together was a small decrease of the summer mean O\textsubscript{3} concentrations in the city. In addition, VOC emissions from local shipping compensate up to 4 ppb decreasing of summer O\textsubscript{3} concentrations due to the NO titration effect. For PM\textsubscript{2.5}, the local ship emissions contributed with 0.1 µg m\textsuperscript{-3} only 3 \% to the annual mean concentrations on the city-in model domain average, while regional shipping was under 2012 conditions a larger contributor to the local PM\textsubscript{2.5} than the local shipping, with an annual mean contribution of 0.5 µg m\textsuperscript{-3} 11 \% on the city-domain average.

Based on the modelled local and regional shipping contributions, the health effects of PM\textsubscript{2.5}, NO\textsubscript{2} and ozone were assessed using the ALPHAA-RiskPoll (ARP) model. An effect of the shipping-associated PM\textsubscript{2.5} exposure in the modelled area was a mean loss of the life expectancy by 0.015 years per person. The relative contribution of the local shipping to the impact of total PM\textsubscript{2.5} was 2.2 \% which can be compared to 5.3 \% contribution from the local road traffic. The relative contribution of the regional shipping was 10.3 \%. The mortalities due to the exposure to NO\textsubscript{2} associated to shipping were calculated to be 2.6 premature deaths/year. The relative contribution of the local and the regional shipping to the total exposure to NO\textsubscript{2} in the reference simulation was 14 \% and 21 \%, respectively. The shipping related ozone exposures were due to the NO titration effect, leading to negative number of premature deaths. Our study shows that overall health impacts of regional shipping can be more important than those of local shipping, emphasizing that abatement policy options on city-scale air pollution require close cooperation across governance levels.
Our findings indicate that the strengthened Sulphur Emission Control Areas (SECA) fuel sulphur limit from 1 % to 0.1 % in 2015, leading to strong decrease in formation of secondary particulate matter on regional scale, has been an important step in improving of the air quality in the city.
1 Introduction

Shipping is an important source of air pollutants, both on the global and European level (Corbett et al., 1999; Eyring et al., 2005; Cofala et al., 2007). The most important species emitted are sulphur oxides (SOx), nitrogen oxides (NOx), particulate matter (PM) and to some extent carbon monoxide (CO) and volatile organic compounds (VOC). Since nearly 70 % of ship emissions occur within 400 km of coastlines (Corbett et al., 1999), the largest contributions of shipping to air pollution are concentrated to coastal regions with intensive ship traffic and to harbours, where emissions from harbour operations add further to the air pollution generated by ships. The primary air pollutants from shipping contribute to the formation of secondary air pollutants, mainly ozone and secondary particulate matter. On average shipping emissions contributed with 9.4 % to concentrations of primary PM2.5 (particulate matter with a median aerodynamic diameter less than or equal to 2.5 µm) and with 12.3 % to concentration of secondary inorganic particulate matter over the Europe during 1997–2003 (Andersson et al., 2009).

Emissions from the international shipping are controlled through International Maritime Organisation (IMO) and regulations included in the International Convention on the Prevention of Pollution from Ships (MARPOL 73/78) and its annexes. The MARPOL Annex VI- “Regulations for the Prevention of Air Pollution from Ships” sets limits for emissions of SOx and NOx. Sulphur is regulated through maximum allowed sulphur content in the fuel used, while NOx is regulated through Tier limits for maximum specific emissions of NOx from each engine on board. The limits depend on the nominal rotation speed of an engine and different Tiers apply for ships built or substantially re-built in different time periods (2000–2011 Tier 1, after 2011 Tier 2). For fuel sulphur content (FSC) a global limit of 0.5 % applies since 1st January 2020, before it was 3.5 %. However, the Baltic Sea, the North Sea and the English Channel are so called Sulphur Emission Control Areas (SECA) where stringier limits apply: in July 2010 it was decreased from 1.5 % to 1.0 %, which is also the limit that applies in this study. In 2015 the fuel sulphur limit was decreased further to 0.10 %. In addition, since 2010 a sulphur content limit of 0.10 % for fuels used by ships at berth for a period longer than 2 hours applied for all EU ports. Sweden has also introduced economic incentives for reduction of the shipping emissions in form of differentiated fairway and port fees with a discount for ships using emission control technologies, contributing to a relatively large share of ships with NOx abatement technology in the region. In 2020 the global cap for the FSC will be decreased to 0.50 %. In 2021 a Nitrogen Emission Control Area (NECA) will enter in force in this area with mandatory Tier 3 standard (80 % reduction comparing to Tier 1) for ships built in 2021 and later operating in the region.

In the Baltic Sea and the North Sea, an intensive ship traffic results in high emissions of air pollutants, and contributes to high atmospheric concentrations of particularly of NOx in and around several major ports (Jonson et al., 2015). The relative contribution of shipping in the North Sea and Baltic Sea to coastal NO2 concentrations are highest along the coasts of southern Sweden, the south-western coast of Finland, and the coast of Estonia, 25–40 % on annual average (Jonson et al., 2019). Jonson et al. (2019) found that the Baltic Sea and North Sea shipping contributed significantly also to concentrations of particulate matter (highest contributions 6–12 %, allocated to similar areas as NOx) and to the deposition of sulphur (highest contributions 10–20 %), before the strengthening of SECA fuel sulphur limit. They have also shown that the strengthened limit on the fuel S content in 2015 from 1.0 % to 0.10 % brought a significant decrease in emissions as well as contributions of shipping to air pollution by SO2 and to S deposition (maximum contribution about 2 %) and to a reduction of shipping contribution to the concentrations of PM. Aulinger et al. (2016) and Matthias et al. (2016) studied impacts of the current and future (2030) North Sea shipping on air pollution and found contributions consistent with Jonson et al. (2019) (highest NO2 contributions 25 % and 15 % in summer and winter, respectively, ozone increased by 10% along Scandinavian coast). By 2030, the contribution of shipping to the NO2 and O3 concentrations was estimated to increase by more than 20 % and 5 %, respectively, due to the expected enhanced traffic, if no regulation for further emission reductions is implemented in the North Sea area (Matthias et al., 2016).

Several studies have assessed impacts of shipping on human exposure to air pollutants and associated to health impacts. Andersson et al. (2009) assessed impacts of different source regions and also of emissions from international shipping on personal exposure and to the relative increase of death rates from exposure to
The city of Gothenburg is located on the western coast of Sweden, with about 0.57 million inhabitants and an area of 450 km². The dominant wind direction in Gothenburg is south-west with average wind speed of 3.5 m s⁻¹, indicating the major transport path from sea to the land, especially in summer. The geomorphology of the Gothenburg area is described as a fissure valley landscape dominated by a few large valleys in north-south and east-west directions. The major air pollution sources in Gothenburg are above all road traffic and industry, wood burning, shipping, agriculture, working machines and long-range transport (LRT) from the European continent and other parts of Sweden. The harbour and shipping activities are important emission sources and directly influences the urban air quality. The centre of the city is situated on the southern shore of the river Göta älvs.
600–700 ships pass to and from ports upstream and on the Göta älv. The port annually handles approximately 900,000 containers, 20 million tonnes of petroleum, and half a million Roll-on/Roll-off (RoRo) units (Winnes et al., 2015). Passenger traffic in Gothenburg is also very busy with 1.5 million passengers who ferry to and from Gothenburg to Denmark, Germany etc. on Stena Line ferries each year. This makes the port the largest cargo port in Scandinavia.

Comparing with other European cities, the air pollution levels in Gothenburg are low and the air quality has become better and better since the 70s because of the effective emission control addressing industry and road traffic. The trends of SO$_2$, NO$_x$, and NO$_2$ have been continuously decreasing from 1990 to 2015, except for the areas close to major roads (Miljöförvaltningen, 2017). O$_3$ exhibits an increasing trend and there is also a slowly increasing trend for PM$_{10}$ in Gothenburg (Olstrup et al., 2018). The annual means for NO$_2$, PM$_{10}$, and PM$_{2.5}$ (particulate matter with aerodynamic diameter less than or equal to 10 µm and 2.5 µm, respectively) during the period 2000–2017 are 12.5 ppb, 16.3 µg/m$^3$ and 7.9 µg/m$^3$, respectively, at an urban background site in Gothenburg. The decreased levels of NO$_x$ and NO$_2$ during the period 1990–2015 in Gothenburg were estimated to increase the life expectancy by up to 12 months and 6 months respectively, and the slight increased trend of O$_3$ and PM$_{10}$ have relatively little impact on life expectancy (~2 month and ~1 month) (Olstrup et al., 2018). In terms of exposure to PM$_{10}$ and PM$_{2.5}$ from different source categories in Gothenburg, Segersson et al. (2017) calculated that the largest part was due to the long-range transport and the dominating local sources were road traffic and residential wood combustion, while the contribution from local shipping was small, 0.04 µg/m$^3$ population weighted annual mean PM$_{2.5}$. The exposure of PM$_{2.5}$ from shipping in other harbour cities in Sweden are lower than in Gothenburg, with 0.02 µg/m$^3$ in Stockholm and 0.01 µg/m$^3$ in Umeå (Segersson et al., 2017).

This study has been conducted within the BONUS SHEBA project (Shipping and Environment of the Baltic Sea Region) where the impact of current and scenario emissions from ships on air quality have been investigated as a part of a holistic assessment framework for impacts of shipping on marine and coastal environment. The shipping-related air pollution has been investigated on a range of spatial scales with several chemistry-transport models: coarse spatial scale resolution was used for simulations in the European domain, finer resolution was used for the Baltic Sea (Karl et al., 2019a, c), and city-scale simulations using high spatial resolution were used for several harbour cities (Ramacher et al., 2019a). The present study (Part I) evaluates the contributions of regional and local shipping to the concentrations of SO$_2$, NO$_2$, PM$_{2.5}$, O$_3$ and secondary PM, as well as the human exposure and the associated health impacts in Gothenburg for year 2012. Health impact studies for the shipping emissions in cities are rare, mainly because the spatial resolution of the regional CTM (Chemical Transport Model) does not allow for the city-scale resolution. This study provides the city-scale Health Impact Assessment (HIA) and identifies and addresses potential health impacts associated to local and regional shipping. The studied year (2012) has been considered as a present-day “normal year” for Baltic Sea Region in terms of meteorological conditions in BONUS-SHEBA. In terms of ship emission regulations, the study presents a situation with 3.5 % FSC global limit, 1.0 % FSC limit in the SECA area whereas 0.1 % FSC limit applies for ships berthing in the port of Gothenburg or operating within the Göta älv estuary. Several alternative shipping scenarios in year 2040 are discussed further in Ramacher et al., 2019b (Part II).

2 Methodology
2.1 Model set-up
For the city-scale chemistry transport model (CTM), the prognostic meteorology-dispersion model TAPM (The Air Pollution Model) (Hurley et al., 2005; Hurley, 2008) was used. TAPM consists of a meteorological component and an air quality component (pollution components). The meteorological component of TAPM is an incompressible, non-hydrostatic, primitive equation model with a terrain-following vertical sigma coordinate for 3-D simulations. Using predicted meteorology and turbulence from the meteorological component, the air pollution component solves prognostic equations for concentrations and cross-correlation of concentrations by the Eulerian grid module. The model solves the momentum equations for horizontal wind components, the incompressible continuity equation for vertical velocity, and scalar equations for potential virtual temperature and specific humidity of water vapour, cloud water/ice, rain water and snow. The turbulence terms in these equations have been determined by solving equations for turbulence kinetic energy.
and eddy dissipation rate, and then using these values to represent vertical fluxes by a gradient diffusion
approach (Hurley, 2008b). Using predicted meteorology and turbulence from the meteorological component,
TAPM applies Eulerian grid module in its air pollution component which consists of nested grid-based
solutions of the Eulerian concentration mean equations representing advection, diffusion, chemical reactions
and emissions. Dry and wet deposition processes for gases and PM are also included. It includes gas-phase
photochemistry based on the Generic Reaction Set (Azzi et al., 1999), gas- and aqueous-phase chemical
reactions for SO$_2$, formation of ozone from NO$_x$ and NMVOC (treated as VOC reactivity) and simple
formation of secondary inorganic and organic aerosol. The model treats also dry and wet deposition processes
of gases and PM. The photochemistry mechanism captures also the important features of the secondary
particle formation, i.e. formation of sulphate and nitrate following the SO$_2$ and NO$_2$ oxidation, as well as
formation of secondary organic aerosol as a fixed part of the degraded smog reactivity representing VOC
species in the reaction scheme of TAPM (Hurley, 2008b).

In this study, the meteorological component of TAPM was driven by the recently published ECMWF ERA5
synoptic meteorological reanalysis ensemble means with 30 vertical layers, 0.3° × 0.3° horizontal and three-
hour temporal resolution. For a time period of 2012, five nested domains have been simulated with the
synoptic meteorological component with the inner-most meteorological fields of a 30 km × 30 km domain
with 500 m resolution. (Figure 3a). In addition, the observed wind fields at four meteorological sites were
assimilated to nudge wind speed and wind direction calculations in the inner-most domain. In TAPM, an
Exner pressure function is integrated from mean sea level to the model top (10 Pa in this study) to determine
the top boundary condition. The Exner pressure function is determined from the sum of the hydrostatic
component and non-hydrostatic component (Hurley, 2008). The number of vertical grid levels was 30 in this
study. Twenty of these layers are below approximately 2 km; the lowest layer extends to ca. 10 m above
ground.

The spatial resolution of the city-scale CTM was 250 m × 250 m with local coordinate system SWEREF 99
1200 and the size of the CTM domain was about 25 km × 25 km, covering the city of Gothenburg and the
harbour area along shores of the Göta älv running through the city (Fig. 31a).

The chemical boundary conditions were taken from the Community Multi-scale Air Quality Modelling
System (CMAQ) (Byun and Ching, 1999; Byun and Schere, 2006). CMAQ model simulations on a 4 km × 4
km grid (Karl et al., 2019c), which were used for the chemical boundary conditions, (Fig. 1b), were driven by
the high-resolution meteorology meteorological fields of the COSMO-CLM (Rockel et al., 2008) version 5.0
using the ERA-Interim re-analysis as forcing data. Chemical boundary conditions for the CMAQ model
simulations were provided through hemispheric CTM simulations, from a SILAM model (Sofiev et al., 2006)
run on a 0.5° × 0.5° grid resolution, which was provided by Finnish Meteorological Institute (FMI). Land
based emissions for the regional-scale simulations were represented by hourly gridded emissions calculated
with SMOKE-EU emission model (Bieser et al., 2011). The SMOKE-EU emission data is based on reported
annual total emissions from the European point source emission register (EPER), the official EMEP
(www.ceip.at) emission inventory and the EDGAR HTAP v2 database (EPER, 2018; CEIP, 2018; Olivier et
al., 1999). For shipping emissions, the model used an inventory calculated with the STEAM model consistent
with the inventory used by TAPM for the city-scale, calculated with 2 km × 2 km grid resolution (STEAM3,
Johansson et al., 2017), more details are given in the next section. The STEAM model version used for the
CMAQ simulations was, however, not including VOC emissions. As chemical boundary conditions, vertical
model layer seven with a mid-layer height of approximately 385 m above ground was selected. CMAQ
simulations with and without ship emissions in the Baltic Sea and the North Sea included were used in TAPM
simulation runs. Since the TAPM allows just 1-d boundary concentration fields with hourly time resolution,
the TAPM boundary concentrations were calculated using the horizontal wind components on each of the four
lateral boundaries for weighting the upwind boundary concentrations around the TAPM model domain
(Fridell et al., 2014). The city-scale model set-up are summarised in Table 1.

2.2 Emission inventory
2.1.1 Regional and local shipping emissions
Shipping emissions were calculated with the Ship Traffic Emission Assessment Model taking into account individual vessel characteristics and vessel activity data (STEAM; Jalkanen et al., 2012; Johansson et al., 2017) based on detailed information of technical parameters of individual vessels and position data of individual ships taken from reports from the Automatic Identification System (AIS) of the Helsinki Convention (HELCOM) member states. The STEAM model calculated fuel consumption and emissions as a function of vessel activity, the STEAM3 version of the model has been used (Johansson et al., 2017), with an additional module for calculation of VOC emissions. The emission inventory includes combustion emissions from all engines and appliances on ships (boilers, auxiliary and main engines). The emission inventory for the local shipping around Gothenburg consists of hourly emissions from ships on 250 m × 250 m grid resolution. The STEAM model provided shipping emissions for year 2012 for the compounds NOx, SOx, CO, CO2, VOC and PM2.5. PM2.5 is further divided into Elemental Carbon (EC), Organic Carbon (OC), SO2 and mineral ash. Ship emissions were provided in two vertical layers with emissions below and above 36 m height in order to differentiate between emissions from large ships with high stacks and the smaller vessels with lower stack heights (Fig. 42). The stack release heights were attributed to the corresponding mid-points of model layers in TAPM, 15 m for the emissions below 36 m height and layer 36 m for emission above 36 m.

2.2.2 Road traffic emissions

The road traffic emissions were calculated from traffic activity data and emission factors. The basic set of emission factors from road vehicles was extracted from HBEFA v. 3.2 (HandBook Emission FAvors for Road Transport, Rexeis et al., 2013). HBEFA comprehends emission factors for different classes of road vehicles based on type of vehicle (e.g. motorcycles, light-duty, heavy-duty vehicles), technology or fuel (e.g. petrol, diesel, hybrid) and emission standard (pre-Euro–Euro 6). For each of those also a number of road categories and driving patterns that affect emissions are specified within the vehicle sub-segments. These emission factors include also emissions of wear particles as well as evaporative VOC emissions. The emission factors for light-duty and heavy-duty vehicles and busses in Gothenburg were calculated using the Swedish national database on car fleet composition and national, vehicle-type specific, activity data in 2012. Road traffic emissions were finally calculated using traffic activity data for Gothenburg (vehicle kilometres for light duty vehicles plus motorcycles, heavy duty vehicles and busses on road links with specified type, speed and congestion hours) from the database of the Environmental Administration, City of Gothenburg (Miljöförvaltningen), and corresponding emission factors calculated in the HBEFA database. These data were applied as line emission sources in the model. Resuspension of the road dust is not covered in the model.

2.2.3 Other emissions

Ten large point sources from industrial processes are present in the city-scale model domain, these are all fugitive emissions from fuel handling and refineries. For technical reasons these were considered as area sources in the model with release heights corresponding to the stack heights allocated to these sources. The emission factors from these industrial sources were obtained from Swedish Environmental Emission Data (SMED 2015) for 2012.

Emissions from the following sectors were geographically distributed on 1 km × 1 km grid and assigned with emission height: ‘Manufacture of solid fuels and other energy industries’, ‘Combustion in industry for energy purposes’, ‘Stationary combustion in agriculture/forestry/fisheries’, ‘Energy and heat production (commercial/institutional)’, ‘Residential plants (boilers), domestic heating, working and off-road machinery’, ‘Use of paints and chemical products in households and enterprises’, ‘Agriculture, waste and sewage’, as well as ‘Other transports’ (the landing and take-off emissions from aviation, trains and military). They stem also from the SMED database and were obtained gridded on the 1 km × 1 km grid from SMHI (Swedish Meteorological and Hydrological Institute). These emissions were applied as gridded sources in the model.
The local emissions of NO\textsubscript{x}, SO\textsubscript{2}, PM\textsubscript{10} and VOC from the above-mentioned sectors in the model domain are shown in Fig. 2. The local shipping was the dominant emission source of SO\textsubscript{2} contributing with 61\% to the total SO\textsubscript{2} emissions (502 ton/year) in the model domain. Further, local shipping contributed with 41\% of the NO\textsubscript{x} emissions in the model domain, which was comparable to contribution from the road traffic (47\%), to total NO\textsubscript{x} emissions of 5072 ton/year. However, the road traffic was the major contributor for PM\textsubscript{10} in the model domain (45\% of 357 ton/year), while the local shipping and industry contributed with approximately 25\% each. For VOC (7457 ton/year), about 92\% of emissions were released from the industrial sector and only 0.3\% from the local shipping.

2.3 Design of model simulations
Several model simulations were performed to investigate the influence of shipping within the city domain and influence of the regional shipping outside the city on air pollution in 2012:

(1) A simulation including complete emission inventory both in the city-scale simulation and in the CMAQ simulation supplying the chemical boundary conditions: “Base scenario”;

(2) A simulation excluding the local shipping in the TAPM city domain but including regional shipping chemical boundary conditions in the CMAQ simulation: “No local shipping scenario” and;

(3) A simulation excluding both local shipping in the TAPM city domain and shipping in the chemical boundary conditions in the CMAQ simulation: “No local and regional shipping scenario”.

In addition, three sensitivity studies were performed within this study:

(1). “No NMVOC from local shipping” had the same emission input as the “Base scenario”, but without NMVOC from local shipping emissions. The difference between “Base” and “No NMVOC from local shipping” was used to investigate the impact of VOC emissions from local shipping, which is often neglected in the emission inventories due to its small proportion and as these has only recently been included in the STEAM model;

(2). “Primary PM from local shipping” had the same emission input as the “Base scenario”, but for the local shipping only the primary PM emissions as calculated by the STEAM model were included, all emissions of the gaseous species were excluded preventing formation of the secondary PM from local shipping. The difference between “Base scenario” and “Primary PM from local shipping” reflects the formation of secondary PM from SO\textsubscript{2}, NO\textsubscript{x} and VOC emitted by the local shipping;

(3). “No road traffic” had the same emissions as the “Base scenario”, but without road traffic emissions. It was used to compare the contributions of shipping emissions as well as the health impact of shipping with emissions from the city traffic.

2.4 Model evaluation
Model evaluations were carried out for both meteorological and air pollution parameters. The simulated meteorological parameters (temperature, relative humidity, wind fields and precipitation) were evaluated with measurements at four stations: Femman (57.70° N, 11.97° E, 30 m a.s.l.), Göteborg A (57.72° N, 11.99° E, 3 m a.s.l.), Vinga A (57.63° N, 11.60° E, 18 m a.s.l.) and Landvetter (57.68° N, 12.29° E, 154 m a.s.l.). The urban background site Femman is located on a rooftop in the city center and the local meteorological variables as well as the air quality data are continuously measured by the Environmental Administration in Gothenburg (Miljöförvaltningen), while the other three meteorological stations are driven by SMHI. For the air pollution evaluation, Femman, Haga and Mölndal were included. Haga is located in a one-sided street canyon in central Gothenburg with a park to the east of the station. Mölndal is located in southern part of Gothenburg on a rooftop about 30 m above ground and corresponds to a traffic station. NO\textsubscript{2} is measured by a reference chemiluminescent method at Femman and Haga and by Differential Optical Absorption Spectrometry (DOAS) method at Mölndal. PM\textsubscript{10} mass concentrations are measured by TEO (Tapered Element Oscillating Microbalance, model 1400ab) instrument at all three stations. Ozone instrument at Femman was Teledyne (model T400) and at Mölndal a DOAS from OPVIS. Hourly averaged air quality data for NO\textsubscript{2}, PM\textsubscript{10}, PM\textsubscript{2.5} and O\textsubscript{3} at the three air quality stations were used to evaluate the model performance.
The FAIRMODE DELTA Tool version 5.4 was used for the evaluation of the model results for the city of Gothenburg. DELTA Tool is an IDL (Interface Definition Language)-based statistical evaluation software which allows to perform diagnostics of air quality and meteorological model performance (Thunis et al., 2012, Pernigotti et al., 2014). The tool focuses on the air pollutants regulated in the Air Quality Directive 2008 (AQD 2008) and calculates statistical performance indicators such as Mean, Exceed, Normalized Mean Bias (NMB), Normalized Mean Standard Deviation (NMSD) and High Percentile (Hperc+) (see section S1 in the Supplement). Moreover, a performance criteria can be calculated, which is combining the statistical performance indicators with fixed parameters to evaluate whether the model results have reached a sufficient level of quality for a given policy support application (Pernigotti et al. 2014). According to the DELTA tool, the capability of a model to reproduce measured concentrations is good when more than 90% of the stations fulfill the performance criteria. We applied the Delta Tool to concentrations of NO2, PM2.5, PM10 and O3 measured at the available urban background sites and road traffic sites, compared them with concentrations calculated by our model system and calculated both, statistical performance indicators and the model performance criteria.

2.5 Health impact assessment

The health impacts of exposure of population in Gothenburg to shipping-related air pollutants were assessed with the ALPHA-RiskPoll (ARP) methodology (Holland et al., 2013) providing for calculation of a wide range of air-pollutant specific health effects based on the population weighted concentrations, national population statistics on age distribution of population, mortality and morbidity data and effect-specific exposure-response relationships. The methodology has been developed and used for quantification and assessments of the benefits of air pollution controls in Europe for the UN/ECE Convention on Long Range Transport of Air Pollution and is based on work for the Clean Air For Europe (CAFÉ) Program and on EU project Modelling of Air Pollution and Climate Strategies (EC4MACS). Following the WHO recommendations (WHO 2013a) and the Clean Air for Europe (CAFÉ) cost-benefit analysis methodology for assessment of health impacts of air pollutants, impacts of exposure to PM2.5, ozone and NO2 have been considered in the analysis. The exposure to these three pollutants is considered most harmful by the World Health Organization (WHO, 2013b). In this study only the most serious impacts, i.e. losses of lives, are presented, taking into account impacts of long-term exposure to PM2.5 and short-term exposures to ozone and NO2, i.e. the impacts marked A* in the HARPIE study (WHO 2013a). For ozone the indicator SOMO35 is used, representing the 8-hourly mean ozone concentrations accumulated dose above a threshold of 35 ppb during a year. The health impacts of some pollutants are correlated and that is why the premature deaths attributed to each pollutant cannot simply be added up. In particular, it has been estimated that adding premature deaths attributed to PM2.5 to those attributed to NO2 could result in double counting of around 30% (WHO 2013a). The health impacts calculated with the ARP model are presented as premature deaths and YOLLS per year, using the ER function of the model, i.e. 1.0062 (95% CI 1.004–1.008) per µg/m³ (WHO, 2013a).

The concentrations fields of PM2.5, O3 and NO2 were calculated by the coupled high resolution (250 m x 250 m) modeling system, as described above. Annual means and SOMO35 were calculated from hourly concentrations for each grid. Population data on 1 km x 1 km resolution were obtained from Statistics Sweden (SCB) for 2015 with a population of 572 779 in the city of Gothenburg. As there are no significant changes in population density between 2012 and 2015, the population data for 2015 were used. Population-weighted average concentrations (PWC) for the model domain were calculated multiplying the modelled annual mean concentration of the pollutant on each grid-cell by the population in the same grid-cell as weight for the modelled concentration.

To calculate the health risks, further information needed is the ERF and the baseline health statistics including the life expectancies, the death rates and morbidity data for estimating the impacts on mortality and morbidity. To estimate YOLLS, the age at which the premature deaths occurred should also be considered. In the ARP model, the ERFs used are those from WHO (2013a): 6.2 % (95% confidence interval 4.0–8.3%) relative risk increase per 10 µg/m³ increased exposure for the PM2.5 exposure, 0.29 % (95% confidence interval 0.14–0.43%) relative risk increase per 10 µg/m³ increased exposure for the ozone exposure and 0.27% (95% confidence interval 0.16–0.38%) relative risk increase per 10 µg/m³ increased exposure for the NO2.
exposure. ARP uses linear ERFs, recognizing the limited range of pollutant exposures in Europe. The YOLLs are calculated per year, applying the relative risk within national life tables. This is done through the relation between the life years lost per 100 000 population per unit PM2.5 concentration and the life expectancy of the population developed by Miller et al. (2003) based on analysis of the life tables. The premature deaths are calculated using the ERF for all-cause mortality and the total national mortality rates. This methodology is justified for European countries with health status and proportion of natural mortality of population corresponding to population studied in the epidemiological studies which brought forward these ERFs. For regions with high concentration levels of PM2.5 the HIA studies need to use different form of ERFs and for populations with different health status and proportion of natural mortality comparing to the US and Western Europe, cause-specific rather than all-cause mortalities need to be used. In this study the analysis was made separately for the population exposure related to the different pollutants from local and regional shipping.

3 Results and discussion

3.1 Model evaluation

The model evaluation was conducted for both meteorology and air pollution in the inner-most model domain. The comparison between hourly measured and modelled local meteorological parameters (temperature, relative humidity, total solar radiation, wind speed, wind direction and precipitation) shows high correlation and low bias, averaged over all available stations, temperature and wind speed are slightly underestimated with -0.46°C and -0.18 m/s, respectively. A detailed analysis can be found in Table S1 in the Supplement. The application of ERA5 datasets in the model shows significant improvements from the default reanalysis datasets. Nevertheless, the predictions of the meteorological parameters such as wind fields flow get better with wind field assimilation, for more detail see Ramacher (2018). For example, the differences between observed and simulated wind rose at Femman in January and July indicate a good model capability to reproduce local wind field except for missing about 30% of low wind speeds (0–2.5 m s⁻¹) from the north (Fig. 4), which may introduce some underestimations in high pollutant concentrations at ground due to accumulation in the boundary layer. Nevertheless, the total frequency of northerly winds at Femman station is low in January (8–17%) and very low in July (1–8%).

The evaluation of ambient pollutants was conducted through the major statistical parameters, (Table S2 in the Supplement). At urban background site Femman, the estimation of NO₂, PM₁₀ and PM₁₀,PM₂.₅ concentrations were satisfactory in summer with lower bias, (-0.16 µg/m³), however the model tended to underestimate NO₂ and, PM₁₀ and PM₂.₅ concentrations in winter, (-15.35 µg/m³). Ozone evaluation was carried out at station Femman and Möldal, and underestimation of daily maximum of the 8-hour means was also detected, which could be caused by low resolution of local NO sources and hence more smoothed titration of ozone. The summary statistics according to the FAIRMODE model evaluation tool shows that less than 90% of daily PM₁₀ concentrations at road site Haga fulfill the performance criteria for the statistic indicator Hperc (Fig. 5). The indicator Hperc indicates the model capability to reproduce extreme events, represented by selected high percentile for modelled and observed values. A detailed evaluation of simulated concentrations in form of scatter plots of modeled versus measured daily concentrations can be found in Fig. S1 in the Supplement.

The underestimation of NO₂ and PM₁₀ especially at road sites demonstrate impact of too coarse spatial resolution (250 m × 250 m) not capturing high concentrations at street level, possible missing or insufficient cover of local emissions like resuspension particulate matters from traffic sources, or incomplete chemical reactions in the model etc. As pointed out by Karl (2019b), recent nested model approaches have not resolved the details in the emission processing and near-field dispersion at the street and neighborhood level. However, shipping emissions are, when reaching the exposed population, more dispersed and the 250 × 250 m² grid resolution should be sufficient to assess their impact. Nevertheless, the other statistic indicators (Mean, exceedances, Normalized Mean Bias, Normalized Mean Standard Deviation, Correlation coefficient, etc.) of model performance in Fig. 5 show a satisfactory performance of the used city-scale model for Gothenburg.

3.2 Impact of ship emissions on local air quality

3.2.1 SO₂

The modelled annual average SO₂ concentrations from all sources, including local and regional shipping are shown in Fig. 6. The study was performed for year 2012 conditions, when the sulfur content in marine fuels was limited to 1 % in the region and 0.1 % for ships at berth. With these fuel Sulphur limits the local shipping
is still the dominant local emission source of SO\(_2\) (60 %) and influences the area around main shipping routes and city ports (Fig. 6). The calculated annual mean concentration of SO\(_2\) from all sources in the model domain is 0.4 ppb and the local shipping contributes with 0.05 ppb (13 %) on the model-domain average and between 0.3 ppb up to 0.6 ppb in a wide area around the main shipping routes and ports. An additional increase of 0.1 ppb SO\(_2\) is detected when considering the regional ship emissions. The impacts were higher in summer months (JJA), local and regional shipping contribute on model domain average with 0.1 ppb to SO\(_2\) concentrations and with 0.8 ppb in maximum than in winter (Fig. S1S2 in the Supplement), being result of higher shipping emission in summer as well as of differences in meteorological situation. The highest SO\(_2\) contributions (maximum 0.67 ppb on annual mean and 0.8 ppb in summer) were found around the major ports: Älvsborgshamnen, Skandiahamnen, Skarvikshamnen, Ryahamnen, Lindholmshamnen, and Frihamnen in the northern bank of the Göta älv (Fig. 6d). In addition, two busy ferry terminals located on the southern bank of the Göta älv can contribute to the high SO\(_2\) concentrations on the opposite river side due to the dominant south-westerly winds. The regional ship emissions outside the model domain contribute with additional 0.06 ppb (15 %) on the model-domain average. In difference from the local-shipping, this contribution is distributed rather evenly over the model domain.

The modelled SO\(_2\) concentrations in Gothenburg are relatively low and Fig. 6 shows highest concentrations around the city ports as well as around industrial areas north of Göta älv. The dominated south-westerly winds transport emissions from the shipping routes and port areas farther inlands to the north. Eriksberg, located on the north waterfront of Göta älv, is today a modern residential and commercial center built in place of an old dockyard area. We have selected this place to study relative impact of shipping in more detail. The shipping-related monthly contributions to SO\(_2\) concentrations at Eriksberg were 47 % on average and over 60% on June-August. Figure S3 in the supplement shows the modelled monthly mean relative contributions at Eriksberg.

3.2.2 NO\(_2\)

NO\(_2\) is mainly emitted as nitrogen oxide (NO), in the STEAM model the NO\(_2\)/NO\(_x\) ratio is 5%. In atmosphere NO is quickly converted to NO\(_2\) in reaction with ozone, so further from the source the atmospheric NO\(_2\) is dominated by NO\(_2\), approaching a photo-stationary state driven by the NO+O\(_3\) reaction and NO\(_2\) photolysis. Maps of modelled annual mean atmospheric concentrations of NO\(_2\) over the Gothenburg area are shown in Fig. 7. The annual mean concentration of NO\(_2\) in the Base simulation is 3.7 ppb as the model-domain average (Fig. 7a), and the model-domain mean contribution from local shipping to the annual mean concentrations is 0.5 ppb (14 %) and up to 3.3 ppb in areas with high contribution (Fig. 7b, 7c). The relative contribution of local shipping to the NO\(_2\) concentrations in the model domain in Gothenburg is comparable with 11 % in Riga (Latvia), 16 % in Gdańsk-Gdynia (Poland) and 22 % in Rostock (Germany) in 2012 (Ramacher et al. 2019). The calculated model-domain mean contribution to NO\(_2\) concentrations from local and regional shipping together is 1.04 ppb (44 %+26 %), and up to 4.12 ppb in most heavily impacted areas (Fig. 7c). The seasonal differences in NO\(_2\) concentrations are driven by emissions, atmospheric chemistry and atmospheric mixing. Maps of modelled air concentrations of NO\(_2\) over the Gothenburg area in winter and summer month are shown in Fig. S2 in the supplement. The higher contribution of NO\(_2\) (Fig. 7c), which is larger than local shipping in contributions. The total shipping-related relative contribution in Gothenburg to the NO\(_2\) annual averaged grid mean in the model domain is 40 %. In summer the contribution reaches 49 % (17 % from local shipping + 32 % from regional shipping) to the sum and lower average NO\(_2\) concentration in the model domain and influenced areas is due to expands to further inland. This is result of 20 % higher summer emissions comparing to winter, different photochemical state as well as different local meteorological conditions. The dominated south-westerly winds in summer transport NO\(_2\) from the shipping routes and port areas farther inlands. Again, the highest level of NO\(_2\) is around the port Skandiahamnen (Fig. S4 in the Supplement).

Eriksberg is a modern residential Nearly 90 % of NO\(_2\) emissions in Gothenburg are from road traffic (47 %) and commercial center built in place of an old dockyard area. The daily influence from local and regional shipping is obvious and illustrated by (11 %). Impact of the calculated daily mean NO\(_2\) concentrations from the three model simulations “Base”, “No local shipping is concentrated in areas inside the harbor along the Göta älv and decreases with growing distance to the port areas. Fig.” and “No 8 presents impacts of the local and regional shipping” (Fig. 8). The local shipping contributes with, as well as of the road traffic and all other anthropogenic sources (including NO\(_2\) coming from the model domain boundary) on monthly levels at
Eriksberg. The modelled annual mean NO\textsubscript{2} concentration from all sources is 7.5 ppb at Eriksberg, in which 2.5 ppb (33\%) originates from local shipping, 1.0 ppb (13\%) from regional shipping and 2.1 ppb (28\%) from road traffic. The maximum relative contributions from local shipping and regional shipping to the monthly mean concentrations of NO\textsubscript{2} on reach 43\% in July and 16\% in June. The monthly average, and up to 11.1 ppb in March. Meanwhile the regional shipping contributes with 1.0 ppb to daily mean NO\textsubscript{2} concentrations on average and reaches 9.2 ppb at most. For comparison, the daily average contributions of NO\textsubscript{2} from road traffic are also presented in Fig. 8. In total, 210 days in 2012 show contributions to daily mean concentrations from the local and regional shipping higher together are larger than those comparable to the contributions from the road traffic at Eriksberg in all months. Even though the road traffic is the major contributor to the NO\textsubscript{2} concentrations in urban environment, the local ship emissions should not be neglected are of major concern, especially in areas close to the city ports.

3.2.3 \textit{O}_3

\textit{O}_3 is formed in photocatalytic cycles involving NO\textsubscript{x}, ozone and hydrocarbons, through the photolysis of NO\textsubscript{2} in sunlight. The same cycle involves also titration of ozone by the reaction with NO forming the NO\textsubscript{2}. Maps of modelled atmospheric concentrations of ozone over the Gothenburg area in 2012 are shown in Fig. 9, with focus on summer months (JJA). The regional background concentration of ozone at a regional background station, station Råö close to Gothenburg area is 37 ppb in the summer of 2012. Modelled summer ozone levels in the model domain are in the 15–30 ppb range (domain average is 28.6 ppb) (Fig. 9a). Since NO\textsubscript{2} is mainly emitted as NO, the emissions from local shipping cause local reduction of ozone concentrations due to the titration of O\textsubscript{3} by NO\textsubscript{2} (Fig. 9b). The O\textsubscript{3} depletion pattern along the north riverbank of the Göta älv is 4 ppb (~14\%) in maximum due to the local shipping with both NO\textsubscript{2} and NMVOC emissions (Fig. 9b), while regional shipping emissions tend to increase the ozone concentrations by 1 ppb over the land. This ozone increase can be compared to 4–6 ppb increase caused by the shipping emissions over the remote ocean due to a result of large-scale summer ozone production in summer (found by Huszar et al. (2010, Fig. 9c).

In the local STEAM inventory, the non-methane volatile organic compounds (NMVOC) from shipping are also available were introduced. These NMVOC serve as precursors of O\textsubscript{3} and enhance photochemical ozone production. TAPM model uses concept of VOC reactivity instead of individual NMVOCs, producing pool of peroxy radicals which take part in the ozone-production photocatalytic cycle. A sensitivity run was performed to study the impact of NMVOC\textsubscript{VOC} emissions from the local shipping on ozone concentrations in the city by excluding the local shipping NMVOC\textsubscript{VOC} emissions from the simulation. Fig. 9d shows the impact of the NMVOC\textsubscript{VOC} emissions: the O\textsubscript{3} concentrations increased by up to 2 ppb (~7 \%) along the main shipping routes and the port areas, which means that the negative titration effects of NO\textsubscript{2} emissions from local shipping on the ozone concentrations was 6 ppb in maximum when NMVOC\textsubscript{VOC} emissions were excluded, comparing to 4 ppb in the Base simulation. Sensitivity of ozone formation to VOC emissions also clearly indicates that the city centre is most often in VOC-limited regime. Further details of impact of the shipping emissions on ozone formation are illustrated in Figure S5 in the Supplement, showing summer ozone formation from regional and local shipping as well as from the local shipping VOC emissions at Eriksberg. At this location the local shipping emissions lead almost always to ozone depletion. On contrary, VOC emissions from local shipping cause the increase of ozone concentrations, confirming that the location is in a VOC-limited photochemical regime. The regional shipping tends to increase the local ozone concentrations in most of days (78 days under June–August). Inspecting details of the diurnal variation of ozone contributions (Figure S5b-d), one can see that during the rare occasions without ozone depletion by the local shipping, there is a small ozone formation from the local shipping emissions and no ozone formation from the local shipping VOC emission, indicating presence of NO\textsubscript{x}-limited regime (Fig. S5b), while during most of the studied days the local shipping emissions have an ozone depletion effect at daytime while the local shipping VOC emission have ozone formation effect peaking in the morning and sometimes also in the afternoon (Fig. S5c). The regional shipping increases the ozone concentrations in all three depicted cases, showing maxima in the afternoon.

3.2.4 Particulate matter

Particulate matter includes primary, directly emitted particles and secondary particulate matter formed upon further processing of emissions in the atmosphere. At the urban background site Femman, close to the city harbour, the measured annual mean PM\textsubscript{2.5} concentration was 7.9 μg m\textsuperscript{-3} in 2012. The calculated annual mean
PM$_{2.5}$ in the Base simulation was 4 µg m$^{-3}$ as the model domain average (Fig. 10a). The local ship emissions contributed with 0.1 µg m$^{-3}$ (3 %) to the annual mean as the model-domain average (Fig. 10b,10b), which had same relative contribution as 3% in Gdansk-Gdynia and higher than 1% in Rostock and Riga in 2012 (Ramacher et al., 2019). Regional shipping was under 2012 conditions a larger contributor to the local PM$_{2.5}$ than the local shipping with annual mean average contribution of 0.4 µg m$^{-3}$ (11 %) (Fig. 10c). In summer, the area with major influence of shipping emissions extended to the north of the Göta älv with maximum contributions from local plus regional shipping of 1.1 µg m$^{-3}$ at Skandahamnen (A3 in Appendix). At the near-harbour residential area Eriksberg, the contribution from the local and regional shipping was in range 0.2–1.1 µg m$^{-3}$ on monthly mean, representing about 5–29 % contribution to the calculated monthly mean PM$_{2.5}$ concentrations and contribution from the regional shipping dominated in the months from winter to summer (Fig. 11). The total ship-related relative contribution to the annual averaged PM$_{2.5}$ concentrations in model domain was 14 %, and reached in summer to 27 % (4 % from local shipping + 23 % from regional shipping) to the summer averaged PM$_{2.5}$ concentrations in model domain (Fig. S6 in Supplement). At the near-harbour residential area Eriksberg the modelled annual mean PM$_{2.5}$ concentration from all sources is 4.5 µg m$^{-3}$. The calculated annual mean contributions from local shipping and regional shipping are 0.2 µg m$^{-3}$ (~4 %) and 0.4 µg m$^{-3}$ (~9 %) respectively. The maximum monthly relative contribution from the local and regional shipping was about 29 % in July, in which 21% from regional shipping (Fig. 11). Road traffic, the largest local source of PM$_{10}$, contributed up to 5% of monthly PM$_{2.5}$ mean concentrations. The large contribution of PM$_{2.5}$ from regional shipping is agree with the character of source apportionment in Gothenburg. An early study shows that the main sources types of PM$_{2.5}$ in Gothenburg were long-range transport (LRT) (about 50 %), followed by ship emissions (20 %) and local combustion (19 %) for year 2008–2009 (Molnár et al., 2017).

In the chemistry mode of TAPM, simplified chemical reactions for the secondary PM are included and the secondary particulate matter consists of organic carbon, reactive nitrogen and sulfate. While in summer more intensive photochemistry favors formation of precursors to the secondary PM, the air temperature and humidity-controlled gas/particle partitioning of ammonium nitrate causes higher PM nitrate concentrations during periods of the year with cold and wet weather. Many city-scale models do not involve chemistry and thus neglect formation of the secondary PM. The secondary aerosol formation in TAPM is heavily parameterized, however, captures the important features of the secondary particle formation, i.e. formation of sulphate and nitrate following the SO$_2$ and NO$_2$ oxidation, as well as formation of SOA as a fixed part of the degraded smog reactivity representing VOC species in the reaction scheme of TAPM (Hurley et al. 2008b).

On urban scale, formation of secondary PM is usually supressed as the radical pool is depleted by the primary emissions and many urban models do not consider the secondary PM at all. Therefore, a sensitivity run was performed to investigate the role of the formation of secondary PM from local shipping on the city scale where only emissions of the primary PM were introduced, without emissions of the gas-phase pollutants from the local shipping. Modelled secondary PM concentrations from shipping were calculated as the difference between the base run and this sensitivity run. They were found relatively low, with maximum 0.0009 µg m$^{-3}$.

Fig. S7 in Supplement shows contributions of max. 2 % of the PM related to the local shipping in Gothenburg in winter, months and spread out since negligible contributions in summer. The secondary PM is mainly formed far from the sources. The secondary PM tends to disperse and accumulate to the east part of Gothenburg due to the prevailing wind directions (Fig. A4 in the Appendix).

4 Calculation of exposure and health effects from ship emissions

The contribution of emission sources to population exposure depends on the relationship between population density and air pollution levels. The areas with relatively high exposure to PM$_{2.5}$ due to local and regional shipping are city ports and areas around, especially north of the Göta älv. Figure 12 presents the population weighted annual mean concentrations of NO$_2$, PM$_{2.5}$ and SOMO35 at each model grid for the base simulation and for contributions of the local plus regional shipping, as well as for contributions of the road traffic. The spatial patterns of PM$_{2.5}$ exposure from shipping are dominated by gradients in the concentration fields around the city ports to the north of Göta älv. PM$_{2.5}$ exposure from shipping is higher than exposure from road traffic in a larger city area since regional shipping-related PM$_{2.5}$ exposure is evenly distributed over the city (Fig. A58 in Appendix the Supplement). The sum of PWC of PM$_{2.5}$ from the local plus regional shipping is 0.51 µg m$^{-3}$ in the model domain, to which the regional shipping contributes with 82 %, comparing to 0.22 µg m$^{-3}$ associated to road traffic (Table 2). The total exposure to PM$_{2.5}$ is dominated by particles transported to the city with the background air. The sum of PWC of NO$_2$ from regional and local shipping was 1.65 ppb, similar to that from the road traffic (1.75 ppb), with gradients in the concentration fields north of the Göta älv.
Because of the effect of local O₃ titration by the shipping emitted NO, the exposure to SOMO35 from shipping was negative along the Göta älv. However, SOMO35 exposure due to regional shipping was positive with sum of PWC 70.9 ppb x h in the model domain and showed relatively high level in areas with high population density.

The PWC for these pollutants were then used in the health impact calculations and results are presented as life years lost per year and loss of life expectancy (years of lifetime lost per person, YOLLs pers.⁻¹) for PM₂.₅ and as premature deaths for ozone and NO₂. The estimated loss of life expectancy (YOLLs pers.⁻¹) due to PM₂.₅ from local shipping was 0.003 while from the regional shipping it was 0.014. For comparison, impact of exposure to PM₂.₅ from the road traffic was calculated to be 0.007 YOLLs pers.⁻¹ and to all PM₂.₅ in the Base simulation to be 0.14 YOLLs pers.⁻¹ (Table 3). In all, shipping contributed with 12% to the calculated health impacts from the total exposure to PM₂.₅ in the city and the impact was more than 2 times larger than that of the local road traffic, the regional shipping being a larger risk for human health than the local shipping (> 80%) in Gothenburg. The exposure to ozone related to shipping emissions reduced the acute mortality by 0.4 premature deaths per year due to the NO titration effect. This effect included additional 0.03 deaths attributed to ozone formed from the regional shipping emissions (Table 3). Exposure to NO₂ related to shipping emissions caused additional 2.6 premature deaths year⁻¹, impact of the local shipping being similar to the regional one. This impact corresponded to 35% of the impact of the NO₂ exposure in the Base simulation and was similar to the impact of the road traffic.

5 Assessment of uncertainties and comparison with other studies

Addressing uncertainties in human health risk assessment is a critical issue when evaluating the effects of contaminants on public health due to the complex associations between environmental exposures and health. Uncertainties are introduced with the calculated pollutant concentrations, the grid resolution when assessing the population exposure, the general shape of concentration-response function and transferability problems of the function from region to region. Hammingh et al. (2012) presented an estimate of the uncertainty in the calculations of YOLLs, which may stem from the methodology used in the YOLL calculations and from the spatial resolution. To compare results of Jonsson et al. (2015) with results of this study, the YOLLs pers.⁻¹ from PM₂.₅ exposure calculated in ARP were multiplied by the life expectancy of population above the age of 30, i.e. 50 years, and divided by the population in the model domain. The health impacts of PM₂.₅ were also calculated using the RAINS methodology directly on the calculated PM₂.₅ exposures. Results of both methods are presented, giving very similar results.

The largest uncertainties are associated with the exposure response functions (ERF) as such. In this study impacts for the mean values of ERFs are presented, the 95% confidence interval for these functions is given in the Methods chapter. The ERFs used here are those recommended in WHO (2013a), for PM₂.₅ ERFs with higher values for spatial analyses of air pollution and mortality were found by project Escape for European cohorts (Beelen et al., 2014) as well as for mortalities in Los Angeles (Jerrett et al., 2005, 17 % per 10 µg m⁻³, 95% confidence interval 5–30 %). These ERFs are of very similar value and those of Beelen et al. (2014) were used as alternative functions for estimates of broader uncertainty limits by Barregård et al. (2019). There are two In ARP a linear form of ERFs is applied which is justified by a rather narrow interval of PM exposure levels in Europe. In terms of impact of the total exposure to PM₂.₅ on natural mortality, the linear and log-linear form of the functions give similar results within the concentration range of 10–30 µg m⁻³, the linear model giving slightly lower relative risks in this range and higher relative risks below and above (Ostro et al., 2004). The PM2.5 levels found in pour study falls below 10 µg m⁻³. For regions with high PM2.5 levels different ERF models need to be applied and for global HIA studies or studies in other regions of the world but Western Europe or North America also ERFs for cause-specific mortalities, rather than natural mortalities are usually used. There are two further important issues regarding the uncertainties associated with the ERFs. First, the air pollution represents a complex mixture and individual gases and particles are often correlated. The impacts on mortality calculated for the different pollutants therefore cannot be simply summed up. Second, the ERFs assume that all particulate matter has the same impact. There is increasing evidence of different ERFs for some compounds, primarily elemental and organic carbon (WHO 2013b).
The most robust relation between the air pollution and effects on human health is for particulate matter (WHO 2013b). In Swedish cities, also in Gothenburg, the main contribution to concentrations of PM$_{2.5}$ comes from the background air (Segersson et al., 2017; Gustafsson et al. 2018). Accurate modelling of the total concentration of particulate matter is, however, very difficult as the processes affecting them are extremely complex and many of them not well quantified. These include natural and anthropogenic emissions, formation of secondary particulate matter in complex photochemical processes as well as dry and wet deposition processes that need to be described on the whole range of relevant geographical and time scales. Many regional- and global-scale models tend to underestimate the simulated PM$_{2.5}$ concentrations, especially in summer, when formation of secondary PM is stronger due to the high photochemical activity and the impact of primary PM is lower due to the more intensive mixing and smaller anthropogenic emissions of primary PM in summer (Karl et al., 2019a). Also, the modelled PM concentrations used as the boundary conditions in this study showed underestimates of PM$_{2.5}$ by 60 % and 17 %, on summer average and on annual average, respectively (Karl et al., 2019a). Two studies addressing impacts of shipping on air pollution in Gothenburg (Segersson et al., 2017; Repka et al., 2019) assessed the total concentration levels and contribution of shipping to these, none of them are, however, calculated for year 2012 assessed in this study. Segersson et al. (2017) shows annual mean background PM$_{2.5}$ concentrations for year 2011 of about 5 µg m$^{-3}$ for Gothenburg, reaching concentrations $> 8$ µg m$^{-3}$ in polluted parts of the city. An annual mean concentration map presented in Repka et al. (2019) for year 2016 shows similar concentration levels with background concentrations of about 6 µg m$^{-3}$ and maximum concentrations $> 8$ µg m$^{-3}$. Both studies used PM$_{10}$ monitoring data at the urban background station ‘Femman’ to inversely derive the boundary conditions for PM$_{2.5}$, Jonson et al. (2015 and 2019) studied impacts of the Baltic Sea and North Sea shipping with the EMEP model for year 2010 and 2016 and found in both cases annual mean concentration levels at the Swedish West coast about 4–5 µg m$^{-3}$. This concentration should correspond to the background levels of the city-scale simulations and year 2016. Jonson et al. (2019) also compared the modelled concentrations with background measurements from the station Råö, situated 20 km south of the city, and found a model underestimation of 0.7 ppb for the annual mean. The concentration levels of PM$_{2.5}$ found in this study were lower than in Segersson et al. (2017) and Repka et al. (2019), but they agree reasonably well with Jonson et al., 2015 and 2019. Segersson et al. (2017) addressed health effects of PM$_{2.5}$, PM$_{10}$ and black carbon in three Swedish cities, among them Gothenburg using gaussian model SIMAIR. The population weighted exposure to PM$_{2.5}$ for Gothenburg was calculated to 6.5 µg m$^{-3}$, which was associated with c.a. 150–290 premature deaths from exposure to PM$_{2.5}$. The lower premature death number in Segersson et al. (2017) comes from calculations using the same ERF as in this study while the higher number uses the ERF presented by Jerrett et al. (2005) for PM$_{2.5}$ from the city sources. The values can be compared to the population weighted exposure to PM$_{2.5}$ of 4.1 µg m$^{-3}$, associated to c.a. 140 premature deaths found in this study. Jonson et al. (2015) calculated impact of shipping emissions in the Baltic Sea and the North Sea using the EMEP model and a map presenting geographical distribution of life expectancy loss shows approximately 0.2 YOLLs pers.$^{-1}$ around the West coast of Sweden. This agrees reasonably well with our estimate of 0.18 YOLLs pers.$^{-1}$ in Gothenburg.

It is important to bear the uncertainties in total concentrations of PM and other air pollutants in mind when assessing the relative contribution of shipping to the overall impact of air pollution. Assessments of impacts of selected anthropogenic sources are, however, associated with smaller uncertainties compared to the impact of the total concentrations as some large uncertainties, e.g. those regarding the natural and agriculture sources, cancel out. The study of Segersson et al. (2017) found contribution of shipping to the population weighted annual PM$_{2.5}$ concentration to be 0.04 µg m$^{-3}$ and contribution of the road traffic exhaust emissions to be 0.27 µg m$^{-3}$ which can be compared to 0.09 µg m$^{-3}$ from shipping and 0.22 µg m$^{-3}$ from road traffic found in this study, however, bearing in mind that the studies assessed two different years.

6 Conclusions

The impact of local and regional ship emissions on in the city of Gothenburg was investigated by a multi-model system for the year 2012. The model evaluation against monitoring data demonstrated fairly good agreement in meteorological parameters and acceptable estimation of hourly air pollutant concentrations.
The city-scale model simulations with and without local and regional shipping in the emission inventory revealed that impacts from shipping on air quality in Gothenburg were substantial. The calculated contribution from local shipping to NO\textsubscript{2} was 0.5 ppb on annual average, representing 14 % of calculated annual mean NO\textsubscript{2} concentration. -Including contribution from regional shipping in the North Sea and the Baltic Sea, the total shipping contribution reached 1.5 ppb representing 41 % of calculated NO\textsubscript{2} concentrations. The contribution from regional and local shipping was higher than that from road traffic around the area of the city ports. In an analysis of exposure from different sources using population weighted concentrations, the contribution of regional and local shipping was similar to that of the road traffic in the city.

The model results of ozone concentrations have shown that titration by NO dominated the overall impact of local shipping on ozone concentration levels in Gothenburg. The maximum impact from local NO\textsubscript{X} and NMVOC emissions on summer seasonal mean ozone concentration was calculated to -4 ppb (~14 %). The negative effect of solely NO\textsubscript{X} emissions from local shipping on the ozone concentrations was up to -6 ppb (~21 %), net negative even in the summer, when photochemical activity and potential for ozone formation are high. The emissions of NMVOC from local shipping as such increased the ozone formation in the city with the highest contribution of 2 ppb (~7 %) as seasonal summer mean. In terms of urban air quality control, reduction in anthropogenic NMVOC could result in a significantly greater decrease in O\textsubscript{3} relative to the same reduction in NO\textsubscript{X} (Karl, 2019b).

The simulated emissions from local and regional shipping contributed 0.5 \mu g m\textsuperscript{-3} on model/domain average and at highest 1.1 \mu g m\textsuperscript{-3} to the annual mean concentration of PM\textsubscript{2.5}. Regional shipping is a larger contributor than local shipping to local PM\textsubscript{2.5} concentrations, corresponding to 11 % of the local PM\textsubscript{2.5} concentrations on average. Also its contribution to the PWC was higher, contributing with 0.4 \mu g m\textsuperscript{-3} (10 % of the total PWC for PM\textsubscript{2.5}). Contribution from the local shipping was 0.1 \mu g m\textsuperscript{-3} (2 % of the total).

The calculated health impacts have shown the most serious effects from shipping in Gothenburg to be associated with exposure to PM\textsubscript{2.5}. Local and regional shipping together reduce life expectancy by 0.015 years per person, of which more than 80 % are associated with the regional shipping in the North and the Baltic Sea. The shipping impact is more than twice as high as the modelled impact of PM\textsubscript{2.5} associated with the local road traffic. Impacts from exposure to NO\textsubscript{2} and ozone were calculated in terms of premature deaths per year and 2.6 additional cases year\textsuperscript{-1} were calculated for exposure to NO\textsubscript{2}, regional and local shipping contributing with 59 % and 41 % respectively. Impacts from exposure to ozone were of opposite magnitude. The decrease of ozone due to the NO titration reduced the calculated mortalities by 0.4 cases year\textsuperscript{-1}. The impact of the exposure to PM\textsubscript{2.5} from shipping calculated as premature deaths was 18 cases year\textsuperscript{-1}. The implementation of the more stringent SECA regulations on FSC in year 2015 is not likely to have changed impacts from NO\textsubscript{2} and ozone. According to the study of Jonson et al. (2019), approximately 35 % reduction of the impact from the regional shipping contribution to PM\textsubscript{2.5} could be expected around Gothenburg while a much smaller change can be expected in emission from the local shipping since hotelling and inland shipping already use a fuel with 0.1 % FSC in the model. This would mean similar reductions of the impacts related to PM\textsubscript{2.5} in the city of Gothenburg. Impact of the global cap of 0.5 % for FSC which entered into force the 1\textsuperscript{st} of January 2020 will not have any significant impact on further reduction of shipping-related air pollution in Gothenburg comparing to situation after 2015. Global study of Sofiev et al. (2018) shows that around the Swedish West coast decrease of PM\textsubscript{2.5} due to the global cap would be below 1 %. The more serious health effects induced by regional shipping indicate that close cooperation across governance levels is required to effectively reduce the air pollution in the city.

Impacts of the local shipping emissions on air quality and human health are further discussed in part II paper (Ramacher et al., 2019, the Part II paper), presenting study of several future shipping scenarios for year 2040 adopting changes in shipping emissions due to changes in ship traffic volumes, legislation on emissions of air pollutants at sea, on energy effectivization as well as introduction of shore side electricity in shipping.

This article is part of the special issue “Shipping and the Environment 2017”
**Code availability**
The TAPM model is a commercial software available at CSIRO, Australia (www.csiro.au). STEAM model is intellectual property of the Finnish Meteorological Institute and is not publicly available. ARP is commercial software available from arirabl.com.

**Data availability**
The model output data are available upon request from the corresponding authors.

**Author contribution**
LT, MOPR, JM and VM designed the model simulations. LJ and JPJ calculated ship emissions with the STEAM model and contributed with text about the shipping emissions. LT, MG and JM prepared emission data from other sources. MK and AA prepared data from the regional-scale simulation used for the boundary conditions and MK contributed with text about these simulations. LT and MOPR prepared the model set-up and other input data, performed the model simulations and evaluated the model results. LT calculated exposures and JM and KY calculated the health impacts. LT and JM wrote the major part of the text with assistance from MOPR and VM.

**Competing interests**
JM is associated editor of the special issue Shipping and Environment.

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**References**


SMED: Description of Methods and Quality of Spatially Distributed Emissions to Air, 2015. Contract no 309 1235.


Figure 1: (a) Five nested meteorological model domains with their sizes and spatial resolutions. The fifth domain with air pollution grid (250 x 250 m²) is pointed out in the figure, showing location of the three air quality monitoring sites Femman, Haga and Mölndal, as well as Eriksberg, a residential area close to the harbour; (b) The boundary conditions of air pollution in TAPM, example of summer mean (JJA) year 2012 NO₂ concentrations [ppbV] in the regional-scale CMAQ simulation (4 x 4 km²). Base map credits: © OpenStreetMap contributors (openstreetmap.org). Distributed under a Creative Commons BY-SA License.
Figure 2: Annual local shipping emissions of (a) NOx and (b) PM₁₀ (equal to PM₂.₅) from small vessels with stack height below 36 m (assumed 15 m) and (c) NOx and (d) PM₁₀ from large vessels with high stack height above 36 m (assumed 36 m) in the Gothenburg area. Base map credits: © OpenStreetMap contributors (openstreetmap.org). Distributed under a Creative Commons BY-SA License.
Figure 2: Proportions of different source categories in the local emission inventory for the city-scale model domain in year 2012. The total emissions were 502 tons year\(^{-1}\) for SO\(_2\), 5072 tons year\(^{-1}\) for NO\(_x\), 357 tons year\(^{-1}\) for PM\(_{10}\) and 7457 tons year\(^{-1}\) for VOCs.

Figure 3: (a) Wind rose plots of measurements at Femman station.
Figure 4: Comparison between measured and modelled winds at Femman station: (a) the observed wind rose in January and July; (b) the differences between observed and simulated wind rose, showing the BIAS in the wind speed based on the difference simulated wind speed – measured wind speed. E.g. A positive BIAS from 0 to 2.5 m s⁻¹ in wind direction N has a frequency of almost 30 %.
Figure 5: Summary statistics of model performance for the annual mean values of (a) $NO_2$ (hourly values), (b) $PM_{10}$ (daily values), (c) $PM_{2.5}$ (daily values), and (d) $O_3$ (daily maximum of the 8-hour means), including days of exceedances. Stations (blue dots) within green bar: performance criteria satisfied, stations within orange bar: performance criteria satisfied, error dominated by the corresponding indicator. Green light: > 90 % of the stations fulfil the performance criteria. Red light: < 90 % of the stations fulfil the performance criteria. The indicator $H_{perc}$ indicates the model capability to reproduce extreme events, represented by selected high percentile for modelled and observed values.
Figure 6: Simulated atmospheric concentrations of SO\textsubscript{2} (in ppb) for year 2012: (a) Annual mean concentrations in Base case simulation; (b) Annual mean contribution of the local shipping; (c) Annual mean contribution of the local and the regional shipping; and (d) Same as (c) with main ports along the Göta älv as well as Eriksberg. Base map credits: © OpenStreetMap contributors (openstreetmap.org). Distributed under a Creative Commons BY-SA License.
Figure 7: Simulated atmospheric concentrations of NO₂ (in ppb) for year 2012: (a) Annual mean concentrations in Base case simulation; (b) Annual mean contribution of the local shipping; (c) Annual mean contribution of the local and the regional shipping. Base map credits: © OpenStreetMap contributors (openstreetmap.org). Distributed under a Creative Commons BY-SA License.
Figure 8: Time series of simulated modelled monthly mean contributions of the local shipping and regional shipping, local road traffic and other anthropogenic emissions (including contribution from the boundary conditions) to the NO$_2$ daily mean concentrations at Eriksberg, located at the north of the Göta älv for the year 2012. Modelled local shipping contributions (black line) deduced from the scenarios “Base” and “No local shipping”, regional shipping contributions (red line) deduced from the scenario “No local shipping” and “No local and no regional shipping”, and road traffic contributions (blue line) deduced from the scenario “Base” and “No road traffic” are presented.
Figure 9: Modelled summer mean (JJA) ozone concentrations (ppb) in year 2012 and contributions of local and regional shipping to the summer mean concentration in year 2012: (a) Modelled $O_3$ concentrations in the Base model simulation; (b) Modelled contribution of emissions from local shipping; (c) Modelled contribution of emissions from local and regional shipping; (d) Modelled contributions of NMVOC emissions from local shipping. Base map credits: © OpenStreetMap contributors (openstreetmap.org). Distributed under a Creative Commons BY-SA License.
Figure 10: Modelled annual mean PM$_{2.5}$ concentrations ($\mu g\,m^{-3}$) and contributions of shipping to the annual mean concentrations in year 2012: (a) Modelled annual mean concentrations in Base model simulation; (b) Modelled annual mean contribution of local shipping; (c) Modelled annual mean contributions of local and regional shipping. Base map credits: © OpenStreetMap contributors (openstreetmap.org). Distributed under a Creative Commons BY-SA License.
Figure 11: Modelled monthly mean contributions from local shipping, regional shipping and other sources (including contribution from the boundary condition) to PM$_{2.5}$ concentrations ($\mu$g m$^{-3}$) at Eriksberg, Göta älvs for the year 2012. Modelled local shipping contributions (black bar) are deduced from the differences between the scenarios “Base” and “No local shipping” and modelled regional shipping contributions (red bar) are deduced from the difference between the scenario “No local shipping” and “No local and no regional shipping”.
Figure 12: The population weighted annual mean concentrations for NO$_2$ (ppb × capita), PM$_{2.5}$ (µg m$^{-3}$ × capita) and SOMO35 (ppb × h × capita): (a) NO$_2$ in the Base simulation; (b) PM$_{2.5}$ in the Base simulation; (c) SOMO35 in the Base simulation; (d) NO$_2$ from local and regional shipping; (e) PM$_{2.5}$ from local and regional shipping; (f) SOMO35 from local and regional shipping; (g) NO$_2$ from road traffic; (h) PM$_{2.5}$ from road traffic and (i) SOMO35 from road traffic. Base map credits: © OpenStreetMap contributors (openstreetmap.org). Distributed under a Creative Commons BY-SA License.
### Table 1: City-scale model setup.

<table>
<thead>
<tr>
<th>Source</th>
<th>Domain</th>
<th>Spatial resolutions</th>
<th>Model / Database</th>
</tr>
</thead>
<tbody>
<tr>
<td>Meteorology</td>
<td>30 km × 30 km</td>
<td>500 m</td>
<td>ECMWF ERA5 0.3’ × 0.3’, 21 layers</td>
</tr>
<tr>
<td>Background concentrations</td>
<td>160 km × 96 km</td>
<td>4 km × 4 km</td>
<td>CMAQ</td>
</tr>
<tr>
<td>Local shipping emissions</td>
<td>30 km × 30 km</td>
<td>250 m × 250 m meters (line sources)</td>
<td>STEAM2</td>
</tr>
<tr>
<td>Local traffic emissions</td>
<td>30 km × 30 km</td>
<td>1 km × 1 km</td>
<td>Miljöförvaltningen and HBEFA v. 3.2</td>
</tr>
<tr>
<td>Local industrial, machines, wood burning and aviation <em>etc.</em></td>
<td>30 km × 30 km</td>
<td></td>
<td>SMED</td>
</tr>
</tbody>
</table>

### Table 2: Population weighted annual mean concentrations of NO\(_2\), PM\(_{2.5}\) and SOMO\(_{35}\) associated to all sources, road traffic and local, regional shipping in city of Gothenburg for year 2012.

<table>
<thead>
<tr>
<th>Sources</th>
<th>NO(_2) (ppb)</th>
<th>PM(_{2.5}) (µg m(^{-3}))</th>
<th>SOMO(_{35}) (ppb × h)</th>
</tr>
</thead>
<tbody>
<tr>
<td>All sources PWC in Base simulation</td>
<td>4.70</td>
<td>4.12</td>
<td>19 698</td>
</tr>
<tr>
<td>Road traffic</td>
<td>1.75</td>
<td>0.22</td>
<td>12</td>
</tr>
<tr>
<td>Local and regional shipping</td>
<td>1.65</td>
<td>0.51</td>
<td>-1 115</td>
</tr>
<tr>
<td><strong>Local shipping</strong></td>
<td>0.68</td>
<td>0.09</td>
<td>-1 186</td>
</tr>
<tr>
<td><strong>Regional shipping</strong></td>
<td>0.97</td>
<td>0.42</td>
<td>71</td>
</tr>
</tbody>
</table>

### Table 3: Health impacts calculated for \(O_3\), NO\(_2\) and PM\(_{2.5}\) contributions of the local and regional shipping and the local road traffic to air pollution in the city of Gothenburg as well as of the total exposure to these pollutants in the city. The health impacts calculated with the ARP model and with the RAINS methodology are presented.

<table>
<thead>
<tr>
<th>Pollutant</th>
<th>Impact</th>
<th>Unit</th>
<th>Local shipping</th>
<th>NSBS regional shipping</th>
<th>All shipping</th>
<th>Local road traffic</th>
<th>Total exposure</th>
</tr>
</thead>
<tbody>
<tr>
<td>(O_3)</td>
<td>Acute Mortality (All ages)</td>
<td>Premature deaths</td>
<td>-0.5</td>
<td>0.03</td>
<td>-0.4</td>
<td>0.005</td>
<td>7.6</td>
</tr>
<tr>
<td>NO(_2)</td>
<td>Acute Mortality (All ages)</td>
<td>Premature deaths</td>
<td>1.06</td>
<td>1.52</td>
<td>2.59</td>
<td>2.73</td>
<td>7.35</td>
</tr>
<tr>
<td>PM(_{2.5})</td>
<td>Chronic Mortality (All ages)</td>
<td>Life years lost</td>
<td>31</td>
<td>143</td>
<td>174</td>
<td>74</td>
<td>1393</td>
</tr>
<tr>
<td></td>
<td>Chronic mortality (All ages, ARP)</td>
<td>YOLLS pers.(^{-1})</td>
<td>0.003</td>
<td>0.013</td>
<td>0.015</td>
<td>0.007</td>
<td>0.12</td>
</tr>
<tr>
<td></td>
<td>Chronic Mortality (Age 30+, RAINS)</td>
<td>YOLLS pers.(^{-1})</td>
<td>0.003</td>
<td>0.014</td>
<td>0.018</td>
<td>0.008</td>
<td>0.14</td>
</tr>
<tr>
<td></td>
<td>Chronic mortality relative to that from the total exposure</td>
<td>-</td>
<td>2 %</td>
<td>10 %</td>
<td>12 %</td>
<td>5 %</td>
<td></td>
</tr>
</tbody>
</table>