Original Article

Transient modifications of the olfactory bulb on MR follow-up of COVID-19 patients with related olfactory dysfunction

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ABSTRACT

Background: Olfactory dysfunction (OD) has been reported with a high prevalence on mild to moderate COVID-19 patients. Previous reports suggest that volume and signal intensity of olfactory bulbs (OB) have been reported as abnormal on acute phase of COVID-19 anosmia, but a prospective MRI and clinical follow-up study of COVID-19 patients presenting with OD was missing, aiming at understanding the modification of OB during patients’ follow-up.

Methods: A prospective multicenter study was conducted including 11 COVID-19 patients with OD. Patients underwent MRI and psychophysical olfactory assessments at baseline and 6-month post-COVID-19. T2 FLAIR-Signal intensity ratio (SIR) was measured between the average signal of the OB and the average signal of white matter. OB volumes and obstruction of olfactory clefts (OC) were evaluated at both evaluation times.

Results: The psychophysical evaluations demonstrated a 6-month recovery in 10/11 patients (90.9%). The mean values of OB-SIR significantly decreased from baseline (1.66±0.24) to 6-month follow-up (1.35±0.27), reporting a mean variation of -17.82±15.20 %. The mean values of OB volumes significantly decreased from baseline (49.22±9.88 mm3) to 6-month follow-up (43.70±9.88 mm3), (p=0.006).

Conclusion: Patients with demonstrated anosmia reported abnormalities in OB imaging that may be objectively evaluated with the measurement of SIR and OB volumes. SIR and OB volumes significantly normalized when patient recovered smell. This supports the underlying mechanism of a transient inflammation of the OB as a cause of Olfactory Dysfunction in COVID-19 patients.

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Introduction

Coronavirus disease-2019 (COVID-19), linked to the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), has been associated with many clinical pictures, ranging from simple rhinitis to acute respiratory distress syndrome. Frequently, the infection has been linked to smell and taste dysfunctions, reported in 50 to 86% of mild-to-moderate cases. The olfactory dysfunction (OD) may develop through nasal obstruction, such as in common cold, or be related to neurological viral spread through the olfactory cleft mucosa. In the first paper reporting OD as a key symptom of the COVID-19, more than 80% of patients presented with OD without associated rhinorrhea or nasal obstruction. Volume and signal intensity of OB have been reported as abnormal on acute phase of COVID-19 anosmia. Although the symptoms remain most of the time regressive, MRI findings in patients with prolonged COVID-19-induced OD was significantly associated with lower olfactory bulbs (OB) heights, compared to controls. These findings support the perception that SARS-CoV-2 may invade the central nervous system through the olfactory pathway, inflicting immediate and potentially irreversible damage to neuronal structures like the OB. However, a prospective longitudinal study was missing to demonstrate the existence of OB modifications...
in terms of volume and aspect in parallel with the clinical follow up of patients recovering or not from COVID-19 anosmia. We here report a prospective MRI and clinical follow-up study of COVID-19 patients presenting with OD, aiming at understanding the modification of OB during patients’ follow-up.

Material and methods

Ethics, patient and setting

The study was approved by the IRB of Foch Hospital (ID-RCB: 2020-A00832-37). Written informed consent was obtained for participants.

From March to May 2020, COVID-19 patients with self-reported sudden OD were prospectively recruited from two University Hospitals (Foch Hospital and Garches Hospital, APHP, Paris, France). Patients had demonstrated mild to moderate COVID-19 through serology or reverse transcription polymerase chain reaction (RT-PCR) findings. Patients with a history of OD before the pandemic, nasal surgery, chronic rhinosinusitis, head and neck trauma, degenerative neurological disease, or contraindication to MRI were excluded from the study.

Olfactory assessments

Patients benefited from psychophysical olfactory assessments (Sniffin’Sticks tests, Medisense, Groningen, Netherlands). The Sniffin’-Sticks test is a validated psychophysical olfactory test using 16 smell pens. Each pen was presented to the individual who had to choose the adequate smell between four given options. The final score ranges from 0 (no olfaction) to 16 (perfect olfaction). Hyposmia was defined by a score ranging from 9 to 11 and anosmia if <9. The objective evaluations were made within the two or four weeks following the onset of the olfactory disease and throughout the follow-up.

The olfactory cleft specific Lund-Kennedy scoring (OC-LK) was performed to evaluate cleft obstruction, this endoscopic score being correlated with the olfactory function in patients with smell disorder.5

The impact of COVID-19 on sinonasal symptoms was evaluated through the French version of the sino-nasal outcome test-22 (SNOT-22), which is a validated patient-reported outcome questionnaire summarized in Table 1.

Imaging acquisition

Imaging studies were performed on a 3 Tesla device (Discovery TM MR750, General Electric, Milwaukee, WI, USA) with a 20-elements head and neck coil. The sequences performed in order to assess the OB signal were 3D-FLAIR-CUBE and 3D-T2-FIESTA acquired respectively in coronal and sagittal planes. The 3D-FLAIR sequence parameters were: TR/TE 8000/133ms; section thickness 2mm; matrix 240 × 240; FOV 230 × 230mm; flip angle 90° and acquisition time = 4min-36sec. The 3D-T2 FIESTA sequence parameters were: TR/TE 5.9/2.5ms; section thickness 0.6mm; matrix 320 × 320; FOV 180 × 180mm; flip angle 55° and acquisition time=4min-32sec. The same protocol was conducted at the initial stage of the infection and at 6-months follow-up.

Image analysis

Two experienced neuroradiologists, blinded to clinical data, independently reviewed the images. A third neuroradiologist resolved potential discordances. Volumes of OB were measured on 3D-T2 sequence using a post-processing application dedicated to segmentation: ITK-SNAP8 3.8 (www.itksnap.org).

Olfactory Cleft (OC) obstruction was evaluated on T2/FLAIR images and was defined partial if it didn’t affect the entire anteroposterior length.

Quantitative analysis was performed on T2/FLAIR image by adjusting contours of a ROI centered on the OB on a coronal plane, in order to measure the average OB signal intensity. A signal intensity ratio (SIR) was then calculated between the average signal of the OB and the average signal of a ROI placed in the frontal white matter of normal appearance at the same slice level.

Results

Eleven anosmic patients were included. The mean age was 41.5 years (range 24-75 years). Patients’ characteristics are summarized in Table 1.

Olfactory assessments

Whereas all patients were initially anosmic, only 1 patient remained hyposmic on the 6 months follow-up (9.09%). The latter had a partial recovery with SniffiSticks test improving from 3 to 8.

At the initial examination, OC-LK was normal (score 0) in 8/11 patients (72.72%).

The mean SNOT-22 was 18.45±19.65 (0-61) on the first exam, and was 10.36±16.36 (range 0-47) on the follow-up examination. The hyposmic patient at the second examination had an improvement of the SNOT-22 score from 43 to 0.

Imaging analysis

Five patients (45.54%) had partial OC obstruction, bilateral in 3 patients on the initial MRI. These obstructions were regressive in all cases on the control MRI.

As for SIR and volume measurements of the OB, the intra-class correlation coefficient was good for SIR measurement (r=0.898, 95%CI [0.770–0.956], p<0.001) and excellent for OB volumes measurement (r=0.996, 95%CI [0.991–0.998], p<0.001).

Table 1

| Patients characteristics | Patients (N-%) |
|-------------------------|---------------|
| Age (mean ± SD) – years old | 41.5 ± 19.5 |
| Gender (female/male) | 7/4 |
| Rhinological history | |
| Patients with seasonal allergy | 3/11 (27.3) |
| Septoplasty | 2/11 (18.2) |
| Other antecedents | |
| Dysthyroidism | 2/11 (18.2) |
| Hypertension | 1/11 (9.1) |
| Cardiac stent | 1/11 (9.1) |
| Autoimmune disease | 1/11 (9.1) |
| Endometriosis | 1/11 (9.1) |

SD: standard deviation.
The mean values of OB-SIR signficantly normalized from baseline (1.66 ± 0.24) to 6-month follow-up (1.35 ± 0.27), reporting a mean variation of -17.82 ± 15.20 % (p < 0.001; Figure 1).

The mean values of OB volumes significantly decreased from baseline (49.22 ± 10.46 mm³) to 6-month follow-up (43.70 ± 9.88 mm³), with a mean variation of -10.30 ± 13.01 % (p=0.006; Figure 2).

Concerning the patient with persistent hyposmia on the follow-up examination, MRI showed a severe drop in signal intensity ratio (-23.96% and -27.72%) and of volumes (-27.03% and -28.38%) in the right and left OB respectively.

Note that one patient presented multiple sclerosis-like lesions at first MRI, the disease being not previously known.

Discussion

MR imaging in COVID-19 anosmic patients showed initial increase in signal intensity and volume of olfactory bulbs with a normalisation on 6 months-MRI-follow-up whilst 90% of our patients clinically recovered smell. This supports the underlying mechanism of a transient inflammation of the OB as a cause of OD in COVID-19 patients who recover. Initial signals and volumes would then be increased due to inflammation, followed by a normalization of these measures over time.

Laurendon et al.9 reported similar findings in a 27-year-old COVID-19 anosmic patient reporting edema and transient swelling of the OB. Initial inflammatory reaction in the OB suggest a neural spread of the SARS-CoV-2 into the OC mucosa. Many viruses, including previous forms of coronaviruses, are known to spread from the nasal epithelium to the olfactory bulb and brain but in smaller proportion of cases.10,11 Interestingly, the brain spread of the virus was associated with concomitant brain damage in two recent studies.12,13 Accordingly, COVID-19 RNA was identified in the brains of patients, supporting the neurological pattern of the virus.14 The virus spreads through the angiotensin converting enzyme 2 receptor (ACE2) and transmembrane serine protease 2 (TMPRSS2), which are both expressed in OC epithelium and in the sustentacular cells of the OB.10,15 The inflammatory reaction starts in the neuroepithelium, which may appear obstructed on the initial MRI. Eliezer et al.16 found that OD and OC obstruction are present at the early stage of the disease, and improved both at one-month follow-up, suggesting that OD in Covid-19 patients was caused, at least in part, by reversible

Fig. 1. Signal intensity ratio (SIR) measurement
Coronal Flair-weighted at the same level slices through the olfactory bulbs (OB) in a 32-year-old anosmic female at the initial exam (A) and at the follow-up exam (B), performed respectively at day 4 and 147 from onset of anosmia. The SIR between the average signal of the OB (manual ROI) and the average signal of a ROI placed in the frontal same level white-matter (circle) went from 1.63 to 1.05 on the right and from 1.68 to 0.96 on the left (fall of 35.26% and 42.45% respectively). There was a partial obstruction of right olfactory cleft (dotted arrow) which regressed on control. Note the isointense signal OB (arrows) compared to normal brain cortex (head of arrows).

Fig. 2. Measurement of the olfactory bulbs (OB) volumes
Same level reformatted axial, frontal and sagittal 3D-T2-weighted slices through the olfactory bulbs (OB) in a 26-year-old anosmic female at the initial exam (A, B, C) and at the follow-up exam (D, E, F), performed respectively at 7 and 147 days from onset of anosmia. OB volumes varied from 54.91 to 41.80 mm³ on the right and from 57.73 to 42.7 mm³ on the left (decrease of 41.0% and 37.85% respectively).
inflammatory changes in the OC. In a recent systematic review,\textsuperscript{17} opacification of OC (60/218 − 27.5%) together with normal OB morphology and signal intensity (68/218 − 31.2%) were the most common imaging findings in patients with olfactory dysfunction secondary to COVID-19. In our study, we found only partial OB obstruction in only 4/11 patients on the initial MRI, totally regressive during follow-up, explaining only partly OD. By adding a quantitative analysis of OD signal intensity, our study supports the presence of an initial inflammation of OB as a cause of OD. OB involvement in COVID-19 OD was also mentioned by Aragão et al.\textsuperscript{18} who found a hypersignal and/or an enhancement of OB on T1-weighted images, explaining the OD by the presence of microbleeding and/or blood-brain barrier break. Kandemirli et al.\textsuperscript{19} also suggested the potential microhemorrhagic complications by finding a punctate hypointense T2-weighted pattern within the OB in 4/23 patients with persistent COVID-19 OD. Those abnormalities may represent a subsequent marker of severity of inflammatory lesion of the OB.

Interestingly, patients with prolonged OD may develop OB atrophy. Tsivgoulis et al.\textsuperscript{20} demonstrated that prolonged SARS-CoV2-induced OD is significantly associated with lower OB heights bilaterally compared to controls. Concerning our patient with persistent hyposmia on the follow-up examination, MRI showed a severe decrease in volumes, corroborating this hypothesis.

Certain limitations of the present study should be highlighted, including the small sample size of our population and the fact that a vast majority of our patients recovered from anosmia, which was not predictable at the initial time point.

**Conclusion**

COVID-19 anosmic patients reported initial increase in signal intensity and volume of olfactory bulbs, followed by a normalization at 6-months-MRI-follow-up whilst patient recovered smell. This supports the underlying mechanism of a transient inflammation of the OB as a cause of Olfactory Dysfunction in COVID-19 patients who then recover. In a context where the clinical examination of infected patient is still discouraging because of aerosolization risk, our study supports that the investigation of anosmia through MRI is feasible.

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**Data availability statement**

The data that support the findings of this study are available on request from the corresponding author. The data are not publicly available due to privacy or ethical restrictions.

**Declarations of Competing Interest**

None.

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