Would like to thank the reviewer for the job that has been done with the review in the case of constructive improvement proposals. Below follow the comments and our answers highlighted in yellow, and the manuscript with track changes.

**General comments**

The authors have addressed all raised issues and have substantially expanded the discussion in their manuscript, which now I consider much improved. A few final points to correct before proceeding with publication.

**Specific comments**

Figures 2-4: Maybe you could specify that the monthly average values are after deseasonalization.

*It is added in the captions to Figures 2-4 that the monthly averages are based on deseasonalization.*

Page 17, line 19: Check parentheses in the Johansson citation.

*It is changed to Johansson et al. (2008).*

Page 17, line 23: ...trend diminishes...

*Changed to diminishes.*

Section 4.1: A reasonable answer has been provided in the response, regarding different weekday-weekend variability between NOx and PM10. See if you can incorporate this along with the references in the manuscript.

*The following sentences have been incorporated in section 4.1.3: The pronounced weekday-weekend pattern associated with NOx and NO2 is not shown for PM10. Despite that PM10 is mainly related to traffic, other factors also affect this pattern. Since the emissions of road dust highly depend on the wetness of the roads, as shown by Johansson et al. (2007), the diurnal cycles will not follow the same pattern as vehicle exhaust from traffic.*

Page 18, line 10: Replace “photochemistry” with “ozone cycle”.

*Photochemistry is replaced with ozone cycle.*

Page 18, line 20: Replace “perfect” with “pronounced”.
Perfect is replaced with pronounced.

Page 20, line 25: The argument regarding passive vs. active sampling is fair. However, when the authors claim that the ESCAPE field campaigns were of limited extent or even inconsistent, the context has to be explained. I presume that they refer to the infrequent sampling. However, these data have been adjusted to annual means following a specific methodology. They should bear in mind that the same schedule was followed for PM sampling as well, that provided the PM10 RR that they have used in this study. I suggest to change the wording here.

Regarding the ESCAPE Study, we consider passive NO2 sampling less valid than PM monitoring with Harvard impactors. Since we don’t have data on PM absorbance, we remove the sentence in section 4.4 “In addition, Beelen et al. (2014) did not find any significant association with PM2.5 absorption, possibly indicating problems to reflect exposure to vehicle exhaust”. Further down in section 4.4, where the choice of RR for PM10 is explained, the sentence is rewritten as: For PM10, we use the RR value of 1.04 (95 % CI 1.00–1.09) per 10 µg m-3 increase from Beelen et al. (2014), where the meta-analysis in ESCAPE is based on measurement campaigns with Harvard impactors.

Page 21, line 16: Replace “accuracy” with “suitability”.

Accuracy is replaced with suitability.

Page 23, line 10: ...sectoral...

It is changed to sectoral.

Page 24, line 16: “…where NO has the capability of scavenging free oxygen atoms wherein NO2 is formed..”.

Rewrite this as: “..where NO oxidizes to NO2.” or omit the sentence. The titration reaction is between NO and O3 to form NO2.

It is rewritten.

Figure A1-A4: Please also correct separators for the axis values.

The separators are corrected.
Trends in air pollutants and health impacts in three Swedish cities over the past three decades

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Abstract. Air pollution concentrations have been decreasing in many cities in the developed countries. We have estimated time trends and health effects associated with exposure to NOx, NO2, O3, and PM10 in the Swedish cities Stockholm, Gothenburg, and Malmö from the 1990’s to 2015. Trend analyses of concentrations have been performed by using the Mann-Kendall test and the Theil-Sen method. Measured concentrations are from central monitoring stations representing urban background levels, and they are assumed to indicate changes in long-term exposure to the population. However, corrections for population exposure have been performed for NOx, O3, and PM10 in Stockholm, and for NOx in Gothenburg. For NOx and PM10, the concentrations at the central monitoring stations are shown to overestimate exposure when compared to dispersion model calculations of spatially resolved population-weighted exposure concentrations, while the reverse applies to O3. The trends are very different for the pollutants that are studied; NOx and NO2 have been decreasing in all cities, O3 exhibits an increasing trend in all cities, and for PM10, there is a slowly decreasing trend in Stockholm, a slowly increasing trend in Gothenburg, and no significant trend in Malmö. Trends associated with NOx and NO2 are mainly attributed to local emission reductions from traffic. Long-range transport and local emissions from road traffic (non-exhaust PM emissions) and residential wood combustion are the main sources of PM10. For O3, the trends are affected by long-range transport, and there is a net removal of O3 in the cities. The increasing trends are attributed to decreased net removal, as NOx emissions have been reduced.

Health effects in terms of changes in life expectancy are calculated based on the trends in exposure to NOx, NO2, O3, and PM10, and the relative risks associated with exposure to these pollutants. The decreased levels of NOx are estimated to increase the life expectancy by up to 11 months for Stockholm and 12 months for Gothenburg. This corresponds to up to one fifth of the total increase in life expectancy (54–70 months) in the cities during the period 1990–2015. Since the increased concentrations in O3 have relatively small impact on the changes in life expectancy, the overall net effect is increased life expectancies in the cities that have been studied.
I. Introduction

Air pollution exposure is clearly recognised as an important global risk factor for the development of a large number of diseases and disabilities (Cohen et al., 2017). In 2015, exposure to PM$_{2.5}$ (particles with an aerodynamic diameter smaller than 2.5 micrometer) was ranked number five among all risk factors for premature deaths, where smoking, diet, and high blood pressure are included. It is estimated to have caused 4.2 million premature deaths among the world’s population. This can be compared to smoking, with a corresponding value of 6.4 million premature deaths for the year 2015 (State of Global Air, 2017).

In view of the apparent major health impact associated with exposure to air pollutants, it is of great importance to analyse the trends in air pollution concentrations, and how these affect the public health.

Worldwide, many cities, especially in high-income countries, show substantial decreasing trends in air pollution concentrations (e.g. WHO, 2016a; Geddes et al., 2016; Colette et al., 2011). Considering the changes in the total emissions in Europe and in the U.S., respectively, there are different conditions. In Europe (EU–27), during the period 2002–2011, the emissions of nitrogen oxides (NO$_x$ = sum of NO and NO$_2$) have decreased by 27 %, and the emissions of primary PM$_{10}$ (particles with an aerodynamic diameter smaller than 10 micrometer) have decreased by 14 % (Guerreiro et al., 2014). In the U.S., the NO$_x$ emissions in eight cities have decreased between 25 % and 48 % during the period 2005–2012 (Tong et al., 2015). Considering the trends in the U.S. regarding all sector-specific emissions during the period 1990–2010, NO$_x$ exhibits a decrease of 48 %, while PM$_{10}$ exhibits a decrease of 50 % (Xing et al., 2013). So, the NO$_x$ and PM$_{10}$ emissions have decreased almost equally in the U.S., while in Europe, the emissions regarding NO$_x$ exhibit a sharper decline compared to PM$_{10}$ (Guerreiro et al., 2014). However, it should also be noted that there are large between-country variations in NO$_x$ emission trends, partly reflecting that some countries have had problems meeting the original National Emission Ceilings and the Air Quality directives (EEA, 2017).

For ozone (O$_3$) in Europe, the concentrations tend to increase at urban sites, especially during the cold season, as observed by e.g. Sicard et al. (2016) for cities in France, and for most cities in Europe by Colette et al. (2011). This rise can be explained by increases in imported O$_3$ by long-range transport, and also by a decreased titration by nitrogen monoxide (NO), due to the reduction in local NO$_x$ emissions (Colette et al., 2011; Sicard et al., 2016). However, trends in O$_3$ are also different for summer and winter, with mainly decreasing trends in the summer and increasing trends in the winter, and there are also some variations between cities in the EU (EEA, 2016). Furthermore, considering different statistical metrics, the trends for O$_3$ at both urban and rural sites in the UK are different during the period 1993–2011, depending on the metric that is used; mean and median trends are positive, while the maximum trend is negative (Munir et al., 2013). In Northern Alberta in Canada, where measurement results from four urban locations were analysed, the mean concentrations of O$_3$ have increased at most stations during the period 1998–2014 (Bari and Kindziersky, 2016).

This means that the general population exposure, and presumably also associated health effects of the urban populations, have been reduced for NO$_x$ and PM$_{10}$, but increased for the mean O$_3$ concentrations. So far, there are, however, rather few studies that have assessed the net health gain or health loss associated with the trends. Henschel et al. (2012) has examined
intervention studies focusing on improvements in air quality, and associated health benefits for the assessed population. Some studies have focused on trends and effects associated with particulate matter (Tang et al., 2014; Keuken et al., 2011; Correria et al., 2013), and some studies have also included ozone (Fann and Risley, 2013; Gramsch et al., 2006). Correria et al. (2013) estimated that for the most urban communities in the USA, as much as 18% of the increase in life expectancy from 2000 to 2007 was attributable to the reduction in PM$_{2.5}$. When health improvements associated with PM$_{2.5}$ trends are compared to those for O$_3$, the health benefits associated with PM$_{2.5}$ are much greater (Fann and Risley, 2013).

In a few studies, air quality dispersion models together with gridded population data and exposure-response functions have been used to assess health impacts of changing air pollutant emissions. E.g. for Rotterdam (The Netherlands), the health benefits associated with decreasing trends for EC (elemental carbon) and PM$_{10}$ have been calculated for the period 1985–2008, and the average gain in life expectancy was 13 and 12 months per person for PM$_{10}$ and EC, respectively (Keuken et al., 2011). Fann and Risley, (2013) estimated changes in ozone and PM$_{2.5}$ concentrations in the U.S. during the period 2000–2007, and they estimated the impact on the number of premature deaths by using health-impact functions based on short-term relative risk estimates for O$_3$ and long-term relative risk estimates for PM$_{2.5}$. Data from monitoring stations were spatially interpolated. Overall, they found net benefits in the number of premature deaths ranging from 22 000 to 60 000 for PM$_{2.5}$, and from 880 to 4 100 for ozone, but interestingly, they found opposing trends in premature mortality associated with PM$_{2.5}$ and ozone at some locations, and a considerable year to year variation in the number of premature deaths. Tang et al. (2014) evaluated the health benefits associated with coal-burning factory shutdowns and accompanied decreasing levels of PM$_{10}$ in Taiyuan in China during the period 2001–2010. They used PM$_{10}$ measurements from monitoring stations, but without considering spatial variations in the exposures, and they used the Chinese national standard of 40 µg m$^{-3}$ as a threshold level. The number of premature deaths associated with PM$_{10}$ levels dropped from 4 948 in 2001 to 2 138 in 2010, and they did not include other pollutants in their analysis.

The studies that have been referred above have used different methodologies of estimating the exposure; by simply taking the concentration at the monitoring stations, and making spatial interpolations between several monitoring stations, or by using dispersion modelling to estimate the spatial distribution of the exposure concentrations. They have also used different relative risks for mortality, and different ways to apply the baseline mortality and the health-effect exposure threshold.

The objective of this study is to quantify and compare the changes in life expectancy, resulting from changes in different pollutants, namely NO$_x$, NO$_2$, O$_3$, and PM$_{10}$, based on measurements during 25 years in the three largest cities in Sweden. We show that the health impacts related to change in different pollutants are different, and we aim to discuss how different trends in different air pollutants affect the life expectancy assessment.
2. Methods

2.1 The choice of air pollutants for trend analysis

Our main trend analyses are based on simultaneous, continuous measurements of NO\textsubscript{x}, NO\textsubscript{2}, O\textsubscript{3}, and PM\textsubscript{10}, during the period 1990 to 2015 in Stockholm, Gothenburg, and Malmo; the three largest cities in Sweden. Based on these trends, we calculate the health impacts associated with changes in the population exposure. The changes in life expectancy are calculated with the mean values and the 95 % confidence intervals of the relative risks, while for the trends and the population-weighted exposure concentrations, only the median and the mean values, respectively, have been used, but without considering their confidence intervals. This means that the ranges of the confidence intervals are narrower than they would have been if the confidence intervals of the trend-lines and the population-weighted exposure concentrations were also included in the calculations. NO\textsubscript{2}, O\textsubscript{3}, and PM\textsubscript{10} have been regulated in the EU directives, and these regulations also set methods, quality assurance, and control (EU, 2008), making these pollutants relevant to analyse in our study.

2.2 Measuring sites and instrumentation

In all analysed cities, the measuring site is located in the city centre and represents the urban background (Fig. 1). Stockholm is the capital and the largest city in Sweden and has a temperate climate with four distinct seasons. Gothenburg is the second largest city located on the west coast of Sweden. It is like Stockholm located within the west-wind belt, and the proximity to the Atlantic Ocean means a slightly milder climate compared to Stockholm. Malmo is located in the southernmost part of Sweden. General information about the three cities regarding population structure, life expectancy at birth, and baseline mortality is presented in Table A1.

The measuring station in Stockholm is located at Torkel Knutssonsgatan on a roof 20 m above ground level, in Gothenburg it is located on a roof 30 m above ground level in the neighbourhood Östra Nordstan, and in Malmo it is located on a roof 20 m above ground level at the city hall (Rådhuset) in the city center (Fig. 1). They are all regulatory monitoring urban background stations using reference methods. In all stations, PM\textsubscript{10} has been measured by using Tapered Element Oscillating Microbalance (TEOM 1400A, Thermo Fisher Scientific, USA). However, in Gothenburg, the Continuous Dichotomous Ambient Air Monitor (Thermo Scientific 1405-DF TEOM) has also been used. Nitrogen oxides are measured in all stations by using chemiluminescense (AC 32M, Environnement SA., France). In Stockholm, ozone is measured by using UV-absorption (O342M, Environnement S.A, France). In Gothenburg, ozone has been measured by using non-dispersive UV photometry (EC9811, Ecotech Pty Ltd, Australia), and also by using CLD 700 AL, Ecophys, Switzerland, and in recent years by using T200, Teledyne, USA. In Malmo, ozone has been measured by using UV-absorption (Thermo Environmental Instruments Model 49C, USA).
2.3 Statistical analysis of the trends

The changes in air pollution concentrations during the period 1990–2015 have been calculated by using the Openair package (Carslaw and Ropkins, 2012). For the trend analyses, the Mann-Kendall test and the Theil-Sen method have been used. The Mann Kendall test is a non-parametric trend test, which is based on the ranking of observations (Hirsch et al. 1982). The Theil-Sen method is used to calculate the median slope of all possible slopes that may occur between the data points (Theil, 1992; Sen, 1968). In our calculations, the trends are based on monthly averages, and they are adjusted for seasonal variations, as these can have a significant effect on monthly data. The Theil-Sen method is regarded as more suitable than the linear-regression method, as it gives more accurate confidence intervals with non-normal distributed data, and it is not affected as much by outliers.

2.4 Health impact and life expectancy calculations

Trends in urban background concentrations reflect the change in population exposure over time (see further below), and they are used to calculate changes in health impacts, presented as changes in life expectancy. As basis for the calculations, we use the population size, age distribution, and mortality rate in Stockholm, Gothenburg, and Malmo according to the year 1997, where data in different age groups have been taken from Statistics Sweden (SCB, 2017), and from the Swedish National Board of Health and Welfare (Socialstyrelsen, 2017). Mortality data before 1997 are not available. However, our own test runs have shown that the calculations in life expectancy give very similar results regardless of the year (1997–2015) in which the population structure and mortality statistics are based on. To illustrate the health benefits associated with decreasing trends, the increase in life expectancy at birth has been calculated in AirQ+ according to a log-linear function (Eq. 1). Similarly, the decrease in life expectancy at birth associated with increasing trends has been calculated in AirQ+ according to the same log-linear function (Eq. 1) (WHO, 2016b). We have applied relative risks obtained from previous epidemiological studies, where
the relationships between mortality and exposure to NO\textsubscript{x}, NO\textsubscript{2}, O\textsubscript{3}, and PM\textsubscript{10} have been analysed. The concept relative risk (RR) represents a log-linear exposure-response function, where the ratio of the incidence in an exposed group is compared to the incidence in a non-exposed, or a less exposed, group. The mortality rate associated with a change in exposure to pollutant x is calculated as:

\[ RR(x) = e^{\beta(x-x_0)} \]  

(1)

where the beta-coefficient (\( \beta \)) indicates the linear relationship between the health impact and the change (x – x\textsubscript{0}) in exposure.

For NO\textsubscript{x}, we apply the RR 1.06 (95 % CI 1.03–1.09) per 10 \( \mu g \) m\textsuperscript{-3} increase, based on the results from Stockfelt et al. (2015), representing all-cause mortality associated with long-term exposure to NO\textsubscript{x} in a cohort with men in Gothenburg.

For NO\textsubscript{2}, we apply the RR 1.066 (95 % CI 1.029–1.104) per 10 \( \mu g \) m\textsuperscript{-3} increase. This RR is based on pooled estimates of mortality associated with long-term exposure to NO\textsubscript{2} (Faustini et al., 2014).

For O\textsubscript{3}, we apply the RR 1.02 (95 % CI 1.01–1.04) per 10 ppb increase, corresponding to 1.01 (95 % CI 1.005–1.02) per 10 \( \mu g \) m\textsuperscript{-3} at 25 °C and 1 atm, based on a large prospective study examining the associations between long-term ozone exposure, and all-cause and cause-specific mortality (Turner et al., 2016).

For PM\textsubscript{10}, we apply the RR 1.04 (95 % CI 1.00–1.09) per 10 \( \mu g \) m\textsuperscript{-3} increase, which is based on a meta-analysis of 22 European cohorts (Beelen et al., 2014). This value is also in line with the RR estimate of 1.043 (95 % CI 1.026–1.061) per 10 \( \mu g \) m\textsuperscript{-3} increase, since many years used in impact assessments (Künzli et al., 2000), based on cohort studies in the U.S.

All RRs described above are based on calculations for the population aged 30 years and over, and therefore, the changes in life expectancy, calculated in AirQ+, are also based on the age group 30 years old and over. The calculations have also been performed by assuming no threshold under which no effect occurs.

### 2.5 Relationship between urban-background concentrations and population-weighted exposure concentrations

The trends measured at urban background monitoring stations may not be representative for the trends in exposure to the entire population in those areas, due to the position of the urban-background measuring stations, the spatial variation in air pollution concentrations, and the variability in the population density within the cities. In order to assess the health effects of the population, associated with changes in air pollution exposure in each metropolitan area, we estimated the relations between the concentrations at the urban-background monitoring stations, and the population-weighted exposure concentrations. This is assessed by comparing model-calculated annual population-weighted exposure concentrations with annual-mean urban-background concentrations. We have done this for NO\textsubscript{x} in Stockholm and Gothenburg, and for O\textsubscript{3} and PM\textsubscript{10} in Stockholm, but due to lack of data, these relations have not been possible to calculate other than for the above-mentioned pollutants and cities.

Geographically resolved annual-mean concentrations of NO\textsubscript{x} in Stockholm were calculated by using a wind model and a Gaussian air-quality dispersion model as a part of the Airviro system (Airviro, 2017). Details on the modelling and emission data are described in Johansson et al. (2017). The same modelling of NO\textsubscript{x} was done for Gothenburg by using a Gaussian model (Aermod, US EPA) as a part of the EnviMan AQ Planner (OPSIS, Furulund, Sweden) as described in Molnár et al. (2015).
Spatially resolved O\textsubscript{3} concentrations in Stockholm are calculated from a combination of measurements and dispersion modelling of NO\textsubscript{x} concentrations. The modelled NO\textsubscript{x} concentrations are converted to O\textsubscript{3} based on the measured NO\textsubscript{x} concentrations at an urban background site in Stockholm, and the difference between O\textsubscript{3} at an urban background site and at a rural background site. The basic chemistry behind this is that the O\textsubscript{3} concentration within a city is controlled by the transport from the surrounding areas into the city, and by the removal of O\textsubscript{3} due to the reaction with NO (nitrogen monoxide). It is further set that if O\textsubscript{3} at the urban background is higher compared to the rural background, then it is set to the same value as the urban background, and if it becomes less than zero, it is set to zero. More details of this method is described in Olsson et al. (2016).

Population-weighted exposure concentrations (C\textsubscript{pop}) are obtained by multiplying the calculated concentration (C\textsubscript{i}) in each grid-square cell with the number of people in the corresponding grid-square cell (P\textsubscript{i}), and summing all products, and dividing the sum by the total population (Eq. 2). This procedure has been used in several previous studies (e.g. Johansson et al., 2009; Orru et al., 2015).

\[
C_{pop} = \frac{\sum C_i P_i}{\sum P_i}
\]  

(2)

The relationships between urban-background levels and population-weighted exposure concentrations are presented in Appendix A Fig. A1–A4.

3. Results

3.1 Overview of trends

Fig. 2–4 show the trends in concentrations of NO\textsubscript{x}, NO\textsubscript{2}, O\textsubscript{3}, and PM\textsubscript{10} measured at urban background sites in Stockholm, Gothenburg, and Malmo during the period 1990–2015. During the given time periods, NO\textsubscript{x} and NO\textsubscript{2} exhibit decreasing trends in all cities, whereas O\textsubscript{3} exhibit increasing trends in all cities, and for PM\textsubscript{10}, the trends are less clear and consistent. For PM\textsubscript{10} in Stockholm and Malmo, the data from the measuring stations only include the period 1997–2015 and 1996–2015, respectively. In several cases, the trends are not perfectly linear throughout the periods (Fig. 2–4).

For NO\textsubscript{x}, NO\textsubscript{2}, and PM\textsubscript{10}, the trends are based on the monthly average concentrations, and for O\textsubscript{3}, they are based on rolling eight-hour daily maximum concentrations. The reason for this division is that the relative risk of O\textsubscript{3}, used in our health-impact calculations, is based on eight-hour daily maximum values. In Fig. 2–4, the median slopes are calculated according to the Theil-Sen method (Theil, 1992; Sen, 1968).
Figure 2. Stockholm: Trends in NO\textsubscript{x}, NO\textsubscript{2}, O\textsubscript{3}, and PM\textsubscript{10}, measured from 1990–2015. For PM\textsubscript{10}, data are only for the period 1997–2015. The blue rings are the deseasonalized monthly averages, and the calculated deseasonalized trends using the Theil-Sen method are shown as the red thick lines. Unit for the trends is µg m\textsuperscript{-3} year\textsuperscript{-1}, and the values in parentheses are 95% confidence intervals. The stars (*) represent the significance level of the trend-lines, where three stars means that p < 0.001, and one star means that 0.01 < p < 0.05.
Figure 3. Gothenburg: Trends in NO\textsubscript{x}, NO\textsubscript{2}, O\textsubscript{3}, and PM\textsubscript{10}, measured from 1990–2015. The blue rings are the deseasonalized monthly averages, and the calculated deseasonalized trends using the Theil-Sen method are shown as the red thick lines. Unit for the trends is µg m\textsuperscript{-3} year\textsuperscript{-1}, and the values in parentheses are 95 % confidence intervals. The stars (*) represent the significance level of the trend-lines, where three stars means that p < 0.001, and one star means that 0.01 < p < 0.05.
Figure 4. Malmo: Trends in NO\textsubscript{x}, NO\textsubscript{2}, O\textsubscript{3}, and PM\textsubscript{10}, measured from 1990–2015. For PM\textsubscript{10}, data are only for the period 1996–2015. The blue rings are the deseasonalized monthly averages, and the calculated deseasonalized trends using the Theil-Sen method are shown as the red thick lines. Unit for the trends is µg m\textsuperscript{-3} year\textsuperscript{-1}, and the values in parentheses are 95% confidence intervals. The stars (*) represent the significance level of the trend-lines, where three stars means that \( p < 0.001 \), and one star means that \( 0.01 < p < 0.05 \).

In Figure 5, we show the trends for the first period 1990–2007, and for the second period 2008–2015 separately (except for PM\textsubscript{10} which for Stockholm is from 1997–2007 and from 2008–2015, and for Malmo from 1996–2007 and from 2008–2015).

In Stockholm, the NO\textsubscript{x} and NO\textsubscript{2} concentrations have decreased significantly from 1990–2007, but for 2008–2015, there is even a tendency for increasing concentrations. The patterns are somewhat different in Gothenburg and Malmo.
Figure 5: The trends for the measured pollutants in µg m⁻³ year⁻¹ (95 % CI) for the whole period, and divided into two different periods.

There are also significantly different trends during weekdays and weekends in the three cities. In Fig. 6–8, the trends have been divided into weekdays and weekends, reflecting the importance of local emissions compared to non-local emissions, where local emissions from traffic are more prominent during weekdays compared to weekends. Fig. 6–8 are otherwise designed in the same manner as Fig 2–4. For NOₓ and NO₂, the downward trends in all cities are more prominent during weekdays compared to weekends, indicating that local emission reductions, mainly from traffic, have had the greatest impact. For O₃, the increasing trends in Gothenburg and Malmo are slightly higher during the weekends compared to the weekdays, while this does not apply to Stockholm. The trends related to PM₁₀ are less clear, and they are also in many cases not statistically significant. Possible reasons for the appearances of the trends are further analysed in the discussion section.
Figure 6. Stockholm: The trends for NO\textsubscript{x}, NO\textsubscript{2}, O\textsubscript{3}, and PM\textsubscript{10}, divided into weekdays and weekends.
Figure 7. Gothenburg: The trends for NO$_x$, NO$_2$, O$_3$, and PM$_{10}$, divided into weekdays and weekends.
Figure 8. Malmo: The trends for NO\textsubscript{x}, NO\textsubscript{2}, O\textsubscript{3}, and PM\textsubscript{10}, divided into weekdays and weekends.

All the trends, except for PM\textsubscript{10} in Malmo, are statistically significant within a 95 % CI. It can be noted that the rate of decreasing NO\textsubscript{x} is highest in Stockholm, and then comes Gothenburg and last Malmo, whereas the opposite order is for the rate of increasing O\textsubscript{3} concentrations. If the change in O\textsubscript{3} was only associated with local NO titration, the opposite would be expected, so this indicates that other factors are also important for the O\textsubscript{3} trends, like the O\textsubscript{3} concentrations at the regional background sites outside the cities.

3.2 Health impact assessment associated with the trends in air pollution concentrations

3.2.1 Change in life expectancy

In order to estimate the health impacts associated with changing concentrations of NO\textsubscript{x}, NO\textsubscript{2}, O\textsubscript{3}, and PM\textsubscript{10} in Stockholm, Gothenburg, and Malmo, the concentration changes during the given time periods for the pollutants, presented in Table 1, and the relative risks, presented in section 2.4, have been used. Adjustments for population-weighted exposure concentrations have also been performed in the cases where data are available (Fig. A1–A4). The changes in life expectancy are calculated with...
the mean values and the 95% confidence intervals of the relative risks, while for the trends and the population-weighted exposure concentrations, only the median and the mean values, respectively, have been used, but without considering their confidence intervals. NO\textsubscript{x} and NO\textsubscript{2} exhibit decreasing trends in all cities, which means an increase in life expectancy (positive values). The opposite applies to O\textsubscript{3}, with increasing trends in all cities. For PM\textsubscript{10}, there is a decreasing trend in Stockholm, with an increase in life expectancy, and an increasing trend in Gothenburg, with a decrease in life expectancy. For PM\textsubscript{10} in Malmo, however, no life expectancy change has been possible to calculate, due to the lack of a significant trend (see Fig. 4).

The detailed results are presented in Table 1.

The largest increase in life expectancy is expected due to the reduction in NO\textsubscript{x}, and somewhat less due to the reduction in NO\textsubscript{2} concentrations. The decrease in NO\textsubscript{x} levels corresponds to up to about 20% of the total population life expectancy increase of between 54 and 70 months during the period 1990–2015 (Table A1.). In opposite, the increase in ozone exposure might have caused the life expectancy to decrease with one to two months. For PM\textsubscript{10}, the effects are very mixed; in Stockholm, there might have been a small increase in life expectancy, whereas in Gothenburg, a small decrease has appeared, and for Malmo, no life expectancy change has been possible to calculate.

Table 1: Change in life expectancy in months with 95% CI in brackets, caused by change in exposure during the measured periods. Decreasing trends are associated with an increase in life expectancy, and increasing trends are associated with a decrease in life expectancy (minus signs). The change in life expectancy, adjusted for population-weighted exposure concentrations (see Section 2.5 and Fig. A1–A4), are presented in bold below the ordinary values. Note that the trend of PM\textsubscript{10} in Stockholm is only for the period 1997–2015.

<table>
<thead>
<tr>
<th>Pollutant</th>
<th>Stockholm</th>
<th>Gothenburg</th>
<th>Malmo</th>
</tr>
</thead>
<tbody>
<tr>
<td>NO\textsubscript{x}</td>
<td>21 (11–32)</td>
<td>17 (9–26)</td>
<td>13 (7–19)</td>
</tr>
<tr>
<td>NO\textsubscript{2}</td>
<td>10 (5–15)</td>
<td>11 (5–16)</td>
<td></td>
</tr>
<tr>
<td>O\textsubscript{3}</td>
<td>13 (6–21)</td>
<td>6 (3–9)</td>
<td>7 (3–11)</td>
</tr>
<tr>
<td>PM\textsubscript{10}</td>
<td>-1 (-0.5 – -2)</td>
<td>-2 (-1 – -4)</td>
<td>-2 (-1 – -4)</td>
</tr>
</tbody>
</table>

4 Discussion

4.1 Trends in concentrations

4.1.1 NO\textsubscript{x} and NO\textsubscript{2}

In all the three cities, the NO\textsubscript{x} and NO\textsubscript{2} trends for urban background concentrations are decreasing. The emissions of NO\textsubscript{x} and PM from different sectors during 1990–2015 have been quantified for Stockholm and Gothenburg as part of a Swedish research programme called SCAC (www.scac.se) (Segersson et al., 2017; Stockfelt et al., 2017). These studies show that the emissions of NO\textsubscript{x} in both Stockholm and Gothenburg are dominated by the contribution from road traffic. The emissions dropped by 60
to 70 % from 1990 to 2011 in Gothenburg (inferred from Figure 3a in Stockfelt et al., 2017). A similar trend in road traffic emissions is calculated for Stockholm based on reported emission data for 1993 and 2017 (SLB, 1995 and SLB, 2018). This is mainly due to decreased vehicle emissions associated with the renewal of the vehicle fleet. The effect of changes in the number of vehicle kilometres is relatively small compared to the effect of lower emissions per kilometre. Other local sources have minor impact on NOₓ, as shown for Stockholm in Johansson et al., 2008). The rural background concentrations of NOₓ outside the cities are low (2–8 µg m⁻³ or between 6 % and 14 % of the urban background concentrations), and not very important for the overall trend (Figure 9 in IVL, 2016). The importance of road traffic is also consistent with the clear difference between the trends measured during weekdays and weekends (Fig. 6–8).

In the case of Stockholm, the downward trend diminishes almost entirely from 2008 onwards, and exhibit rather an increasing trend (Fig. 5). Based on statistics on the share of new registrations of diesel vehicles in the three cities during the period 2008–2015, the proportion of new registrations of diesel vehicles has been greatest in the Stockholm area, which possibly can explain these slightly increasing NOₓ and NO₂ trends seen in Stockholm from 2008 onwards, but not in Gothenburg and Malmo (BilSweden, 2018).

Comparing NOₓ and NO₂ in the three cities, the decreasing trends for NOₓ are much more pronounced compared to NO₂.

The proportion of NOₓ that is NO₂ has been shown to be increasing at kerbside sites in many cities, and this has partly been attributed to more directly emitted NO₂ due to the increase in diesel vehicles equipped with oxidation catalysts (Carslaw, 2005) and particle filters (Grice et al., 2009). However, for urban background sites, the observed increased share of NO₂, and associated slower decline in concentrations compared to NOₓ, are mainly related to the change in O₃/NOₓ equilibrium, caused by decreased NOₓ and increased O₃ concentrations (Keuken et al., 2009) and not to the direct NO₂ emissions.

### 4.1.2 Ozone

The 8-hour daily maximum O₃ levels exhibit increasing trends in all the three cities during the period 1990–2015. Since the concentrations of O₃ are lower at central urban-background sites in the Cities compared to outside the cities, the net effect of the ozone cycle photochemistry is that ozone is consumed, mainly due to the titration involving its reaction with NO. Reduced emissions of NO mean that less O₃ is consumed, due to a reduction of the NO titration, and thereby arises an increasing trend, consistent with many other cities in Europe (Sicard et al., 2016).

The increasing trend of O₃ in Stockholm, based on 8-hour daily maximum values, is weaker compared to Gothenburg and Malmo, even though the NOₓ trend in Stockholm exhibits a sharper decline compared to Gothenburg and Malmo. This may, however, be explained by considering the trends in the regional background concentrations of O₃. The regional background concentrations of O₃ are measured at Aspvreten 60 km SW of Stockholm, at Råö 45 km S of Gothenburg, and at Vavihill 50 km N of Malmo. The trend for O₃ at Aspvreten is more sharply decreasing in comparison with the trends at Råö and Vavihill (SEPA, 2017a). This means that the site transport of O₃ into Stockholm during 1990–2015 decreases more strongly in relation to Gothenburg and Malmo, which explains the weakest increasing O₃ trend in Stockholm, despite the sharpest decreasing NOₓ.
trend. The differences between the regional background trends in O₃ may explain the absence of pronounced perfect anti-correlations between NOₓ and O₃ in the cities.

4.1.3 PM₁₀

The trends for PM₁₀ are not as clear as those for NOₓ and O₃. Only Stockholm exhibits a significant downward trend for the entire period. PM₁₀ is a mixture of locally produced particles and long-range transport of particles from emissions in other countries. The main local (primary) emissions of PM₁₀ are non-exhaust particles from road traffic, ranging from 25 % to 55 %, and biomass burning ranging from 30 % to 35 % (Segersson et al. 2017; Stockfelt et al., 2017). Primary vehicle exhaust particles contribute with less than 10 % of the total mass (Johansson et al., 2007). Road wear due to the use of studded winter tyres is the main source of non-exhaust PM₁₀. The decreasing trend of PM₁₀ in Stockholm can be explained by the reduced use of studded tires, and also by dust binding that has been introduced in the recent years (SLB, 2015).

Unlike Stockholm, Gothenburg exhibits increasing trends for PM₁₀ during the measurement period (Fig. 3 and 7), however, not significant during weekdays. According to Fig. 5, the trend for PM₁₀ in Gothenburg from 2008 onwards is negative, but not significant. During 2006, measures were established in order to reduce the levels of PM₁₀. Dissemination of dust-binding agents, prohibiting the use of studded tires, and the introduction of congestion charges during 2013 may have contributed to this changing trend (Gothenburg City, 2015; Gothenburg Annual Report, 2015).

For Malmo, the trends are not significant, neither for weekdays, weekends, or for the entire period (Fig. 4 and 8).

The pronounced weekday-weekend pattern associated with NOₓ and NO₂ is not shown for PM₁₀. Despite that PM₁₀ is mainly related to traffic, other factors also affect this pattern. Since the emissions of road dust highly depend on the wetness of the roads, as shown by Johansson et al. (2007), the diurnal cycles will not follow the same pattern as vehicle exhaust from traffic.

The trend in concentrations measured at the regional background outside Stockholm (Aspvreten) is decreasing during the period, but it is fragmented with lack of data during parts of the period. For Gothenburg (Råö) and Malmo (Vavihill), there are only data for very limited parts of the period, and firm conclusions about their impact on the urban background concentrations cannot be made (SEPA, 2017b).

4.2 Comparisons between our trends and the trends in U.S. and Europe as a whole

Regarding PM₁₀, the trends in the Swedish cities do not exhibit the same decrease as the trends in the U.S. or in Europe as a whole. An important reason for this is the large contribution of mechanically generated road-dust particles in the Swedish cities. In Stockholm, up to 90 % of the mass fraction of PM₁₀ is generated from road abrasion, which is mostly caused by the use of studded tires during winter-time, and the concentration trend for PM₁₀ is not significantly influenced by the reduction of exhaust emissions (Johansson et al., 2007). Similar conditions regarding the composition of PM₁₀ prevail in Gothenburg (Grundström et al., 2015). In Europe as a whole, the PM₁₀ concentrations exhibit a decreasing trend during the period 1990–2010, and this is mainly caused by reduced emissions of both primary PM and precursors of secondary PM within the European countries (EEA, 2017).
According to Geddes et al. (2016), where the global population-weighted annual mean concentrations of NO$_2$ from 1996–2012 were analysed, the average downward trend in North America (Canada and U.S.) is 4.7 % per year, which is the most heavily declining trend compared to all the other regions in the world. For Western Europe, the average downward trend per year is, according to Geddes et al. (2016), 2.5 % per year. This downward trend is in the same magnitude as the NO$_2$ trends in Stockholm, Gothenburg, and Malmo (Fig. 2, 3, and 4).

For O$_3$, the comparison is a little bit more complicated, since the trends may differ depending on the measure that is used. In the U.K., during the period 1993–2011, the mean and median concentration trends were positive, while the maximum concentration trend was negative (Munir et al., 2013). In Europe as a whole, the O$_3$ concentrations are influenced by both local emissions, intercontinental inflow, and the meteorological conditions. The O$_3$ concentrations, and their changes in Europe during the period 1990–2010, are very different depending on time and place, and no unambiguous explanation or interpretation is possible to do (EEA, 2017). Considering the O$_3$ trends in Fig. 2–4, where these are based on 8-hour daily maximum values, all the trends are increasing. However, the O$_3$ concentrations associated with monthly averages also show increasing trends for all the three cities during 1990–2015, and as mentioned in section 4.1, the increasing O$_3$ concentrations in the Swedish cities are probably largely caused by a reduced titration effect, which in turn is caused by decreasing NO concentrations.

### 4.3 Impacts on life expectancy

In general, the impact of NO$_x$ and NO$_2$ on life expectancy among the populations is much larger compared to the impact of O$_3$ and PM$_{10}$. The calculated gain in life expectancy, associated with decreasing NO$_x$ trends (Table 1), contributes up to as much as about 20 % of the total gain in life expectancy during the period 1990–2015 (Table A1). However, since the O$_3$ concentrations exhibit increasing trends during the same period, and thereby giving rise to a loss in life expectancy, the summarized effects of NO$_x$ exposure and O$_3$ exposure may be relevant to consider. For Stockholm, where population-weighted exposure concentrations are available for both NO$_x$ and O$_3$, the mean value of the gain in life expectancy for NO$_x$, which is ten months, should be summed up with the mean value of the loss of life expectancy for O$_3$, which is one month, resulting in a net gain of 9 months. The very small corresponding gain in life expectancy of 0.2 months, associated with exposure to PM$_{10}$, can also be taken into account. However, the relatively weak trends associated with PM$_{10}$, with a significance level of 0.01 < p < 0.05 for Stockholm and Gothenburg, and a corresponding level of p > 0.05 for Malmo, cannot be compared with the trends for the others pollutants, which in all cases exhibit significance levels of p < 0.001.

### 4.4 Uncertainties associated with the measurements and the relative risks

In general, the results are highly sensitive to the relative risks obtained from previous epidemiological studies. For NO$_x$, we use the RR 1.06 (95 % CI 1.03–1.09) per 10 µg m$^{-3}$ increase, based on a recent cohort study in Gothenburg (Stockfelt et al., 2015). Considering the similarities between Gothenburg, Stockholm, and Malmo regarding climate, vehicle fleet, and type of city, this RR may be representative for all these three Swedish cities. The other option could have been to choose the RR 1.08
(95 % CI 1.06–1.11), which comes from on a previous long-term cohort study in Oslo (Nafstad et al., 2004), but we decided to implement the RR from the Swedish study, with exposure levels closer to the actual situation. For NO₂, we use the pooled RR estimate of 1.066 (95 % CI 1.029–1.104) per 10 µg m⁻³ increase, from a meta-analysis (Faustini et al., 2014). The other option could have been to choose the RR 1.05 (95 % CI 1.03–1.08) from Hoek et al. (2013), which is also based on a meta-analysis from several studies in different countries and continents, but we considered the study group in Faustini et al. (2014) as more relevant for the Swedish situation, since it is based only on European studies. A RR for NO₂ based on European conditions, has also been calculated in the ESCAPE study (Beelen et al., 2014). In this study, the results for NO₂ were based on a limited passive sampling campaign for each cohort, and differ from the consistent results in the two meta-analyses (Faustini et al., 2014 and Hoek et al., 2013), where the studies with active monitoring were also included. In addition, Beelen et al. (2014) did not find any significant association with PM₂.₅ absorption, possibly indicating problems to reflect exposure to vehicle-exhaust. Because of this, we consider the RR in Faustini et al. (2014) as more relevant in comparison with the RR in Beelen et al. (2014).

Due to the influence of photochemistry, NOₓ is a better indicator of health risks associated with vehicle exhaust emissions than NO₂. This might explain the differences in relative risks for NOₓ and NO₂. For the same reason (photochemistry), the trends in NO₂ are to some extent influenced by the trends in O₃.

For O₃ exposure, the relative risk of 1.01 (95 % CI 1.005–1.02) per 10 µg m⁻³ increase, that has been used in our calculations, is based on Turner et al. (2016), which is a large prospective long-term cohort study performed in the U.S. The other option could have been to choose the RR 1.014 (95 % CI 1.005–1.024) from Jerret et al. (2009), which is very close to the RR in Turner et al. 2016, but considering the time periods during which the studies have been conducted, Turner et al. (2016) is more in line with the trend analysis period in this study. The number of epidemiological studies focusing on long-term exposure to O₃ are relatively few, and no meta-analyses are available. This means that the RR values associated with O₃ exposure are based on relatively less amount of data compared to the RR values for NO₂ and PM₁₀, which are obtained from meta-analyses.

For PM₁₀, we use the RR value of 1.04 (95 % CI 1.00–1.09) per 10 µg m⁻³ increase from Beelen et al. (2014), where the meta-analysis in ESCAPE is based on measurement campaigns with Harvard impactors, which is a meta-analysis based on 22 European cohorts. An important issue to consider in these Swedish cities is that the mass of PM₁₀ to a relatively small percentage consists of combustion-related particles. In the Swedish cities, non-exhaust traffic generated particles are the largest contributors to the mass of PM₁₀, which is clearly shown for Stockholm by Johansson et al. (2007), and for Gothenburg by Grundström et al. (2015). In Beelen et al. (2014), there is also a RR value of 1.04 (95 % CI 0.98–1.10) per 10 µg m⁻³ increase associated with exposure to PM₁₀. This value is very similar to the corresponding value for PM₁₀ mentioned above. Since PM₁₀ in the Swedish cities to a large extent consists of mechanically generated particles in the coarse fraction, this gives additional support for the suitability of the RR that is applied in the calculations for change in life expectancy.

This means that NOₓ and NO₂, which both primarily indicate locally produced combustion-related traffic emissions, are in Sweden clearly separated from PM₁₀, which rather indicates non-exhaust emissions. From a health perspective, PM₁₀ is a reflective indicator of the effects of exposure to the coarse fraction of urban aerosols. The health effects related to exposure to
the coarse fraction of particles differ from exposure to the fine (combustion related) fraction. Previous studies indicate an association between exposure to the coarse fraction of PM₁₀ and respiratory admissions, such as asthma and chronic obstructive pulmonary disease, while cardiovascular diseases are more closely linked to exposure to the fine fraction (Brunekreef and Forsberg, 2005).

4.5 Correlations between the different pollutants

Calculating the health effects associated with exposure to these different pollutants may be problematic, considering that exposures can occur simultaneously. When conducting health-impacts assessments, it is important to avoid double calculations when gain or loss in life expectancy are calculated as a result of decreasing or increasing trends. NOₓ and NO₂ are highly correlated, and the health impacts assessments, calculated as a gain in life expectancy, cannot be summed up, but they can rather be considered as different indicators of air pollutants, mainly originated from combustion processes. On the other hand, for these three Swedish cities, the correlation between NOₓ or NO₂ and PM₁₀ is very low, since the mass proportion of exhaust particles in PM₁₀ is very small (Johansson et al., 2007), and the health impacts associated with exposure to NOₓ and PM₁₀, respectively, can therefore largely be assumed to be independent of each other. Moreover, when considering simultaneous exposure to NO₂ and PM₂.₅, the relative risks associated with these pollutants are not significantly affected when included in a two-pollutant model (Faustini et al., 2014).

The relative risks associated with O₃ exposure are not significantly affected when they are included in two-pollutant models. The increased risks of circulatory and respiratory mortality associated with exposure to O₃ remained stable after adjustment for PM₂.₅ and NOₓ (Turner et al., 2016). Consequently, the health effects associated with exposure to O₃ can be assumed to be separated from the health effects from the other three pollutants, and the change in life expectancy can therefore be added or subtracted from the change in life expectancy associated with NOₓ, NO₂, and PM₁₀.

4.6 The differences between urban-background concentrations and population-weighted exposure concentrations

As shown in Section 2.5 and Figure A1–A4, the average population-weighted exposure concentrations may differ substantially from the concentrations measured at the measuring stations. For NOₓ and PM₁₀, the average exposure concentrations for the urban population are lower compared to the measured urban-background concentrations. The reason for this is that the people who live in the outskirts of the urban area are exposed to lower concentrations compared to those living close to the measuring station in the central parts of the cities, where both exhaust emissions and the formation of mechanically generated particles are most apparent. For O₃, the average exposure concentrations are higher compared to the measured urban-background concentrations. This can be explained by the interaction between NOₓ and O₃, where the O₃ reaction with NO dominates over the formation of new O₃, and this reaction is more apparent in the central part of a city, where the NO concentration is relatively higher. This explains the higher average population-exposure to O₃ compared to the measured urban-background concentrations at a central monitoring station.
The population distribution within the cities are not constant over time, and the meteorological conditions may also vary over time. Consequently, to calculate a corresponding change in population-weighted exposure, based on an air pollution trend during 25 years, is connected with uncertainties. Linear regression between the concentrations at the measuring stations, and the corresponding population-weighted exposure concentrations for the available years (Fig. A1–A4), are the best estimates that can be performed. A disadvantage is the limited number of years of calculated relationships, but where the $R^2$-values in the range of 0.7–0.9 may be considered as good.

Due to lack of data, the relations between urban-background concentrations and population-weighted exposure concentrations have not been possible to obtain for NO$_2$ in Stockholm, and not for NO$_2$, O$_3$ and PM$_{10}$ in Gothenburg, and not for any of the pollutants in Malmo. Without these relations, the population-weighted exposure concentrations have to be based on the data obtained directly from the measuring stations. However, based on the already established relations (Fig. A1–A4), it can be assumed that the life expectancy values associated with NO$_x$ and NO$_2$ exposure, but without adjustment for population-weighted exposure, are overestimated, while the opposite applies to O$_3$ exposure.

4.7 Policy implications

From a policy point of view, it is important to know the health impacts of local air pollution emissions (i.e. the primary emissions in the city from e.g. wood burning and road traffic), versus long-range transport of non-local origin. The low rural compared to the urban background concentrations outside the cities shows that the trends in local primary emissions of NO$_x$ and NO$_2$ are most important for the decreasing trends, and thereby also most important for the increase in life expectancy of the populations. This implies that policies addressing local emissions in the cities are important in order to improve the public health. NO$_x$ may be regarded as an indicator of vehicle exhaust emissions, which consist of many toxic compounds (Künzli et al., 2000). For PM$_{10}$, long-range transport of mainly secondary PM is more important than local emissions. However, primary emissions may still be more important for the health effects associated with PM, as clearly demonstrated by Segersson et al. (2017). They calculated the sectorial contributions to PM$_{10}$, PM$_{2.5}$ and black carbon (BC) exposure, and the related premature mortalities in three Swedish cities, including Gothenburg and Stockholm, by using dispersion models with high spatial resolution. They showed that although the main part of the PM$_{10}$ and PM$_{2.5}$ exposures were due to long-range-transport, the major part of the premature deaths were related to local emissions, with road traffic (dominated by non-exhaust wear emissions) and residential wood combustion having the largest impacts. For black carbon, the emissions from local road traffic (exhaust emissions) and residential wood combustion were more important both for exposures and health risks, as compared to long-range transport. This means that local policies aiming at reducing emissions from road traffic, both non-exhaust and exhaust emissions, and residential wood combustion, will be most important for reducing health risks associated with urban air pollution in these cities.

The large impact on life expectancy, which both NO$_x$ and NO$_2$ represent, makes these pollutants important to consider. NO$_x$ and NO$_2$ are both indicators of combustion-related air pollutants, as indicated by the high correlation between NO$_x$ and particle number concentration at kerbside sites (Johansson et al., 2007; Gidhagen et al., 2004), and they may therefore be good
indicators of population exposure to exhaust emissions from road traffic. Additionally, the change in life expectancy can be regarded as the tip of the iceberg, since reductions in NO\textsubscript{x} concentrations are also expected to cause reductions in cardiovascular and respiratory morbidity (Johansson et al., 2009).

Our study also shows that although the reduced NO\textsubscript{x} emissions have caused increased ozone concentrations, which contribute to adverse health effects, the net effect will be a longer life expectancy in the population (Table 2). This may not be true if other health outcomes are analysed. Our focus on increased mortality should be seen as a limitation and underestimation of the total impact. The regulated air pollutants have been associated not only with respiratory and cardiovascular morbidity and mortality, but more recently also with systemic effects and metabolic diseases, pregnancy and developmental outcomes, as well as CNS and psychiatric effects (Thurston et al., 2017). Many of these effects are poorly represented by the studies of mortality.

Health impact assessments from other studies clearly suggest that public health could largely benefit from better air quality (e.g. Künzli, 2002). However, estimates of health benefits of reducing air pollution are dependent on the indicator that is used, and on the shape of the concentration-response functions (Pope et al., 2015). In addition, Pope et al. (2015) also discusses recent evidence that the shape of the concentration-response function may be supra linear (dose-response relationship with a negative second derivative) across wide ranges of exposure, as has been shown in the case of exposure to PM\textsubscript{2.5}. It means that incremental pollution-abatement efforts may yield greater benefit in relatively clean areas compared to highly polluted areas. This motivates actions to be taken even in areas with relatively low levels of air pollutants.

5. Conclusions

The air pollution trends regarding NO\textsubscript{x}, NO\textsubscript{2}, and O\textsubscript{3} in Stockholm, Gothenburg, and Malmo exhibit significant (95 % CI) tendencies during the measured time periods. For PM\textsubscript{10}, the trends in Stockholm and Gothenburg are significant, but the trend in Malmo is non-significant. The slopes of the trends are very different, where the NO\textsubscript{x} and NO\textsubscript{2} concentrations in all cities exhibit decreasing trends, while O\textsubscript{3} in all cities exhibits increasing trends. For PM\textsubscript{10}, the trends are less clear with a decreasing trend in Stockholm, an increasing trend in Gothenburg, and no significant trend in Malmo. When the trends are divided into weekdays and weekends, it follows that the trends for NO\textsubscript{x} and NO\textsubscript{2} in all cities exhibit large differences, where the trends associated with weekdays exhibit sharper declines compared to the weekends. This phenomenon is consistent with the fact that local emission reductions mostly explain those declines, since the traffic intensity is more prominent during weekdays. An anti-correlation between NO\textsubscript{x} and O\textsubscript{3} can be seen in all cities, which can be attributed to an increased or decreased titration effect, where NO oxidizes to NO\textsubscript{2} has the capability of scavenging free oxygen atoms wherein NO\textsubscript{2} is formed. When the trends in this article are compared to the trends in other studies, the trends associated with NO\textsubscript{x} and NO\textsubscript{2} are in the same magnitude as those measured in the U.S. and Europe as a whole, but the trends associated with PM\textsubscript{10} are completely different, mostly caused by the large contribution of mechanically generated particles in the Swedish cities.

The change in life expectancy, associated with the air pollution trends, is most obvious for NO\textsubscript{x} and NO\textsubscript{2}, where up to about 20 % of the total increase in life expectancy during the period 1990–2015 can be attributed to decreasing NO\textsubscript{x} trends. Since
NO₂ and NOₓ are indicators of combustion-related air pollutants in general, an overall conclusion is that exposure to these air pollutants are particularly important in terms of health effects and health benefits. Our study indicates that policies aiming at reducing local emissions from road traffic have been most important for the increased life expectancy associated with urban air pollution in these three Swedish cities.

6. Acknowledgements

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7. Competing interests

The authors declare that they have no conflict of interest.

References


Appendix A

Table A1: General facts about the three cities regarding population structure, life expectancy at birth, and baseline mortality in terms of the number of deaths per 100 000 inhabitants. The number of deaths per 100 000 inhabitants are age standardized according to the average population for the year 2000.

<table>
<thead>
<tr>
<th>Stockhol</th>
<th>Gothenburg</th>
<th>Malmo</th>
</tr>
</thead>
<tbody>
<tr>
<td>Population size</td>
<td>1 040 907</td>
<td>1 515 017</td>
</tr>
<tr>
<td>Density of population (inhabitants per km²)</td>
<td>3 601</td>
<td>4 935</td>
</tr>
<tr>
<td>Expected life expectancy at birth</td>
<td>76.8</td>
<td>82.6</td>
</tr>
<tr>
<td>----------------------------------</td>
<td>------</td>
<td>------</td>
</tr>
<tr>
<td>Baseline mortality (all causes, age standardized), number per 100 000 inhabitants.</td>
<td>1 157*</td>
<td>833</td>
</tr>
</tbody>
</table>

* Extrapolated from a linear regression based on data from 1997–2015

Figure A1. Relationship between measured urban-background concentrations and population-weighted exposure concentrations of NO\textsubscript{x} in Stockholm. Slope and intercept are specified with 95% confidence intervals.
Figure A2. Relationship between measured urban-background concentrations and population-weighted exposure concentrations of NOx in Gothenburg. Slope and intercept are specified with 95% confidence intervals.

\[ y = (0.59\pm0.69) \times + (2.04\pm30.66) \]
\[ R^2 = 0.71 \]

Figure A3. Relationship between measured urban-background concentrations and population-weighted exposure concentrations of O3 in Stockholm. Slope and intercept are specified with 95% confidence intervals.

\[ y = (1.17\pm0.50) \times - (6.83\pm25.70) \]
\[ R^2 = 0.82 \]
Figure A4. Relationship between measured urban-background concentrations and population-weighted exposure concentrations of PM$_{10}$ in Stockholm. Slope and intercept are specified with 95% confidence intervals.