Otolaryngology manifestations of COVID-19: a contemporary viewpoint

Since the first documented case of pneumonia of unknown origin hailed from Wuhan city, Hubei province, China, back in December 2019, the novel SARS-CoV-2 leading to COVID-19 has swept across the globe afflicted the people in every corner. As of 21 March 2021, the total case number has culminated to over 122 million worldwide, with a mortality rate of 2.2%.1 11 March 2021 marks the first anniversary since WHO declared the COVID-19 outbreak a pandemic. With the vaccine challenges that lie ahead, the light at the end of the tunnel remains elusive.

We learnt that the transmission routes are via respiratory droplet, aerosols and by contact that may be indirect, which succinctly impose a remarkably high risk to the head and neck surgeons who often deal with aerosol-generating procedures. In fact, the first doctor who succumbed to COVID-19 in Wuhan was an otolaryngologist.2 The acknowledgement of a heterogeneous spectrum of clinical manifestations of SARS-CoV-2 viral infection, ranging from asymptomatic to severe disease with multiple organ failure, is of paramount importance. Apart from the overall presentation being fever, myalgia and lower respiratory tract symptoms of cough and dyspnoea, extrapulmonary symptomatology of COVID-19 manifested through a variety of ear, nose, throat (ENT) complaints have been gradually recognised.

In conjunction with hypogeusia, smell deficit constitutes the earliest reported ENT manifestation of the novel coronavirus. Olfactory dysfunction was identified in a substantial fraction of patients who suffered from COVID-19, with a prevalence of 47.85% worldwide.3 The idiosyncratic characteristics of smell disturbance attributed to COVID-19 include its sudden onset, transient duration, and in most cases, rapid recuperation. Additionally, smell deficit may manifest as a sole nasal symptom without concomitant rhinorrhea or congestion.4

Other sinonasal manifestations, namely nasal congestion and rhinorrhea, have been reported at a much lower occurrence rate. Paradoxically, a significantly lower amount of viral entry receptors, was observed in patients with allergic airway diseases, of which the diagnostic symptoms include nasal discharge and/or blocked nose.5 The prevalence of chronic rhinosinusitis in patients with COVID-19 was comparable with the general population, implying that sinusitis may not increase viral infection susceptibility. The underlying chronic eosinophilic inflammation with long-term exposure to corticosteroid in allergic airway diseases was found to downregulate the viral entry gene expression, inferring the possibility of a protective role against COVID-19 infection.6

How about the otological presentation? The theory of virus-induced aural dysfunction has always been a familiar centre in the field of medicine. Viruses can damage the inner ear structures by initiating a host-immune response. A diversity of overlapped manifestations, including hearing impairment, vertigo, peripheral facial palsy and tinnitus with various onset and severity, have been described. There is scattered but emerging evidence of COVID-19-related peripheral facial palsy. The correlation between the two, however, remains undetermined. Understandably, the SARS-CoV-2 possess neurotropic and neuroinvasive properties, ergo the predilection of a viral-induced peripheral nerve injury. The postulated pathogenesis of facial palsy includes direct infection injury, ischaemia of the vasa nervorum, and local inflammation giving rise to demyelination.6

The SARS-CoV-2 infection may manifest itself through inner ear complaints such as vertigo and hearing loss. Of note, dizziness was among the most typical neurological manifestation of COVID-19 disease.7 Akin to the sequelae of virus infection on the olfactory sense’s neural pathway, the proposition of aural dysfunction due to vascular hypoperfusion and local inflammation on the audiovestibular centres caused by SARS-CoV-2 infection is plausible.8 The viruses may intrude on the haematological path by binding the erythrocytes via the β chain. The ensuing viral deposit onto the ACE2 receptor protein at the temporal lobe and hearing centre then trigger a cascade of a local inflammatory process, paving the way to hearing loss and imbalance.9 Intriguingly, the risk of hearing damage seems to be reciprocating the severity of COVID-19 infection, granted that the fact patient might also complain of hearing impairment as the sole symptom.10 It is crucial to establish the dichotomy between non-vertiginous dizziness and vertigo. While dizziness may be related to a myriad of causes, vertigo, on the other hand, is almost always caused by a vestibular pathology. Pertinent history, including the characteristic of a false sense of motion, the duration and frequency, and most importantly, other associated otological complaints such as hearing loss and tinnitus, should be carefully evaluated. It is worthy of note that vertigo, although a non-specific COVID-19 symptom, should not be taken lightly for its detrimental repercussions.

SARS-CoV-2 has also been identified in the saliva due to the high expression of ACE2 in the salivary gland’s lining epithelial cells. There have been anecdotal reports of orofacial manifestation link to COVID-19. Interestingly, such clinical features were reported to be the first clinical sign of SARS-CoV-2 infection.10 The archetypal presentation of a unilateral parotid swelling associated with pain and non-specific ulcerative or vesiculobullous lesion of the oral mucosa indifferent from other inflammatory or infective cause then raised the concern of the virus as a potential great masquerader.

The battle against COVID-19 is not over. Early recognition of atypical manifestation of SARS-CoV-2 infection remains pivotal to avert misdiagnosis and delay in management. Therefore, we urge the treating physicians to be well informed of the growing spectrum of clinical presentation of COVID-19 while remaining vigilant for potentially disastrous consequences.

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