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Modelling chemistry and transport in urban street canyons: Comparing offline multi-box models with large-eddy simulation

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Highlight

- NO_x-O₃-VOC chemical reactions are coupled with multi-box canyon models.
- The multi-box models reproduce flow characteristics in regular and deep canyons.
- Reactive species concentrations in street canyons are well captured in <1% of the run time of computational fluid dynamics.
- The impacts of segregation on reactive species can be investigated.

Abstract

Computational fluid dynamics models are resource-intensive, particularly when complex chemical schemes are implemented, and this computational expense limits their use in sensitivity analyses. We propose a flexible multi-box model that permits spatial disaggregation of sources and depositions to simulate the transportation and distribution of chemical species in street canyons with any aspect ratios for which a large eddy simulation (LES) of the flow exists. The spatial patterns of reactive species in the multi-box simulations are in good agreement with those from the LES, especially for the deep canyon from which air escapes more slowly. The overestimation of the LES simulation

31 worsens somewhat due to segregations when the chemistry of volatile organic compounds (VOCs) is
32 included but the overall pattern is captured in a modelling framework. By reducing computational
33 costs by several orders of magnitude, the multi-box model allows more sensitivity testing than the
34 LES, and is an effective approach to investigate spatial pattern of fast non-linear chemistry or
35 microphysics at the street scale.

36 *Keywords:* Air quality; Urban air pollution; Box models; Street canyon; Nitrogen dioxide; Ozone

37 **1. Introduction**

38 Street canyons typically combine to build up a semi-enclosed urban environment with high
39 concentrations of anthropogenic pollutants trapped inside, leading to persistent higher exposure risk
40 for pedestrians near the roadside (Ahmad et al., 2005; Oke, 1988; Vardoulakis et al., 2003). Canyons
41 can be divided into three types in terms of the ratio of building height (H) to street width (W), namely
42 aspect ratio (AR): wide canyons ($AR < 0.3$), regular canyons ($AR \approx 1$), and deep canyons ($AR > 1.3$)
43 (Vardoulakis et al., 2003). Oke (1988) classified the canyon flows into three isothermal regimes
44 according to AR and L/W (L is the building length along the span-wise direction). These regimes
45 include skimming flow ($0.66 < AR < 1.57$), wake interference flow ($0.1 < AR < 0.66$) and isolated
46 roughness flow ($AR < 0.1$). Besides the isolated roughness flow, wake interference flow, and
47 skimming flow, there is the fourth flow regimes, i.e., multi-vortex flow regime in deep street canyons.
48 If Reynolds (Re) number independence is satisfied, there is only one-vortex when AR is 1 and 3, but
49 two main vortexes appear when AR is 5 or more (Yang et al., 2020). However, in wind-tunnel-scale
50 street canyons ($H = 6$ cm, $Re \sim 1.2 \times 10^4$), if Re number is not sufficiently large, there are two contra-
51 rotative vortexes when $AR = 2$ and three to five vertically aligned vortexes when AR is 3-5 (Li et al.,
52 2008). The main difference between these two groups is whether the requirement of Re-number
53 independence is satisfied or not. Chew et al. (2018) proposed that the widely adopted criterion $Re >$
54 11,000 for ensuring Reynolds-number-independence (Re-independence) is not applicable for 2D
55 street canyons as $AR > 1.5$. They discovered that only one primary vortex appeared when $AR = 2$ if
56 $Re > 8.7 \times 10^4$. Moreover, Yang et al. (2021) found that isothermal urban airflows for full-scale deep
57 canyons can be independent of Re when Re exceeds 1×10^6 and 1×10^7 when AR is 3 and 5.

58 Because of much of the human exposure to outdoor air pollutants occurs at the pedestrian level in
59 street canyons, understanding airflow characteristics and distributions of pollutants is of vital

60 importance in evaluating the pollutant health risk and in making policy for targeted air pollution
61 alleviation. The concentration of a passive scalar (PS, an idealised chemically inert substance that
62 negligibly interferes with local fluid dynamics through effects such as buoyancy) can exhibit sharp
63 gradients at the pedestrian level of the street canyons (Fellini et al., 2020; Lietzke and Vogt, 2013;
64 Murena et al., 2009). Furthermore, the dispersion of atmospheric pollutants within the canyon is
65 accompanied by complex non-linear chemical reactions, evolving on the timescale comparable to the
66 canyon circulation and residence timescale. During the past two decades, studies have focused on the
67 investigation of time-evolution and spatial variations of reactive species, for example, nitric oxide
68 (NO), nitrogen dioxide (NO₂), and ozone (O₃) in street canyons, using coupled computational fluid
69 dynamics (CFD) such as Reynolds-Averaged Navier-Stokes (RANS) and Large-Eddy Simulation
70 (LES) models with different chemical schemes. Baker et al. (2004) integrated an LES model with a
71 simple NO_x-O₃ cycle (two reactions in sunlight) to simulate the dispersion and spatial distribution of
72 reactive species in a regular street canyon. The impacts of dynamics on the chemistry had been
73 investigated by introducing the concept of the photo-stationary state defect (PSSD). Low PSSD values
74 indicate equilibrium-like chemistry and were found at the centre of the primary vortex and on the
75 windward corner at the street level. High PSSD values indicate rapidly-changing chemistry and were
76 found at the location near NO_x sources, along the leeward facet (where pollutants were escaping the
77 street canyon), and along the windward wall on the outer edge of the vortex (where fresh air was
78 entrained into the canyon). Baik et al. (2007) adopted the renormalised k - ϵ RANS model coupled with
79 the simple NO_x-O₃ photochemistry to simulate pollutant dispersion with the effects of street-bottom
80 heating; Kwak and Baik (2012) further incorporated Volatile organic compounds (VOC) chemistry
81 into the model and discussed the sensitivity of O₃ concentrations to NO_x and VOC emissions. Kim et
82 al. (2012) examined reactive species in a regular canyon using RANS with a comprehensive
83 tropospheric NO_x-O₃-VOC chemistry from GEOS-Chem. They found a substantial influence of
84 different chemical schemes on O₃ levels and highlight the importance of more explicit chemistry
85 simulation. However, Bright et al. (2013) found that VOCs could contribute additional but modest
86 NO₂ and O₃ formation (about 12%) in the regular canyon, which is consistent with Garmory et al.
87 (2009). Zhong et al. (2017) extended the LES model with the NO_x-O₃-VOC chemistry to simulate the
88 spatial distribution of reactive species in an idealised deep urban street canyon. They revealed that
89 volume-averaged NO₂ and the total oxidants (O_x) concentrations significantly increased due to VOCs-
90 relating chemistry. Their works provided a better understanding of the combined effects of insufficient

91 mixing and non-linear reactions in street canyons with a higher aspect ratio. Zhang et al. (2020)
92 conducted RANS simulations with a simple $\text{NO}_x\text{-O}_3$ cycle in street canyons with $\text{AR} = 1, 3$ and 5 , but
93 their work neglected the effect of organic free radicals, which is likely important in determining the
94 concentration of reactive species inside deep street canyons. Recently, Wu et al. (2021) developed a
95 platform that integrated a CFD model (i.e., OpenFOAM) and a photochemical mechanism including
96 VOCs for pollutant dispersion in the regular canyon. Although these CFD-based studies provide
97 promising methods in order to simulate physical dispersion and chemical transformation of reactive
98 species within canyons, modelling scenarios in reality are much more sophisticated in terms of, for
99 example, emissions (Wu et al., 2021), wind conditions, roof shapes (Takano and Moonen, 2013),
100 ground or wall heating (Cai, 2012), and the presence of green infrastructures such as trees with
101 varying leaf area density (Abhijith et al., 2017; Gromke et al., 2008; Gromke and Ruck, 2007). The
102 domain/model configurations need to be adjusted from case to case, and using a CFD model is, in
103 general, very computationally expensive for the study of such a wide range of scenarios, especially
104 when explicit VOC chemical reactions are included.

105 A more convenient and efficient way is adopting offline simulations, as is routinely employed at
106 regional and global scales (e.g., Jacobson and Jacobson (2005); Kukkonen et al. (2012)). The zero-
107 dimensional one-box model can easily adopt complex chemistry without intensive computational
108 resources. It also exhibited satisfactory performance compared to the LES (Bright et al., 2013),
109 despite relatively higher modelled NO , NO_2 and hydroxyl radical (OH) concentrations. Zhong et al.
110 (2016) implemented a coupled two-box model with $\text{NO}_x\text{-O}_3\text{-VOC}$ chemistry for simulating pollutants
111 in a deep canyon. They highlight that the one-box treatment would miscalculate flow structure and
112 pollutant gradients, and, hence, underestimate the exposure risk of pedestrians to NO_2 in the deep
113 canyon. However, these simplified box models are still too coarse to evaluate air quality conditions
114 at the pedestrian level. They would systematically neglect substantial concentration contrasts near the
115 centre of the carriageway and cannot capture the horizontal distribution of pollutants, which is as
116 important as vertical features in street canyons. The lack of available process-based methods makes
117 it difficult to investigate systematically coupled chemistry-transport effects in street canyons.

118 In this study, a multi-box model with a flexible number of boxes and flexible chemical schemes has
119 been developed for air pollution simulations in street canyons. The model design, mathematical
120 formulation, and configurations for testing are described in Section 2. In Section 3, the modelling

121 results of reactive species from the multi-box models with a reduced NO_x-O₃-VOC chemistry and a
 122 simple NO_x-O₃ cycle are evaluated against the published modelling data from the LES dynamical
 123 models at the box grid resolution, for idealised regular and deep street canyons. The time-evolution
 124 of concentrations, segregation effects due to overly fast chemistry in the multi-box case, and spatial
 125 variations inside canyons are discussed in detail. The conclusions and future perspective are
 126 summarised in Section 4. Although not the focus of the current paper, we note that aerosol
 127 microphysics introduces non-linear processes in street canyons in a similar way to chemistry (Gelbard
 128 and Seinfeld, 1980; Jacobson and Seinfeld, 2004; Jacobson et al., 1996; Zhong et al., 2020a, b; Zhong
 129 et al., 2018); the modelling framework described below could be extended to include detailed size-
 130 dependent aerosol microphysics in the future.

131 2. Methods

132 2.1. Description of models

133 2.1.1 The multi-box model

134 The principle of the multi-box model is to split the volume of street canyons into several boxes, where
 135 each box ideally reflects a resolved airflow arising from the aspect ratio or physical obstructions in
 136 street canyons (e.g., Fig. 1). In the two-dimensional framework, boxes inside the model are indexed
 137 based on their locations in Cartesian coordinates. Assuming the background wind above the roof level
 138 blows across the street canyon from left to right, starting from the bottom-left to the top-right, e.g.,
 139 Box_[1,1] represents the leeward corner of a canyon. Pollutant transfer between adjacent boxes is
 140 determined by the mean wind advection and by turbulent diffusion across the mesh interface. Vertical
 141 (denoted by capital “*G*”) and horizontal (denoted by capital “*F*”) mixing-ratio fluxes (ppb m s⁻¹) for
 142 pollutant, *q*, into Box_[k,i] (i.e., “*k*” represents vertical index position, “*i*” represents horizontal index
 143 position) can be formulated as:

$$144 F_{e,[k,i]} = u_{e,[k,i]} (C_{q,[k,i-1]} - C_{q,[k,i]}) \quad (1)$$

$$145 F_{a,[k,i]} = \begin{cases} U_{a,[k,i]} C_{q,[k,i-1]}, U_{a,[k,i]} \geq 0 \\ U_{a,[k,i]} C_{q,[k,i]}, U_{a,[k,i]} < 0 \end{cases} \quad (2)$$

$$146 \quad G_{e,[k,i]} = w_{e,[k,i]} (C_{q,[k-1,i]} - C_{q,[k,i]}) \quad (3)$$

$$147 \quad G_{a,[k,i]} = \begin{cases} W_{a,[k,i]} C_{q,[k-1,i]}, W_{a,[k,i]} \geq 0 \\ W_{a,[k,i]} C_{q,[k,i]}, W_{a,[k,i]} < 0 \end{cases} \quad (4)$$

148 where $G_{a,[k,i]}$ (ppb m s⁻¹) and $F_{a,[k,i]}$ (ppb m s⁻¹) are mixing-ratio fractional fluxes due to advective
 149 transfer (i.e., flow resolved by the LES); $G_{e,[k,i]}$ (ppb m s⁻¹) and $F_{e,[k,i]}$ (ppb m s⁻¹) are mixing-ratio
 150 fluxes due to turbulent diffusion formulated by the Fick's law. $W_{a,[k+1,i]}$ (m s⁻¹) and $U_{a,[k+1,i]}$ (m s⁻¹) are
 151 the advective transfer velocities in the vertical and horizontal directions, respectively; and $w_{e,[k+1,i]}$ (m
 152 s⁻¹) and $u_{e,[k+1,i]}$ (m s⁻¹) are transfer velocities due to turbulent diffusion. By assuming all fluxes as
 153 vectors with positive values along the coordinate directions, the concentration in Box_[k,i] can be
 154 calculated from the following equation:

$$155 \quad \frac{dC_{q,[k,i]}}{dt} = E_{q,[k,i]} - \frac{G_{a,[k+1,i]} + G_{e,[k+1,i]}}{l_i} + \frac{G_{a,[k,i]} + G_{e,[k,i]}}{l_i} - \frac{F_{a,[k+1,i]} + F_{e,[k+1,i]}}{h_k} + \frac{F_{a,[k,i]} + F_{e,[k,i]}}{h_k} + \Delta S_{q,[k,i]} + \Delta V_{q,[k,i]} \quad (5)$$

156 where $C_{q,[k,i]}$ (ppb) is the mixing-ratio of the q^{th} species in Box_[k,i], $E_{q,[k,i]}$ (ppb s⁻¹) is the emission rate
 157 of the q^{th} species into Box_[k,i], h_k (m) and l_i (m) are the box height and box width respectively, $\Delta S_{q,[k,i]}$
 158 (ppb s⁻¹) is the net production rate of the q^{th} species due to chemistry in Box_[k,i], and $\Delta V_{q,[k,i]}$ (ppb s⁻¹)
 159 is the net deposition term of the q^{th} species in Box_[k,i]. Allowing computations with more species, and
 160 where the associated $\Delta S_{q,[k,i]}$ terms reflect more complicated non-linear chemistry, is one of the prime
 161 motivations for the development of the box model. By default, the boundary layer above the street
 162 canyon is assumed as one compartment, representing relatively steady background conditions over a
 163 long period (e.g., 1 hour). The 4th order Runge-Kutta method is adopted in the multi-box model to
 164 solve the ordinary differential equations (ODEs) numerically. To facilitate a systematic investigation,
 165 we use two dimensionless ratios to represent the size of grid boxes to the entire street canyon:

$$166 \quad \alpha_k = \frac{h_k}{h_0} \quad (6)$$

$$167 \quad \beta_i = \frac{l_i}{l_0} \quad (7)$$

168 where h_0 (m) and l_0 (m) are the canyon height and street width, respectively. Then the volume-
 169 averaged concentrations of q^{th} species ($C_{q,m \times n\text{-box}}$) for the entire canyon with $m \times n$ boxes is:

$$170 \quad C_{q,m \times n\text{-box}} = \sum_{k=1; i=1}^{k=m; i=n} \alpha_k \beta_i C_{q,[k,i]} \quad (8)$$

171 If the volume is equal for all the boxes in street canyons, that is, i.e., $\alpha_1 = \alpha_2 = \dots = \alpha_k = \frac{1}{m}$ and

172 $\beta_1 = \beta_2 = \dots = \beta_i = \frac{1}{n}$, then the equation (8) can be rewritten to:

$$173 \quad C_{q,m \times n\text{-box}} = \frac{\sum_{k=1; i=1}^{k=m; i=n} C_{q,[k,i]}}{m \times n} \quad (9)$$

174 Additionally, a dimensionless factor $\gamma_{q,[k,i]}$ is adopted to account for the heterogeneous on-road
 175 emission of the q^{th} species:

$$176 \quad \gamma_{q,[k,i]} = \frac{\alpha_k \beta_i E_{q,[k,i]}}{\sum_{k=1; i=1}^{k=m; i=n} \alpha_k \beta_i E_{q,[k,i]}} \quad (10)$$

177 Up to n continuous line sources could be added into the k^{th} ($k = 1, 2, \dots, m$) layer of the canyon, which
 178 is helpful for elevated road or rail sources and for biogenic emissions from street trees. $\gamma_{q,[k,i]} = 0$ or
 179 $\gamma_{q,[k,i]} = 1$ indicates that no emission or all vehicle emissions have been injected into $\text{Box}_{[k,i]}$.

180 In order to derive net chemical terms especially for short-lived reactive species such as hydroxyl
 181 radical (OH) and hydroperoxyl radical (HO_2), the ODEs of a chemical system for q^{th} species can be
 182 written as:

$$183 \quad \frac{d}{dt} C_{q,[k,i]} = P_{q,[k,i]} - L_{q,[k,i]} C_{q,[k,i]} \quad (11)$$

184 where $P_{q,[k,i]}$ and $L_{q,[k,i]}$ are the chemical production and loss rates in the specific $\text{Box}_{[k,i]}$. If those

185 chemical kinetics remain constant during a given timestep, Δt , equation (11) may be solved
 186 numerically with the quasi-steady-state approximation (QSSA):

$$187 \quad C_{q,[k,i],t_0+\Delta t} = \frac{P_{q,[k,i],t_0}}{L_{q,[k,i],t_0}} + \left(C_{q,[k,i],t_0} - \frac{P_{q,[k,i],t_0}}{L_{q,[k,i],t_0}} \right) e^{-L_{q,[k,i],t_0}\Delta t} \quad (12)$$

188 where t_0 represents the starting point of each time interval during simulations. **T**he chemical lifetime
 189 of q^{th} species, $\tau_{q,[k,i]}$, in the Box_[k,i] is:

$$190 \quad \tau_{q,[k,i]} = \frac{1}{L_{q,[k,i]}} \quad (13)$$

191 If $\tau_{q,[k,i]} < 0.1\Delta t$, chemical reactions are extremely fast compared to Δt , which means the chemical-
 192 steady-state can be adopted:

$$193 \quad C_{q,[k,i],t_0+\Delta t} = \frac{P_{q,[k,i],t_0}}{L_{q,[k,i],t_0}} \quad (14)$$

194 If $\tau_{q,[k,i]} > 100\Delta t$, chemical reactions take through much slower compared to Δt , and the forward
 195 Eulerian formula can be used:

$$196 \quad C_{q,[k,i],t_0+\Delta t} = C_{q,[k,i],t_0} + \left(P_{q,[k,i],t_0} - L_{q,[k,i],t_0} C_{q,[k,i],t_0} \right) \Delta t \quad (15)$$

197 If $0.1\Delta t < \tau_{q,[k,i]} < 100\Delta t$, then the chemical timescale has a comparable magnitude with Δt , and
 198 equation (12) is employed for the calculation. However, solving equation (12) incurs substantial
 199 computational costs in practice. Alexandrov et al. (1997) proposed an alternative way for the
 200 optimisation of the QSSA algorithm, which rationally expands the exponential term based on the
 201 Taylor expansion in the second order:

$$202 \quad e^{-L_{q,[k,i],t_0}\Delta t} \approx \frac{1}{1 + L_{q,[k,i],t_0}\Delta t + 0.5(L_{q,[k,i],t_0}\Delta t)^2} \quad (16)$$

203 and equation (12) can be reformatted as:

$$C_{q,[k,i],t_0+\Delta t} = \frac{C_{q,[k,i],t_0} + (1 + 0.5L_{q,[k,i],t_0} \Delta t) P_{q,[k,i],t_0} \Delta t}{1 + L_{q,[k,i],t_0} \Delta t + 0.5(L_{q,[k,i],t_0} \Delta t)^2} \quad (17)$$

In the one- and multi-box models, reactive species have been divided into two categories in terms of their chemical lifetime. For the regular and deep canyons, an empirical timestep value of 0.03 s was used with equation (15) for numerical integration of long-lived species (e.g., NO, NO₂, O₃), and a value of 0.003 s with the equation (17) was used for short-lived species (e.g., HO, HO₂). They are the same as those used in the LES (Bright et al., 2013; Zhong et al., 2015) based on the timescale of the turbulent eddies, which also indicates that no species are fully in steady state in urban street canyon environment.

The multi-box model is written in R version 3.6.2 (R Core Team, 2019) and Fortran 90 (using the Intel Fortran (IVF) Compiler) in the origin version 1.0, including three modules: the main program, the dynamical submodule, and a chemical submodule. Modularisation allows the model to be easily modified or updated for various research purposes, e.g., to investigate the impact of different chemical mechanisms on air quality in street canyons or to investigate in-canyon particle microphysics (cf., Nikolova et al. (2016)).

Additionally, a typical one-box model is used as a reference in this study, and the mathematical expression is (Liu and Leung, 2008):

$$\frac{dC_{q,0}}{dt} = E_{q,0} - \frac{w_{t,0}}{h_0} (C_{q,0} - C_{q,b}) + \Delta S_{q,0} + \Delta V_{q,0} \quad (18)$$

where a subscript “0” indicates that signs have the same meaning as those in the multi-box model but for a whole space of the street canyon in volume. The parameter $w_{e,0}$ (m s⁻¹) represents the “exchange velocity” between the street canyon and boundary layer above the rooftop. A two-box model is also used only for simulations in the deep street canyon, the mathematical expressions are (Murena, 2012; Zhong et al., 2016):

$$\frac{dC_{q,L}}{dt} = E_{q,L} - \frac{w_{t,L}}{h_L} (C_{q,L} - C_{q,U}) + \Delta S_{q,L} + \Delta V_{q,L} \quad (19)$$

$$\frac{dC_{q,U}}{dt} = \frac{w_{r,L}}{h_U} (C_{q,L} - C_{q,U}) - \frac{w_{r,U}}{h_U} (C_{q,U} - C_{q,b}) + \Delta S_{q,U} + \Delta V_{q,U} \quad (20)$$

where subscripts “L” and “U” indicate that signs have the same meaning as aforementioned but for the lower and upper compartments (i.e., including 16 boxes in the red box to solve the main vortexes) of the deep street canyon in volume, respectively. $w_{e,L}$ (m s^{-1}) represents the “exchange velocity” between two in-canyon compartments, and $w_{e,U}$ (m s^{-1}) represents the “exchange velocity” between the upper compartment and the overlying background.

2.1.2 Large-eddy simulation

The LES results of reactive species were taken from Bright et al. (2013) for an idealised regular canyon, and from Zhong et al. (2017) for a deep canyon. Their studies applied OpenFoam v2.1.1 (Jasak et al., 2007) to resolve turbulence at large spatial and temporal scales and to simulate incompressible airflow with a high Reynolds number ($\text{Re} \sim 1 \times 10^6$) in street canyons under the neutral atmosphere. The unresolved sub-grid scales (SGS) processes were treated using the one-equation SGS turbulence model, and the logarithmic law of the rough-wall (Schlichting and Gersten, 2016) was used for the near-wall treatment. Symmetry boundary conditions were used for the domain overlying the canyon, and cyclic boundary conditions were employed in x- and y-directions. A constant pressure gradient in the upper background was assumed to produce a flow perpendicular to the canyon axis. The simulations of the LES were conducted with only dynamics for about 5 h in order to obtain a dynamical-steady flow field, which was further adopted as the initial turbulence condition.

For the regular canyon, the resolved turbulent kinetic energy (TKE) has been evaluated against wind-tunnel experiments (Cui et al., 2004) and the scalar has been validated by Cai et al. (2008); for the deep canyon, the flow field agreed well with water-channel (Li et al., 2008) and wind tunnel (Kovar-Panskus et al., 2002) experiments. The deposition of air pollutants is not considered in the LES modelling. The method for integrating VOC chemistry in the LES modelling is similar to that of the multi-box simulations described above and has been detailed in Zhong et al. (2017). The methods for initialising and processing the LES (e.g., emission, time step) are also adopted directly for the multi-box model and are described in the following section.

254 2.2. Model configurations

255 2.2.1 Street canyon geometry

256 The LES domains adopted for regular ($AR = 1$) and deep ($AR = 2$) canyons were presented in Fig.
257 1(a) and Fig. 1(b), respectively. In the LES, mesh resolutions were 0.3 m, 1.0 m and 0.3 m in the x ,
258 y , z directions, respectively, inside street canyons. That is, there are respectively 3,600 and 7,200 cells
259 inside regular and deep canyons in the LES simulations. For the layer above the canyon in the LES
260 simulation, Δz gradually increased by a factor of 1.15 from the rooftop to the domain top ($z_0 = 18$ -90
261 m and $z_0 = 36$ -112 m for regular and deep canyons). Fig. 1(c) and Fig. 1(d) are the respective
262 counterparts in multi-box models for the cross-section canyons. The building height and street width
263 are 18 m ($h_0 = l_0$) for the regular street canyon, and are 36 and 18 m ($h_0 = 2l_0$) for the deep street
264 canyon. The red frame in Fig. 1(d) presents a two-box model with $h_L = h_U = 18$ m. One primary
265 clockwise vortex forms in the simulations of the regular canyon as the background skimming wind
266 perpendicularly blows along the x -direction across the canyon axis; however, in the deep street
267 canyon, a clockwise vortex in the upper compartment and a weak counter-clockwise vortex in the
268 lower compartment are formed. In the multi-box model, the in-canyon volumes have been divided
269 into identical 16 and 32 boxes for the regular and deep canyons, respectively (namely the 16- and 32-
270 box models). This implies that each grid has a volume of $4.5 \text{ m} \times 1 \text{ m} \times 4.5 \text{ m} = 20.25 \text{ m}^3$.

271 2.2.2 Dynamical parameters for air mass exchange

272 The exchange velocity ($w_{e,0}$) for the one-box model, and the advective and turbulent velocities for the
273 multi-box model, are of the utmost importance in determining the intensity of in-canyon mixing, and
274 transport and rates of escape of atmospheric pollutants to the overlying background. The values of
275 $w_{e,0} = 0.022 \text{ m s}^{-1}$ and $w_{e,0} = 0.012 \text{ m s}^{-1}$ are respectively adopted for the cross-section $18 \text{ m} \times 18 \text{ m}$
276 regular and $18 \text{ m} \times 36 \text{ m}$ deep canyons (as in Fig. 1) in the one-box model, corresponding to a
277 reference wind velocity of 2 m s^{-1} in the above rooftop layer under the neutral conditions (Cai, 2012a;
278 Zhong et al., 2017). The values of $w_{e,L} = 0.0229 \text{ m s}^{-1}$ and $w_{e,U} = 0.0156 \text{ m s}^{-1}$ are used for the two-box
279 simulation in the deep canyon, respectively.

280 In order to quantify the transport due to advection ($W_{a,[k,i]}$ and $U_{a,[k,i]}$, denote “advective velocities”
281 in this study) and mixing due to turbulence ($w_{e,[k,i]}$ and $u_{e,[k,i]}$, denote “turbulent velocities”) in the

282 multi-box model, firstly vertical mixing-ratio fluxes including advective fluxes and turbulent fluxes
283 of PS at the interface of boxes were extracted from the LES modelling results, and the averaged values
284 over a period of the last 60 minutes were used for calculations. The total horizontal fluxes were then
285 derived based on the flux balance of each grid on the Eulerian coordinates under an assumed
286 equilibrium state over the period. These LES-derived fluxes describe the transmission of PS between
287 any two adjacent boxes in street canyons. However, the velocities from the LES simulations
288 (representing “real” wind conditions) cannot be used directly, because they need to be adjusted to
289 calculate the mass transfer between coarser grids. Therefore, pollutant advective velocities ($U_{a,[k,i]}$ and
290 $W_{a,[k,i]}$) are obtained using the vertical advective fluxes ($F_{a,[k,i]}$ and $G_{a,[k,i]}$) and the grid concentrations
291 (i.e., an volume-averaged value at the grid resolution of the multi-box model) following Equation (2)
292 and (4), and vertical mixing-ratio turbulent velocities ($w_{e,[k,i]}$ and $u_{e,[k,i]}$) are obtained using vertical
293 turbulent fluxes ($F_{e,[k,i]}$ and $G_{e,[k,i]}$) and concentration gradients based on Equation (1) and (3). In this
294 way, the mass fluxes of the multi-box and LES models are consistent at the same box interface under
295 the equilibrium state. It is noted that the derivation for horizontal fluxes may have uncertainties
296 because emissions from a single line source are simply assumed to be injected into one grid in the
297 multi-box model, whereas a Gaussian distribution over the carriageway was assumed in the LES
298 model. A coarser grid resolution can reduce this type of error. Therefore, in this study, the 16- and
299 32-box models are used to evaluate reactive species in regular and deep urban street canyons, and the
300 details of advective and turbulent velocities are presented in the support information (Table S1 and
301 Table S2). Model performance under higher horizontal resolutions is left to be evaluated in future
302 work.

303 Due to the very similar magnitude of advective fluxes entering and escaping the street canyon in the
304 neutral atmosphere (Salizzoni et al., 2009), the escaping canyon fluxes due to vertical advection are
305 rotated into the horizontal direction, and thus the multi-box model considers only turbulent terms at
306 the rooftop. Turbulent velocities as described above could be negative (although usually of small
307 magnitude in such cases), indicating counter-gradient turbulent diffusion under the box-model
308 framework (Figs. 1c and 1d). This may cause the model to crash during a box model timestep if
309 concentrations become negative. In order to address this issue, a minimum positive value of 1.0×10^{-4}
310 m s^{-1} is applied to the turbulent velocity, implying a very small turbulent flux compared to the
311 advective flux for the interface under such conditions.

2.2.3 Emissions and chemical mechanism

Traffic-related pollutants were emitted by two consecutive line-sources at 1 m height and at 2.5 m to the left and right of the street centre axis for the LES models. Therefore the emissions were evenly injected into $\text{Box}_{[1,2]}$ and $\text{Box}_{[1,3]}$ at the street level for the multi-box models ($\gamma_2 = \gamma_3 = 0.5$). The emission rates of NO, NO₂, carbon monoxide (CO), ethene (C₂H₄), propene (C₃H₆), formaldehyde (HCHO) and acetaldehyde (CH₃CHO) were calculated based on the UK Road Vehicle Emission Factors (Boulter et al., 2009), which were 558, 62, 1356, 56, 24, 32 and 15 g km⁻¹ hr⁻¹, representing a typical weekday traffic scenario of 1500 vehicles per hour with an average speed of 30 miles per hour (mph), which equate to 900, 100, 3593, 347, 150, 96 and 98 ppb s⁻¹ for the LES cell; to 4.0, 0.44, 16, 1.55, 0.67, 0.88 and 0.42 ppb s⁻¹ for the 16- (and 32-) box grid; and to 0.252, 0.028, 1.0, 0.097, 0.042, 0.055 and 0.026 ppb s⁻¹ for the one-box grid in the regular canyon and for the two-box grid in the deep canyon. Moreover, PS that only undergoes physical processes was injected at the same rate with NO_x (= NO + NO₂) to investigate the sole effect of canyon dynamics on the model performance.

The Reduced Chemical Scheme (RCS) was developed by Bright et al. (2013), based on the Common Representative Intermediates mechanism version CRI v2-R5 (Watson et al., 2008). The RCS retains the compounds that have important effects on core chemical intermediates in urban street canyons and includes 51 gas-phase species and 136 chemical reactions. The chemical kinetics are calculated at 20°C under a standard atmosphere pressure (e.g., photodegradation rate of 9.20×10^{-3} s⁻¹ for NO₂), and they are adopted for simulating the daytime chemistry in the present study. The comparison between RCS and the benchmark Master Chemical Mechanism (12691 chemical reactions of 4351 species for MCMv3.0) (Saunders et al., 2003) showed that the maximum differences were 3%, 13%, 16%, and 12% for NO, NO₂, O₃ and OH during a four-hour simulation, which was comparable to, or smaller than, the errors from emissions and detection techniques (Boulter et al., 2009; Heard and Pilling, 2003). The total computation time of the multi-box model with RCS is about 6 minutes, which is higher than that of the one-box (~1 s) and two-box models (~8 s), but is significantly faster compared to those of the LES (around 10 days) (Zhong et al., 2017).

2.2.4 Model initialisation and output post-processing

The initial concentrations of the multi-box model are consistent with those in LES conditions for two types of canyons, which were taken from the field study of the Tropospheric Organic Chemistry

341 (TORCH) experiment (Lee et al., 2006). These observations represent a typical atmospheric condition
 342 in rural London, UK, during the summer of 2003. The models have been operated for 30 min “spin-
 343 up” period without any emission in order to initialise the chemical intermediates, then traffic
 344 emissions are switched on and concentrations of all species at $t = 30$ min are used as the cyclic “fixed”
 345 background conditions for a next 210 min modelling duration (i.e., $t = 30$ -240 min) in a time step of
 346 0.03 s. The solar radiation intensity remains constant during the model operation. “zero background”
 347 for PS is assumed during the LES and box operations.

348 The concentrations from box models were stored in an interval time step of 1 min for time-evolution
 349 analysis. The final hour of the modelling results (i.e., $t = 180$ -240 min) was extracted for the
 350 calculation. Details about LES outputs pre-processing can be found in Bright et al. (2013) (for the
 351 regular canyon) and Zhong et al. (2015) (for the deep canyon), respectively. Subsequently, the
 352 modelling results of LES were averaged equivalent to the coarse resolution of box models for the
 353 purpose of evaluating the spatial distributions of air pollutants with the multi-box models. In order to
 354 investigate the impacts of incomplete mixing on the sub-grid scale variability due to chemistry, a
 355 widely-used dimensionless parameter *intensity of segregation* (Krol et al., 2000) is adopted in this
 356 study:

$$357 \quad I_{S(q_1+q_2)} = \frac{\langle q_1^* q_2^* \rangle}{\langle q_1 \rangle \langle q_2 \rangle} \quad (21)$$

358 where $I_{S(q_1+q_2)}$ represents the intensity of segregation between chemical species q_1 and q_2 , angle
 359 brackets refer to the volume-averaged conservation quality, defined by Equation (8) for the multi-box
 360 model, $\langle q_1^* q_2^* \rangle$ represents the volume-averaged covariance between q_1 and q_2 , and an asterisk means
 361 the deviation from the averaged value. So, for a general case, we have $\langle q_1 \rangle = \sum \alpha_k \beta_1 C_{q_1, [k,i]}$,

$$362 \quad \langle q_2 \rangle = \sum \alpha_k \beta_1 C_{q_2, [k,i]}, \quad q_{1, [k,i]}^* = C_{q_1, [k,i]} - \langle q_1 \rangle, \quad q_{2, [k,i]}^* = C_{q_2, [k,i]} - \langle q_2 \rangle, \quad \text{and} \quad \langle q_1^* q_2^* \rangle = \frac{1}{16} \sum q_{1, [k,i]}^* q_{2, [k,i]}^* . \quad \text{The}$$

363 volume-averaged second-order reaction rate ($\langle k_{(q_1+q_2)} \rangle$) can be written as:

$$364 \quad \langle k_{(q_1+q_2)} \rangle = k_{(q_1+q_2)} (1 + I_{S(q_1+q_2)}) \quad (22)$$

365 where $k_{(q_1+q_2)}$ is the original reaction rate in the sufficiently well-mixed one-box model. Therefore,
366 $I_{S(q_1+q_2)}$ can also be thought of as quantifying the deviation from chemical equilibrium due to the
367 spatial segregation associated with atmospheric dynamics. For any species in the one-box model,
368 $I_{S(q_1+q_2)}$ equals to zero because there is no spatial segregation inside the box. A positive $I_{S(q_1+q_2)}$
369 indicates that $\langle k_{(q_1+q_2)} \rangle$ in the 16-box model is larger than $k_{(q_1+q_2)}$ in the one-box model because of the
370 segregation effect. When $q_1 = q_2$, $I_{S(q_1+q_2)}$ represents the spatial variability of any specific pollutant in
371 respect to its canyon volume-averaged concentration.

372 **3. Results and discussion**

373 3.1. Temporal evolution and the intensity of segregation within the canyon

374 Fig. 2 shows temporal evolution of the volume-averaged pollutant mixing-ratios of the LES, and
375 multi- and one-box models under the same raw emissions, meteorological conditions, and RCS
376 chemistry as the LES model, in idealised regular (a, b) and deep (c, d) street canyons, respectively.
377 Modelling results of the two-box model are available only for the deep canyon. The LES-RCS data
378 for the regular and deep canyons are from Bright et al. (2013) and Zhong et al. (2017), respectively.
379 The box models produce less temporal variability; reactive species slowly move toward a chemical-
380 transport equilibrium. The O₃ concentrations drop sharply after zero-emission “spin-up” period due
381 to NO and O₃ titration, accelerating the formation of NO₂ in street canyons. The OH and HO₂
382 concentrations rapidly relax to equilibrium when traffic emissions are switched on, the maintenance
383 of a steady-state HO₂ by oxidation of VOCs contributes to an additional NO₂ fraction. The chemical
384 species in the regular canyon achieve a transport-chemistry balance more quickly (i.e., about 90 min
385 for NO and NO₂) than in the deep canyon (e.g., around two hours for NO and NO₂) in both one- and
386 multi-box models, and the LES, because of more effective ventilation for canyons with a lower AR.
387 The concentrations of selected species at the equilibrium state over the final hour of model operation
388 are discussed below for a better understanding of the coupled dynamical and chemical processes in
389 street canyons.

390 Table 1 illustrates the time-averaged mixing-ratios of PS, NO, NO₂, O₃, OH, HO₂, NO_x (= NO +
391 NO₂), O_x (= NO₂ + O₃) and NO₂/NO ratios from the models coupled with the RCS chemistry during
392 the final simulation period ($180 \leq t \leq 240$ min) in the regular and deep street canyons, respectively.

393 For the regular canyon, LES outputs are slightly different from those in Bright et al. (2013), because
394 of the dynamics-driven variability in concentrations and the difference in averaging times (i.e., $180 \leq$
395 $t \leq 240$ min in the current study vs. $150 \leq t \leq 210$ min). The modelled mean PS concentrations of box
396 models are in good agreement with those of the LES-based models (differences within $\pm 0.5\%$). The
397 spatially-averaged NO concentrations over the canyon are underestimated up to 4% by the box
398 models, while levels of other species are all overestimated to different extents, in particular with OH
399 levels (overestimated up to around 38%). The overestimations of O₃ are about 5.7% and 2.8% by the
400 one- and 16-box models compared to more accurate LES modelling results.

401 Previous modelling results (Bright et al., 2013) showed that the NO_x modelled by the zero-
402 dimensional one-box model were about 8 ppb (3.3%) higher than by the LES, O₃ concentrations were
403 underestimated by 6%, and NO was overestimated by around 1%. One explanation for the difference
404 between the results of the, present study and the previous study is that the value of $w_{e,0}$ adopted for
405 representing the canyon ventilation is ~5% higher in this study (i.e., $w_{e,0} = 0.22$) compared to their
406 work (i.e., $w_{e,0} = 0.21$), resulting in a higher abundance of O₃ in the street canyon due to inward
407 transport in from the background. The NO and NO₂ concentrations would be changed accordingly. It
408 should be noted that NO_x increases by about 1.6% in the box models, which means NO_x loss processes
409 (e.g., production of nitric acid and organic nitrates) are more effective in the LES model. The O_x
410 levels are respectively 9.0% and 11.1% higher in the 16- and one-box models because the efficient
411 mixing of background O₃ and in-canyon NO produces more NO₂ than under less efficient mixing
412 conditions, which is further discussed in the following section.

413 For the deep canyon, it is noted that the coarse resolution leads to a systematic underestimation of
414 about 3.0% in PS by the one-box model and of about 2% by the two-box model (Table 1); additionally,
415 Table 2 presents the mixing-ratio of pollutants from the LES, 32-box and two-box models in the upper
416 and lower compartments of the deep canyon, respectively. The 32-box model performs better than
417 the two-box model in simulating PS in both compartments. This may be attributed to the definition
418 of the exchange velocity (e.g., $w_{e,0}$, which is calculated based on the gradient between the whole
419 canyon-averaged and the ambient concentrations) and other dynamical parameters. For example, the
420 exchange velocities for the two-box and multi-box models are calculated based on the gradient
421 between the concentrations in the rooftop boxes and the overlying background. This more local

422 concentration gradient provides a better description for the flux balance in the multi-box model and
423 thus, better modelling results (i.e., closer to the LES outputs).

424 The difference in the flux-balanced exchange velocities may also have important impacts on the
425 chemistry in the canyon. The multi-box models include horizontal transport that can compensate
426 errors in modelling the canyon-averaged concentration, as less PS levels (around 1.5%) are
427 underestimated by the 32-box model. The box models overestimate reactive species except for NO
428 compared to the LES, which is consistent with the tendencies in the regular canyon. The absolute
429 errors between the LES and box models are larger for NO, NO₂ and O₃ but are smaller for OH and
430 HO₂ in the deep canyon, partly due to poor ventilation. Moreover, different chemical regimes may
431 exist in the canyon in terms of multi-vortices formed in the canyon, generating a complex outcome
432 for the whole-canyon averages. O_x is overestimated, as in the regular canyon, but NO_x is
433 underestimated partly because of the advective and turbulent velocities used in the model. The
434 concentration differences show a consistent pattern going from the complete-mixing (i.e., one-box
435 resolution) to resolved transport-and-mixing (i.e., LES resolution) conditions for most reactive
436 pollutants. That is, the multi-box model in general presents closer-to-LES results than the one-box
437 model. For example, instant mixing in street canyons accelerates more NO conversion to NO₂. Hence,
438 NO₂ is overestimated around 13% by the one-box model but 10% by the 16-box model in the regular
439 canyon; and it is overestimated about 15% by the one-box model, 13.6% by the two-box model, but
440 8.6% by the 32-box model in the deep canyon (Table 1 and 2). The NO₂/NO ratios gradually reduce
441 from 0.53 to 0.45, and from 0.63 to 0.47 in the regular and deep street canyons from the one-grid
442 approximation to the highest resolution, respectively. For the modelled concentrations of HO_x (= OH
443 + HO₂), the results of all the box models are very close to each other. The rationale behind this
444 similarity is discussed below using the intensity of segregation.

445 Table 3 compares the percentage intensities of segregation between any two selected chemical species
446 from the LES and multi-box models in regular and deep street canyons. It shows that the most
447 spatially variable air pollutant in the canyon is NO (i.e., indicated by $I_{S(\text{NO}+\text{NO}_2)}$) and the least spatially
448 variable pollutant is OH (i.e., indicated by $I_{S(\text{OH}+\text{OH}_2)}$) in general. Not surprisingly almost all intensities
449 of segregation in the deep street canyon are considerably higher (e.g., 26% for $I_{S(\text{NO}+\text{NO}_2)}$) than those in
450 the regular canyon (e.g., 3% for $I_{S(\text{NO}+\text{NO}_2)}$), which are supported by Zhong et al. (2017). It indicates
451 that NO concentrations become more heterogeneous due to the perturbed in-canyon vortexes, but OH

452 is less affected in terms of their abundance and lifetime in street canyons. It is also found that within
453 both types of canyons the values of $I_{S(q_1+q_2)}$ in the multi-box models mostly have a same sign with
454 those in the LES, which is positive for “emitted-inside-canyon” species (i.e., NO, NO₂) and is negative
455 for “entrained-from-background” and “formed in situ” species (i.e., O₃, OH, HO₂), indicating similar
456 chemical behaviours (e.g., generation or depletion) of these species in the LES and multi-scale models.

457 For atmospheric pollutants that directly react with each other, the intensities of segregation of the
458 multi-box models have a trend toward “zero” compared with the LES. For example, NO and O₃
459 titration rates are slower by 2.79% and 10.02% in regular and deep canyons under the “true” condition,
460 but slower by only 1.34% and 9.35% respectively under the “multi-box” approximation at the current
461 resolution. That is, the performance of multi-box model is in between the LES (i.e., close to the “true”
462 flow, insufficient mixing with hundreds of thousands of boxes) and the one-box model (i.e., instant
463 and homogeneous mixing with no segregation). On the contrary, the reactions of O₃ and HO₂
464 producing OH have been accelerated by 1.74% and 2.36% in the LES, but only by 0.42% and 1.56%
465 in the multi-box models.

466 3.2. The spatial variation of pollutants in street canyons

467 Fig. 3 illustrates vertical profiles ($0 < z/l_0 < 0.25$, $0.25 < z/l_0 < 0.5$, $0.5 < z/l_0 < 0.75$, $0.75 < z/l_0 < 1.0$)
468 of time-averaged PS, NO, NO₂, O₃, OH and HO₂ in the regular canyon (black lines), along with the
469 leeward (blue lines) and windward walls (red lines) in order to further ascertain the performance of
470 the multi-box model. In order to compare the model performance of the LES and multi-box model at
471 the same grid resolution, the quantities of the LES were averaged over the entire street width ($-0.5 <$
472 $x/l_0 < 0.5$) and the nearest box grid resolution adjacent to the canyon walls (i.e., $-0.5 < x/l_0 < -0.25$ for
473 the leeward, $0.25 < x/l_0 < 0.5$ for the windward), respectively. The distributions of chemical species
474 from the 16- and 32-box models are presented in the support information. Over the final hour of the
475 simulation, it is clear that the vertical distributions of PS from the 16-box model are in good agreement
476 (within 5 ppb) with those from the LES, with higher concentrations elevated on the leeward side of
477 the street corner (305.0 ppb) and considerably lower concentrations on the windward side (172.2 ppb)
478 due to the in-canyon air circulation. The agreement for PS indicates that the multi-box model with
479 LES-driven dynamics captured the major flow pattern well (i.e., a primary vortex) within the regular
480 street canyon.

481 The agreement for PS also implies that the discrepancies in the abundance of reactive species between
482 the multi-box and the LES models are mainly attributed to NO_x-O₃-VOC chemical reactions, though
483 their vertical and horizontal features are also well-reproduced by the multi-box model. NO
484 concentrations are well-simulated in Box_[1,4] (difference within 1 ppb) but are slightly underestimated
485 at other grid points, especially for the leeward side (difference around 8 ppb, ~3%). Due to less
486 segregation for the 16-box model, there is a more effective O₃ titration in the canyon, which depletes
487 more NO turning to NO₂, thus causing an underestimation of NO and an overestimation of NO₂
488 contents in the 16-box model. Errors in NO₂ concentrations become smaller on the windward side but
489 become larger on the leeward side from the street level to the canyon rooftop. The maximum
490 overestimation of NO₂ is 15.6 ppb (19.1%) at $0 < z/l_0 < 0.25$ of the leeward side (Box_[1,1]) and 9.9 ppb
491 (12.6%) at $0.5 < z/l_0 < 0.75$ of the windward side (Box_[3,4]). The O₃ concentrations are underestimated
492 on the windward side with the maximum box-minus-LES value of 2.2 ppb (24.3%) at $0 < z/l_0 < 0.25$
493 (Box_[1,1]) due to effective titration with recirculated NO under the sufficient mixing condition.
494 However, there is an clear overestimation in O₃ contents on the leeward side with the maximum
495 difference of -3.6 ppb (19.1%) at $0.75 < z/l_0 < 1.0$ (Box_[1,4]). The explanation could be that more O₃
496 accumulated on the leeward side because of the rapid photodegradation of overproduced NO₂ due to
497 segregation effects on the windward side. Moreover, higher O_x levels in the 16-box model may be
498 attributed to overestimated HO_x concentrations (about 0.03 ppt for OH, 0.07 ppt for HO₂) in the street
499 canyon.

500 The Damköhler number (Da) is a widely used ratio of the chemical reaction rate to the diffusion rate
501 (or, equivalently, the ratio of diffusion timescale to reaction timescale) for determining the importance
502 of segregation effects on reactive species (Driscoll et al., 1992). If $Da \ll 1$, the dynamics achieve
503 “equilibrium” much faster than the reaction, leading to minimal segregation effects for the pollutant;
504 if $Da \gg 1$, the reaction may be considered to reach equilibrium instantaneously compared to the
505 relatively slower diffusion rates. That is, dynamics for the pollutants with substantial segregation
506 effects must be considered in a coupled manner (Garmory et al., 2006). Zhong et al. (2017) reported
507 the Da numbers of NO, NO₂, O₃, OH and HO₂ in the street canyon, which were 3.4, 5.8, 69, 1.44×10^5
508 and 4.44×10^4 , respectively. The chemical reaction rates of HO_x are much quicker than the diffusion
509 rates across the model grid ($Da \gg 1$). This explains that HO₂ is 0.02 ppt higher on the windward side

510 compared to the leeward side, and vertical distributions of OH and HO₂ from the 16-box model are
511 not significant compared to the outputs of the LES.

512 Fig. 4 shows the vertical distributions of selected species in the deep canyon and along with the
513 leeward and windward facets. The PS concentrations of the 32-box model are well-matched to those
514 of the LES model in both vertical and horizontal directions, with an underestimation of around 10
515 ppb (2.4%) on the leeward side and of 25 ppb (2.5%) on the windward side. Cumulative traffic
516 emissions produce very high PS mixing-ratios in the lower part of the canyon ($0 < z/l_0 < 1.0$). The
517 concentration of PS decreases smoothly with height on the leeward side but sharply varies on the
518 windward side, e.g., concentrations are higher on the windward side when $0 < z/l_0 < 1.0$ but on the
519 leeward side when $1.0 < z/l_0 < 2.0$. This indicates noticeable segregation between the lower and upper
520 compartments of the deep street canyon due to the presence of two counter-rotating vortices that are
521 captured by the LES and multi-box models. The upper vortex is driven by the ambient wind in the
522 shear layer at the rooftop ($z/l_0 = 2.0$) so that the characteristics of the vertical profiles are similar to
523 those of the regular canyon, while the lower vortex is driven by the upper vortex at approximately z/l_0
524 $= 1.0$ (or slightly lower) (Eliasson et al., 2006; Zhong et al., 2015).

525 Considering the chemically reactive species, NO concentrations are slightly underestimated on both
526 sides of the canyon as for the regular canyon, with the maximum difference of 74.9 ppb (9.5%) in
527 Box_[1,4] on the windward corner. NO₂ is overestimated by the 32-box model, especially for the
528 windward side (~6%). The O₃ concentrations are slightly underestimated by the 32-box model in the
529 upper vortex on the windward side (~5%), and then are overestimated when air parcel moves to the
530 upper leeward side and to the lower vortex (~10%). Although HO_x concentrations are still higher in
531 the 32-box model compared to the LES, the spatial profiles of HO₂ are well reproduced due to a longer
532 diffusion time in the deep canyon in contrast to the regular canyon. Errors in modelled concentrations
533 (solid and dash lines with the same color) always become larger at $0 < z/l_0 < 1.0$ but become closer at
534 $1.0 < z/l_0 < 2.0$. For example, the 32-box model overestimates NO₂ and O₃ by 14.9% and 27.4% on
535 the leeward side, and by 18.1% and 23.1% on the windward side at the street level ($0 < z/l_0 < 0.25$),
536 but those differences decrease to 8.1%, 17.8%, 14.4% and -10.0% at the rooftop level ($0.75 < z/l_0 <$
537 1.0), respectively. Therefore, when applying the grid assumption to simulate reactive species in street
538 canyons, it is necessary to consider the inherent uncertainty due to chemical reactions being too fast
539 in particular at the street level, which may lead to an overestimation of pedestrian exposure risks.

540 3.3. The model performance with the simple NO_x-O₃ cycle

541 The performance of the multi-box models has been further evaluated with the simplified NO_x-O₃
542 chemical reactions:



545 where $h\nu$ represents solar photons. The production and photodegradation coefficients of NO₂ are
546 taken from the RCS chemistry, which are $4.01 \times 10^{-4} \text{ ppb}^{-1} \text{ s}^{-1}$ and $9.20 \times 10^{-3} \text{ s}^{-1}$, respectively. The
547 QSSA is adopted for the calculation. The computation time for the multi-box model using the simple
548 NO_x-O₃ cycle is much quicker (~40 s) than that using RCS chemistry. Table 4 illustrates the time-
549 and spatial-averaged mixing-ratios of NO, NO₂, O₃, NO_x, O_x and NO₂/NO ratios with solely NO_x-O₃
550 reactions during the final hour in the regular and deep canyons. The modelled NO_x concentrations of
551 the LES model are about 1.9% lower in the regular canyon but are slightly higher in the deep canyon
552 compared to the box models, partly because of strong turbulent fluctuations in the LES. As expected,
553 NO₂ concentrations gradually reduce by 10-13% in the regular canyon without involving OH/HO₂
554 chemicals, thereby decreasing O_x contents to a similar magnitude. This reduction is rather significant
555 (37-41%) in the deep canyon, resulting in dramatically lower NO₂/NO ratios, which is demonstrated
556 by Zhong et al. (2017). In comparison with the LES outputs, O₃ concentrations are underestimated
557 by the box models to different extents in both regular and deep canyons, which differs from its trends
558 of overestimation with the RCS chemistry. Overall the multi-box models performed better than the
559 one-box and two-box models in simulating all reactive species except O₃.

560 Modelling differences between the four models shrink when using the very simplified chemical
561 reactions in contrast to the NO_x-O₃-VOC chemical scheme. More specifically, Fig. 5 presents vertical
562 profiles of the difference of NO, NO₂ and O₃ concentrations under different chemical schemes using
563 the multi-box models and the LES, respectively. It clearly exhibits the direct contributions from the
564 VOCs mechanisms spatially, which, in general, shows more substantial influences on the modelling
565 results of the multi-box model compared to those of the LES. The contributions of VOCs are
566 significantly enhanced in the deep street canyon compared to the regular canyon; involving VOCs

567 tends to have greater impacts on local emitted species such as NO and NO₂ in the lower compartments
568 (i.e., $0 < z/l_0 < 1.0$) and on remote species such as O₃ in the upper compartments (i.e., $1.0 < z/l_0 < 2.0$).
569 Although there are some differences between the performance of the multi-box models and the LES,
570 the multi-box models reproduce well the contribution of VOCs to the vertical distribution of NO₂ and
571 O₃.

572 Fig. 6 illustrates the vertical distributions of NO, NO₂, and O₃ in the regular (a, b, c) and deep (d, e,
573 f) street canyons (i.e., horizontal-averaged concentration cross $-0.5 < x/l_0 < 0.5$) and along the canyon
574 walls (i.e., $-0.5 < x/l_0 < -0.25$ and $0.25 < x/l_0 < 0.5$). In the regular canyon, NO is well-simulated with
575 negligible difference between two models. The vertical profile of NO₂ is overestimated by the 16-box
576 model due to segregation. Nevertheless, extent of overestimation in NO₂ becomes smaller compared
577 to the simulations with NO_x-O₃-VOC chemistry, leading to relatively lower O₃ concentrations along
578 the leeward wall. However, underestimation of O₃ becomes larger on the windward side, with a
579 maximum difference of 4.8 ppb at the rooftop ($0.75 < z/l_0 < 1.0$). In the deep canyon, we also obtained
580 satisfied vertical distributions of all three species from the 32-box model compared to those in the
581 LES. Concentration gradients of NO₂ and O₃ lessen when applying the simple NO_x-O₃ cycle,
582 especially for the street level. NO₂ is still overestimated on the windward side and underestimated on
583 the leeward side, but to a much acceptable degree compared to the scenario with the RCS (Fig. 4).
584 The trends of O₃ are the same as that of the regular canyon in the upper vortex ($1.0 < z/l_0 < 2.0$), e.g.,
585 an underestimation on the windward side and overestimation on the leeward side but are consistent
586 with the LES in the lower vortex ($0 < z/l_0 < 1.0$). This indicates that the modelling domain with VOC
587 emissions (e.g., high coverage of vegetation) should consider using air quality models with resolution
588 as high possible to diminish the segregation effects in chemistry.

589 **4. Conclusions**

590 A process-based multi-box photochemical street-canyon model, with a flexible number of boxes and
591 based on a set of transport parameters calculated from the LES outputs is developed in this study. The
592 performance of this street-scale chemical transport model coupled with NO_x-O₃-VOC chemistry (or
593 a simplified NO_x-O₃ cycle), for modelling reactive species in the regular (AR = 1) and deep (AR = 2)
594 street canyons, is compared to published LES data and to one- and two-box simulations. Results show
595 that the model configured with a single box captures the average state of air pollutants in street

596 canyons, and the results are consistent with previous studies (Bright et al., 2013; Zhong et al., 2016).
597 Compared with the benchmark LES simulation, the multi-box model reproduces well the spatial
598 contrast in pollutant concentrations inside the street canyons in particular with a chemically inert
599 passive scalar. Namely, the spatial concentration patterns have been captured for the regular street
600 canyon (i.e., single primary vortex), as well as for the deep street canyon (i.e., two counter-rotating
601 vortices). For the regular canyon, it is found that the NO and O₃ titration reaction becomes more
602 effective due to less segregation along the windward facet, leading to an underestimate of O₃ and an
603 overestimate of NO₂ levels. On the leeward side, the overestimated NO₂ results in an overestimation
604 of O₃ through photolysis. An additional overestimation of NO₂ is attributed to OH/HO₂ chemistry.
605 The impact of segregation effects on the reactive species is substantial, in particular for short-lived
606 species. The OH and HO₂ concentrations are overestimated by the multi-box model, their vertical
607 variations are not very significant compared to the LES because of their very short chemical
608 timescales.

609 In a deep canyon with poor ventilation, the relationship of reactive species in the upper compartment
610 between the 32-box and LES models is consistent with those in the regular canyon. However, in the
611 lower compartments of the canyon, O₃ is always overestimated, and there are obvious variations of
612 OH and HO₂ concentrations along the vertical direction. Additionally, under the simple NO_x-O₃ cycle,
613 the differences between the multi-box model and the LES become smaller particularly for the deep
614 canyon. Although the effects of VOC chemistry on reactive species such as NO₂ are underestimated
615 compared to the LES, the multi-box models capture the vertical contribution of VOCs to these
616 pollutants in street canyons, and are a significant step forward from the simple one- and two-box
617 models. The multi-box model is less computationally efficient than the typical one-box (~1 s) and
618 two-box models (~8 s), but it can offer spatial information on reactive species within street canyons
619 based on coupled chemical-transport processes (with VOC chemistry) in a fairly short computational
620 times (~6 min) compared to the LES (e.g., ~10 days in Zhong et al. (2017)). Overall, the multi-box
621 model enables insightful investigations into the multiple processes as well as their complex
622 interactions and is of practical utility for air quality assessment or pollution mitigation management
623 in street canyons.

624 Further works may focus on adopting the multi-box model to simulate air pollution in street canyons
625 with spatially segregated emissions due to the presence of vegetation. More detailed chemical
626 schemes (e.g., Master Chemical Mechanism, MCMv3.0 (Saunders et al., 2003)) can be incorporated

627 into the model. The evaluation of modelling results with field measurements is recommended. A more
628 challenging task is to merge the effects of airflow parallel to the street axis on the diffusion and
629 transformation of chemical species, which would enable for a wider range of model applications.

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Figures

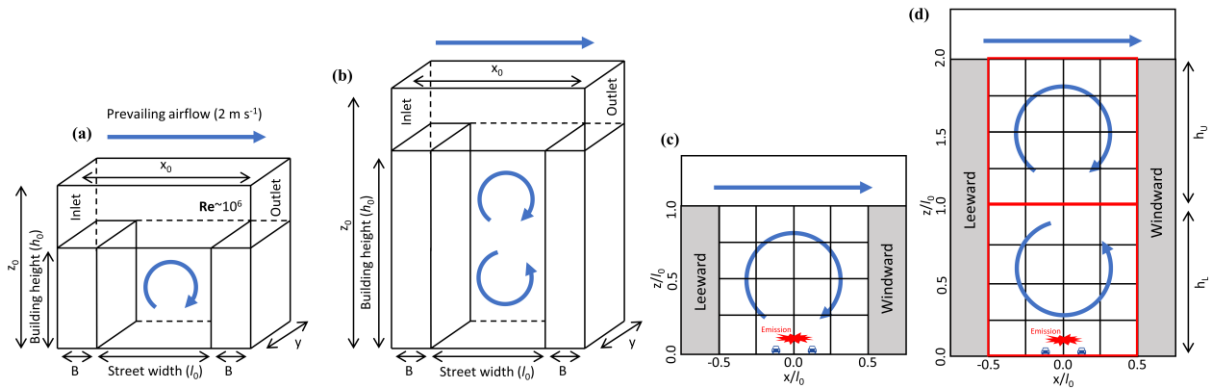


Fig. 1. Schematic diagram of (a) the LES domain for a regular urban street canyon where $x_0 = 24 \text{ m}$, $y_0 = 40 \text{ m}$ and $z_0 = 90 \text{ m}$, and canyon geometries $l_0 = h_0 = 18 \text{ m}$, wall width $B = 3 \text{ m}$, modified from Bright et al. (2013); (b) the LES domain for a deep canyon $x_0 = 36 \text{ m}$, $y_0 = 40 \text{ m}$ and $z_0 = 112 \text{ m}$, and canyon geometries $l_0 = 18 \text{ m}$, $h_0 = 36 \text{ m}$ and $B = 9 \text{ m}$, modified from Zhong et al. (2015); (c) the equivalent multi-box (16) model for the regular canyon; (d) the two-box model denoted by the red frame, where the height of upper (h_U) and lower compartments (h_L) is 9 m , and the multi-box (32) model for the deep canyon. See text for details.

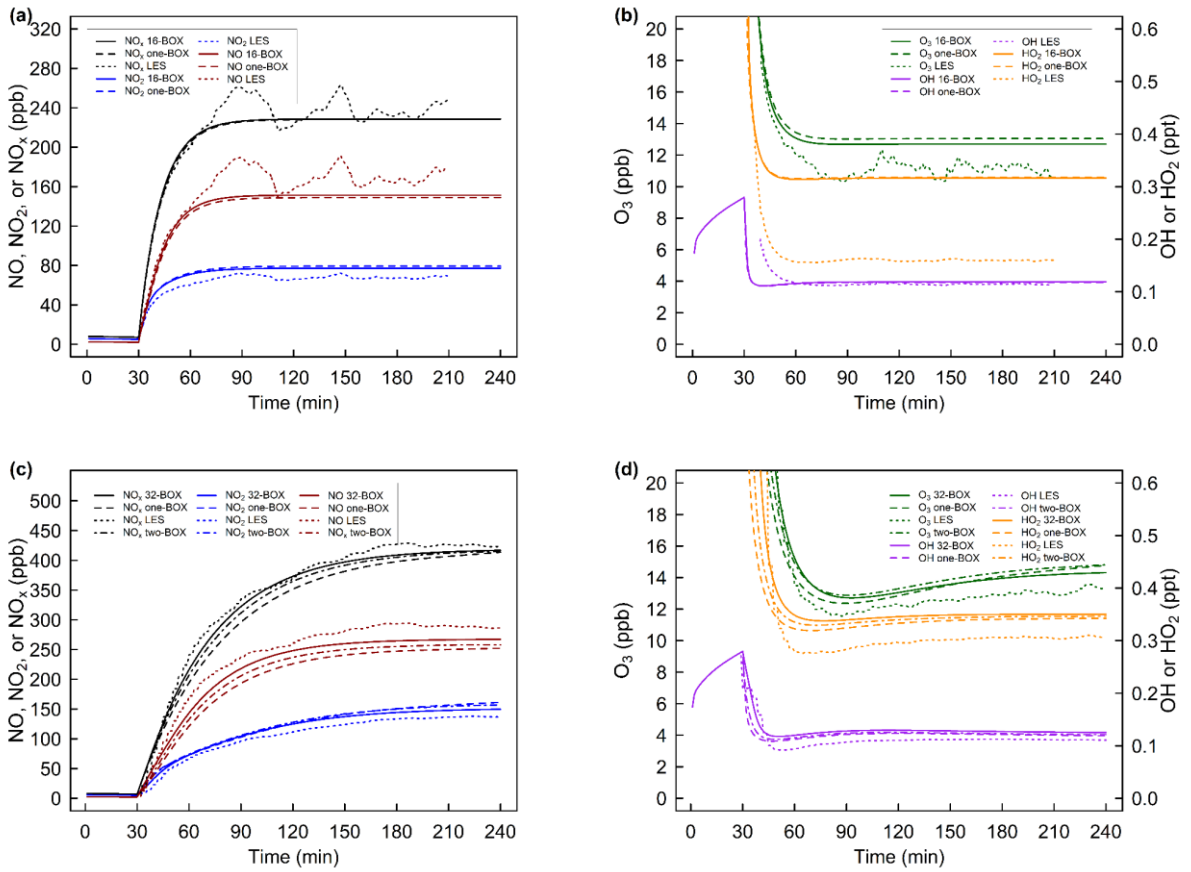


Fig. 2. Temporal variation of the spatially averaged mixing-ratio of NO, NO₂, NO_x, O₃ (ppb), OH and HO₂ (ppt) calculated using the LES, multi-box and a typical one-box models for the regular (a, b) and deep (c, d) street canyons. Two-box simulations are conducted only in the deep canyon, see text for details.

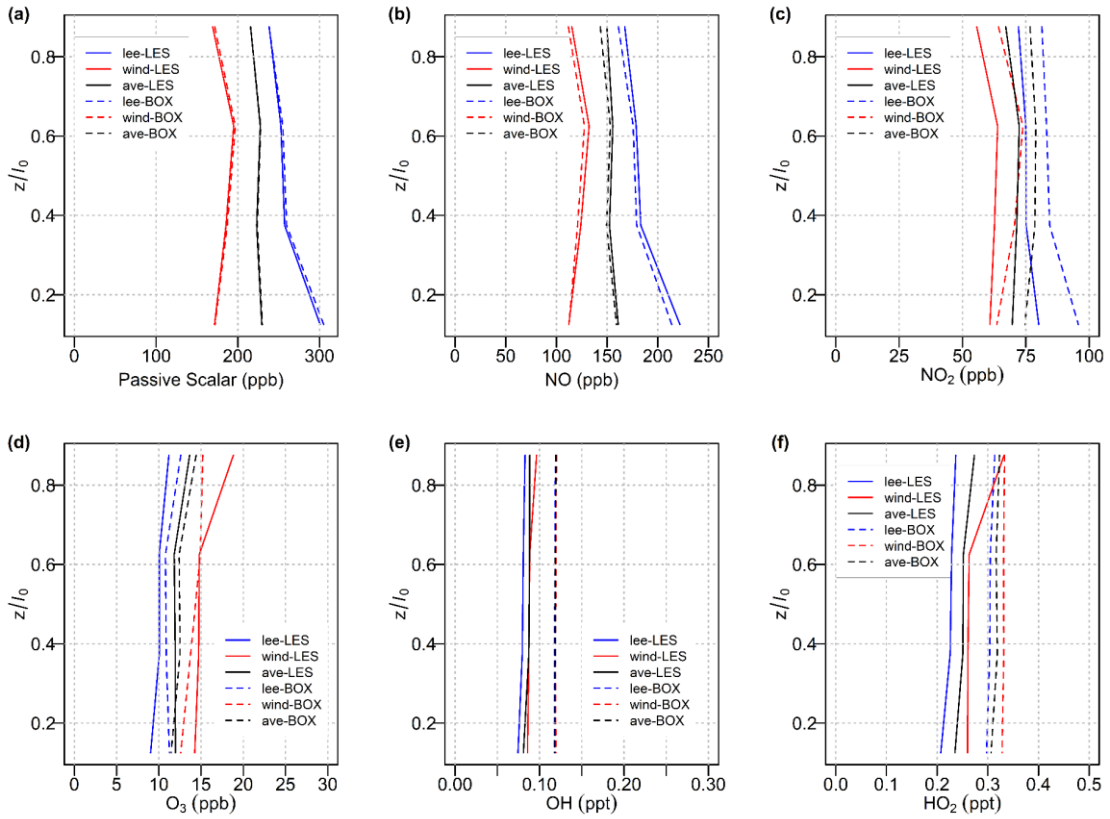


Fig. 3. Vertical profiles of the time-averaged mixing-ratios of PS, NO, NO₂, O₃, OH and HO₂ in the regular street canyon ($-0.5 < x/l_0 < 0.5$) represented by the black lines, along with the leeward wall ($-0.5 < x/l_0 < -0.25$) represented the by blue lines, and along with the windward wall ($0.25 < x/l_0 < 0.5$) represented by the red lines. Solid and dash lines indicate modelling results from LES and multi-box models, respectively.

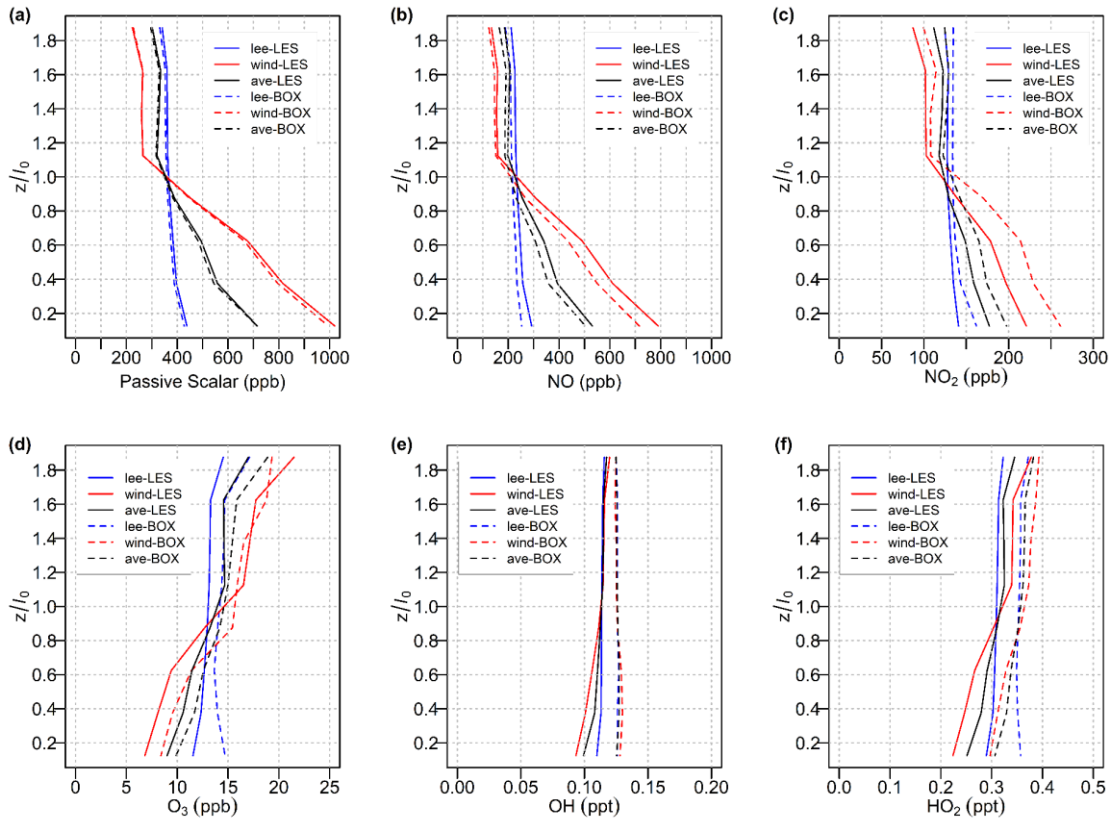


Fig. 4. Vertical profiles of the time-averaged mixing-ratios of PS, NO, NO₂, O₃, OH and HO₂ in the deep street canyon ($-0.5 < x/l_0 < 0.5$) represented by the black lines, along with the leeward wall ($-0.5 < x/l_0 < -0.25$) represented the by blue lines, and along with the windward wall ($0.25 < x/l_0 < 0.5$) represented by the red lines. Solid and dash lines indicate modelling results from LES and multi-box models, respectively.

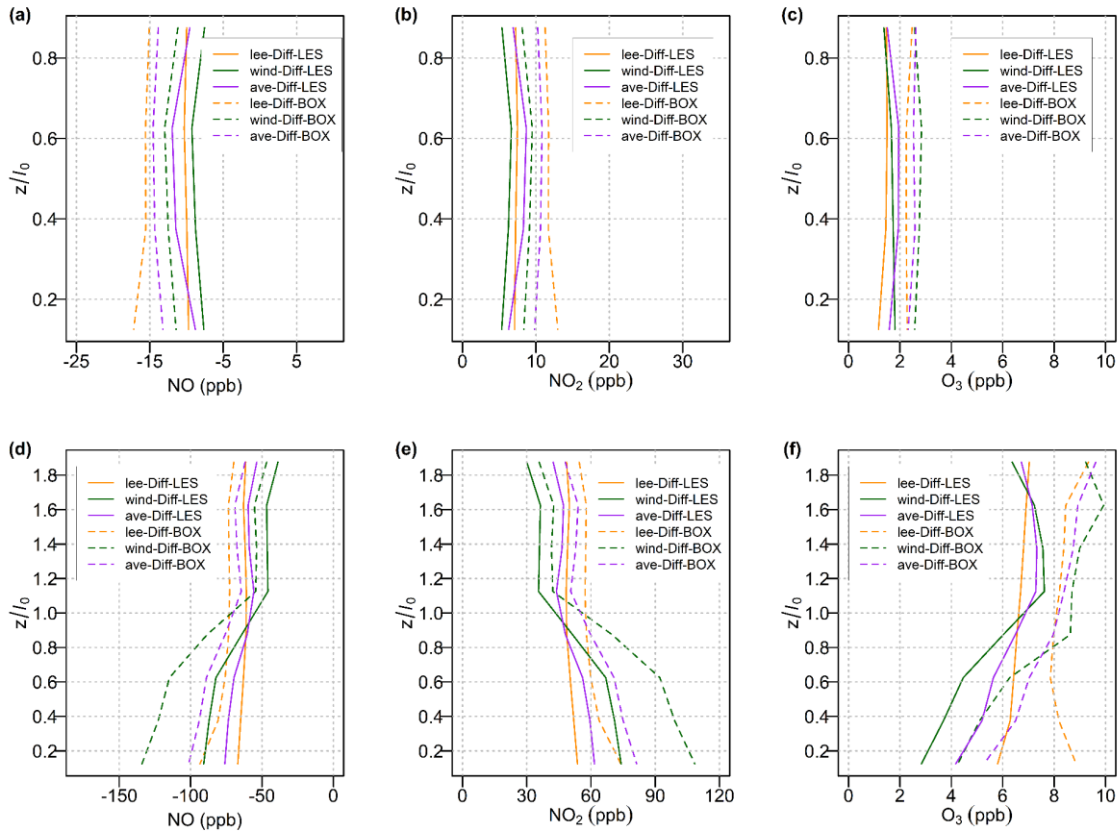


Fig. 5. Vertical profiles of the difference of NO, NO₂ and O₃ concentrations under different chemical schemes (modelling results with the VOC chemistry minus results with the simple NO_x-O₃ chemistry) in regular (a, b, c) and deep (d, e, f) street canyons ($-0.5 < x/l_0 < 0.5$) represented by the purple lines, along with the leeward wall ($-0.5 < x/l_0 < -0.25$) represented by the orange lines, and along with the windward wall ($0.25 < x/l_0 < 0.5$) represented by the green lines. Solid and dash lines indicate modelling results from LES and multi-box models, respectively.

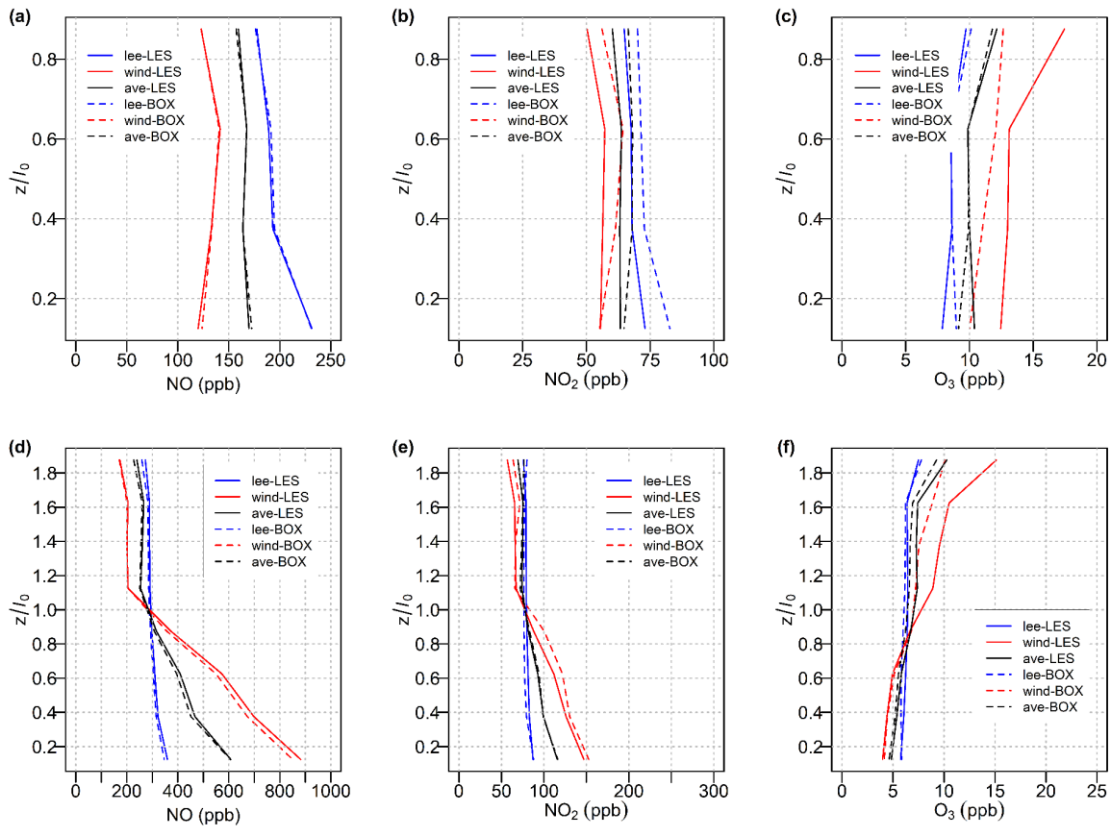


Fig. 6. Vertical profiles of the time-averaged mixing-ratios of NO, NO₂, and O₃ in regular (a, b, c) and deep (d, e, f) street canyons ($-0.5 < x/l_0 < 0.5$) represented by the black lines, along with the leeward wall ($-0.5 < x/l_0 < -0.25$) represented the by blue lines, and along with the windward wall ($0.25 < x/l_0 < 0.5$) represented by the red lines. Solid and dash lines indicate modelling results from LES and multi-box models, respectively.

Tables

Table 1. Time-averaged mixing-ratios from LES, the multi (16/32)-box model, and the one-box and two-box models with RCS chemistry for urban street canyons.

		Mixing-ratio (ppb)							[(b)- (a)]/(a) (%)	[(c)- (a)]/(a) (%)	[(d)- (a)]/(a) (%)
		(a)	(b)	(c)	(d)	(b) - (a)	(c) - (a)	(d) - (a)			
		LES	16/32- box	Two- box	One- box						
The regular street canyon (AR = 1)	PS	223.86	224.33	-	223.85	0.47	-	-0.01	0.21	-	-0.004
	NO	154.66	151.28	-	148.87	-3.38	-	-5.79	-2.19	-	-3.74
	NO ₂	70.12	77.19	-	79.35	7.07	-	9.23	10.08	-	13.16
	O ₃	12.35	12.70	-	13.05	0.35	-	0.70	2.83	-	5.67
	OH*	0.086	0.119	-	0.118	0.033	-	0.032	38.37	-	37.21
	HO ₂ *	0.253	0.316	-	0.318	0.063	-	0.065	24.90	-	25.69
	NO _x	224.78	228.47	-	228.22	3.69	-	3.44	1.64	-	1.51
	O _x	82.47	89.89	-	92.4	7.42	-	9.93	9.00	-	11.05
	NO ₂ /NO	0.45	0.51	-	0.53						
The deep street canyon (AR = 2)	PS	430.23	423.53	421.53	417.06	-6.7	-8.7	-13.17	-1.56	-2.02	-3.06
	NO	289.10	266.48	257.48	250.11	-22.62	-31.62	-38.99	-7.82	-10.94	-13.49
	NO ₂	136.07	147.84	154.57	156.52	11.77	18.5	20.45	8.65	13.60	15.03
	O ₃	13.14	14.19	14.62	14.46	1.05	1.48	1.32	7.99	11.26	10.05
	OH*	0.11	0.13	0.12	0.12	0.02	0.01	0.01	18.18	9.09	9.09
	HO ₂ *	0.31	0.35	0.34	0.34	0.04	0.03	0.03	12.90	9.68	9.68
	NO _x	425.17	414.32	412.05	406.63	-10.85	-13.12	-18.54	-2.55	-3.09	-4.47
	O _x	149.21	162.03	169.19	170.98	12.82	19.98	21.77	8.59	13.39	13.44
	NO ₂ /NO	0.47	0.55	0.60	0.63						

* Mixing-ratio of OH and HO₂ are presented in part per trillion (ppt).

Table 2. Time-averaged mixing-ratios from LES, the 32-box model and the two-box model with RCS chemistry in the deep canyon.

		Mixing-ratio (ppb)					[(b)-(a)]/(a) (%)	[(c)-(a)]/(a) (%)
		(a) LES	(b) 32-box	(c) Two-box	(b) - (a)	(c) - (a)		
The upper compartment	PS	321.23	315.93	313.65	-5.30	-7.58	-1.65	-2.36
	NO	198.14	182.87	178.03	-15.27	-20.11	-7.71	-10.15
	NO ₂	118.44	125.78	129.02	7.34	10.58	6.20	8.93
	O ₃	15.21	16.29	16.96	1.08	1.75	7.10	11.51
	OH*	0.11	0.12	0.12	0.01	0.01	9.09	9.09
	HO ₂ *	0.33	0.36	0.37	0.03	0.04	9.09	12.12
	NO _x	316.58	308.65	307.05	-7.93	-9.53	-2.50	-3.01
	O _x	133.65	142.06	145.97	8.41	12.32	6.29	9.22
NO ₂ /NO	0.60	0.68	0.72					
The lower compartment	PS	539.23	531.13	529.42	-8.10	-9.81	-1.50	-1.82
	NO	380.06	350.09	336.93	-29.97	-43.13	-7.89	-11.35
	NO ₂	153.69	169.90	180.12	16.21	26.43	10.55	17.20
	O ₃	11.06	12.09	12.29	1.03	1.23	9.31	11.12
	OH*	0.11	0.12	0.12	0.01	0.01	9.09	9.09
	HO ₂ *	0.28	0.33	0.33	0.05	0.05	17.86	17.86
	NO _x	533.75	519.99	517.05	-13.76	-16.7	-2.58	-3.13
	O _x	164.75	181.99	192.41	17.24	27.66	10.46	16.79
NO ₂ /NO	0.40	0.48	0.53					

* Mixing-ratio of OH and HO₂ are presented in part per trillion (ppt).

Table 3. The percentage intensities of segregation between pairs of reactive species from the LES and multi-box models (BOX) for the regular and deep canyons. Bold symbols represent species that directly react with each other in the models, and negative values are shown in red.

		LES					BOX				
		NO	NO ₂	O ₃	OH	HO ₂	NO	NO ₂	O ₃	OH	HO ₂
The regular street canyon (AR = 1)	NO	3.02	-	-	-	-	2.97	-	-	-	-
	NO ₂	1.38	0.83	-	-	-	1.61	1.15	-	-	-
	O ₃	-2.79	-1.64	3.54	-	-	-1.34	-0.36	1.61	-	-
	OH	-0.81	-0.27	0.76	0.43	-	-0.08	-0.03	0.07	0.004	-
	HO ₂	-1.43	-0.71	1.74	0.60	1.12	-0.56	-0.20	0.42	0.02	0.15
The deep street canyon (AR = 2)	NO	26.15	-	-	-	-	26.35	-	-	-	-
	NO ₂	10.21	4.18	-	-	-	11.50	5.58	-	-	-
	O ₃	-10.02	-4.35	5.09	-	-	-9.35	-3.86	4.45	-	-
	OH	-2.75	-1.06	1.09	0.31	-	0.22	0.15	-0.09	0.01	-
	HO ₂	-5.13	-2.12	2.36	0.57	1.15	-3.54	-1.45	1.56	-0.03	0.57

Table 4. Time-averaged mixing-ratios from LES, the multi (16/32)-box model, and the one-box and two-box models with the simple NO_x-O₃ chemistry for urban street canyons.

		Mixing-ratio (ppb)							[(b)- (a)]/(a) (%)	[(c)- (a)]/(a) (%)	[(d)- (a)]/(a) (%)		
		(a) LES	(b) 16/32- box	(c) box	Two-	(d) box	One-	(b) - (a)				(c) - (a)	(d) - (a)
The regular street canyon (AR = 1)	NO	165.10	165.27	-		163.24		0.17	-	-1.86	0.10	-	-1.13
	NO ₂	62.58	66.81	-		68.72		4.22	-	6.14	6.75	-	9.81
	O ₃	10.60	10.19	-		10.43		-0.41	-	-0.17	-3.89	-	-1.60
	NO _x	227.69	232.08	-		231.97		4.39	-	4.28	1.93	-	1.88
	O _x	73.18	77.00	-		79.16		3.81	-	5.97	5.21	-	8.16
	NO ₂ /NO	0.38	0.40	-		0.42							
The deep street canyon (AR = 2)	NO	352.57	344.33	336.85		333.14		-8.24	-15.72	-19.43	-2.34	-4.46	-5.51
	NO ₂	85.43	86.30	90.29		92.03		0.87	4.86	6.60	1.02	5.69	7.72
	O ₃	6.89	6.38	6.62		6.45		-0.51	-0.27	-0.44	-7.35	-3.95	-6.43
	NO _x	438.00	430.63	427.14		425.17		-7.37	-10.86	-12.83	-1.68	-2.48	-2.93
	O _x	92.32	92.69	96.91		98.48		0.37	4.59	6.15	0.40	4.97	6.67
	NO ₂ /NO	0.24	0.25	0.27		0.28							