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DOI:  
[10.1016/j.atmosenv.2012.10.056](https://doi.org/10.1016/j.atmosenv.2012.10.056)

*Document Version*  
Peer reviewed version

*Citation for published version (Harvard):*

Bright, VB, Bloss, WJ & Cai, X 2013, 'Urban street canyons: Coupling dynamics, chemistry and within-canyon chemical processing of emissions', *Atmospheric Environment*, vol. 68, pp. 127-142.  
<https://doi.org/10.1016/j.atmosenv.2012.10.056>

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# Urban street canyons: coupling dynamics, chemistry and within-canyon chemical processing of emissions

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## Abstract

Street canyons, formed by rows of buildings in urban environments, are associated with high levels of atmospheric pollutants emitted primarily from vehicles, and substantial human exposure. The street canyon forms a semi-enclosed environment, within which emissions may be entrained in a recirculatory system; chemical processing of emitted compounds alters the composition of the air vented to the overlying boundary layer, compared with the primary emissions. As the prevailing atmospheric chemistry is highly non-linear, and the canyon mixing and predominant chemical reaction timescales are comparable, the combined impacts of dynamics and chemistry must be considered to quantify these effects. Here we report a model study of the coupled impacts of dynamical and chemical processing upon the atmospheric composition in a street canyon environment, to assess the impacts upon air pollutant levels within the canyon, and to quantify the extent to which within-canyon chemical processing alters the composition of canyon outflow, in comparison to the primary emissions within the canyon. A new model for the simulation of street canyon atmospheric chemical processing has been developed, by integrating an existing Large-Eddy Simulation (LES) dynamical model of canyon atmospheric motion with a detailed chemical reaction mechanism, a Reduced Chemical Scheme (RCS) comprising 51 chemical species and 136 reactions, based upon a subset of the Master Chemical Mechanism (MCM). The combined LES-RCS model is used to investigate the combined effects of mixing and chemical processing upon air quality within an idealised street canyon. The effect of the combination of dynamical (segregation) and chemical effects is determined by comparing the outputs of the full LES-RCS canyon model with those obtained when representing the canyon as a zero-dimensional box model (i.e. assuming mixing is complete and instantaneous). The LES-RCS approach predicts lower (canyon-averaged) levels of  $\text{NO}_x$ , OH and  $\text{HO}_2$ , but higher levels of  $\text{O}_3$ , compared with the box model run under identical chemical and emissions conditions. When considering the level of chemical detail implemented, segregation effects were found to reduce the error introduced by simplifying the reaction mechanism. Chemical processing of emissions within the canyon leads to a significant increase in the  $\text{O}_x$  flux from the canyon into the overlying boundary layer, relative to primary emissions, for the idealised case considered here. These results demonstrate that within-canyon atmospheric chemical processing can substantially alter the concentrations of pollutants injected into the urban canopy layer, compared with the raw emission rates within the street canyon. The extent to which these effects occur is likely to be

dependent upon the nature of the domain (canyon aspect ratio), prevailing meteorology and emission / pollution scenario considered.

*Keywords:* Street Canyons; Air Pollution; Large-eddy simulation; NO<sub>x</sub>; Ozone

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## 1. Introduction

Urban street canyons, formed by parallel rows of buildings enclosing a vehicular roadway, represent a unique atmospheric environment. Street canyons are commonly the location of substantial primary pollutant emissions, usually dominated by traffic sources. They are also locations where substantial receptor exposure occurs, for pedestrians, road-users and occupants of adjacent buildings which may source their ventilation from the canyon environment. The semi-enclosed form of many urban street canyons can give rise to re-circulatory air flow, reducing the ventilation of the canyon to the overlying urban boundary layer, and further increasing air pollution levels, to potentially harmful levels with serious implications for public health, vegetation and the built environment (Vardoulakis et al., 2003). Atmospheric composition within street canyons is determined by a combination of the composition of background air mixed in from above the canyon, vehicle exhaust and other emissions from within the street, and the mixing and chemical processing of pollutants within the canyon. The interaction of these factors determines both the pollutant concentrations experienced within the canyon, their spatial and temporal variation, the extent to which emissions may undergo chemical processing within the canyon prior to their release into the overlying boundary layer. As the atmospheric chemical reaction system which describes the interaction between nitrogen oxides, ozone and organic compounds is highly non-linear, and the timescale of many of the chemical processes is comparable to the timescale of the large-scale canyon circulation and mixing (seconds to minutes), it is necessary to consider both dynamical and chemical effects to understand the abundance and distribution of reactive species within a street canyon. The aim of this work is to investigate this chemical - dynamical coupling on the street canyon scale, and to gain insight into the oxidative nature of such an environment to understand the potential effects upon within-canyon atmospheric composition (and hence pollutant exposure), and how the canyon outflow composition may differ from the primary emissions, as a result of within-canyon chemical processing.

The airflow characteristics of street canyon dynamics are dependent on the overlying wind direction and speed, and the canyon geometry. In the most simple case, when the direction of the prevailing wind is perpendicular to the canyon axis and the street canyon aspect ratio (defined as the ratio of canyon height ( $H$ ) to canyon width ( $W$ )) is unity ( $H/W = 1$ ), skimming flow is observed in which a large proportion of the flow does not enter the canyon and instead skims over the top of buildings (Oke, 1987). This flow regime results in the formation of a large primary canyon vortex (Fig. 1) in which recirculation effectively decouples the canyon atmosphere from the background air above,

73 resulting in reduced exchange and poor ventilation. As a result, skimming flow is relatively  
74 ineffective in removing pollutants from within the canyon (Hunter et al., 1992). Strong intermittency  
75 in the street canyon re-circulation was observed using roof level measurements by Louka et al. (2000)  
76 who concluded that the mean flow within the canyon was merely a residual of an unsteady turbulent  
77 re-circulation. It was proposed that the strong intermittency observed in the recirculation may be due  
78 to the intermittent nature of the mechanism that couples the flow within the canyon with that above  
79 i.e. the shear layer at roof level (Fig. 1). This provides an effective ventilation mechanism allowing air  
80 to escape and efficient mixing to occur when the shear layer moves to an upward position. Such  
81 events, known as sweeps and ejections, are the major turbulent processes that govern pollutant  
82 removal from street level to the background atmosphere above (Cheng and Liu, 2011).

83  
84 Primary pollutants such as nitric oxide (NO), nitrogen dioxide (NO<sub>2</sub>) and volatile organic compounds  
85 (VOCs) are frequently present in significantly higher concentrations within the street canyon when  
86 compared to the overlying background atmosphere, as expected considering proximity to (traffic)  
87 emission sources. The oxides of nitrogen (NO<sub>x</sub> = NO + NO<sub>2</sub>), released predominantly in vehicle  
88 exhaust, dominate gas phase chemistry in urban environments. In polluted urban air, levels of the  
89 hydroxyl radical (OH) are strongly dependent on chemical cycling, which in turn is dependent on  
90 levels of NO<sub>x</sub>. OH initiated VOC degradation in the presence of NO<sub>x</sub> can result in the formation of  
91 secondary pollutants such as ozone (O<sub>3</sub>). O<sub>3</sub> is of particular concern in terms of canyon atmospheric  
92 composition due to its detrimental effects on human health, vegetation, and the built environment.  
93 Within such canyons, O<sub>3</sub> levels are often reduced compared with the overlying air (and more  
94 generally, within urban regions compared with the surrounding areas) due to titration by primary NO,  
95 an effect known as the urban decrement (AQEG, 2009). Under sunlit conditions, NO<sub>x</sub> chemistry is  
96 governed by the photochemical steady state (PSS) chemistry in which NO, NO<sub>2</sub> and O<sub>3</sub> establish  
97 equilibrium. Interactions with peroxy radicals lead to additional NO-to-NO<sub>2</sub> conversion and result in  
98 the net production of O<sub>3</sub>. As such, it is valuable to assess the extent to which pollutants may be  
99 processed prior to their emission into the wider urban atmosphere, in order to gain an insight into the  
100 importance of the oxidative environment of the street canyon.

101  
102 Long lived pollutants such as carbon monoxide (CO) and volatile organic compounds (VOCs) are  
103 unlikely to show a substantial variation in concentration within a street canyon due to chemical  
104 processing alone, with their distributions governed near-exclusively by canyon dynamics. Chemical  
105 processing of NO<sub>x</sub> and O<sub>3</sub>, which have shorter chemical lifetimes, occurs on similar timescales to  
106 those of the key dynamical processes within the urban canopy layer, thereby causing these species to  
107 exhibit a marked variation within the canyon. Very short lived species, such as OH and HO<sub>2</sub>, with  
108 atmospheric lifetimes of seconds, are highly variable within the canyon and respond rapidly to  
109 changes in chemical composition on the canyon scale.

Canyon dynamics and composition have been the subject of field measurement campaigns, wind tunnel experiments and numerical modelling investigations. As road traffic represents the dominant emission source within typical street canyons, in situations where a single primary vortex forms, a large gradient of pollutant concentration is observed across the street with the highest concentration of many pollutants evident toward the leeward wall. A number of field studies have shown that the concentration of pollutants on the leeward side of the street can be significantly greater than that on the windward side and that a vertical decrease in concentration on both sides of the street is observed (Baker et al., 2004; Berkowicz et al., 2002; DePaul and Sheih, 1985; Tomlin et al., 2009; Xie et al., 2003). Although important in terms of model validation, field studies are often relatively sparse in terms of their resolution / spatial coverage and can be influenced considerably by the prevailing meteorological conditions and complex geometry of the surrounding urban environment. Reduced scale physical models (wind tunnel experiments) have also been used to study pollutant dispersion and canyon dynamics and have provided evidence for significant horizontal and vertical gradients in the concentration of passive pollutants within and above the canyon (Gromke et al., 2008; Kastner-Klein and Plate, 1999; Pavageau and Schatzmann, 1999; Salizzoni et al., 2009; Tomlin et al., 2009); however scale effects, and the high spatial and temporal resolution of observations required for full understanding, provide substantial opportunity to use numerical models to investigate canyon dynamics and composition.

Model studies focussing on high resolution representation of canyon fluid dynamics and pollutant transport/dispersion processes have utilised Reynolds-Averaged Navier-Stokes (RANS) models such as large-eddy simulation (LES) to simulate canyon turbulent flow and associated pollutant dispersion; a number of these have been reviewed by Li et al. (2006). Most research has involved the simulation of passive pollutant dispersal (Cai et al., 2008; Cheng and Liu, 2011; Li et al., 2008; Li et al., 2009; Liu and Barth, 2002; Liu et al., 2004; Salim et al., 2011; So et al., 2005). In contrast, relatively little attention has been given to modelling the dispersion of reactive pollutants with most practical applications tending to focus on the dispersion of a passive scalar, or including only a limited number of chemical reactions (Baik et al., 2007; Baker et al., 2004; Garmory et al., 2009; Wang and Mu, 2010). Baker et al. (2004) investigated the turbulent dispersion and transport of reactive pollutants within a street canyon using a large-eddy simulation model with very simple  $\text{NO}_x$ - $\text{O}_3$  titration chemistry (a two-reaction system) applied, to investigate the deviation from PSS arising from dynamical effects. Substantial deviations from the expected bulk photochemical steady state were found, which were quantified as the photostationary state defect (PSSD). The highest values of the PSSD and therefore greatest deviation from chemical equilibrium, were observed well above the canyon ( $z/H \approx 1.3$ ) corresponding the outer extent of the escaping canyon plume – i.e. where polluted canyon air meets less polluted background air flowing over the canyon. A significant variation in the

photostationary state defect within the canyon was also found to occur with the highest ‘within canyon’ values observed downwind of the emission source and toward ground level on the windward wall, due to entrainment of air by the canyon vortex. The lowest values of the passive scalar co-emitted with  $\text{NO}_x$  were observed in the centre of the canyon vortex. The same limited reaction scheme was subsequently used by Grawe et al. (2007) to investigate the effect of local shading on pollutant concentrations. Baik et al. (2007) used the same simple representation of the photochemistry to study the dispersion of reactive pollutants within the street canyon, using a RANS-based dynamical model. A similar variation in the photostationary state defect was found, with the largest deviation from chemical equilibrium found above roof level where canyon outflow meets background air, and the region closest to chemical equilibrium found to be the within the canyon vortex. The limited chemical mechanisms present in these studies however neglects the impact of more detailed atmospheric chemistry, for example, peroxy-radical mediated NO to  $\text{NO}_2$  conversion with associated  $\text{O}_3$  (or total oxidant) production, or  $\text{NO}_x$  removal through formation of reservoir compounds.

Few studies have investigated the dispersion of reactive pollutants by applying more comprehensive photochemical reaction schemes to dynamical models. Due to limitations in computing power it is not practical to include the full range of chemical species and reactions that occur in the urban atmosphere (even should all such emissions and reactions be known), particularly when combined with computationally expensive dynamical models such as LES. Explicit representation of the oxidation processes taking place within the canyon may contain several thousand chemical species and over 20,000 reactions (Dodge, 2000). Owing to the computational expense of such models it is still impractical to include *near*-explicit chemical schemes (such as the Master Chemical Mechanism – see below) as a true representation of the canyon chemistry. As a result a number of *reduced* chemical mechanisms, including those reviewed by Dodge (2000), have been developed that reasonably accurately represent the chemical environment of urban canyons, and may be effectively and affordably applied to photochemical / dynamical models.

The work of Garmory et al. (2009) used the Stochastic Fields (SF) method to simulate turbulent reacting flows and the dispersion of reactive scalars within the street canyon. This research applied the same simple chemical scheme as used by Baker et al. (2004), Grawe et al. (2007) and Baik et al. (2007), and utilised a number of statistical methods to characterise atmospheric processing within the canyon. The results of this initial study were in close agreement with those of Baker et al. and Baik et al., with lowest values of the photostationary state defect located within the canyon, and greatest values observed just above roof level, within the mixing layer. In addition to the simple scheme the more detailed Carbon Bond Mechanism (CBM-IV; Gery et al., 1989) was used. Comparing both mechanisms, the effect of segregation was quantified by evaluating the Damköhler number,  $Da$ , defined as the ratio of the mixing timescale to the chemical timescale. A value of  $Da \gg 1$  indicates

that the chemistry / dynamical interaction is important and that segregation effects must be accounted for (Garmory et al., 2006; Krol et al., 2000). If  $Da \ll 1$ , species become well mixed much more rapidly than their chemical processing timescale, and hence segregation effects are minimal. For intermediate- and long-lived species, including important species such as  $O_3$  and  $NO$ , Garmory et al. found the effect of segregation to be minimal; however for a number of the shorter lived radical species this work, using the CBM-IV mechanism, demonstrated significant differences in predicted concentration in the mixing region above the canyon when segregation effects were considered.

A six-species chemical reaction scheme was applied using both box and LES model frameworks by Krol et al. (2000), with a focus upon the larger atmospheric boundary layer scale (rather than the street canyon domain). Akin to Garmory et al. (2009), this research investigated the deviation from chemical equilibrium, as a result of the turbulent nature of the convective boundary layer, in terms of the intensity of segregation. It was found that turbulence inherent in the convective boundary layer results in large concentration fluctuations and that these give rise to a divergence from chemical equilibrium in contrast to that obtained using box model calculations. When species are emitted uniformly the volume averaged concentrations were found to deviate only slightly from the box model concentrations, however when reactive hydrocarbons were emitted non-uniformly (as is likely to occur in reality) segregation effects are increased, with volume averaged LES model results showing that the rate of destruction of reactive hydrocarbons (RH - representing all reactive hydrocarbons and intermediate species) may be reduced by up to 30 % when compared to that calculated using the box model – i.e. the box model dynamical framework substantially underestimates the chemical lifetime of emitted species within the canyon. It was also found that if both the turbulent timescale and the chemical timescale of a compound are comparable, the integrated flux of RH through the RH-OH reaction will be reduced due to the chemistry-turbulence interaction. Pugh et al. (2011) also investigated boundary layer segregation effects using field measurements taken above a tropical rainforest in South-East Asia. The effect of segregation on the reaction between OH and isoprene was determined using high temporal resolution isoprene concentration data. It was found that the reduction in the effective rate constant for the reaction of isoprene with OH due to segregation effects was typically less than 15 %; an intensity of segregation considerably lower than that needed to explain observed inconsistencies between measured and modelled OH concentrations produced by global and box models of atmospheric chemistry in isoprene rich environments (e.g. Lelieveld et al., 2008).

Most recently, Kwak & Baik (2012) reported CFD simulations of street canyon atmospheric composition incorporating the CBM-IV mechanism, and explored the sensitivity of within-canyon ozone levels to perturbations in  $NO_x$  and VOC emissions. Ozone was found to be negatively correlated with  $NO_x$ , reflecting the impact of NO titration dominating over  $NO_2$  photolysis as a source

of ozone; this was characterised as a negatively  $\text{NO}_x$ -sensitive regime. Kim et al. (2012) report comparison of CFD simulations using the chemical code from the GEOS-Chem model (Bey et al., 2001) with field observations in a street canyon in Guangzhou, China. The model successfully reproduced the observed levels of species which are essentially passive on the canyon timescale (e.g. CO), but substantially overestimated NO (factor of three). In contrast to the work of Garmory et al., Kim et al. found that modelled canyon  $\text{O}_3$  levels varied substantially with the chemical mechanism applied – in comparison with the full reaction scheme, application of a reduced mechanism ( $\text{NO}_x$ - $\text{O}_3$  photostationary steady state only) led to increases in within-canyon  $\text{O}_3$  levels of up to 150 %. The importance of within-canyon processing of emissions, the level of chemical detail necessary to satisfactorily account for such processing, and quantification of the resulting pollutant export to the urban atmosphere, remain open questions.

In the present work, we describe the development of a new model combining a validated representation of street canyon dynamics, together with a sufficiently detailed chemical mechanism to fully assess the impacts of chemistry - dynamics coupling upon atmospheric composition and atmospheric oxidation rates within a street canyon regime. This model combines an LES dynamical treatment with a newly developed and computationally affordable chemical mechanism (termed the reduced chemical scheme, RCS). The RCS mechanism itself was developed from a near-explicit chemical mechanism (the Master Chemical Mechanism; Jenkin et al., 1997), and is validated against the MCM (which in turn has been validated against field and chamber observations), under conditions and timescales relevant to the street canyon regime. In contrast to our previous work, (Baker et al., 2004; Grawe et al., 2007) the level of chemical detail incorporated in the RCS allows the within-canyon VOC oxidation, associated net ozone (or oxidant) production, and  $\text{NO}_x$  removal through conversion to reservoir species, to be evaluated – all processes well known to be of importance in the real atmosphere, but which the more simple chemical mechanisms cannot address. We apply the combined LES-RCS model to evaluate impacts of the combined chemical and dynamical processes upon within-canyon atmospheric composition and its variability (and hence pollutant exposure), and to evaluate how the canyon outflow differs from the primary emissions within the canyon, as a consequence of within-canyon chemical processing. The effects of chemistry and dynamics are separated by comparing the canyon-average results from full LES-RCS simulations, with those performed under identical chemical conditions, using a box model alone (which corresponds to uniform and instantaneous mixing). A representative flux of processed pollutants out of the street canyon into the overlying boundary layer is determined.

## **2. Development of the reduced chemical scheme (RCS)**

### *2.1 Chemical Mechanism*



In order to develop a suitable model to study street canyon atmospheric composition a representation of the chemistry to be applied within the dynamical model is required. To achieve this, a zero-dimensional box model was used as an efficient tool to develop a chemical reaction mechanism which was sufficiently computationally affordable when implemented within the (computationally expensive) LES dynamical model. The most accurate representation of gas-phase tropospheric chemistry can be achieved through the use of near-explicit chemical mechanisms such as the Master Chemical Mechanism (MCM) (Jenkin et al., 1997) or the Generator for Explicit Chemistry and Kinetics of Organics in the Atmosphere (GECKO-A) (Aumont et al., 2005). The MCM v3.1 describes the degradation of 135 volatile organic compounds (VOCs) including the major UK anthropogenic emissions and biogenic species including isoprene and the monoterpenes  $\alpha$ - and  $\beta$ -pinene, including over 5,900 chemical species and 13,500 chemical reactions (Bloss et al., 2005; Jenkin et al., 1997; Jenkin et al., 2003; Saunders et al., 2003). The MCM has been evaluated using high quality datasets obtained from experiments carried out in large outdoor environmental reaction chambers and ambient field observations. As the MCM is too computationally expensive to incorporate directly into the LES framework, a subset of the MCM, the Common Representative Intermediates mechanism version CRI v2-R5 (Jenkin et al., 2008; Watson et al., 2008), which includes 19 emitted anthropogenic VOCs to represent full speciation, 196 chemical species and 555 reactions, was used as a starting point for further scheme reduction. The reduced chemical scheme derived from this (hereafter referred to as the RCS) was then compared with the full MCM simulation, the latter providing the standard for RCS evaluation. The reduced form of the CRI has itself been evaluated in comparison to the MCM and other mechanisms, and found to replicate well both integrated ozone production on timescales of days, and (of more relevance here) OH levels on timescales of hours under polluted (industrial) conditions (Emmerson and Evans, 2009).

## 2.2 Mechanism Development

In order to reduce the CRI v2-R5, the scope of simulations was initially limited to daytime scenarios, allowing night time only chemistry to be removed (parent VOC degradation by  $\text{NO}_3$  and inorganic  $\text{NO}_3/\text{NO}_y$  reactions were retained). Further reduction was achieved by eliminating parent compounds, and any unique daughter products, which had little effect on the key chemical intermediates under street canyon conditions, while scaling the abundance of other (parent) compounds to retain the same OH reactivity. The metrics used to assess each simplification were concentrations of OH, NO,  $\text{NO}_2$ ,  $\text{O}_3$  and hydroperoxy radical ( $\text{HO}_2$ ), the key criterion being to maintain OH levels within 10 % of those predicted by the (full) MCM using the (full) parent VOC set. The explicit inorganic chemistry from the MCM was fully retained. The initial concentrations of pollutants used in developing the reduced chemical scheme were determined using observations taken during the Tropospheric ORganic CHemistry experiment (TORCH) field campaign carried out in suburban London (Lee et al., 2006), with the  $\text{NO}_x$  range extended to cover a range from 15 and 513 ppb, to better represent the conditions

that may occur within a street canyon. The number of parent VOCs considered was reduced, using VOC reactivity (with respect to OH) as a proxy for ozone production potential on the short timescales relevant to the street canyon residence time. Physically the OH reactivity of a VOC represents the inverse of the lifetime of OH due to loss by reaction with that species; OH reactivity provides a suitable measure of the overall potential for VOC oxidation and subsequent formation of organic peroxy radicals and hence O<sub>3</sub> formation. The remaining parent VOC abundance was adjusted to maintain the total OH reactivity (Equation 1), at the (observed) value of 3.4 s<sup>-1</sup> (Lee et al., 2006).

$$k'_{OH} = \sum k_{1(OH+VOC_1)}[VOC]_1 + k_{2(OH+VOC_2)}[VOC]_2 + k_{3(OH+VOC_3)}[VOC]_3 \dots \quad (1)$$

The final RCS includes 51 chemical species and 136 reactions; with methane and 8 parent non-methane hydrocarbons (NMHCs) included in the mechanism (Table 1: isoprene, ethene, propene, formaldehyde, acetaldehyde, methanol, ethanol and peroxyacetylnitrate).

### 2.3 Evaluation of the RCS

The accuracy of the RCS was assessed in comparison to the evolution of species concentrations calculated using the full MCM for a single air parcel, initialised using the starting concentrations listed in Table 1. Over a four hour period, the maximum percentage difference in OH between the RCS and the MCM was approximately 6 %, which is within the bounds of the smallest errors associated with the measurement of OH (7 – 16 %) (Heard and Pilling, 2003). For NO, NO<sub>2</sub>, O<sub>3</sub> and HO<sub>2</sub> the largest differences between the RCS and the MCM, which occur toward the end of the four hour time period, are 15 %, 7 %, 4 % and 14 % respectively. At the 30 minute time point, more relevant to canyon residence times, smaller differences of 0.4 %, 0.1 %, 0.2 %, 1.1 % were observed respectively, with 0.7 % for OH. The RCS and MCM were also compared under elevated NO<sub>x</sub> conditions (NO = 1000 ppb and NO<sub>2</sub> = 120 ppb) which may be experienced within canyons due to proximity to vehicle exhausts. The maximum differences observed over a four hour period were 3 %, 13 %, 16 %, and 4 % for NO, NO<sub>2</sub>, O<sub>3</sub> and HO<sub>2</sub> respectively with 12 % for OH. At  $t = 30$  minutes the modelled differences were 0.3 %, 1.7 %, 2.1 %, 3.0 % and 2.4 % for NO, NO<sub>2</sub>, O<sub>3</sub>, HO<sub>2</sub> and OH respectively. These values are significantly smaller than the uncertainty associated with emissions and with the measurement of such pollutants (Boulter et al., 2009; Lee et al., 2006).

### 3. Configuration of models used to simulate urban street canyon composition

Two principal approaches are applied here to simulate atmospheric composition and pollutant processing in street canyons. The primary tool is the full LES-RCS model, which represents the most comprehensive treatment of the combined effects of dynamics, chemistry and emissions; the LES-RCS domain included both the street canyon and the overlying boundary layer (described below, section 3.1). The LES-RCS output was compared with simulations in which the street canyon was approximated by a zero-dimensional box model (“Box-RCS”), corresponding to a scenario in which

concentrations are homogeneous (*i.e.* the volume of air is assumed to be instantaneously and completely mixed). In this scenario, the box model corresponds to the within-canyon region ( $z/H \leq 1$ ), and exchange with the overlying boundary layer was parameterised using an exchange velocity expression, as described below.

### 3.1 Configuration of the LES-RCS model

The LES dynamical model was used to simulate atmospheric motion and turbulent flow in and above an idealised street canyon with an aspect (height/width) ratio of one. The model is based on the Regional Atmospheric Modelling System (RAMS), described in more detail in Cui *et al.* (2004). The prevailing wind direction in the model remains constant and perpendicular to the canyon axis, which is representative of a worst case scenario from the perspective of pollutant accumulation, in which canyon ventilation is minimal. The wind speed was initially set to zero below the roof level and increased logarithmically to a maximum speed,  $U_{max}$ , of  $2.5 \text{ m s}^{-1}$  at the top of the domain. Mesh resolution in the  $x$ ,  $y$  directions were  $\Delta x = 0.3 \text{ m}$ ,  $\Delta y = 1.0 \text{ m}$  respectively. In the  $z$  direction,  $\Delta z = 0.3 \text{ m}$  within the canyon and gradually stretched by a factor of 1.15 above roof level ( $z = 18.0 \text{ m}$ ) to a maximum of  $5.0 \text{ m}$  at the top of the domain ( $z = 94 \text{ m}$ ). Cyclic boundary conditions were applied to all three velocity components along  $x$ - and  $y$ -directions. The temperature was defined as  $293 \text{ K}$  throughout the whole model domain, representative of neutral conditions. Fig. 2a illustrates the street canyon domain included in the LES model, while Fig. 2b shows the developed flow field within and above the canyon. The dynamical part of the LES model for a canyon with an aspect ratio  $H/W$  of 1 has been validated by Cui *et al.* (2004), through comparisons of the mean wind and resolved-scale turbulent kinetic energy (TKE) against wind-tunnel experiments, while the scalar part of the model under various  $H/W$  ratios has been validated by Cai *et al.* (2008), in which the normalized fluxes were compared with wind-tunnel measurements.

In each experiment, the model was run without chemistry for 30 minutes in order for the turbulent dynamics to reach a quasi-equilibrium state (Cai *et al.*, 2008). At this time, concentrations of all 51 chemical species (initial concentration as listed in Table 1; intermediate and product concentrations as calculated from a 30-minute model integration) were inserted in the whole model domain uniformly. This set of values was also used as the inlet boundary conditions above the upwind building

throughout the simulation. At the outlet boundary, the advective condition,  $\frac{\partial c_i}{\partial t} + u \frac{\partial c_i}{\partial x} = 0$ , was applied to all chemical species, where  $c_i$  represents the concentration of species,  $i$  and  $u$  is horizontal velocity. For the intermediate- and longer-lived chemical species included in the chemical mechanism, for example,  $\text{NO}$ ,  $\text{NO}_2$ ,  $\text{O}_3$  and  $\text{CO}$ , a timestep of  $10^{-2} \text{ s}$  was used in the numerical integration, while for shorter lived species such as  $\text{OH}$ ,  $\text{HO}_2$  and  $\text{RO}_2$  a timestep of  $10^{-3} \text{ s}$  was

employed. These timestep values were empirically chosen to balance the requirement for stable output / convergence against integration time.

Emissions within the canyon were represented by two line sources centred at 2.5 m to the left and right of the canyon centre, signifying two lanes of traffic. Each of the line sources was considered to have a Gaussian distribution (where  $\sigma_x = 3$  m and  $\sigma_y = 1$  m), which were located at 1.0 m above the road as illustrated in Figure 1, and were continuous in nature. The emission rates included in the model were determined using the UK Road Vehicle Emission Factors (2009 - Boulter et al., 2009) and are representative of moderate weekday traffic (1500 vehicles per hour) for an urban road with cars travelling at an average speed of 30 mph. The total emissions for NO, NO<sub>2</sub>, CO, ethene (C<sub>2</sub>H<sub>4</sub>), propene (C<sub>3</sub>H<sub>6</sub>), formaldehyde (HCHO) and acetaldehyde (CH<sub>3</sub>CHO) used were 101, 17, 377, 36, 24, 11 and 16  $\mu\text{g m}^{-1} \text{s}^{-1}$ , which equate to 900, 100, 3593, 347, 150, 96 and 98 ppb emitted into one LES model cell (0.3 m  $\times$  0.3 m  $\times$  1 m) per second, respectively. Photolysis frequencies were calculated offline using the Tropospheric Ultraviolet and Visible (TUV) Radiation Model v4.1 (Madronich and Flocke, 1998), using conditions representative of those for a street canyon in Birmingham, UK (52° 29' N; -1° 54' W) at 12.00 UTC on 1<sup>st</sup> August, giving a calculated solar zenith angle of 34°. The ground elevation was specified as 0.12 km and the atmosphere was assumed to be cloud free. A surface albedo of 0.15 was defined i.e. typical of an urban street surface (DFT, 2009; Liu et al., 2011). Photolysis rates were held constant over the modelling period in order to reduce the computational expense once implemented in the LES, reflecting the short residence time (minutes) of a typical street canyon air parcel (Baker et al., 2004; Vardoulakis et al., 2003). Following the initiation of the emissions and chemistry, the combined effects of emission, mixing and chemical processing on atmospheric composition could then be simulated by the LES-RCS model, and the results examined to provide an insight into the processes affecting atmospheric composition over a 180 minute period.

Using the LES-RCS model, 3-D data was obtained over the period from 30 to 210 minutes at 5 s time intervals. Results were averaged along the y-axis over the length of the canyon ( $L_y$  - along which the resolved-scale turbulence is homogeneous), across the width of the canyon (the x-axis;  $W$ ) and over the height of the canyon (the z-axis;  $H$ ) to give a volume averaged (0-D) within-canyon concentration as a function of time, i.e.:

$$\overline{c_i}(t) = \frac{1}{W \cdot H \cdot L_y} \int_{-0.5W}^{0.5W} \int_0^H \int_0^{L_y} c_i(x, y, z, t) dx dy dz. \quad (2)$$

The LES results were averaged along the length of the canyon in the y axis ( $L_y$ ), across the width of the canyon in the x axis ( $W$ ), over the height of the canyon in the z axis ( $H$ ) and averaged over the final hour of the simulation ( $150 \leq t \leq 210$  min, i.e. after 30 minutes of dynamical and 120 minutes of

chemical spin-up) to give a time and volume averaged within canyon concentration, for comparison with the box-model scenario.

$$\bar{c}_i = \frac{1}{W \cdot H \cdot L_y \cdot (t_2 - t_1)} \int_{t_1}^{t_2} \int_{-0.5W}^{0.5W} \int_0^H \int_0^{L_y} c_i(x, y, z, t) dx dy dz dt. \quad (3)$$

The LES results were also averaged along the length of the canyon in the y axis ( $L_y$ ) and over the final 60 minutes of the averaging period to give 2-D time averaged concentrations ( $150 \leq t \leq 210$  min) i.e.:

$$\langle \varphi \rangle(x, z) = \frac{1}{L_y \cdot (t_2 - t_1)} \int_{t_1}^{t_2} \int_0^{L_y} \varphi(x, y, z, t) dy dt, \quad (4)$$

where  $\varphi$  can be  $w$  or  $c_i$ . Following Eq. (4),  $\tilde{\varphi} = \varphi - \langle \varphi \rangle$  represents the resolved fluctuations of  $\varphi$  about  $\langle \varphi \rangle$ . Thus the following quantities are defined: the resolved-scale vertical turbulent flux,  $F_{turb} = \langle \tilde{w} \tilde{c}_i \rangle$ , the vertical advective flux,  $F_{adv} = \langle w \rangle \langle c_i \rangle$ , and total resolved-scale vertical flux  $F_{total} = F_{turb} + F_{adv}$ . These are 2D quantities showing the spatial pattern of these variables in the  $x$  and  $z$  domain. For the purposes of analysis, vertical mixing ratio profiles were extracted from the 2-D time averaged concentrations at five sites across the canyon. In addition, the horizontally-averaged vertical profile of the passive scalar's flux inside and above the canyon was derived (Equation 5):

$$F(z) = \frac{1}{W} \int_{-0.5W}^{0.5W} F(x, z) dx. \quad (5)$$

### 3.2 Configuration of the zero-dimensional box-RCS model

As one aim of this work is to compare canyon-average concentrations predicted by the full LES-RCS model using Equation 2, with their equivalents determined using a zero-dimensional box model, under identical chemical and emission conditions, treatment for the exchange between the street canyon and the overlying boundary layer was required. In the case of the LES-RCS model, the modelled domain (Figure 2) includes both within-canyon and above-canyon regions, and so implicitly incorporates exchange between the canyon and the overlying boundary layer. For the box model scenario, mixing with an overlying boundary layer was achieved by implementing a suitable exchange velocity ( $\omega$ ) within the model, defined as:

$$\omega_t = \frac{F_c}{\bar{c} - c_B} \quad (6)$$

where  $F_c$  is the pollutant flux  $F_i$  at roof level ( $z/H = 1$ ) and the denominator is the difference between the mean concentration within the canyon ( $\bar{c}$ ) and the background concentration above the canyon ( $c_B$ ). In the case of the box model simulations, the composition of the overlying boundary layer was assumed to be constant, and set equal to the LES-RCS domain inlet composition.

The value of  $\omega_t$  was determined using LES simulations of a passive scalar (a non-reactive emitted species which is conserved within the model, and whose concentrations are therefore determined by dynamics alone) to evaluate mean rate of exchange between the canyon and boundary layer in the LES simulations. A mean value of  $\omega_t = 0.021 \text{ m s}^{-1}$  was determined, averaged over the final hour of the simulation ( $150 \leq t \leq 210 \text{ min}$ ). Use of a single mean value removes the variability in the LES simulations (arising from periodic sweep / ejection events) – apparent in the comparisons of mean concentrations discussed below. The chemical scheme, photolysis treatment and emissions were identical in the LES-RCS and Box-RCS model cases.

## 4. Results and discussion

### 4.1 Spatial variation

Fig. 3 illustrates the mean mixing ratios of the passive scalar (subject solely to dispersion / mixing) and of a number of chemical species, averaged over the final 60 minutes of the 210 minute simulation (logarithmic colour scales for  $\text{O}_3$ , OH and  $\text{HO}_2$ ). Major features apparent are the primary vortex that spans the canyon and the shear layer at roof level that increases in amplitude and becomes increasingly turbulent downwind, trapping pollutants toward the leeward wall and allowing greater exchange toward the windward wall. Increased levels of NO,  $\text{NO}_2$  and the passive scalar are observed within the street canyon compared to the background atmosphere above roof level, as expected. The highest concentrations of these species are found at low level toward the leeward wall, a result of the limited dispersion and chemical processing of emissions before they are transported downwind from their sources located towards the centre of the street. In the case of  $\text{NO}_2$ , increased levels observed toward the leeward wall at street level arise through secondary formation through reaction of NO with entrained  $\text{O}_3$ , i.e. through the oxidation of emitted NO, in addition to that emitted directly.  $\text{HO}_x$  levels are much lower within the canyon than in the background air, with a local maximum in the centre of the vortex, a semi-isolated region of entrained background air. Elevated OH within this region (relative to the periphery of the canyon) primarily reflects a reduced OH sink, in particular reaction with NO – see below.

Fig. 4 illustrates the vertical mixing ratio profiles of  $\text{O}_3$ , NO,  $\text{NO}_2$ , OH and  $\text{HO}_2$  shown using a logarithmic scale at five across street locations within the canyon ( $z/H \leq 2$ ). The concentrations of  $\text{O}_3$ , OH and  $\text{HO}_2$  increase near roof level on the approach to the less polluted background atmosphere above. The smallest transition between within canyon and background concentrations of all chemical species occurs toward the windward wall due to the increase in exchange of air between the canyon and the background air above in this region associated with the turbulent nature of the shear layer at this point. Towards the leeward wall a sharp contrast exists between the canyon and the background atmosphere, marked by a large change in concentration with height at roof level as pollutants become

effectively trapped by the relatively impermeable shear layer that exists in this region. Fig. 4(a) illustrates a significant change in the concentration of NO and NO<sub>2</sub> across the canyon with the concentration of NO at street level toward the leeward wall more than double (2.3 x) that observed on the windward side, whilst NO<sub>2</sub> levels are over a third higher (1.4 x). The highest mixing ratios of ozone within the canyon are evident toward the windward wall and in particular toward roof level where ozone rich air is brought into the canyon from aloft. Toward the leeward wall the mixing ratios of both NO and NO<sub>2</sub> show a significant decrease with height above roof level where levels rapidly approach those of the background atmosphere. Moving toward the windward wall of the canyon again results in a much more gradual transition between within canyon and the background atmosphere as the influence of the shear layer spans a greater distance in this region causing the air well above the canyon to be mixed with that escaping from within.

The change in mixing ratio of OH and HO<sub>2</sub> is illustrated in Fig. 4(b). Within the canyon the greatest concentrations of both OH and HO<sub>2</sub> are observed toward roof level at the windward wall where concentrations are slightly higher for OH and over a third greater for HO<sub>2</sub> when compared to that at street level toward the leeward wall. OH and HO<sub>2</sub> levels are higher in the centre of the primary canyon vortex. Close to roof level ( $z/H \approx 1.1$ ) toward the leeward wall of the canyon OH and HO<sub>2</sub> approach their background mixing ratios of 0.22 and 1.54 ppt respectively, while these levels are only achieved well above at  $z/H \approx 1.5$  on the windward side of the canyon. The reductions in OH and HO<sub>2</sub> within the canyon compared to the background atmosphere are 64 % and 85 % respectively (averaged across the canyon). Mean within canyon levels of NO and NO<sub>2</sub> of 168 and 68 ppb respectively (Table 3) are considerably greater than the overlying urban boundary layer, while within-canyon O<sub>3</sub> levels are on average 11.2 ppb lower than those in the background atmosphere.

Figure 5(a) illustrates the mixing ratio of the sum of organic peroxy radicals (i.e.  $\Sigma \text{RO}_2$ , here excluding HO<sub>2</sub>) determined by the LES-RCS, averaged over the final 60 minutes of the simulation (logarithmic colour scale). The RO<sub>2</sub> abundance exhibits a similar pattern to that of OH, with a local minimum toward the lower leeward wall and maxima observed within the primary vortex, close to the mixing layer at roof level and toward the windward wall, and a tongue of elevated RO<sub>2</sub> accompanying the entrained air into the vortex on the windward side. To further investigate the importance of HO<sub>x</sub> sources and sinks within and above the canyon, the chemical rate of production / loss for OH was evaluated, using time averaged ( $150 \leq t \leq 210$  min) reactant concentrations determined for three locations (indicated in Fig. 5a): (V) within the vortex ( $z/H = 0.5$ ,  $x/W = 0$ ), (L) toward the leeward wall ( $z/H \approx 0.08$ ,  $x/W \approx -0.3$ ) and (B) in the background atmosphere above the canyon ( $z/H \approx 1.2$ ,  $x/W \approx -0.3$ ). The change in OH with respect to time is governed by the rate of production and loss, as below (neglecting transport, negligible for OH):

$$\frac{\partial[\text{OH}]}{\partial t} = P_{\text{OH}} - \sum_X k[\text{OH}][X]. \quad (7)$$

Where  $P_{OH}$  is the total rate of production of OH, and  $-k [OH] [X]$  represents the chemical loss of OH through reaction with X. The dominant OH chemical production and loss terms for each location (rates of reaction, here expresses in ppt s<sup>-1</sup>) are given in Table 2. Primary OH production ( $O_3$  photolysis) predictably falls with reduced  $O_3$  abundance from the background into the canyon (NO titration), but comprises a small fraction of the total production rate. NO-driven radical cycling dominates OH production, with a significant (up to 22 %) contribution from HONO photolysis. The lifetime of HONO is comparable to the timescale of the vortex circulation and mixing (photolysis lifetime of 8 minutes), thus HONO must be explicitly considered in understanding OH abundance (cannot be assumed to be in steady state). No heterogeneous sources of HONO were implemented within this model – see discussion below. Going from the background to leeward (within-canyon) locations, the OH sink increases from 2.8 s<sup>-1</sup> to 96 s<sup>-1</sup>, a factor of 30, while the dominant OH production rate,  $HO_2 + NO$ , increases from 1.1 to 11.7 ppt s<sup>-1</sup> (total OH production increases by a factor of 8.8); accordingly within-canyon OH levels are much lower than in the overlying boundary layer. The local maximum in OH in the vortex centre arises from the semi-isolated nature of this part of the domain; at the centre, OH production rates are comparable with those at the leeward site (total production rate of 11.5 vs. 13.6 ppt s<sup>-1</sup>, dominated by  $HO_2 + NO$ ), but the OH sink is much lower than that directly downwind of the emission source – 61 vs. 96 s<sup>-1</sup> – and OH levels are correspondingly *ca.* 31 % higher.

Within the street canyon, gas phase chemistry is dominated by  $NO_x$  and as such it is often useful to consider the temporal and spatial variation in  $NO_x$  levels. The chemical interaction of  $NO_x$  with  $O_3$  plays a key role in determining  $NO_2$  levels observed in the urban environment. As a result of this interaction, another useful measure to consider is defined as the total oxidant ( $O_x = O_3 + NO_2$ ). Considering only NO- $NO_2$ - $O_3$  reactions,  $O_x$  is conserved whilst partitioning between the component forms of  $O_3$  and  $NO_2$  is determined by overall levels of  $NO_x$ ,  $O_3$  and solar radiation. Figs. 5(b) and (c) illustrate the spatial variation in  $NO_x$  and  $O_x$  within and above the canyon domain. The mixing ratios of  $NO_x$  and  $O_x$  are both greater within the canyon (236 and 79 ppb respectively) compared to the background atmosphere where  $NO_x = 9$  ppb and  $O_x = 50$  ppb. The spatial distribution of  $NO_x$  (Fig. 5(b)) is similar to that of the passive scalar (Fig. 3(d)) demonstrating the effective conservation of  $NO_x$  on the canyon residence timescale. The change in mixing ratio of  $NO_x$  and  $O_x$  with height within the canyon ( $0.0 \leq z/H \leq 2.0$ ) at five sites across the street is illustrated in Fig. 6 using a logarithmic scale. The highest concentration of  $NO_x$  occurs at street level toward the leeward wall downwind of the two line emission sources located in the centre of the street. Figs. 5 and 6 also indicate that the lowest levels of  $O_x$  occur toward street level on the upper windward wall with the highest concentrations close to street level toward the leeward wall. Elevated  $O_x$  in this region arises from the 5 % primary  $NO_2$  emission within the  $NO_x$  source term. The lowest ratios of  $NO_2$  to NO (Fig. 5d) occur at street level downwind of the emission sources toward the leeward wall. The



decrease in the NO<sub>2</sub>/NO ratio within the canyon primarily reflects the predominance of NO in the primary NO<sub>x</sub> emission term. Table 3 illustrates that the ratio of NO<sub>2</sub> to NO is higher for the box model case indicating greater (mean) simulated conversion of NO to NO<sub>2</sub> when dynamical effects are neglected (i.e. assuming instant mixing). This arises as a number of cells within the LES model have very little or no O<sub>3</sub> present (Fig. 3) hence NO to NO<sub>2</sub> conversion is precluded at many locations.

#### 4.2 Atmospheric composition and exchange rate effects

The change in the canyon averaged concentration of the passive scalar over time is compared between the LES and box model results (Fig. 7). Fluctuations in the concentration of the averaged passive scalar inherent in the LES results are caused by large scale variations of the flow and the variable nature of canyon ventilation caused by the unsteady fluctuations in the shear layer at roof level, as observed by Louka et al. (2000) and reproduced by the model. The optimum value of exchange velocity,  $\omega_t$ , can be determined by minimising the difference between the passive scalar results for the LES and box model. As shown in Fig. 7, for the final 60 minute averaging period a value of  $\omega_t = 0.021 \text{ m s}^{-1}$  applied to the box model best represents the LES values and is therefore applied to the box model simulations for comparison with canyon averaged LES results of other chemical species.

The sensitivity of the canyon averaged concentrations derived from the box model output to the value of the exchange velocity,  $\omega_t$ , between the canyon air and the background atmosphere was investigated. The effect of increasing  $\omega_t$  from  $0.021 \text{ m s}^{-1}$  to  $0.022 \text{ m s}^{-1}$  is illustrated in Table 4. This increase in the exchange velocity results in a decrease in mean within canyon averages of NO<sub>x</sub> and O<sub>x</sub> by 4.5 % and 2 % respectively. The concentration of O<sub>3</sub> increases with  $\omega_t$ , partially due to an increase in O<sub>3</sub> rich background air entering the canyon from above, and partly due to lower NO levels reducing the titration of O<sub>3</sub> to NO<sub>2</sub>. For OH and HO<sub>2</sub>, increasing  $\omega_t$  reduces the modelled levels due to a reduction in the concentration of VOCs, as increased mixing leads to increased ventilation out of the canyon. In the case of the passive scalar, we have  $\omega_t \cdot (\bar{c} - c_B) = \text{constant}$ , and thus

$$\frac{\bar{c}^{(b)} - c_B}{\bar{c}^{(a)} - c_B} = \frac{\omega_a}{\omega_b}, \quad \text{and} \quad \frac{\bar{c}^{(b)} - \bar{c}^{(a)}}{\bar{c}^{(a)}} \cong -0.045.$$

Whereas for NO<sub>x</sub> where  $c_B \ll \bar{c}^{(a)}$  then  $\frac{\bar{c}^{(b)} - \bar{c}^{(a)}}{\bar{c}^{(a)}} \cong -\frac{\omega_b - \omega_a}{\omega_a} = -0.048$ .

The similarity of these results (and those in table 4) indicates that (on this timescale) NO<sub>x</sub> is approximately passive in nature.

#### 4.3 Temporal changes and segregation effects within the canyon

The variation with time of the spatially averaged ‘within canyon’ concentrations of a number of species simulated by the LES-RCS model is compared with their equivalents simulated using the box

model (with equal emissions and net external mixing applied) in Fig. 8. Significant differences between the concentrations of key chemical species simulated using the box and LES approaches are apparent. In general, the LES results show much greater dynamically-driven variability, and with some net deviations from the within-canyon mixing ratios taken from the box model. For NO, over the final 60 minutes of the simulation, the box model results are around 1 % higher than those of the LES simulation. Levels of NO<sub>2</sub> are also higher in the box model than the LES, throughout the simulation, with a mean difference of 10 % over the final hour of the modelled period ( $150 \leq t \leq 210$  min), while ozone levels are correspondingly lower, by 6 % over the final hour of the simulation. Fig. 8(a) illustrates the change in mixing ratio of NO<sub>x</sub> and O<sub>x</sub> over time. Over the final hour of the model run, levels of NO<sub>x</sub> and O<sub>x</sub> simulated by the box model are higher than those from the LES by 3 and 8 % respectively (Table 3) – thus the changes in abundance reflect chemical impacts upon NO<sub>x</sub> and O<sub>x</sub> abundance, rather than solely perturbations to the NO-NO<sub>2</sub>-O<sub>3</sub> photochemical steady state partitioning. Following the initial spin up of each model run, a large initial peak in both OH and HO<sub>2</sub> is observed directly after emissions are introduced. These peaks are followed by a rapid decline to equilibrium, which is achieved *ca.* 30 minutes after emission initiations. In contrast to the LES, the box model simulations approach equilibrium much more rapidly, reflecting the slower mixing processes inherent in the LES when compared to the fast and perfectly mixed conditions of the box model – as would be expected, segregation effects cause a reduction in the rate at which canyon-averaged concentrations approach the equilibrium levels simulated in a single-compartment model, i.e. street canyons respond more slowly to perturbations than a single-box model would suggest. OH and HO<sub>2</sub> levels simulated by the box model in steady state are higher than those in the LES simulation, with mean differences of 11 % and 8 % respectively over the final 60 minutes of the simulation. The assumption of instant mixing inherent to the box model leads to overestimates of the concentrations of NO, NO<sub>2</sub> and OH, and an underestimate for O<sub>3</sub>, relative to the LES-RCS approach. Segregation effects, spatial inhomogeneity in composition due to incomplete mixing, reduce the canyon-averaged rate at which O<sub>3</sub> reacts with NO to produce NO<sub>2</sub> (i.e. the dominant pathway for NO to NO<sub>2</sub> conversion), due to limited or near-zero quantities of O<sub>3</sub> in a number of cells within the LES model domain, as is apparent in Fig. 3. In terms of HO<sub>x</sub>, it is clear that segregation effects also play an important role in determining composition; the higher OH abundance within the box model implies an overestimate of the extent of OH-driven processing of reactive emissions within the canyon, compared with the more accurate LES scheme. While the comparison shown in Fig. 8(b) suggests a deviation of the order of 11 %, the actual difference will be greater, as the OH levels experienced by the majority of emitted air parcels within the canyon will reflect the circumference of the vortex, rather than the centre (where OH levels are approximately 30 % higher).

#### 4.4 VOC oxidation chemistry and atmospheric composition: RCS v a simple chemistry case

To further ascertain the effect of the detailed VOC oxidation chemistry, the results obtained using the RCS mechanism (51 chemical species and 136 reactions) were compared with those obtained using a 3 reaction scheme ( $\text{O}_3\text{-NO}_x$  chemistry alone, *i.e.* reactions 8 - 10, below), as used by previously by Baik et al. (2007), Baker et al. (2004) and Grawe et al. (2007).



Comparisons were performed using both the LES and box-model dynamical frameworks. The initial conditions ( $\text{NO}_x$  and  $\text{O}_3$  levels) were identical to those applied using the full chemical scheme in both cases, and within the LES construct the simulated dynamics were identical to those used with the full chemical scheme. It is important to note that photochemical steady state was not in fact achieved (or assumed) in any cells of the model domain – as the dynamical residence time was too short (typically fractions of a second, *vs.* the 1 - 2 minute time constant for the  $\text{NO-NO}_2\text{-O}_3$  system under sunlit conditions). Fig. 9 shows a comparison of the  $\text{O}_3\text{-NO}_x$ -only case with the full RCS reaction scheme. Going from the simple to the full scheme, levels of  $\text{NO}_2$  and  $\text{O}_3$  are higher and  $\text{NO}$  lower, reflecting additional  $\text{NO}$  to  $\text{NO}_2$  conversion (and net ozone production) in the more detailed chemistry. Correspondingly, levels of  $\text{O}_x$  are higher, while  $\text{NO}_x$  levels are slightly lower (again, going from the simple to the full scheme), reflecting the presence of  $\text{NO}_x$  loss processes (e.g. formation of nitric acid,  $\text{HNO}_3$ ) and partitioning to other  $\text{NO}_y$  species (e.g.  $\text{HONO}$ ,  $\text{HO}_2\text{NO}_2$ ). The differences between the full RCS scheme, and  $\text{O}_3\text{-NO}_x$ -only case, are similar but not identical between the box and LES dynamical frameworks – the changes in  $\text{NO}$ ,  $\text{NO}_2$ ,  $\text{NO}_x$ ,  $\text{O}_3$  and  $\text{O}_x$  are all less between the two LES models (dashed lines in Fig. 9), than the two box models (solid lines in Figure 9). For example, for  $\text{NO}$  the box models show a decrease of ca. 8 % going from the  $\text{O}_3\text{-NO}_x$ -only to the full chemical approaches, while the reduction is only 5 % for the LES models. For  $\text{NO}_2$ , the corresponding increases are ca. 18 % for the box models and 12 % for the LES approaches; changes in ozone are (proportionately) similar. Within the LES dynamical framework, the system is less sensitive to the additional chemical processes included in the RCS, compared with the  $\text{O}_3\text{-NO}_x$ -only scheme. In much of the domain,  $\text{O}_3$  levels are very low / zero, such that the  $\text{NO}:\text{NO}_2$  ratio is increased, and impacts of  $\text{NO}_x$  removal and partitioning in the full chemical scheme are reduced, while ozone production is immediately repartitioned into  $\text{NO}_2$  – effectively, segregation effects in the LES simulations make the canyon-averaged model composition less sensitive to the chemical simplifications attendant in moving from the RCS to the basic  $\text{O}_3\text{-NO}_x$  only chemical mechanism. These results are consistent with those of Garmory et al. (2009), who found only a modest sensitivity of the predicted  $\text{NO}_x$  and  $\text{O}_3$  levels towards the level of detail in the chemical mechanism used, but do not agree with the findings of Kim et al. (2012), who reported substantial (> 100 %) changes in ozone levels with the inclusion of VOC oxidation chemistry, in comparison with an  $\text{O}_3\text{-NO}_x$  only type approach. It is surprising that Kim et al. observed such substantial increases in ozone, as observed photochemical ozone production

rates are typically of the order of 5-20 ppb per hour (e.g. Cazorla et al., 2012), so increases of the order reported would imply canyon residence times of the order of hours. The study of Kim et al. successfully reproduced observed levels of essentially conserved species such as CO, suggesting that the general canyon circulation / residence time / ventilation rate were well simulated, but noted that levels of NO were substantially greater than those observed, pointing to differences in emissions from those occurring in reality – substantial discrepancies between predicted and observed vehicular emissions, particularly for NO<sub>x</sub>, are widely observed (e.g. Carslaw et al., 2011). Alternatively, this may reflect different model approaches to implementing photochemical steady state within the model – if it is assumed that photochemical steady state is actually achieved within each cell (in contrast to the present work), a different distribution between NO + O<sub>3</sub> and NO<sub>2</sub> would result – one which, relative to the approach used here, would be expected to favour NO<sub>2</sub> in the near-source region within the canyon, and NO + O<sub>3</sub> in the canyon outlet. It is not clear from the descriptions given if photochemical steady state was in fact adopted by Kim et al.

#### 4.5 Within-Canyon Chemical Processing

Fig. 10 illustrates the change in mixing ratio of O<sub>3</sub>, NO, NO<sub>2</sub>, NO<sub>x</sub> and O<sub>x</sub> with height at the canyon inlet ( $x/W = -0.5$ ) and canyon outlet planes ( $x/W = +0.5$ ). Increases are observed in the level of NO, NO<sub>2</sub>, NO<sub>x</sub> and O<sub>x</sub> leaving the canyon, indicating the combined effect of primary emissions and chemical processing on the abundance of pollutants escaping to the wider background atmosphere. In order to evaluate the extent of within-canyon processing further, the change in vertical flux of a number of species with height was calculated based on Equation 5 for the final hour of the model simulation (shown in Fig. 11). The calculated resolved-scale flux near roof level ( $z/H = 1$ ) can be used to determine a representative flux of pollutants out of the canyon and into the background atmosphere. This may then be compared with raw emission rates to evaluate the within-canyon processing (although given the turbulent nature of the canyon-background interface (e.g. Figs 1, 3) the choice of height at which to evaluate this is somewhat arbitrary). A peak in the resolved-scale turbulent flux profile is apparent at  $z/H \approx 0.1$  and a decrease in total flux is seen for  $z/H < 0.1$  in all flux profiles. This is the result of the elevation of the line emission source located within the centre of the canyon 1 m above street level. For the passive scalar, the profile differs from that expected a priori for a conserved quantity which should remain constant with height. At roof level ( $z/H \approx 1$ ) the flux of the passive scalar into the background atmosphere is equal to 933 ppb m<sup>-2</sup> s<sup>-1</sup> i.e. 93 % of that emitted is escaping into the wider atmosphere. The maximum flux of the passive scalar of 1000 ppb m<sup>-2</sup> s<sup>-1</sup> is observed slightly below roof level. This observed decrease in the flux of passive scalar with height arises from the sub-grid scale turbulent dispersion not resolved explicitly within the LES model (the sub-grid scale flux is not included here). Because of this, the fluxes of NO<sub>x</sub> and O<sub>x</sub> out of the canyon discussed below are obtained at a height slightly below the roof level,  $z_f = 0.933 H$ , where the

contribution of sub-grid scale dispersion is minimised and at which height the flux of passive scalar reaches 99.6% of its theoretical value.

A positive upward total flux of both NO and NO<sub>2</sub> from within the street to the overlying background atmosphere is observed with a negative (downward) flux of O<sub>3</sub>, with a significant effect of within-canyon chemical processing upon pollutant flux escaping the canyon apparent for NO and NO<sub>2</sub>. The maximum total flux of NO (798 ppb m<sup>-2</sup> s<sup>-1</sup>) occurs near street level ( $z/H \approx 0.2$ ). At  $z = z_f$ , the flux of NO is equal to 752 ppb m<sup>-2</sup> s<sup>-1</sup> compared to a raw emission rate equivalent to 900 ppb s<sup>-1</sup>. Therefore there is an approximately 16.5 % chemical conversion of NO within the canyon when compared to the raw emission rate. The maximum flux of NO<sub>2</sub> (248 ppb m<sup>-2</sup> s<sup>-1</sup>) occurs just below roof level and is approximately 2.5 times that of the raw emission rate of 100 ppb s<sup>-1</sup>, again indicating within-canyon processing affecting the level of NO<sub>2</sub> escaping into the wider urban boundary layer. In terms of O<sub>3</sub> the maximum downward flux into the canyon of 135 ppb m<sup>-2</sup> s<sup>-1</sup> occurs just above roof level as O<sub>3</sub> rich background air enters the canyon from above, and O<sub>3</sub> is removed within the canyon by reaction with NO. At  $z = z_f$  (near roof level), the flux of NO<sub>x</sub> is 989 ppb m<sup>-2</sup> s<sup>-1</sup> or 1.1% lower than that emitted; O<sub>x</sub> is 130 ppb m<sup>-2</sup> s<sup>-1</sup> at  $z = z_f$ , or 30% higher than the 100 ppb s<sup>-1</sup> of NO<sub>2</sub> emitted. Therefore NO<sub>x</sub> release into the boundary layer is almost the same as that emitted, but oxidant release increases significantly, in part as a result of the chemical processing taking place within the canyon. These findings demonstrate the value in considering the modelled (and observed) levels of NO<sub>x</sub> and of oxidant (O<sub>x</sub>), alongside NO / NO<sub>2</sub> / O<sub>3</sub>, which can be used to disaggregate the effects of NO<sub>x</sub>-O<sub>3</sub> photochemical steady state from net photochemical ozone production. Garmory et al. (2009) report values of simulated OH levels along a vertical profile through the centre of the canyon of the order of 0.003 – 0.006 ppt, significantly lower than the mean within-canyon value determined here (0.08 ppt; Table 3); however differences in the emissions profiles (total NO<sub>x</sub> emission used here *ca.* 60 % of that of Garmory et al. (with differing fractions of primary NO<sub>2</sub>: 5 % vs. 1 % respectively), and a much lower VOC:NO<sub>x</sub> emission ratio (*ca.* 10 % of that used by Garmory et al., by volume, excluding CO) probably accounts for much of this difference – lower OH levels reflecting the much larger VOC sink, not compensated for by equivalently enhanced NO-driven HO<sub>x</sub> cycling. Kim et al. (2012) showed that OH-driven oxidation (of SO<sub>2</sub> and NO<sub>2</sub>) could result in measurable secondary aerosol production of up to 0.7 µg m<sup>-3</sup> within the street canyon, which may be compared with roadside PM<sub>10</sub> levels of 25 – 47 µg m<sup>-3</sup> (whole UK roadside site annual average, and Marylebone roadside annual average - AQEG, 2005). Kim et al. do not report their simulated OH levels, and no condensed phase processes were considered in this work; however an upper estimate for comparison may be obtained: The mean flux through the OH + NO<sub>2</sub> reaction (Table 2, *i.e.* neglecting that only a fraction of this will partition to the particulate phase within the canyon) equates to 56 µg m<sup>-3</sup> of HNO<sub>3</sub> production, supporting the

conclusion of Kim et al. that significant secondary aerosol production may occur on the canyon timescale.

Several studies have examined the partition of total flux of a scalar ( $F_{total}$ ) between mean advective flux ( $F_{adv}$ ) and turbulent flux ( $F_{turb}$ ) for a street canyon. Based on the results of their RANS renormalisation group (RNG)  $k-\varepsilon$  turbulence model for a passive scalar, Liu et al. (2011) analysed roof level fluxes, *i.e.*,  $F(z=H)$ , where  $F(z)$  as defined by Equation 5 can be either  $F_{adv}$  or  $F_{turb}$  (denoted by  $\overline{PCH}$  or  $PCH''$  in their paper). Our results (Fig. 11) demonstrate that both  $F_{adv}(z)$  or  $F_{turb}(z)$  are very sensitive to height near the roof level. The 2D fields of  $F_{adv}(x,z)$  and  $F_{turb}(x,z)$  for a passive scalar derived from LES simulations conducted by Cheng and Liu (2011) support this observation. Therefore the analysis of vertical profiles of  $F_{adv}$  and  $F_{turb}$  is necessary and useful. As seen in Fig. 11, the dominant vertical flux observed within the canyon from the street level up to  $z/H \approx 0.8$  is the mean advective flux which transports all pollutants (except for  $O_3$ ) upwards toward roof level throughout the canyon. Between  $z/H \approx 0.8$  and roof level, the mean advective flux decreases rapidly to become negative just above the top of the canyon ( $1.0 \leq z/H \leq 1.2$ ) indicating that the mean flow (averaged across the domain) acts to entrain pollutants downwards toward the canyon at this height. For all species included in Fig. 11, a large increase in the resolved-scale turbulent flux is observed toward roof level indicating the importance of the shear layer associated turbulent processes in pollutant exchange at this level. For all species except  $O_3$ , the resolved-scale turbulent flux at roof level is at a maximum and is positive, indicating that there is more pollutant escaping out of the canyon at this height through turbulent transport than entering from the background atmosphere above. The mean flux of all species excluding  $O_3$  and  $O_x$  becomes close to zero at a height of  $z/H \approx 1.4$  *i.e.* toward the free flowing boundary layer above, which is unaffected by the dynamical processes taking place within and just above the canyon itself. For  $O_3$  however, a negative mean flux is observed until well above roof level indicating net downward transport of  $O_3$  rich air towards the canyon. These results are consistent with those of Baik et al. (2007), who adopted an RNG  $k-\varepsilon$  turbulence model but applied it to simple  $NO_x$ - $O_3$  chemistry; they showed 2D fields of  $-\partial F(x,z)/\partial z$  instead of  $F(x,z)$  for the purpose of budget analysis, in which  $F(x,z)$  was either  $F_{adv}(x,z)$  or  $F_{turb}(x,z)$  of  $NO$ ,  $NO_2$ , and  $O_3$ .

## 5. Conclusions

A reduced chemical scheme has been developed based upon a subset of a near explicit chemical mechanism, and implemented within an LES simulation of urban street canyon dynamics. The resultant LES-RCS model has been used to investigate urban street canyon atmospheric composition by simulating the combined effects of emissions, mixing and chemical processing on pollutant concentration within an idealised canyon. Pollutants within and above the canyon were found to show a clear spatial variation, with  $NO_x$  levels close to the leeward wall over double those of the

windward wall; such variations are of importance in assessing the potential exposure of receptors to air pollutants. Through comparison of simulations using the LES dynamical framework with those using a simple zero-dimensional box model approach, the effects of segregation on canyon atmospheric chemistry and composition are evident. Compared with a single-box canyon model, the LES scheme responds more slowly to chemical perturbations, and (after quasi-equilibrium is established) the box model simulated levels of NO, NO<sub>2</sub> and OH were found to be higher than their (canyon-averaged) equivalents in the more realistic LES scheme, while levels of O<sub>3</sub> were underestimated compared with the LES approach. The assumption of instant mixing inherent to the box model leads to overestimates of the concentrations of NO, NO<sub>2</sub> and OH, and an underestimate for O<sub>3</sub>, relative to the LES approach. Segregation effects, due to spatial inhomogeneity in composition due to incomplete mixing, reduce the canyon-averaged rate at which O<sub>3</sub> reacts with NO to produce NO<sub>2</sub> (i.e. the dominant pathway for NO to NO<sub>2</sub> conversion), due to limited quantities of O<sub>3</sub> present in a number of cells within the LES model domain (Fig. 3). Segregation effects also affect HO<sub>x</sub> levels; the higher OH abundance within the box model implies an overestimate of the extent of OH-driven processing of reactive emissions within the canyon, compared with the more realistic LES approach. While the comparison shown in Fig. 8(b) suggests a deviation of the order of 11 %, the actual difference will be greater, as the OH levels experienced by the majority of emitted air parcels within the canyon will reflect the circumference of the vortex, rather than the centre (where OH levels are approximately 30 % higher). Through comparison of the comprehensive RCS and O<sub>3</sub>-NO<sub>x</sub>-only chemical mechanisms, a clear effect of the inclusion of detailed oxidation chemistry is also evident. Going from the O<sub>3</sub>-NO<sub>x</sub> only system to the RCS mechanisms, levels of NO<sub>2</sub> and O<sub>3</sub> are higher and NO lower, reflecting additional NO to NO<sub>2</sub> conversion (and net ozone production) under the more detailed chemistry. Segregation effects reduced the sensitivity of the model outputs to the increase in chemical complexity when comparing the box model dynamical framework to the LES approach.

Chemical processing of emissions takes place within the canyon, contributing to an increase in O<sub>x</sub> (O<sub>3</sub> + NO<sub>2</sub>) of 30 % (compared to the primary NO<sub>2</sub> component of the emission source), for the moderately polluted emission scenario considered. This result shows that the atmospheric “pre-processing” of primary emissions taking place within street canyons can be significant in terms of atmospheric composition and the flux of pollutants from street canyon level to the wider urban boundary layer above. These processes are likely to be dependent upon the nature of the domain (canyon aspect ratio), prevailing meteorology and emission / pollution scenario considered. Further research to average the extent of these effects across a representative parameter space will determine the modification to raw emission rates which might be applied to account for within canyon processing of raw emissions in larger scale regional and neighbourhood models.

## Acknowledgements

786

787 The authors would like to thank Dr Mike Jenkin and Professor Dudley Shallcross for the provision of  
788 the Common Representatives Intermediates mechanism version 2-R5. VB thanks the University of  
789 Birmingham for award of a PhD scholarship. The computations described in this paper were  
790 performed using the University of Birmingham's BlueBEAR HPC service  
791 (<http://www.bear.bham.ac.uk>), which was purchased through HEFCE SRIF-3 funds. The comments  
792 of the anonymous reviewers are gratefully acknowledged.

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## References

- AQEG, 2005. Particulate Matter in the United Kingdom. Report of the UK Air Quality Expert Group, AQEG. Prepared for the Department for Environment, Food and Rural Affairs, the Scottish Executive, the Welsh Assembly Government and the Department of the Environment in Northern Ireland.
- AQEG, 2009. Ozone in the United Kingdom. Report of the UK Air Quality Expert Group, AQEG. Prepared for the Department for Environment, Food and Rural Affairs, the Scottish Executive, the Welsh Assembly Government and the Department of the Environment in Northern Ireland.
- Aumont, B., Szopa, S., Madronich, S., 2005. Modelling the evolution of organic carbon during its gas-phase tropospheric oxidation: development of an explicit model based on a self generating approach. *Atmos. Chem. Phys.* **5**, 2497-2517.
- Baik, J.J., Kang, Y.S., Kim, J.J., 2007. Modeling reactive pollutant dispersion in an urban street canyon. *Atmospheric Environment* **41**, 934-949.
- Baker, J., Walker, H.L., Cai, X.M., 2004. A study of the dispersion and transport of reactive pollutants in and above street canyons - a large eddy simulation. *Atmospheric Environment* **38**, 6883-6892.
- Berkowicz, R., Ketzel, M., Vachon, G., Louka, P., Rosant, J.M., Mestayer, P.G., Sini, J.F., 2002. Examination of traffic pollution distribution in a street canyon using the Nantes'99 experimental data and comparison with model results, in: Sokhi, R.S., Bartzis, J.G. (Eds.), *Urban Air Quality - Recent Advances, Proceedings*, pp. 311-324.
- Bey, I., Jacob, D.J., Yantosca, R.M., Logan, J.A., Field, B.D., Fiore, A.M., Li, Q., Liu, H.Y., Mickley, L.J., Schultz, M.G. 2001. Global modelling of tropospheric chemistry with assimilated meteorology: model description and evaluation. *J. Geophys. Res.* **106**, 23073-23095.
- Bloss, C., Wagner, V., Jenkin, M.E., Volkamer, R., Bloss, W.J., Lee, J.D., Heard, D.E., Wirtz, K., Martin-Reviejo, M., Rea, G., Wenger, J.C., Pilling, M.J., 2005. Development of a detailed chemical mechanism (MCMv3.1) for the atmospheric oxidation of aromatic hydrocarbons. *Atmos. Chem. Phys.* **5**, 641-664.
- Boulter, P.G., Barlow, T.J., Latham, S., McCrae, I.S., 2009. Emission Factors 2009: Report 1 - a review of methods for determining hot exhaust emission factors for road vehicles. TRL, Wokingham.

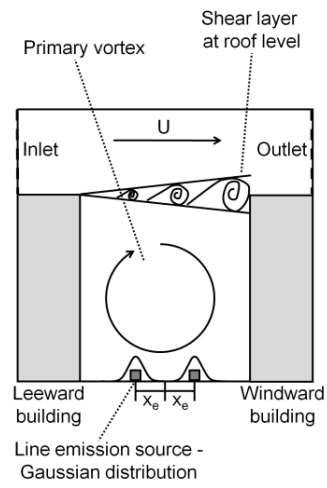
- 828 Cai, X.M., Barlow, J.F., Belcher, S.E., 2008. Dispersion and transfer of passive scalars in and  
829 above street canyons - Large-eddy simulations. *Atmospheric Environment* **42**, 5885-  
830 5895.
- 831 Carslaw, D., Beevers, S., Westmoreland, E., Williams, M., Tate, J., Murrells, T., Stedman, J.,  
832 Li, Y., Grice, S., Kent, A., Tsagatakis, I., 2011. Trends in NO<sub>x</sub> and NO<sub>2</sub> emissions and  
833 ambient measurements in the UK, *Department for Environment Food and Rural*  
834 *Affairs*.
- 835 Cazorla, M., Brune, W.H., Lefer, B. 2012. Direct measurement of ozone production rates in  
836 Houston in 2009 and comparison with two estimation methods, *Atmos. Chem. Phys.* **12**,  
837 1205-1212.
- 838 Cheng, W., Liu, C.-H., 2011. Large-Eddy Simulation of Flow and Pollutant Transports in and  
839 Above Two-Dimensional Idealized Street Canyons. *Boundary-Layer Meteorology*, 1-  
840 27.
- 841 Cheng, W., Liu, C.-H., 2011. Large-eddy simulation of turbulent transports in urban street  
842 canyons in different thermal stabilities, *J. Wind Engineering Ind. Aerodyn.*, **99**, 434-  
843 442.
- 844 Cui, Z.Q., Cai, X.M., Baker, C.J., 2004. Large-eddy simulation of turbulent flow in a street  
845 canyon. *Quarterly Journal of the Royal Meteorological Society* **130**, 1373-1394.
- 846 DePaul, F.T., Sheih, C.M., 1985. A tracer study of dispersion in an urban street canyon.  
847 *Atmospheric Environment* (1967) **19**, 555-559.
- 848 DFT, 2009. Transport Statistics Bulletin, Road Statistics 2008: Traffic, Speeds and  
849 Congestion. Department for Transport.
- 850 Dodge, M.C., 2000. Chemical oxidant mechanisms for air quality modeling: critical review.  
851 *Atmospheric Environment* **34**, 2103-2130.
- 852 Emmerson, K.M., Evans, M.J., 2009. Comparison of tropospheric gas-phase chemistry  
853 schemes for use within global models. *Atmos. Chem. Phys.* **9**, 1831-1845.
- 854 Garmory, A., Richardson, E.S., Mastorakos, E., 2006. Micromixing effects in a reacting  
855 plume by the Stochastic Fields method. *Atmospheric Environment* **40**, 1078-1091.
- 856 Garmory, A., Kim, I.S., Britter, R.E., Mastorakos, E., 2009. Simulations of the dispersion of  
857 reactive pollutants in a street canyon, considering different chemical mechanisms and  
858 micromixing. *Atmospheric Environment* **43**, 4670-4680.
- 859 Gery, M.W., Whitten, G.Z., Killus, J.P., Dodge, M.C., 1989. A photochemical kinetics  
860 mechanism for urban and regional scale computer modeling. *J. Geophys. Res.-Atmos.*  
861 **94**, 12925-12956.

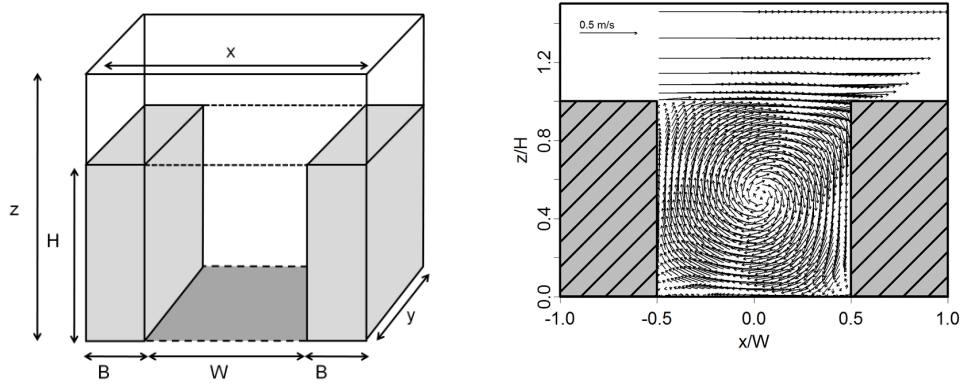
- 862 Grawe, D., Cai, X.M., Harrison, R.M., 2007. Large eddy simulation of shading effects on  
863 NO<sub>2</sub> and O<sub>3</sub> concentrations within an idealised street canyon. *Atmospheric*  
864 *Environment* **41**, 7304-7314.
- 865 Gromke, C., Buccolieri, R., Di Sabatino, S., Ruck, B., 2008. Dispersion study in a street  
866 canyon with tree planting by means of wind tunnel and numerical investigations -  
867 Evaluation of CFD data with experimental data. *Atmospheric Environment* **42**, 8640-  
868 8650.
- 869 Heard, D.E., Pilling, M.J., 2003. Measurement of OH and HO<sub>2</sub> in the troposphere. *Chem.*  
870 *Rev.* **103**, 5163-5198.
- 871 Hunter, L.J., Johnson, G.T., Watson, I.D., 1992. An investigation of three-dimensional  
872 characteristics of flow regimes within the urban canyon. *Atmospheric Environment.*  
873 Part B. Urban Atmosphere **26**, 425-432.
- 874 Jenkin, M.E., Saunders, S.M., Pilling, M.J., 1997. The tropospheric degradation of volatile  
875 organic compounds: A Protocol for Mechanism Development, *Atmospheric*  
876 *Environment* **31**, 81-104.
- 877 Jenkin, M.E., Saunders, S.M., Wagner, V., Pilling, M.J., 2003. Protocol for the development  
878 of the Master Chemical Mechanism, MCM v3 (Part B): tropospheric degradation of  
879 aromatic volatile organic compounds. *Atmos. Chem. Phys.* **3**, 181-193.
- 880 Jenkin, M.E., Watson, L.A., Utembe, S.R., Shallcross, D.E., 2008. A Common  
881 Representative Intermediates (CRI) mechanism for VOC degradation. Part 1: Gas phase  
882 mechanism development. *Atmospheric Environment* **42**, 7185-7195.
- 883 Kastner-Klein, P., Plate, E.J., 1999. Wind-tunnel study of concentration fields in street  
884 canyons. *Atmospheric Environment* **33**, 3973-3979.
- 885 Kim, M.J., Park, R.J., Kim, J.-J. 2012. Urban air quality modelling with full O<sub>3</sub>-NO<sub>x</sub>-VOC  
886 chemistry: Implications for O<sub>3</sub> and PM air quality in a street canyon. *Atmos. Environ.*  
887 **47**, 330-340.
- 888 Krol, M.C., Molemaker, M.J., de Arellano, J.V.G., 2000. Effects of turbulence and  
889 heterogeneous emissions on photochemically active species in the convective boundary  
890 layer. *J. Geophys. Res.-Atmos.* **105**, 6871-6884.
- 891 Kwak, K.-H., Baik, J.-J. 2012. A CFD modeling study of the impacts of NO<sub>x</sub> and VOC  
892 emissions on reactive pollutant dispersion in and above a street canyon. *Atmos.*  
893 *Environ.* **46**, 71-80.
- 894 Lee, J.D., Lewis, A.C., Monks, P.S., Jacob, M., Hamilton, J.F., Hopkins, J.R., Watson, N.M.,  
895 Saxton, J.E., Ennis, C., Carpenter, L.J., Carslaw, N., Fleming, Z., Bandy, B.J., Oram,

- 896 D.E., Penkett, S.A., Slemr, J., Norton, E., Rickard, A.R., Whalley, L.K., Heard, D.E.,  
 897 Bloss, W.J., Gravestock, T., Smith, S.C., Stanton, J., Pilling, M.J., Jenkin, M.E., 2006.  
 898 Ozone photochemistry and elevated isoprene during the UK heatwave of August 2003.  
 899 *Atmospheric Environment* **40**, 7598-7613.
- 900 Li, X.-X., Liu, C.-H., Leung, D., 2008. Large-Eddy Simulation of Flow and Pollutant  
 901 Dispersion in High-Aspect-Ratio Urban Street Canyons with Wall Model. *Boundary-  
 902 Layer Meteorology* **129**, 249-268.
- 903 Li, X.-X., Liu, C.H., Leung, D.Y.C., 2009. Numerical investigation of pollutant transport  
 904 characteristics inside deep urban street canyons. *Atmospheric Environment* **43**, 2410-  
 905 2418.
- 906 Li, X.X., Liu, C.H., Leung, D.Y.C., Lam, K.M., 2006. Recent progress in CFD modelling of  
 907 wind field and pollutant transport in street canyons. *Atmospheric Environment* **40**,  
 908 5640-5658.
- 909 Liu, C.-H., Cheng, W.C., Leung, T.C.Y., Leung, D.Y.C., 2011. On the mechanism of air  
 910 pollutant re-entrainment in two-dimensional idealized street canyons. *Atmospheric  
 911 Environment* **45**, 4763-4769.
- 912 Liu, C.H., Barth, M.C., 2002. Large-eddy simulation of flow and scalar transport in a  
 913 modeled street canyon. *J. Appl. Meteorol.* **41**, 660-673.
- 914 Liu, C.H., Barth, M.C., Leung, D.Y.C., 2004. Large-eddy simulation of flow and pollutant  
 915 transport in street canyons of different building-height-to-street-width ratios. *J. Appl.  
 916 Meteorol.* **43**, 1410-1424.
- 917 Lelieveld, J., Butler, T.M., Crowley, J.N., Dillon, T.J., Fischer, H., Ganzeveld, L., Harder, H.,  
 918 Lawrence, M.G., Martinez, M., Taraborrelli, D. and Williams, J., 2008. Atmospheric  
 919 oxidation capacity sustained by a tropical forest. *Nature* **452**, 736-740.
- 920 Louka, P., Belcher, S.E., Harrison, R.G., 2000. Coupling between air flow in streets and the  
 921 well-developed boundary layer aloft. *Atmospheric Environment* **34**, 2613-2621.
- 922 Madronich, S., Flocke, S., 1998. The role of solar radiation in atmospheric chemistry in:  
 923 Boule, P. (Ed.), *Handbook of Environmental Chemistry*. Springer, New York, pp. 1–26.
- 924 Oke, T.R., 1987. *Boundary layer climates* / T.R. Oke, 2nd ed. Methuen, London.
- 925 Pavageau, M., Schatzmann, M., 1999. Wind tunnel measurements of concentration  
 926 fluctuations in an urban street canyon. *Atmospheric Environment* **33**, 3961-3971.
- 927 Pugh, T.A.M., MacKenzie, A.R., Langford, B., Nemitz, E., Misztal, P.K., Hewitt, N.K.,  
 928 2011. The influence of small-scale variations in isoprene concentrations on atmospheric  
 929 chemistry over a tropical rainforest, *Atmos. Chem. Phys.* **11**, 4121-4134

- 930 Salim, S.M., Buccolieri, R., Chan, A., Di Sabatino, S., 2010. Numerical simulation of  
931 atmospheric pollutant dispersion in an urban street canyon: Comparison between  
932 RANS and LES. *Journal of Wind Engineering and Industrial Aerodynamics* **99**, 103-  
933 113.
- 934 Salizzoni, P., Soulhac, L., Mejean, P., 2009. Street canyon ventilation and atmospheric  
935 turbulence. *Atmospheric Environment* **43**, 5056-5067.
- 936 Saunders, S.M., Jenkin, M.E., Derwent, R.G., Pilling, M.J., 2003. Protocol for the  
937 development of the Master Chemical Mechanism, MCM v3 (Part A): tropospheric  
938 degradation of non-aromatic volatile organic compounds. *Atmos. Chem. Phys.* **3**, 161-  
939 180.
- 940 So, E.S.P., Chan, A.T.Y., Wong, A.Y.T., 2005. Large-eddy simulations of wind flow and  
941 pollutant dispersion in a street canyon. *Atmospheric Environment* **39**, 3573-3582.
- 942 Tomlin, A.S., Smalley, R.J., Tate, J.E., Barlow, J.F., Belcher, S.E., Arnold, S.J., Dobre, A.,  
943 Robins, A., 2009. A field study of factors influencing the concentrations of a traffic-  
944 related pollutant in the vicinity of a complex urban junction. *Atmospheric Environment*  
945 **43**, 5027-5037.
- 946 Vardoulakis, S., Fisher, B.E.A., Pericleous, K., Gonzalez-Flesca, N., 2003. Modelling air  
947 quality in street canyons: a review. *Atmospheric Environment* **37**, 155-182.
- 948 Wang, P., Mu, H.L., 2010. Analysis of Reactive Pollutants Distribution in Damaging Street  
949 Canyon Architectures, in: Luo, Q. (Ed.), *Information Technology for Manufacturing*  
950 *Systems*, Pts 1 and 2, pp. 1115-1120.
- 951 Watson, L.A., Shallcross, D.E., Utembe, S.R., Jenkin, M.E., 2008. A Common  
952 Representative Intermediates (CRI) mechanism for VOC degradation. Part 2: Gas phase  
953 mechanism reduction. *Atmospheric Environment* **42**, 7196-7204.
- 954 Xie, S.D., Zhang, Y.H., Li, Q., Tang, X.Y., 2003. Spatial distribution of traffic-related  
955 pollutant concentrations in street canyons. *Atmospheric Environment* **37**, 3213-3224.

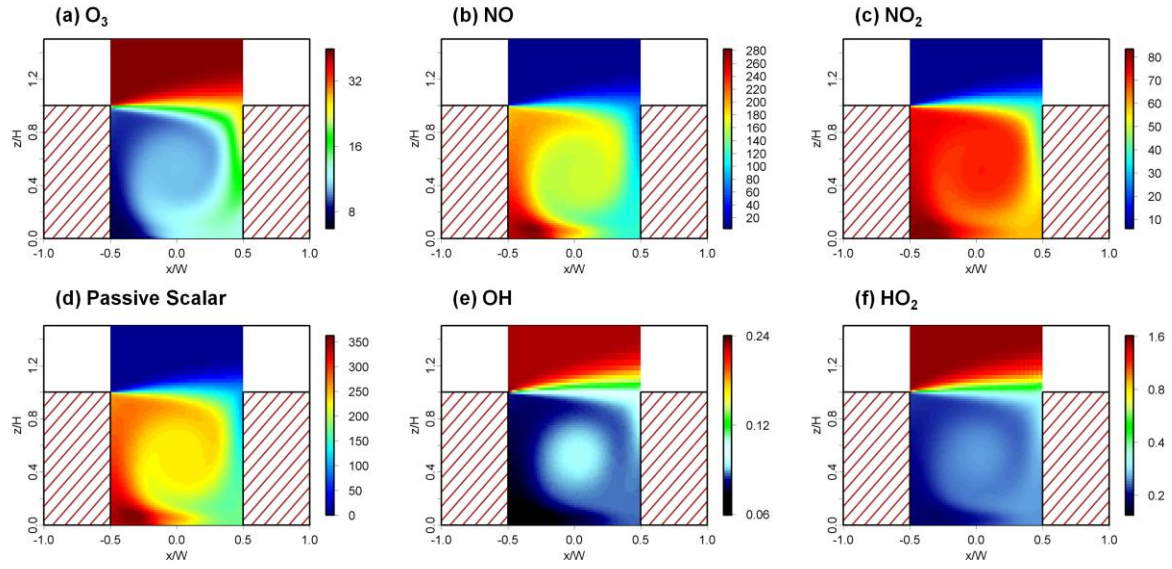


**Figure 1****Fig. 1.** Schematic diagram illustrating the main components of an idealised street canyon, as used in this study.

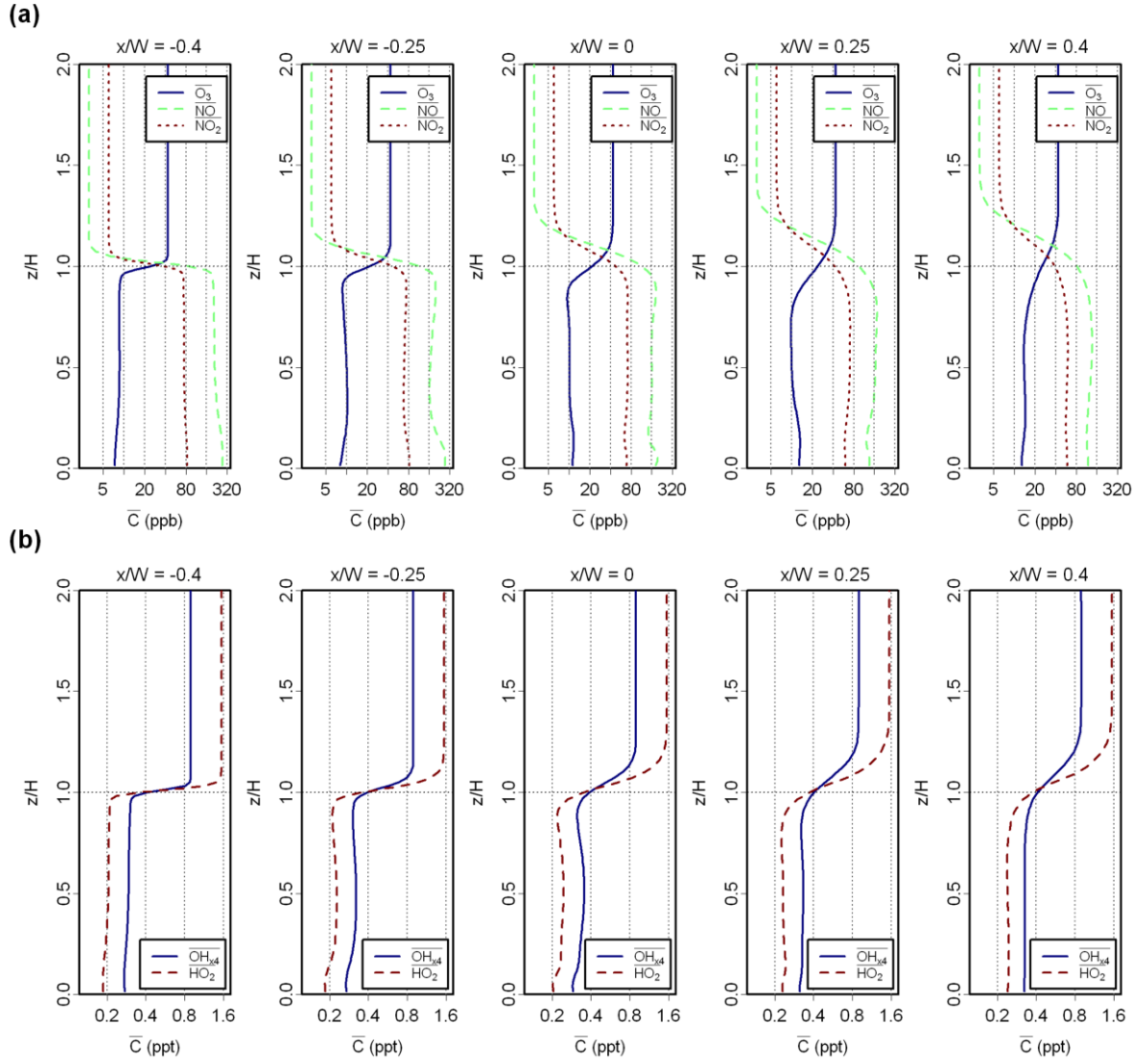
**Figure 2**

**Fig. 2.** (a) Schematic illustration of the LES model domain where  $x = 24$  m,  $y = 40$  m and  $z = 94$  m with canyon dimensions  $W = 18$  m,  $H = 18$  m and  $B = 3$  m and (b) Mean flow field diagram, showing temporally and spatially (along the  $y$  axis) averaged wind vectors ( $u$ ,  $w$ ). Averages were taken over the final hour of the model simulation ( $150 \leq t \leq 210$  min), when the canyon circulation was fully developed.



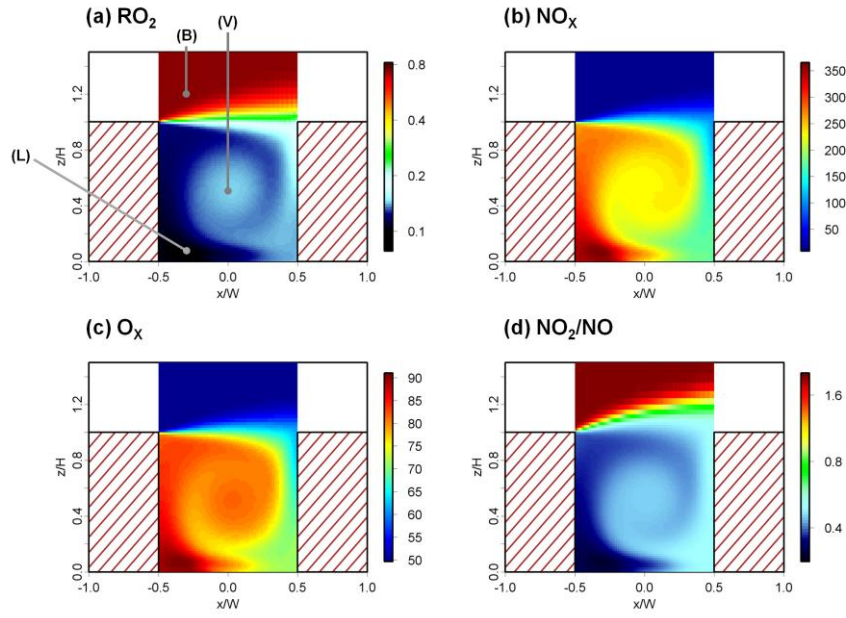
**Figure 3**

**Fig. 3.** Mean (time averaged) mixing ratios (ppb) of (a)  $O_3$ , (b)  $NO$ , (c)  $NO_2$ , (d) passive scalar, (e)  $OH$  (ppt) and (f)  $HO_2$  (ppt) from the LES-RCS simulations. Data averaged over the final hour of the model simulation ( $150 \leq t \leq 210$  min).

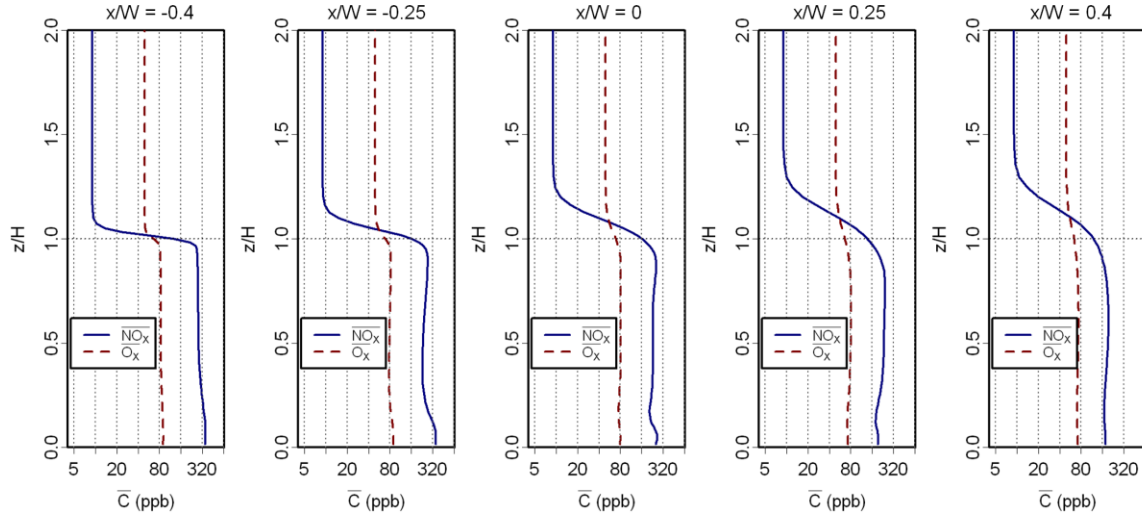
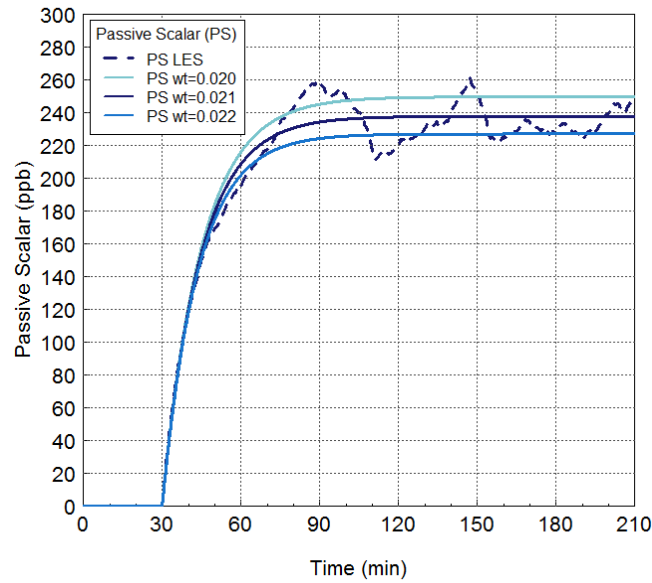
**Figure 4**

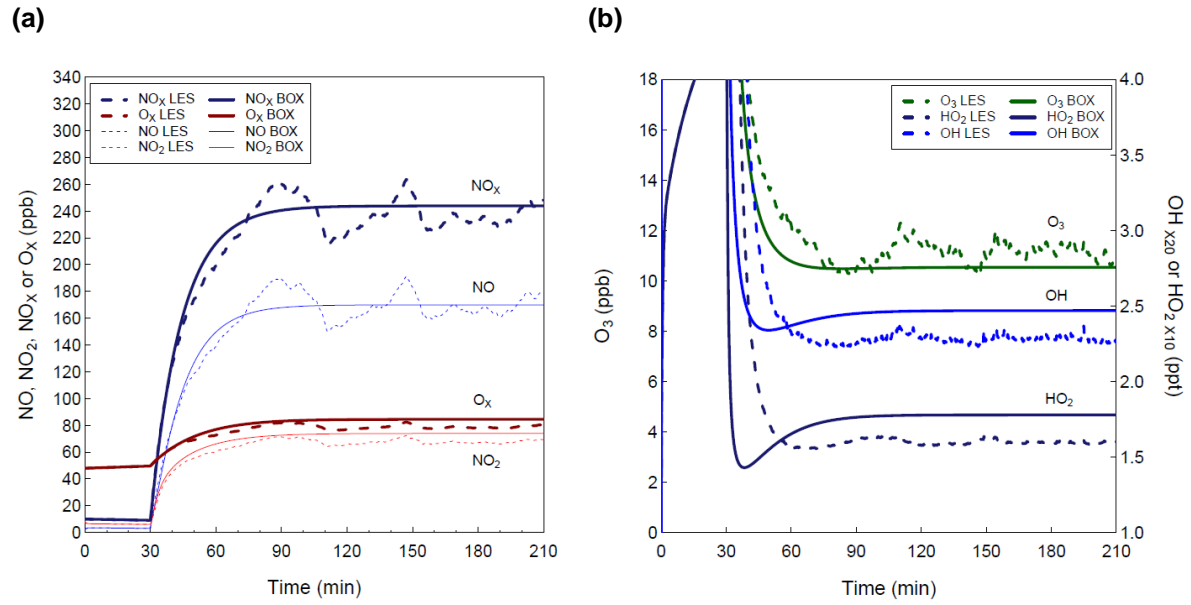
**Fig. 4.** Variation in the mean and time averaged mixing ratios of (a)  $O_3$ ,  $NO$ ,  $NO_2$ , and (b)  $OH_{x4}$  ( $4 \times OH$ ) and  $HO_2$  with height within the canyon ( $0.0 \leq z/H \leq 2.0$ ) at  $x/W = -0.4, -0.25, 0.0, 0.25, 0.4$ , from the LES-RCS simulation.

Figure 5

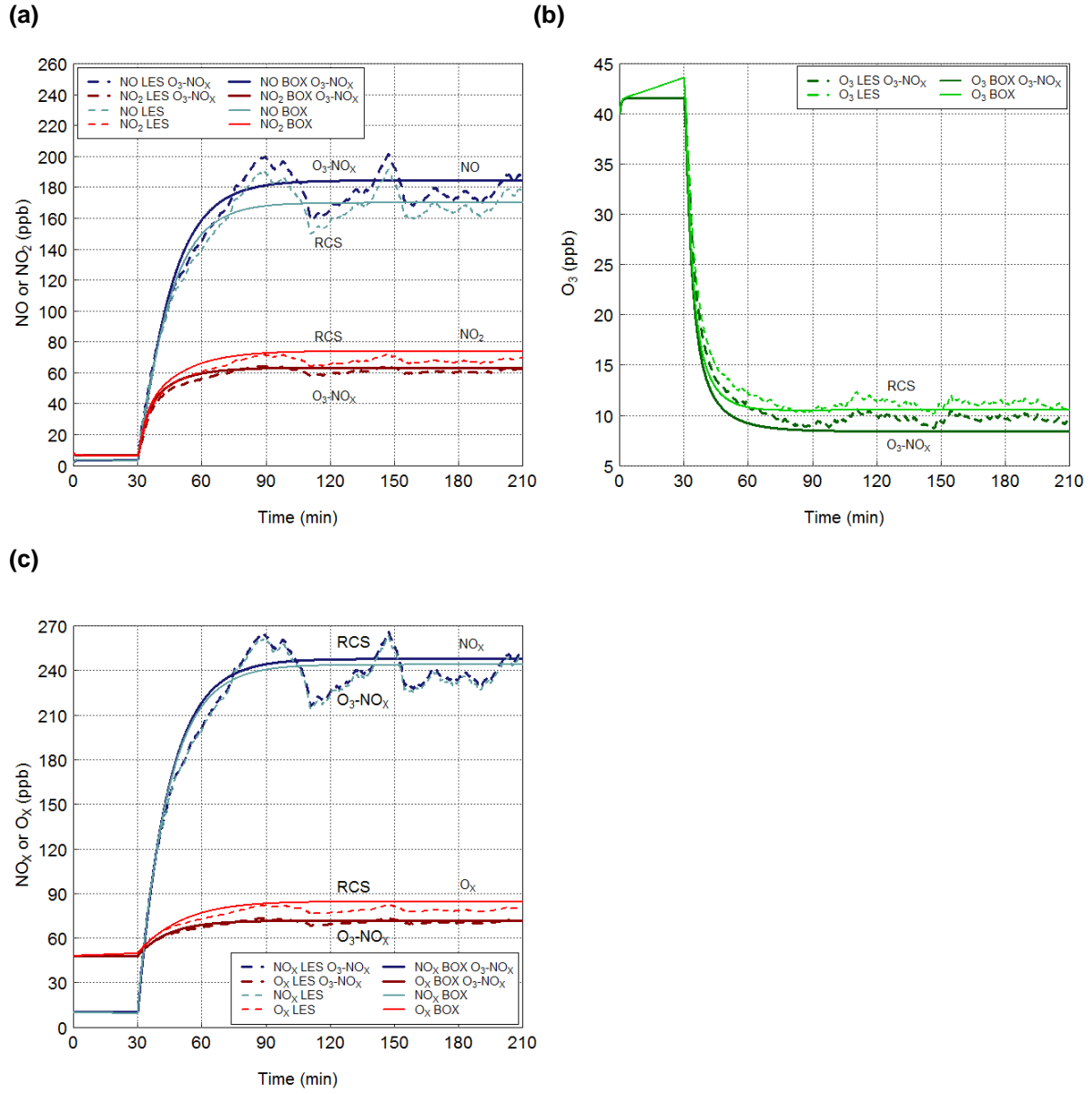


**Fig. 5.** Mean and time averaged mixing ratios of (a)  $RO_2$  (ppt), (b)  $NO_x$  (ppb), (c)  $O_3$  (ppb) and (d)  $NO_2/NO$  ratio from the LES-RCS simulation.

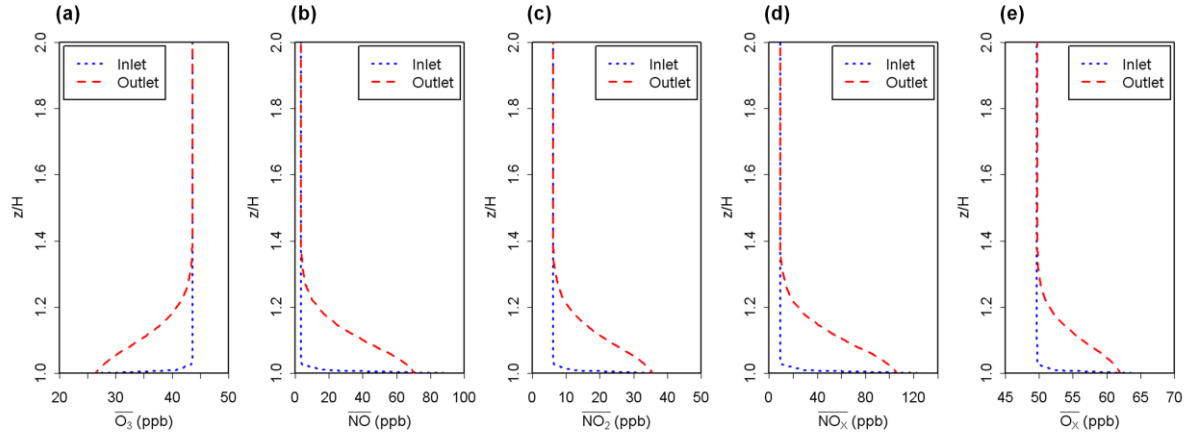
**Figure 6****Fig. 6.** Time averaged vertical mixing ratio profiles on a logarithmic scale of  $\text{NO}_x$  and  $\text{O}_3$  at  $x/W = -0.4, -0.25, 0.0, 0.25, 0.4$  within and above the canyon ( $0.0 \leq z/H \leq 2.0$ ), from the LES-RCS simulation.**Figure 7****Fig. 7.** Time evolution of the canyon averaged mixing ratio of the (chemically conserved) passive scalar calculated using the LES and box model dynamical frameworks, as a function of the exchange velocity ( $\omega_t$ ) – values in  $\text{ms}^{-1}$ .

**Figure 8**

**Fig. 8.** Time evolution of the canyon averaged mixing ratio of (a)  $\text{NO}_x$ ,  $\text{O}_x$ ,  $\text{NO}$ ,  $\text{NO}_2$  (ppb) and (b)  $\text{O}_3$  (ppb),  $\text{OH}_{x20}$  ( $20 \times \text{OH}$ ) (ppt),  $\text{HO}_{2x10}$  ( $10 \times \text{HO}_2$ ), calculated using the LES-RCS and Box-RCS simulations.

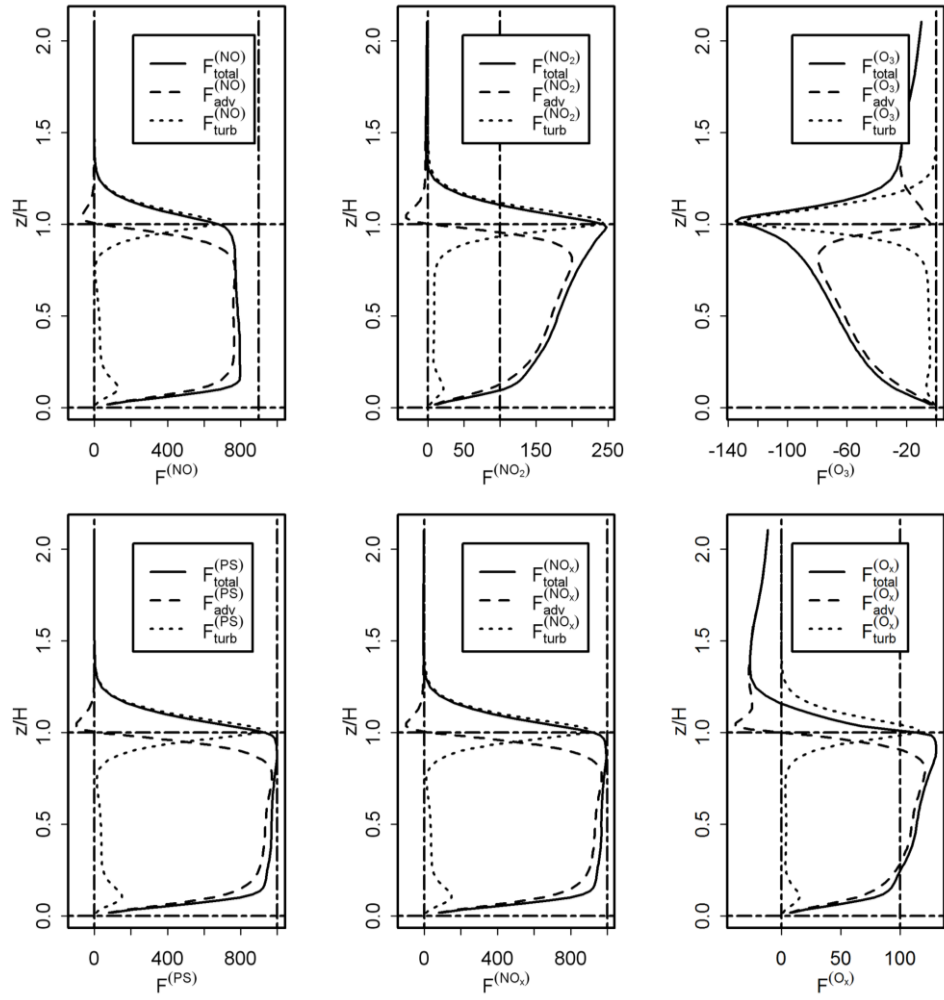


**Fig. 9.** Time evolution of canyon averaged mixing ratio (ppb) of (a) NO, NO<sub>2</sub>; (b) O<sub>3</sub> and (c) NO<sub>x</sub>, O<sub>3</sub>, using the LES and box model dynamical frameworks, and the RCS and O<sub>3</sub>-NO<sub>x</sub>-only chemistry cases.

**Figure 10**

**Fig. 10.** Time averaged vertical mixing ratio profiles ( $1.0 \leq z/H \leq 2.0$ ) of (a)  $O_3$ , (b)  $NO$ , (c)  $NO_2$ , (d)  $NO_x$  and (e)  $O_x$  (ppb) at the canyon inlet ( $x/W = -0.5$ ) and canyon outlet ( $x/W = 0.5$ ), from the LES-RCS simulations.

Figure 11



**Fig. 11.** Vertical profiles of advective, turbulent and total flux (ppb m<sup>-2</sup> s<sup>-1</sup>) averaged across the canyon ( $-0.5 \leq x/W \leq 0.5$ ), from the LES-RCS simulation.



**Table 1**

Initial mixing ratios (ppb) used in the RCS model simulations

Species	Chemical formula	Mixing ratio / ppb
Nitric oxide	NO	2
Nitrogen dioxide	NO <sub>2</sub>	8
Ozone	O <sub>3</sub>	40
Carbon monoxide	CO	200
Nitric acid	HNO <sub>3</sub>	2
Methane	CH <sub>4</sub>	1800
Water vapour	H <sub>2</sub> O	2 %
<b>VOCs</b>		
Ethene	C <sub>2</sub> H <sub>4</sub>	0.91
Propene	C <sub>3</sub> H <sub>6</sub>	0.29
Formaldehyde	HCHO	3.14
Acetaldehyde	CH <sub>3</sub> CHO	2.98
Isoprene	C <sub>5</sub> H <sub>8</sub>	0.28
Methanol	CH <sub>3</sub> OH	7.38
Ethanol	C <sub>2</sub> H <sub>5</sub> OH	2.37
Peroxyacetyl nitrate	PAN	0.46

**Table 2**

The dominant OH production and loss rates calculated for the LES-RCS simulation. Average concentrations taken over the final hour of the model simulation ( $150 \leq t \leq 210$  min) and at three locations (Fig. 5): within the canyon vortex (V), toward the lower leeward wall (L) and in the background atmosphere (B).

	(V) Vortex	(L) Lower leeward wall	(B) Background atmosphere
Rate of production / loss (ppt s <sup>-1</sup> )			
<b>Production</b>			
$\text{HO}_2 + \text{NO} \rightarrow \text{OH} + \text{NO}_2$	100	174	4.8
$\text{HONO} + h\nu \rightarrow \text{OH} + \text{NO}$	27	25	0.74
$\text{O}_3 + h\nu \rightarrow \text{OH} + \text{OH}$	0.79	0.78	1.3
$\text{O}_3 + \text{VOC} \rightarrow \text{OH} + \text{products}$	0.35	0.57	0.0060
<b>Loss</b>			
$\text{OH} + \text{VOC} \rightarrow \text{products}$	-61	-96	-2.8
$\text{OH} + \text{NO} \rightarrow \text{HONO}$	-41	-71	-0.78
$\text{OH} + \text{NO}_2 \rightarrow \text{HNO}_3$	-22	-26	-1.9
$\text{OH} + \text{CO} \rightarrow \text{HO}_2$	-5.0	-7.4	-1.0

**Table 3**

Canyon- and time-averaged mixing ratios calculated using the LES-RCS and Box-RCS model approaches

	(a) LES	(b) Box	(b) - (a)	[(b) - (a)] / (a)
	Mixing ratio (ppb)			%
O <sub>3</sub>	11.2	10.5	-0.7	-5.9
NO	168	170	2	1.2
NO <sub>2</sub>	68	74	6	9.5
OH*	0.08	0.09	0.01	11.3
HO <sub>2</sub> *	0.23	0.25	0.02	8.1
NO <sub>x</sub>	236	244	8	3.3
O <sub>x</sub>	79	85	6	8.0
NO <sub>2</sub> /NO	0.40	0.44		

\*Mixing ratios of OH and HO<sub>2</sub> are given in ppt.

**Table 4**

Effect of exchange velocity: canyon and time averaged mixing ratios for the Box-RCS simulation with  $\omega_t = 0.021 \text{ m s}^{-1}$  and  $\omega_t = 0.022 \text{ m s}^{-1}$ .

	(a) Box $\omega_t = 0.021 \text{ m s}^{-1}$	(b) Box $\omega_t = 0.022 \text{ m s}^{-1}$	(b) - (a)	[(b) - (a)] / (a)
	Mixing ratio (ppb)			%
O <sub>3</sub>	10.54	10.76	0.22	2.1
NO	169.9	161.8	-8	-4.8
NO <sub>2</sub>	74.0	71.6	-2	-3.2
OH*	0.089	0.088	-0.0012	-1.3
HO <sub>2</sub> *	0.247	0.246	-0.0006	-0.3
NO <sub>x</sub>	244	233	-11	-4.5
O <sub>x</sub>	85	82	-3	-3.5

\*Mixing ratios of OH and HO<sub>2</sub> are given in ppt.

**Table 5**

Comparison of mean (canyon- and time-averaged) mixing ratios, for the RCS and O<sub>3</sub>-NO<sub>x</sub>-only mechanisms, using the LES and Box model dynamical frameworks. .

	(a) LES RCS	(b) Box RCS	(c) LES O <sub>3</sub> -NO <sub>x</sub>	(d) Box O <sub>3</sub> -NO <sub>x</sub>	[(a) - (c)] / (c)	[(b) - (d)] / (d)
	Mixing ratio (ppb)				%	
O <sub>3</sub>	11.2	10.5	9.7	8.4	15.5	25.0
NO	168	170	177	185	-5.1	-8.1
NO <sub>2</sub>	68	74	61	63	11.5	17.5
NO <sub>x</sub>	236	244	238	248	-0.8	-1.6
O <sub>x</sub>	79	85	70.7	71.4	11.7	19.0
NO <sub>2</sub> /NO	0.40	0.44	0.34	0.34		