### Coupled Eulerian Wall Film-Discrete Phase model for predicting respiratory droplet generation during a coughing event

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### 17 Abstract

18 Infectious respiratory diseases have long been a serious public health issue, with airborne 19 transmission via close person-to-person contact being the main infection route. Coughing 20 episodes are an eruptive source of virus-laden droplets that increase the infection risk of 21susceptible individuals. In this study, the droplet generation process during a coughing event 22 was reproduced using the Eulerian wall film (EWF) model, and the absorption/expulsion of droplets was tracked using the discrete phase model (DPM). A realistic numerical model that 2324included the oral cavity with teeth features and the respiratory system from the throat to the 25 first bifurcation was developed. A coughing flow profile simulated the flow patterns of a single coughing episode. The EWF and DPM models were coupled to predict the droplet formation, 26 27 generation, absorption, and exhalation processes. The results showed that the large droplet number concentration was generated at the beginning of the coughing event, with the peak 28 29 concentration coinciding with the peak cough rate. Analysis of the droplet site of origin showed 30 that large amounts of droplets were generated in the oral cavity and teeth surface, followed by 31 the caudal region of the respiratory system. The size of the expelled droplets was 0.25-24 µm, 32 with the peak concentration at  $4-8 \mu m$ . This study significantly contributes to the realm on the 33 site of origin and localized number concentration of droplets after a coughing episode. It can 34 facilitate studies on infection risk assessment, droplet dispersion, and droplet generation 35 mechanisms from other sneezing or phonation activities.

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37 Keywords: Computational Fluid Dynamics, Eulerian Wall Film Model, Discrete Phase Model,

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- 38 Coughing, Droplet site origin and number concentration.
- 39 40

### 41 I. INTRODUCTION

42Infectious respiratory diseases have long been a significant public health concern. This 43 includes instances of plagues, measles, tuberculosis, influenza, severe acute respiratory syndrome coronavirus (SARS-CoV), Middle East respiratory syndrome coronavirus (MERS-44 45 CoV), and the most recent SARS-CoV-2 (Churchyard et al., 2017; Piret & Boivin, 2021). 46 Person-to-person airborne transmission of respiratory viruses can occur via direct or indirect contact, respiratory droplets, and droplet nuclei transmission (Dhand & Li, 2020; C. C. Wang 4748 et al., 2021). Infected patients can expel pathogen carriers into the ambient environment via 49 respiratory activities such as breathing, talking, phonating, singing, sneezing, or coughing (Wei 50 & Li, 2016; Stadnytskyi et al., 2021). Once in the surrounding environment, viral-laden 51 droplets can remain airborne for an extended period and transported by indoor airflow to the effective breathing zone of residents, and subsequently inhaled at various exposure levels, as 52 53 reported by previous studies that were comprehensively reviewed by Inthavong (Inthavong, 542020). During the expiratory phase, the infection risks are associated with the (1) droplet 55 number concentration, (2) size distribution, (3) content of infectious agents, and (4) 56 performance frequency (Morawska, 2006). Besides, more recent studies underscore the 57 dependence of disease transmission on infective dose threshold of the virus (SeyedAlinaghi et al., 2022) and the viral shedding rate (Widders et al., 2020). Thus, quantitative studies are 58 59 required to determine the pathogen susceptibility targeting each activity.

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The threat of coughing-related infections has gained public attention, being extensively

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61	studied over the past decades owing to its possibility for the eruptive release of many pathogens
62	in a short period (Stadnytskyi et al., 2021). When coughing, the sequent build-up of expiratory
63	flow velocity, reaching a Reynolds number of $10^4$ (Bourouiba et al., 2014), expulses
64	compressed air through the open mouth. Due to the high speed, droplets are produced in the
65	oral cavity due to shear-induced surface-wave instabilities (Wei & Li, 2016; Pöhlker et al.,
66	2021). The stripped droplet parcels follow the air stream and escape to the environment through
67	the open mouth or are re-absorbed into the mucus layer. Virus-laden droplets are generally
68	deposited in the respiratory tract after inhalation (H. Li et al., 2022; C. C. Wang et al., 2021).
69	The mucus clearance process (i.e., coughing) can trigger the re-emittance of deposited virions
70	or the local transmission of progeny viruses shed by infected cells (Schaefer & Lai, 2022).
71	Therefore, the number concentration and size distribution of droplets from coughing have
72	gained significant attention from the scientific community. A review of studies over the past 20
73	years has indicated significant scatter (Yang et al., 2007; Chao et al., 2009; Morawska et al.,
74	2009; Johnson et al., 2011; Lindsley et al., 2012; Zayas et al., 2012), which can be attributed
75	to the heterogeneity of measurement techniques, sampling methods, or intersubject variability.
76	More recently, facilitated by advanced techniques, the realm of measuring ejected cloud
77	characteristics and generated droplet properties from oral activities in an indoor environment
78	has been embarked upon to deliver a comprehensive understanding during the pandemic epoch

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79	(Wang et al., 2020; Archer et al., 2022; Harrison et al., 2023; Bahramian & Ahmadi, 2023).
80	Drawing upon the properties of generated droplets from experimental studies, computer-aided
81	approaches have been expanded to explore further the deleterious effects of these droplets in
82	the context of disease transmission. Among them, Computational Fluid Dynamics (CFD) has
83	been universally adopted for the numerical investigation of the contagion risk of coughing.
84	Most studies have investigated the cough-jet stream characteristics and fate of expiratory
85	droplets in an enclosed environment (H. Li et al., 2021; Payri et al., 2021; Nie et al., 2022;
86	Aljabair et al., 2023; Nishandar et al., 2023), or the distance-based exposure risks between
87	residents (Calmet et al., 2021; Mariam et al., 2021; Hossain et al., 2023; X. Li et al., 2023). As
88	can be seen, up to date, both experimental and numerical endeavors have extensively advanced
89	our understanding of droplet size distribution and their behavior under varying microclimatic
90	conditions within an enclosed environment. Nonetheless, the site origin and generation
91	mechanism of these droplets in the respiratory tract have yet to be discussed in the listed studies.
92	Against this background, further numerical studies have been conducted on the interactions
93	between high-speed, chaotic exhaled air and the liquid-layer lining the inner surface of the
94	airway (i.e., the mucus layer) during the coughing episode. The two most common methods
95	adopted are the volume of fluid (Paz et al., 2019; Rajendran & Banerjee, 2019; Pairetti et al.,
96	2021; Yi et al., 2021) and Eulerian wall film (EWF) models (Paz, Suárez, Parga, et al., 2017;
97	Paz, Suárez, & Vence, 2017; Ren et al., 2018, 2020, 2022; Anzai et al., 2022). These studies

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98 provide an understanding of the respiratory droplet generation process, the effects of mucus 99 properties on cough clearance efficiency, and the impact of airway deformation. To facilitate 100 the assessment of the infection risks of virus-laden droplets, it is crucial to qualitatively and 101 quantitatively investigate the origin of respiratory droplets during expiratory events (i.e., 102 coughing). Furthermore, the realistic and comprehensive characteristics of the target 103 respiratory airway model should be considered to ensure the accuracy of the cough-jet stream 104 and expelled droplet features. 105 In this study, the EWF model and Lagrangian discrete phase model (DPM) were coupled

106 In this study, the 2007 model and Englangtan discrete phase model (D100) were cooped 106 to characterize the following: (1) fluid flow profiles of coughing; (2) number concentration, 107 size distribution, and locality of generated and expelled droplet particles during coughing; and 108 (3) absorption efficiency of stripped droplets. The results establish a link between the high viral 109 load of the infected respiratory tract and the possibility of such viral pathogens being released 110 into the environment by coughing. Subsequently, the infection risk can be determined for 111 different respiratory viruses.

### 112 II. MATERIALS AND METHODS

### 113 A. Numerical model geometry

The realistic airway model comprised a computed tomography (CT)-based tracheabronchus model and oral cavity produced by the open-source DAZ Studio software (DAZ Productions, Inc.) shown in Fig. 1. During coughing, various oral shapes and sizes were

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117	observed (Dbouk & Drikakis, 2020). In this study, the newly developed oral cavity model
118	mimicked the configuration of a slightly open mouth with dimensions of $L = 4.7$ cm and $H =$
119	0.85 cm (Fig. 1) to similarly match the data provided by experimental measurements from high-
120	speed imaging (Dbouk & Drikakis, 2020). The total area of the open mouth of 3.76 cm <sup>2</sup> was
121	within the range of 4 $\pm$ 0.95 $\text{cm}^{2}$ as stated in a previous study (Seminara et al., 2020). In
122	addition, the teeth attributes were integrated to provide the most realistic traits of the oral cavity
123	(Fig. 1). The realistic trachea-bronchus model was created from CT scans, and details on the
124	process are available in the previous study by Ito (Ito, 2016). This trachea-bronchus model, in
125	combination with the nasal cavity, has been validated by our research group, providing reliable
126	results in diverse research themes (C. Wang et al., 2020; Yoo & Ito, 2022; Kuga et al., 2021,
127	2022, 2023; Murga et al., 2023; Khoa, Li, et al., 2023).
128	B. Grid design information

The discretization process was executed using the poly-hex core elements, which proposed the CFD simulation with higher accuracy at a reduced computational cost (Zore et al., 2019). The accuracy in the vicinity of the wall was enhanced by applying ten prism layers. This hybrid mesh has been successfully used to predict airflow and particle transportation/deposition simulation of the respiratory tract (Khoa, Phuong, et al., 2023; Khoa, Li, et al., 2023). According to the mesh independence test, the mesh size of 15.5 million cells was selected for subsequent simulation in this study, more detailed information can be found in Fig. S1

- 136 (Supplementary Material). This analysis strikes the balance between the computational burden
- 137 and prediction accuracy.

### 138 C. Numerical simulation of airflow pattern

- 139 The unsteady, incompressible, and isothermal fluid flow in the human respiratory tract was
- 140 obtained by solving the Reynolds-averaged Navier–Stokes equations.

$$\frac{\partial \overline{U}_i}{\partial x_i} = 0 \tag{1}$$

$$\frac{\partial \overline{U_i}}{\partial t} + \frac{\partial \overline{U_i} \overline{U_j}}{\partial x_j} = -\frac{1}{\rho_g} \frac{\partial \overline{\rho_g}}{\partial x_i} + \frac{\partial}{\partial x_j} \left[ \left( \nu + \nu_T \right) \left( \frac{\partial \overline{U_i}}{\partial x_j} + \frac{\partial \overline{U_j}}{\partial x_i} \right) \right]$$
(2)

141 where  $\overline{U}$  is the mean velocity; u' is the fluctuating components; and  $p_g$ ,  $\rho_g$ ,  $\nu$ , and  $\nu_T$  are

142 the pressure, density, kinematic viscosity of the fluid, and turbulent viscosity, respectively.

143 This study selected the turbulent model of the shear stress transport (SST) k- $\omega$ . This model has been used to predict adverse pressure gradient flow, strong curvature, and swirling flow in 144 145 airway systems, as shown in previous experiments (Phuong & Ito, 2015; Elcner et al., 2016). The coughing flow profile was obtained from field measurements of Gupta for male subjects, 146 as shown in Fig. 2A (Gupta et al., 2009). From the empirical equation, the coughing flow rate 147 was allocated to four inlets according to the flow weighting of each lobe of the lung proposed 148 149 in (Shelley et al., 2014), and reasonably applied in our study (Khoa, Li, et al., 2023), as shown in Fig. 2B. This procedure was implemented via a user-defined function (UDF) macro in 150 151 ANSYS Fluent. The numerical boundary conditions for the coughing airflow simulation are

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152listed in Table I.

### D. Eulerian Wall Film simulation 153

In a coughing episode, an excessive fluid flow velocity is rapidly produced and subjected 154 to turbulence and high shear stress at the interface between the airstream and thin liquid film 155 (mucus/saliva). Under such conditions, Kelvin-Helmholtz instabilities occur, which cause 156157 waves with an escalating amplitude on the surface of a thin liquid film (Pöhlker et al., 2021). 158 Droplets form from the crest of these waves in a multimodal mode and are carried by the flow. 159 The droplet generation mechanism can be predicted using the EWF model, in which thin 160 liquid mucus layers were hypothesized to be aligned on the inner surface of the oral-tracheal

model. The governing equation of the EWF model is as follows (ANSYS, Inc. 2022): 161

$$\frac{\partial \rho_l h}{\partial t} + \nabla_s \cdot \left( \rho_l h \vec{V}_l \right) = \dot{m}_s \tag{3}$$

$$\frac{\partial \rho_l h \vec{V}_l}{\partial t} + \nabla_s \cdot \left( \rho_l h \vec{V}_l \vec{V}_l + \vec{D}_V \right) = -h \nabla_s P_L + \rho_l h \vec{g}_\tau + \frac{3}{2} \vec{\tau}_{fs} - \frac{3\mu_l}{h} \vec{V}_l + \vec{q}_s$$
(4)

$$\vec{D}_V = \frac{\partial}{\partial s} \int_0^h v_l^2 dy$$
(5)

$$P_L = P_{gas} + P_h + P_\sigma \tag{6}$$

$$P_{h} = -\rho h\left(\vec{n}.\vec{g}\right) \tag{7}$$

$$P_{\sigma} = -\sigma \nabla_{s} \cdot \left( \nabla_{s} h \right) \tag{8}$$

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### where $\rho_l$ is the film density, h is the film height, $V_l$ is the mean film velocity, and $m_s$ is the 163 mass source per unit wall area owing to droplet collection, film separation, film stripping, and

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164	phase change. In Equation 4, $D_V$ is the differential advection term computed based on the
165	quadratic film velocity with fluctuating velocity $v_l(s,y,t)$ , in which s is the horizontal flow
166	direction and y is the vertical direction (Kakimpa et al., 2015). The term $P_L$ is the mucus film
167	pressure, $g_{\tau}$ is the gravity component, $\tau_{fs}$ is the shear force at the film–liquid interface, $\mu_l$ is the
168	viscosity, $q_s$ is the momentum source, $\sigma$ is the surface tension, $P_{\sigma}$ is the pressure exerted by the
169	surface tension, $P_h$ is the gravity component normal to the wall, and $n$ is the normal vector.

The reliability of applying the EWF model in anticipating the interaction between the coughing stream jet and lining fluid is emphasized through supplementary simulations conducted using a simplified airway model. The simulation results were subsequently compared with experimental data, and additional information on this validation process is elaborated upon in Fig. S2 (Supplementary Material).

175 In the context of our primary simulation, the EWF model was included with the fluid flow 176 at the beginning of the coughing episode. The simulation was applied to the total inner surface 177 of the numerical domain, including the teeth surface. In general, the mucus thickness varies 178along the airway system. However, to simplify the simulation, a constant thickness of 30  $\mu$ m 179 was used to represent the mucus layer in the throat, larynx, trachea, and bifurcation, based on previous studies (Paz, Suárez, Parga, et al., 2017; Paz, Suárez, & Vence, 2017; Ren et al., 2022; 180 Anzai et al., 2022). For the oral cavity, the saliva thickness ranged from 11.3 to 68.9 µm, as 181 182 proposed by the research of Assy (Assy et al., 2022). For the teeth surface, the saliva thickness

was assigned based on early experimental data (Collins & Dawes, 1987), which ranged from
2.59 to 4.44 µm following the mandibular, maxillary, left, or right position of teeth. The saliva
and mucus layers were assumed to be water with a density of 998.2 kg/m<sup>3</sup> and viscosity of
0.001 kg/m s.

187 Shear-induced droplet generation was considered through the high velocity and turbulent 188 flow, causing the instability of the lining mucus/saliva layers and leading to droplets peeling 189 from the crests of the formed waves. This process was defined by the initial parameters given 190 in Table II. Among them, the critical shear stress is the main factor that governs the number 191 concentration of generated droplets. A parametric analysis was required prior to the main 192 simulation to determine an appropriate value for the simulation, which is delivered in Fig. S3 193 (Supplementary Material). Notably, this analysis was significantly influenced by the individual 194 morphological characteristics considered in this study. Then, a value of 5 Pa was specified for 195 the critical shear stress, which imposed the limit that any region subjected to shear stress greater 196 than 5 Pa would trigger the shedding of mucus/saliva layers into droplets. In addition, a 197 diameter coefficient, which specifies the droplet size range (ANSYS, Inc. 2022), was also determined by experiencing the parametric analysis with a value of 0.0003 (Fig. S4, 198 199 Supplementary Material). Finally, the film time-step size was automatically assigned by 200 ANSYS Fluent using adaptive time-stepping functions, which controlled the time-step size to 201 be small enough to ensure that the maximum Courant number during the simulation was less

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202 than 1. The continuous generation of mucus/saliva layers beneath the epithelial cells was

203 neglected, which indicated no refill of mucus/saliva layers after being dispossessed.

### 204 E. Discrete Phase model simulation

The EWF model was coupled with the DPM to track the trajectories of droplets stripped from the liquid film. Forces acting on the body were used to predict the droplet transportation, absorption, and exhalation characteristics of the oral-tracheal model. The Lagrangian discrete phase of the particle trajectories was computed by Equation 9.

$$\frac{d\vec{u}_p}{dt} = \vec{F}_D + \vec{F}_G + \vec{F}_S \tag{9}$$

where the subscript p is the droplet phase and  $F_D$  is the drag force per unit particle mass derived from Stokes' drag law, expressed in Equation 10.

$$\vec{F}_D = \frac{18\mu}{\rho_p d_a^2} \frac{C_D \operatorname{Re}_p}{24} \left( \vec{U} - \vec{u}_p \right)$$
(10)

211 where  $\mu$  is the air viscosity,  $\vec{U}$  is the fluid flow velocity,  $\vec{u}_p$  is the droplet velocity,  $d_a$  is 212 the aerodynamic droplet diameter,  $\rho_p$  is the droplet density,  $C_D$  is the drag coefficient, and  $Re_p$ 213 is the particle Reynolds number.

The second term,  $\vec{F}_{G}$ , denotes the gravitational settling. The third term,  $\vec{F}_{s}$ , is Saffman's lift force due to shear on a unit mass basis. The lift force was adapted from a previous study by Li and Ahmadi (A. Li & Ahmadi, 1992), and is a generalization of the expression provided by Saffman (Saffman, 1965), expressed as Equation 11.

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$$\vec{F}_{s} = \frac{5.188 v^{\frac{1}{2}} \rho d_{ij}}{\rho_{p} d_{a} \left( d_{ik} d_{kl} \right)^{\frac{1}{2}}} \left( \vec{U} - \vec{u}_{p} \right)$$
(11)

218 where  $d_{ij}$ ,  $d_{lk}$ , and  $d_{kl}$  are deformation rate tensors.

219 The aerodynamic diameter and number of tracked droplets were determined based on the EWF simulation. The droplets were defined with a unit density (1000 kg/m<sup>3</sup>), equal to that of 220 221 pure water. The fate of the droplets was considered as exhaled via the mouth opening or was re-absorbed into the mucus layer; hence, the "escape" boundary condition was assigned to the 222223 mouth opening, and the "perfect trap" condition was applied as the wall boundary conditions. 224 The evaporation and breakup of the droplets were negligible. The droplets were continuously 225 generated during the simulation; therefore, the velocity, size, and spatial and temporal 226 information of the droplets stripped from the mucus layer were recorded using the UDF macro 227for each time-step.

The generated, exhaled, and absorbed percentages, denoted as  $\eta_{G-i}$ ,  $\eta_{E-i}$ , and  $\eta_{A-i}$ , can be expressed by Equations 12, 13, and 14, respectively.

$$\eta_{G-i} = \frac{N_{G-i}}{N_G} \times 100\%$$
(12)

$$\eta_{E-i} = \frac{N_{E-i}}{N_E} \times 100\%$$
(13)

$$\eta_{A-i} = \frac{N_{A-i}}{N_A} \times 100\%$$
(14)

where  $N_{G-i}$ ,  $N_{A-i}$ , and  $N_{E-i}$  are the number of droplets generated, absorbed, and exhaled that belong to region *i*, which corresponds to the regions defined in Fig. 1.  $N_G$ ,  $N_A$ , and  $N_E$  are the total stripped, absorbed, and exhaled droplets after a single coughing episode (duration of 0.5

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s), respectively. The size distribution of the coughed droplets was multimodal; accordingly, the droplet size bin was identified to establish the size distribution percentage of the generated, exhaled, and absorbed droplets ( $\eta_{G-s}$ ,  $\eta_{E-s}$ , and  $\eta_{A-s}$ ), given in Equations 15, 16, and 17, respectively.

$$\eta_{G-s} = \frac{N_{G-s}}{N_G} \times 100\%$$
(15)

$$\eta_{E-s} = \frac{N_{E-s}}{N_E} \times 100\% \tag{16}$$

$$\eta_{A-s} = \frac{N_{A-s}}{N_A} \times 100\%$$
(17)

where  $N_{G-s}$ ,  $N_{A-s}$ , and  $N_{E-s}$  are the number of droplets generated, absorbed, and exhaled, respectively, which decreases in the size bin, as listed in Table III.

The size distribution of exhaled droplets can be calculated by dividing the number of droplets within the specific size bin by the logarithm of the droplet size class interval ( $dN_{E}$ dLogD). Finally, the exhaled droplet size distribution was normalized with the total cough exhaled volume (1,000 cm<sup>3</sup>) to obtain the number concentration divided by the logarithm of the droplet size class interval ( $dC_{NE-s}/dLogD$ ).

### 244 III. RESULTS

### 245 A. Coughing fluid flow characteristics

The results of the coughing fluid flow features are depicted in Fig. 3 at the start of the coughing episode at 0.01 s, cough peak flow rate (CPFR) at 0.077 s, and near the end of the coughing event at 0.4 s. Notably, the velocity magnitude and distribution herein are described

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249	at the instantaneous times. At the onset of coughing (Fig. 3A), the culminated velocity occurred
250	at the bifurcation, trachea, glottis, and throat regions with a value of 3 m/s. At CPFR, the spatial
251	distribution of the jet stream remained, but the magnitude increased by almost 13-fold up to 40
252	m/s. In the final coughing stage, the coughing velocity rapidly decreased to less than 3 m/s in
253	identical acceleration regions.
254	In the oral cavity (Fig. 3B), the airstream accelerated in the throat region and impacted the

255palate and bends following the curvilinear shape of the oral ceiling. The expulsive flow that 256escaped the oral region was primarily distributed at the bottom of the mouth opening. Reserve 257 flow also formed in the basal region near the mouth opening, which contributed to the swirling flow at the mouth opening (Fig. 3B). Due to the flow features toward the ceiling of the oral 258 259 cavity, the maxillary teeth are expected to endure the high-velocity flow during the cough. Fig. 3C shows the two-dimensional flow distribution for the maxillary teeth, which revealed that 260261 the high-velocity fluid flow attacked the inner surface of the molars, premolars, canines, and 262incisors.

### 263 B. Droplet generation mechanism during coughing

The rapid increase in the flow rate due to coughing induces substantial shear stress on the airway wall, closely linked to the droplet production criteria in the EWF model simulation. The relationship between shear stress, mucus thickness, and stripped droplets is presented in Fig. 4, which describes the instantaneous value of each variable. At 0.01 s (Fig. 4A), the low coughing

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268	velocity resulted in a modest shear stress of less than 5 Pa on the airway wall, which failed to
269	meet the minimum threshold to produce droplets; hence, the droplets were not observed and
270	the mucus/saliva thickness remained in its original state. At CPFR (Fig. 4B), the cough ejection
271	velocity increased to 40 m/s, which induced significant shear stress on the wall surfaces.
272	Multiple airway surfaces experienced shear stress levels greater than 5 Pa, which fulfilled the
273	criterion for droplet generation off the thin liquid film on the oral-airway surfaces. Accordingly,
274	the mucus/saliva thickness decreased to almost 0 µm and droplets simultaneously emerged in
275	the corresponding regions (Fig. 4B). The complex and uneven surface of the oral-airway model
276	produced an uneven distribution of the peak shear stresses. This completely removed the
277	mucus/saliva layers in several specific regions while the remaining areas preserved their initial
278	liquid film thickness. Droplet production decreased significantly at 0.15 s (Fig. 5C) due to the
279	mucus/saliva layers dissipating, following the former intense erosion process during CPFR.

### 280 C. Properties of generated droplets during coughing

281 The relationship between the coughing flow profile and droplet production was correlated 282 to understand the droplet generation process better. The instantaneous number of droplets 283 stripped from the mucus/saliva layers due to the coughing flow rate during the single coughing episode (~0.5 s) is shown in Fig. 5A. The results demonstrate that the coughing flow rate 284 285 rapidly increased in the early stage of coughing. In contrast, droplet generation lagged and 286 didn't start until 0.04 s with an initial rapid rise. The number of droplets produced increased

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287	rapidly in parallel with the coughing flow rate and formed a sharp slope, with the peak almost
288	coinciding with the CPFR (at approximately 0.077 s). After reaching their peaks, the cough
289	flow rate and number of stripped droplets exhibited a downward trend. The number of droplets
290	produced rapidly dropped to almost 0 at 0.25 s, in contrast to the cough flow that gradually
291	decreased to 0 taking 0.5 s. After the single coughing episode, 11,594,566 droplets were
292	generated (Table IV). The instantaneous position of the droplets during different times during
293	the cough are shown in Fig. 5B for time, $t = 0.025$ , 0.04, 0.077, and 0.2 s, where the droplets
294	are colored by diameter. The results indicate an early appearance of droplets in the bifurcation
295	and throat regions (at 0.025 s). At 0.04 s, a dense population of droplets was observed in the
296	bifurcation and oral cavity, which corresponds to the airway curvature of the palate region in
297	line with the flow distribution in the oral cavity. At CPFR at $t = 0.077$ s, the oral-airway model
298	is filled with a very high density of droplets due to the rapid droplet production (shown in Fig.
299	5A). At $t = 0.2$ s, most of the droplets were re-absorbed into the mucus/saliva layers or
300	transported by the coughing flow toward the oral region and released into the external
301	environment via the mouth opening. Hence, fewer droplets are found in the airway compared
302	to the oral cavity. Furthermore, the droplet size was less than 10 $\mu m$ for the entire coughing
303	episode.

304 One advantage of this study is that it provides details on the origin of the stripped droplet 305 during the coughing event, illustrated in Fig. 5C. The oral region was identified as the primary source of droplets (up to 46.8%) during the cough. In the remaining regions, the droplet

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307	production levels were similar (approximately 11.9-14.3%), slightly reducing towards the
308	bifurcation region. The large amount of stripped droplets in the oral cavity is attributed to
309	higher shear stresses (exceeding 5 Pa) than in other regions. Fig. 5D shows the total surface
310	area (cm <sup>2</sup> ) of each region subjected to shear stress of >5 Pa, where the oral cavity exhibited the
311	highest surface area. Therefore, more droplets were produced from the saliva film in the oral
312	airway region.
313	The percentage of droplets generated per droplet diameter size bin can be estimated using
314	Equation 15, as shown in Table IV. Most droplets produced were in the size interval of 4–8 $\mu m$
315	(approximately 78.01%), followed by 2–4 and 8–16 $\mu$ m, respectively. The droplets in the size

bin of  $<1 \mu m$  had a low percentage of 0.078–0.181%, while for the larger size bins (>16  $\mu m$ ),

the rate was only 0.169%.

### D. Properties of exhaled droplets during coughing 318

319 An analysis of the expelled droplet concentration for different droplet diameters was 320 performed and compared with experimental measurements of Yang (Yang et al., 2007). In the 321 experimental work, 54 volunteers of varying ages and genders coughed into a sampling bag 322 with a well-controlled relative humidity; thus, the coughed droplets retained their original size. 323 Fig. 6A shows the number concentration of droplets at the mouth opening plotted against the 324 average droplet size and compared with Yang (Yang et al., 2007). The EWF model closely

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325	matched the profile with the measured exhaled number concentration. There was a sharp
326	increase in the number concentration for droplet sizes approximately at 2 $\mu$ m before reaching
327	a peak concentration (approximately 2,500/cm <sup>3</sup> ) at 4–8 $\mu$ m. After the peak, a downward trend
328	was observed, and the cutoff diameter where no droplets were expelled was >10 $\mu m.$
329	The source of droplets expelled to the environment is shown in Fig. 6B, where the oral
330	cavity (including the teeth surface) was responsible for the largest amount of droplets exhaled
331	into the environment (up to 75%). The amount gradually reduced for the geometry moving

posteriorly toward the caudal airway; specifically, 12.3% originated from the throat, followed
by the larynx (approximately 8.3%), trachea, and bifurcation (approximately 2.2%),
respectively.

Most exhaled droplets, 73.1%, were between 4-8  $\mu$ m due to the highest percentage of droplets produced in this range (Table IV). For smaller droplets, the exhalation rates were 12.8% and 8.9% for size bins of 2–4  $\mu$ m and 8–16  $\mu$ m, respectively. Despite the low production rate of droplets in the size bin of <1  $\mu$ m, the appearance of these small droplets in the exhaled breath was 0.49–0.79%.

The spatial distribution of the droplets that escaped through the mouth during the coughing is illustrated in Fig. 6C, where the different droplet colors denote the source location. The results show that the oral cavity is the primary location of the expelled droplets, and the droplets exit through the entire space of the mouth opening. For the other airway regions (throat, larynx,

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349	E. Absorbed efficiency of droplets during coughing
348	and the exhaled jet stream found in the lower half of the mouth opening (shown in Fig. 3).
347	distribution in the lower half was consistent. This distribution was due to the oral cavity shape
346	Despite the significant variation in the vertical distribution of droplets expelled, the horizontal
345	mouth opening. There was a small scattering of droplets in the upper half of the mouth opening.
344	trachea, and bifurcation), the expelled droplets mainly dispersed through the lower half of the

The total number of droplets absorbed onto the mucus/saliva layers was 10,128,559 (Table IV), accounting for approximately 87.3% of the total droplets produced by the cough. The primary absorption region was the oral cavity (including the teeth surface), with 47.8%. In contrast, in the remaining regions, the absorption efficiency significantly decreased by approximately four-fold in the 11.2–14.1% range. Most of the generated droplets re-absorbed into the oral region's saliva layer can be associated with the complex morphology of this region, which prevented the smooth movement of the droplets.

Fig. 7B shows the droplet absorption efficiency categorized by the region where the droplets originated from (e.g. slice color indicates the droplet source location). In general, droplets were immediately re-absorbed back into its own region where they were generated. For example, 87.5% of droplets produced in the oral cavity re-absorbed in its region, and that of the throat, larynx, trachea, and bifurcation, the re-absorption rate was 88.2%, 91.9%, 93.1%, and 100%, respectively.

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363	The cough-jet stream began from the tracheal bifurcation, and the oral region had the
364	greatest exposure to all droplets originating from lower regions, including the throat (5.3%),
365	larynx (4.2%), trachea (1.8%), and bifurcation (1.3%). This geometry and flow feature explains
366	the lack of absorbed droplets from geometrically lower regions (e.g. upstream flow) than the
367	region itself since the jet flow transports the droplets from the bifurcation to the oral cavity.
368	Droplets with sizes of 4–8 $\mu m$ (Table IV) were the most re-absorbed due to the greater number
369	of generated droplets in this size bin. For the other size bins, the absorption rate was similar to
370	its generation rate.
371	Fig. 8 shows the deposition pattern on specific airway regions based on where the droplets
372	were produced. Deposition in the oral cavity showed that the focal absorption region occurred
373	in the palate regardless of the droplet origin. Most droplets in the throat tended to accumulate
374	in the upper and branching regions. In the larynx and trachea, there was a high rate of re-

absorption from itself. Only the droplets produced by itself were re-absorbed for the bifurcation,

and most were observed in the left bifurcation.

377 Generally, the cough-jet stream influenced droplet deposition in the oral airway, and the 378 droplet absorption patterns coincided with the high-velocity regions, as discussed in the 379 previous section.

### 380 F. Role of teeth surface in the simulation of droplet generation during coughing

381 The results reveal the critical role of the oral cavity in droplet generation, emission, and

absorption. This raises the question of the contribution of the teeth surface to the areas of

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383 interest in this study. In this section, the droplet generation, absorption and exhalation from the 384 oral cavity were divided into the teeth surface and the remaining portion of the oral airway 385 surface (shown in Fig. 9A). The total of 46.8% of droplets that were produced from the oral 386 cavity (from Fig. 5C), were found to originate evenly between the teeth surface and remaining 387 portion (approximately 23%). The instantaneous shear stress at 0.04 s and 0.077 s is given in Fig. 10A. Shear stress >5 388 389 Pa was observed on the maxillary incisor surface in the early stage of the coughing event (0.04)390 s). A high shear rate was found on the inner surface of the maxillary teeth, including the incisors, 391 canines, and molars, at the CPFR (0.077 s). Consequently, many droplets were stripped from 392 the saliva layers along the inner side of the maxillary teeth surface (Fig. 10B). This 393 phenomenon is closely associated with the cough-jet stream, which impacted the upper jaw, as

394 discussed in the previous section.

For the expelled droplets (Fig. 9B), approximately 40% was derived from the teeth's surface. This was attributed to the droplets that formed on the teeth surface, especially on the incisors, which travelled a short distance without any obstacles to the mouth opening. For the reabsorption capacity, the larger surface area of the remaining portion of the oral region caused a larger number of droplets to re-absorb (approximately 27%). In comparison, 21% of droplets re-absorbed onto the teeth surface.

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401	Figure 11 presents both the quantitative analysis and spatial distribution of droplets
402	absorbed on the teeth surface. Primarily, the quantitative analysis serves to delineate the
403	proportion of total absorbed droplets on the teeth surface, which was generated from distinct
404	regions within the model. The findings revealed that out of the total number of droplets
405	absorbed on the teeth surface, 81.02% of absorbed droplets originated from this area.
406	Subsequently, droplets emanating from the oral cavity constituted 14.79% of the total absorbed
407	droplets on the teeth surface. The percentage gradually reduced for the posterior regions and
408	reached 0.45% for the bifurcation, which implies a limited quantity of droplets stemmed from
409	this region absorbed onto the teeth surface. In addition, the spatial distribution of the droplet
410	deposition on the teeth surface indicates that most droplets tended to settle on the inner surface
411	of the maxillary teeth or partially in the mandibular molars and canines, regardless of their
412	origin.

### 413 IV. DISCUSSIONS

The coupled EWF-DPM model was applied to explore droplet generation and flow behavior during a single coughing event, where the origin and local number concentration of the generated, absorbed, and expelled droplets were determined. This study first analysed the coughing flow rate, which provides insight into the droplet generation process during coughing. The rapid development of airflow velocity along the airway and oral cavity subjected the surface walls to high shear stress, which caused the stripping of mucus/saliva layers into

droplets. The airstream accelerated through the narrow airway lumen (e.g., trachea, glottis, and

throat regions), consistent with a previous study (Kou et al., 2018). Thus, the predicted fluid

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422 flow is recognized as a constant feature of coughing in terms of the velocity distribution but 423 would vary in magnitude depending on the lumen diameter of individual airway structures. In 424 the oral cavity, this study included the teeth geometry, which has been lacking in reported 425 simulation studies of the fluid flow characteristics during coughing. Our results indicate a 426 strong interaction between the cough-jet stream and maxillary teeth surface, which could 427 influence droplet generation and absorption/exhalation. 428 The shear-induced droplet generation due to the high-speed velocity is one of the four 429 generation mechanisms that mainly occurs in the main bronchus, trachea, and larynx regions 430 (Pöhlker et al., 2021). The results revealed an essential connection between the shear stress and 431 the number concentration of generated droplets, summarized as follows. The higher the 432 coughing flow rate, the greater the shear stress, and the larger the number of droplets generated. 433 This relationship indicates the uncertainty of the generated droplet number concentration owing 434 to the coughing flow profile, particularly at the CPFR. For instance, the significant diversity 435 between individuals and genders has been recorded for coughing parameters, such as the peak

436 velocity, peak velocity time, and coughing duration by field measurements (Han et al., 2021).

437 These variations are expected to affect droplet generation. In addition, the fidelity of the

438 numerical domain needs to be considered. Earlier simulation studies used an idealized model

that proposed a smooth shear stress distribution on the wall surface (Anzai et al., 2022). The

study by Ren et al. (Ren et al., 2022) reported a nonuniform distribution of shear stress on a

realistic lower airway wall. The results showed heterogeneities in spatial distribution and

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442	magnitude of shear stress compared with ours; consequently, the droplet generation differed.
443	Nevertheless, the coughing flow profile, realistic attributes, and intersubject variability
444	presented challenges, and the simulation results need to be benchmarked against the
445	experimental results.
446	For infection risk assessment, it is crucial to understand the origins of droplets during
447	expiratory activities within a specific area and to quantify the droplets generated from each
448	source (Morawska, 2006). For coughing, droplet-released sources are well established and
449	consist of the lungs, trachea, nasopharynx-larynx, and nasal and oral passages (Stadnytskyi et
450	al., 2021; Zhou & Zou, 2021). However, the exact location of droplet generation/exhalation
451	and their localized number concentration remain uncertain. Our droplet generation analysis
452	indicates that the most likely sources of droplets produced during coughing were the oral cavity
453	and teeth surface. Although common respiratory viruses primarily occur in the epithelial cells
454	of the respiratory system (Alexander-Brett & Holtzman, 2015), evidence suggests high viral
455	loads of SARS-CoV-2 in saliva samples from an asymptomatic cohort and the active replication
456	of infected cells in the oral cavity (Huang et al., 2021). In addition, previous clinical studies
457	have detected SARS-CoV-2 in patient saliva (To et al., 2020; Wölfel et al., 2020; Wyllie et al.,

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458	2020). Thus, from our results, the high infection risk of SARS-CoV-2 may be associated with
459	a high droplet concentration in the oral cavity. Apart from the oral cavity, the deposition site
460	can be determined in the airway system of the potential host (C. C. Wang et al., 2021), but this
461	depends on the aerosols or virus-laden droplet size. Droplets from the deposition regions may
462	cause the re-emission of progeny viruses shed by infected cells (Schaefer & Lai, 2022).
463	Therefore, the possibility of transmission droplets originating from the respiratory system was
464	indicated by our analytical data, where considerable quantities of exhaled droplets were
465	recorded from the throat, larynx, trachea, and bifurcation. The results showed that the
466	generation sites of droplets tended to be in a particular position rather than evenly distributed.
467	The site of origin affected the potential direction of the droplet cloud during a cough. Our
468	results showed that exhaled droplets from the oral cavity escaped to the surroundings at all
469	locations and angles from the mouth opening; hence, the droplets travelled further and
470	dispersed widely. Meanwhile, exhaled droplets stripped from the respiratory airway exhibited
471	a downward direction as they exited the mouth into the ambient environment. This
472	phenomenon is expected to direct the droplets to the ground outside. Thus, our location-specific
473	results can inform studies on the threat of coughing-related infections in indoor environments.
474	In addition to identifying the origin site, the local number concentration and size
475	distribution of droplets exhaled during coughing were determined. Based on the literature
476	review, these two parameters were estimated for a varied population and measurement

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477	techniques. As summarized in Fig. S5 (Supplemental Material), the field measurement data
478	were scattered broadly regarding the size distribution and number concentration. In particular,
479	the peak number concentration was recorded in the 4–8 $\mu m$ range proposed by Yang and Chao
480	(Yang et al., 2007; Chao et al., 2009). This peak shifted to 0.75–2 $\mu$ m in the experiment by
481	Morawska and Johnson (Morawska et al., 2009; Johnson et al., 2011). The highest number
482	concentration was recorded at a much smaller size, at 0.3 $\mu m$ measured by Lindsley and Zayas
483	(Lindsley et al., 2012; Zayas et al., 2012). In addition, the number concentration among the
484	references indicated significant variations. Therefore, our analysis was validated by recruiting
485	one specific experimental data from Yang et al (2007). Both experimental and simulated data
486	showed that a 4–8 $\mu m$ droplet size was formed and exhaled at a high number concentration
487	during coughing. Within this size range, the expelled droplets would travel further into the
488	environment and suspend for longer, enhancing the exposure risk of residents in a confined
489	space (Jones & Brosseau, 2015; Dhand & Li, 2020; Bourouiba, 2021). Nevertheless, the
490	lifetime and size of droplets have a complex relationship with many environmental factors,
491	including temperature, humidity, and ventilation mechanisms (Jayaweera et al., 2020;
492	Bahramian, 2023). Once inhaled by a susceptible person, fine aerosols (<5 $\mu m$ ) have a high
493	possibility of escaping the defence mechanism of the upper airway and penetrating the lower
494	airway, which is often associated with higher severity, morbidity, and fatality (Zuo et al., 2020;
495	Sosnowski, 2021). A more significant amount of viral genomes of common respiratory viruses

496 have been revealed in fine aerosols (<5 μm) compared with larger ones (Gralton et al., 2013;</li>
497 Yan et al., 2018). Information regarding the site origin and local number concentration was
498 revealed by our analysis, which may not be possible in field measurements with volunteers
499 owing to ethical and technical barriers.

500 Our simulation data showed that one issue that should be addressed in droplet generation, 501 absorption, and exhalation studies is the inclusion of detailed oral cavity and teeth features. 502 The oral cavity was a significant source of droplet expulsion; however, due to its complex 503 anatomy with the existence of the teeth, the region provided a source for droplet absorption, 504 thereby contributing to the mitigation of droplet emissions from caudal respiratory areas. By 505 eliminating realistic features or simplifying the oral cavity, the high local concentrated 506 absorption regions may shift to the larynx-throat region, according to our data and a previous simulation (Guo et al., 2020), allowing more droplets to be expelled. Thus, the realistic 507 508 anatomy of the oral cavity alters the number concentration and spread angle of ejected droplets, 509 thereby changing the dispersion and migration characteristics of the expelled droplets.

510 Throughout the discussion points, it is noteworthy that the data was obtained upon the 511 assumption of one single cough event. Concerns may arise regarding whether introducing a 512 successive coughing process would impact the results. In this context, the previous 513 experimental research revealed the second cough characteristics with a homogeneous profile 514 but weakened mechanical effectiveness, such as cough peak flow rate and expired volume

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(Gupta et al., 2009; Hegland et al., 2013). This weakening was prolonged until the end of the
cough epoch. Upon these conditions, outcome variation can be anticipated as following points:
(i) Within the coughing flow patterns, continuous accelerations are expected to occur
during each coughing episode, with the highest magnitude in the first cough and a
precipitous decline in the subsequent cough events.

- (ii) Regarding the generated droplet number concentration, although a number
  concentration increase is conceivable, it would not anticipated to yield a significant
  deviation from the current results. This expectation is rooted in the observation that
  the mucous membrane almost diminished after the initial coughing event, as indicated
  in Fig.4. Additionally, the WSS exerted on the respiratory wall may fall below the
  given CSS and be insufficient to trigger the droplet stripping.
- (iii) In terms of droplet behaviors (absorbed or expelled), consecutive acceleration of fluid
   flow can result in the variation of the absorption rate due to the inertia, subsequently
- 528 leading to the corresponding alternations in the escaped rate.

Henceforth, subjecting the current research to successive cough simulations while maintaining consistent initial conditions is expected to yield modest variations. Hence, it may be reasoned that the applicability of the results may be regarded as akin to the current findings under the assumption of a single cough process.

### 533 V. CONCLUSIONS

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534	Detailed information on the droplet generation, exhalation, and absorption behavior in a
535	realistic oral-tracheal model was elucidated using the coupled EWF-DPM model. The main
536	points of the analysis are summarised as follows:
537	• The EWF model reliably predicted the interaction between the free stream and
538	liquid film lining by validating it against measured data. Coupled with the DPM
539	model, the concentration profile of exhaled droplets from the mouth opening after
540	a coughing episode could represent experimental data.
541	• The morphometry fidelity of the oral-tracheal model and the coughing flow rate
542	affects the cough-jet stream and is expected to alter droplet generation
543	transportation/absorption, and exhalation.
544	• Shear stress at the liquid film-air interface is a critical factor governing the drople
545	generation properties. Therefore, appropriate values as the input criteria for the
546	EWF and benchmarking against experimental data are required.
547	• The oral and teeth surface were primary locations for droplet generation. While
548	other regions of the respiratory system showed a much lower droplet production
549	rate. The rates gradually decreased toward the posterior regions.
550	• Most exhaled droplets were 4–8 $\mu$ m in diameter. Droplet sizes of <1 $\mu$ m and >10
551	$\mu m$ were also detected but were much fewer upon exhalation through the mouth
552	opening.

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**VI. FIGURES** Teeth features Lateral view Oral cavity Oropharynx Larynx Trachea Bifurcation Frontal view Mouth opening H = 0.85

FIG. 1. Outline of the simulation model



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L = 4.7



FIG. 2. (a) Coughing flow rate following the previous field measurement. (b) The description of inlet boundary conditions.



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FIG. 3. (a) Instantaneous streamline velocity distribution at the specific time during coughing. (b) Instantaneous 2D flow features in the oral cavity and the mouth opening, and (c) in the vicinity of the maxillary teeth surface at the CPFR (0.077 s)

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FIG. 4. The immediate distribution of wall shear stress (left), mucus/saliva thickness (middle), and initial position of stripped droplets (right) at (a) 0.01 s, (b) 0.077 s, and (c) 0.15 s.

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FIG. 5. (a) The instantaneous number of generated droplets during the cough event associated with the coughing flow rate. (b) Instantaneous positions of droplets at different times during the cough. (c) The percentage of droplets produced from their origin. (d) The surface area of each airway region that experienced wall shear stresses greater than 5 Pa.



FIG. 6. (a) Validation of total concentration of exhaled droplets after one single cough event.(b) The percentage of exhaled droplets following their site origin. (c) Spatial distribution of exhaled droplets on the mouth opening after a single coughing episode.

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FIG. 7. (a) Total absorbed efficiency of the generated droplets on the inner surface of the airway region. (b) The absorption efficiency rate for different droplet source locations.

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FIG. 8. Visualization of the absorbed droplets in local regions of the oral-tracheal model (column) derived from a specific location (row).

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FIG. 9. (a) Outline of the oral cavity, including teeth surface and the remaining regions. (b)The total percentage of generation, exhalation, and absorption of droplets collapsed for the teeth surface and remaining portion.

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FIG. 10. (a) The instantaneous wall shear stress distribution on the teeth surface at 0.04 s (left) and 0.077 s (right). (b) Generated location and distribution of droplets on the teeth surface during coughing.

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Trachea

FIG. 11. 3D visualization of deposited droplets on the teeth surface colored by where the droplet was produced.

Bifurcation

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### 597 VII. TABLES

### 598 TABLE I. Numerical boundary conditions for the coughing airflow simulation.

Parameter	Information	
	SIMPLE (Semi-Implicit Method for	
Algorithm	Pressured Linked Equations	
Convection scheme	Second order upwind	
Density (kg/m <sup>3</sup> )	1.185	
Viscosity (kg/m s)	1.81x10 <sup>-5</sup>	
Averaged cough peak flow rate (L/s)	5.75	
Averaged cough exhaled volume (cm <sup>3</sup> )	1,000	
Total simulation time (s)	0.5	
Time step size (s)	0.001	

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### TABLE II. Initial parameter for Eulerian Wall film model.

-	
Parameter	Value
Critical shear stress – CSS (Pa)	5
Diameter coefficient	0.0003
Mass coefficient	0.25
Surface tension (N/m)	0.0589

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### 611 TABLE III. Classification of droplet size bin.

Droplet size bin (µm)	Averaged size (µm)
0.1 - 0.25	0.175
0.25 - 0.5	0.375
0.5 - 0.75	0.625
0.75 - 1.0	0.875
1.0 - 2.0	1.5
2.0 - 4.0	3.0
4.0 - 8.0	6.0
8.0 - 16.0	12.0
16.0 – 24.0	20.0
24.0 - 32.0	28.0

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	Generated	Exhaled	Absorbed
Droplet size bin(µm)	percentage- <i>η<sub>G-s</sub></i> <sup>a</sup>	percentage- <i>n<sub>Es</sub></i> <sup>b</sup>	percentage- <i>ŋ</i> <sub>4-s</sub> <sup>c</sup>
	(%)	(%)	(%)
0.1 - 0.25	0.078	0.49	0.044
0.25 - 0.5	0.133	0.79	0.078
0.5 - 0.75	0.149	0.65	0.108
0.75 - 1.0	0.181	0.50	0.157
1.0 - 2.0	1.94	2.60	1.94
2.0 - 4.0	11.81	12.82	11.58
4.0 - 8.0	78.01	73.06	78.47
8.0 - 16.0	7.53	8.93	7.47
16.0 - 24.0	0.15	0.15	0.15
24.0 - 32.0	0.019	0.01	0.003
Total percentage	100	100	100
Total number count	11,594,566 (N <sub>G</sub> )	938,206 (N <sub>E</sub> )	10,128,559 (N <sub>A</sub> )

TABLE IV. Generated, exhaled, and absorbed efficiency of droplets during the coughing eventfollowing the droplet size bin.

\*a,b,c The percentage was calculated using Equations 15, 16 and 17

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### 648 VIII. SUPPLEMENTARY MATERIAL

The supplementary material contains details and discussions regarding the optimization processes for the total mesh counts and the initial parameters for the EWF model, such as critical shear stress – *CSS* and diameter coefficient – *F*. In addition, detailed boundary conditions and results, which serve to reinforce the reliability of employing the EWF model to simulate the interaction between the free-stream flow and the lining fluid on the surface, are provided.

### 655 IX. LIMITATIONS

This study focused on a specific oral-tracheal model, which did not cover the individualrelated variability. Besides, it is notable that the initial parameters for the EWF model including critical shear stress – *CSS* and diameter coefficient – F were emphasized by the individual structure and numerical boundary conditions of this study. In addition, a rigid airway model was assumed, which is the opposite of the elastic nature of the respiratory tract and ignores glottis deformation during coughing.

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962	CONFLICT OF INTEREST STATEMENT
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963	The authors declare that they have no competing financial interests or personal
964	relationships that could have influenced the work reported in this paper.
965	DATA AVAILABILITY



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966 The data supporting this study's findings are available from the corresponding author upon

967 reasonable request.