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Large eddy simulation of reactive pollutants in a deep urban street canyon: Coupling dynamics with $O_3$-$NO_x$-VOC chemistry

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Abstract:

A large eddy simulation (LES) model coupled with $O_3$-$NO_x$-VOC chemistry is implemented to simulate the coupling effect of emissions, mixing and chemical pre-processing within an idealised deep (aspect ratio = 2) urban street canyon under a weak wind condition. Reactive pollutants exhibit significant spatial variations in the presence of two vertically aligned unsteady vortices formed in the canyon. Comparison of the LES results from two chemical schemes (simple NO$_x$-$O_3$ chemistry and a more comprehensive Reduced Chemical Scheme (RCS) chemical mechanism) shows that the concentrations of NO$_2$ and O$_3$ inside the street canyon are enhanced by approximately 30-40% via OH/HO$_2$ chemistry. NO, NO$_x$, O$_3$, OH and HO$_2$ are chemically consumed, while NO$_2$ and O$_3$ (total oxidant) are chemically produced within the canyon environment. The within-canyon pre-processing would increase oxidant fluxes released from the canyon to the overlying boundary layer, and this effect is more crucial for deeper street canyons (as found in many traditional European urban centres) than shallower (lower aspect ratio) streets. There is clear evidence of distinct behaviours for emitted chemical species and entrained chemical species, and positive (or negative) values of intensities of segregations are found between pairs of species with a similar (or opposite) behaviour. The simplified two-box model underestimated NO and O$_3$ levels, but overestimated NO$_2$ levels for both the lower and upper canyon compared with the more realistic LES-chemistry model. This suggests that the segregation effect due to incomplete mixing reduces the chemical conversion
rate of NO to NO$_2$. This study reveals the impacts of nonlinear O$_3$-NO$_x$-VOC photochemical processes in the incomplete mixing environment and provides a better understanding of the pre-processing of emissions within canyons, prior to their release to the urban boundary layer, through the coupling of street canyon dynamics and chemistry.

Capsule:

The impacts of nonlinear O$_3$-NO$_x$-VOC photochemical processes in an incomplete mixing environment are revealed by an LES-chemistry model.

Keywords:

Large eddy simulation; Street canyon; Nonlinear photochemistry; Segregation effect.
1 Introduction

Urban air pollution associated with road transport is a major environmental issue (Murena et al., 2009). Air pollutants in urban areas may be classified into two categories. The first category is primary air pollutants (Mayer, 1999), which are released directly into the atmosphere from emission sources (normally resulting from the combustion of fuels) and consist of nitrogen oxides (NO\textsubscript{x}) - primarily NO, volatile organic compounds (VOCs), carbon monoxide (CO) and components of particulate matter (PM) (Dunmore et al., 2015). The second category is secondary air pollutants, which are produced into the atmosphere while primary air pollutants undergo chemical reactions (Jacobson, 2005). As one of the important secondary air pollutants, ozone (O\textsubscript{3}) is formed by chemical reactions involving primarily the oxidation of VOCs and NO\textsubscript{x} while the sunlight is present. Fast-reacting chemical radicals (e.g. the hydroxyl radical (OH), and hydroperoxy radical (HO\textsubscript{2})) govern the cycle of the chemical degradation of VOCs (O\textsubscript{3} precursors) converting NO to NO\textsubscript{2} and hence O\textsubscript{3} formation (Bloss, 2009).

The street canyon is normally the basic geometric unit in the built environment of many cities and typically describes a place with surrounding buildings on both sides of the street (Li et al., 2008b). Buildings are the artificial obstacles to urban atmospheric flow (Salim et al., 2011) and give rise to limited ventilation, especially for deep street canyons. The deterioration of urban air quality occurs due to the combined effects of traffic emissions, dynamics and chemistry within such an atmospheric compartment (Li et al., 2008b). Investigation of urban air pollution in street canyons has become a focus of environmental research (Zhong et al., 2016). A variety of approaches, from full-scale field measurements, reduced-scale physical modelling (e.g. water-channel experiments and wind-tunnel experiments), numerical modelling (e.g. Computational fluid dynamics (CFD) models) to parametric (operational) modelling (e.g. street canyon box models), have been undertaken over recent years to investigate street canyon air quality in urban areas (Vardoulakis et al., 2003; Ahmad et al., 2005; Li et al., 2006; Yazid et al., 2014). Details of individual aspects of these approaches can be found in several reviews, e.g. Vardoulakis et al. (2003) focused on full-
scale field measurements and parametric (operational) modelling; Ahmad et al. (2005) examined wind tunnel experiments; Li et al. (2006) used CFD models to understand the dynamical processes and Yazid et al. (2014) carried out field measurements and modelling studies. Recently, Zhong et al. (2016) presented a comprehensive review of numerical modelling studies that couple street canyon dynamics with chemistry of reactive pollutants. Coupling dynamics and chemistry accommodates the mixing, and photo-chemical processes of emissions. The CFD approach (including mainly Reynolds-Averaged Navier-Stokes (RANS) and Large-eddy Simulation (LES) models) has become a powerful tool to investigate high spatial and temporal resolution simulations of wind and pollutant fields within urban street canyons (e.g. Bright et al., 2013; Kwak et al., 2013; Li et al., 2012). RANS only provides the information about the mean time-averaged properties, while LES can predict the additional unsteadiness and intermittency of turbulence (Li et al., 2006). However, RANS is normally computationally faster than LES (Li et al., 2006).

Zhong et al. (2015) employed an LES model to investigate the dispersion and transport of the reactive species NO, NO2 and O3 in a deep urban street canyon by coupling with simple NOx-O3 chemistry. The turbulent flow derived from the LES model was evaluated against a water-channel experiment (Li et al., 2008a) and the model agreed well with the experiment in terms of the velocities, turbulent intensities and flow structure (Zhong et al., 2015). There were spatial distributions of reactive pollutants driven by the two canyon vortices. A simplified two-box model coupled with the NOx-O3 chemistry was then developed in the presence of two vortices. This study further extends the framework used in Zhong et al. (2015), with a focus upon the coupling of the O3-NOx-VOC chemistry, rather than simple NOx-O3 chemistry.

2 Methodology

2.1 Large-Eddy simulation

The numerical model employed to simulate the incompressible turbulent flow under neutral conditions in the canyon is based on the LES technique, which computes the larger, grid-resolved
eddies explicitly and parameterises the smaller, unresolved eddies (Liu et al., 2005). The one-equation sub-grid scale (SGS) turbulence model is adopted to model the SGS motions. The near-wall treatment is based on the logarithmic law of the rough-wall model (Schlichting and Gersten, 2000). The LES model is solved by OpenFoam v2.1.1 (OpenFOAM, 2012). The detailed numerical model is described by Zhong et al. (2015).

2.2 Coupling with O₃-NOₓ-VOC chemistry

The filtered governing equations for the transport of reactive pollutants are:

\[
\frac{\partial \overline{c_i}}{\partial t} + \frac{\partial}{\partial x_j} (\overline{u_j c_i}) = \frac{\partial}{\partial x_j} \left( \left[ \nu + \nu_{\text{SGS}} \right] \cdot \frac{\partial \overline{c_i}}{\partial x_j} \right) + \Delta S_i + E_i
\]  

(1)

Here, \( \overline{c_i} \) is the resolved-scale concentration of the \( i^{th} \) chemical species; \( Sc (=0.72) \) is the Schmidt number (Liu et al., 2005); \( \Delta S_i \) is the chemical source term of the \( i^{th} \) chemical species; \( E_i \) is the emission of the \( i^{th} \) chemical species; \( j (=1, 2, 3) \) denotes the Cartesian component. Due to the computational limitation of LES, a reduced chemical scheme (RCS) with 51 chemical species and 136 chemical reactions (Bright, 2013) is employed as the chemical mechanism in the present study. The RCS was developed and reduced based on the Common Representative Intermediates mechanism version CRI v2-R5 (Jenkin et al., 2008), which included 555 chemical reactions of 196 species. This CRI v2-R5 is a subset of the near-explicit chemical mechanism, i.e. the Master Chemical Mechanism (MCM) (Jenkin et al., 1997) which involves about 13,500 chemical reactions of 5,900 species for MCM v3.1 (Pinho et al., 2007). The MCM has been evaluated against multiple chamber and field datasets (Bloss et al., 2005; Saunders et al., 2003; Jenkin et al., 1997; Jenkin et al., 2003) and can be a benchmark mechanism for the development and evaluation of reduced chemical mechanisms. Both the RCS and CRI have been evaluated regarding the abundance of key oxidants for the daytime scenarios compared to the MCM (Bright, 2013; Bright et al., 2013). The largest difference of OH between the RCS (reduced from CRI) and MCM was about 6% over a four
hour box model simulation (Bright, 2013; Bright et al., 2013), which was acceptable compared with the errors associated with the OH measurement of 7-16% (Heard and Pilling, 2003).

A challenge to solve the transport equations for reactive pollutants (especially fast-reacting chemical radicals, e.g. OH and HO$\textsubscript{2}$) is to derive the chemical source terms. According to Hertel et al. (1993), the ordinary differential equations of a chemical system can be described as follows:

$$\frac{dc_i}{dt} = P_i - L_i c_i \quad i = 1, 2, ..., nc$$  \hspace{1cm} (2)

Here, $nc$ is the total number of the chemical species, $i$ represents the $i^{th}$ chemical species, $c_i$ is the concentration, $P_i$ is the chemical production term and $L_i$ is the chemical loss rate ($L_i c_i$ therefore represents the chemical loss term). Both $P_i$ and $L_i$ are non-negative functions of concentrations, i.e.

$$P_i = P_i(t, c_1, c_2, ..., c_{nc})$$  \hspace{1cm} (3)

$$L_i = L_i(t, c_1, c_2, ..., c_{nc})$$  \hspace{1cm} (4)

The chemical timescale of the $i^{th}$ chemical species $\tau_i$ (Neophytou et al., 2004) is defined as follows:

$$\tau_i = \frac{1}{L_i}$$  \hspace{1cm} (5)

The chemical system is normally stiff due to the variability (from very fast to rather slow) of the chemical time scale (Verwer and Simpson, 1995). The QSSA (quasi-steady-state approximation) algorithm has been widely used to handle stiff chemical systems in air pollution modelling (e.g. Hesstvedt et al., 1978; Verwer and Vanloon, 1994). The QSSA algorithm is described below.

$c_i^n$ denotes the concentration of the $i^{th}$ chemical species at $t = t_n$. It is assumed that $P_i$ and $L_i$ in Equation (2) are constant over a given time step $\Delta t$ (Hertel et al, 1993),

$$t_{n+1} = t_n + \Delta t$$  \hspace{1cm} (6)
Equation (2) may be solved analytically by the following formula:

\[ c_i^{n+1} = \frac{P_i^n}{L_i^n} + (c_i^n - \frac{P_i^n}{L_i^n})e^{-\Delta t L_i^n} \]  

(7)

Depending on the chemical timescales and time steps, three categories of formulae are derived (Alexandrov et al., 1997). (i) If \( \tau_i < \frac{\Delta t}{10} \), it means that the chemical reaction is very fast over the given time step. The steady state at the end of the time step can be assumed and Equation (7) can be expressed by the following approximation,

\[ c_i^{n+1} = \frac{P_i^n}{L_i^n} \]  

(8)

(ii) If \( \Delta t/10 \leq \tau_i \leq 100\Delta t \), it means that the chemical reaction is at a medium rate over the given time step and Equation (7) is applied. (iii) If \( \tau_i > 100\Delta t \), it means that the chemical reaction is rather slow over the given time step and the forward Eulerian formula can be employed,

\[ c_i^{n+1} = c_i^n + (P_i^n - L_i^n c_i^n)\Delta t \]  

(9)

The QSSA algorithm has simple formulae and can be easily employed in large air pollution models. However, there are also some drawbacks. At each cell for each species, there are three formulae to be conditionally determined. It demands more computational time for the air pollution models with a huge number of cells. In addition, the computational cost to compute the exponential function in Equation (7) is expensive. Therefore, an attempt has been conducted (Alexandrov et al., 1997) to improve the performance of the QSSA algorithm. The exponential function can be rationally approximated by the following expression based on the Taylor expansion in the second order:

\[ e^{-\Delta t L_i^n} \approx \frac{1}{1 + L_i^n \Delta t + 0.5(L_i^n \Delta t)^2} \]  

(10)

Then Equation (7) can be rewritten as:
In this study, the chemical species in the RCS chemical mechanism can be separated into two groups, i.e. slower chemical species (e.g. NO\textsubscript{x} and O\textsubscript{3}) and faster chemical species (e.g. OH and HO\textsubscript{2}). For slow chemical species, a time step of 0.03 s (Bright et al., 2013) is adopted and Equation (9) is used. For fast chemical species, the above-mentioned QSSA algorithm with Equation (11) (with a smaller time step of 0.003 s) (Bright et al., 2013) is adopted.

\[ c_i^{n+1} = c_i^n + \frac{(1 + 0.5L_i^2\Delta t)P_i^n\Delta t}{1 + L_i^2\Delta t + 0.5(L_i^2\Delta t)^2} \] (11)

### 2.3 Model setup

The computational domain in the study is same as that in Zhong et al. (2015), shown as Figure S1 (in the Supplementary Materials), which consists of an idealised deep street canyon (AR=2, i.e. \( H=36 \) m and \( W=18 \) m). A constant pressure gradient above the canyon (perpendicular to the street axis) is used to drive the street canyon flow. Symmetry boundary conditions are used at the domain top for both the flow and pollutants. Cyclic boundary conditions are used in both the \( x \)- and \( y \)-directions for the flow. For pollutants, the inlet adopts “fixed-value” boundary conditions. A photochemical box model (including the RCS mechanism) is run without emissions for 30 mins in order to achieve a photochemical pseudo-equilibrium condition. Then concentrations of all chemical species at \( t=30 \) min are used as their inlet boundary conditions. For the outlet, the advective boundary condition

\[ \frac{\partial \tilde{c}_i}{\partial t} + \bar{u} \frac{\partial \tilde{c}_i}{\partial x} = 0 \] (12)

is employed. On the solid boundaries, the near-wall treatment (Schlichting and Gersten, 2000) is employed for the flow and zero-gradient conditions are used for pollutants (assuming no pollutant deposition). At \( t=30 \) min, the chemistry mechanism and emissions modules are switched on in the presence of canyon dynamics, i.e. a statistically steady turbulent flow derived from the LES model (Zhong et al., 2015), for a 210 min duration window (\( t = 30 \) to 240 min).
Emissions are represented by two traffic lanes with a Gaussian distribution located at $z = 1$ m and at 2.5 m from both sides of the street centre (Zhong et al., 2015). Based on the UK Road Vehicle Emission Factors (Boulter et al., 2009), emission rates for NO$_x$, VOCs and CO of 620, 128 and 1,356 g km$^{-1}$ hr$^{-1}$ (equivalent to 1,000 ppb s$^{-1}$, 3,593 ppb s$^{-1}$, and 691 ppb s$^{-1}$ respectively released into a typical LES model cell, i.e. $0.3 \times 1 \times 0.3$ m in the $x$-, $y$- and $z$-directions) were used. This emission scenario approximately represents urban continuous road traffic of 1,500 vehicles hr$^{-1}$ with an average speed of 30 mph (Boulter et al., 2009). The fractional NO$_x$ emission rate by volume is 90% for NO and 10% for NO$_2$ (Baker et al., 2004). The fractional VOCs emission rate by volume is 44% for ethane (C$_2$H$_6$), 19% for propene (C$_3$H$_6$), 25% for formaldehyde (HCHO) and 12% for acetaldehyde (CH$_3$CHO) (Bright et al., 2013).

Computation of the LES model coupling the RCS mechanism (LES-chemistry) was performed in the University of Birmingham's Linux-based High Performance Computing (HPC) cluster. In this research, the total number of cores used was 32, i.e. the LES-chemistry model was run in parallel on 2 nodes, each of which consists of 16 cores with 64-bit 2.2 GHz processors and 32 GB of memory. The total computation time for the LES-chemistry was about 10 days.

### 3 Post-processing of LES output

The 3-D fields (at each grid) for wind and concentrations were stored in interval time step of 3s and the final hour of the simulation ($t = 180$ to 240 min) was used to analyse the resolved-scale turbulent statistics. The temporal average over $t \in [t_1, t_2]$ and spatial average over $y \in [0, L_y]$ of any resolved-scale quantity $\tilde{\phi}$ gives a 2D function $\langle \tilde{\phi} \rangle(x, z)$, i.e.

$$
\langle \tilde{\phi} \rangle(x, z) = \frac{1}{L_y (t_2 - t_1)} \int_{t_1}^{t_2} \int_{0}^{L_y} \tilde{\phi}(x, y, z, t) dy dt
$$

(13)
and $\phi(x,y,z,t) = \bar{\phi}(x,y,z,t) - \langle \tilde{\phi} \rangle(x,z)$ denotes the resolved-scale fluctuation component of $\bar{\phi}$ about $\langle \tilde{\phi} \rangle$.

3.1 Decomposition of contributions from emission and chemistry

For a passive scalar (i.e. a non-reactive scalar), abundance within a street canyon is determined by the emission rate and the background concentration of the passive scalar once the street canyon turbulent flow is given. We denote its $t$- and $y$-averaged spatial field as $\langle \bar{C}_{ps} \rangle$. For a reactive scalar (e.g. NO$_2$), in addition to the contribution from its emission rate and background concentration, the contribution from chemistry also affects its abundance within the street canyon. If the mean concentration for any chemical species inside an idealised 2D street canyon derived from LES is denoted by $\langle \bar{C} \rangle(x,z)$, we can then define the contribution to its abundance from chemistry, denoted by $\langle \bar{C}_{chem} \rangle$, using the following equation (without loss of generality, the overbar and the angle brackets of all terms are dropped for the purpose of presentation):

$$C_{chem}(x,z) = C(x,z) - C_{ps}(x,z) \quad (14)$$

Further,

$$C_{ps}(x,z) = C_b(x,z) + C_{emn}(x,z) \quad (15)$$

Here, ‘b’ is for ‘background’ and ‘emn’ is for ‘emission’, $C_{emn}(x,z)$ is the field of the passive scalar induced by the given emission and corresponding to ‘zero background’. $C_b(x,z)$ is the spatial distribution induced by an upwind background concentration of $C_0$. It is assumed that

$$C_b(x,z) = C_0 = const \quad (16)$$

This assumption is true if the system is allowed to achieve a steady state; air inside the canyon will be in balance with the above-roof concentration and should be constant. Thus,
\[ C_p(x, z) = C_0 + C_{em}(x, z) \]  

(17)

It is assumed that \( C_{em}(x, z) \) linearly scales with the emission rate. It is further defined that \( C_{em,1}(x, z) \) is the concentration for a unit emission rate and \( C_{em,1}(x, z) \) can be derived from the LES simulation for a passive scalar (with zero background). Therefore,

\[ C_{em}(x, z) = E \cdot C_{em,1}(x, z) \]  

(18)

where \( E \) is the emission rate for this species. In such a way, a spatial pattern of a passive scalar can be used to reconstruct the pattern for any other passive scalars, or the emission-induced component of a non-passive scalar.

Therefore, the contribution from chemistry to any non-passive scalar can be diagnosed by following equation:

\[ C_{chem}(x, z) = C(x, z) - E \cdot C_{em,1}(x, z) - C_0 \]  

(19)

in which both \( C(x, z) \) and \( C_{em,1}(x, z) \) are derived from LES with prescribed \( E \) and \( C_0 \). This formula can be applied for all emitted species (e.g. NO, NO\(_2\), NO\(_x\) and O\(_x\)). In particular, for any non-emitted species (e.g. O\(_3\), OH and HO\(_2\)), the contribution from emission is effectively ignored and therefore the contribution from chemistry is simply obtained:

\[ C_{chem}(x, z) = C(x, z) - C_0 \]  

(20)

A negative (positive) value of \( C_{chem} \) means the chemical consumption (production). A zero value of \( C_{chem} \) means no chemical consumption or production. For higher-order reactions, \( C_{chem} \) is also a function of the background concentrations and the emission rates of relevant chemical species. This complexity due to the nonlinearity of the chemical processes, however, does not limit the
application of such analysis for the purpose of diagnosis under a given set of conditions (i.e. background concentrations and emission rates).

The Damköhler number (Da), defined as the ratio of the turbulent mixing timescale ($T_{mix}$) to the chemical timescale ($T_{chem}$), can be used to investigate the combined effect between dynamics and chemistry (Auger and Legras, 2007). When Da $\ll$ 1, chemical processes are rather slow compared with dynamical processes and chemical species may be regarded as well-mixed scalars with minimal segregation effects. When Da $\gg$ 1, chemistry is very fast and can achieve a chemical equilibrium before the flow mixes together chemical species. In such situations, the interaction between dynamics and chemistry is very important with substantial segregation effects.

3.2 Vertical advective and turbulent fluxes of pollutants

The vertical advective flux of any species at the resolved-scale is defined:

$$ F_{adv}(x, z) = \langle w \rangle(x, z) \langle \phi \rangle(x, z) $$  \hspace{1cm} (21)

and the vertical turbulent flux is defined:

$$ F_{turb}(x, z) = \left\langle w' \phi' \right\rangle(x, z) = \frac{1}{L_y(t_2 - t_1)} \int_{t_1}^{t_2} \int_{y_1}^{y_2} w'(x, y, z,t) \phi'(x, y, z,t) dy dt $$  \hspace{1cm} (22)

Thus the vertical total flux is obtained as follows:

$$ F_{total}(x, z) = F_{turb}(x, z) + F_{adv}(x, z) $$  \hspace{1cm} (23)

These quantities of fluxes represent the 2D spatial variation. For the purpose of discussion, these quantities are further averaged horizontally and vertical profiles are derived:

$$ F(z) = \frac{1}{W} \int_{-0.5W}^{0.5W} F(x, z) dx $$  \hspace{1cm} (24)
3.3 Intensity of segregation

In order to characterise the segregation effect due to incomplete mixing of chemical species, a widely used dimensionless number, the intensity of segregation (Krol et al., 2000) between two chemical species A and B, \( I_{S(A+B)} \), is introduced and defined as:

\[
I_{S(A+B)} = \frac{1}{W \cdot H} \int_{-0.5W}^{0.5W} \int_{0}^{H} \frac{A'(x,z) \cdot B'(x,z)}{[A] \cdot [B]} \, dx \, dz
\]  

(25)

where \([A]\) and \([B]\) represent the canyon averages of \( \langle A \rangle(x,z) \) and \( \langle B \rangle(x,z) \), respectively, in which \( \langle A \rangle(x,z) \) and \( \langle B \rangle(x,z) \) are derived from Equation (13); the prime \( A'(x,z) = \langle A \rangle(x,z) - [A] \) (or \( B'(x,z) = \langle B \rangle(x,z) - [B] \)) denotes the local spatial deviation from the canyon-averaged concentration, and \( A'(x,z) \cdot B'(x,z) \) stands for the spatial covariance between A and B. The intensity of segregation is a proper measure of the effect of spatial segregation on nonlinear chemical processes (Hilst, 1998) and represents the deviation from a well-mixed environment due to the coupling between dynamics and chemistry (Zhong et al., 2014). For a second-order reaction \( A+B \rightarrow C \) in a heterogeneous system, the rate of formation of C (Vinuesa and de Arellano, 2005) evaluated for the whole volume of air inside the street canyon in the framework of one-box model can be described as follows,

\[
\frac{d[C]}{dt} = k_{eff(A+B)}[A][B]
\]  

(26)

where \( k_{eff(A+B)} \) is the effective second-order rate constant for formation of C in the heterogeneous system, i.e.

\[
k_{eff(A+B)} = k_{(A+B)}(1 + I_{S(A+B)})
\]  

(27)
where $k_{(A+B)}$ is the original rate constant of the reaction in a well-mixed system. Such a constant is normally obtained from laboratory experiments in a well-mixed chamber. If $I_{S(A+B)} = 0$, it means that species A and B can be regarded as well-mixed; If $I_{S(A+B)} > 0$ or $I_{S(A+B)} < 0$, it implies that $k_{(A+B)}$ in the heterogeneous system is larger or smaller than $k_{(A+B)}$ in the well-mixed system due to the effect of segregation. Segregation effect for the street canyon environment will be investigated in this study.

4 Results and discussion

4.1 Spatial variation of reactive pollutants

4.1.1 Spatially and temporally averaged concentrations

Figure 1 illustrates the spatial variation of (a) $\langle NO \rangle$, (b) $\langle NO_2 \rangle$, (c) $\langle O_3 \rangle$, (d) $\langle NO_x \rangle$, (e) $\langle O_x \rangle$, (f) $\langle NO \rangle/\langle NO_2 \rangle$, (g) $\langle OH \rangle$ and (h) $\langle HO_2 \rangle$ (See Equation (13); averaged from 180 to 240 min).

These plots show the influence of two primary vortices on pollutant dispersion as observed by Zhong et al. (2015) with simple NO$_x$-O$_3$ chemistry. Figure 1a-c show similar spatial patterns of NO, NO$_2$ and O$_3$ as those in Zhong et al. (2015) (Figure 4a-c in their study). There are sinks of NO$_x$ and sources of O$_x$ in the current study (with the O$_3$-NO$_x$-VOC chemistry). However, NO$_x$ and O$_x$ are effectively conserved in Zhong et al. (2015) (with simple NO$_x$-O$_3$ chemistry). NO$_x$ plays a key role in the street-canyon atmospheric chemistry. NO$_2$ levels are largely determined by the within-canyon processing through the chemical reactions of NO with other species (e.g. O$_3$ titration and also OH/HO$_2$ chemistry). Both NO$_x$ and O$_x$ are useful measures of the street-canyon atmospheric chemistry (Figure 1d-e). The ratio of NO/NO$_2$ (Figure 1f) is a useful indicator of chemical interactions within the street canyon, reflecting the conversion of NO to NO$_2$ through chemistry.

The NO/NO$_2$ ratio also had a similar pattern driven by two vortices, ranging from about 3.6 (vs 6 in Zhong et al., 2015) at the right of lower canyon to about 1.4 (vs 3 in Zhong et al., 2015) at the
The NO/NO\textsubscript{2} ratio within the street canyon was much lower compared to the raw emission ratio of NO/NO\textsubscript{2} (assumed as a value of 9). This reflected the contributions of directly emitted NO\textsubscript{2} and chemical oxidation of emitted NO to increased levels of NO\textsubscript{2}. The fast reacting chemical radicals (OH and HO\textsubscript{2}) played a key role in the additional conversion of NO to NO\textsubscript{2} through OH/HO\textsubscript{2} chemistry. The spatial distributions of OH and HO\textsubscript{2} (Figure 1g-h) had similar patterns to that of O\textsubscript{3}, in which their background mixing ratios were much higher than those within the canyon (averagely by a factor of about 2 for OH, 5 for HO\textsubscript{2} and 3 for O\textsubscript{3}).

### 4.1.2 Simple NO\textsubscript{x}-O\textsubscript{3} chemistry vs O\textsubscript{3}-NO\textsubscript{x}-VOC chemistry

Figure 2 illustrates spatial variations of the overestimation of the spatially and temporally averaged concentrations (%) by simple NO\textsubscript{x}-O\textsubscript{3} chemistry (Zhong et al., 2015) compared with the RCS mechanism (O\textsubscript{3}-NO\textsubscript{x}-VOC chemistry) in this study for (a) \(\langle\text{NO}\rangle\), (b) \(\langle\text{NO}\textsubscript{2}\rangle\), (c) \(\langle\text{O}\textsubscript{3}\rangle\), (d) \(\langle\text{NO}\textsubscript{x}\rangle\), (e) \(\langle\text{O}\textsubscript{x}\rangle\) and (f) \(\langle\text{NO}\rangle/\langle\text{NO}\textsubscript{2}\rangle\). Generally, simple NO\textsubscript{x}-O\textsubscript{3} chemistry overestimates the levels of NO, NO\textsubscript{x} and NO/NO\textsubscript{2}, but underestimates the levels of NO\textsubscript{2}, O\textsubscript{3} and O\textsubscript{x}. Such findings suggest that using simple NO\textsubscript{x}-O\textsubscript{3} chemistry may provide a reasonable prediction of air pollution in street canyon (for NO\textsubscript{2} - i.e. predicted levels are biased low) while in reality the NO\textsubscript{2} level may exceed the air quality standards, which may mislead a policy-maker to make an inappropriate decision with respect to air quality management. There are some common features for these overestimations (biases) in Figure 2. At the canyon roof level, a sharp decrease of the magnitudes of those biases was observed and those values approach to zero for the wider background. The largest values of the magnitudes of overestimation (about 30\% for NO, about -38\% for NO\textsubscript{2}, about -52\% for O\textsubscript{3}, about 4\% for NO\textsubscript{x}, about -40\% for O\textsubscript{x} and about 115\% for NO/NO\textsubscript{2}) were found close to the centre of the upper vortex.

For air pollution problems related to higher levels of NO\textsubscript{2} in urban areas, an underestimation of NO\textsubscript{2} by 40\% could be a substantial issue (Defra, 2008). In the lower part of the canyon, the magnitudes of those overestimations were comparatively low and generally decrease down to the street ground - this is of significance as the region in which receptor / population exposure occurs (at heights of 1 - 2 m). It is noted that there was a slight overestimation for NO\textsubscript{x} by the simple chemistry and this was...
due to the extra sink of NO, to other N-contained species (such as Nitric acid (HNO₃) and Nitrous acid (HONO)) presented in the more comprehensive RCS. It is interesting that there is a large underestimation for the oxidants (NO₂, O₃ and Oₓ) by the simple chemistry. This is attributed to the additional conversion of NO to NO₂ by the OH/HO₂ chemistry in the RCS, exacerbated by limited exchange in this deep canyon scenario.

4.1.3 Vertical profiles of concentrations

Figure 3 depicts vertical profiles of (a) \( \langle NO \rangle \), (b) \( \langle NO₂ \rangle \), (c) \( \langle NOₓ \rangle \), (d) \( \langle Oₓ \rangle \), (e) \( \langle NO \rangle / \langle NO₂ \rangle \), (f) \( \langle C \rangle / C_{b} \) along the leeward and windward buildings, respectively. For the upper (or lower) vortex, NO, NO₂, NOₓ, Oₓ and NO/NO₂ at the vicinity of the leeward building were generally higher (lower) than those of the windward building (Figure 3a-e). But for O₃, the situation is reversed (Figure 3f).

For the AR=1 case, there are higher concentrations of emitted pollutants towards the leeward building at the pedestrian level (e.g. Baker et al., 2004; Kwak and Baik, 2012). However, for the AR=2 case, higher concentrations of emitted pollutants (e.g. NOₓ) are observed towards the windward building at the pedestrian level (due to the opposite direction between the upper vortex and lower vortex) (Figure 3a-e). It was also interesting to note that just above the canyon roof level (z/W=2), there were much higher levels of pollutants (e.g. NOₓ) at the windward side (i.e. the canyon outlet) than those at the leeward side (i.e. the canyon inlet) (Figure 3a-e). This reflected increased levels of pollutants transferred from the canyon to the wider ambient environment, which highlighted the importance of the coupling effect of emissions, mixing and chemical pre-processing within the street canyon. The windward side was the main location of this street canyon ventilation system, potentially taking ambient air into buildings. The vertical profiles of OH and HO₂ have similar patterns to that of O₃, in which their levels along both the leeward and windward walls increased with the vertical height and approach to their corresponding background concentrations at approximately z/W=2.4. In the upper (or lower) part of the canyon, the mixing ratios of OH and HO₂ along the windward wall were slightly higher (or lower) than those along the leeward wall. For
comparison, levels of O$_3$, OH and HO$_2$ are normalised by their background concentrations (Figure 3f). It was observed that HO$_2$ had the sharpest drop near the canyon roof level while O$_3$ decreased the least. This suggested that either HO$_2$ or OH is more rapidly consumed (sharp shift) than O$_3$ at the canyon roof level as the above canyon air is entrained into the canyon. Within the canyon, there was a similar consumption rate for OH and HO$_2$, but much slower than that for O$_3$. The rapid O$_3$ consumption inside the canyon can be explained by NO$_2$ (NO) emissions from the street level, which had a strong titration effect thereby leading to the rapid consumption of O$_3$.

### 4.1.4 Contributions from emission and chemistry

Figure 4 illustrates the spatial variations of (a) $C_{\text{emn},1}$ and $C_{\text{chem}}$ of (b) NO, (c) NO$_2$, (d) NO$_x$, (e) O$_3$, (f) O$_3$, (g) OH and (h) HO$_2$ (See Equations (18), (19) and (20)). The spatial pattern of a passive scalar with a unit emission rate is depicted in Figure 4a. It was observed that the distribution of the passive scalar can be characterised by the two unsteady vortices formed inside the street canyon (Cross-sectional flow structure can be seen from Fig. 3 of the LES simulation in Zhong et al. 2015). The emission of the passive scalar was mainly trapped inside the lower vortex closer to the windward wall. This unit emission rate scenario was used to reconstruct the spatial pattern based on Equation (19) for the chemistry-induced component of a non-passive scalar in Figure 4 b-e. It was found that NO, NO$_x$, O$_3$, OH and HO$_2$ were chemically consumed (negative values of $C_{\text{chem}}$). For NO$_2$ and O$_x$, however, chemical production occurred inside the street canyon (positive values of $C_{\text{chem}}$). The chemical consumption of NO and O$_3$ was largely caused by the titration effect and by the (slow, but important near emission source) NO + NO + O$_2$ reaction. The concentration contributed from the chemical consumption for NO$_x$ was about 14% of that for NO. The slight chemical consumption of NO$_x$ indicated that there was a sink of NO$_x$ in the chemical processing, but this rate was relatively slow on the canyon timescale. The concentration contributed to the chemical production for O$_x$ was about 67% of that for NO$_2$. This was partially attributed to the chemical consumption of O$_3$ in the canyon. The chemical production of O$_x$ was due to VOC
oxidation processes through the chemistry associated with fast radicals (e.g. HO\(_x\)) which effectively convert NO to NO\(_2\). It was observed that \(C_{\text{chem}}\) for OH and HO\(_2\) were negative, i.e. chemically consumed in the street canyon environment. The spatial patterns for these \(C_{\text{chem}}\) were dependent upon the vortex structure inside the street canyon. There were also sharp gradients at the canyon roof level and a clear separation at the interface between the lower and upper vortices. It is interesting to note that the greatest magnitudes of either positive or negative values were observed close to the windward wall in the lower vortex. These may be explained by the trapped emissions due to the anti-clockwise vortex in the lower canyon and the relatively longer retention time than that in a much shallower canyon with an AR=1. These magnitudes for the upper canyon were slightly lower than those for the lower canyon. It was also noted that in the background atmosphere above the canyon, there was no chemical production or consumption (the values of \(C_{\text{chem}}\) being close to zero). This was due to that the background atmosphere was already in the quasi-equilibrium state, in the simulation approach adopted here (although not necessarily in reality, with upwind heterogeneity). The turbulent mixing timescale (\(T_{\text{mix}}\)) can be estimated as the length scale of the canyon divided by its velocity scale. The length scale of the canyon is \(H\) (=36 m) and the velocity scale of turbulent mixing is estimated as 0.058 m s\(^{-1}\), i.e. the square root of the mean resolved-scale turbulent kinetic energy (Salizzoni et al., 2009). The turbulent mixing timescale was calculated as 621 s. The chemical timescales (\(T_{\text{chem}}\)) within the canyon (calculated based on Equation (5) using the canyon averaged quantities over the last 60 min period) was estimated as 185 s for NO, 108 s for NO\(_2\), 9 s for O\(_3\), 0.0043 s for OH and 0.014 s for HO\(_2\). The Damköhler number (Da) was calculated as 3.4 for NO, 5.8 for NO\(_2\), 69 for O\(_3\), 1.44×10\(^5\) for OH, and 4.44×10\(^4\) for HO\(_2\). This reflected that the chemical production or consumption for these species was limited by dynamical processes in the street canyon environment (Figure 4 b-h). In other words, the local chemistry dominated the relative abundance of HO\(_x\) and probably O\(_3\) (Da >> 1), but for NO and NO\(_2\), the interaction between dynamics and chemistry was of vital importance (the turbulent mixing timescale was comparable to the chemical timescale, as Da was in the same order as 1).
4.2 Pre-processing of emitted pollutants

Figure 5 shows vertical profiles of the horizontally averaged total (Equation 23), turbulent (Equation 22) and advective (Equation 21) fluxes, for (a) NO, (b) NO\textsubscript{2}, (c) O\textsubscript{3}, (d) NO\textsubscript{x}, (e) O\textsubscript{x} and (f) NO/NO\textsubscript{2}. The relative total fluxes for non-passive scalars were reconstructed based on a passive scalar with a unit emission rate (Figure 4a) and were denoted by the red solid lines. The departure of the total fluxes (black solid lines) away from the red solid lines represents chemically induced fluxes. Negative (or positive) values of fluxes mean that pollutants are entrained downward (or upward) to the street canyon. It was observed that advective fluxes were higher than turbulent fluxes for both upper and lower vortices, but lower for shear layers, which indicated that advective fluxes acted as a dominant mechanism for the transport of pollutant within a vortex while turbulent fluxes played an important role for the exchange of pollutants within the zone between the vortices.

There was also clear evidence that both advective fluxes and turbulent fluxes change rapidly close to the canyon roof level and the level where two vortices formed in the deep street canyon interacted. This sensitivity to the vertical height at the canyon roof level was also found by Cheng and Liu (2011a), in which LES simulations of a passive scalar in the street canyon with AR=1 were conducted. A positive (upward) total flux was observed for emitted species (e.g. NO and NO\textsubscript{2}) while a negative (downward) total flux was observed for entrained species (e.g. O\textsubscript{3}). Total fluxes for emitted species (e.g. NO and NO\textsubscript{2}) increased rapidly from the ground to the level at z/W=0.1 (where the centre of the elevated emissions is located), and then decreased with height in the canyon for NO, but increased with height for NO\textsubscript{2}. This reflects the chemical conversion of NO to NO\textsubscript{2} through within-canyon pre-processing of emissions. The NO/NO\textsubscript{2} ratio of total fluxes was about 1.7 (vs 4 in Zhong et al., 2015) at the canyon roof level, smaller than the NO/NO\textsubscript{2} ratio of the emission fluxes (assumed as 9:1). Therefore, the within-canyon processing resulted in increased levels of NO\textsubscript{2} through the chemical conversion of NO to NO\textsubscript{2} and changed the partitioning of total NO\textsubscript{x} emissions at the canyon roof level. This indicated that apart from the emitted NO\textsubscript{2}, the chemical processing within the canyon had a substantial contribution to the high level of NO\textsubscript{2}
released to the overlying canopy layer. For NO\textsubscript{x}, the total flux remained almost constant with height (about 5 ppb m s\textsuperscript{-1}) except a rapid increase near the ground level, which was ascribed to the simulation approach here, in which the near-vehicle dispersion was assumed to exhibit a Gaussian distribution. For O\textsubscript{x}, the total flux increased with an increase in the vertical height up to about 1.4 ppb m s\textsuperscript{-1} at the canyon roof level, which was about 2.8 times its raw emission flux (about 0.5 ppb m s\textsuperscript{-1}). This ratio was higher than that found by Bright et al. (2013) for the AR=1 case (about 1.3 times). This was due to the HO\textsubscript{x} chemistry, which converted NO to NO\textsubscript{2} and thereby resulting in an increase in O\textsubscript{x}. Without the HO\textsubscript{x} chemistry, O\textsubscript{x} flux would be nearly constant in the canyon environment because titration does not change the abundance of O\textsubscript{x}. It was the longer retention time of pollutants in the deep street canyon (AR=2) that allowed the accumulation of O\textsubscript{x} generated from the HO\textsubscript{x} chemistry. This was very different for the case with an AR=1 (Bright et al., 2013) with a much shorter retention time (residence time) for pollutants. The fluxes increased further with the vertical height for the deep street canyon (AR=2) in this study (with the canyon roof fluxes around 2 times compared to those for the canyon with an AR=1). These findings suggested that the within-canyon pre-processing results in an increase in the oxidant flux and this effect is more significant for the deeper street canyon.

4.3 Segregation effects

Table 1 lists intensities of segregation (in percentage) between selected pairs of chemical species for the street canyon. It is interesting to note that \( I_{S(A=B)} \) for A=B are positive, with the largest value of 28.49 % for \( I_{S(NO,NO)} \), and the smallest value of 0.36 % for \( I_{S(OH,OH)} \). This can be explained by the fact that the auto-covariance of any chemical species was always positive if the chemical species was not homogenously distributed within the canyon. \( I_{S(A=B)} \) (where A=B) reflected the spatial variability of the chemical species within the canyon relative to its mean concentration.

It was found that there were positive values for intensities of segregation between NO, NO\textsubscript{2} and VOCs, indicating that ‘emitted chemical species’ have similar correlations and are driven by the
dynamical processes acting upon emissions. The highest value was found to be 22.32 % for $I_{S(NO+VOCs)}$. These emitted chemical species were carried by the canyon vortices and removed from the canyon roof level to the background atmosphere. Positive values of intensities of segregation between O$_3$, OH and HO$_2$ were also clearly observed, but these magnitudes were lower (below 3%) . This can be explained by considering that O$_3$, OH and HO$_2$ are ‘entrained chemical species’ with higher levels in the background environment than those inside the street canyon and thereby exhibiting similar behaviour. This implied that segregation effect would enhance the rate of a reaction between pairs of species with similar origins.

It was also noted that negative values are found for intensities of segregation between emitted and entrained chemical species. This was attributed to the opposite origin of those chemical species. Negative correlations between those species were therefore expected. As shown in Table 1, these pairs of both emitted and entrained chemical species generally undergo the chemical reactions within the canyon. The average chemical reaction rates across the canyon domain were expected to be reduced due to the incomplete mixing in such an environment. Segregation effects were relatively larger between O$_3$ and emitted species than those between OH (or HO$_2$) and emitted species. It is noted that intensity of segregations are -11.09 % between NO and O$_3$ (featured by O$_3$ titration) and -2.37 % between VOCs and OH (linked by fast radical reactions), which were about twice those calculated for a regular street canyon with an AR= 1 in Bright et al. (2013). It is not surprized that the segregation effect for a deep street canyon with two primary vortices is more significant than that for a regular street canyon with only one primary vortex (Bright et al., 2013). These findings showed that the NO and O$_3$ titration to generate NO$_2$ within the street canyon was reduced by 11.09 % and the conversion rate of NO to NO$_2$ by the VOCs oxidation chemistry via the OH/HO$_2$ chemistry was reduced by 2.37 % due to segregation effects, compared with a wellmixed system. In other words, if the whole street canyon was treated by a single well-mixed box, effective chemical reaction rates (Equation (25-27)) should be adopted, e.g. 11.09% smaller for NO and O$_3$ and 2.37 % smaller for VOCs and OH, than the standard kinetic values for a well-mixed
system (or for the individual reactions in isolation). Such effective chemical reaction rates would compensate for the immediate and total mixing assumption inherent in the single box approach - but the change required would vary for each scenario (emissions, background chemistry, aspect ratio) due to the non-linearity. Auger and Legras (2007) suggested that due to the nonlinear nature of chemical processes, even a small value for intensity of segregation (e.g. 1 %) may lead to substantial effects on the mean concentrations, especially where the pollutant residence time is short.

4.4 A coupled two-box model approximation

Zhong et al. (2015) developed a simplified two-box model coupled with simple NO$_x$-O$_3$ chemistry to capture the concentration contrast between the lower canyon (box) and the upper canyon (box), reflecting the potential segregation effect caused by the two counter-rotating vortices. By using a plane at the level of $z/H = \alpha$ (where $\alpha$ is the box height ratio determined by the flow structure with the street canyon; a value of 0.25 was approximated based on their LES results), the two-box model framework for the deep street canyon was developed (Figure S2 in the Supplementary Materials). The mathematical description of the two-box model is as follows:

$$\frac{dC_{i,L}}{dt} = -\frac{w_{i,L}}{H_L} (C_{i,L} - C_{i,U}) + E_{i,L} + \Delta S_{i,L}$$  \hspace{1cm} (28)$$

$$\frac{dC_{i,U}}{dt} = \frac{w_{i,L}}{H_U} (C_{i,L} - C_{i,U}) - \frac{w_{i,U}}{H_U} (C_{i,U} - C_{i,b}) + \Delta S_{i,U}$$  \hspace{1cm} (29)$$

where $C_i$ (ppb) represents the concentration of $i^{th}$ species; $t$ (s) is the time; $H$ (m) represents the height; $w_i$ (m s$^{-1}$) represents the exchange velocity; $E_i$ (ppb s$^{-1}$) represent the emission rate of $i^{th}$ species; $\Delta S_i$ represents the chemical source term of $i^{th}$ species; “L” represent the properties for the lower box while “U” represent the properties for the upper box. The exchange velocities employed in the two-box model are 0.018 m s$^{-1}$ for $w_{i,L}$ and 0.014 m s$^{-1}$ for $w_{i,U}$, which are derived based on the LES model for the exchange of a passive scalar in Zhong et al. (2015). In this study, these
chemical sources terms in Equations (28) and (29) are derived from the RCS O$_3$-NO$_x$-VOC chemistry rather than simple NO$_x$-O$_3$ chemistry in Zhong et al. (2015).

4.4.1 Time evolution of volume-averaged concentrations

Figure 6 shows time evolution of volume-averaged concentrations of NO, NO$_2$, O$_3$, NO$_x$, O$_x$, OH and HO$_2$ for the lower and upper canyon (box) calculated by the LES-chemistry model and the two-box model, respectively. In Figure 6, it is interesting that there were fluctuations for the LES approach due to its inherently unsteady (dynamical) nature. It was observed that there were rapid changes in mixing ratios when the emissions were released into the street canyon from 30 min.

Compared with the LES-chemistry model over the period of 180-240 min (volume- and time-averaged mixing ratios listed in Table 2), the two-box model underestimated NO levels by about 5.25 % and 5.8 % for, but overestimated NO$_2$ levels by about 8.47 % and 5.94 % for the lower and upper boxes respectively. Levels of O$_3$ derived from the two-box model were about 1.97 % and 1.83 % lower than those derived from the LES-chemistry model for the lower and upper boxes respectively. These differences were small, suggesting that the two-box approach performs pretty well compared with the “more realistic” LES-chemistry model. These results also showed that segregation effects caused by incomplete mixing (i.e. spatial inhomogeneity represented by the LES-chemistry model) reduced the conversion rate of NO to NO$_2$ through chemistry (dominated by NO and O$_3$ titration with an additional pathway through VOCs chemistry), which was consistent with negative values of intensities of segregation between NO and O$_3$, and between OH and VOCs (Table 1). It was also observed that NO$_2$/NO ratios in the two-box model were generally higher than those in the LES-chemistry model, i.e. about 14.47 % for the lower box and about 12.50 % for the upper box. Therefore, there were higher levels of O$_3$ and NO, but lower levels of NO$_2$ in the LES-chemistry model than those in the two-box model for both lower and upper boxes. The LES-chemistry model had slightly higher levels of NO$_x$ (about 1.59 % for the lower box and 1.69 % for the upper box) compared with the two-box model, which suggests that segregation effects slightly reduced the NO$_x$ loss rate to other species (e.g. HNO$_3$ and HONO). This was also consistent with
negative values of intensities of segregation between OH and NO$_2$, and between OH and NO (Table 1). Lower levels of O$_x$ were observed in the LES-chemistry model compared with the two-box model, i.e. about 7.89 % for the lower box and 5.15 % for the upper box. This indicated that segregation effects generally reduced the rate of oxidation chemistry for both the lower and upper boxes. It was also observed that the two-box model slightly underestimated levels of both OH and HO$_2$ (generally around 1%) compared with the LES-chemistry model. This may be explained as levels of OH and HO$_2$ were low within street canyons and their reactions with other chemical species were very fast. Segregation effects can reduce the rate for some of these chemical reactions, but increased the rate for other chemical reactions (Table 1). The total segregation effect may be somewhat balanced out - depending upon the metric under consideration, i.e. which species is concerned. In terms of overall performance, the two-box model generally matched the LES approach in the volume averaged concentrations of pollutants for both the lower and upper boxes.

5 Conclusions

An LES model coupled with O$_3$-NO$_x$-VOC chemistry (the RCS mechanism) was implemented to simulate the coupling effect of emissions, mixing and chemical pre-processing within a deep urban street canyon (AR=2). There were significant spatial variations of reactive pollutants in the presence of two vertically aligned unsteady vortices formed in the canyon. Compared with the RCS mechanism, simple NO$_x$-O$_3$ chemistry overestimated NO level, but underestimated levels of NO$_2$ and O$_3$, indicating the additional conversion of NO to NO$_2$ through OH/HO$_2$ chemistry. NO, NO$_x$, O$_3$, OH and HO$_2$ were chemically consumed, while NO$_2$ and O$_x$ (oxidants) were chemically produced within the canyon environment. The within-canyon pre-processing would lead to chemical conversion of NO to NO$_2$ and an increase in the oxidant fluxes released from the canyon to the overlying canopy layer, and this effect was more significant for the deeper street canyon than the regular canyon (AR=1) (Bright et al., 2013). Consequently, urban canopy layer air quality in cities will vary systematically with the street architecture (aspect ratio), with greater release of NO$_2$
and O\textsubscript{3}, for the same traffic emissions, where street canyons are taller - for example in many traditional European city centres, in contrast to more open suburban regions. Such findings can be of importance in guiding the development of atmospheric pollutant flux parameterisation schemes for larger scale (e.g. city or regional scale) models, and for urban planning considerations. There was clear evidence of two distinctive behaviours for emitted chemical species and entrained chemical species. Positive (or negative) values of intensities of segregation were found between pairs of species with a similar (or opposite) behaviour. Such findings indicated that segregation effects were of importance in the incomplete mixing environment (e.g. the street canyon) with chemical processing involved. The simplified two-box model underestimated NO and O\textsubscript{3} levels, but overestimated NO\textsubscript{2} levels for both the lower and upper boxes compared with the LES-chemistry model. NO\textsubscript{2}/NO ratios in the two-box model were found to be much higher than those in the LES-chemistry model. Segregation effects due to incomplete mixing may reduce the conversion rate of NO to NO\textsubscript{2} through chemistry.

**6 Implications and future research**

The LES-chemistry model in this study reveals the impacts of nonlinear O\textsubscript{3}-NO\textsubscript{x}-VOC photochemical processes in the incomplete mixing environment (e.g. street canyons) and provides a better understanding of the pre-processing of emissions in the presence of both the street canyon dynamics and chemistry. This research may guide the location of new urban air quality stations, to ensure these are representative of human exposure and/or understand the measurement bias that may accrue from a particular location within a canyon. Because of the high computational cost, this study was limited to one typical emission scenario, with relatively poor air ventilation under neutral conditions. Also, due to the simple assumption of the idealised street canyon geometry under perpendicular ambient wind, flow field within the canyon was dominated by flow recirculation (i.e. two vortices). However, the current LES model of idealised scenarios does not capture lateral channelling flow (e.g. Longley et al., 2004) or even helical flow (e.g. Dobre et al., 2005; Barlow et
al., 2009) present in real, complex urban street canyons (Smalley et al., 2008). Future studies may consider the effects of other factors, such as complex urban configurations, wind speed, oblique wind directions, emissions and thermal/shading effects (e.g. Li et al., 2015; Li et al., 2016; Cheng and Liu, 2011b), on both the dynamic and chemical processing of reactive pollutants.

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### Table 1
Intensities of segregation (in percentage) between pairs of chemical species for the street canyon. Values shown in parentheses and bold denote those pairs of chemical species that react directly with each other in the RCS \( \text{O}_3\)-\( \text{NO}_x\)-\( \text{VOC} \) chemistry. Positive (negative) values mean that these pairs of chemical species have similar (opposite) behaviours.

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<th>( \text{O}_3 )</th>
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<th>( \text{NO}_2 )</th>
<th>( \text{VOCs} )</th>
<th>( \text{HO}_2 )</th>
<th>OH</th>
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<td>-</td>
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### Table 2
Volume- and time-averaged (over the period of 180-240 min) mixing ratios in the lower and upper boxes derived from the LES-chemistry model (LES-RCS) and the two-box model (BOX-RCS), respectively. Positive (negative) values represent the amount of overestimation (underestimation) by BOX-RCS compared with LES-RCS.

<table>
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<th>Mixing ratio (ppb) for Upper Box</th>
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<td>(B) Box-RCS</td>
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<td>( \text{OH} )(ppt)</td>
<td>0.103619</td>
<td>0.1023</td>
</tr>
<tr>
<td>( \text{HO}_2 )(ppt)</td>
<td>0.265364</td>
<td>0.2640</td>
</tr>
<tr>
<td>( \text{NO}_x )</td>
<td>630.6373</td>
<td>620.5910</td>
</tr>
<tr>
<td>( \text{O}_x )</td>
<td>177.9566</td>
<td>191.9999</td>
</tr>
<tr>
<td>( \text{HO}_x )</td>
<td>0.3690</td>
<td>0.3664</td>
</tr>
<tr>
<td>( \text{NO}_2/\text{NO} )</td>
<td>0.3636</td>
<td>0.4163</td>
</tr>
</tbody>
</table>
Figure 1 Spatial variation of (a) $\left< NO \right>$ (ppb), (b) $\left< NO_2 \right>$ (ppb), (c) $\left< O_3 \right>$ (ppb), (d) $\left< NO_3 \right>$ (ppb), (e) $\left< O_3 \right>$ (ppb), and (f) $\frac{\left< NO \right>}{\left< NO_2 \right>}$, (g) $\left< OH \right>$ (ppt) and (h) $\left< HO_2 \right>$ (ppt). Logarithmic colour scales are applied for $\left< OH \right>$ and $\left< HO_2 \right>$.
Figure 2 Spatial variations of the overestimation of the spatially and temporally averaged concentrations (%) by simple NO$_x$-O$_3$ chemistry compared with the RCS chemical mechanism for (a) $\langle NO \rangle$, (b) $\langle NO_x \rangle$, (c) $\langle O_3 \rangle$, (d) $\langle NO \rangle$, (e) $\langle O_3 \rangle$ and (f) $\langle NO \rangle/\langle NO_x \rangle$. 
Figure 3 Vertical profiles of (a) $\langle NO \rangle$, (b) $\langle NO_2 \rangle$, (c) $\langle NO_x \rangle$, (d) $\langle O_x \rangle$, (e) $\langle NO \rangle/\langle NO_2 \rangle$, and (f) $\langle C \rangle/C_0$ (for O$_3$, OH and HO$_2$ normalised by their background levels) along the leeward and windward walls, represented by the dash and solid lines respectively.
(a) $C_{\text{chem},1}$

(b) $C_{\text{chem}}$ for NO

(c) $C_{\text{chem}}$ for NO$_2$

(d) $C_{\text{chem}}$ for NO$_x$

(e) $C_{\text{chem}}$ for O$_x$

(f) $C_{\text{chem}}$ for O$_3$

(g) $C_{\text{chem}}$ for OH

(h) $C_{\text{chem}}$ for HO$_2$
Figure 4 Spatial variation of (a) $C_{\text{em}}$ (ppb) and $C_{\text{chem}}$ of (b) NO (ppb), (c) NO$_2$ (ppb), (d) NO$_x$ (ppb), (e) O$_3$ (ppb), (f) O$_3$ (ppb), (g) OH (ppt) and (h) HO$_2$ (ppt). Logarithmic colour scales are applied for OH and HO$_2$. Note very different colour scales for different species.

Figure 5 Vertical profiles of the horizontally averaged total, turbulent and advective fluxes for (a) NO, (b) NO$_2$, (c) O$_3$, (d) NO$_x$, (e) O$_3$ and (f) NO/NO$_2$. The total, turbulent and advective fluxes for each quantity are represented by the black solid, dash and dotted lines, respectively. The total fluxes for non-passive scalars assuming no chemical reactions (i.e. reconstructed based on a passive scalar with a unit emission rate) are denoted by red solid lines.
Figure 6 Time evolution of the volume averaged concentrations of (a) NO, (b) NO₂, (c) O₃, (d) NOₓ and Oₓ, (e) OH and (f) HO₂ derived from the LES-chemistry model (LES-RCS) and the two-box model (BOX-RCS), respectively. ‘L’ represents the lower box while ‘U’ represents the upper box.
References:


