Air quality simulations for London using a coupled regional-to-local modelling system

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Abstract. A coupled regional-to-local modelling system comprising a regional chemistry–climate model with 5 km horizontal resolution (EMEP4UK) and an urban dispersion and chemistry model with explicit road source emissions (ADMS-Urban) has been used to simulate air quality in 2012 across London. The study makes use of emission factors for NOx and NO2 and non-exhaust emission rates of PM₁₀ and PM₂.₅ which have been adjusted compared to standard factors to reflect real-world emissions, with increases in total emissions of around 30 % for these species. The performance of the coupled model and each of the two component models is assessed against measurements from background and near-road sites in London using a range of metrics concerning annual averages, high hourly average concentrations and diurnal cycles. The regional model shows good performance compared to measurements for background sites for these metrics, but under-predicts concentrations of all pollutants except O₃ at near-road sites due to the low resolution of input emissions and calculations. The coupled model shows good performance at both background and near-road sites, which is broadly comparable with that of the urban model that uses measured concentrations as regional background, except for PM₂.₅ where the under-prediction of the regional model causes the coupled model to also under-predict concentrations. Using the coupled model, it is estimated that 13 % of the area of London exceeded the EU limit value of 40 µg m⁻³ for annual average NO₂ in 2012, whilst areas of exceedances of the annual average limit values of 40 and 25 µg m⁻³ for PM₁₀ and PM₂.₅ respectively were negligible.

1 Introduction

Poor air quality has long been recognized as having adverse effects on health. Particulate pollution in the UK has been assessed as causing a loss of life expectancy from birth of approximately 6 months (COMEAP, 2010), while air pollution in the WHO European Region was estimated to cause 600 000 premature deaths in 2010 (WHO, 2015). Improved understanding of these health effects requires additional information about air quality, especially in urban areas where high pollutant concentrations coincide with high population densities.

Continuous air quality measurements, for example from the UK Automatic Urban and Rural Network (AURN, Defra, 2017), are typically carried out at a limited number of fixed locations in an urban area and are expected to be representative of several square kilometres for urban background locations (EC Directive, 2008). In addition, short-term intensive campaigns making use of specialist monitoring equipment, as for example carried out for the ClearfLo project (Clean Air for London; Bohnenstengel et al., 2015), are of great value for detailed assessment of model performance and underlying processes, whilst sampling equipment can also be carried by moving vehicles or individuals for short-term detailed studies. In contrast to measurements, air quality or atmospheric chemistry transport models, evaluated with the above data, allow pollutant concentrations to be simulated with complete spatial-temporal coverage leading to detailed calculations of population exposure (Smith et al., 2016).

Air quality models require accurate input emissions data to make reliable predictions of ambient concentrations. How-
ever, in the last decade, it has become clear that measured NO\textsubscript{x} and NO\textsubscript{2} concentrations have not decreased as fast as would have been anticipated from published emission factors (Carslaw et al., 2011). Several measurement techniques for direct assessment of on-road tailpipe emissions, as reported by Carslaw and Ryhs-Taylor (2013) and O’Driscoll et al. (2016), have confirmed differences from the official emissions estimates (EFT, Defra, 2016). In-service emissions performance evaluation of Euro 6/VI vehicles (Moody and Tate, 2017) indicated that, whilst Euro VI heavy-duty vehicles and Euro 6 petrol light-duty vehicles are performing broadly as predicted, Euro 6 diesel light-duty vehicles emit NO\textsubscript{2} at rates exceeding the published data, by factors of up to 4.5.

There is further uncertainty in the rates of particulate emissions from road vehicles due to wear of tyre, brake, and road surfaces and resuspension of pre-existing particulates (Thorpe and Harrison, 2008). Particulate exhaust emissions have decreased considerably in recent years, primarily due to the introduction of diesel particulate filters, so the relative contribution of non-exhaust PM\textsubscript{10} and PM\textsubscript{2.5} to total traffic emissions is now considerable, of the order of half of the total exhaust (Grigoratos and Martini, 2014).

Atmospheric chemistry models that simulate air quality vary in complexity in terms of the scales and processes represented. Global and regional models use gridded emissions data to calculate transport and chemistry over global or regional modelling domains, such as EMEP MSC-W (EMEP, Simpson et al., 2012) used in this study, CAMx (Ramboll Environment and Health, 2016), CMAQ (Byun and Schere, 2006) and WRF-Chem (Grell et al., 2005). Models on a smaller scale apply detailed transport and fast chemistry processes to individual sources, such as ADMS-Urban (Owen et al., 2000) also used in this study, the US EPA model AERMOD (Cimorelli et al., 2004) and CAR-FMI (Kukkonen et al., 2001).

Global and regional air quality models typically use detailed chemistry schemes whilst urban models typically only represent fast chemistry, such as O\textsubscript{3}–NO\textsubscript{x} chemistry which is relevant for pollution concentration gradients across urban areas. Some hybrid process–statistic-based approaches have also been developed, where measured concentration data are used to constrain modelled concentrations in order to reduce uncertainties, for example those described in Stedman et al. (2001) and Sokhi et al. (2008).

Urban dispersion models typically use measured upwind rural concentrations to represent long-range transport. This is a successful approach for modelling historic periods but has limited applicability for assessing future scenarios, including those related to climate change or the local effects of regional emissions changes. The use of a limited number of upwind monitoring sites can also make the upwind concentration data less representative and does not allow for variations over a large urban area. An alternative method (Stocker et al., 2012, 2014) is to combine regional modelling with urban local modelling in order to take into account both short-range and long-range transport and chemistry effects, whilst avoiding double counting of the gridded and explicit emissions. The balance between regional and local influences differs according to the pollutant lifetime. For example, concentrations of ozone and particulates, which have lifetimes at the surface of days to weeks, are strongly influenced by regional emissions and transport, whereas concentrations of NO\textsubscript{2}, with a surface lifetime of around 1 day, are primarily related to the dispersion and chemical transformation of local emissions.

The overall methodology for the detailed evaluation of air pollution concentrations across London for 2012 using a coupled regional–urban model is described in Sect. 2 with details of the measurement data, models, emissions and statistical parameters. Section 3 gives the results of the model evaluation against measured concentrations while Sect. 4 discusses the results in relation to air quality in London and the different modelling methods.

2 Methods

This paper presents a detailed evaluation of air pollution concentrations across London for 2012 using a coupled regional–urban model (described in Sect. 2.4), which comprises a regional version of the EMEP atmospheric chemistry transport model, EMEP4UK (Sect. 2.2), and the ADMS-Urban local dispersion and chemistry model (Sect. 2.3). The coupled model is evaluated alongside the stand-alone implementations of the two underlying models. The evaluation exercise compares hourly modelled concentrations of NO\textsubscript{x}, NO\textsubscript{2}, O\textsubscript{3}, CO, PM\textsubscript{10} and PM\textsubscript{2.5} with measured hourly concentrations from up to 42 automatic monitoring sites within Greater London for the year 2012, described in Sect. 2.1. The model simulations use road vehicle emissions of NO\textsubscript{x}, NO\textsubscript{2} and particulates which have been adjusted in line with real-world emissions measurements. The emissions data, including the raw 2012 emissions, adjustments and time variation, are described in Sect. 2.5. Definitions of the statistics used for model evaluation are given in Sect. 2.6.

2.1 Measurement data

The monitoring sites selected for the model evaluation were those from the London Air Quality Network (LAQN, Mittal et al., 2017) located within Greater London that had at least 70% data capture of hourly data during 2012 for PM\textsubscript{10}, PM\textsubscript{2.5}, CO or at least two of NO\textsubscript{x}, NO\textsubscript{2} and O\textsubscript{3}. A summary of site numbers by type and their average heights is given in Table 1; with the exception of two background monitors at 5.5 and 10 m all monitors are between 2 and 4 m above ground level (a.g.l.). The site locations of the NO\textsubscript{2} and O\textsubscript{3} monitors are presented in Fig. 1. Note that all the map plots in this paper adopt the polar stereographic coordinate system as used in EMEP4UK, with an approximately 30° anticlock-
Table 1. Number of monitoring sites per pollutant and site type, with average heights over all sites and by site type. “Background” includes both suburban and urban background sites while “near-road” includes sites classified as roadside and kerbside. Note that all the selected O₃ monitoring sites have co-located NO₂ and NOₓ measurements.

<table>
<thead>
<tr>
<th>Pollutant</th>
<th>Number of sites</th>
<th>Average site inlet height (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total</td>
<td>Background</td>
</tr>
<tr>
<td>NOₓ</td>
<td>42</td>
<td>15</td>
</tr>
<tr>
<td>NO₂</td>
<td>42</td>
<td>15</td>
</tr>
<tr>
<td>O₃</td>
<td>20</td>
<td>11</td>
</tr>
<tr>
<td>PM₁₀</td>
<td>33</td>
<td>12</td>
</tr>
<tr>
<td>PM₂.₅</td>
<td>11</td>
<td>5</td>
</tr>
<tr>
<td>CO</td>
<td>7</td>
<td>3</td>
</tr>
</tbody>
</table>

Figure 1. Locations of NO₂ (black) and O₃ (pale grey) monitoring sites in Greater London, with square symbols for background sites and round symbols for near-road sites. The London borough extents and boundaries are shown for reference, alongside the extent of the locally modelled emissions and the location of the measured meteorological data at Heathrow airport.

2.2 Regional-scale modelling: EMEP4UK

EMEP4UK is a nested regional application of the EMEP MSC-W (European Monitoring and Evaluation Programme Meteorological Synthesizing Centre-West) model, focused specifically on air quality in the UK. The main EMEP MSC-W model has been widely used for both scientific studies and for policymaking in Europe, with references to evaluation and application studies available in Simpson et al. (2012), Schulz et al. (2013) and at http://www.emep.int (last access: 1 August 2018). EMEP4UK is described in Vieno et al. (2010, 2014, 2016a, b); the version used here is based on EMEP MSC-W rv4.5. It uses one-way nesting from a 50 km × 50 km resolution greater European domain (standard EMEP domain) to an inner 5 km × 5 km domain which covers the British Isles and nearby parts of continental Europe, both in a polar stereographic projected coordinate system, as shown in Fig. 2. An intermediate 10 km × 10 km resolution domain is used for WRF (Weather Research and Forecasting Model) to ensure numerical stability, but is not required for the chemistry transport calculations (Vieno et al., 2010). Full details of the WRF model domains are given in Table A1. The model has 21 vertical levels extending from the surface to 100 hPa, with the lowest vertical layer 50 m thick, meaning that modelled surface concentrations represent a height of around 25 m. This is a finer vertical resolution than was available in the standard EMEP model rv4.5. Hourly average output concentrations are available from each cell for the full 2012 modelling period.

The gaseous chemical scheme used in EMEP4UK in this study is the CRI-v2-R5 mechanism (Watson et al., 2008), which has 220 species and 609 reactions. Five classes of fine and coarse particles, with differing size and deposition properties, are used in EMEP4UK (Simpson et al., 2012) along with the MARS (Model for an Aerosol Reacting System) equilibrium module for gas–aerosol partitioning of secondary inorganic aerosol (Binkowski and Shankar, 1995; Simpson et al., 2012) and a treatment of secondary organic aerosol formation using the volatility basis set approach (Bergström et al., 2012; Ots et al., 2016). Dry (including stomatal) and wet deposition of gases and particles are simulated. Sixteen land cover classes are used for dry deposition modelling and for the calculation of biogenic emissions. Ozone boundary conditions for the outer European domain are based on the approach in Simpson et al. (2012), scaling a monthly climatology with clean air measurements at Mace Head. Initial and boundary conditions of all other species for the European domain are specified as fixed functions of latitude and time of year.

The chemistry transport model was driven by meteorological output from the WRF version 3.6.1 (Skamarock et al., 2008; NCAR, 2008) including data assimilation of 6-hourly meteorology from the European Centre for Medium Range Weather Forecasting ERA-Interim reanalysis (Dee et al., 2011). The option of input meteorological data from WRF has been developed for EMEP4UK and is used in both the European and UK domains. The WRF configuration was
as follows: Purdue Lin for microphysics, Grell-3 for cumulus parameterization, Goddard shortwave for radiation physics and Yonsei University (YSU) for planetary boundary layer (PBL) height. Land use categories were based on the MODIS IGBP classification. This WRF configuration is similar to that discussed in Vieno et al. (2010), where it is shown to perform well in comparison with measurements. An evaluation of the WRF-EMEP4UK modelling system against measured gaseous and particulate pollutant concentrations across the UK for 2001 to 2010 is given by Lin et al. (2017), while Ots et al. (2016) compares WRF-EMEP4UK air quality simulations with detailed measurements of secondary organic aerosols made in London during the 2012 ClearLo campaign.

2.3 Urban-scale modelling: ADMS-Urban

The Atmospheric Dispersion Modelling System (ADMS, Carruthers et al., 1994) is a quasi-Gaussian plume air dispersion model able to simulate a wide range of passive and buoyant releases to the atmosphere. The dispersion calculations are driven by hourly meteorological profiles of wind speed and direction, among other parameters, which are characterized using Monin–Obukhov length similarity theory; meteorological input data may be derived from measurements or output from a mesoscale model such as WRF. ADMS is a local dispersion model, which is able to resolve concentration gradients that occur in the vicinity of a range of emission source types, including point, jet, line, area and volume sources. The modelling of dispersion and chemistry for source emissions is independent of the output grid resolution.

The ADMS-Urban model has been used to simulate air quality within cities worldwide; applications include testing of emission-reduction scenarios and forecasting (Stidworthy et al., 2017). Emissions from all sources within the model domain are included, either explicitly with detailed time-varying profiles, for instance major road and industrial sources, or as grid-averaged emissions, representing diffuse sources such as those from heating and minor roads as a grid of regular volume sources, with simpler time variation.

The flow field that drives dispersion of pollutants within an urban area is inhomogeneous. On the neighbourhood scale, buildings displace the upwind wind speed profile and reduce in-canopy wind speeds. ADMS-Urban has an “urban canopy” flow field module, which calculates wind speed and turbulence flow profiles that relate to the spatial variation of the surface roughness length, $z_0$. Locally, if street canyons are formed by densely packed tall buildings, it is important to model the complex combination of recirculating and channelled flows. The ADMS-Urban advanced street canyon module is able to model (i) the channelling of flow along and circulation of flow across a street canyon, (ii) asymmetric street canyons, (iii) the effect of pavements within a canyon, and (iv) the effect of a street canyon on the surrounding area. The module has been validated extensively by comparison with measurements from monitoring networks in Hong Kong and London (Hood et al., 2014). In the present modelling the
advanced street canyon option has been used for all roads in the modelling domain with adjacent buildings.

For this project 3-D building data and road centreline locations from Ordnance Survey MasterMap (Ordnance Survey, 2014) were processed for use in ADMS-Urban, as described in Jackson et al. (2016), although using the EMEP4UK polar stereographic projected coordinate system. The inputs to ADMS-Urban take two forms: (i) gridded building height and density parameters for urban canopy flow field calculations and (ii) street canyon properties for each side of explicitly modelled road sources. Figure 3 shows the variation of average building height over the Greater London area, which is used to determine the local roughness length for flow calculations.

In this study ADMS-Urban version 4.0.4 was used for the stand-alone runs, with emissions covering the Greater London area, defined by the London Atmospheric Emissions Inventory (LAEI) emissions extent. The stand-alone runs use hourly measured meteorological data from Heathrow airport (location shown in Fig. 1) for the whole domain, with valid data available for over 95% of hours in 2012. Long-range transport of NO, NO$_2$, O$_3$, PM$_{10}$, PM$_{2.5}$ and SO$_2$ is represented by hourly measured background concentrations from rural sites upwind of London in the AURN monitoring network (Ricardo-AEA, 2013): Wicken Fen (north of London, gaseous pollutants only); Lullingston Heath (south, gaseous pollutants only); Harwell (west, all pollutants); and Rochester (east, all pollutants), providing boundary conditions to the local modelling. No monitoring data are available for CO, so a constant background concentration for CO of 220 µg m$^{-3}$ was obtained from the annual mean background map published by Defra (2013b). Output concentrations are given at hourly resolution and post-processed to calculate long-term averages as required. The modelling also uses the ADMS-Urban NO$_x$ photolytic chemistry module, which accounts for fast, near-road oxidation of NO by O$_3$ to form NO$_2$ (Smith et al., 2017), and simple sulphate chemistry for conversion of SO$_2$ to PM$_{2.5}$ and PM$_{10}$. The options for flow over complex terrain (Carruthers et al., 2011), and gaseous and particulate wet and dry deposition, are not used in this study.

### 2.4 Coupled regional-to-urban-scale model

At short times after release of a pollutant from a source, concentration gradients due to releases from that source are high and a street-scale resolution model such as ADMS-Urban is needed to capture the fine details of dispersion and fast chemistry, for instance at roadside locations. At longer times after release, pollutant concentration gradients are reduced by mixing and a gridded regional model that accounts for long-range transport and detailed chemical transformations simulates these processes adequately. These models may be coupled within a single system. However, the computational linkage process is non-trivial in order to avoid double counting of emissions and to ensure that the chemical processes are accounted for at all timescales.

The underlying concept for coupling the regional- and urban-scale models, described in Stocker et al. (2012), is to use the local urban model to represent the initial dispersion of emissions up to a mixing time, typically 1 h, after which the emissions are considered well mixed on the scale of the regional model grid. In general, Gaussian plume models such as ADMS-Urban treat plumes as continuing for all time, but within the coupled system the calculations are truncated at the mixing time $\Delta t$. An ADMS-Urban run with gridded emissions, limited to the mixing time ($C_{\text{grid}}(\Delta t)$), is used to represent the regional model calculations within the mixing time and is subtracted from the regional model output, $C_{\text{RM}}$, in order to avoid double counting emissions. The final concentrations from the coupled model, $C_{\text{coupled}}$, are then calculated by adding the output from an ADMS-Urban run with explicit emissions, also limited to the mixing time ($C_{\text{expl}}(\Delta t)$), with the overall expression given as

$$C_{\text{coupled}} = C_{\text{RM}} - C_{\text{grid}}(\Delta t) + C_{\text{expl}}(\Delta t).$$ (1)

Additional steps calculate local background concentrations from the regional model which are used as input to the subsequent ADMS-Urban runs to ensure that the long-range transport and chemical environment is adequately represented for local chemical processes.

The initial implementation of an automated coupled system using ADMS-Urban and the CAMx regional model for Hong Kong is described in Stocker et al. (2014), with evaluation against monitoring data. For the work described in this paper, the coupled ADMS-Urban Regional Model Link (RML) system was further developed to allow the ADMS-Urban runs to be carried out using the ARCHER UK National Supercomputing Service. In the coupled system, meteorology and background concentrations are extracted and used as inputs for separate ADMS-Urban runs for each 5 km $\times$ 5 km EMEP4UK grid cell, leading to spatially vary-
ing meteorology and background concentrations across the 
modelling domain.

The version of ADMS-Urban (3.4.6) used within the cou-
ped system was slightly older than for the stand-alone runs 
(4.0.4); the older version was modified for compatibility with 
the ARCHER supercomputer. There are no relevant differ-
ences in terms of dispersion or chemical modelling between 
ADMS-Urban versions 3.4.6 and 4.0.4.

2.5 Emissions data

2.5.1 2012 emissions

EMEP4UK uses anthropogenic emissions of NO\textsubscript{x}, NH\textsubscript{3}, 
SO\textsubscript{2}, primary PM\textsubscript{2.5}, primary coarse PM (PM\textsubscript{2.5–10}), CO 
and non-methane VOCs (volatile organic compounds). Emis-
sions from the UK are derived from the National Atmo-
spheric Emission Inventory (NAEI, Tsagatakis et al., 2016) 
for 2012 at 1 km resolution and aggregated to the model’s 
5 km x 5 km grid. Within Greater London these NAEI emis-
sions are replaced by the emissions prepared for ADMS-
Urban as described below. Outside the UK, EMEP4UK uses 
2012 anthropogenic emissions provided by the EMEP Centre 
at/; last access: 1 August 2018) at 50 km resolution. Ship-
ping emission estimates for seas around the UK are derived 
from ENTEC (2010), projected to 2012. The anthropogenic 
emissions are distributed vertically within the model accord-
ing to their Selected Nomenclature for sources of Air Pollu-
tion (SNAP) sector; for example road transport emissions 
(sector 7) are assigned to the lowest layer while power sta-
tion emissions (sector 1) are assigned to layers between 184 
and 1106 m (Simpson et al., 2012, Supplement). Biogenic 
emissions of isoprenes and monoterpene are calculated at 
each time step according to insolation and surface tempera-
ture (Guenther et al., 1995). Emissions of wind-driven sea 
salt and NO\textsubscript{x} from soils are also calculated interac-
tively as described by Simpson et al. (2012), whereas lightning NO\textsubscript{x} 
emissions are prescribed. Import of Saharan dust is treated 
using a monthly dust climatology as a model boundary con-
dition. Resuspension of settled dust by wind is not included.

For ADMS-Urban the emissions for all sources except 
road traffic have been taken from the London Atmospheric 
Emissions Inventory 2010 (GLA, 2013) for the LAEI do-
main, which covers the area bounded by the M25 orbital motor-
way. The emissions have been projected from the LAEI 
base year 2010 to the modelling year 2012. Road traffic 
emissions have been calculated using activity data from the 
LAEI. The emission factors used to calculate emission rates 
are based on the UK NAEI 2014, which includes speed-
emissions data from the COPERT 4 version 10 software tool 
(Katsis et al., 2012). However, due to uncertainties in NO\textsubscript{x} 
emissions factors for some diesel vehicles and non-exhaust 
particulate emission factors, adjustments have been made to 
the published factors to improve consistency with real-world 
emissions measurements. The adjustments are discussed fur-
ther in Sect. 2.5.2 and their effects on the modelled concen-
trations examined in Sect. 3.1.

The NAEI and LAEI emissions are supplied as a regular, 
orthogonal 1 km resolution grid in the OSGB coordinate sys-
tem. The use of the EMEP4UK model in this study requires 
a conversion to the polar stereographic coordinate system, 
with reaggregation onto a grid with a different orientation. 
This causes some loss of precision in the location of emis-
sions, which is more acute for the ADMS-Urban runs with 
1 km gridded emissions than for the EMEP4UK runs with 
5 km grid resolution. The average 1 km gridded emissions 
are reduced by around 5% as a result of the re-gridding pro-
cess. Within the coupled modelling system, this reduction of 
average emissions makes little difference as it only affects 
the ADMS-Urban run including explicit sources, where con-
centrations are dominated by the unaffected explicit emis-
sions due to running a limited spatial extent. The change is 
also small relative to the real-world adjustments and other 
sources of uncertainty in the emissions. For consistency, the 
stand-alone local model runs have used the same coordinate 
system as EMEP4UK and the coupled system in this study.

In ADMS-Urban the road source emissions are modelled 
with a standard initial mixing height of 2 m a.g.l., although 
they may be distributed further upwards due to street canyon 
effects. Aggregated emissions, represented as gridded vol-
ume sources, have a depth of 100 m in the runs without 
explicit sources, in order to match the behaviour of the 
EMEP4UK modelling, and a depth of 10 m in the run with 
explicit sources, where individual point sources are modelled 
with release heights of 30–200 m.

2.5.2 Road traffic emissions factor adjustments

A significant cause of the discrepancies in NO\textsubscript{x} and NO\textsubscript{2} 
emission rates between published figures and real-world 
measurements is the difference in driving conditions between 
standard test cycles and real journeys, especially those in 
congested urban traffic (Franco et al., 2013). This issue was 
highlighted in 2015 when it became apparent that Volkswa-
gen had installed software in their diesel cars that automati-
cally reconfigured the engine during emissions testing (Old-
enkamp et al., 2016). The discrepancies in European ve-
hicle emission rates are expected to begin to decrease due 
to recent legislative changes (Commission Regulation (EU) 
2016/646) which require the use of urban driving cycles and 
real-world assessment for emissions testing. Emission fac-
tor adjustments are still likely to be necessary for modelling 
older vehicles which will remain in the active fleet.

Measured volume ratios of NO\textsubscript{x} and NO\textsubscript{2} to CO\textsubscript{2} emis-
sions (a proxy for fuel usage) have been compiled for a 
range of vehicles, categorized by Euro emission standard 
and size, with corresponding speeds by Carslaw and Rhys-
Tyler (2013). Measurements were taken at four sites, repre-
senting roads in central and outer London. Additional data
from bus monitoring campaigns are provided in Carslaw and Priestman (2015) and used for buses running with compressed natural gas fuel. For this study, to make use of these measured data to improve road traffic emissions, the emissions inventory toolkit (EMIT, CERC, 2015) software was used to calculate standard ratios of NO\textsubscript{x} to CO\textsubscript{2} emissions from the raw NAEI data set for different vehicle types and Euro classes, for average speeds as available in the measured data. A total of 22 vehicle categories were used for light vehicles and 17 categories for heavy vehicles, with scaling factors calculated from the measured data ranging from 0.80 for Euro II buses to 3.32 for Euro IV buses with scaling factors calculated from the measured data. A total of 22 vehicle categories were used for Euro classes, for average speeds as available in the measured data. The following statistics are used to evaluate the modelled concentrations \( M \) in relation to the observed concentrations.

The adjusted emissions that reflect real-world conditions as well as possible are hereafter referred to as real-world emissions. The total emissions for the LAEI area are summarized by sector in Table 2, including the effects of the adjustments to road transport emissions. Note that CO emissions are unaffected by the road traffic adjustments. A graphical representation of the emissions used in ADMS-Urban is shown in Fig. 4.

2.5.3 Time variation of emissions

In addition to annual average emission rates, it is important for models to capture the temporal variation of emissions in order to represent the short-term variation of concentrations. Within EMEP4UK, the 2012 annual total anthropogenic emissions derived from the inventories are resolved to hourly resolution using prescribed monthly, day-of-week and diurnal hourly emissions factors (the latter differing between weekdays, Saturdays and Sundays) for each pollutant and for each of the SNAP sectors (Simpson et al., 2012).

The stand-alone local model implementation uses an hourly time-varying profile for weekdays, Saturdays and Sundays for all explicit road sources and for aggregated emissions. This time-varying profile is based on a long-term analysis of NO\textsubscript{x} measurements in central London (Beever et al., 2009). The ADMS-Urban runs with gridded emissions within the coupled system use a simplified version of the EMEP4UK monthly and hourly time-varying profiles for NO\textsubscript{x} and PM, combined using a weighting by total emissions for each sector, while runs with explicit emissions use the same profile as in the stand-alone implementation for explicit road sources.

2.6 Model evaluation statistics

The non-exhaust particulate components are relatively unrefined, for example the EMEP CORINAIR non-exhaust factors use a linear speed-emissions profile and a maximum of 10 vehicle categories, in contrast to the hundreds of vehicle categories used for exhaust emissions classification. Analyses of roadside measurements demonstrate that the contribution from brake wear in particular is considerably higher than the published factors (GLA, 2016).

Non-exhaust particulate emission factors were adjusted based on work by Harrison et al. (2012), who analysed measurements of speciated and size-segregated particulates at the Marylebone Road monitoring site and nearby urban background sites, made during four month-long campaigns between 2007 and 2011. Non-exhaust emissions were found to contribute 77% of the total traffic-related particulate emissions, with 55% of the non-exhaust attributable to brake wear and smaller proportions from resuspension of road dust and tyre wear. Assuming that the standard exhaust emission factors are reliable, the non-exhaust emission factors were scaled in EMIT in order to make up 77% of the total traffic emissions and to have the correct proportionality between the different components. This is consistent with the approach taken in the LAEI 2013 (GLA, 2016). Applying these adjustments increases the total annual PM\textsubscript{10} emissions from road traffic sources by 45%. Basing the adjustment of all road non-exhaust emissions on measurements from one site is an approximation, but it is still expected to improve the overall estimates of non-exhaust emissions due to the substantial uncertainty in the standard factors.
Table 2. Total emission rates within the LAEI area by SNAP sector, including the effects of real-world emission adjustments for NO\textsubscript{x} and PM from road transport (sector 7). Italic font indicates the road transport and total emissions with unadjusted (raw) emission factors.

<table>
<thead>
<tr>
<th>SNAP sector</th>
<th>Description</th>
<th>Total emission rate (Mg yr\textsuperscript{-1})</th>
<th>NO\textsubscript{x}</th>
<th>NO\textsubscript{2}</th>
<th>PM\textsubscript{10}</th>
<th>PM\textsubscript{2.5}</th>
<th>CO</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Energy production</td>
<td>7886</td>
<td>394</td>
<td>307</td>
<td>0</td>
<td>918</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Domestic and commercial combustion</td>
<td>3887</td>
<td>194</td>
<td>99</td>
<td>60</td>
<td>5428</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Industrial combustion</td>
<td>4796</td>
<td>240</td>
<td>192</td>
<td>115</td>
<td>3367</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>Production processes</td>
<td>948</td>
<td>47</td>
<td>227</td>
<td>22</td>
<td>435</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Fossil fuel extraction and distribution</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>Solvent use</td>
<td>0</td>
<td>0</td>
<td>28</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>Road transport (raw)</td>
<td>32</td>
<td>147</td>
<td>8520</td>
<td>2724</td>
<td>1420</td>
<td>48738</td>
</tr>
<tr>
<td>7</td>
<td>Road transport (adjusted)</td>
<td>49</td>
<td>673</td>
<td>11878</td>
<td>3943</td>
<td>1916</td>
<td>48738</td>
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<td>8</td>
<td>Other transport</td>
<td>8439</td>
<td>587</td>
<td>197</td>
<td>150</td>
<td>29</td>
<td>173</td>
</tr>
<tr>
<td>9</td>
<td>Waste treatment and disposal</td>
<td>1647</td>
<td>82</td>
<td>168</td>
<td>150</td>
<td>508</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>Agriculture</td>
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<td>0</td>
<td>15</td>
<td>1</td>
<td>0</td>
<td></td>
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<tr>
<td>11</td>
<td>Nature</td>
<td>96</td>
<td>5</td>
<td>106</td>
<td>76</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Total with raw road transport</td>
<td>59845</td>
<td>10070</td>
<td>4063</td>
<td>1996</td>
<td>88</td>
<td>568</td>
<td></td>
</tr>
<tr>
<td>Total with adjusted road transport</td>
<td>77371</td>
<td>13429</td>
<td>5282</td>
<td>2491</td>
<td>88</td>
<td>568</td>
<td></td>
</tr>
<tr>
<td>Change in total due to adjustments (%)</td>
<td>29</td>
<td>33</td>
<td>30</td>
<td>25</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
</tbody>
</table>

\(O\); \(n\) is the number of pairs of modelled and observed concentrations, a bar indicates the mean value (e.g. \(\overline{M}\)) and a subscript indicates a single parameter value ranked between unity and \(n\) (e.g. \(M_i\)).

Fractional bias (Fb) is a measure of the mean difference between the modelled and observed concentrations:

\[
Fb = \frac{\overline{M} - \overline{O}}{0.5 \left( \overline{O} + \overline{M} \right)}. \tag{2}
\]

Normalized mean square error (NMSE) is a measure of the mean difference between matched pairs of modelled and observed concentrations:

\[
NMSE = \frac{(M - O)^2}{\overline{M}\overline{O}}. \tag{3}
\]

Pearson’s correlation coefficient (\(R\)) is a measure of the extent of a linear relationship between the modelled and observed concentrations:

\[
R = \frac{1}{n-1} \sum_{i=1}^{n} \left( \frac{M_i - \overline{M}}{\sigma_M} \right) \left( \frac{O_i - \overline{O}}{\sigma_O} \right). \tag{4}
\]

where \(\sigma_O\) is the standard deviation of observed concentrations and \(\sigma_M\) is the standard deviation of modelled concentrations.

Fraction of modelled hourly concentrations within a factor of 2 of observations (Fac2) is given by the fraction of model predictions that satisfy

\[
0.5 \leq \frac{M_i}{O_i} \leq 2.0. \tag{5}
\]

The model quality indicator (MQI, Thunis and Cuvelier, 2016) has been developed through the Forum for Air quality Modelling in Europe (FAIRMODE, Janssen et al., 2017) as an overall metric of model performance which depends on the measurement uncertainty. The MQI is defined as the ratio between the model bias and twice the measurement uncertainty (RMS\textsubscript{U}, scaled from the estimated measurement uncertainty at the relevant limit value); lower values reflect better model performance and values of the MQI less than 1 are considered to fulfill the modelling quality objective, in which case model bias is less than twice the measurement uncertainty. This statistic is not defined for NO\textsubscript{x} or CO as there are no EU limit values for NO\textsubscript{x}, whilst CO is typically well below the EU limit value, so it is not normally assessed. On the assessment target plot, the MQI represents the distance between the origin and a given station point as follows.

\[
MQI = \frac{RMSE}{2RMS_U}, \tag{6}
\]

where

\[
RMSE = \sqrt{\frac{1}{n} \sum_{i=1}^{n} (O_i - M_i)^2}, \tag{7}
\]

and the ordinate and abscissa correspond to the bias (\(\overline{M} - \overline{O}\)) and CRMSE (centred root mean square error):

\[
CRMSE = \sqrt{\frac{1}{n} \sum_{i=1}^{n} [(M_i - \overline{M}) - (O_i - \overline{O})]^2}, \tag{8}
\]

where both statistics are normalized by twice the measurement uncertainty.

The robust highest concentration (RHC) gives an indication of the performance of the model for high hourly concentrations and is defined as

\[
RHC = \chi(j) + (\chi - \chi(j)) \ln \left( \frac{3j - 1}{2} \right), \tag{9}
\]

where \(\chi\) is the number of pairs of modelled and observed concentrations.
where $j$ is the number of values considered as the upper end of the concentration distribution, $\chi$ is the average of the $j-1$ largest values and $\chi(j)$ is the $j$th largest value. The value of $n$ is set to 26, as used in Perry et al. (2005).

3 Results

Section 3.1 assesses the impact of the emissions adjustments on simulated concentrations using the stand-alone ADMS-Urban local model. Section 3.2 presents the spatial variation of annual average NO$_2$, O$_3$ and PM$_{2.5}$ concentrations across London predicted by the coupled modelling system while Sect. 3.3 gives detailed evaluation statistics for the regional, local and coupled models based on hourly concentration data for all modelled species. Section 3.4 presents additional analysis of the annual average modelled and measured concentrations while Sect. 3.5 concerns the hourly average concentrations and diurnal cycles for NO$_x$, NO$_2$ and O$_3$. The regulatory standards for NO$_2$, which are defined for annual average and maximum hourly concentrations, have driven this study’s focus on these two averaging periods.

### 3.1 Impact of emission adjustments on modelled concentrations

The effect of the adjustment of road traffic NO$_x$ and PM emissions to reflect real-world conditions on all simulated species is shown for background and near-road site types across London in Table 3. This comparison was performed as a preliminary assessment using simulations from the stand-alone ADMS-Urban local model since for this model measured background concentrations are utilized, leading to lower uncertainty in the long-range transport component of concentrations in the stand-alone model than in the coupled system, where the long-range transport contribution is also modelled. The reduced uncertainty means that model errors are the most closely associated with local emissions for this model. The statistics presented are fractional bias, normalized mean square error and correlation coefficient.

The CO concentration results show negligible changes due to the adjustment of emissions, as expected, since CO emissions were not changed. For NO$_x$, NO$_2$, O$_3$ and PM$_{10}$ the emission adjustments result in substantial concentration changes and improvements in Fb and NMSE, especially for near-road sites. For NO$_x$ the concentrations are increased, with Fb values reduced from around −0.3 to close to zero for near-road sites and NMSE reduced substantially; there are smaller concentration and statistics changes for NO$_2$. The change in NO$_x$ concentrations at background sites (+23%) is similar to the change in total emissions (+29%, Table 2), reflecting the direct link from emissions to concentrations for NO$_x$. The change in NO$_2$ concentrations at background sites (+18%) is smaller than both the NO$_2$ emissions change (33%) and the NO$_x$ concentration change, since emitted NO$_2$ contributes only a relatively small amount to total NO$_x$ and due to the time required for chemical processes to convert NO to NO$_2$, which means the response of NO$_2$ concentrations to NO$_x$ emissions is less than linear.

For O$_3$ the impact of the adjusted NO$_x$ and NO$_2$ emissions leads to lower concentrations and reduces the Fb from 0.16 to 0.02 for near-road sites, although there is little change in the NMSE. For PM$_{10}$ concentrations are higher, so the magnitude of the negative Fb values is smaller when using the adjusted emissions, whilst NMSE values are lower over near-
road sites but not background sites, which are dominated by regional PM. The large relative contribution of regional PM also causes the concentration changes (2 %–15 %) to be substantially smaller than the emissions changes (25 %–30 %) for these pollutants. For PM$_{2.5}$ the small overestimate of concentrations is increased by the emissions adjustment: Fb increases at near-road sites from 0.02 to 0.09. For all species the correlation coefficients remain very similar when using adjusted emission compared to the raw emissions, consistent with the correlation being influenced mainly by the variation in the relative magnitude of concentrations over time, not by their absolute magnitude.

All remaining model results presented in this section use the adjusted road traffic emissions.

### 3.2 Spatial variation of NO$_2$, O$_3$ and PM across London

Annual average contour plots of concentrations for NO$_2$, O$_3$ and PM$_{2.5}$ produced from the hourly coupled regional-to-urban model output using the adjusted emissions data are shown in Figs. 5–7. The influence of the M25 London orbital motorway is clearly visible for all three species. The corresponding monitored data are overlaid as coloured points. For NO$_2$, the highest concentrations (over 100 µg m$^{-3}$ in central London) are found near busy roads, while away from roads the concentrations increase from around 20 µg m$^{-3}$ outside the M25 to around 50 µg m$^{-3}$ in the centre of the urban area. The average NO$_2$ concentration calculated by the coupled model at urban background monitoring sites is 36 µg m$^{-3}$, just below the EU annual average limit value of 40 µg m$^{-3}$ for NO$_2$, while at near-road sites it is 60 µg m$^{-3}$, substantially above the limit value; corresponding measured values are 35 µg m$^{-3}$ for background sites and 61 µg m$^{-3}$ for near-road sites. The modelled fraction of NO$_x$ which is NO$_2$ increases from 0.43 at near-road sites to 0.59 at urban background sites, due to chemical conversion of emitted NO to NO$_2$. Across London, 333 km$^2$ (13 %) of the 2690 km$^2$ urban area within the M25 motorway, excluding road carriageways, exceeds the NO$_2$ annual average limit value, as shown by the yellow, orange and red colours in Fig. 5.

Annual average O$_3$ concentrations show an inverse pattern to NO$_2$, with low concentrations near busy roads (< 25 µg m$^{-3}$) and in the centre of the urban area, due to the effects of titration of O$_3$ by NO. There is no relevant limit
value for annual average O$_3$ for comparison. PM$_{2.5}$ concentrations show more uniform background concentrations of less than 10 µg m$^{-3}$ throughout the urban area, with steep increments near roads. The average PM$_{2.5}$ concentration calculated at urban background monitoring sites is 8.9 µg m$^{-3}$ and at near-road sites is 11.2 µg m$^{-3}$, both substantially below the annual average limit value of 25 µg m$^{-3}$ for PM$_{2.5}$. The increase in average concentrations at near-road sites over background sites is similar to the corresponding measured value, although the overall values are under-predicted (measured 13.7 µg m$^{-3}$ at background sites and 15.7 µg m$^{-3}$ at near-road sites, still well below the limit values). The absolute and relative concentration increment at near-road sites over urban background sites is smaller for PM$_{2.5}$ than for NO$_x$; this difference is captured by the coupled modelling system. A negligible fraction of the urban area (0.003 %) exceeds the annual average limit value of 25 µg m$^{-3}$ for PM$_{2.5}$, as shown by the predominantly blue and green colours in Fig. 7. A corresponding plot for PM$_{10}$ concentrations, showing very similar patterns to PM$_{2.5}$ and negligible exceedances of the annual average limit value of 40 µg m$^{-3}$, is given in Fig. A1.

Plots of annual average concentration of NO$_2$ and O$_3$ against site height, calculated from hourly observations and hourly coupled model output for monitors where both NO$_2$ and O$_3$ are available, are given in Fig. A2. They show generally good agreement between the modelled and observed concentrations, with the increased NO$_2$ and reduced O$_3$ at near-road sites compared to background sites captured by the model. There is no clear relationship between the concentrations and the site height, especially at the background sites where there is a slightly wider range of site heights.

Overall, the modelled pollutant distributions are closely related to the locations of explicit emissions sources and are also in good agreement with the spatial variation of observed concentrations, especially when viewed at street-scale resolution. The comparisons between modelled and monitored concentrations are discussed in more detail in the following sections.

3.3 Evaluation statistics for NO$_2$, O$_3$, CO, PM$_{10}$ and PM$_{2.5}$ for regional, local and coupled models

The performance of the regional EMEP4UK, local ADMS-Urban and coupled models has been assessed using evaluation statistics calculated from hourly concentrations of each pollutant. Table 4 gives statistics for NO$_x$ and NO$_2$ at all of the 42 background and near-road sites at which they are measured, while Table 5 gives statistics for O$_3$, NO$_x$, and NO$_2$ at the 20 sites where O$_3$ is measured, in order to allow detailed analysis of these closely related pollutants at a consistent set of sites. Table 6 gives corresponding statistics for CO and for the particulate pollutants PM$_{10}$ and PM$_{2.5}$. An additional visual representation of model performance, plots of NMSE against fractional bias, is given in Fig. A3. The statistics include those presented for the emissions adjustments in Table 3 (Sect. 3.1) as well as the fraction of modelled hourly concentrations within a factor of 2 of observations (Fac2) and the model quality indicator. The observed and modelled values of robust highest concentrations are also presented. If this value is calculated from all observed or modelled data, it is likely to be dominated by the highest values at a single site, so the approach of averaging individual site values over all sites has been taken in order to calculate more representative values for high observed and modelled concentrations.

The average measured NO$_x$ and NO$_2$ concentrations are lower for sites with O$_3$ measurements (shown in Table 5) compared to all sites (Table 4), as there is a lower proportion of near-road sites with O$_3$. However, the general findings are the same for both sets of sites. The regional EMEP model underestimates NO$_x$ and NO$_2$, as expected for a model using 5 km × 5 km gridded emissions. The stand-alone ADMS-Urban model and coupled ADMS-Urban RML system show broadly similar performance for the gaseous pollutants, indicating that the regional model is performing well at simulating the local background gaseous concentrations. For NO$_x$ and NO$_2$, the Fb and NMSE values are much lower when simulated by the stand-alone and coupled models than for
the regional model, due to the dominant influence of local emissions in determining concentrations for these short-lived species. Correlation coefficients are higher, with values of around 0.68 for both species for the stand-alone and coupled model simulations, as well as a similar increase in Fac2. The MQI values for all models except EMEP are less than 1 for NO\textsubscript{2}, indicating achievement of the FAIRMODE model quality objective. The modelled RHC shows both stand-alone and coupled models have good performance in the prediction of peak NO\textsubscript{2} concentrations. However, these models underestimate peak NO\textsubscript{x} values and have values of Fb for NO\textsubscript{2} greater than those for NO\textsubscript{x}, suggesting some over-prediction of NO\textsubscript{2} relative to NO\textsubscript{x} in general. This is at least in part likely to be due to an overestimate of the assumed fractions of NO\textsubscript{x} emitted as NO\textsubscript{2} (Carslaw et al., 2016).

The Fb values for O\textsubscript{3} concentrations from the urban and coupled models are also low (0.001–0.02), whilst the NMSE, R values and Fac2 results are fairly similar when comparing all three models to measurements. This reflects the significant contribution of regional background O\textsubscript{3} concentrations to the local concentrations within the urban area. The lower values of MQI for the urban and coupled models show improved overall model performance due to the inclusion of explicit sources. All three modelled RHC values are lower than the observations, indicating that, although the annual average O\textsubscript{3} concentrations are underestimated, the highest hourly concentrations are overestimated. This is likely to be due to additional short-term chemistry effects, for instance those caused by large increases in biogenic emissions in hot conditions (Guenther et al., 2006), which are not well captured by the models.

Although no adjustment was applied to the emission rates for CO there is reasonable agreement between model and observations, with particularly good values of Fb and Fac2 for the coupled model. The EU air quality standard for CO is 10 mg m\textsuperscript{-3} for maximum daily 8 h mean, whereas the maximum hourly observed concentration is around 2 mg m\textsuperscript{-3}, consistent with no observed exceedances of this standard.

For the particulate pollutants the ADMS-Urban model with measured background concentrations performs markedly better than the coupled model due to poorer performance of the regional model for these pollutants.
Table 6. CO model evaluation statistics calculated at 7 sites and particulate pollutants statistics calculated at 33 sites (PM$_{10}$) and 11 sites (PM$_{2.5}$) from regional (EMEP), local (ADMS-Urban) and coupled modelled hourly concentrations. Statistics as defined for Table 4. Note that MQI is not defined for CO.

<table>
<thead>
<tr>
<th>Poll</th>
<th>Model</th>
<th>Annual mean ($\mu$g m$^{-3}$)</th>
<th>Model evaluation statistics</th>
<th>Average RHC ($\mu$g m$^{-3}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Obs</td>
<td>Mod</td>
<td>Fb</td>
</tr>
<tr>
<td>CO</td>
<td>EMEP</td>
<td>318.8</td>
<td>232.8</td>
<td>−0.312</td>
</tr>
<tr>
<td>CO</td>
<td>ADMS-Urban</td>
<td>318.8</td>
<td>359.5</td>
<td>0.120</td>
</tr>
<tr>
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<td>Coupled</td>
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<td>317.3</td>
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<td>23.2</td>
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<td>0.139</td>
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<td>PM$_{2.5}$</td>
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<td>0.511</td>
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<tr>
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<td>ADMS-Urban</td>
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<td>15.8</td>
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<tr>
<td>PM$_{2.5}$</td>
<td>Coupled</td>
<td>14.7</td>
<td>10.0</td>
<td>−0.377</td>
</tr>
</tbody>
</table>

3.4 Annual average concentrations for NO$_2$, O$_3$ and PM$_{2.5}$

Some pollutants are closely connected by chemical or physical processes. Here the model performance for NO$_2$ and O$_3$ concentrations is evaluated concurrently. Figure 8 compares the annual average fractional bias for NO$_2$ and O$_3$ for each model for background and near-road site locations. For many sites the fractional bias of modelled concentrations for both pollutants from each model is within an estimated measurement uncertainty of 15%, as shown on the plot by the square of dotted lines. This is the maximum uncertainty allowable in continuous measurements reported to the EU (EC Directive, 2008). The remaining points, especially those for the regional model at near-road sites, most commonly show that overestimates of O$_3$ are associated with underestimates of NO$_2$ while underestimates of O$_3$ are associated with overestimates of NO$_2$, as expected from the fast O$_3$ titration chemistry that usually prevails in the urban high-NO$_x$ environment. There are near-road sites where the coupled model overestimates both NO$_2$ and O$_3$, which does not fit the general pattern.

Assessment target plots (developed as part of the DELTA tool within FAIRMODE, Janssen et al., 2017) allow model performance to be evaluated with an allowance for the measurement uncertainty (Pernigotti et al., 2013), which is particularly relevant for particulate pollutants because of their higher measurement uncertainty compared to gaseous pollutants. Figure 9 shows the coupled model results for PM$_{10}$ and PM$_{2.5}$ presented on target plots showing the normalized bias against the centred root mean square error (CRMSE) for each monitoring site, such that the distance of points from the origin gives the value of the MQI and allowance is made for measurement uncertainty. Equivalent plots for O$_3$ and NO$_2$ are given in Fig. A4. The quadrant in which the points are located depends on the magnitude of the relative contributions of any lack of correlation and standard deviation to the model.
error. Note that the correlation here is calculated with a consideration of measurement uncertainty and is different from the values given in Table 6. The area of the plot with green shading shows where the model errors are within a factor of 2 of the measurement uncertainty, leading to a value of MQI below 1. All of the PM$_{2.5}$ sites and all except one of the PM$_{10}$ sites lie within the green shading, which indicates achievement of FAIRMODE’s model quality objective. The plots show that the errors that occur are mainly associated with negative bias (underestimate), as noted for Fb values in Table 6 (Sect. 3.3), and lack of correlation.

### 3.5 Hourly concentrations and diurnal cycles for NO$_2$ and O$_3$

In this section an evaluation of hourly data and of diurnal cycles is performed. Figures 10–12 present frequency scatter plots of hourly NO$_x$, NO$_2$ and O$_3$ concentrations respectively, over all the sites where O$_3$ is measured, split by model and site type. The NO$_3$ plots show a large spread for high concentrations from the regional model at background sites, which is reflected in the coupled model. This may indicate an inaccuracy in the diurnal variation of emissions used in the regional model. The stand-alone local and coupled models capture the large range of observed hourly concentrations; however, as expected and noted from the evaluation statistics, the regional model underestimates NO$_3$ at near-road sites.

The scatter of NO$_2$ concentrations is substantially smaller than that for NO$_x$ concentrations, since the dependence of NO$_2$ on NO$_x$ is less than linear on account of the proportion of NO$_3$ that is NO$_2$ increasing with distance from an emission source, due to the chemical reaction of NO with O$_3$, whilst the concentration of NO$_3$ decreases due to mixing and dilution. A high density of points (indicated by the red and orange colours) lying close to the $y = x$ line indicates that a model accurately captures the complex balance between chemical and dispersion processes; Fig. 11 shows that the local and coupled models perform well at both background and roadside sites, but the regional model is unable to represent near-road NO$_2$. For a small number of hours NO$_2$ is overpredicted by the local and coupled models; this may be due to some overestimate of the fraction of NO$_x$, which is emitted in the form of NO$_2$ (primary NO$_2$), and to limitations in the local chemistry scheme in these cases. The plots for O$_3$ (Fig. 12) show generally good agreement between modelled and observed hourly concentrations for all models at all sites but they show an under-prediction of the peak observed values. Some of this may relate to the corresponding over-prediction of NO$_2$, suggesting that the rate of local O$_3$ production through NO$_2$ photolysis is underestimated. The under-prediction of peak background O$_3$ concentrations by the regional model is reflected in the coupled model results. The under-prediction of peak urban background concentrations by the stand-alone local model using measured rural upwind O$_3$ also indicates an underestimate of the local generation of O$_3$ through photochemistry within the urban area, for example due to an underestimate of biogenic VOCs in hot conditions (Malkin et al., 2016).
Mean diurnal profiles and 95% confidence intervals for NO$_x$, NO$_2$ and O$_3$ averaged over background and near-road sites are shown in Fig. 13, alongside a specific near-road site (BT4) in order to demonstrate the variability in individual sites. The BT4 site is located alongside the inner orbital North Circular Road, with annual average daily traffic of 108 000 vehicles spread across 6 lanes of traffic, and a neighbouring car park. The stand-alone urban and coupled models which include explicit road source emissions (ADMS-Urban and ADMS-Urban RML) typically capture the diurnal cycle of NO$_x$, NO$_2$ and O$_3$ concentrations for the different site types, whilst for the regional model this is only the case for background sites. The diurnal cycle for NO$_2$ strongly reflects NO$_x$ emissions at all site types, showing morning and afternoon traffic-related peaks, but also a dip around midday driven by a peak in NO$_2$ photolysis at this time. O$_3$ peaks around midday but concentrations are lower when NO$_2$ traffic-related peaks occur. Diurnal cycles are similar at both...
Figure 12. Frequency scatter plots for each model and site type showing the distributions of hourly average modelled and observed O$_3$ concentrations, where the colour represents the density of points for a given combination of measured and modelled values.

Figure 13. Diurnal temporal variations of NO$_x$, NO$_2$ and O$_3$ concentrations for the average over all background and near-road sites, and for an individual near-road site, with observations and modelled concentrations from each model. Note different concentration axis limits for each plot. The shaded area around the central line shows the 95% confidence interval in the mean.
background and near-road sites, although the NO$_x$ and NO$_2$ peak-to-peak concentration ranges are lower and O$_3$ higher at background compared to near-road sites (as noted for the annual average concentrations in Sect. 3.3). The observed diurnal cycle of NO$_x$ concentrations at the BT4 site has a notably higher morning peak than the cycle for the average over all near-road sites; this is less pronounced for NO$_2$ concentrations.

It is apparent that all of the models tend to overestimate O$_3$ when underestimating NO$_2$, especially for near-road sites, as noted for annual-average comparisons in Sect. 3.3. At BT4, the time-variation profile of NO$_x$ is not well captured by the models, but appears closer for NO$_2$ and O$_3$.

4 Discussion and conclusions

This study presents a regional-to-local air quality modelling system which couples the regional EMEP4UK model with the fine-scale urban model ADMS-Urban. Model simulations of NO$_x$, NO$_2$, O$_3$, CO, PM$_{10}$ and PM$_{2.5}$ using the coupled system are compared with the regional and urban models run separately and with measurements from background and near-road sites across London for 2012. This choice of base modelling year has allowed detailed assessment of the model chemistry schemes with the ClearfLo summer and winter intensive observation data (Bohnenstengel et al., 2015; Malkin et al., 2016). During the summer of 2012 London hosted the Olympic and Paralympic Games, but no effects from the games’ periods were apparent in a comparison of modelled and monitored concentrations at the monitoring site closest to the Olympic Park (now Queen Elizabeth Olympic Park).

The simulations make use of an emissions inventory in which road traffic emissions of NO$_x$, NO$_2$, PM$_{10}$ and PM$_{2.5}$ were adjusted to represent real-world conditions. Using the stand-alone version of ADMS-Urban these were shown to substantially improve both the fractional bias and normalized mean square error but had little effect on correlations with measured data as these depend on the relative changes in emissions hour by hour which were unaffected by the adjustment.

From the results using the coupled model it is estimated that 13 % of the area of London exceeds the EU annual average limit value of 40 µg m$^{-3}$ for NO$_2$ in 2012. This is consistent with the UK report to the EU of the Greater London urban area exceeding both the annual average and hourly average limit values (Defra, 2013a). In contrast, concentrations of PM$_{2.5}$ and PM$_{10}$ in London are estimated to have negligible exceedances of the annual average limit values of 25 and 40 µg m$^{-3}$ respectively.

The performance of the different modelling approaches used in this study varies depending on the relative importance of regional and local emissions, chemistry and transport processes for different pollutants and site types. The concentrations of the gaseous pollutants NO$_x$, NO$_2$ and CO are dominated by local emissions. This is particularly clear for NO$_x$ and NO$_2$, with large absolute and relative increments in concentration between background and near-road sites. The regional model (EMEP4UK) performs well at background sites but underestimates concentrations of these gases at near-road sites, due to the low resolution of its input emissions data which does not represent individual road sources. The urban (ADMS-Urban) and coupled models both show good agreement compared to measurements at both site types due to the inclusion of explicit source emissions. This means that the coupled model system can be used with confidence for locations or time periods where rural upwind measurements are not available for use in ADMS-Urban or for assessment of impacts of future emissions or climate change.

The model performance statistics for NO$_2$ are generally better than those for NO$_x$ for all models. This is in part due to the reduced sensitivity to NO$_x$ emissions of NO$_2$ concentrations relative to NO$_x$ concentrations, as exemplified by the analysis of the emissions adjustments. However, the clear inverse relationship between model biases for NO$_2$ and O$_3$ is consistent with the local chemistry generally being well modelled, with uncertainty in NO$_2$ and O$_3$ being related to uncertainty in NO$_x$. Comparison between the coupled and stand-alone urban models and measurements of average diurnal profiles of NO$_x$ concentrations suggest the models are capturing the measured features of the profiles, although there is some underestimation in NO$_x$ at roadside around midday.

The concentrations of PM$_{10}$, PM$_{2.5}$ and O$_3$ show more influence from long-range transport than the other gaseous pollutants. Hence, the coupled model results for these species are strongly affected by the regional background and the regional model simulation of PM$_{10}$, PM$_{2.5}$ and O$_3$. For the coupled model, simulated O$_3$ agrees well with measured O$_3$ at the background sites, but simulated PM$_{10}$ and PM$_{2.5}$ are largely underestimated compared to background site measurements. However, the coupled model still shows a significant improvement compared to the regional model for simulated particulate concentrations, especially at near-road sites, because it includes an explicit representation of local source emissions. The average increment in PM$_{2.5}$ concentrations between background sites and near-road sites is much smaller than for NO$_2$ but is well represented by the coupled system.

The ability of the models to simulate high concentrations has also been investigated. In general the high concentrations are well simulated by the three models and for all pollutants examined in this study except for PM$_{10}$, where the highest concentrations are due to local short-term emission effects, for example construction dust, which is not included in emissions inventories. For PM$_{2.5}$, the urban model gives a reasonable value of the RHC metric and hence of high concentrations, which indicates that these are due to long-range transport, such as from forest fires, which is captured by the measured upwind rural background concentrations but not necessarily by the regional model. The general tendency of
the EMEP4UK regional model to underestimate particulate concentrations, both long-term and episodic, has been identified previously (Lin et al., 2017; Vieno, 2016b), while all models are affected by local emission effects and uncertainties in measured particulate concentrations (Pernigotti et al., 2013).

Representing the time variation of emissions accurately, including the variation between sites and pollutants, is a challenge for all models and particularly affects the correlation values. In the current work a single time-varying profile was used for all road emissions in the urban and coupled models but the modelling would be improved if more detailed profiles were included. However, no time variation data are currently associated with the LAEI. A further model performance evaluation that would be of great interest would be to assess the models against measurements over a wider range of heights, as model predictions are routinely used to calculate building facade concentrations within street canyons. Some very high measurements were carried out during the ClearfLo project, at a height of 180 m on the BT tower, but the authors are not aware of any measurements covering the range of average building heights in central London of 20–40 m (as shown in Fig. 3).

Overall, this study has shown the benefit of coupling a regional atmospheric chemistry transport and dispersion model with a local model in order to calculate detailed spatiotemporal distributions of air pollutants. Such detailed pollutant spatial distributions have applications in health-related exposure analysis (Smith et al., 2016). The coupled system could also be used to assess the effects of air quality policies at a range of scales. An extension to the current study would be to process the hourly model output to assess the exceedance of short-term objectives and combine the results with population data to calculate exposure. The work presented in this paper provides a framework for more detailed examinations of urban atmospheric chemistry, in particular the effects of additional species which interact with \( \text{NO}_x \), \( \text{NO}_2 \) and \( \text{O}_3 \), and for studies of the effects of the urban heat island and future climate on urban air quality and chemistry.

**Data availability.** The authors intend to make the high-resolution hourly output data from the coupled model available on the British Atmospheric Data Centre (BADC). The data may be accessed in the meantime by contacting the authors.

AURN data are available via http://uk-air.defra.gov.uk (last access: 7 December 2017, Defra, 2017). LAQN data were obtained from the Environmental Research Group of Kings College London (http://www.erg.kcl.ac.uk), using data from the London Air Quality Network (http://www.londonair.org.uk/london/asp/datadownload.asp, last access: 2 August 2018, Mittal et al., 2015). Both of these data sets are licensed under the terms of the Open Government Licence.
Appendix A

Table A1 gives the definitions of the WRF modelling domains used for the EMEP4UK modelling system. The top half of the table gives details of how the WRF domains are defined and the bottom half presents the coordinate system parameters.

Contours of annual average PM$_{10}$ concentrations predicted by the coupled model, overlaid with the observed annual average concentrations, are shown in Fig. A1. The pattern is similar to the plot for PM$_{2.5}$ (Fig. 7), with negligible exceedance of the annual average standard of 40 µg m$^{-3}$, as shown by the mostly blue and green colours. However, these model predictions need to be treated with some caution due to the model’s general underestimate of PM$_{10}$ concentrations, demonstrated by the statistics given in Table 6.

Plots of annual average concentration against site height for NO$_2$ and O$_3$, calculated from observations and coupled model predictions, are shown in Fig. A2. For background site types, there does not seem to be a clear relationship between concentration and height above ground in the range 2–10 m. The model captures the general differences between near-road and background sites, with increased NO$_2$ and reduced O$_3$.

Following Chang and Hanna (2004), the model performance by site type has been assessed visually as shown in Fig. A3. In general all the models show good performance, with points clustered close to the origin of the graph. The regional model represents background sites adequately but has less good agreement for near-road sites, particularly for NO$_x$ and NO$_2$ where concentrations are dominated by local road emissions, as expected. The urban and coupled models, which represent road sources explicitly, show similar performance for background and near-road sites, with some variation between pollutants. The coupled model shows similar performance to the regional model for background sites, especially for the particulate pollutants and CO, showing the greater influence of the regional model at sites where there are fewer explicit sources represented by the local model.

Figure A4 shows target plots for NO$_2$ and O$_3$ for the coupled model (both plots only include sites where O$_3$ is measured); results match the measurements to within the target criterion. The model agreement for O$_3$ is particularly good, with many points within the inner circle and a MQI value less than 0.5 indicating that any difference between the modelled and measured values is less than or equal to the estimated measurement uncertainty.

### Table A1. Definitions of WRF modelling domains used for the EMEP4UK modelling system. Note that domain 2 is only used for meteorological modelling.

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<th>Domain 3</th>
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Figure A1. PM$_{10}$ annual average concentration contours from the coupled model for the whole of Greater London (a) and an area of central London (b), with monitoring data overlaid – round symbols for near-road sites and square symbols for background sites.

Figure A2. NO$_2$ (a) and O$_3$ (b) annual average concentrations from measurements and the coupled model, plotted against site height.

Figure A3. Model evaluation plots comparing NMSE and Fb for near-road or background sites by pollutant, where improved model performance is shown by the results closest to (0,0). The solid parabola indicates the minimum NMSE for a given Fb, while the dashed lines show modelled results within a factor of 2 of the observations. Note that the NO$_x$ plot has different axis limits to the other pollutants due to the outlying EMEP near-road point.
Figure A4. Model assessment target plots for NO$_2$ (a) and O$_3$ (b) for coupled system outputs. Each symbol represents a single station and the distance between the origin and the symbol corresponds to the MQI for that station; the ordinate and abscissa correspond to the bias and CRMSE respectively. Good model performance is indicated by points within the green shading.
Author contributions. IM set up and ran the regional model with support from MV and RD. KJ set up and ran the local stand-alone model with support from JS and CH. CH developed the coupled system with support from JS and DC. IM ran the coupled system with support from CH and RD. KJ and CH processed the model outputs with support from JS. This paper was drafted by CH with input and review by IM, JS, DC, MV and RD.

Competing interests. Christina Hood, Jenny Stocker and Kate Johnson work for and David Carruthers is a director of CERC, who develop and license the ADMS-Urban and ADMS-Urban RML models.

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