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Review

The influenza virus, SARS-CoV-2, and the airways: Clarification for the otorhinolaryngologist

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ABSTRACT

The influenza virus and SARS-CoV-2 cause trivial upper and severe lower respiratory infections (Influenza virus 290,000 to 650,000 deaths/year). These viruses come into contact with the airways either by direct projection, by secondary inhalation of airborne droplets, or by handling (fomites). The objective of this article is to clarify the mechanisms of production and penetration of droplets of secretions emitted during all expiratory phenomena likely to transport these viruses and come into contact with the respiratory mucosa. The droplets > 5 μm follow the laws of ballistics, those < 5 μm follow Brownian motion and remain suspended in the air. The aerosols of droplets are very heterogeneous whether the subject is healthy or sick. During an infectious period, not all droplets contain viral RNA. If these RNAs are detectable around patients, on surfaces, and in the ambient air at variable distances according to the studies (from 0.5 m to beyond the patient's room), this is without prejudice to the infectious nature (viability) of the virus and the minimum infectious dose. There is a time lag between the patient's infectious period and that of RNA detection for both viruses. Subsequently, the inhaled particles must meet the laws of fluid dynamics (filtration) to settle in the respiratory tree. All of this partly explains the contagiousness and the clinical expression of these two viruses from the olfactory cleft to the alveoli.

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1. Introduction

Lower respiratory infections are the leading cause of death in poor countries and the 6th in high income countries (https://www.who.int/news-room/fact-sheets/detail/the-top-10-causes-of-death). Upper respiratory infections are the most frequent sources of morbidity with 2 to 5 colds/adult/year and 7 to 10 colds/infant/year [1,2]. Coronaviruses belong to the Coronaviridae family which has 4 genera: α-β-γ-δ. Genera α and β contain 7 coronaviruses transmissible to humans: Four are responsible for trivial upper and lower respiratory infections, the three others, SARS-CoV-1, MERS-CoV, and SARS-CoV-2 are responsible for severe lower respiratory infections [3]. They have an S surface glycoprotein (Spike) arranged like a crown which allows them to attach themselves to the epithelial receptor angiotensin-converting enzyme 2 (ACE2) and the protease transmembrane TRMPSS2 [4–8]. The A and B influenza viruses (IV) and the seasonal flu belong to the Orthomyxoviridae family and are responsible for 290,000 to 650,000 deaths/year worldwide through respiratory failure [9]. They have a haemagglutinin surface glycoprotein that attaches to sialic acid [10]. The inter-human transmission of viral infection occurs through close direct contact with an infected person, by touching a surface contaminated with short-distance projections, and impaction of droplets of secretion (fomites) [11]. Transmission can occur over a longer distance by airborne droplets [12,13]. The objective of this clarification was to analyse the objective data of patients’ contagiousness by transporting the virus through producing secretions and the possibility of them penetrating the airways. Assessing transmissibility is important for the otorhinolaryngologist who is at the forefront of treating the upper aerodigestive tracts and because many viruses cause ENT manifestations. Finally, the current SARS-CoV-2 pandemic and the annual flu epidemic require professionals to be up to date with the latest knowledge about these mechanisms in order to adapt their practices.

2. Method

A structured search on PubMed was carried out until the end of April 2020. It took publications with abstracts in French and English into consideration. The terms searched were “influenza virus”,

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3. Discussion

The air that we breathe transports particles of different sizes (granulometry) (Table 1) combining pollution, diesel, allergens, and viruses [3,18,20,21]. Its composition varies depending on the place (indoors/outdoors, city/countryside), the day, meteorology (wind, hygrometry), profession, weeks, months, seasons, and countries [16,22–25]. Coughing, sneezing, or simply breathing and speaking produces thousands of droplets whose sizes vary from the microscale to the nanoscale (Table 1) [14]. The largest ones, >5 μm, follow the laws of ballistics and of gravity. Their inertia parameter is defined as the product of the density (ρ) multiplied by the diameter of the particle squared (d^2) and the flow (Q) (inertia parameter = ρd^2Q in g.m^2.s^-1) [26]. A droplet of 100 μm will settle on the ground in 16 s [27]. Finer particles, <5 μm, make up the breathable part of the aerosol, remain airborne (negligible falling speed limit <25 cm/s), and follow Brownian motion [28–31]. There are thus short-range or long-range particles depending on their capacity to remain airborne or not. On the other hand, from their creation, the droplets undergo evaporation (droplet nuclei) which reduces their diameter and increases their capacity to remain airborne, encouraging the maintenance of a long-range contamination source: a particle of 100 μm will take 100 seconds to reach a diameter of 32 μm in an atmosphere with 95% humidity, and less than 2 seconds in an atmosphere with 35% humidity. Inversely, evaporation can make other droplets with smaller diameters disappear [27,28,32].

Table 1
Size of the main particles likely to be inhaled.

| Particles     | Mean diameter | References |
|---------------|---------------|------------|
| When calm expiration | 0–500 μm      | Xie 2009 [14], Tang 2013 [15] |
| When coughing  | 0–1500 μm     |            |
| Grasses (Timothy grass) | 30–40 μm    | Crouzy 2016 [16] |
| House dust mite | 22 ± 6 μm      | Zhang 2019 [17] |
| Pollen         |                | Lapaerta 2003 [18], Ezz 2015 [19] |
| PM10          | < 10 μm       |            |
| Fines         | < 2.5 μm      |            |
| Ultrafines    | < 100 nm      |            |
| Virus          |                | CSG 2020 [3] |
| Rhinovirus    | 28–30 nm      |            |
| Coronavirus    | 80–200 nm     |            |
| Influenza A & B | 80–120 nm   |            |
| Synantial Virus| 150–400 nm    |            |

3.1. Production, diffusion, and penetration of droplets from the patient to the recipient

3.1.1. During normal expiration and speech

The air that we breathe out through our nose or mouth is projected a distance of 0.6 to 0.8 m at an average maximum speed of 1.3 to 3.9 m/s [15,33]. Depending on the studies, the total number of droplets produced is estimated between 112 and 6720 droplets, when the subject is asked to count from 0 to 100 out loud. 90% settle on the ground at a distance of 30 cm [14]. Their size, however, is very heterogeneous and variable from one individual to another between 0 and 500 μm with an average diameter of 16 μm [14,33]. These droplets represent a mass of 18 to 79 mg [14]. More recently, Asadi et al. have demonstrated a production of 4.8 ± 3 particles/s with diameters of 0.5 to 5 μm while speaking, with a production rate that correlates with increasing voice loudness [34].

The digital simulation of droplets <5 μm produced by normal expiration between two people face-to-face showed that, after 200 seconds, 6.2–5.7–0.8 and 0.4% settled on the subject facing them at respective distances of 0.5–1–1.5 and 3 metres [27]. The proportion of inhaled airborne droplets was 0.2–0.6–0.02 and 0% for the same distances [27]. The authors estimated the safe distance at 1.5 m [27]. This distance must be adjusted depending on oral or nasal breathing, the size of the people, the position of the face, the ambient humidity, the room’s ventilation, and the exposure time [27,34].

3.1.2. When sneezing and coughing

The average flow rate of a sneeze is 4.8 L/s, the air is projected 0.6 m at a maximum speed of 4.5 to 8 m/s with an expansion rate of 2 m^2/sec [15,35,36]. For coughing, the air is projected 0.7 m at a maximum speed of 6 to 11.7 m/s with an expansion rate of 1.5 m^2/sec [15,33,35]. It will be noted that these parameters are not very different [15].

In the healthy subject, who by definition does not cough, a cough would produce on average 800 to 2045 droplets with 80% settling on the ground at a distance of more than 50 cm [14,33]. When coughing, the granulometry of the droplets produced has a different distribution compared to that observed when speaking, with diameters ranging from 0 to 1500 μm. However, their average diameters are quite close: 13.5 μm for coughing versus 16 μm for speaking [14,33]. Their mass is between 23 and 85 mg after 20 successive coughs [14].

3.1.3. Which droplets can penetrate the airways?

Large droplets have little chance of reaching the recipient’s airways as these are short-range objects. Safe distances are useful even though they can vary twofold depending on the studies [20,27] and their contaminating properties have not been demonstrated (see below). Airborne droplets can reach the airways: in a calm environment, a 4 μm particle takes 33 minutes to move 1 metre and a 1 μm particle takes 8 hours. However, it must pass through the nasal filter whose filtration capacity is not linear [26]. On the microscale, the bigger the particle the more it is filtered. On the nanoscale, it is the reverse: the smaller the particle the more it is filtered [30,37]. For calm respiration (10–15 L/min), 100% of 20 μm particles settle in the nasal fossa and the cavum, 40% for 9 to 11 μm particles, and only 4% for those from 2.5 μm to 1.3 to 3.9 μm [15,33]. For the same flow rate, 80% of 1 nm particles settle in the nasal fossa, 40% of 4 nm particles, and 10% of those between 10 and 100 nm [30,37]. The deposit rate in the nasal fossa and the cavum is thus the same for particles from 20 μm and from 1 nm. However, as a result of the influence of the inertia parameter or of Brownian motion, the mapping of the deposits is different: particles from 20 μm will follow the main air current and will impact on anatomical structures in the back third of the nasal fossa where the flow changes direction, while particles from
1 nm will be deposited in all directions, all over the nasal mucosa [30,37,38].

Several variation factors play a part: when the density increases, the inertia parameter increases the capacity of microscale particles to impact while it has no effect on nanoscale particles [30]. An increasing airflow will have the same effect on the microscale, while it decreases the deposit rate of nanoscale particles, all the more so if they are smaller [30,38]. This behaviour for nanoscale particles is applicable throughout the nasal fossa except in the olfactory cleft, where this parameter is not modified or only slightly modified [37,39], which is very important for viruses transported by a large heterogeneity of airborne droplets (from 0.3 and 4 μm) [21,40]. After the inspiratory phase, the still airborne particles in reverse flow will increase their deposit rate in the nasal fossa and the olfactory clefts because of upwards air deflection by the turbinates, and increase their residence time through the resistance of the nasal valve [41].

In contrast, particles between 10 μm and 10 nm are not filtered by the nose and can penetrate the rest of the airways (Table 2). These filtration characteristics are already used in daily practice for aerosol therapy since it is recommended to use machines producing particles >5 μm for rhinological pathologies, from 5 to 2 μm for tracheobronchial pathologies, and between 2 and 0.5 μm if intended for the pulmonary alveoli [43,44]. FFP-type respiratory protection masks are tested on inspiration and clased according to their filtration performance with regard to a sodium chloride aerosol composed of particles with a median diameter of 0.6 μm and a paraffin oil aerosol with a median diameter of 0.4 μm (www.inrs.fr/dms/inrs/CataloguePapier/ED/TI-ED-6106/ed6106.pdf), with 3 levels of protection: FFP1 the filter allows only 20% of the particles to penetrate on inspiration with a maximum leakage rate of 22% (interior leakage), FFP2 – the filter allows only 6% of the particles to penetrate with a maximum leakage rate of 8%, and FFP3 – where only 1% of the particles penetrate with a maximum leakage rate of 2%. The mechanisms for stopping the droplets vary depending on the granulometry: from 10 to 1 μm, the inertia parameter and gravity are used; from 1 μm to 300–100 nm, the number, diameter, and weave of the fibres allow for mechanical filtration; below this, the particles are retained by the electrostatic properties of the fibres and diffusion [45,46]. The performance of a mask is a compromise between the difficulty with breathing (pressure loss <30 Pa) and its filtration capacity, which depends on fluid and electrostatic mechanics but also on their fitting to the face [45,46]. “Home-made” masks made of several fabrics (cotton/silk) and several layers (3 to 4) can match the performance of industrial masks [45], knowing that small-diameter fibres increase the electrostatic capture of nanoparticles [46].

### 3.2. Transmission of the influenza virus

During a flu, the air volume of a cough does not change before and after the illness and varies from 2.33 to 2.48 L/cough with a flow of 5.33 to 6.9 L/sec [21,47]. The number of particles on coughing varied widely from one individual to another: while ill, it was 75,400 ± 97,300 droplets/cough, and 52,200 ± 98,600 droplets/cough after recovery (non-significant difference). In contrast, the average volume of aerosol was 38.3 pl/cough while ill versus 26.4 pl/cough afterwards (P < 0.0143) [21]. The size of this aerosol’s droplets varied little during and after the illness, between 0.35 to 2.5 μm for an average of 0.63 μm [21].

Gralton et al. showed, during coughing, the presence of IV, rhinovirus, and respiratory syncytial virus in 12 adults and 41 children, 57% of the particles were >5 μm and 82% were <5 μm [48]. Milton et al., with 33 patients, found that 43% of particles >5 μm carried flu virus RNA, and 92% of particles <5 μm did so [49]. The number of viral copies was only 12 copies/30 minutes in the coarse part of the aerosol while there was on average 560 copies/30 minutes in particles <5 μm. Wearing a surgical mask reduced the risk of diffusion of viral copies by a factor of 25 for particles >5 μm and by 2.8 times for particles <5 μm [49].

In flu season and respiratory syncytial virus infection, Lindsay et al. measured the presence of airborne viral RNA in 264 samples taken from fixed stations and from 21 employees at a healthcare facility treating patients suffering from both illnesses. These RNAs were detected in the air and there was a good correlation between the number of positive samples and the number of positive patients [20]. This was especially the case in poorly ventilated rooms such as examination rooms (door closed) where the concentration was at a maximum. These RNAs were contained in 43% of the particles with a diameter ≤4.1 μm. The maximum of samples was positive in an area 1.8 m around the patient and for higher up stations because of the room’s ventilation system, demonstrating these particles’ capacity to remain airborne [20]. In other facilities, the virus was found airborne in patients’ rooms only in 50% of cases with 162 ± 1.9 copies/m³ in 24% of particles >4 μm, 144 copies/m³ in 6% of particles between 1–4 μm, and none in particles <1 μm [40].

However, the presence of viral RNA does not imply contagiousness: what do these values mean? Does this represent an infectious risk for the individual? No statistically significant association was highlighted between the level of viral excretion and transmission [50,51]; finding viral RNA or a viral load is without prejudice to the viability of the virus and the minimal infectious dose. In a H1N1 and H3N1 IV infection model, it is the loss of positivity of the viral culture that signals the non-contaminant nature of the patient but not the absence of RNA, which coincides with the end of the infectious period. Alford et al. administered the IV to healthy volunteers in order to measure the minimal infectious dose [52]. Via aerosol distributing droplets of 1 to 3 μm, the infectious dose for humans was from 0.6 to 3 TCID₅₀/mL (dose necessary to infect 50% of the cells in a reference cell culture). Administered as nasal droplets, this value was from 127 to 320 TCID₅₀/mL [53]. Patients infected by the aerosol said that their symptoms were in line with the usual ones, while those infected by nasal inoculation had low-intensity symptoms over longer periods [52,53].

In practice, the percentage of infectious IV in droplets produced by coughing or expiration varied from 5 to 42% for particles from 0.3 to 8 μm in diameter where patients had clinical signs for 2 days [47,49,54,55]. Regardless, these figures vary widely depending on the granulometry of the particles. Lindsay et al. did not find infectious virus in particles >5 μm [47]. In contrast, the quantity of infectious virus in airborne particles (<5 μm) was enough to be contaminating [47,52]. However, these studies have low patient populations, involve mild forms of the illness, and do not take into account the kinetics of viral detection or the length of viability of the virus; their results depend on the sensitivity of the viral culture methods implemented to demonstrate infectivity. Lastly, the presence of infectious virus in expectoration does not mean that they will reach their target because these studies do not take into account distance and residence time, the environment (survival of IV A in low rates of hygrometry) [9], air movements, the
recipient’s breathing, their sneezing, their mucociliary clearance, and their immune status.

### 3.3. Transmission of SARS-CoV-2

Guo et al. measured the presence of viral RNA in the air of Intensive Care Units (ICU) and traditional hospitalisation sections in Wuhan. The virus was present in 33% of the samples taken from near air treatment openings, in 44% of samples near patients, and in 12.5% of cases at the entrance to the ICU bedroom (4 m from the patient’s head). The ground was systematically positive even outside of bedrooms, leading to half of employees’ shoe soles being positive. In the hospitalisation sections, only 15.4% of samples were positive at 2.5 m from the patient’s head, and 18.2% near the patient [56]. The presence of viral RNA followed human activity, air movements, and depended on the viral load of the patient, which is often more significant in ICU than in general hospitalisation where there are milder forms [57].

Yu F. et al. showed that, out of 323 samples taken from 76 confirmed COVID+ patients, the most productive biological liquids were, in descending order, spit, oropharyngeal swabs, and nasal swabs. Spit contained the most viral copies (17,429 ± 6920 copies/test versus 2552 ± 1965 copies/test for oropharyngeal swabs, and 651 ± 501 copies/test for nasal swabs, $P<0.001$) [58]. Other authors confirmed this result [59–61]: the viral load in saliva was very high the older than 60 the patients were, if they had 1 to 2 comorbidities, and/or a severe form in the first week. This salivary excretion of the virus seems to correlate with the biomarkers of tissue damage and a strong expression of the ACE2 receptor in the oral mucosa [8,62]. However, others showed the superiority of nasopharyngeal swabs for diagnosing COVID-19 [63]. Given this heterogeneity of results, the American Centre for Disease Control and Prevention left the choice of the type of sample for diagnosing the illness to the discretion of the operator in its recommendations dated 26 April (https://www.cdc.gov/coronavirus/2019-ncov/lab/guidelines-clinical-specimens.html).

Zheng et al. demonstrated that, in 96 COVID+ patients, viral RNA was detected in respiratory secretions and stools up to the 3rd and 4th week after the onset of symptoms, with a net reduction in expression time for mild forms in comparison with severe forms [64]. These time periods were also found by others [60]. However, Xu et al. showed that, from a group of patients in Wuhan who had transmitted the virus to a group in another region, who themselves transmitted the illness to a third group, there was reduction in intensity and in the duration of RNA expression, which could indicate a finite phenomenon: the RNA was no longer detectable in samples taken from the third group 7 days after the onset of symptoms [60].

In any event, with the same tools used for the flu, the same questions are raised for SARS-CoV-2: is detecting viral RNA synonymous with contagiousness? If the viral RNA is detectable in respiratory secretions and stools after the clinical condition for more than a month, the “living” virus could not be detected by culture after the 3rd week: the results of the reverse transcription polymerase chain reaction (RT-PCR) remained positive 6 to 8 days after the loss of transmissibility [65]. He et al., with a contagiousness model for the flu and SARS-CoV-1 and using data from oropharyngeal samples taken from 77 COVID+ patients, estimated peak contagiousness at 2 days before and 1 day after the onset of symptoms, while viral RNA is detectable up to the 21st day after the onset of symptoms [66]. The experimental production of droplets < 5 μm containing SARS-CoV-2 demonstrated the presence of viable virus in the aerosol for 3 hours at a temperature of 21–23 °C and in an atmosphere with 40% humidity [11]. The half-life of SARS-CoV-2 was estimated at 1 hour in air, < 1 hour on copper, 3.8 hours on cardboard, and 5.6 and 6.8 hours on stainless steel and plastic [11].

Finally, if the contamination modes are similar for IV and SARS-CoV-2, all of this data allows us to understand certain features and similarities in the clinical expression of the two viruses. Pneumopathies are common but for COVID+ patients, unlike the flu, inflammatory rhinological manifestations (congestion, nasal obstruction, and rhinorhoea) are rare [67]. On the other hand, both viruses are capable of causing anosmia and ageusia, but the former is rare with associated rhinological inflammatory signs, slowly and slightly reversible, the latter is frequent, with little or no rhinological inflammatory signs but a quicker olfactory recovery of between 7 and 10 days [67,68]. For the flu, sialic acid is expressed on the surface of the ciliated epithelial cells [10]. For SARS-CoV-2, ACE2 and TRMPSS2 are expressed in the nasal mucosa and the olfactory mucosa [5–7]. However, mechanisms causing loss of function should not be similar as, in a population of 262 patients with flu-like symptoms, 70% of COVID+ patients (40 cases) manifested olfactory losses versus 17% of COVID– patients (35 cases) [69]. Other authors confirmed these results [68]. COVID+ patients with olfactory disorders were significantly younger than those who had none [68].

### 4. Conclusion

Droplets are produced during all expiratory phenomena in healthy and ill subjects with wide disparities between individuals and few between the two states. Aerosol droplets < 5 μm seem to be the most problematic because they remain airborne. However, they do not all contain viral RNA. If viral RNA is detectable around patients, on surfaces, and in the surrounding air, this is without prejudice to the viability of the virus and the possibility of transmitting the illness. In this respect, there are a lot of similarities between IV and SARS-CoV-2. Finally, if droplets are inhaled, the minimal infectious dose must be reached. It is not known for SARS-CoV-2 and it is likely that contact time remains a determining factor.

This clarification should allow the ENT specialist to assess the situation in the professional contexts they encounter, to explain it to the general public, and to justify the necessary measures to take with discernment.

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The authors declare that they have no competing interest.

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