

# The treatment of uncertainties in reactive pollution dispersion models at urban scales

A. S. Tomlin,<sup>\*a</sup> T. Ziehn,<sup>†a</sup> P. Goodman,<sup>‡b</sup> J. E. Tate<sup>b</sup> and N. S. Dixon<sup>c</sup>

Received 10th November 2015, Accepted 27th November 2015

DOI: 10.1039/c5fd00159e

The ability to predict NO<sub>2</sub> concentrations ([NO<sub>2</sub>]) within urban street networks is important for the evaluation of strategies to reduce exposure to NO<sub>2</sub>. However, models aiming to make such predictions involve the coupling of several complex processes: traffic emissions under different levels of congestion; dispersion *via* turbulent mixing; chemical processes of relevance at the street-scale. Parameterisations of these processes are challenging to quantify with precision. Predictions are therefore subject to uncertainties which should be taken into account when using models within decision making. This paper presents an analysis of mean [NO<sub>2</sub>] predictions from such a complex modelling system applied to a street canyon within the city of York, UK including the treatment of model uncertainties and their causes. The model system consists of a micro-scale traffic simulation and emissions model, and a Reynolds averaged turbulent flow model coupled to a reactive Lagrangian particle dispersion model. The analysis focuses on the sensitivity of predicted in-street increments of [NO<sub>2</sub>] at different locations in the street to uncertainties in the model inputs. These include physical characteristics such as background wind direction, temperature and background ozone concentrations; traffic parameters such as overall demand and primary NO<sub>2</sub> fraction; as well as model parameterisations such as roughness lengths, turbulent time- and length-scales and chemical reaction rate coefficients. Predicted [NO<sub>2</sub>] is shown to be relatively robust with respect to model parameterisations, although there are significant sensitivities to the activation energy for the reaction NO + O<sub>3</sub> as well as the canyon wall roughness length. Under off-peak traffic conditions, demand is the key traffic parameter. Under peak conditions where the network saturates, road-side [NO<sub>2</sub>] is relatively insensitive to changes in demand and more sensitive to the primary NO<sub>2</sub> fraction. The most important physical parameter was found to be the background wind direction. The study highlights the key parameters required for reliable [NO<sub>2</sub>] estimations suggesting that accurate reference measurements for wind direction should be a critical part of air quality assessments for in-street locations. It also highlights the importance of street

<sup>a</sup>Energy Research Institute, School of Chemical and Process Engineering, University of Leeds, Leeds, UK. E-mail: A.S.Tomlin@leeds.ac.uk

<sup>b</sup>Institute for Transport Studies, University of Leeds, Leeds, UK

<sup>c</sup>School of Earth and Environment, University of Leeds, Leeds, UK

<sup>†</sup> Current address: CSIRO, Oceans and Atmosphere, Aspendale, Australia.

<sup>‡</sup> Current address: School of Civil Engineering and Geosciences, Newcastle University, UK.



scale chemical processes in forming road-side  $\text{NO}_2$ , particularly for regions of high  $\text{NO}_x$  emissions such as close to traffic queues.

## A Introduction

Although European directives to reduce  $\text{NO}_x$  emissions from vehicles have been in operation for well over a decade, many urban areas across Europe are still failing to meet the  $\text{NO}_2$  air quality standards set by the EU Directive 2008/50/EC. Within the UK there are a large number of Air Quality Management Areas (AQMAs) which have been declared on the basis of  $\text{NO}_2$ , a large proportion of which are in highly trafficked urban areas. Strategies must therefore be put in place to address potential reductions in  $\text{NO}_2$  concentrations, particularly focussed on traffic sources. To aid in this AQM process, models can be developed which aim to predict road-side  $\text{NO}_2$  as a function of important traffic characteristics and can therefore be used to assess the potential impact of traffic management or emissions based intervention measures. Such models should also include relevant features of the urban environment that affect pollution formation and dispersion such as wind speed and direction, rapid chemical transformations and street topologies. The latter feature should be included since it is well understood that urban buildings interact with background winds to modify the turbulent flow structures within the streets, restricting the dispersion of traffic related pollutants out of the street network,<sup>1-5</sup> and potentially allowing time for in-street chemical processing of emissions. Models aiming to achieve road-side predictions therefore should contain representation of this potentially circulating flow occurring within the street canyons, as well as vertical and horizontal fluxes into and out of the street network.<sup>3</sup> Representations of chemical transformations are also required, in particular those relevant to the formation of secondary  $\text{NO}_2$ . A number of modelling approaches have been suggested to address the dispersion part of the problem including Gaussian based models such as OSPM (Operational Street Pollution Model),<sup>6</sup> network models such as SIRANE,<sup>7</sup> and compartment based models where pollutant exchange rates are parameterised according to canyon aspect ratios,<sup>8</sup> as well as high resolution computational fluid dynamics (CFD) approaches.<sup>9,10</sup> Several studies have attempted to couple models of the complex turbulent flow with chemical sub-models, albeit for single or small networks of streets.<sup>11-15</sup> These studies have highlighted the influence of incomplete mixing on the formation of secondary pollutants in regions of high primary  $\text{NO}_x$  emissions. This suggests that coarse resolution models (*e.g.* urban air shed models) are unlikely to be of direct relevance for the study of road-side concentrations and exposure, although they may provide boundary conditions for higher resolution studies at the street scale. However, detailed representations of the interplay between emissions, chemical transformations and dispersion within the street network will be critical for near field exposure. Few models, however, have attempted to couple this modelling of dispersion at the micro-scale with both chemical transformation processes and high resolution traffic emissions models. The current work aims to present such an integrated system which couples a micro-scale traffic emissions model with a turbulent reactive dispersion model based on a combined CFD and reactive Lagrangian particle dispersion approach.<sup>16,17</sup>



The use of such models within the AQM framework requires an understanding of the confidence that can be placed in their predictions. Lack of confidence, or uncertainty, can result from a lack of detailed knowledge of the model parameterisations. It follows that model evaluation will benefit from the inclusion of sensitivity studies that highlight the impact of uncertain input parameters on predicted output concentrations. As part of an uncertainty/sensitivity study we might ask questions such as: (i) how robust are predicted road-side NO<sub>2</sub> concentrations, *i.e.* how wide are the predicted distributions if we properly account for uncertainties within the model input parameters, and (ii) taking into account uncertainties in the model parameters, can we trace the effects of potential traffic management strategies such as reductions in demand, reductions in emissions, and changes in emissions profiles, or are these effects swamped by uncertainties due to the model itself?

Within CFD based approaches, the use of Reynolds Averaged Navier Stokes (RANS) models in particular has raised questions as to their suitability for accurately describing turbulent chemical interactions when they contain only averaged representations of turbulent length and time-scales. On the plus side, such models allow the representation of reactive dispersion within complex street topologies at lower computational cost than, for example, Large Eddy Simulations (LES). On the negative side, they require parameterisation of mixing lengths rather than resolving eddies in an explicit way. They do however, provide much more realistic descriptions of in-street processing of pollutants than urban air shed type models which are too coarse to capture complex street geometries and the effects of buildings on in-street flow profiles. Given the potential utility of such RANS models, it is worth considering how robust their simulations are to the parameterisations chosen. We attempt to address some of these questions here and present an approach for the assessment of sensitivities for a complex multi-component model aiming to predict time-averaged road-side concentrations of NO<sub>2</sub> as a function of street topologies, background meteorology, traffic characteristics and temperature dependent parameterisations of chemical reactions.

## B Methodology

### (i) Case study and dispersion model structure

The location modelled in this study is that of Gillygate, York, UK, the site of an extensive measurement campaign<sup>2,18</sup> that has provided observations used in previous evaluations of some of the model components used here.<sup>10</sup> Gillygate is a relatively narrow street with an aspect ratio (building height to street width) of approximately 0.8, leading to cross-street recirculating flow under a range of background wind directions, restricting the dispersion of pollutants out of the street. The traffic flow along Gillygate is quite high with significant periods of congestion, and it therefore represents a potential pollution hot spot. It was included within the stated AQMAs by the City of York Council in 2014.<sup>19</sup> 19 exceedances of the hourly limit value for NO<sub>2</sub> of 200 µg m<sup>-3</sup> were noted in 2011, and annual average [NO<sub>2</sub>] values varied between 30 and 50 µg m<sup>-3</sup> from (2010–2015) based on data from the air quality monitoring site which is situated away from the congested junction at the Northern end of the street. Diffusion tube data from various points within Gillygate<sup>19</sup> showed significant variation in concentrations along the street, with annual average values exceeding 50 µg m<sup>-3</sup> in some



locations. It therefore provides a useful polluted road-side case study on which to demonstrate the approach adopted for model uncertainty analysis in this work.

Fig. 1 (top) shows the grid and the building configuration of Gillygate and the surrounding area that were used for the simulations in this study. The building heights in metres are indicated in the legend. Fig. 1 (bottom) provides an aerial view of the main part of the modelling domain. The basis for the underlying flow and turbulence model under consideration is the  $k$ - $\epsilon$  RANS model MISKAM.<sup>20</sup> This model was chosen on the basis that it is commonly used as an operational model<sup>9</sup> and has undergone previous evaluation for street canyon case studies.<sup>10</sup> It has been shown to substantially improve the representation of dispersion in built up areas due to improved representation of turbulent flow patterns compared to more general dispersion models such as those used in compliance assessment.<sup>21</sup> In this study, a non-equidistant grid was used to enable a higher resolution within the area of interest.

Fig. 1 shows the two locations G3 and G4 that were used in the original measurement campaign as well as the mast location that was used to obtain reference wind speed and direction. The experimental measurements were of concentrations of the non-reactive tracer carbon monoxide. Unfortunately no measurements of  $[\text{NO}_2]$  are available from the campaign but previous evaluations with respect to  $[\text{CO}]$  highlighted the ability of the model to capture the main flow and dispersion characteristics within the street.<sup>10</sup> The uncertainty analysis here allows additional investigation into the robustness of the model with respect to the chemical parameterisations. We use the same locations here as in the previous study for investigating  $[\text{NO}_2]$  predictions, as well as three other sites on each side of the street at 20 m intervals to the South of G3 and G4. A wind direction of  $0^\circ$  represents channelled flow from North to South along the street canyon. The wind directions sampled in the case study ( $110$ – $130^\circ$ ) represent oblique flow over the building adjacent to G3 towards the North of the domain and lead to a helical in-street recirculating flow with a northerly channelled component.<sup>2</sup> We focus on a single wind sector here due to the computational cost of the random sampling approach required for the global sensitivity calculations. In reality of course, many different wind directions would be used within an air quality assessment. Since model validation is usually performed using concentrations normalised by a reference wind speed, the inflow wind speed is kept to a constant value of  $5 \text{ m s}^{-1}$  at a height of 50 m, with the vertical profile then determined using a log-law which is based on the sampled values of the roughness length for the incoming flow.

The output from MISKAM is used as the underlying turbulent flow structure for a dispersion model based on the Lagrangian stochastic particle dispersion approach with micro-mixing and chemical sub-models (for a full description and evaluation see ref. 16 and 17). The complex dispersion modelling system was used previously to investigate a reactive plume of nitrogen oxides ( $\text{NO}_x$ ) released into an approximately homogeneous turbulent grid flow doped with ozone ( $\text{O}_3$ ) for comparison against wind tunnel experiments.<sup>17</sup> The chemical and micro-mixing sub-models used here are the same as those specified in the photolysis extended case described in Ziehn *et al.*<sup>17</sup> The chemical reactions included are detailed in Table 1. In summary, only simple reactions between  $\text{NO}$ ,  $\text{NO}_2$  and  $\text{O}_3$  are included in the chemical model but these include the photolysis of  $\text{NO}_2$  and  $\text{O}_3$ . Quite broad ranges have been included for photolysis rates reflecting variation in daytime conditions. No organic reactions are included in the analysis. Rather, any reactions



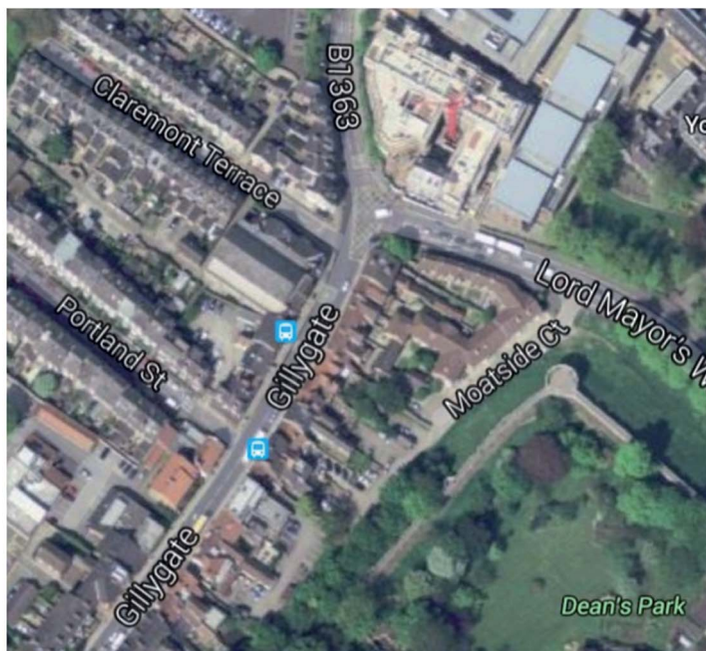
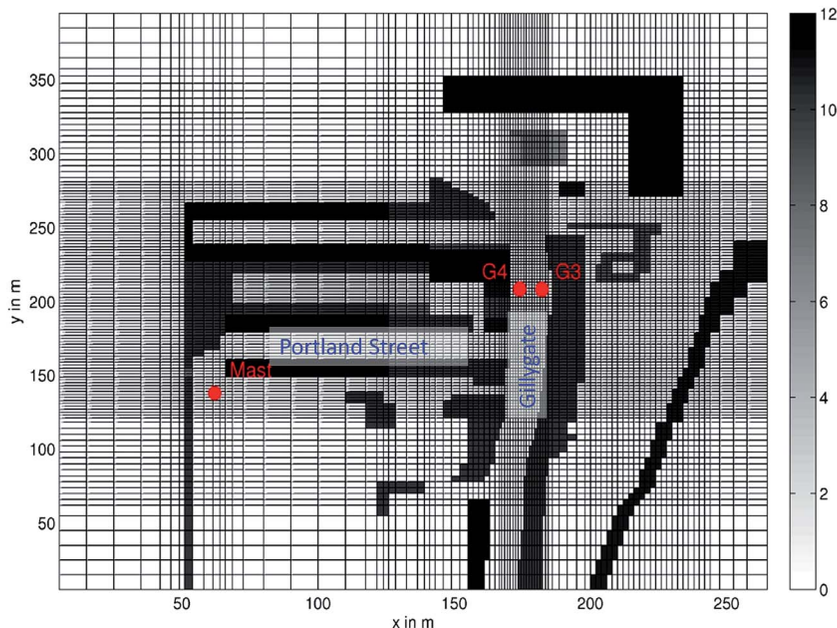


Fig. 1 Top: Site schematic for the York Gillygate site showing the grid and building configuration as used in MISKAM. Bottom: Aerial view of the site. ©2015 Infoterra Ltd. & Bluesky, ©2015 Google.

occurring at longer time-scales are represented implicitly within the description of above-roof  $O_3$  concentration which is considered as an uncertain parameter. A high sensitivity with respect to this parameter would indicate the importance of long



**Table 1** Chemical scheme parameter ranges used within the uncertainty analysis. Concentrations of  $[N_2] = 2.03 \times 10^{19}$  molecule  $\text{cm}^{-3}$  and  $[O_2] = 5.45 \times 10^{18}$  molecule  $\text{cm}^{-3}$  have been used

Parameter	Nominal value	Minimum	Maximum	Unit	Source
A factor, R1 $O + O_2 + N_2 = O_3 + N_2$	$5.60 \times 10^{-34}$	$4.991 \times 10^{-34}$	$6.283 \times 10^{-34}$	$\text{cm}^6$ per molecule <sup>2</sup> per s	33
A factor, R2 $O + O_2 + O_2 = O_3 + O_2$	$6.00 \times 10^{-34}$	$5.348 \times 10^{-34}$	$6.732 \times 10^{-34}$	$\text{cm}^6$ per molecule <sup>2</sup> per s	33
A factor, R3 $O + O_3 = 2O_2$	$8.00 \times 10^{-12}$	$6.654 \times 10^{-12}$	$9.618 \times 10^{-12}$	$\text{cm}^3$ per molecule per s	33
A factor ( $k_0$ ), R4 $O + NO + M = NO_2 + M$	$1.00 \times 10^{-31}$	$0.794 \times 10^{-31}$	$1.259 \times 10^{-31}$	$\text{cm}^6$ per molecule <sup>2</sup> per s	33
A factor R4 ( $k_1$ )	$3.00 \times 10^{-11}$	$1.504 \times 10^{-11}$	$5.986 \times 10^{-11}$	$\text{cm}^3$ per molecule per s	33
A factor, R5 $O + NO_2 = NO + O_2$	$5.50 \times 10^{-12}$	$4.790 \times 10^{-12}$	$6.315 \times 10^{-12}$	$\text{cm}^3$ per molecule per s	33
A factor, R6 $NO + O_3 = NO_2 + O_2$	$1.40 \times 10^{-12}$	$1.165 \times 10^{-12}$	$1.683 \times 10^{-12}$	$\text{cm}^3$ per molecule per s	33
Photolysis rate $J O_3$ , R7 $O_3 = O + O_2$	$2.75 \times 10^{-5}$	$1.0 \times 10^{-5}$	$4.5 \times 10^{-5}$	$\text{s}^{-1}$	34
Photolysis rate $J NO_2$ , R8 $NO_2 = NO + O$	0.0075	0.004	0.011	$\text{s}^{-1}$	35
$n$ for reaction R1	2.6	2.1	3.1	—	33
$n$ for reaction R2	2.6	2.1	3.1	—	33
$E/R$ for reaction R3	2060	1860	2260	$\text{K}^{-1}$	33
$n_0$ for reaction R4	1.6	1.3	1.9	—	33
$n_\infty$ for reaction R4	-0.3	-0.6	0	—	33
$E/R$ for reaction R5	-188	-268	-108	$\text{K}^{-1}$	33
$E/R$ for reaction R6	1310	1110	1510	$\text{K}^{-1}$	33

range chemical processes for the net formation of  $\text{NO}_2$ . For the current study the coupled dispersion model is further linked to a traffic micro-simulation model and a zero background concentration of  $[\text{NO}_2]$  is assumed so that the modelled concentrations represent road-side increments above background.

## (ii) Traffic micro-simulation modelling

Vehicle flows within the study area were modelled using an established, commercial traffic micro-simulation package AIMSUN 5.1.10 (ref. 22) which represents the movement of individual vehicles through a road network using discrete time intervals of the order of one second. Individual components within the micro-simulation govern the interaction of vehicles with one another, the interaction of vehicles with traffic signals, how vehicles make lane-changing manoeuvres and how vehicles accept gaps in traffic streams. Within each time-



step individual components are called, using information from the previous time step, to assign new kinematic information (acceleration, speed and position) to every vehicle. Vehicle and driver parameters which are considered static within a given run (*e.g.* maximum vehicle acceleration rates) are generated on vehicle entry to the network, and sampled from appropriate distributions. Given the fine spatial and temporal resolution of traffic micro-simulation, output statistics may be aggregated over a wide variety of scales, for use within appropriate environmental models. A substantial body of literature already exists on methodologies detailing such approaches, *e.g.* ref. 23 and 24.

### (iii) Traffic emissions modelling

Using the AIMSUN model, total NO<sub>x</sub> emissions were calculated using the polynomial emissions functions proposed by Int Panis *et al.*<sup>25</sup> based on vehicle type, instantaneous speed, and acceleration parameters. Instantaneous emissions rates (g s<sup>-1</sup>) were calculated for each vehicle at each model time-step (~1 s) using polynomial functions. These were converted to mass values, based on initial vehicle velocity, and the assumption of linear acceleration over the time-step. Given the position of the vehicle at the start and end of a time-step, contributions were then apportioned to the individual sections, through which the vehicle had passed. The final output from the bespoke software was the emission rates (g h<sup>-1</sup>) for each 10 m road section, based on the integrated total of all contributions from vehicles passing through the sections in that period. These were then converted to g m<sup>-1</sup> s<sup>-1</sup> for input into the dispersion model.

The specific fraction of NO<sub>2</sub> within the total NO<sub>x</sub> was treated as an uncertain parameter as discussed later. Calculated emissions were then linked by vehicle position to a particular 10 m section of road giving spatial-profiles of emissions along Gillygate *via* bespoke software external to AIMSUN.<sup>26</sup> The overall traffic network consisted of 4 km of roads surrounding Gillygate and 8 intersections, including 2 which were signalised. Four categories of vehicles were considered: cars, vans, HGVs and buses, for compatibility with Int Panis *et al.*<sup>25</sup> The dynamic demand in the network (the number of vehicles desiring to travel through the network within the simulated hour) was varied over two sets of normalised ranges. The first is an “off-peak” case from 0.8–1.2 with the mean value of 1.0 representing ‘typical’ inter-peak demand. The second was a “peak” case with demand varying from 1.2–1.6. Each simulation run therefore represented 1 h at a particular level of demand using a random sampling approach within the specified ranges.

The normalised demand level was derived from a year of traffic flow data obtained from York's urban traffic control system equating to a two-way flow of ~880 veh per h along Gillygate. Additional to the dynamic demand was a fixed level of demand from buses based on timetable information. At the base demand level, the network is considered as busy, but in an ‘under-saturated’ state, *i.e.* able to cope with the level of demand, with only transient queues forming at junctions. At demand levels above 1.1, modelled speeds begin to decline rapidly from ~20 km h<sup>-1</sup> to ~10 km h<sup>-1</sup> at a demand of 1.4. At these higher demand levels, substantial over-saturated queues form as vehicles are unable to clear signalised junctions within a single signal period. For off-peak, under capacity periods, total emissions increase in a slightly non-linear fashion with the volume of traffic as shown for a typical section of Gillygate in Fig. 2a. Some of the non-linearity may be explained by the increasing



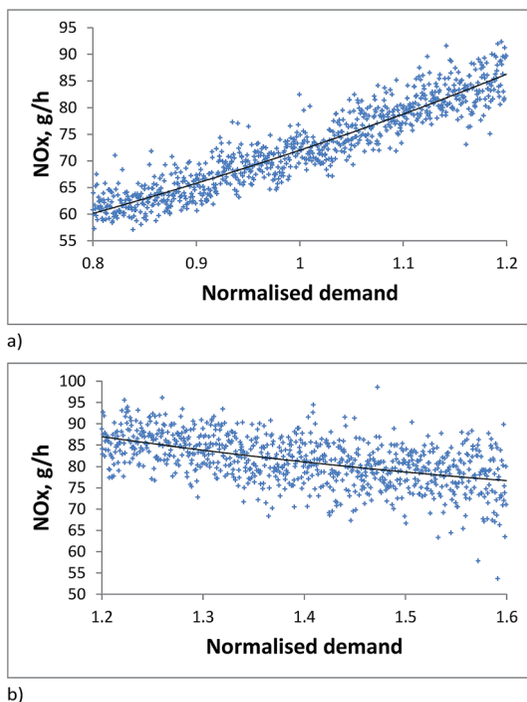


Fig. 2 Total exhaust emissions of NO<sub>x</sub> for a sample section of Gillygate for (a) off-peak and (b) peak demand scenarios.

relative fraction of HGVs present, whose contribution to NO<sub>x</sub> emissions starts to dominate those of passenger cars. This phase is followed by a transitional period as demand approaches and exceeds network capacity, where emissions stabilise at a high overall level as shown in Fig. 2b. The influence of these characteristics on modelled road-side mean NO<sub>2</sub> levels is discussed in the next section.

#### (iv) Model parameterisations and ranges of uncertainty

Close to a surface the effects of turbulence in a RANS model need to be modelled using boundary conditions that reflect the surface roughness. MISKAM represents these boundary conditions using an idealised log-law based on three surface roughness lengths  $z_0$  for the incoming boundary flow, the urban surface and wall surfaces, respectively. It follows that parameterisations of these roughness lengths will be a possible source of uncertainty in the final output predictions since they determine the near surface velocity profiles and therefore influence both dispersion and deposition<sup>27</sup> processes. A study by Benson *et al.*,<sup>28</sup> based on the application of MISKAM to a street canyon scenario, showed that overall, the uncertainty in the predicted mean and turbulent flow fields due to roughness lengths was small in comparison to the mean outputs. The study concluded that the model was well defined even with large ranges of input parameter uncertainty. In general, at a particular receptor, the closest surface was found to be most influential on the model output except when a recirculating canyon flow pattern was strongly present. In this latter case background wind angle was found to dominate.



The turbulent flow field is used as an input to the Lagrangian particle dispersion model where it is assumed that the velocity of a fluid element, for times larger than the Kolmogorov time-scale, is a Markov process that is a continuous function of time.<sup>29</sup> Within the Lagrangian particle model framework, the two important parameterisations are the Lagrangian structure coefficient  $c_0$ , and the mixing time-scale coefficient  $\alpha$ . The Lagrangian structure function is defined as the ensemble average of the square of the change in Lagrangian velocity and the definition of  $c_0$  is therefore important in determining the effective turbulent diffusion in velocity space. There is some debate within the literature as to whether its value can be universally defined for all types of turbulent flows with a range of values between 2 and 10 quoted from different studies.<sup>30–32</sup> It is interesting to establish therefore how sensitive concentration predictions are to the chosen value. Within the model tested, a simple particle mixing model is adopted, that of interaction by exchange with the mean (IEM) concentration.<sup>29</sup> In order to provide generality, the mixing model uses a coefficient  $\alpha$  which defines the relationship between the turbulent time-scales (total turbulent kinetic energy and its dissipation rate) and the mixing time-scale at every point in the flow as defined by the following equation:

$$t_m = \alpha \frac{k}{\varepsilon}$$

The specification of  $\alpha$  should also be considered to be uncertain and the relative time-scales of the chemical processes compared to the mixing time-scale could have an important influence on the formation of secondary species such as  $\text{NO}_2$ .

Uncertainties in the traffic emissions model have been adopted for the level of traffic demand as discussed above, and the  $\text{NO} : \text{NO}_x$  ratio for the emissions source which determines the fraction of  $\text{NO}$  vs. primary  $\text{NO}_2$  assumed at source. The range adopted was chosen to reflect levels of primary  $\text{NO}_2$  estimated for UK vehicle fleets,<sup>36</sup> taking into account a range of possible fuel types as well as engine exhaust treatment methodologies (15–25% primary  $\text{NO}_2$ ). The 26 model parameters varied within the sensitivity/uncertainty analysis can therefore be summarised as:

- Velocity structure function coefficient  $c_0$  [4–6].
- Mixing time-scale coefficient  $\alpha$  [0.6–3].
- Surface roughness length  $z_0$  for inflow, surface and wall [5–50, 0.5–50, 0.5–10 cm].
- Temperature dependent Arrhenius rate parameters for  $\text{NO}/\text{NO}_2/\text{O}_3$  reactions, photolysis rate parameters for  $J \text{O}_3$  and  $J \text{NO}_2$  [see Table 1 for details].
- Background wind direction  $\theta$  [110–130°].
- Temperature [273–298 K].
- Background ozone concentration [ $7.35 \times 10^{11}$  to  $1.23 \times 10^{12}$ , molecules  $\text{cm}^{-3}$  or 30–50 ppb].
- $\text{NO} : \text{NO}_x$  ratio for traffic emissions [0.75–0.85].
- Normalised traffic demand [off peak 0.8–1.2, peak 1.2–1.6].

Where the ranges used are shown in the square brackets except for the Arrhenius and photolysis parameters which were detailed in Table 1. Of these parameters, most relate to uncertainties in the model formulation. For a given urban morphology, it is mainly the last three, background  $\text{O}_3$  concentration,  $\text{NO} : \text{NO}_x$



ratio and traffic demand, that could be affected by the implementation of pollution mitigation strategies. Therefore if the model were to be highly robust, we would like to see the overall sensitivities of the model dominated by these last three parameters. In a wider study it would of course be possible to assess the impact of changes to urban form including “urban greening<sup>8</sup>” which may impact on roughness lengths and therefore dispersion and deposition.

### (v) Global uncertainty and sensitivity analysis methods

Within a global uncertainty analysis the aim is to determine the possible range of predicted outputs (in this case time-averaged road-side NO<sub>2</sub> concentration increments) given the ranges of uncertainty within the model input parameters. Often a Monte Carlo type approach is adopted where random or quasi-random samples of the inputs are used within the adopted ranges of uncertainty, and many model simulations are performed; one for each random sample. The result is a predicted distribution of outputs instead of a single value. The width of this distribution gives the modeller an idea of the robustness of the model given the model input uncertainties. Statistical parameters such as standard deviation can be used to provide simple quantified measures of the output uncertainty in cases where the distribution does not deviate too far from being Gaussian. Otherwise higher moments may also be useful. The aim of improving model robustness is to provide better and better quantification of the model inputs such that the predicted output distribution becomes as narrow as possible *i.e.* the predicted output uncertainty is low.

In order to assist in the model improvement processes it is useful to determine which of the uncertain model inputs have the largest influence of predicted output uncertainties. This is achieved here through a global sensitivity analysis based on the ANalysis Of VAriance (ANOVA) approach.<sup>37</sup> In such methods, the output variance is decomposed into component functions representing the effects of individual and groups of parameters whose importance can then be ranked according to global sensitivity indices.

The global sensitivity analysis has been achieved using the RS-HDMR (Random Sampling High Dimensional Model Representation) method introduced by Rabitz *et al.*<sup>38</sup> to express the input–output relationship of complex models with large numbers of input parameters, and further developed into a user friendly Matlab package GUI-HDMR by Ziehn and Tomlin.<sup>39</sup> The mapping between input parameters  $x_1, \dots, x_n$  and output variables  $f(x) = f(x_1, \dots, x_n)$  in the domain  $R^n$  is written in the form:

$$f(x) = f_0 + \sum_{i=1}^n f_i(x_i) + \sum_{1 \leq i < j \leq n} f_{ij}(x_i, x_j) + \dots + f_{12\dots n}(x_1, x_2, \dots, x_n) \quad (1)$$

where  $f_0$  denotes the mean effect (or zeroth-order term), which is a constant. The function  $f_i(x_i)$  is a first-order term (or first-order component function) giving the effect of parameter  $x_i$  acting independently (although generally nonlinearly) upon the output  $f(x)$ . What this component function shows is the independent influence of parameter  $x_i$  on the output right across the domain of uncertainty for  $x_i$ . The function  $f_{ij}(x_i, x_j)$  is a second-order term describing the co-operative effects of the parameters  $x_i$  and  $x_j$  upon the output  $f(x)$ . The higher-order terms reflect the co-operative effects of increasing numbers of input parameters acting together to influence the output  $f(x)$ .



Due to its formulation as a set of hierarchical component functions, the HDMR expansion provides the possibility to determine sensitivity indices for each of the input parameters in an automatic way for selected model outputs. For given input parameter ranges, these indices indicate the relative contribution of each parameter to the predicted output variance. Thus they can be directly used to rank the importance of each individual parameter in determining the model output variance and to explore parameter interactions. The HDMR expansion is computationally very efficient if higher order input parameter interactions are weak and can therefore be neglected. For many systems a HDMR expression up to second-order already provides satisfactory results and a good approximation of  $f(x)$  (e.g. ref. 28).

In RS-HDMR, a number of model simulations are performed using a quasi-random set of input samples. This set of model simulations is then used to fit polynomial expressions for each component function in eqn (1). The sensitivity coefficients for individual parameters or for interaction terms can then be easily calculated from the coefficients of the polynomial expansion (see Ziehn and Tomlin<sup>39</sup> for details). For the current studies, the 26 dimensional input space is sampled 512 times using a quasi-random approach from uniform distributions within the parameter ranges specified. The RS-HDMR meta-model fit is then generated where the output function  $f(x)$  represents the NO<sub>2</sub> concentration at the 8 in-street locations discussed above. In practice, the larger the sample size, the better the fitted representation of the component functions and therefore the more accurate the sensitivity indices will be. In reality sample size is often limited by computational resources, particularly for complex high resolution models with substantial individual run times. When using a limited sample size such as used here, it is very important to assess the accuracy of the functional fit to eqn (1) and the GUI-HDMR code provides the facility to do this. Previous applications of the method in chemical kinetics problems<sup>40</sup> has shown that large sample sizes (>1000) are usually only needed where significant second-order terms are present (*i.e.* important parameter interactions exist).

## C Results and discussion

### (i) Overall uncertainty of predicted [NO<sub>2</sub>] and accuracy of HDMR fits

The predicted [NO<sub>2</sub>] distributions for sites G3 and G4 are shown in Fig. 3 for the off-peak case study. The most frequently predicted [NO<sub>2</sub>] at G3 is around  $1 \times 10^{12}$  molecules cm<sup>-3</sup> which equates to  $\sim 76 \mu\text{g m}^{-3}$  and is twice that at G4 confirming the influence of the in-street recirculation on the concentration profiles across the canyon. Direct comparison with high time resolution measured data is not possible but data from the diffusion tube network operated by the City of York Council offers a qualitative comparison.<sup>19</sup> In 2013, their measurement site in Gillygate opposite Portland street (close to G3, see Fig. 1) gave an annual average for [NO<sub>2</sub>] of  $48.4 \mu\text{g m}^{-3}$  compared to only  $21 \mu\text{g m}^{-3}$  at various sites along Portland street itself. It is clear therefore that due to the in-street dispersion processes large variations in NO<sub>2</sub> concentration can be seen over quite short distances. The distributions show that, given the uncertainties adopted for the input parameters, the predicted [NO<sub>2</sub>] on both sides of the street can vary by up to a factor of 2. This suggests that more accurate parameterisations of the inputs are necessary to improve the robustness of the model predictions.



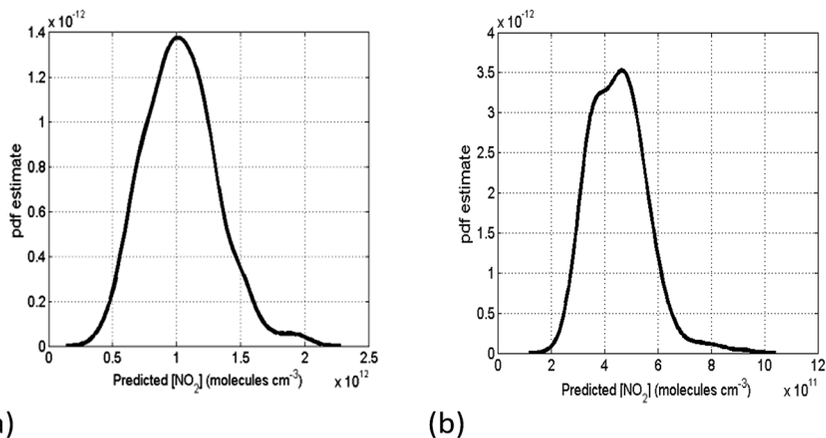


Fig. 3 Predicted  $[\text{NO}_2]$  distributions for sites (a) G3 and (b) G4 based on a quasi-random sample of 512 runs and the uncertainty limits described in Section B.

By decomposing the predictive variance into contributions from each of the uncertain inputs, the global sensitivity methodology allows us to target such model improvement strategies on the most important parameters. However, in order to exploit the HDMR component functions for sensitivity analysis purposes, it is first important to establish that the HDMR meta-model gives a reasonable fit to the outputs from the full model runs. This test is especially important for the current example since the combined model simulation time was of the order of an hour and therefore the sample size of 512 was limited by available computer resources. The coefficient of determination or  $R^2$  values comparing the fitted second-order HDMR meta-model with the data from the full model simulations vary between 0.905 and 0.951 illustrating that the second-order meta-model gives a good fit despite the limited sample size. This provides confidence in the accuracy of the HDMR component functions and the sensitivity results derived from them. The percentage of total variance accounted for by first-order effects ranges from 79–94% for the different sites indicating that the variance is dominated by sensitivities to individual parameters, and that the effects of parameter interactions are quite small. Where second- or higher-order effects do exist they tend to lead to tails in the predicted output distributions which can be seen to some extent in Fig. 3.

## (ii) Sensitivity coefficients – overall parameter importance

**Turbulent flow parameterisations.** The sensitivity coefficients calculated using HDMR provide the relative influence of each parameter on the variance within distributions such as that shown in Fig. 3. They are scaled between 0 and 1, where 1 represents 100% contribution to the output variance. Fig. 4 presents the average sensitivity coefficients across the 8 sample locations for predicted road-side time-averaged  $[\text{NO}_2]$ . The mixing time-scale coefficient  $\alpha$  and other parameters with low importance are not shown on the plot. The lack of sensitivity to  $\alpha$  is important and shows that the mean concentration predictions are not greatly affected by the mixing time-scale and that the simple IEM mixing model is valid in this context. The Lagrangian structure function coefficient  $c_0$  also exhibits a low sensitivity. This



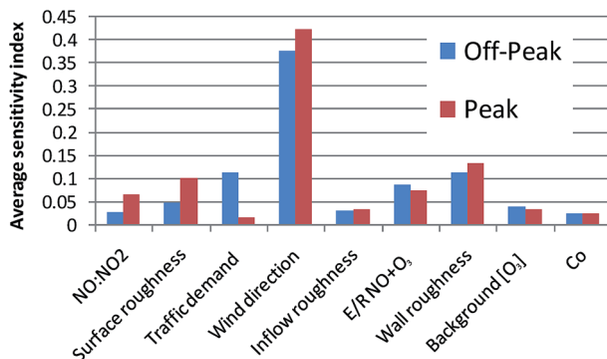


Fig. 4 The average first-order sensitivity coefficients across all 8 sites for the peak and off-peak normalised traffic demand studies.

is an encouraging result and suggests that the simulated concentrations are not highly sensitive to the chosen turbulence model parameterisations. There are however, sensitivities to the parameterisation of the velocity profiles at the model surfaces (*i.e.* to the chosen values of roughness lengths  $z_0$ ). The lowest sensitivity is to the inflow roughness, which is used to define the incoming logarithmic velocity profile at the boundary of the domain. The low sensitivity indicates that the computational domain was large enough to reduce the influence of the inflow boundary conditions on the overall dispersion profiles. The wall roughness however, exhibits a mean sensitivity of over 10%. Detailed calculations show that for site G3, this can be as high as 30%. This suggests that predicted  $[\text{NO}_2]$  close to the street canyon walls (within 2 m in the case of G3) can be highly sensitive to the near wall flow parameterisations as was previously suggested to be the case by Benson *et al.*<sup>28</sup> for velocity and turbulence fields in these locations. We will return to this point in the next section where sensitivity coefficients for individual sites are discussed in more detail.

**Physical and chemical parameterisations.** Wind direction is the major physical parameter which dominates the prediction of  $[\text{NO}_2]$  at all locations. On average it accounts for around 40% of the variance in predicted  $[\text{NO}_2]$ . This implies that a reliable reference measurement of background wind conditions is an essential input for air quality modelling systems, particularly those aiming to represent recirculating flows within urban street canyons and to assess road-side exposure. The sensitivity of predicted mean  $[\text{NO}_2]$  to background  $[\text{O}_3]$  is on average quite low, which may be surprising given that the main formation route for  $\text{NO}_2$  on short time-scales that are comparable with the in-street recirculation times, is *via* the reaction of  $\text{NO}$  with  $\text{O}_3$ . However, a higher sensitivity to the activation energy for the reaction of  $\text{NO}$  with  $\text{O}_3$  is seen. This suggests that there is a higher sensitivity to the rate of this short time-scale reaction occurring within the street than to the amount of ozone mixing into the canyon due to longer range processes. This, coupled with the importance of wind direction, indicates that if we wish models to capture exposures at the road-side, then detailed models of flow, reaction and dispersion on very short spatial scales are necessary. More accurate parameterisation of the temperature dependence of the reaction  $\text{NO} + \text{O}_3$ , using for example, *ab initio* calculations, would be critical to improving model robustness.



In terms of the influence of traffic characteristics, there are clear differences between the two modelled demand scenarios. For off-peak conditions, there is clearly a response to the levels of traffic demand with an average contribution of  $\sim 11\%$  to the predicted  $[\text{NO}_2]$ . Under peak conditions where the network saturates, road-side  $[\text{NO}_2]$  is relatively insensitive to changes in demand and more sensitive to the primary  $\text{NO}_2$  fraction (see Fig. 4). The results show that in order to have a substantial effect on road-side  $\text{NO}_2$  through traffic demand reduction measures, it would be necessary to reduce demand to the lower end of the sensitive region, *i.e.* by 60% or more.

### (iii) Sensitivity coefficients – site to site variation

In this section we focus on the off-peak traffic demand scenario and investigate the responses to changes in selected parameters at individual sites. First we compare sites G3 and G4 which lie on opposite sides of the canyon close to the signalled junction on the leeward and windward sides of the canyon, respectively. Fig. 3 shows that G3 displayed the higher  $\text{NO}_2$  concentrations due to the in-canyon helical circulation present. At G3 the highest sensitivity ( $S_i = 0.3$ ) is to the wall roughness length  $z_0$ , with the response shown in Fig. 5a. The greater the assumed roughness length, the higher the predicted  $[\text{NO}_2]$ , since a higher roughness leads to lower in-canyon flow velocities as the boundary is approached. In these figures the red line displays the first-order component function for the selected parameter (see eqn (1)). Were the response to depend only on a single parameter with an  $S_i$  of 1, then all points would lie on the red line. In reality, several parameters contribute to the output variance and hence the scatter points in the plots represent the effects of the other important parameters. For the wall  $z_0$ , the range spanned by the component function for this parameter (shown in red) is about  $5 \times 10^{11}$  molecules  $\text{cm}^{-3}$  *i.e.* almost  $40 \mu\text{g m}^{-3}$  and suggests that the near wall flow is highly influential on pollutant exposure within street canyon type geometries. Depending on the type of wall, windows, street furniture *etc.* the  $z_0$

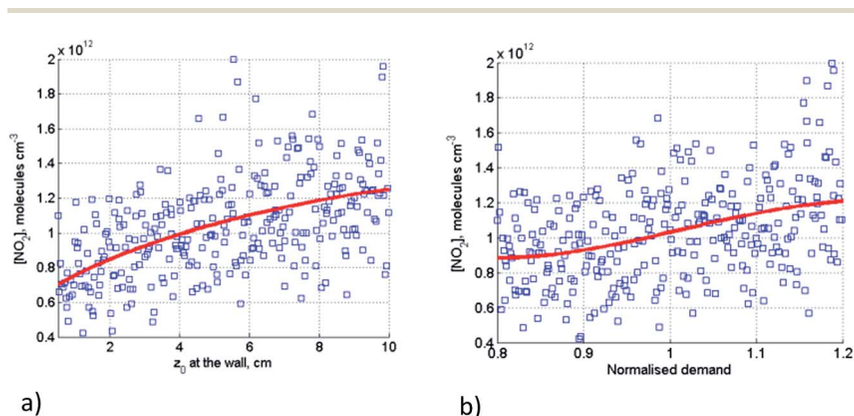


Fig. 5 Scatter plots and first-order component functions highlighting the response to changes in individual parameters on time-averaged  $[\text{NO}_2]$  at receptor G3 (a) wall roughness length  $z_0$  (b) normalised traffic demand. The scatter points show the overall response based on a random sample size of 512. The red lines show the first-order component functions *i.e.* the individual response to each parameter.



value may vary significantly from site to site and is a difficult parameter to estimate causing problems for improving model robustness.

The second most influential parameter is traffic demand ( $S_i = 0.15$ ) as shown in Fig. 5b. Changes in demand of 40% lead to a range of  $\text{NO}_2$  concentrations of a width of about  $3 \times 10^{11}$  molecules  $\text{cm}^{-3}$  *i.e.*  $23 \mu\text{g m}^{-3}$ . This illustrates that within the off-peak flow regime, reductions in demand could lead to substantial reductions in road-side mean  $[\text{NO}_2]$  levels. The figure shows that as the demand levels increase, the slope of the sensitivity decreases which is consistent with the fact that overall emissions flatten off under highly congested conditions. The payoff is therefore greater for the lower demand scenarios.

Lower  $[\text{NO}_2]$  concentrations are seen at G4 since the flow has circulated around the canyon before reaching this receptor point and hence primary emissions of  $\text{NO}_x$  have dispersed to a certain extent. At G4 the most dominant parameter is now the normalised demand ( $S_i = 0.25$ ) which suggests that the model is relatively robust to the flow and turbulence parameterisations at this receptor. Chemical processes are, however, more dominant at this windward location than they were at G3. The activation energy for reaction  $\text{NO} + \text{O}_3 = \text{NO}_2 + \text{O}_2$  has a high sensitivity ( $S_i = 0.24$ ) at this site, as does the background ozone concentration ( $S_i = 0.08$ ). The component functions and scatter plots for these two parameters are shown in Fig. 6a and b respectively. As the activation energy is lowered, the production of  $\text{NO}_2$  across the canyon increases and assumed uncertainties in this parameter can account for differences in predicted  $[\text{NO}_2]$  of around  $15 \mu\text{g m}^{-3}$  based on the predicted range of the component function (shown in red). The influence of background  $[\text{O}_3]$  is about half as strong based on the assumed uncertainties.

Wind direction was not highly influential at sites G3 and G4 since the most important dispersion process is the recirculating flow which is driven by a strong cross-street wind component for all reference directions tested. However, as we move down the canyon away from the congested junction, then it begins to play a more and more dominant role as it affects the along street dispersion of emissions from the heavily trafficked junction. At sites A and B, 60 and 40 m to the

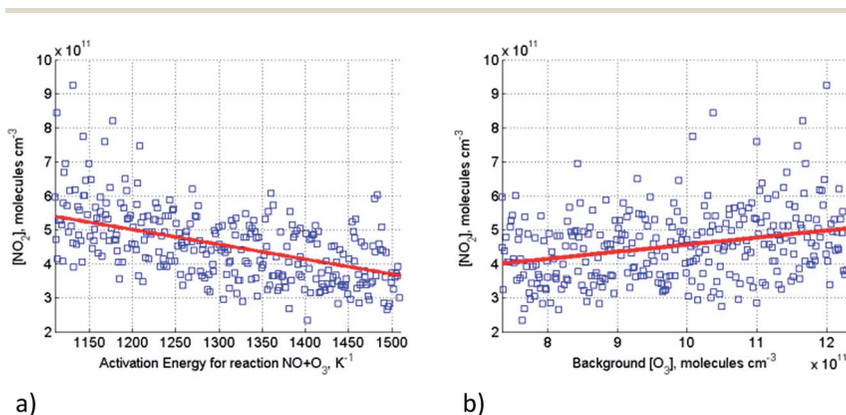


Fig. 6 Scatter plots and first-order component functions highlighting the response to changes in individual parameters on time-averaged  $[\text{NO}_2]$  at receptor G4, (a) activation energy for  $\text{NO} + \text{O}_3$  reaction and (b) background ozone concentration. The scatter points show the overall response based on a random sample size of 512. The red lines show the first-order component functions *i.e.* the individual response to each parameter.



South of G4 respectively, the sensitivity coefficients with respect to wind angle are  $>0.7$ , and the response to background wind direction is highly non-linear as shown in Fig. 7 for site A. The complex interplay between the topology of the street network and the flow patterns established means that the concentrations of  $\text{NO}_2$  at site A are mostly dominated by primary emissions, and therefore as the wind direction takes on an increasingly southerly component, the concentration decreases. The relative importance of in-street chemical processes will be discussed further in the next section.

#### (iv) Discussion of chemical processes

In-street  $\text{NO}_2$  concentration increments above background will be affected by both primary  $\text{NO}_2$  emissions being dispersed from vehicle exhausts, and the formation of secondary  $\text{NO}_2$  from primary  $\text{NO}$  through its reaction with ozone which is mixing into the street from the background. The relative importance of these processes will vary with location within the street and can be explored *via* analysis of the sensitivity coefficients. In this section we therefore also consider the sensitivity of in-street time-averaged ozone concentrations ( $[\text{O}_3]$ ) to the model parameterisations.

The main first-order sensitivity indices for sites G3 (leeward close to junction), G4 (windward close to junction), A (windward 60 m South of G4), and D (leeward 60 m South of G3) are shown in Fig. 8 for both ozone and  $\text{NO}_2$  concentrations. In-street ozone is mainly controlled by the background concentration except for at site G3 where high concentrations of  $\text{NO}_x$  are expected due to its close proximity to queuing traffic and the in-street recirculation patterns. However, this sensitivity to background ozone is not seen for  $[\text{NO}_2]$  which suggests that concentration increments of  $\text{NO}_2$  above background within congested street networks of the street canyon type are unlikely to be dominated by long range transport of ozone. Instead, in-street  $\text{NO}_2$  tends to be dominated by the effects of wind direction, traffic demand, local secondary  $\text{NO}_2$  formation rates, and near wall velocity

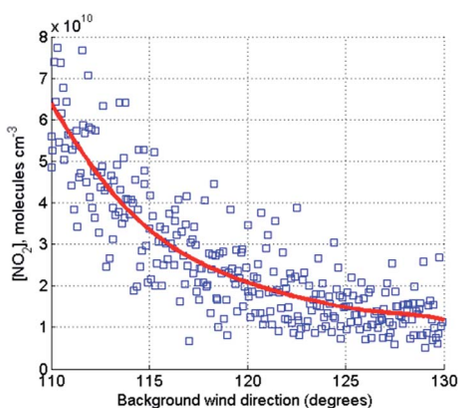


Fig. 7 Scatter plot and first-order component function highlighting the response to changes in reference wind direction on time-averaged  $[\text{NO}_2]$  at a receptor 60 m to the South of G4. The scatter points show the overall response based on a random sample size of 512. The red lines show the first-order component functions *i.e.* the individual response to each parameter.



profiles governed by roughness parameterisations. The importance of local  $\text{NO}_2$  formation is indicated by the importance of the rate parameters for the inorganic chemical reactions with the activation energy for  $\text{NO} + \text{O}_3$  being the dominant parameter. This parameter is particularly influential at sites G3 and G4 close to the junction where  $\text{NO}_x$  concentrations are high. It is also the dominant parameter affecting in-street  $[\text{O}_3]$  at site G3 since a higher rate for this reaction increases the titration of ozone by  $\text{NO}$ .

### (v) Uncertainties in model structure and issues of “uniqueness of place”

The approach for uncertainty assessment used here is parameter based and therefore assumes that the chosen model structure is appropriate for the intended application. It assesses uncertainties in prediction due to lack of knowledge of parameter values but not due to missing processes within the model. There are a number of challenges in choosing an appropriate model structure. The first is computational cost. It is unlikely that a high resolution CFD model that resolves all eddy structures formed within the complex geometry (*e.g.* using LES or direct numerical simulation), as well as all possible chemical transformations, can be used within the context of compliance assessment due to computational requirements. Even such highly detailed models require accurate boundary and initial conditions. At the other end of the spectrum, semi-empirical models such as those based on a Gaussian or multi-compartment approaches are based on parameterisations informed by street topologies, rather than detailed representations of in-street flow characteristics. The use of parameters based on geometric features such as canyon width to height ratios has provided a way of generalising the use of such models within different city geometries. However, site specific

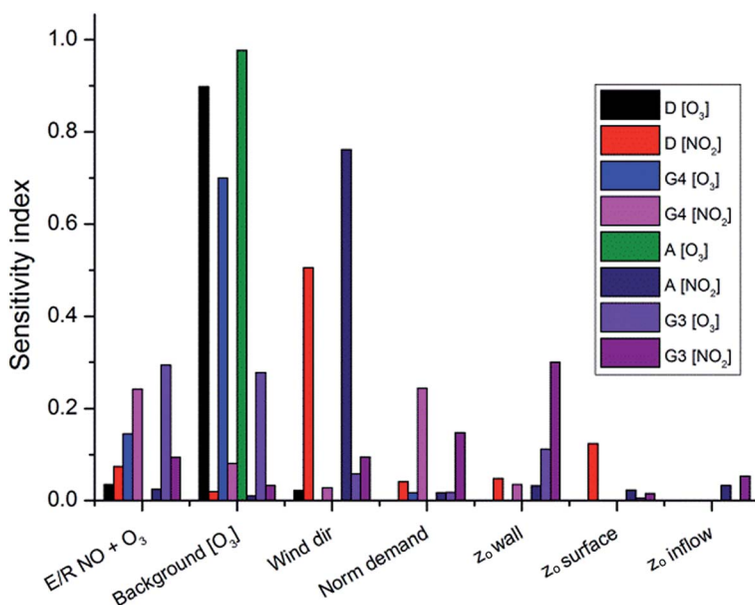


Fig. 8 Main first-order sensitivity indices for sites G3, G4, A and D. An index of 1 implies that a single parameter totally determines the variance in predicted concentrations.



features such as complex intersections and roof shapes cannot be captured by such models and yet the DAPPLE (Dispersion of Air Pollution and its Penetration into the Local Environment) field campaigns highlighted the importance of a number of these features in determining in-street flow and dispersion.<sup>41,42</sup> Such models then may contain structural uncertainties that limit their generality. This raises the issue of “uniqueness of place” for models that has been previously highlighted within the field of hydrological modelling.<sup>43</sup> LES models may require less site specific parameterisation, but may be unaffordable from a computational perspective.

The RANs CFD approach adopted in this work attempts to seek the middle ground, but each simulation still takes of the order of 10 minutes to an hour on a standard PC, and hence the use of such models is difficult within the context of rapid screening. These models, however, can capture the average effects of flows within complex geometries that lead to spatial variation in pollutant concentrations over small scales. A key question that remains is whether uncertainties within their parameterisations can be reduced significantly or whether they still suffer from issues of “uniqueness of place”, where parameterisations are highly site specific and difficult to quantify. Background wind direction ranked highly amongst the sensitive parameters affecting in-street  $[\text{NO}_2]$ . In principle, above-roof measurements could be obtained for all sites of interest but in reality this would be practically difficult. The need for a reference site free of interference from local flow features was discussed by Barlow *et al.* for the case of London.<sup>44</sup> Roughness lengths were also important, in particular those of the nearest surface to the receptor site studied. Roughness lengths cannot be measured directly and whilst approaches have been recently suggested to estimate these for complex city-scale surfaces,<sup>45</sup> in reality canyon wall roughness lengths will always have to be estimated and may be site specific.

## D Conclusions

A global uncertainty and sensitivity study was used to investigate the robustness of predictions of road-side time averaged  $\text{NO}_2$  concentrations based on a Reynolds averaged flow model coupled with a reactive Lagrangian particle dispersion model using  $\text{NO}_x$  emission rates calculated from a traffic micro-simulation model. Overall, predicted  $[\text{NO}_2]$  at a number of in-street sites was uncertain to within approximately a factor of 2. The predictions were fairly robust with respect to the parameterisations of turbulent dispersion and mixing time-scales. Uncertainties in the parameterisation of these processes did not greatly affect the time-averaged predictions. At near wall sites close to the high emission traffic junction, however, predictions were strongly influenced by the parameterisation of near boundary flows determined by the wall roughness length  $z_0$ . This is a difficult parameter to estimate but could strongly influence the prediction of in-street exposure within narrow street canyons such as the one studied here. Further away from the traffic junction, reference wind direction was the most dominant parameter affecting predicted  $[\text{NO}_2]$ . Whilst this parameter could be measured above-roof, it is not a routine measurement that is carried out within UK urban areas. Rather, reference meteorological measurements are commonly obtained from nearby rural sites or local airports. The direction of above-roof winds has been shown to strongly affect in-street flow and dispersion processes<sup>5,44</sup> and thus



it is not surprising that it influences the transport of  $\text{NO}_x$  emissions from the congested junction to other in-street receptors in this case. However, obtaining long term appropriate reference measurements is a difficult challenge for most cities.

With respect to chemical processes, it is the short time-scale, in-street processes that seem to most strongly affect road-side  $[\text{NO}_2]$ . In particular the activation energy for the reaction of  $\text{NO}$  with  $\text{O}_3$  is the most critical parameter. Better quantification of the rate of this reaction would help to improve model robustness.

In terms of mitigation strategies, two patterns emerge. Under peak traffic conditions the model suggests that even moderate reductions in traffic demand are unlikely to reduce in-street  $[\text{NO}_2]$ . For such congested conditions, the fraction of  $\text{NO}_2$  in total  $\text{NO}_x$  was shown to be more influential than demand, indicating that reducing primary  $\text{NO}_2$  could be a key factor in reducing  $\text{NO}_2$  in congested street canyon situations. For lower demand scenarios, there were demonstrable benefits to reducing traffic demand, mainly at sites close to the traffic queue. Overall, despite the scatter in the data due to uncertainties within the model parameterisations, there were clearly discernible effects of possible demand or emissions management measures suggesting that the model set up was robust enough to be useful within the context of exposure mitigation.

## Acknowledgements

This work was supported by EPSRC grant GR/S90881/01.

## References

- 1 S. K. Park, S. D. Kim and H. Lee, *Sci. Total Environ.*, 2004, **323**, 263–271.
- 2 J. W. D. Boddy, R. J. Smalley, N. S. Dixon, J. E. Tate and A. S. Tomlin, *Atmos. Environ.*, 2005, **39**, 3147–3161.
- 3 P. Louka, S. E. Belcher and R. G. Harrison, *Atmos. Environ.*, 2000, **34**, 2613–2621.
- 4 A. S. Tomlin, R. J. Smalley, J. E. Tate, J. F. Barlow, S. E. Belcher, S. J. Arnold, A. Dobre and A. Robins, *Atmos. Environ.*, 2009, **43**, 5027–5037.
- 5 A. Dobre, S. J. Arnold, R. J. Smalley, J. W. D. Boddy, J. F. Barlow, A. S. Tomlin and S. E. Belcher, *Atmos. Environ.*, 2005, **39**, 4647–4657.
- 6 R. Berkowicz, *Environ. Monit. Assess.*, 2000, **65**, 323–331.
- 7 M. Carpentieri, P. Salizzoni, A. Robins and L. Souhac, *Environ. Modell. Softw.*, 2012, **37**, 110–124.
- 8 T. A. M. Pugh, A. R. MacKenzie, J. D. Whyatt and C. N. Hewitt, *Environ. Sci. Technol.*, 2012, **46**, 7692–7699.
- 9 M. Ketzler, R. Berkowicz and A. Lohmeyer, *Environ. Monit. Assess.*, 2000, **65**, 363–370.
- 10 N. S. Dixon, J. W. D. Boddy, R. J. Smalley and A. S. Tomlin, *Atmos. Environ.*, 2006, **40**, 958–972.
- 11 J. J. Baik, Y. S. Kang and J. J. Kim, *Atmos. Environ.*, 2007, **41**, 934–949.
- 12 A. Garmory, I. S. Kim, R. E. Britter and E. Mastorakos, *Atmos. Environ.*, 2009, **43**, 4670–4680.
- 13 V. B. Bright, W. J. Bloss and X. M. Cai, *Atmos. Environ.*, 2013, **68**, 127–142.



- 14 J. Baker, H. L. Walker and X. M. Cai, *Atmos. Environ.*, 2004, **38**, 6883–6892.
- 15 J. Zhong, X. M. Cai and W. J. Bloss, *Environ. Pollut.*, 2015, **200**, 42–52.
- 16 N. S. Dixon and A. S. Tomlin, *Atmos. Environ.*, 2007, **41**, 8114–8127.
- 17 T. Ziehn, N. S. Dixon and A. S. Tomlin, *Atmos. Environ.*, 2009, **43**, 5978–5988.
- 18 J. W. D. Boddy, R. J. Smalley, P. S. Goodman, J. E. Tate, M. C. Bell and A. S. Tomlin, *Atmos. Environ.*, 2005, **39**, 3163–3176.
- 19 City of York Council, *2014 Air Quality Progress Report for City of York Council*, 2014.
- 20 J. Eichhorn and A. Kniffka, *Meteorol. Z.*, 2010, **19**, 81–90.
- 21 M. O. Letzel, T. Flassak and D. Angel, *Gefahrstoffe - Reinhalt. Luft*, 2012, **72**, 329–334.
- 22 TSS, *AIMSUN – The Integrated Transport Modelling Software*, <http://www.aimsun.com/wp/>.
- 23 F. An, M. Barth, J. Norbeck and M. Ross, *Transp. Res. Rec.*, 1997, **1587**, 52–62.
- 24 M. Zallinger, J. Tate and S. Hausberger, *16th International Transport and Air Pollution Conference*, Graz, Austria, 2008.
- 25 L. I. Panis, S. Broelcx and R. H. Liu, *Sci. Total Environ.*, 2006, **371**, 270–285.
- 26 P. S. Goodman and G. Rhys Tyler, *6th AIMSUN User Group Meeting*, Capri, Italy, 2008.
- 27 M. W. Gallagher, E. Nemitz, J. R. Dorsey, D. Fowler, M. A. Sutton, M. Flynn and J. Duyzer, *J. Geophys. Res.: Atmos.*, 2002, 107.
- 28 J. Benson, T. Ziehn, N. S. Dixon and A. S. Tomlin, *Atmos. Environ.*, 2008, **42**, 1874–1891.
- 29 B. L. Sawford, *Flow, Turbulence and Combustion*, 2004, **72**, 133–160.
- 30 D. Anfossi, G. Degrazia, E. Ferrero, S. E. Gryning, M. G. Morselli and S. T. Castelli, *Bound.-Lay. Meteorol.*, 2000, **95**, 249–270.
- 31 S. M. Du, *Bound.-Lay. Meteorol.*, 1997, **83**, 207–219.
- 32 A. M. Reynolds, *Bound.-Lay. Meteorol.*, 1998, **89**, 161–170.
- 33 R. Atkinson, D. L. Baulch, R. A. Cox, J. N. Crowley, R. F. Hampson, R. G. Hynes, M. E. Jenkin, M. J. Rossi and J. Troe, *Atmos. Chem. Phys.*, 2004, **4**, 1461–1738.
- 34 R. E. Shetter, C. A. Cantrell, K. O. Lantz, S. J. Flocke, J. J. Orlando, G. S. Tyndall, T. M. Gilpin, C. A. Fischer, S. Madronich, J. G. Calvert and W. Junkermann, *J. Geophys. Res.: Atmos.*, 1996, **101**, 14631–14641.
- 35 R. E. Shetter, A. H. McDaniel, C. A. Cantrell, S. Madronich and J. G. Calvert, *J. Geophys. Res.: Atmos.*, 1992, **97**, 10349–10359.
- 36 D. C. Carslaw, *Atmos. Environ.*, 2005, **39**, 4793–4802.
- 37 I. M. Sobol, *Math. Comput. Simulat.*, 2001, **55**, 271–280.
- 38 H. Rabitz, O. F. Alis, J. Shorter and K. Shim, *Comput. Phys. Commun.*, 1999, **117**, 11–20.
- 39 T. Ziehn and A. S. Tomlin, *Environ. Modell. Softw.*, 2009, **24**, 775–785.
- 40 T. Turányi and A. S. Tomlin, *Analysis of Kinetic Reaction Mechanisms*, Springer, 2014.
- 41 A. A. Balogun, A. S. Tomlin, C. R. Wood, J. F. Barlow, S. E. Belcher, R. J. Smalley, J. J. N. Lingard, S. J. Arnold, A. Dobre, A. G. Robins, D. Martin and D. E. Shallcross, *Bound.-Lay. Meteorol.*, 2010, **136**, 489–513.
- 42 C. R. Wood, S. J. Arnold, A. A. Balogun, J. F. Barlow, S. E. Belcher, R. E. Britter, H. Cheng, A. Dobre, J. J. N. Lingard, D. Martin, M. K. Neophytou, F. K. Petersson, A. G. Robins, D. E. Shallcross, R. J. Smalley, J. E. Tate, A. S. Tomlin and I. R. White, *Bull. Am. Meteorol. Soc.*, 2009, **90**, 955–970.



- 43 K. J. Beven, *Hydrol. Earth Syst. Sci.*, 2000, **4**, 203–213.
- 44 J. F. Barlow, A. Dobre, R. J. Smalley, S. J. Arnold, A. S. Tomlin and S. E. Belcher, *Atmos. Environ.*, 2009, **43**, 5536–5544.
- 45 J. T. Millward-Hopkins, A. S. Tomlin, L. Ma, D. B. Ingham and M. Pourkashanian, *Bound.-Lay. Meteorol.*, 2013, **146**, 447–468.

