

Numerical investigations of reactive pollutant dispersion and personal exposure in 3D urban-like models

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Accepted Version

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Zhang, Y., Yang, X., Yang, H., Zhang, K., Wang, X., Hang, J., Luo, Z. and Zhou, S. (2020) Numerical investigations of reactive pollutant dispersion and personal exposure in 3D urban-like models. Building and Environment, 169. 106569. ISSN 0360-1323 doi: https://doi.org/10.1016/j.buildenv.2019.106569 Available at http://centaur.reading.ac.uk/87490/

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To link to this article DOI: http://dx.doi.org/10.1016/j.buildenv.2019.106569

Publisher: Elsevier

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- 1 To be submitted to Building and Environment 2019
- 2

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1

20 Abstract

With satisfactory validation by experimental data, we perform computational 21 fluid dynamic(CFD) simulations with the standard $k-\varepsilon$ model to investigate how 22 NO-NO₂-O₃ photochemistry and turbulent mixing influence reactive pollutant 23 dispersion and vehicular NO_x exposure in 21-row(neighborhood-scale~1km) 24 three-dimensional(3D) medium-dense urban models with an approaching wind 25 parallel(perpendicular) to the main(secondary) streets. Personal intake fraction P_iF 26 and its spatially-averaged values for the entire building (i.e. building intake fraction 27 $\langle P_iF \rangle_B$) are adopted for reactive/passive exposure analysis with/without 28 29 NO_x-O₃-photochemistry.

Some meaningful findings are proposed: 1) There are flow adjustment processes 30 coupling turbulent mixing and chemical reactions through urban areas(i.e. secondary 31 Street 1 to 20). NO-NO₂-O₃ photochemistry induces O₃ depletion and NO conversion 32 into NO₂ producing significant increase in NO₂ exposure and slight decrease in NO 33 exposure compared with passive dispersion. 2) With span-wise NO_x sources, Street 10 34 in the fully-developed region experiences weaker wind and subsequently greater 35 $\langle P_iF \rangle_B(0.207 \text{ppm})$ than Street 3(0.135 ppm) in the upstream flow-adjustment region. 36 37 $\langle P_iF \rangle_B$ descends exponentially from the target building toward downstream, and Street 10 experiences quicker decay rates. 3) With stream-wise NO_x sources along the 38 main reaches 39 street. $\langle P_iF \rangle_B$ first ascends, then equilibrium values (e.g.0.046-0.049ppm for passive). 4) If background O₃ concentration [O₃] rises from 40 20ppbv to 40 and 100ppbv, more NO is oxidized by O₃ to generate NO₂. As 41 [O₃]=20ppbv, if NO-NO₂ emission ratio decreases from 10 to 5, NO₂ exposure is 42

partly offset but NO exposure change little. Present methodologies are confirmed
effective to investigate impacts of more complicated meteorological conditions and
chemical mechanisms on exposure in urban districts.

46

Keywords: NO-NO₂-O₃ photochemistry; reactive pollutant dispersion; personal intake fraction (P_iF); building intake fraction ($\langle P_iF \rangle_B$); computational fluid dynamic (CFD) simulation; three-dimensional (3D) urban models

50

51 **1. Introduction**

52 Following the ongoing urbanization worldwide, vehicular pollutant emissions have become one of the major sources in urban air pollution[1-3]. Heavy traffic flows, 53 compact urban configurations and unfavorable meteorological conditions are the main 54 55 reasons of large pollutant exposure and adverse health impacts on city dwellers[4]. On average, people spend more than 90% of their time indoors. Outdoor air pollutants in 56 urban areas can penetrate into indoor via doors, windows, building cracks and other 57 ventilation duct systems[5-7]. Particularly, vehicular pollutant exposure to urban 58 residents living near busy roads should be paid more attention, because they suffer 59 from higher health risks than other urban microenvironments^[7-10]. Apart from 60 reducing vehicular pollutant emissions, sustainable urban design with better 61 understanding the influence of urban layouts and atmospheric conditions on the flow 62 and passive/reactive pollutant dispersion in urban areas can help enhancing pollutant 63

dilution and mitigating traffic-related pollutant exposure[11-15].

As reviewed by the literature [16-29], in the past decades, a number of 65 computational fluid dynamic (CFD) simulations, outdoor field observations and wind 66 tunnel experiments have been widely performed to clarify turbulent flow 67 characteristics and pollutant dispersion in urban models from street-scale (~100m) to 68 neighborhood-scale (~1km). Street aspect ratios (H/W) are reported as the most key 69 urban parameters in two-dimensional (2D) street canyons[16-18, 25-26, 29-32]. 70 Realistic urban districts are usually three-dimensional (3D) with pollutant exchange 71 72 across street roofs and lateral/stream-wise urban boundaries. Generally, the building planar area index λ_p (i.e. the ratio between the planar area of buildings viewed from 73 74 above and the total floor area) and the frontal area index λ_f (i.e. the ratio of the frontal area of buildings to the total floor area) are typical building packing density indexes 75 and key parameters of 3D urban areas[33-38]. Moreover, other significant factors 76 have been also verified, such as building height variations[38-40], ambient wind 77 78 directions[40-41], elevated building design[42-44], tree planting[46-49] etc. In addition, thermal stratification and buoyancy forces induced by wall heating and solar 79 shading also significantly influence or dominate the flow and pollutant dispersion if 80 81 wind speed is relatively small and Richardson number is large[50-56].

Recently, several researches [7, 44, 57] introduce personal intake fraction (P_iF) to quantify street-scale pollutant exposure induced by vehicle emissions. In contrast to population intake fraction (*IF*) for a specific population [4, 58], P_iF is independent of population density and size which represents the fraction of pollutants inhaled averagely by each person of a population to the total pollutant emissions. For example, 1ppm (part per million or 10^{-6}) means the inhalation of 1mg pollutants if 1kg pollutants being emitted. By performing CFD simulations validated by experimental data, Hang et al.[7] estimated spatial mean P_{iF} (i.e. $\langle P_{iF} \rangle \sim 1-5$ ppm) of passive pollutant (i.e. CO) in shallow 2D street canyons (H/W=0.5-1). Later, $\langle P_{iF} \rangle$ in 3D urban district models were confirmed one order smaller (~0.1ppm) than 2D models with similar aspect ratios (H/W=0.5-1)[41, 57].

Besides the dynamic dispersion of passive pollutants, there are chemical 93 94 processes of reactive pollutants in urban streets, such as the NO_x -O₃ photochemistry [59-64], NO_x-O₃-VOCs chemical mechanisms[65-69] etc. Among them, the impacts 95 of different heating scenarios and building configurations on reactive dispersion are 96 extensively investigated through LES or RANS approaches, such as various shading 97 settings[61], wall heating scenarios[60, 62], aspect ratios[63] etc. However, most 98 studies so far mainly examine reactive pollutant dispersion in 2D street canyons while 99 100 investigations on reactive pollutant dispersion and the related exposure in 3D urban models are still rare. Therefore, this study incorporates NO-NO₂-O₃ chemistry into 101 102 CFD simulations and numerically investigates reactive pollutant dispersion and exposure in urban models. As a start, the impacts of various ground-level source 103 locations and reactant proportions (NO:NO2:O3) in 3D medium-dense urban models 104 (*H*/*W*=1, $\lambda_p = \lambda_f = 0.25$) are studied under neutral meteorological conditions. 105

The sketch of this paper is organized as follows: Section 2 introduces the indexesfor pollutant dispersion and exposure. Section 3 illustrates model setups and all test

108	cases in CFD simulations. Section 4 presents the flow and pollutant dispersion
109	validations by wind tunnel data. Section 5 shows results and discussions, and Section
110	6 draws the conclusions.

112 2. Indexes for pollutant dispersion and exposure

113 **2.1** Personal intake fraction (P_iF) and building intake fraction $(\langle P_iF \rangle_B)$

Population intake fraction (*IF*) and personal intake fraction (P_iF) are effective indexes to quantify vehicular pollutant exposure in local streets or neighborhoods[41, 44, 57]. Both exposure indexes are defined as Eq. (1):

117
$$IF = \sum_{i}^{N} \sum_{j}^{M} P_{i} \times Br_{i,j} \times \Delta t_{i,j} \times Ce_{j}/m$$
(1a)

$$P_iF = IF / \sum_{i}^{M} P_i$$
 (1b)

where, N and M are the number of age groups and microenvironments, P_i represents 119 the population size for the age group i, $Br_{i,i}$ (m³/s) and $\Delta t_{i,i}$ (s) are the average 120 breathing rate and the individual time spent for the *i*th age group in the *j*th 121 microenvironment, Ce_i (kg/m³) denotes pollutant concentration in microenvironment j 122 and \dot{m} (kg) means total emissions released from vehicles. It is worth mentioning that 123 *P_iF* is independent of pollutant release rates as well as population density and size, 124 but can be influenced by building configurations, meteorological conditions and 125 126 pollutant source settings etc.

According to the literature [4, 70-71], the population data are categorized into

three subgroups(N=3, Fig. 1a): Children(21.2%), Adults(63.3%) and Elders(15.5%). 128 Moreover, the time activity patterns for each subgroup are divided into four 129 130 microenvironments (M=4, Fig. 1b): indoors at home (j=1), other indoor locations (j=2), in or near vehicles (i=3), and other outdoor locations (i=4). Table 1 lists activity time 131 patterns and breathing rates in each subgroup for indoors at home(j=1). In this study, 132 all present building models are supposed to residential type and only j=1 (indoors at 133 home) is adopted to calculate P_{iF} [44, 57]. Especially, the area-averaged P_{iF} of the 134 building wall surface is denoted as wall intake $fraction(\langle P_iF \rangle_W)$. The 135 136 spatially-averaged $P_{i}F$ of the entire building surfaces is represented as building intake fraction ($\langle P | iF \rangle_B$), which means the fraction of total traffic emissions inhaled 137 averagely by each person living in this roadside building. 138

139

140 **2.2** Photostationary state defect (d_{ps} , unit: %)

141 Referring to previous researches[59-60], the photostationary state defect (d_{ps}) is 142 an effective indicator to measure the departure degree from photochemical 143 equilibrium and can be expressed in the following form:

144
$$d_{ps} = \left(\frac{k_I [\text{NO}][\text{O}_3]}{J_{NO_2} [\text{NO}_2]} - 1\right) \times 100$$
(2)

here, k_I [NO][O₃] and J_{NO_2} [NO₂] represent the depletion and generation rates of ozone. d_{ps} is positive value if ozone depletion rate greater than its formation rate, and vice versa. Especially, d_{ps} equals zero when chemical reactions reach equilibrium state.

150 **3.** CFD setups and case descriptions

151 **3.1 Numerical approaches**

With advances in computer technologies, CFD as a powerful modelling tool has 152 153 been widely employed to reproduce turbulent flow structure as well as to predict pollutant dispersion and transport in urban districts. Though large eddy simulations 154 (LES) have been confirmed to be more accurate in predicting turbulence than 155 Reynolds-Averaged Navier-Stokes (RANS) models[27, 73-75], RANS approaches are 156 still extensively used, since LES models require expensive computational expenses 157 and have challenges in selecting sub-grid scale models and specifying appropriate 158 159 boundary conditions. Moreover, among the RANS models (e.g. various k- ε and k- ω models), the standard k- ε model shows good agreements with experimental data and 160 has been widely adopted [7-8, 37-38, 45, 57-58, 76-79], although it has limitations in 161 predicting turbulent kinetic energy in strong-wind regions. Therefore, by considering 162 model performance and computational loads, the standard k- ε model is selected to 163 solve the steady-state isothermal flow field. The governing equations for the flow and 164 turbulent quantities implemented are as below[80]: 165

166 The mass continuity equation:

$$\frac{\partial \overline{u}_j}{\partial x_i} = 0 \tag{3}$$

168 The momentum equation:

169
$$\overline{u}_{j}\frac{\partial\overline{u}_{i}}{\partial x_{j}} = -\frac{1}{\rho}\frac{\partial\overline{p}}{\partial x_{i}} + \frac{\partial}{\partial x_{j}}\left(\nu\frac{\partial\overline{u}_{i}}{\partial x_{j}} - \overline{u_{i}'u_{j}'}\right)$$
(4)

170 The transport equations of turbulent kinetic energy (*k*) and its dissipation rate (ε):

171
$$\overline{u}_{i}\frac{\partial k}{\partial x_{i}} = \frac{\partial}{\partial x_{i}}\left[\left(\nu + \frac{\nu_{t}}{\sigma_{k}}\right)\frac{\partial k}{\partial x_{i}}\right] + P_{k} - \varepsilon$$
(5)

172
$$\overline{u}_{i}\frac{\partial\varepsilon}{\partial x_{i}} = \frac{\partial}{\partial x_{i}}\left[\left(\nu + \frac{\nu_{i}}{\sigma_{\varepsilon}}\right)\frac{\partial\varepsilon}{\partial x_{i}}\right] + C_{\varepsilon I}\frac{\varepsilon}{k}P_{k} - C_{\varepsilon 2}\frac{\varepsilon^{2}}{k}$$
(6)

173 where, \overline{u}_j is mean velocity components ($\overline{u}_j = \overline{u}, \overline{v}, \overline{w}$ as j=1, 2, 3); v and $v_t = C_\mu \frac{k^2}{\varepsilon}$ 174 ($C_\mu = 0.09$) represent the kinematic viscosity and the eddy viscosity, respectively; the 175 Reynolds stress tensor $-\overline{u_i'u_j'}$ and the turbulence production term P_k are defined as:

176
$$-\overline{u_i'u_j'} = v_t \left(\frac{\partial \bar{u}_i}{\partial x_j} + \frac{\partial \bar{u}_j}{\partial x_i}\right) - \frac{2}{3}k\delta_{ij}$$
(7)

177
$$P_{k} = v_{t} \times \frac{\partial \overline{u}_{i}}{\partial x_{j}} \left(\frac{\partial \overline{u}_{i}}{\partial x_{j}} + \frac{\partial \overline{u}_{j}}{\partial x_{i}} \right)$$
(8)

178 Note that δ_{ij} is the Kronecker delta whose value is 1 when i=j and otherwise is 0.

179 The NO_x - O_3 simple photochemical mechanism is described as follows[59-60]:

180
$$NO_2 + hv \rightarrow NO + O$$
, J_{NO_2} (9a)

181
$$O + O_2 + M \rightarrow O_3 + M$$
, k_2 (9b)

$$182 O_3 + NO \rightarrow NO_2 + O_2 , k_1 (9c)$$

Here, *M* denotes a third molecule, for example O_2 or N_2 , to absorb excess energy and stabilize O_3 molecule formed; J_{NO_2} , k_2 and k_1 represent rate constant for each reaction, respectively. Since the oxygen atom (O) is highly reactive, it combines rapidly with O_2 once O is produced from NO_2 photolysis. This is so-called pseudo-steady-state approximation that the depletion and production rates of O_3 are 188 nearly equal[81]:

189 $k_2[O][O_2][M] = J_{NO_2}[NO_2]$ (10)

190 Therefore, Eq. (9a) is the rate control step for O_3 formation. Based on the above 191 assumption, the transport equations for reactive pollutants can be defined as[59-60]:

192
$$\overline{u}_{j}\frac{\partial[\text{NO}]}{\partial x_{j}} = D_{m}\frac{\partial^{2}[\text{NO}]}{\partial x_{j}\partial x_{j}} + \frac{\partial}{\partial x_{j}}\left(D_{e}\frac{\partial[\text{NO}]}{\partial x_{j}}\right) + J_{NO_{2}}[\text{NO}_{2}] - k_{I}[\text{O}_{3}][\text{NO}] + S_{NO}$$
(11a)

193
$$\overline{u}_{j} \frac{\partial [\text{NO}_{2}]}{\partial x_{j}} = D_{m} \frac{\partial^{2} [\text{NO}_{2}]}{\partial x_{j} \partial x_{j}} + \frac{\partial}{\partial x_{j}} \left(D_{e} \frac{\partial [\text{NO}_{2}]}{\partial x_{j}} \right) - J_{NO_{2}} [\text{NO}_{2}] + k_{I} [\text{O}_{3}] [\text{NO}] + S_{NO_{2}}$$
(11b)

194
$$\overline{u}_{j} \frac{\partial [O_{3}]}{\partial x_{j}} = D_{m} \frac{\partial^{2}[O_{3}]}{\partial x_{j} \partial x_{j}} + \frac{\partial}{\partial x_{j}} \left(D_{e} \frac{\partial [O_{3}]}{\partial x_{j}} \right) + J_{NO_{2}} [NO_{2}] - k_{I} [O_{3}] [NO]$$
(11c)

where, D_m and D_e are the molecular and eddy diffusivity; the third terms on the right-hand side of Eq. (11) represent chemical reaction term; S_{NO} and S_{NO_2} denote the source terms of NO and NO₂; the Schmidt number $Sc_i = v_t/D_e$ is specified as 0.7[33, 39, 46, 48, 78]. Furthermore, the photolysis rate J_{NO_2} and rate constant k_I are calculated by [59-60]:

200
$$J_{NO_2} = 8.14 \times 10^{-3} \left\{ 0.97694 + 8.3700 \times 10^{-4} (T - 273.15) + 4.5173 \times 10^{-6} \times (T - 273.15)^2 \right\}$$
 (12a
201)

202
$$k_1 = 4.405 \times 10^{-2} exp(-\frac{1370}{T})$$
 (12b)

Here, *T* is temperature in K; the units of J_{NO_2} and k_I are s⁻¹ and ppbv⁻¹s⁻¹. Additionally, the temperature-dependent rate constant is not considered in isothermal flow of this study, thus J_{NO_2} and k_I are 8.1×10^{-3} s⁻¹ and 4.450×10^{-4} ppbv⁻¹s⁻¹ by assuming reactive pollutants undergo chemical processes under the isothermal condition with a fixed *T* of 298.15K. All governing equations(Eqs. (3-6, 11)) are discretized by a finite volume method (FVM) with the second-order upwind scheme. The SIMPLE algorithm is employed for pressure and velocity coupling. The under-relaxation factors for pressure, momentum, k and ε terms are specified as 0.3, 0.7, 0.8 and 0.8. Numerical simulation does not stop until the absolute residuals of all variables are less than 10⁻⁶.

213

214 **3.2** Three-dimensional (3D) urban model setups in CFD

The building layouts, as depicted in Fig. 2a, are based on the idealized 3D 215 medium-dense urban clusters (i.e. street aspect ratio H/W=1; building packing density 216 $\lambda_p = \lambda_t = 0.25$). To better illustrate model configurations, x, y and z are described as the 217 218 stream-wise, span-wise and vertical directions, respectively. The cubic building 219 models(H=B=30m) with a uniform spacing($W_s=W_m=30m$) are constructed in x and y directions. Moreover, the approaching wind is parallel to the main streets(*x* direction) 220 and perpendicular to the secondary streets(y direction). x/H=0 means the cross section 221 in windward street opening and y/H=0 denotes the central cross section of the main 222 street (Fig. 2c). 223

As verified by the literature[38-40, 82-83], the airflow in the middle column is hardly affected by lateral boundaries if the lateral width of the urban model is sufficiently large. Hence, only half of the middle column with two lateral symmetry boundaries are considered during CFD simulations to reduce computational efforts (Fig. 2b). Following the CFD guidelines[84-86], the distances between urban

boundaries and the domain top, domain inlet, domain outlet are 9H, 6.7H and 32.3H, 229 respectively. Furthermore, the zero normal gradient boundary condition is adopted at 230 231 two lateral boundaries and the domain top (i.e. symmetry) and the domain outlet (i.e. outflow). Several recent studies [5-6, 8-9, 44, 57-58] reported that, pollutant 232 concentration on the wall surfaces of near-road buildings can be treated as indoor 233 concentration originated from outdoor pollutants since the indoor/outdoor (I/O) ratio 234 of pollutant concentration is nearly one for naturally-ventilated buildings[5, 72]. 235 Therefore, by assuming all present building models are naturally-ventilated type, the 236 237 flow and pollutant dispersion within indoor space of buildings are not taken into account to reduce the grid numbers and computational costs in CFD simulations. The 238 239 literature have applied such technique to effectively quantify vehicular pollutant 240 exposure in 2D street canyons or 3D urban models[5-6, 8-9, 44, 57-58].

For the domain inlet, the power-law time-averaged velocity profile $U_0(z)$ in the upstream free flow is adopted (Eq. 13a)[40-41, 57, 82-83] which is scaled to that in wind tunnel experiments[87]. Base on the CFD guideline[84-86], vertical profiles of k(z) and $\epsilon(z)$ are given by Eqs. (13b-c):

245
$$U_0(z) = U_{ref} \times (z/H)^{0.16}$$
 (13a)

246
$$k(z) = u_*^2 / \sqrt{C_{\mu}}$$
 (13b)

247
$$\varepsilon(z) = C_{\mu}^{3/4} k^{3/2} / (\kappa_{\nu} z)$$
 (13c)

where, U_{ref} is the reference velocity at the building height in upstream free flow ($U_{ref}=3.0$ m/s at z = H); the friction velocity $u_*=0.24$ m/s[40-41, 57, 82-83]; von Karman constant $\kappa_v = 0.41$, $C_{\mu} = 0.09$. Furthermore, vertical profiles of Eq. (13) represent neutral atmospheric boundary layer with a full-scale surface roughness $z_0=0.1m[93]$ and have been adopted in previous studies[40-41, 57, 82-83].

In addition, Fig. 2c demonstrates the overhead and lateral views of mesh 253 distribution for test cases. The minimum grid size of 0.2 m near the wall surfaces and 254 the stretching ratio between adjacent grids of 1.15 (about 3.2 million hexahedral cells) 255 are applied to ensure sufficiently fine grid at the pedestrian level (0-1.5m) and near 256 building surfaces. As the normalized distance y^+ ($y^+ = yu_r/v$) ranges from 30 to 600 at 257 258 most regions of wall surfaces, standard wall function with no slip boundary condition is set on all wall surfaces. According to the literature[13, 40, 88-89], a specific 259 roughness modification is assigned to the upstream and downstream ground to obtain 260 a horizontally homogeneous ABL surrounding urban regions. Especially, the grid 261 independence tests are presented later in subsection 4.1. 262

263

3.3 Pollutant source settings and model description of test cases

Three kinds of pollutant source locations are considered in this study, i.e. the span-wise (y direction) emission sources in the 3^{rd} or 10^{th} secondary street is denoted as S3 and S10 (Fig. 3a-b), and the stream-wise (x direction) emission sources along the main street is represented by Sm (Fig. 3c). The reactive pollutants involved in present photochemistry are NO, NO₂ and O₃. Among them, NO_x is assumed to be emitted from vehicles into the street canyon while background O₃ concentration is

specified at the domain inlet and entrained by approaching wind into urban districts. 271 The background concentrations for NO and NO₂ at the domain inlet are zero in this 272 273 study. Following the literature [59-60], typical automobile emission ratio of NO to $NO_2(R_{NO/NO_2})$ of 10 is adopted. NO and NO₂ are released from the lowest grid cell 274 (z=0-0.2m) at rates of 100 and 10 ppbv s⁻¹, which corresponds to emission strengths 275 of 1.227×10^{-7} and 1.88×10^{-8} kg m⁻³s⁻¹ (*P*=1 atm, *T*=298.15K). For S3 or S10 sources, 276 as the street width (W_s) is 30m, this NO_x emission intensity of 849 µg m⁻¹s⁻¹ is 277 equivalent to a traffic volume of about 6100 vehicles per hour when considering a 278 NO_x emission of 0.5g km⁻¹ per vehicle[59-60]. Particularly, we are mainly concerned 279 with the proportion among NO and NO₂ rather than the assigned emission rate for 280 each vehicular pollutant. 281

Furthermore, four O₃ background concentrations (i.e. $[O_3] = 1, 20, 40$ and 100 ppbv) with $R_{NO/NO_2} = 100:10$ are investigated to study the effects of different $[O_3]$ on reactive dispersion. In addition, Carslaw[90] verified that there is an increasing NO₂/NO_x emission ratio in road traffic emissions referring to observation in real cities. Thus, three emission ratios of NO to NO₂ (i.e. $R_{NO/NO_2} = 100:10, 50:10$ and 100:20) with $[O_3] = 20$ ppbv are considered to examine the impacts of various R_{NO/NO_2} on reactive pollutant dispersion.

Overall, total 11 test cases are described as Case P [Source, R_{NO/NO_2}] or Case R [Source, R_{NO/NO_2} , [O₃]] and summarized in Table 2. Here 'P' means passive dispersion without chemical reactions, 'R' denotes reactive dispersion with NO-NO₂-O₃ chemistry; 'Source' contains three pollutant source locations, i.e. S3, S10 and Sm, respectively; $R_{\text{NO/NO}_2}$ represents the emission ratios of NO to NO₂; [O₃] is background O₃ concentration (mole fraction, unit: ppbv).

295

296 4 Validation study of flow and pollutant dispersion in 3D urban models

297 4.1 Flow validation by wind tunnel data

In this subsection, the performances of various steady *k-ε* models (standard,
RNG and Realizable) with standard wall function are evaluated by wind tunnel data.
Moreover, the grid independence tests are also implemented.

As shown in Fig. 4a-b, idealized 3D urban models in wind tunnel experiment[87] consist of 7 rows and 11 columns of regularly-aligned cubic buildings (H=B=W=15cm). Vertical profiles of velocity components (\bar{u} , \bar{w}) and turbulence kinetic energy (k) are measured at points of V_i (i=1-6), which are central positions in the *i*th secondary street located at y/H=1 and x/H=1.5H, 3.5H, 5.5H, 7.5H, 9.5H, 11.5H respectively (Fig. 4a-b).

In CFD validation study, the similar full-scale model configurations (H=B=W=30m, Fig. 4c) are reconstructed with the scale ratio of 200:1 to wind-tunnel-scale models. Moreover, all CFD setups including computational domains, boundary conditions and convergence criteria in this full-scale CFD validation study are similar with those in subsection 3.2 except that the distance between urban boundaries and domain outlet is 40.3H(Fig. 4c). Based on the building height(H=0.15m or 30m) and reference velocity($U_{ref}=3m/s$), the reference Reynolds number $Re = \frac{U_{ref}H}{v} \approx 30000$ and 6×10^6 for wind-tunnel-scale and full-scale urban models, which are much larger than 11000 satisfying Reynolds number independence requirement. Furthermore, Fig. 4d depicts the coarse, medium and fine grid arrangements with hexahedral cells of about 1, 2 and 3.3 million and the minimum grid size of 0.4m, 0.2m and 0.1m, respectively.

Fig. 5 first shows results of grid independence test at V1(Fig. 5a-b), then depicts 319 vertical profiles of time-averaged stream-wise velocity $\bar{u}(z)$ at Points V1, V4 and V6 320 (Fig. 5c-e), vertical velocity $\overline{w}(z)$ (Fig. 5f) and turbulence kinetic energy k(z) (Fig. 5g) 321 322 at Point V1 between numerical results and experimental data. There is little difference in numerical results between the coarse, medium and fine grid, and thus medium grid 323 is selected for case studies to reduce computational loads. Besides, the standard k- ε 324 model with the medium grid shows better agreements with wind tunnel data than the 325 RNG and Realizable k- ε models. 326

To further quantify the modeling accuracy and reliability of the standard $k-\varepsilon$ 327 model with the medium grid arrangement, several statistical performance metrics are 328 applied, including the mean value, the standard deviation, the factor 2 (FAC2), the 329 normalized mean square error (NMSE), the fraction bias (FB) and the correlation 330 coefficient (R)[91]. Among which, the closer NMSE value to zero, the smaller 331 difference between experiment data and CFD results; FAC2 value larger than one 332 means over-prediction against experiment data while smaller than one represents 333 334 under-prediction; Similarly, the negative and positive FB values denote overestimation and underestimation. Results of $\bar{u}(z)$ at Points V1, V4 and V6 as well 335

as $\overline{w}(z)$ and k(z) at Point V1 are listed in Table 3. Referring to COST Action 732's 336 recommended criteria^[91], a credible CFD model should satisfy the following 337 statistical metrics standards: $0.5 \le FAC2 \le 2$, NMSE ≤ 1.5 and $-0.3 \le FB \le 0.3$. 338 Overall, metrics lie in the recommended criteria, particularly values of $\overline{u}(z)$ meet 339 well. However, values of $\overline{w}(z)$ and k(z) reveal relatively poorer performance than 340 those of $\overline{u}(z)$, which is largely attributed to the limitation of the standard k- ε model. 341 Conclusively, the validation study shows that present CFD methodologies applying 342 standard k- ε model with medium grid have credible numerical accuracy in predicting 343 344 urban turbulent flow and can be employed for further case studies.

345

4.2 Validation of pollutant dispersion by wind tunnel data

Experimental and numerical studies have been extensively performed to 347 investigate passive pollutant dispersion in the idealized street canyons. Unfortunately, 348 there are currently little experimental data to directly validate the present CFD model 349 coupled with chemistry[59-60, 63-64]. However, the reactive pollutants considered in 350 this paper are effectively passive and can be divided into two subsets, i.e. total 351 352 nitrogen oxide ($NO_x=NO+NO_2$) and total oxidant ($OX=NO_2+O_3$), since reactions (Eq. 9) interconvert NO with NO₂, and O₃ with NO₂ but without redundant productions. In 353 addition, the chemical reaction terms cancel out when Eqs. (11a-b) and Eqs. (11b-c) 354 355 are added, which indicates that the transport and dispersion of NO_x and OX can be deemed as passive scalar. Consequently, in this subsection, Standard k- ε model with 356

standard wall function has been implemented and validated against wind tunnel
experiment to evaluate the reliability of numerical simulation in predicting passive
pollutant distribution.

The configurations of the wind tunnel measurement[92], as depicted in Fig.6a-b, 360 are consisted of nine rectangular building models (Lx=27.6cm, Ly=18.4cm, H=8cm) 361 with three rows and three columns (3×3) and uniform street widths (W=8cm, H/W=1). 362 Moreover, tracer $gas(C_2H_6)$ is released from a line source (0.5cm in width, dx and 363 18.4 cm in length, Ls) at a velocity of $w_{source}=0.01$ m/s which is paralleled with y 364 direction and locates in the central street canyon in front of building No.2(Fig. 6a-b). 365 Furthermore, C₂H₆ concentration profiles are measured in the middle of leeward and 366 windward walls near the line source as well as on the central line of roof surface in 367 the building No.2(Fig. 6a-b). 368

In full-scale pollutant dispersion validation study, similar model configurations 369 (Lx=138m, Ly=92m, H=40m, in Fig. 6c) are rebuilt with the scale ratio of 500:1 to 370 wind-tunnel-scale models. CFD setups are similar with subsection 3.2, but the 371 distances between urban boundaries and the domain top, domain side, domain inlet, 372 373 domain outlet are 9H, 5H, 5H and 15H, respectively. The approaching wind is parallel to the main streets(x direction) and perpendicular to the secondary streets(y direction). 374 Furthermore, vertical profiles of stream-wise velocity $\bar{u}(z)$, turbulence kinetic energy 375 k(z) and turbulent dissipation rate $\varepsilon(z)$ fitted by measured data in wind tunnel 376 377 experiments^[92] are adopted at the domain inlet (Fig. 6d-f). In addition, to compare CFD results with measured data, the normalized C_2H_6 concentration K is defined as 378

below:

380
$$K=CHU_{ref}/w_{sourced}x$$
(14)381Here the height of building (*H*), the reference velocity (U_{ref}) and the line source382emission strength (w_{source}, dx) are applied.383As shown in Fig. 7, the agreements of *K* between wind tunnel data and CFD384results are well confirming the standard k - ε model has sufficient modeling accuracy in385predicting passive pollutant dispersion within 3D urban district models.386538753885.1 Flow patterns in 3D urban district models389Fig. 8a depicts velocity distribution in the plane of $z=1.5m$ (the pedestrian level).390Obviously, the flow adjustment process can be observed through the entire building391clusters, in which wind speed decreases toward downstream from Street 1 to Street 6,392and then reaches a comparatively flow equilibrium from Street 7 to Street 18. Fig.3938b-c further depict velocity magnitude and 2D streamlines in the plane of $y=30m$ (the394center plane of the target street canyon) and $z=1.5m$ for Street 3 and Street 10.395Moreover, the corresponding 3D streamlines are displayed in Fig. 8d. For both street396units, 3D downward helical vortices are produced inside the secondary streets (Fig.3978b-d). The lateral flow direction near street ground (i.e. $z=1.5m$) are from the398secondary streets to the main streets and from the downwind building (No.4 and 11)

to upwind building (No. 3 and 10) (Fig. 8c). In addition, Street 3 apparently
experiences greater wind speed than Street 10.

In particular, the following analysis (subsection 5.3) emphasizes more on reactive pollutant dispersion in the fully-developed region (e.g. Street 10) than the flow-adjust region (e.g. Street 3).

404

405 5.2 Impacts of source locations (S3, S10 and Sm) on reactive pollutant dispersion

This subsection considers the impacts of source locations (i.e. S3, S10 and Sm in 406 Fig. 3) on reactive pollutant dispersion(with chemical reactions, R-type) under the 407 specific pollutant proportion (i.e. $R_{NO/NO_2}=100:10$, [O₃]=20ppbv). Because the present 408 photochemical mechanism contains the interconversion of nitrogen oxides(i.e. 409 $NO_x=NO+NO_2$) and oxidants (i.e. $OX=NO_2+O_3$), The following discussions 410 (subsection 5.2 and 5.3) concentrate more on NO_2 to simplify analysis. Moreover, 411 passive dispersion (without chemical reactions, P-type) are also presented to 412 investigate the sole role of turbulent mixing. 413

414

415 **5.2.1 NO₂ concentration distribution in 3D urban-like models**

Fig. 9a-d exhibit NO₂ concentration between P-type(passive, left) and R-type (reactive, right) cases in y=30m and z=1.5m for Street 3 and 10 fixed with span-wise sources(i.e. S3 and S10). Moreover, Fig. 9e compares NO₂ concentration at the

pedestrian level between passive and reactive cases with NO_x sources along main 419 streets (i.e. Sm). For passive dispersion with S3 or S10 sources (P-type), due to source 420 421 emissions and turbulent transports, NO₂ concentration near the upwind building (No.3 and 10) is higher than that near the downwind building(Fig. 9a-b). Furthermore, a 422 large amount of NO₂ pollutants accumulate in the intersection of main street and 423 secondary street(Fig. 9c-d). In an overall view, NO2 concentration in Street 10 is 424 slightly higher than Street 3. For P-type case with Sm sources(Fig. 9e), NO₂ 425 concentration first rises toward downstream streets, then reaches an approximate 426 427 equilibrium from Street 7 to 18. Such findings are similar with the flow adjustment as discussed in subsection 5.1. 428

With chemical reactions, as verified by Fig. 9, NO₂ concentration in R-type cases
considerably exceeds that in P-type cases. Oppositely, passive NO concentration is
higher than that in R-type cases.

432

433 5.2.2 d_{ps} distribution in Street 3 and 10

Fig. 10 shows d_{ps} distribution in y=30m and z=1.5m in local target streets with S3 and S10 sources. Here, the distribution of photostationary state defect(d_{ps}) is emphasized below the roof level (z/H<1) and toward downstream domains (i.e. x/H>5and x/H>19 for Street 3 and 10, respectively).

As introduced in subsection 2.2, smaller d_{ps} value represents the less departure degree from photochemical equilibrium. In the centre plane of secondary streets

(y=30m, Fig. 10a), the local small d_{ps} values emerge near the roof of the upwind 440 building (No.3 and 10) while the large d_{ps} values appear near the roof of the 441 442 downwind building (No.4 and 11) and the ground level close to NO_x emissions. At the pedestrian level(Fig. 10b), downstream areas of the main streets (x/H>6 and x/H>20) 443 experience small d_{ps} values while the junction regions of the main street and 444 secondary street near NO_x source locations obtain large d_{ps} values, particularly in 445 Street 3. In summary, d_{ps} value is usually smaller in regions with weaker wind and 446 turbulence, where reactive pollutants have more time to mix and react. 447

448

449 5.2.3 Concentration, $\langle P_iF \rangle_W$ and $\langle P_iF \rangle_B$ on building wall surfaces

Based on spatial mean concentration at the entire building surfaces, we calculate wall intake fraction($\langle P_iF \rangle_W$) and building intake fraction($\langle P_iF \rangle_B$) to analyze overall vehicular pollutant(NO_x) exposure in near-road buildings. Especially, 1ppm represents 1 mg inhaled averagely by each person living in the near-road building if 1kg pollutants emitted out.

Fig. 11a-b first compare NO₂ concentration on the leeward and windward walls between P-type and R-type cases in target street units with S3 and S10 sources. No matter with or without chemical reactions, NO₂ concentrations on the leeward wall are always higher than those on the windward wall. Once NO_x -O₃ photochemical reactions are conducted, an increase of NO₂ concentration emerges on the upwind and downwind walls. Such results are similar with the aforementioned discussion in 461 subsection 5.2.1.

Then, Table 4 lists $\langle P | iF \rangle_W$ of NO and NO₂ at leeward and windward walls 462 adjoining target street in cases with S3 and S10 (positions as described in Fig. 3a-b). 463 Obviously, in both P-type and R-type cases, $\langle P_iF \rangle_W$ of NO_x at leeward wall(Table 4, 464 row 1-4, column 2-3) are greater than those at windward wall(Table 4, row1-4, 465 column 4-5). Regarding P-type cases as the references, $\langle P_iF \rangle_W$ of NO₂ in R-type 466 cases rise nearly 90%-160%, i.e. 0.660 to 1.372ppm and 0.853 to 1.643ppm at 467 leeward wall for S3 and S10(Table 4, column 2), 0.180 to 0.473ppm and 0.230 to 468 0.610ppm at windward wall for S3 and S10(Table 4, column 4); while $\langle P | iF \rangle_W$ of 469 NO reduces about 9%-16%, i.e. 0.660 to 0.588ppm and 0.853 to 0.775ppm at leeward 470 wall for S3 and S10(Table 4, column 3), 0.180 to 0.151ppm and 0.230 to 0.193ppm at 471 windward wall for S3 and S10(Table 4, column 5). Because NO_x emission ratio 472 released from vehicles is $R_{\rm NO/NO_2}=10$, the present photochemical processes lead to a 473 significant increase in NO₂ exposure and a slight decrease in NO exposure. 474

Furthermore, NO₂ concentrations on the entire building wall surfaces with Sm 475 sources are presented in Fig. 11c. Both P-type and R-type cases experience the NO₂ 476 concentration adjustment processes toward downstream buildings. To quantify NO_x 477 exposure adjustments in P-type and R-type cases with various sources (S3, S10 and 478 Sm), the horizontal profiles of building intake fraction $\langle P_iF \rangle_B$ of NO and NO₂ are 479 shown in Fig. 12. It is found that $\langle P_iF \rangle_B$ with S3 or S10 sources descends 480 481 exponentially toward downstream buildings(Fig. 12a-b), instead, $\langle P | iF \rangle_{\rm B}$ with Sm sources first ascends quickly from building No.1 to 8, then reaches an approximate 482

equilibrium(Fig. 12c). Besides, R-type cases experience larger NO₂ exposure than P-type cases, i.e. 0.420-0.108ppm against 0.135-0.020ppm for S3(Fig. 12a), 0.605-0.160ppm against 0.207-0.030ppm for S10(Fig. 12b), 0.005-0.090ppm against 0.002-0.049ppm for Sm(Fig. 12c). Oppositely, NO exposure in P-type cases are greater than R-type cases, i.e. 0.135-0.020ppm than 0.106-0.010ppm for S3(Fig. 12a), 0.207-0.030ppm than 0.168-0.017ppm for S10(Fig. 12b) and 0.002-0.049ppm than 0.002-0.045ppm for Sm(Fig. 12c).

In addition, the decay function expressed in $\langle P_iF_n \rangle_{\mathbf{B}} = a \times \langle P_iF_t \rangle_{\mathbf{B}} \times e^{(t-n)/b}$ is 490 employed to further quantify the $\langle P_iF \rangle_B$ decay processes from target building unit (t 491 =4 or 11) toward downstream building (n=t to 21) in S3 and S10 cases. Note that, 492 smaller decay factor b means relatively sharper descending processes of $\langle P_iF \rangle_B$ 493 curves. Table 5 summarizes the $\langle P_iF_{t}\rangle_B$ of building "No.t" and the exponential 494 decay factors b in S3 and S10 cases. Obviously, compared with those in P-type case 495 (b=7.46 and 3.96 in Table 5, row 1 and 3), R-type cases with S3 and S10 obtain larger 496 b for NO₂ (10.80 and 6.84 in Table 5, row 2 and 4) and smaller b for NO (5.92 and 497 2.91). Moreover, $\langle P_iF \rangle_B$ curves in S10 cases(Table 5, row 3-4) decline more 498 sharply toward downstream regions than S3 cases(Table 5, row 1-2). Particularly, as 499 shown in Fig. 12c, $\langle P_iF_t \rangle_B$ in Sm cases are calculated by the mean $\langle P_iF \rangle_B$ from 500 building No.9 to 21, i.e. 0.048ppm in P-type case, 0.044ppm and 0.088ppm in R-type 501 case for NO and NO₂. 502

In summary, with $R_{NO/NO_2} = 100:10$ and $[O_3] = 20$ ppbv, the present NO_x-O₃ titration interactions result in the production of NO₂ and depletion of O₃ and NO. By focusing on $\langle P_iF_i \rangle_B$ values, NO₂ exposure in R-type cases are greater than P-type cases about 3.1 times for S3, 2.9 times for S10 and 1.8 times for Sm while NO exposure in R-type cases are nearly 21%, 19% and 8% smaller than P-type cases for S3, S10 and Sm, respectively.

509

510 **5.3 Impacts of reactant proportions** (NO:NO₂:O₃) **on reactive pollutant dispersion**

In this subsection, based on S10 sources, we discuss the effects of different reactant proportions (NO:NO₂:O₃) on the interaction of turbulent mixing and photochemical processes in urban districts. Additionally, P_iF and $\langle P_iF \rangle$ are independent on source emission strength in passive pollutant dispersion, therefore Case P[S10,100:10] is treated as the reference case to compare with the cases with other reactant proportions.

517

518 5.3.1 Impacts of O₃ background concentration ([O₃])

519 With the same emission ratio of NO to NO₂ (i.e. $R_{\text{NO/NO}_2}$ =100:10), the impacts of 520 four O₃ background concentrations(i.e. [O₃] =1, 20, 40 and 100ppbv) on reactive 521 pollutant dispersion are examined.

It is apparent that the formation of NO by photolyzing NO₂ is slightly dominant in photochemistry when $[O_3]$ is 1ppbv. In contrast to the reference values(0.853 and 0.230ppm in Table 4, row 3), $\langle P_iF \rangle_W$ of NO₂ at the leeward and windward walls slightly decrease(i.e. 0.816 and 0.211ppm in Table 4, row 5) while $\langle P_iF \rangle_W$ of NO increase a little(i.e. 0.858 and 0.233ppm). Besides, such phenomenon is distinctly observed in $\langle P_iF \rangle_B$ curves(Fig. 13) between the reference case and Case R [S10,100:10,1], i.e. 0.207-0.030ppm against 0.187-0.020ppm for NO₂(Fig. 13a) and 0.207-0.030ppm against 0.209-0.031ppm for NO(Fig. 13b).

However, if [O₃] rises from 20ppbv to 40 and 100ppbv, more NO is oxidized by 530 O₃ to generate NO₂. Based on the reference values (0.853 and 0.230ppm in Table 4, 531 row 3), $\langle P_iF \rangle_W$ of NO₂ become about 1.9-5.2 times greater (1.643, 2.454 and 532 533 4.442ppm in Table 4, column 2) at leeward wall and 2.6-6.7 times larger (0.610, 0.944 and 1.534ppm in Table 4, column 4) at windward wall; while $\langle P_iF \rangle_W$ of NO 534 reduces approximately 9%-42% (0.775, 0.694 and 0.495ppm in Table 4, column 3) at 535 leeward wall and 16%-57% (0.193, 0.159 and 0.100ppm in Table 4, column 5) at 536 windward wall. Furthermore, Fig. 13 and Table 5 display the corresponding $\langle P_{i}F \rangle_{\rm B}$ 537 curves, $\langle P_i F_t \rangle_B$ values and decay factors b under different [O₃]. As depicted in Fig. 538 13, compared with Case R[S10,100:10,20], Case R[S10,100:10,100] and R 539 [S10,100:10,40] obviously attain much larger $\langle P_iF \rangle_B$ of NO₂ (1.573-0.275 and 540 541 0.952-0.218ppm in Fig. 13a) and smaller $\langle P_iF \rangle_B$ of NO (0.071-0.005 and 542 0.133-0.011ppm in Fig. 13b). Additionally, both decay factors of NO₂ and NO are smaller (b=6.02 and 2.33 for [O₃]=40ppbv, b=4.74 and 1.87 for [O₃]=100ppbv in 543 Table 5, row 6-7) than those in Case R[S10,100:10,20] (b=6.84 and 2.91 in Table 5, 544 row 4), which implies higher $[O_3]$ induces the quicker decay of $\langle P_iF \rangle_B$ curves for 545 NO_x with S10 sources toward downstream building units. By concentrating on 546

 $< P_i F_t >_B$, NO₂ exposure in R-type cases surpass the reference case nearly 2.9, 4.6 and 7.6 times for $[O_3]=20$, 40 and 100ppbv, respectively(Table 5, column 2). Correspondingly, $< P_i F_t >_B$ of NO in these $[O_3]$ cases are about 19%, 36% and 66% smaller than the reference values, respectively(Table 5, column 4). It clearly indicates that increasing $[O_3]$ would aggravate NO₂ exposure within urban clusters but is conductive to the mitigation of NO exposure.

553

554 5.3.2 Effects of emission ratio of NO to $NO_2(R_{NO/NO_2})$

The effects of source emission ratios ($R_{\text{NO/NO}_2}$ =100:10, 50:10 and 100:20) on reactive pollutant dispersion are investigated with the same [O₃] of 20ppbv.

It is shown that decreasing NO or increasing NO₂ emissions based on the 557 reference case can mildly change the fraction of NO converting into NO₂. For 558 example, reducing $R_{\text{NO/NO}2}$ from 100:10 to 50:10 and 100:20, $\langle P_iF \rangle_W$ of NO 559 varies from 0.775 to 0.744 and 0.784ppm at the leeward wall(Table 4, column 3), and 560 from 0.193 to 0.186 and 0.197ppm at windward wall(Table 4, column 5). Furthermore, 561 $\langle P_iF \rangle_W$ of NO₂ drops from 1.643 to 1.406 and 1.205ppm at the leeward wall(Table 562 4, column 1), and from 0.610 to 0.453 and 0.400ppm at windward wall(Table 4, 563 column 4). In addition, Fig. 14 presents $\langle P_iF \rangle_B$ curves of NO₂ and NO in P-type and 564 R-type cases with three NO-NO2 emission ratios. Obviously, photochemical reactions 565 in these R-type cases are still dominated by the depletion of O₃ with NO to produce 566 NO₂. 567

568	As displayed in Fig. 14a and Table 5, Case R[S10,50:10,20] and R
569	[S10,100:20,20] obtain smaller $\langle P_iF \rangle_B$ and decay factor of NO ₂ (i.e. 0.446-0.091
570	ppm, <i>b</i> =5.41 and 0.384-0.088ppm, <i>b</i> =6.00) than those in Case R[S10,100:10,20] (i.e.
571	0.605-0.160ppm, b=6.84). In contrast to Case R[S10,100:10,20], $\langle P_iF_i \rangle_B$ of NO in
572	Case R[S10,50:10,20] and R[S10,100:20,20] reduce nearly 26% and 37%. However,
573	$\langle P_iF \rangle_B$ curves and decay factors b of NO are quite close between three R-type cases,
574	i.e. 0.172-0.018, 0.168-0.018 and 0.160-0.017ppm; b=3.05, 2.91 and 3.11(Fig. 14b
575	and Table 5). It is confirmed that the decrement of $R_{\text{NO/NO}_2}$ (from 10 to 5) can partly
576	offset NO ₂ exposure but have much less impacts on NO exposure.

577 Overall, the NO_x-O₃ photochemical processes dependent on the initial proportion 578 of reactive pollutants are toward satisfying the photostationary state relationship 579 (i.e. k_I [NO][O₃]= J_{NO_2} [NO₂]).

580

581 **5.4 Limitations and future work**

Since the 3D urban district models, photochemical reactions and meteorological 582 conditions adopted in this study are fairly simplified, the present exposure results may 583 change if more realistic factors are taken into account, such as more realistic urban 584 configurations(e.g. with variations of building height and street width), more 585 complicated chemical mechanisms(e.g. $VOCs-NO_x-O_3$) and 586 more realistic atmospheric conditions etc. It is worth mentioning that the chemical processes 587 dependent on reaction rates are highly associated with the reactive pollutant 588

589 concentration and ambient air temperature. Moreover, realistic atmospheric conditions 590 include the unsteady temporal and spatial variations of wind speed and direction as 591 well as various atmospheric stabilities and solar radiation conditions. Thus, further 592 unsteady CFD simulations will be performed to examine the integrated impacts of 593 urban turbulence and solar radiation on reactive pollutant dispersion in 3D urban 594 districts.

595

596 6 Conclusions

Urban residents in near-road buildings commonly suffer from high exposure risk 597 of vehicular pollutants in which NO_x(NO and NO₂) act as primary reactive pollutants. 598 599 With satisfactory full-scale CFD validation of flow and pollutant dispersion by experimental data, this study first focuses on the impact of turbulent transport 600 combined with NO_x-O₃ photochemical reactions on reactive pollutant dispersion in 601 neighborhood-scale(21-row, ~1km) three-dimensional(3D) medium-dense urban 602 clusters (*H*/*W*=1, $\lambda_p = \lambda_f = 0.25$). Ground-level emission sources of NO and NO₂ are 603 considered in the presence of background O₃. The approaching wind is parallel to the 604 605 main streets and perpendicular to the secondary streets. As a start, the influences of various source locations and reactant proportions(NO:NO₂:O₃) on pollutant dispersion 606 are investigated under neutral meteorological condition. Personal intake fraction $P_{i}F_{i}$, 607 608 its spatially-averaged values for a building wall($\langle P_iF \rangle_W$) and the entire building surfaces (i.e. building intake fraction $\langle P_iF \rangle_B$) are adopted to quantify pollutant 609

exposure with and without NO-NO₂-O₃ reactions(i.e. reactive and passive).

611 Some meaningful findings are summarized as below:

1) There are flow adjustment processes coupling turbulent mixing and chemical 612 reactions through urban building clusters(Street 1 to Street 20 toward 613 downstream). With span-wise sources, the secondary street of Street 10 located in the 614 fully-developed region(i.e. S10 case) experiences weaker wind and subsequently 615 greater $\langle P_iF \rangle_B$ than the secondary Street 3 located in the upstream flow-adjustment 616 region (i.e. S3 case). Consequently, in contrast to S3 case, photostationary state defect 617 (d_{ps}) is smaller in S10 case since reactive pollutants have more time to mix and react 618 in Street 10. 619

620 2) With source emission ratios of NO to NO₂ of $10(R_{NO/NO2}=100:10)$ and background O₃ concentration of 20ppbv([O₃]=20ppbv), NO-NO₂-O₃ photochemistry 621 leads to production of NO₂ and depletion of O₃ and NO, inducing a significant 622 increase in NO₂ exposure and a slight decrease in NO exposure when compared to 623 corresponding passive dispersion which only considers the sole role of turbulent 624 transport. With span-wise pollutant sources, 3D downward helical flows transport 625 more NO_x to the leeward side, inducing much greater leeward-side $\langle P_iF \rangle_W$ than the 626 windward-side. Moreover, by defining exponential decay function expressed 627 in $\langle P_iF_n \rangle_B = a \times \langle P_iF_t \rangle_B \times e^{(t-n)/b}$, it is found that $\langle P_iF \rangle_B$ descends exponentially 628 629 from target building ($\langle P_iF_t \rangle_B=0.135$ ppm or 0.207ppm, t=4 or 11 for S3 or S10) to downstream buildings (n=t to 21). Especially, $\langle P_iF \rangle_B$ curves decline more sharply 630

from Street 10 toward downstream than that from Street 3. However, if stream-wise sources fixed along the main streets, $\langle P_iF \rangle_B$ first ascends quickly from building No.1 to 8, then reaches approximate equilibrium values of $\langle P_iF \rangle_B$ = 0.046-0.049ppm.

3) Furthermore, the O_3 background concentration ($[O_3]$) and source emission 635 ratios of NO to NO₂($R_{\rm NO/NO_2}$) are confirmed as key factors on NO_x-O₃ reactive 636 dispersion. The formation of NO by photolyzing NO₂ is slightly dominant in 637 photochemistry when $[O_3]$ is 1ppby. However, if $[O_3]$ rises from 20ppby to 40 and 638 639 100ppbv, more NO is oxidized by O_3 to generate NO₂, which would aggravate NO₂ exposure within urban clusters but is conductive to the mitigation of NO exposure. 640 Under [O₃] of 20ppbv, results show that the decrement of $R_{\text{NO/NO}_2}$ from 10 to 5 can 641 partly offset NO₂ exposure but have much less impacts on NO exposure. 642

Although further investigations are still required to provide practical guidelines, this paper is one of the first attempts to quantify how reactive pollutant source locations and reactant proportions influence reactive pollutant exposure in 3D urban districts, which can present meaningful references for urban planning. The effective methodologies are proposed for reactive pollutant exposure assessment in more complicated urban districts with various meteorological conditions and chemical mechanisms.

650

651 Acknowledgements

This study was financially supported by National Key R&D Program of China 652 [2016YFC0202206, 2016YFC0202205 and 2016YFC0202204], the National Science 653 654 Fund for Distinguished Young Scholars (No. 41425020) and National Natural Science Foundation--Outstanding Youth Foundation (No. 41622502), STINT 655 (dnr CH2017-7271) and the National Natural Science Foundation of China (No. 656 51811530017 and 41875015). 657

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Figure list:
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- Fig.1 (a) The population census data of Hong Kong [4]; (b) Time activity patterns for 938 each subgroup in four microenvironments [70]. 939
- 940 Fig.2 (a) Idealized 3D urban district models $(H/W=1, \lambda_p=\lambda_f=0.25);$ (b) Computational domains and boundary conditions for test cases; (c) The overhead 941 and lateral views of mesh distribution. 942

943 Fig.3 Model setups of vehicular emission sources: (a) S3, (b) S10 and (c) Sm.

- Fig.4 (a-b) The side and top views of measured points and model configurations in 944 wind tunnel experiment [87]; (c) Computational domains and boundary 945 conditions in CFD validation study; (d) The mesh arrangements in grid 946 sensitivity test. 947
- 948 Fig.5 Results of grid independence test: (a) $\overline{u}(z)$, (b) $\overline{w}(z)$ at Point V1; Comparison of vertical profiles between wind tunnel data and CFD results: (c-e) $\bar{u}(z)$ at Points 949 V1, V4 and V6, respectively; (f-g) $\overline{w}(z)$ and k(z) at Point V1. 950

951	Fig.6 (a-b) The lateral and overhead views of experiment settings in concentration
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953	profiles of stream-wise velocity, turbulence kinetic energy and turbulent
954	dissipation rate in domain inlet.
955	Fig.7 Comparison of K between wind tunnel data and CFD results applying standard
956	k - ε model: (a) on the leeward and windward walls; (b) on the central line of roof
957	surface.
958	Fig.8 (a) Velocity distribution in the pedestrian level of $z=1.5m$; (b-c) Velocity
959	magnitude and 2D streamlines for Street 3 (left) and 10 (right) in the plane of
960	y=30m and $z=1.5m$, respectively; (d) 3D streamlines in Street 3 and 10.
961	Fig.9 NO ₂ concentration distribution between P-type (left or above) and R-type (right
962	or below) cases: (a-b) in the plane of <i>y</i> =30m with S3 and S10, respectively; (c-d)
963	in the pedestrian level of $z=1.5m$ with S3 and S10, respectively; (e) in the
964	pedestrian level of $z=1.5$ m with Sm.
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966	y=30m and $z=1.5m$, respectively.
967	Fig. 11 NO ₂ concentration between P-type (left or above) and R-type (right or below)
968	cases: (a-b) on the leeward and windward walls with S3 and S10, respectively; (c)

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972	Fig. 13 $\langle P_iF \rangle_B$ curves of (a) NO ₂ and (b) NO in P-type and R-type cases under four
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975	kinds of NO-NO ₂ emission ratios.
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977	Table list:
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979	Table 2 Summary of all test cases
980	Table 3 Comparison between wind tunnel experiment and CFD simulation
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982	street in S3 and S10 cases
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