From Microscopic Droplets to Macroscopic Crowds: Crossing the Scales in Models of Short-Range Respiratory Disease Transmission, with Application to COVID-19

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Short-range exposure to airborne virus-laden respiratory droplets is an effective transmission route of respiratory diseases, as exemplified by Coronavirus Disease 2019 (COVID-19). In order to assess the risks associated with this pathway in daily-life settings involving tens to hundreds of individuals, the chasm needs to be bridged between fluid dynamical simulations and population-scale epidemiological models. This is achieved by simulating droplet trajectories at the microscale in numerous ambient flows, coarse-graining their results into spatio-temporal maps of viral concentration around the emitter, and coupling these maps to field-data about pedestrian crowds in different scenarios (streets, train stations, markets, queues, and street cafés). At the individual scale, the results highlight the paramount importance of the velocity of the ambient air flow relative to the emitter's motion. This aerodynamic effect, which disperses infectious aerosols, prevails over all other environmental variables. At the crowd's scale, the method yields a ranking of the scenarios by the risks of new infections, dominated by the street cafés and then the outdoor market. While the effect of light winds on the qualitative ranking is fairly marginal, even the most modest air flows dramatically lower the quantitative rates of new infections.

1. Introduction

The theoretical combat against respiratory infections stretches over a whole gamut of lengthscales: These diseases are caused by viruses or bacteria, which measure several tens of nanometers ($10^{-8}$ m) and around one micron ($10^{-6}$ m) of diameter, respectively; these germs are (directly or indirectly) transmitted from person to person, using as carriers respiratory droplets (ranging from several hundred nanometers to several hundred microns in diameter) that are mostly expelled through the centimetric ($10^{-2}$ m) mouth gap and transported over some $\approx 10^{-1} - 10^0$ m. The daunting fundamental challenge of bridging so many scales to model the transmission of these diseases has also become an imperative practical necessity since a new coronavirus, the “severe acute respiratory syndrome coronavirus 2” (SARS-CoV-2) was identified during a first epidemic outbreak in Wuhan, China, in December 2019. Since then, it has spread all over the world and caused the Coronavirus Disease 2019 (COVID-19) pandemic, which is officially accountable for more than 750 million cases and 6.8 million deaths as of March 2023.[1]

Regarding the transmission pathways of respiratory diseases, direct impact of droplets of respiratory fluids on the nasal or oral mucosa of the susceptible individual as well as contact with droplet-contaminated surfaces (the so-called fomites) have long been identified as possible routes. However, it may be that these pathways only play a lesser role,[2] as the susceptible individual can also get infected after inhaling pathogen-laden aerosols exhaled by a contagious person, a mechanism termed airborne transmission.[3,4] Here, the term aerosol refers, and will henceforth refer, to all respiratory droplets small enough to dwell in the air for at least a few seconds and to be inhaled by somebody through their nose or mouth. The prevalence of this airborne transmission route has been increasingly acknowledged,[2,4–7] especially in crowded indoor environments that led to well documented superspreading events.[2,8–10] The alarm has also been raised with respect to crowded outdoor settings[11] (e.g., at mass sports events), where accumulation of virus-laden aerosols in the air is implausible but short-range exposure can occur; nevertheless, the actual risks that they present have been a bone of contention.[12]

To assess how the disease may spread in crowds, modeling the emission, transport, and inhalation of respiratory droplets is an appealing option and has been widely used in COVID-19-related studies. However, modeling transmission is a major challenge, owing to the sensitivity of droplet propagation to
environmental factors such as temperature, humidity, and wind.\textsuperscript{[13–15]} as well as the uncertainty about the sizes of emitted droplets \textsuperscript{[16]} or the person-to-person variability.\textsuperscript{[17]} among others. Moreover, microscopic studies of droplet propagation, supposed to describe the evolution of droplets most accurately, are generally limited to static scenarios involving two people facing each other. At the other extreme, most macroscopic models focusing on indoor transmission assume a well-mixed environment\textsuperscript{[9,18–20]} and thus overlook short-range exposure, which is the main source of risks outdoors. An interesting compromise has been proposed by Löchner et al.\textsuperscript{[21]} by simplifying the geometry, the boundary conditions, and the transport of the droplets, and resorting to coarse meshes, these researchers are able to perform full-scale simulations involving tens of pedestrians in motion. Nevertheless, each scenario, although tractable, remains computationally intensive.

Here, we endeavor to bridge the gap between detailed micro-environment studies and their macroscopic counterparts, by building on the framework outlined in ref. \textsuperscript{[22]}. In this framework, field data about pedestrian behavior (including the inter-pedestrian distances, interaction durations, head orientations, etc.) are coupled to concentration maps of virus-laden particles exhaled by a (supposedly contagious) individual in the crowd in order to assess the number of susceptible people that this individual would infect. Unfortunately, these concentrations maps were so far largely ad hoc and rested on crude modeling assumptions. In this paper, the connection with the microscale is fully established thanks to genuine computational fluid dynamical (CFD) simulations of droplet propagation (performed using large-eddy computations to account for flow turbulence) and converted into concentrations maps via a transparent coarse-graining method. A variety of ambient conditions, notably air flow velocities, are considered, which enables us to quantify the effect of ambient air flows, the walking speed, as well as the pedestrians’ activity (breathing or talking). Overall, the framework provides an unprecedented means to assess the risks of new infections via short-range exposure in arbitrary (real or hypothetical) crowd settings. Incidentally, while we have here chosen model parameters corresponding to SARS-CoV-2, the framework can easily be generalized to any pathogen with airborne transmission.

In the next section, the scientific context of the work with regard to airborne transmission is further clarified. Next, Section 3 describes our methodology, from the macroscopic model to assess the risk of new infections to the microscopic simulations of droplet transport. Section 4 then exposes the risks of transmission.

2. Scientific Context of the Study

Whenever one breathes, talks, pants, coughs, or sneezes, droplets of respiratory fluids possibly containing pathogens are expelled through one’s mouth and, to a much lesser extent, nose.\textsuperscript{[2–4,23]} In the case of COVID-19, the largest droplets thus produced had initially been thought to be liable for disease transmission. However, airborne transmission by inhalation of their smaller counterparts is now supported by robust evidence and was acknowledged by the World Health Organization (WHO) in Spring 2021, after months of controversy\textsuperscript{[12,24,25]} SARS-CoV-2’s ability to be transmitted via aerosols is now well established.\textsuperscript{[6–10]} It follows that closed, poorly ventilated spaces are particularly propitious for transmission,\textsuperscript{[18,26,27]} insofar as the smallest aerosols, of less than a few microns, can linger in the air for hours and accumulate in rooms. This opens the door for long-range airborne transmission, which cannot be avoided by social distancing. Nevertheless, airborne transmission may also occur at short distances, when a susceptible person inhales infected aerosols close to the emitter, where they are more concentrated. Large aerosols, with diameters $d_a$ up to 100 microns according to the recent literature,\textsuperscript{[2,10,25,28]} may also be inhaled before their sedimentation. The sedimentation speed $v_s$ in quiescent air may be estimated by balancing the gravity $g$ and drag forces at low Reynolds numbers. If one neglects the density of the air (of viscosity $\eta$) compared to that of the droplet, $\rho_d$, the Stokes law reads:

$$v_s \approx \frac{D_d \rho_d g}{18\eta}$$

A droplet of fixed diameter $d_a = 100 \mu m$ thus sediments at a speed $v_s \approx 0.3 \text{ m s}^{-1}$ (it will thus hit the ground in 5 s if it falls from a height of 1.5 m).\textsuperscript{[29]} To gauge whether it can be lifted up by an inhaling flow, bear in mind that the latter has a typical speed of a few tens of centimeters per second around the nostrils ($0.22 \text{ m s}^{-1}$ in ref. \textsuperscript{[30]}).

To what extent is the scenario of indoor transmission altered by outdoor settings? The most obvious difference is that aerosols are dispersed outdoors, which wards off the risk of long-range airborne transmission\textsuperscript{[2]} and ascertainment the mitigation efficiency of social distancing. On the other hand, the risks due to short-range exposure persist: one may inhale the small respiratory droplets emitted by a sick person in one’s immediate vicinity, the definition of which depends on the expiratory activity (for instance, uncovered sneezes propel droplets several meters ahead of the emitter$^{[1,13,11,32]}$). Besides, short-range exposure outdoors may differ from indoors because, all variations in temperature and humidity conditions left aside, it involves stronger wind and air flows. Note, however, that (moderate) air currents may also be worth considering indoors, where they are also present.\textsuperscript{[51]} In the context of the COVID-19 pandemic, transport by the wind has alternatively been thought to favor transmission by extending the spatial reach of droplets and to inhibit it by quickly dispersing pathogens.\textsuperscript{[14,31,34]}

In practice, for prevention policies, the risks incurred in crowded indoor environments have been highlighted by famous superspreading events.\textsuperscript{[7–10]} Outdoor infections have also been documented,\textsuperscript{[12,35,36]} but very generally looked down upon as secondary. Still, crowded outdoor settings are still listed among the risky configurations, for example, in WHO’s animation for public information (accessed in July 2022).\textsuperscript{[11]} In particular, mass outdoor events such as sports games have been suspected of promoting viral spread in periods of low viral prevalence,\textsuperscript{[37,38]} but the specific contribution of outdoor transmission in these
occurrences remains unclear, notably because many such events mix indoor and outdoor settings. In addition, retrospective studies may be biased toward an overestimation of the impact of specific large events, which are more closely monitored. Despite these difficulties, the question of the regulation of these events is vested with special interest, given their huge economic and social role; assessing the transmission risks that they present is thus of paramount importance to hit the right balance between public safety and social impact.

To this end, some randomized controlled trials have been conducted, in particular for indoor concerts, but general conclusions can hardly be reached from the small pool of such studies. Numerical studies provide a means to circumvent these limitations; indeed, their replicability enables researchers to test assumptions, investigate the effect of different parameters, relate behaviors to transmission risks and build a mechanistic picture of the risks in such contexts. The COVID-19 pandemic has prompted an unprecedented effort from the fluid mechanics community to probe the transport of respiratory droplets after their emission, in particular using CFD, which has shed light on the sensitivity of this propagation to the environment. Simulations have thus considered diverse environmental settings, as well as diverse expiratory events, including coughing, sneezing, speaking, and breathing. Coughs, in particular, have received a lot of attention, but in this paper we put the focus on talking and breathing through the mouth, because we have deemed that direct exposure to coughs (not covered by the emitter’s hand and directed toward the receiver’s face) is fairly rare and, in addition, talking for 1 min produces approximately as many droplets (i.e., a few thousand altogether) as one cough.

Risk assessment must also involve a model for inhalation. In simulations, specific areas (nose, mouth, eyes) that can be impacted or traversed by droplets may be marked in the simulation domain, in order to gauge the relative risks raised by droplet impact and inhalation or to quantify the protective effect of the exhalation of the susceptible person in a conversation. For instance. The inhalation volume of a passive scalar may also be used to assess the risk. Leaving aside inhalation, the local concentration of virus in a region of interest may be monitored, as a proxy for the infection risk, the need to simulate the susceptible person at each position that they may occupy is thus bypassed. Time may be involved by comparing the quantity of infected virus over time to an infectious dose via a dose-response model. This quantity can be measured in absolute terms, as a number of viral copies, which requires specifying the viral titer in the emitter’s respiratory fluids and the minimal infectious dose, or in terms of quanta of infection, if the number of emitted virus is rescaled by the infectious dose. In either case, an additional step is required to bridge the gap between such studies of very specific settings with CFD and a risk assessment at a larger scale in a variety of scenarios.

3. Methodology

Assessing the risks of viral spread via respiratory droplets in a crowd requires connecting the macroscopic configuration of the crowd and the activity of the attendants to the microscopic dynamics of droplet propagation. Here, we take up the method of our recent work to derive the number of new infections caused by an index patient on the basis of field data about crowds and mesoscale models of viral transmission (briefly recalled in Section 3.1), but here we aim to fully bridge the scales by anchoring the mesoscale models in a bona fide coarse-graining of microscopic simulations of droplet transport that take account of ambient air flows (see Sections 3.3 and 3.4), instead of resorting to mostly ad hoc models. The full algorithm is summarized in Appendix A.

3.1. Assessing Transmission Risks in a Crowd: General Principle

We will assess risks in a variety of crowd scenarios, each corresponding to a video recording collected and analyzed by García et al. For each scenario (stages, stations, markets, and more static scenes such as queues and street cafés), groups of pedestrians were tracked and, in the non-static scenarios, the infection risks within groups were discarded, assuming that a contagious individual is more likely to have infected the people walking in their company elsewhere.

For a given scene, one of the pedestrians, denoted by index \( i \) is supposed to be contagious and to expel virus-laden droplets. The algorithm is run once for each pedestrian of the scene to gather statistics. Under the independent action hypothesis, each inhaled virus has the same probability to cause an infection, independently of the others. The transmission risks, expressed as a number \( C_i^{(C)} \) of new cases that agent \( i \) transmitted to the pedestrians \( j \) that crossed his/her path in the interval \( [t_0, t_0 + \tau] \) during which he/she was filmed (his/her group \( G_i \) excluded, except at the cafés; there were no groups in the other static scenario, the waiting line at a screening center), can then be calculated using a Wells–Riley-like equation:

\[
C_i^{(C)} = \sum_{j \in G_i} S_j^0 \times \left( 1 - e^{-N_j} \right)
\]

Here, \( N_j = \int_{t_0}^{t_0 + \tau + \tau_{max}} v_j(t) \, dt \) is the cumulative transmission risk, with \( v_j(t) \) the instantaneous rate of transmission between the infected person \( i \) and a susceptible person \( j \). Manifestly, \( v_j(t) \) is the key quantity of the model and it will be the focus of the next paragraphs. \( S_j^0 \) is the probability that \( j \) is susceptible (i.e., not already infected) at the beginning of the observation interval. In principle, this quantity is not known. However, assuming that \( S_j^0 = 1 \) gives an upper bound on \( C_i^{(C)} \). Prior interactions with \( i \) in the scenario may already have caused \( j \)’s infection and thus reduced \( S_j^0 \) below 1, but only to a certain degree, consistent with the total number of new infectious caused by \( i \), which derives from \( C_i^{(C)} \); this criterion of self-consistency leads to a lower bound on \( C_i^{(C)} \). In practice, the lower bound is very close to the upper bound so in the following we shall not mention upper and lower bounds, but only estimated infection rates.

Finally, note that droplets emitted in the interval \( [t_0, t_0 + \tau] \) may take some time to reach a susceptible individual and be inhaled after the end of the interval; in practice, a maximum delay \( \tau_{max} = 21 \) s is imposed between emission and inhalation (we have ascertained that this maximum delay is large enough to not affect our quantitative estimates).
To do so, we suppose that on each frame in which $E$ is visible he/she emits a small set of droplets, whose evolution is then tracked for $t \in [t_i; t_i + \tau_{\text{max}}]$ or equivalently over time delay $\tau \in [0; \tau_{\text{max}}]$. For $\tau \leq \tau_{\text{max}}$, the spatio-temporal field of viral concentration that $R$ may inhale (depending on his/her position and head orientation) needs to be known; it will be expressed in the lab frame centered on $E$’s position at the instant of emission $t_i$, as a function of $\tau$, $r$, and $\theta_R$. Of course, the concentration map depends on several parameters: some of them can be extracted from the field measurements, such as the emitter’s walking velocity vector $\vec{v}_m$ and the orientation of the emission (or equivalently the angle between the head and the walking direction); others are unknown, namely, the wind velocity $\vec{v}_w$ (direction and magnitude) and the characteristics of the exhalation, and will be left as free parameters, whose effect will be assessed.

### 3.3. CFD Database: Parametrization

Were our computational means truly unbounded, we would run one computation for each set of the parameters and each micro-environment, and construct a concentration map for each of these. This is not computationally feasible in the real world. To circumvent this problem, we build a finite database of CFD simulations and concentration maps in a limited number of situations, from which the required maps will be derived through a suitable change of variables (under certain approximations) or interpolated.

Note, in particular, that the ambient flow around the emitter, which appears in the arguments of $\vec{v}$ in Equation (3), depends on the relative wind speed, but also on the details of the emitter’s surroundings, notably on the aerodynamic perturbations induced by the people around. In Appendix B, we explain that, while these perturbations do alter $\vec{v}$, their impact is moderate in practice for our risk assessments over the whole crowd. Therefore, the emitter’s surroundings will be discarded for the rest of the paper. This assumption is essential to allow the calculation of spatio-temporal field of viral concentration independently of the pedestrians’ configuration.

Let $(x, y)$ be the earthbound frame centered on the emitter $E$ at time $t_i$ and let $\vec{v}_m$ be $E$’s vectorial velocity (walking) at that time $t_i$ and $\vec{v}_w$ be the wind velocity; we denote $\vec{e}_m$ and $\vec{e}_w$ the corresponding unit vectors (directions). Note that the head direction $\vec{e}_h$ does not necessarily align with $\vec{e}_m$ in practice, for instance during a conversation, as illustrated in Figure 2a. We make the following assumptions:

1. The wind is uniform in space and constant in time during the relevant delay after the emission; velocity gradients in the height direction are neglected. As the relevant transport occurs at the height of human heads, neglecting the boundary layer profile is not expected to impact the results.
2. A walker’s motion is a plain translation in the walking direction, at constant speed; idiosyncrasies of the human gait are neglected.
3. The walking direction is aligned with the head orientation, so that $\vec{v}_{m} = \vec{v}_{w} \vec{e}_h$, as represented in Figure 2b. Our empirical data show that this assumption is inaccurate, but the associated angular differences are fairly small, with

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**Figure 1.** Snapshot of a single interaction filmed at a train station, with an alleged emitter $E$ and a receiver $R$. $r$ is the separation distance and $\theta_E$ and $\theta_R$ are the directions of emission and inhalation with respect to the line of connection.

In order to compare different scenarios, $C_i$ is recast into a rate of infections per hour: $C_i \triangleq C_i^{(\Delta T)} = \frac{\Delta t}{\Delta T}$ with $\Delta T = 1$ h, with the assumption that the recorded videos are representative. Note that in static scenarios (the cafes and waiting lines), the treatment is slightly different to account for the fact that interactions occur with a limited number of people, and always the same during the recording. Equation (2) is applied with $S_j^0 = 1$ (everyone except the infected person are susceptible) and the hourly rate is directly computed by using $N_{ij} = \int_{t_i}^{t_i+\Delta T} f(t) v(t) \, dt$.

### 3.2. Instantaneous Rate of Transmission between Two Individuals

Let us consider the droplets emitted by an infected person $i$ (the emitter $E$) and inhaled by another person $j$ (the receiver $R$). At time $t$, $R$ may inhale droplets emitted at different times. In the model, a double decomposition in time is performed. The emission time interval coincides with agent $i$’s observation period $[t_j, t_j + \tau]$, and for each emission at time $t_j$, droplets may be inhaled or “received” at time $t_e$. $e(t_j, t_j + \tau_{\text{max}})$.

More precisely, the instantaneous transmission rate due to droplets emitted at $t_j$ and inhaled at $t > t_j$ is expressed as

$$v(t_j, t) = T_0^{-1} \frac{\Delta t}{\Delta T} f(t_j) v(t) \int_{t_j}^{t_j+\Delta T} f(t) v(t) \, dt$$

where the characteristic time for infection $T_0 \propto n_{\text{inf}}/c_i$ is related to the specifics of the disease (namely, the minimal infectious dose $n_{\text{inf}}$ and the viral titer $c_i$ in the respiratory fluid, which makes it possible to account for a variable viral load), whereas the function $\hat{v}$ accounts for the fluid dynamics of droplet emission and transport. $r$ is the horizontal distance between the individuals’ heads and $\theta_E$ and $\theta_R$ are the orientations of the emitter’s and receiver’s heads, respectively, relative to the direction of the vector that connects them (see Figure 1). While simple ad hoc expressions for the function $\hat{v}$ were proposed in ref. [22], here we strive to compute $v(t_j, t)$ thanks to resolved CFD.
a standard deviation of 26°, and, above all, dwindle with the walking speed (see Figure 8), so much so that they reach the experimental uncertainty (19°)[22] for \( v_m \geq 1 \text{ m s}^{-1} \). Large deviations are mostly observed for static people, for which \( v_m \approx 0 \) and the assumption is theoretically justified. (We have checked that relaxing this assumption in favor of the opposite one, namely, aligning the emissions with the walking direction, leads to similar results, except in the static scenarios; see Figure D3).

Under such assumptions, the wind \( \vec{v}_w \) and the walk \( \vec{v}_m \) play a symmetric role: as far as air flows are concerned, walking at 1 m s\(^{-1}\) is equivalent to facing a head wind of 1 m s\(^{-1}\). Accordingly, CFD simulations are performed in the frame \( (x',y') \) attached to the emitter, located at \( (x', y') = (0, 0) \), with the basis vector \( \vec{e}_h \) coinciding with the head orientation \( \vec{e}_h \). This choice entails that the co-moving frame is rotated by an angle \( \theta_h = (\vec{e}_h, \vec{e}_m) \) and translated at a constant speed \( \vec{v}_m \), with respect to the lab frame.

In the co-moving frame, sketched in Figure 2c, the wind blows with a velocity \( \vec{v}_w' = (\vec{v}_w - \vec{v}_m) \); it is modeled by imposing a uniform velocity field \( \vec{v}_w' \), parallel to the ground, as a boundary condition in the far field. Note that \( \vec{v}_w' \) may result from the emitter’s motion, the wind, or from both of them. Finally, we denote \( v \) the angle between \( -\vec{v}_w' \) and the direction of emission \( \vec{e}_m \). Thus, \( v = |\vec{v}_w'| \) and \( \varphi \) fully parameterize the CFD database. For instance, in this database, the above situations in which an emitter walks at 1 m s\(^{-1}\) in still air and a static emitter faces a headwind blowing at 1 m s\(^{-1}\) both correspond to \( \varphi = 0 \) and \( v = 1 \text{ m s}^{-1} \).

### 3.4. CFD Database: Simulation Details

The CFD simulation protocol is detailed in Appendix B. In short, a still-standing manikin mimics a man who is breathing through the mouth, at a rate of 20 breaths of 1 L per minute, that is, one breath every 3 s, with an equal time for exhalation and inhalation. This signal was originally designed to replicate the breathing flow rate of a walking person, but also applies for speaking. Large-eddy simulations are performed along the same lines as refs. [22, 31], using the incompressible version of the Navier–Stokes equations, which was found to provide the best compromise between cost and accuracy of the simulations. Each simulation starts with three cycles to establish the flow, followed by four cycles during which statistics are collected.

Echoing Abkarian et al.,[31] we remark that the unsteady starting jets close to the mouth tend to form a main jet whose characteristics far from the mouth resemble those of a steady jet, with a limited influence of the details of the exhalation signal. This is why we use the same aerodynamic simulations to model mouth breathing and speaking. However, the number and sizes of emitted droplets will differ between breathing and speaking.

Overall, the database consists of 25 microscopic simulations for ambient relative velocity \( v (\text{m s}^{-1}) \in S_v \approx [0.1, 1.0, 2.0] \) and \( \varphi \in S_{\varphi} \approx [0, \frac{\pi}{4}, \frac{\pi}{2}, \frac{3\pi}{4}, \pi] \), plus the case \( v = 0.0 \text{ m s}^{-1} \), for which \( \varphi \) is undetermined (the simulations at \( v = 0.1 \text{ m s}^{-1} \) were used for control exclusively). The simulation cost increases with \( v \). For \( v = 2.0 \text{ m s}^{-1} \), a simulation takes more than \( \approx 80 \text{ h on 5 AMD EPYC Rome 7H12 bi-sockets nodes (640 cores) of the IRENE-AMD partition of Joliot-Curie cluster (TGCC/CEA, France). Contrary to alternative CFD works quantifying transmission risks in crowds, the choice has been made to use high-fidelity simulations to minimize the influence of modeling assumptions in the treatment of turbulence and of particle dynamics. Compared to other approaches using a passive scalar field to mimic the concentration of virus in the air,[21] it also offers the possibility to study larger droplets whose dynamics is not only determined by the flow. This choice is made possible by the assumptions of our approach, which limit the dimensionality of the parameter space (see Section 3.3)."

"Test-particles," that is, droplets of diameters uniformly distributed in \( d_0 \in [0.1 \mu\text{m}, 1 \text{ mm}] \), are injected into the airflow exhaled by the emitter. Importantly, the number of injected droplets (about 60 000 per breath) is not intended to be consistent with the empirical data for breathing [17,62], but merely to collect sufficient statistics in terms of particles behavior over a few cycles; since these droplets have very weak mutual aerodynamic interactions in the puff, at any reasonable concentration, this statistical contrivance will play virtually no role in the results.

Indeed, any actual distribution of emitted droplet sizes can be obtained by suitably resampling the simulated distribution, that is, assigning appropriate weights to the droplets depending on their size so as to match the desired distribution. In practice, following ref. [63] superpositions of log-normal distributions of
droplet diameters $d_p$, whose cumulative functions $P$ obey

$$dP(d_p) \propto \frac{1}{\ln\sigma} \times e^{-\frac{1}{2} \left( \frac{\ln d_p}{\ln \sigma} \right)^2} \, d\ln d_p$$

Breathing features only one such mode, with $D = 0.8 \mu m$ and $\sigma = 1.3$, whereas vocalization (speech) features one mode at $D = 0.8 \mu m$ and $\sigma = 1.3$, multiplied by a coefficient 0.069, and one mode at $D = 1.2 \mu m$ and $\sigma = 1.66$, multiplied by a coefficient 0.085. The third mode associated with speech is peaked at $D = 217 \mu m$, with $\sigma = 1.795$ and a coefficient 0.001; it thus corresponds to large droplets unlikely to be inhaled (see Section 4.5). The coefficient associated with the breathing mode is found by recalling that breathing produces $\approx 20$ times fewer droplets than normal speech.\(^{[27]}\)

Note that the foregoing sizes correspond to those measured by Johnson et al.\(^{[63]}\) prior to the application of their corrective factors, which notably account for evaporation. Indeed, small respiratory droplets are expected to undergo quick partial evaporation, which makes it sensible to consider their propagation with their evaporated diameters; besides the (slight) effect on droplet transport, we expect no further incidence of the application of a constant corrective factor on the diameters, thanks to our renormalization with the characteristic infection time.

### 3.5. Coarse-Grained Dynamic Maps of Viral Concentration

Once resampled, the detailed configuration of the simulated droplets is coarse-grained into dynamic maps of viral concentration $c(r, \theta, \tau)$, where $(r, \theta)$ are polar coordinates in the earth-bound frame centered on the emitter’s mouth at the instant of emission $t_e$, and $\tau$ is the delay since emission of the droplets. $\theta = 0$ is the direction of emission. These maps $c(r, \theta, \tau)$ are obtained by binning droplets in space and time, with a resolution $\Delta r = 20 \text{ cm}$ on $r$, $\Delta \theta = \frac{\pi}{12}$ on $\theta$, and $\Delta \tau = 0.2 \text{ s}$ on the delays $\tau$, that is, the lifetimes of droplets, and then computing the total volume of droplets in each spatiotemporal cell, within a 40-cm-thick horizontal slice centered on the emitter’s mouth (the yellow box displayed in Figure 3), and dividing it by the cell volume. This relies on the assumption that viral copies are homogeneously distributed in respiratory fluids and each raises the same risk of infection (regardless of the droplet size), which is classical in modeling but possibly underestimates the viral load in small particles.\(^{[2,28]}\) The resulting maps are then symmetrized with respect to the $\theta = 0$ axis ($\bar{c}_0$), if such a symmetry is expected to hold, that is, for head and tail winds. Finally, for any relative wind velocity $\vec{v}$, the emitter can be assigned a walking speed $v_w$, without additional CFD simulations, by simply translating the origin of the concentration maps with the time delay $\tau$ at a speed $v_w$ in the direction opposite to the emitter’s head orientation, that is, along $-\vec{c}_0$. Note that the question of the normalization of the concentration maps, that is, their scale, will be circumvented by setting a characteristic infection time $T_{in} = 15 \text{ min}$ for someone standing face-to-face, a distance $r = 50 \text{ cm}$ away from a static speaking emitter. In other words, the quantum is defined as the quantity of virus inhaled in 15 min while standing 50 cm away from a static speaking person, without wind; this volume of droplets is used to normalize the transmission risks.

### 4. Results: Transmission Risks Generated by an Emitter

In this section, we inspect the spatiotemporal pattern of risks (i.e., virus-laden aerosols) emitted by an index patient, depending on the environmental conditions.

#### 4.1. Propagation of the Droplets Simulated with CFD

Figure 3 shows the propagation of an arbitrary number of droplets of less than 10 microns in diameter, emitted by the

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**Figure 3.** Examples of the variation of the exhaled puffs with the wind and the walking speed. (Top row) Photographs. (Bottom row) CFD simulations, displayed at time 0.75 s of the sixth cycle, when the exhaled flow is close to its maximum. Droplets of less than 10 microns in diameter are displayed to mimic tracer particles. They are colored in blue for those exhaled over the sixth cycle (0.75 s before and less) and in red for those exhaled during cycles 1 to 5. The region with a yellow overlay materializes the box within which droplets will be counted (see the main text). First column: exhalation in still air; second column: exhalation during walk (at 1.0 m s$^{-1}$ in the CFD) in still ambient air; and third and fourth columns: exhalation in headwind and crosswind, respectively (at 2.0 m s$^{-1}$ in the CFD). Pictures are illustrative, as walking speed and wind speed do not match, and exhalation has not been characterized.
manikin in the CFD simulations, for distinct incident velocities ($v = 0.0, 1.0, \text{ and } 2.0 \text{ m s}^{-1}$) and angles $\varphi$. Interestingly, both the relative wind (generated by walking) and the external wind have a dominant effect on the propagation, after a short first regime during which the direction of emission prevails. This holds true even at very low wind speeds, which would not even be qualified as “light breeze” ($6–11 \text{ km h}^{-1}$) on the Beaufort scale. For example, a wind of $1.0 \text{ m s}^{-1}$ ($3.6 \text{ km h}^{-1}$) corresponds to the air flow felt when walking in a still environment. To illustrate the consistency of these numerical results, one subject gave his consent to be photographed while smoking an e-cigarette. This experiment consisted in several exhalations indoors while walking or not, and outdoors in the wind, whose speed was not measured. Thus, the walking speed and the wind speed could not be matched between the experiments and the simulations; the comparison mostly has an illustrative purpose. The whole experiment lasted a few minutes and did not lead to a substantial change in the subject’s consumption of his e-cigarette.

Although qualitative, the comparison highlights the singularity of the case without relative wind (left), where a long jet may develop without being perturbed. Another observation is the similarity of the pictures/results obtained while walking in still air (second column) or being static in headwind (third column): the spatial extent of the puff in front of the subject is substantially smaller than in the static case without wind (fourth column) and the puff disperses behind the subject’s head. We observe that the wind may transport droplets farther over shorter periods of time, but we will need to turn to concentration maps to determine if this heightens transmission risks.

### 4.2. Dynamic Maps of Viral Concentration

Following the coarse-graining method exposed in Section 3.5, the detailed output of the CFD simulations is converted into dynamic maps of viral concentration, that is, spatiotemporal diagrams of risks centered on the emitter at the moment when droplets are shed.

We first comment on the maps obtained at zero walking speed $v_w$, shown in Figure 4. The values displayed in these diagrams, say at position $(r, \theta)$ and after a delay $\tau$, for a relative wind $\vec{v}_w$, can naturally be interpreted as the risks incurred by a non-moving receiver located at $(r, \theta)$ relative to static emitter under an external wind $\vec{v}_w$, up to an inhalation coefficient, but let us mention that it also describes the situation in which the emitter and the receiver are walking at the same velocity $\vec{v}_m$ when the wind blows at $\vec{v}_w = \vec{v}_w + \vec{v}_m$. We notice, once again, that in the first fractions of a second the puff tends to follow the line of emission, but it is then steered by the wind, while remaining fairly compact, and it is swept more than $2 \text{ m}$ away from the emitter in a matter of seconds at any finite wind speed ($v \geq 0.3 \text{ m s}^{-1}$), compared to more than a dozen seconds in windless conditions. This further underlines the singularity of the windless case as far as one is concerned with droplet transport.

Not surprisingly, maximal risks are incurred in the immediate vicinity of the emitter in the time-cumulative diagrams (where the peak for $r = 20 \text{ cm}$ away from the emitter is shown as a small yellow dot), but the azimuthal position is influenced by the relative wind. For example, it lies behind the emitter for a head wind blowing at $1.0 \text{ m s}^{-1}$ (Figure 4 (bottom)).

Besides, it is now apparent that the major effect of the wind, besides steering the puff, is to quickly disperse the emitted droplets and thus to lower transmission risks. We illustrate this in Figure 5 in the case of a side wind by plotting the radial decay of the maximum concentration of viral particles over all azimuthal directions for different wind speeds. A wind speed as low as $0.3 \text{ m s}^{-1}$ reduces the peak value at $50\text{-cm distance}$ by a factor of three, roughly speaking. Even the slightest, almost imperceptible draft, at $v_w = 0.1 \text{ m s}^{-1}$, has a significant effect on the transmission risks. This demonstrates how singular are the stagnant air conditions ($v_w = 0$) often used to model droplet transport. The local maximum observed in the $v_w = 0$ case is notably due to inhalation, which removes the last droplets exhaled from the near-mouth region. Such droplets are swept out in the presence of wind.

Let us now make the emitter move while breathing or speaking. Figure 6 shows that the air flow generated by walking drags the puff forward, along the emitter’s path; similarly to an external wind, this drag also tends to sweep away the emitted droplets. Incidentally, recall that the concentration maps are shown in the Earth’s frame, and not in the walker’s co-moving frame, which can explain why no clear detachment transition of the puff is observed as the walking speed increases.\[64\]

### 4.3. Robustness to Simulation Details Related to the Flow Dynamics

In view of the many recent reports highlighting the sensitivity of droplet transport to physical and numerical details, before proceeding with the assessment of risks, we examine the influence of these details on our results (more information in Appendix B, notably in Figure B4).

i) Moderate variations of droplet sizes have little impact on droplet propagation. Indeed, the concentration maps obtained for subsets of droplet sizes between 0.5 and 20.0 $\mu$m are qualitatively very similar; this is due to the fact that results are gathered on regions of 40 cm height, while sedimentation velocity of 20.0 $\mu$m droplets is of the order of the centimeter per second.\[29\] In addition, the Stokes number remains small for such droplets, which means that particles essentially follow the air flow.

ii) Buoyancy effects, due to the different temperature of the puff compared to the ambient air, have an impact, but a moderate one, as we see when these effects are introduced into CFD simulations. The buoyancy generated by the thermal plume surrounding human bodies (which are hotter than the environment)\[65\] has not been directly simulated. As stressed by Nielsen and Xu in a recent review,\[66\] the air inhaled indoors mainly comes from the lower part of the body and is lifted to the nose by the thermal plume. To quantify its effect, we have compared the reference concentration maps with their counterparts measured when the region of interest is shifted downward by 20 cm, with limited differences.

iii) Evaporation does not affect the results much. Crucial in this lack of incidence is the fact that, because of their finite fraction $c$ of non-water content, droplets do not fully evaporate,
but shrink into residues (also called droplet nuclei), of final size of the order of $c^1$ times their initial size.\cite{29,67}

All these effects are already weak with no external wind, but are even further reduced in the presence of wind, which always ends up dominating transport far from the mouth.

4.4. Interpolation of Spatiotemporal Diagrams

Given that the CFD database only contains a limited number of cases $(v, \phi) \in S_v \times S_\phi$, interpolation is needed to obtain the spatiotemporal diagram corresponding to given relative wind conditions $(v, \phi)$ at a delay $\tau$. We select the two closest values $\phi_1$ and $\phi_2$ ($\phi_1 \leq \phi \leq \phi_2$) in $S_\phi$ and $v_1$ and $v_2$ ($v_1 \leq v < v_2$) in $S_v$. Now, since $\phi$ controls the rotation of the exhaled air puff under the wind and $v$ affects the propagation dynamics, a naïve linear interpolation over $\phi$, and $v$ would perform poorly.

Instead, noticing from Figure 4 and its kin that the puff is first transported in the direction of emission, and then in the wind direction, we handle (very) short delays $\tau \leq \tau^*(v) \approx 0.12/v$ distinctly from longer delays $\tau > \tau^*(v)$. For the former, the directions of emission are already aligned with $\vec{e}'_x$ in the spatiotemporal diagrams, so no rotation is needed. For the latter, the diagrams corresponding to $\phi_i$, $i = 1, 2$, will be rotated by an angle $\phi - \phi_i$ prior to interpolation.

Turning to the speed variable $v$, noticing that (for a given wind direction) increasing $v$ has an effect somewhat comparable to “fast-forwarding the movie,” that is, inspecting the diagram at a shorter delay $\tau$, the diagrams corresponding to speeds $v_i$, $i = 1, 2$, are probed at delays $\tau_i = \tau v_i/v$ (if $v_i > 0$; $\tau$ otherwise) and their values (which represent the transmission risk over a fixed time interval $\delta\tau$) are rescaled by multiplication with $v_i/v$.

To sum it up, for each $v_i$, we interpolate linearly between the diagrams corresponding to $\phi_1$ and $\phi_2$, after suitably rotating them if $\tau > \tau^*$, and linear interpolation between the resulting diagrams at $v_i$ and $v_j$ yields the final diagram. No interpolation is needed on the walking speed $v_m$ because, from the CFD output at $(v, \phi)$, we are able to generate and store dynamic maps for a wide range of walking speeds (in practice, $v_m = 0, 0.1, 0.2, ..., 2$ m s$^{-1}$). The example shown in Figure 7 (and Figure D2), in which a genuine

![Figure 4. Dynamic maps of viral concentration associated with talking: Effect of the external wind. (Top row) no wind, (middle row) lateral wind blowing at 0.3 m s$^{-1}$, and (bottom row) head wind blowing at 1 m s$^{-1}$. The emission is arbitrarily aligned along the x-axis and the color bar saturates at the arbitrarily imposed top value.](image-url)
Figure 5. Radial decay of the maximum concentration of viral particles emitted by a static pedestrian over all azimuthal directions, in the cumulative concentration maps associated with speaking. An external wind is assumed to blow perpendicularly to the head orientation ($\varphi = \frac{\pi}{2}$), at the speed $v_w$ indicated in the legend. The concentrations were normalized to one at a distance $r_c = 50$ cm in windless conditions.

concentration map computed for $\varphi = \frac{\pi}{4}$ is compared to its interpolated counterpart, demonstrates that this method produces quite reasonable results.

4.5. Inhalation and the Case of Large Oral Droplets

Once respiratory droplets have been emitted, they must be inhaled by a receiver to bring on a risk of transmission. So far the dynamic concentration maps have been established irrespective of the receiver and her head orientation. We now take care of the latter by inserting a multiplicative factor $v_R$ accounting for inhalation in the transmission rate function $\hat{v}$ in Equation (3), on top of the concentration at the receiver’s location. We consider two possibilities; in the first one the incident puff can be inhaled only if it “hits” the side of the receiver’s head containing the mouth and nose, hence

$$v_R(\theta_R) = \begin{cases} 1 & \text{if } \theta_R \in [-\frac{\pi}{2}, \frac{\pi}{2}] \\
0 & \text{otherwise} \end{cases}$$

where $\theta_R$ was defined in Figure 1. The second option, isotropic inhalation, is inspired by the steady-flow situation in which the concentration becomes homogeneous all around the head, including the “dead-waters” zone located downstream, so that

$$v_R(\theta_R) = 1$$

In doing so, we assume that the breathing activity of the receiver has little incidence on the transport of the droplets produced by the emitter. While it is true that under specific face-to-face conditions the receiver’s expiration can significantly perturb the emitter’s expiration flow (and possibly act as a shield), more generally, this assumption sounds reasonable, especially during the receiver’s inhalation, provided that the emitter is not too close to the receiver.

Besides, the droplets must be small enough for inhalation to be possible. This will always be the case for the breathing and vocalizing modes in the micron range, but the question is worth discussing for the third mode (“oral” mode), peaked at 217 µm. Indeed, the typical sedimentation speed for a droplet of diameter $d_p = 100$ µm (resp. $d_p = 217$ µm) is $v_g \approx 0.3$ m s$^{-1}$ (resp. 1.5 m s$^{-1}$), which is larger than the magnitude of inhalation velocity measured by Murakami, for instance. This is confirmed by our

Figure 6. Dynamic maps of viral concentration associated with speaking (in the lab frame): Effect of the walking speed in windless conditions. Walking speed $v_m = (\text{top})$ 1.0 m s$^{-1}$, (bottom) 2.0 m s$^{-1}$. It should be recalled that the color bar saturates at the arbitrarily imposed top value.
own simulations of nose breathing presented in Appendix C (see Figure C2 in particular). Therefore, inhalation of the droplets of that mode is deemed rather unlikely and in the following we will focus on the smaller droplets. Nevertheless, in Appendix C the possibility to inhale these larger droplets is restored and leads to a distinct global picture.

5. Results: Risks of New Infections at the Macroscale

Moving on to macroscopic crowds, we now couple the coarse-grained dynamic maps of viral concentration obtained in the previous section with field data about crowds in daily-life situations.

5.1. Empirical Crowd Dynamics

We begin with a presentation of the empirical scenarios that will be used as test-cases in our macroscopic risk assessments. They are listed in Table 1 and include fairly busy streets (and riverbanks), metro and train stations, an outdoor market, and street cafés. All scenarios are in outdoor settings or in large, well ventilated areas and the data were collected in the metropolitan area of Lyon, France, during the COVID-19 pandemic, between July 2020 and January 2021. Note that these data are openly available on the Zenodo platform.[68] More details about the data acquisition and pedestrian tracking protocols can be found in Garcia et al.[22] In short, the pedestrians’ positions and head orientations are marked on the videos every 0.5 s; the temporal resolution is then increased to a point every 0.1 s through linear interpolation. By double tracking pedestrians, the experimental uncertainty on the absolute positions was estimated to below or around 20 cm while the error on the head orientations had a standard deviation of 19°. The accuracy of the angular data is well evidenced by our ability to capture the tendency of pedestrians to look more and more straight ahead as they walk faster, as reflected in Figure 8 by the variation of the standard deviation of \( \delta \theta \), the angular difference between the head orientation and the walking direction, with the walking speed.

Finally, to compensate for the narrow field of view and the interactions thus missed, a reweighting process, based on an estimation of the number of missed contacts, was proven to effectively correct the bias toward shorter-ranged interactions.[22]

5.2. Different Perspectives for the Assessment of Risks

We now apply the methodology exposed in Section 3 to the field data, recalling that the static scenarios (queue and street cafés) are handled slightly differently from their moving counterparts: People are assumed to keep interacting with the same neighbors over the whole period \( \Delta T = 1 \text{ h} \) in the former, whereas in the latter they will interact with new people. Moreover, in the moving scenarios, the risks of infecting one’s co-walkers are overlooked, because we are interested in the supplemental risks generated by the scenario under consideration and the co-walkers probably interacted with the index patient in more risky places, such as enclosed settings; all the other people are considered susceptible.[22,69] By contrast, no social groups are taken into account at street cafés.

Besides, it should be underlined that risks are here quantified by the number of new cases \( \overline{C}(\Delta T) \) expected in each setting when an index patient is present on the premises for a duration \( \Delta T = 1 \text{ h} \), and not the total rate of new infections in the scenario or the risks incurred by a typical attendant. We claim that this perspective, centered on the infected person, is the relevant one at the collective scale, for policy-making: It
| Scenario / Location                                      | GPS coord (WGS-84) | Date and time | Comments                      | #ped | Density ($\text{m}^{-2}$) | Snapshot |
|--------------------------------------------------------|---------------------|---------------|-------------------------------|------|--------------------------|----------|
| Pedestrian banks of the Rhône river                    | 45.7685, 4.8406     | July 2020     | Sunny and windy day           | 164  | 0.042                    |          |
| Square in front of Perrache hub                        | 45.7476, 4.8250     | July 2020     | Nice weather                  | 1021 | 0.038                    |          |
| Shopping street (Grande rue de la Croix Rousse)         | 45.778, 4.8322      | January 2021  | Cold and cloudy day           | 420  | 0.06                     |          |
| Busy street (rue du Dr. Bouchut)                       | 45.7595, 4.8545     | July 2020     | Hot and sunny day             | 800  | 0.05                     |          |
| Saint-Jean street in the Old Town                      | 45.7622, 4.8275     | Sept. 2020    | Hot and cloudy day            | 481  | 0.11                     |          |
| COVID-19 testing site                                  | 45.7241, 4.8281     | October 2020  | Outdoor and indoor waiting lines (cold day) | 66   | /                        |          |
| Part-Dieu train station - Ground level                 | 45.7607, 4.8596     | July 2020     | Indoors                       | 875  | 0.22                     |          |
| Bellecour subway station - Platform                    | 45.7576, 4.8336     | July 2020     | Indoors                       | 849  | 0.26                     |          |
| Main alley of an outdoor market                        | 45.7739, 4.8270     | October 2020  | Sunny Sunday                  | 183  | 0.46                     |          |
| Street cafés and restaurants on Place des Terreaux     | 45.7677, 4.8337     | October 2020  | Sunny day, cool weather       | 30   | /                        |          |
| Street cafés on Croix-Rousse boulevard                 | 45.7745, 4.8308     | September 2020| Nice weather                  | 13   | /                        |          |

enables the decider to compare the infection potentials of the different activities in which an infected person would engage. In this sense, and contrary to the total rate approach, a massive gathering will be deemed to present higher risks than a number of smaller gathering only if it leads to higher $\overline{C}(\Delta T)$ than in the smaller gatherings. Compared to the perspective centered on the receiver, it is true that, if a given fraction $\gamma$ of the crowd is contagious (irrespective of the scenario), the infecter-centered risk assessment also reflects the average risk per hour incurred by an individual in the crowd, $\gamma \overline{C}(\Delta T)$, but the two perspectives
may have very dissimilar distributions (for instance with a low risk for a large number of people or a high risk for a small number of people). We put the focus on the infecter-centered risk assessment.

5.3. Rate of New Infections in Perfectly Windless Conditions

Figure 9a shows the rates of new infections in windless conditions for the different scenarios, when all pedestrians are supposed to be constantly talking. Cafés present the highest risks by far, followed by the outdoor market and, further down, the metro and train platforms and halls (filmed in the midst of the pandemic), whereas the risks raised by fairly busy streets are comparatively quite low.

Reassuringly, these trends are in line with those found previously on the basis of various ad hoc models, which totally discarded the relative winds generated by walking, among other aspects.\[22\] But, here, the account is more quantitative, given that

![Figure 8. Standard deviation of the difference between the head orientation $\vec{e}_h$ and the walking direction $\vec{e}_m$, as a function of walking speed.](image)

![Figure 9. Estimated risks of infection associated with speaking in each scenario, for different wind speeds. Pay attention to the widely different scales in each panel.](image)

![Figure 10. Estimated mean risks of infection in each scenario under an external wind blowing at $1 \text{ m s}^{-1}$, for a) speaking, b) speaking with an isotropic inhalation coefficient and c) breathing through the mouth. Pay attention to the widely different scales in the panels. The reader is referred to Table 1 for the correspondence between colors and scenarios.](image)
our transmission models are rooted in fluid dynamical models; the only adjustable variable is the characteristic time of infection $T_0$ (once rescaled by this time, the rates of infections vary little with $T_0$, within reasonable bounds).

Switching to breathing through the mouth instead of talking does not affect the ranking in the slightest way (Figure 10), which makes sense, given the relative insensitivity of droplet transport to small variations in droplets sizes (Figure B4). But it dramatically lowers the risks, by a factor of order 100, consistently with the lower volume of respiratory droplets produced in this case.

In reality, people will carry out a mix of respiratory activities; the risks should then be computed by an average of the risks raised for each type of activity, weighted by the proportion of time spent for each activity; breathing through the nose may be considered to raise no risks. In practice, this weighting will further enhance the risks associated with cafes, insofar as talking will probably occupy a larger fraction of time in this scenario than in the other ones.

### 5.4. Effect of Modest Winds or Ambient Air Flows

Introducing an external wind alters the foregoing picture to some extent (Figures 9 and 11). Most strikingly, the absolute levels of risks are strongly depressed, for example, by a factor of 4 in the case of the market, for a wind speed of $v_w = 2 \text{ m s}^{-1}$. This is in line with the dispersal effect of wind established in Figure 5.

Focusing on specific scenarios in Figure 11, we observe that the risks are systematically lowered with wind speed in the scenarios which display significant levels of risks, whereas for example in fairly busy streets this mitigation effect becomes clear only beyond a finite wind speed; below this value, the drag due to the wind (which carries droplets farther, toward other groups of people) tends to compensate the dispersal effect, because the crowd is not so dense.

Besides, the risk gap between street cafes and the outdoor market lessens with increasing $v_w$, so much so that the two scenarios become about as risky under external winds blowing at 2 m s$^{-1}$, which is still calm air. (Recall however that this comparison only holds if similar expiratory activities are performed in all scenarios; otherwise, the activities should be reweighted; see Figure 10). This effect is easily understood as the wind bends the particularly unfavorable propagation of droplets in the case of face-to-face conversations, and favors transmission in isotropically dense settings. The distribution of rates of new infectious caused by the different individuals that were filmed is presented in Figure 12.

How robust are these results to variations in the simulated conditions? In principle, at equal wind speed, the wind direction is not expected to impact the results. This is verified in most scenarios, but not all. Some effect is observed for one of the street cafes and the queue at the screening center: In these cases, the specific settings that we filmed displayed preferential directions (whether it is the alignment of the table or that of the queue), which may couple with the wind direction.
Turning to the directional dependence of inhalation, the anisotropic inhalation coefficient $\nu_\theta$ given by Equation (5) was used so far. Replacing it with a fully isotropic one, Equation (6), tends to enhance the risks (Figure 10b), since it allows more directions for inhalation. This is most acute in the windless queuing scenario, where the face-to-back alignment of people in the queue used to make inhalation inefficient.

5.5. A Practical Guide to Assessing Risks in Real or Test Scenarios

We close this section with a succinct summary of the practical way in which risks of viral transmission can be assessed using our framework:

i) Collect or simulate the $(x, y)$-trajectories of people in the scenario; the head orientations can be measured or assumed to align with the walking direction;
ii) If needed, interpolate between points in order to increase the temporal resolution;
iii) Feed these trajectories into the Python scripts (which can be found under https://github.com/an363/InfectiousRisksAcrossScales) coupling them to the dynamic concentration maps to get the mean rates of new infections $C(a, T)$ for each expiratory activity and wind velocity under study;
iv) Estimate the fraction of time $\alpha_t$ spent talking and the fraction of time $\alpha_b$ spent breathing through the mouth;
v) Compute the rate of new infections as a weighted average\textsuperscript{[70]} of the $C(a, T)$ over the expiratory activities $a$ (if talking is not negligible, then $C(a, T) \simeq C(\alpha_t, T)$) at the typical external wind speed $\nu_w$, or averaged over the distribution of different wind speeds and wind directions.

6. Conclusions

In summary, we have put forward and implemented a methodological framework to assess the risks of viral transmission via short-ranged exposure in crowds. It provides an unprecedented connection between the fluid dynamical propagation of respiratory droplets at the microscale and field-data about macroscopic crowds, using spatio-temporal maps of viral concentration.

These concentration maps give insight into the transmission risks in binary interactions, highlighting the paramount effect of the respiratory activity, owing to the much larger volume of respiratory droplets expelled while talking, as compared to mouth-breathing; the effect of (even very modest) winds or ambient air flows is also apparent. The impact of air flows had been reported previously in case studies of specific (mostly indoor) settings\textsuperscript{[33,50,52,71–73]} its generality is underscored here. Our study further shows how the walking velocity of pedestrians also contributes to decreasing the risks, through the same mechanism.

By coupling these binary transmission risks with pedestrian trajectories, the rates of new infections generated by a contagious individual wandering for 1 h on the premises was generated. We put this to the test using field data collected in diverse daily-life scenarios, which were thus ranked by risks. Consistently with our previous findings with much coarser models, street cafés present the highest risks among the investigated scenarios in windless conditions, due to the configuration of the crowd (even if the larger fraction of time spent talking is left aside), followed by the observed outdoor market and, further down the list, train and metro stations (at the peak of the pandemic). However, our finer models also stress the dramatic quantitative effect of the wind on these results, which strongly depresses transmission risks and tends to reduce the gap between street cafés and the busy outdoor market. The work thus contributes to explaining why overall outdoor settings appear to raise substantially fewer risks of viral transmission than enclosed spaces, besides the negligible risk of long-range air-borne transmission in non-confined settings.

Note that the critical influence of air currents even at low speed also urges one to reconsider short-range transmission risks indoors. Indeed, indoor drafts have a typical speed of a few tens of centimeters per second,\textsuperscript{[24]} which implies that they can dominate droplet propagation after the exhaled puffs lose their initial momentum. This should be taken into account in microscopic studies of droplet transport, which are usually performed in stagnant air.

To conclude, on the bright side, it is worth underlining the generality of the proposed framework. The approach is on no account restricted to COVID-19 and it enables one to test a diversity of scenarios, for example, to explore the efficiency of redesign strategies aimed at mitigating viral spread, in a stadium or at any other mass gathering. All this is brought within reach by the reduction of the computational time by several orders of magnitude for a given scenario, thanks to the recourse to coarse-grained viral concentration maps. This expedites the assessment of risks, compared to alternative numerical methods based on CFD, such as the one proposed by Löhner et al.,\textsuperscript{[21]} for which each test necessitates a full simulation. An essential assumption to reduce the computational burden is to neglect interactions between the emitter and the environment, allowing the computation of viral concentration maps independently of the neighboring pedestrians. Thanks to the reduced computational burden, any technical service has the possibility to test different scenarios or redesigns and assess the risks that they raise with strictly minimal computational needs. This may be particularly relevant for screening purposes, that is, to short-list suitable options, possibly prior to a deeper inspection with finer approaches,\textsuperscript{[21]} notably in which the influence of the local environment (including the other pedestrians) on the transport of infectious material is directly accounted for.

On a less positive note, we must plainly acknowledge that our approach, which provides a unique way to span six orders of magnitude in lengthscales in order to address transmission risks, rests on several serious approximations. Relaxing them would overcome some limitations of the model, possibly at the expense of more costly simulations. First, the CFD simulations could be refined to more accurately reflect the exogenous turbulence of the wind (before it interacts with the manikin’s body and the expiration flow). Along similar lines, the influence of the receiver’s exhalation and inhalation flow on the transport of droplets generated by an emitter\textsuperscript{[49]} could be taken into account, even though this may require many more CFD simulations. In that case, a compromise may be to lower the level of fidelity of the CFD simulations in order to span a larger parameter space.
Height differences between people could also be incorporated, and should be so if the crowd mixes standing and seated people. As a final word, we recall that the framework is well suited for outdoor configurations, but was not designed for confined settings featuring strong variations of airflows in space and/or vertical motions due to ventilation, for instance. In such cases, full CFD simulations remain the most versatile approach.

Appendix A: Algorithm for the Calculation of Transmission Risks

Algorithm A1 presents a pseudo-code for the risk assessment from a series of images. One person is considered as an index patient in the scenario, then the risk is calculated for each of the other people in the movie potentially interacting with i. i is then varied for ensemble averaging. The wind velocity $\vec{v}_w$ is a free parameter that has not been measured.

Appendix B: Numerical Method and Simulations Details

This appendix details the microscopic numerical simulations used in the paper.

Algorithm A1 Algorithm to assess the transmission risk associated with one contagious person in a movie

B.1 Physical Model and Flow Solver

3D numerical simulations are performed in the idealized case of non-buoyant jets, neglecting temperature effects. To account for turbulence, we use large Eddy simulations (LES) as reported and described in ref. [31]. The fluid flow is governed by the incompressible versions of the filtered continuity and Navier–Stokes equations

$$\frac{\partial \tilde{u}_i}{\partial x_i} = 0$$  \hspace{1cm} (B1)

$$\rho \frac{\partial \tilde{u}_i}{\partial t} + \rho \frac{\partial \tilde{u}_i \tilde{u}_j}{\partial x_j} = -\frac{\partial \tilde{p}}{\partial x_i} + \mu \frac{\partial^2 \tilde{u}_i}{\partial x_j^2} + \frac{\partial \tau_{ij}}{\partial x_j}$$ \hspace{1cm} (B2)

where $\tilde{u}_i$ is the filtered fluid velocity component in the $i$th direction, $\tilde{p}$ the filtered pressure, $t$ the time, $x_i$ the spatial coordinate in the $i$th direction, $\rho$ the constant air density, and $\mu$ the constant dynamic viscosity. $\tau_{ij} = \rho(u_i u_j - \tilde{u}_i \tilde{u}_j)$ is the residual stress-tensor coming from the subgrid-scale unresolved contribution, for which a closure needs to be provided. Here we use the so-called sigma model which has notably been built to yield zero extra dissipation in laminar flows, so that it is well adapted to
situations at moderate Reynolds numbers where transition to turbulence occurs,\cite{76,77} which is the case here.

The considered fluid is assumed to represent air at constant ambient temperature. The kinematic viscosity is fixed at $\nu = \mu / \rho = 1.5 \times 10^{-5}$ m$^2$ s$^{-1}$.

For the present work, we used the flow solver YALES2.\cite{31,77,78} The fluid equations are discretized using a fourth-order finite-volume scheme, adapted to unstructured grids.\cite{79,80}

Exhaled droplets are represented as spherical Lagrangian particles, tracked individually with a point-particle Lagrangian approach. One-way coupling is used as the concentration of droplets is small (of the order of a few particles per liter\cite{17}). The droplet motion is obtained by advancing them along the flow

$$\frac{dx_p}{dt} = \bar{u}_p$$

where $x_p$ is the position of the droplet and $\bar{u}_p$ its velocity.

Conservation of momentum is given by Newton’s second law:

$$\frac{d}{dt} (m_p \bar{u}_p) = \bar{F}_p^C + \bar{F}_p^D$$

where $m_p$ is the mass of the droplet, $\bar{F}_p^C$ is the buoyancy force and $\bar{F}_p^D$ the drag force. The buoyancy force and drag force read:

$$\bar{F}_p^C = (\rho_p - \rho) \frac{d_p^3 \bar{g}}{6}$$

and

$$\bar{F}_p^D = m_p \frac{1}{\tau_p} (\bar{u}_p - \bar{u})$$

where $\rho_p$ is the droplet density, $\rho$ the local gas density, $d_p$ the droplet diameter and $\bar{g}$ the gravitational acceleration. $\tau_p$ is the characteristic drag time. It is modeled with the empirical correlation of Schiller and Naumann\cite{81} for moderate values of the droplet Reynolds number. This correlation tends to the Stokes law at low Reynolds numbers.

**B.2 Computational Domain and Grid**

The computational domain is a 3 m-high, 6-m long box in the wind direction; its width is 4 m. The manikin’s head is oriented in the $x'$ direction and the ambient wind blows in a variable direction as a function of the incident angle $\varphi$ (see Figure 2).

The grid is initially refined around the head of the manikin with a spatial resolution of 1 mm and coarsened further away. A dynamic mesh adaptation algorithm is used to refine the grid wherever needed. To do so, a passive scalar is injected at the mouth. Any location where the concentration of this passive scalar is non-zero is identified as a meaningful region and the grid is subsequently refined during the calculation, with a target grid size of 8 mm. Figures B1 and B2 prove that CFD simulations with a finer grid size of 4 mm yield virtually identical coarse-grained concentration maps.

**B.3 Boundary Conditions**

The CFD database is parametrized by the incident air speed $v$ and the angle $\varphi$ between minus the incident velocity vector and the mouth direction (Figure 2). The inflow boundary condition mimics an ambient wind, where a uniform flow of $\bar{v}_w = (-v \cos \varphi, v \sin \varphi, 0)$ is imposed; the outflow boundary condition is applied on the other side of the domain, and slipping wall boundary conditions are applied to the lateral boundaries.

The breathing flow is injected at the manikin’s mouth, which was delimited by hand as the surface covered by the lips of the

![Figure B1. Comparison of the dynamic concentration maps generated at distinct time delays $\tau$ for a static pedestrian in windless conditions, by coarse-graining CFD simulations performed with (top row) the original meshgrid, (bottom row) a refined meshgrid.](image)
Figure B2. Comparison of the dynamic concentration maps generated at distinct time delays $\tau$ for a static pedestrian in a head wind blowing at 0.3 m s$^{-1}$, by coarse-graining CFD simulations performed with (top row) the original meshgrid, (bottom row) a refined meshgrid.

Figure B3. Flow rate imposed at the mouth of the manikin at each cycle of duration 3.0 s.

manikin, whose mouth is initially closed. This yields a non-planar surface of 4.7 cm$^2$ on which a uniform velocity is imposed, parallel to the ground and in front of the manikin.

The time signal is periodic (Figure B3), with a period of 3.0 s, and was designed to mimic a breathing signal$^{[82]}$ with a short period typical of the breathing pace while walking. Each breath is of 1 L of volume, so the breathing rate is 20 L min$^{-1}$.

B.4 Simulations

Simulations are first performed over four cycles to install the flow. Then, four cycles are computed to collect the statistics presented in Figure 4 and the following maps. Solutions are stored every 0.25 s (12 per cycle) for statistical accumulation.

B.5 Alternative Physical Modeling

The typical speeds of expiratory droplets and ventilation in conditions where there is a concern for contamination are well within the regime of incompressibility, that is, low Mach numbers. However, thermal effects are present, which cannot be modeled with the incompressible Navier–Stokes equations presented in the previous sections. To test the influence of these effects, we performed a limited number of simulations with the variable density formulation of the Navier–Stokes equations applicable at low Mach number, as detailed in ref. [83]. In these simulations, the air is represented as a mixture of nitrogen, oxygen, argon, carbon dioxide, and water, and its temperature is set to 20 °C, with a relative humidity of 50%. The exhaled puff is warmer and more humid, with a temperature set to 35 °C and a relative humidity of 90%. The second row of Figure B4 illustrates the (moderate) impact of these thermal effects: In comparison with the incompressible model, droplets evade the region of interest somewhat closer to the point of emission, under windless conditions, because of buoyancy.

Next, the evaporation of droplets during their propagation was also introduced in the model. For that purpose, droplets were considered to be composed of pure water, until they shrunk to a diameter of one third of the original diameter, at which point evaporation was halted, to reflect the presence of a non-water content in the droplet.$^{[29,67]}$ The model for evaporation follows the approach developed by Spalding,$^{[84]}$ similar to the one described by Bale et al.$^{[14]}$ The impact of evaporation on the concentration maps, as compared to those with thermal effects only, is barely noticeable (Figure B4). It should be noted that finer models for the evaporation of droplets of respiratory fluids (instead of water) are available in the literature,$^{[67]}$ but were not implemented in our tests. They would certainly further restrict the effect of evaporation by slowing down the decrease of diameter in time.

Finally, the thermal plume due to the receiver’s body heat may lift droplets upward and was claimed to facilitate the airborne transmission of the virus in enclosed spaces.$^{[65]}$ Our dynamic concentration maps are oblivious to the presence of a receiver,
but we investigated the possible impact of these ascending flows by shifting the region of interest downward by 20 cm (to compensate for these potentially overlooked ascending flows). Figure B4 (bottom row) ascertains that this shift only has a moderate effect on the concentration maps.

So far, the effects of these physical details of the CFD simulations were assessed in windless conditions and found to be modest. As soon as an external wind is added, these effects become slighter (Figure B4 (right)), as the droplets are swept away by the wind.

Turning to the impact of the emitter’s surroundings (in particular, the people around) on the incident air flow, we have appraised it by randomly sampling crowd configurations in a fairly dense scenario (Bellecour metro platform) and, each time, identified the nearby pedestrian most likely to perturb the air flow around the emitter, given a random wind direction. In more than 70% of the case, it was clear that this perturbation would be very small, given the distance and position of the person. For the remaining cases, we have isolated one particular configuration and performed CFD simulations of the exhalation flow with and without the pedestrian “obstacle.” We have found that, for this specific example (in which the emitter stood 1.15 m away from the “obstacle,” directly in its wake), the resulting viral concentration maps are altered by the obstacle, but the impact is relatively moderate. Nevertheless, one should bear in mind that in very dense situations, our approach may overestimate the effect of the wind in that it overlooks the wakes of surrounding people.

Appendix C: Transmission via Large Droplets

C.1 Inhalation of Large Droplets

The viral concentration for different sizes of droplets is displayed in Figure C1 with and without wind. In the main text, the focus
was put on micron-sized respiratory droplets, which are much more easily inhaled than their larger counterparts (≥50 μm after evaporation) originating from the oral cavity and lips. \[63\]

Indeed, an inhalation flow speed of around 0.3 m s\(^{-1}\) in the vertical direction is needed to overcome the sedimentation of a ≈100 μm-droplet. A dedicated simulation has been performed to delineate the region in space where the vertical flow velocity exceeds 0.3 m s\(^{-1}\) during inhalation. A flow rate of 0.55 L s\(^{-1}\) is imposed, which is a rather high value for calm breathing. \[82\] Even so, Figure C2 shows that such a vertical speed of 0.3 m s\(^{-1}\) is only reached extremely close to the nostrils, which makes them less plausible candidates for inhalation than smaller droplets.

C.2 Dynamic Maps of Concentration

Nevertheless, we computed the dynamic maps of viral concentration associated with the emission of these large droplets, the so-called oral mode; they are shown in Figure C3. Clearly, these maps differ from those obtained for smaller aerosols: In windless conditions, they are transported over less than 1 m before sedimenting. The walking motion drags these droplets farther ahead, whereas the external wind impacts them diversely; it hardly affects the mostly ballistic motion of the heaviest ones, while others are carried away by the wind. It follows that the interpolation method proposed in Section 4.4 does not work quite as well on this broad class of droplet sizes, but tests show that this interpolation still yields reasonable concentration maps; it will thus be retained.

**Figure C1.** Comparison of the viral concentration maps corresponding to different ranges of droplet sizes, for two wind speeds. Note that only one yellow dot (denoting the peak concentration) is shown, even when the map exhibits two maxima.

**Figure C2.** Region of the simulation domain where the vertical air flow during inhalation reaches a speed of at least 0.3 m s\(^{-1}\), as required to compensate the downward sedimentation of droplets of diameter about 100 μm. The inhaling flow rate is 0.55 L s\(^{-1}\).
Figure C3. Dynamic concentration maps associated with the oral mode (large droplets) emitted while speaking, in windless conditions. (Top row) static emitter, (bottom row) walking emitter.

Figure C4. Transmission risks associated with the oral mode (large droplets, assumed to be inhalable here) emitted while speaking, in the different scenarios under study.

C.3 Rates of New Infections via Large Droplets

The inclusion of the oral mode substantially alters the ranking of transmission risks across the scenarios under study, with street cafes that become overshadowed by the observed outdoor market (Figure C4). This can be ascribed to the short range of propagation of the large droplets when they are emitted by a static person (see Figure C3), as compared to a moving pedestrian. In parallel, owing to their larger sizes, droplets follow a more ballistic trajectory and are less sensitive to the wind, as reflected by the limited effect of the external wind on the rate of new infections in the different scenarios.

Appendix D: Additional Figures

Additional Figures are given below (Figures D1–D3). Figure D1 displays dynamic maps of viral concentration in the case of mouth breathing and may be compared to the speaking case shown in Figure 4. Figure D2 (as Figure 7) illustrates the performances of the interpolation of concentration maps. Finally,
Figure D1. Dynamic maps of viral concentration associated with breathing through the mouth: Effect of the external wind. (Top row) no wind, (middle row) lateral wind at 0.3 m s$^{-1}$, (bottom row) head wind at 1.0 m s$^{-1}$. Note the very different scale, as compared to the maps associated with speaking.

Figure D2. Interpolation of dynamic concentration maps at for $\nu = 1.2$ m s$^{-1}$ and $\varphi = 115^\circ$. (Top) Coarse-grained map calculated using a bona fide CFD simulation corresponding to these specific conditions. (Bottom) Concentration map obtained by interpolation.
Figure D3 assesses the effect of the assumption regarding the direction of emission with respect to the head orientation and the walking direction (see discussion in section 3.3) on the transmission risks in the different scenarios.

Acknowledgements
The setup of the CFD simulations was designed collectively, with P. Bénard, G. Lartigue, V. Moureau (CORIA Rouen, France), G. Balarac, P. Bégu (LEGI Grenoble, France), Y. Dubief (Univ. Vermont, USA), and R. Mercier (Safran Tech, France). The authors thank their colleague O. Kaplan for generating the smoke jets of Figure 3. CFD simulations were performed using HPC resources from TGCC-IRENE (Grants No. AP010312425, A0100312498, and A0120312498). A.N. acknowledges the help of B. Fray during his internship and thanks the MODCOV group for facilitating the initiation of collaborations. This work was funded by Agence Nationale de la Recherche: projects SeparationsPietons (ANR-20-COV1-0003, A. Nicolas) and TransporTable (ANR-21-CO15-0002, S. Mendez).

Conflict of Interest
The authors declare no conflict of interest.

Data Availability Statement
To allow other researchers to contribute to these prospective improvements and/or apply the proposed approach, our main scripts have been made publicly available on the GitHub repository https://github.com/an363/infectiousRisksAcrossScales; the other scripts can be requested by email.

Keywords
crowd dynamics, epidemiology, fluid dynamics, respiratory droplets

Received: September 13, 2022
Revised: March 14, 2023
Published online: May 3, 2023
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