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### **ORIGINAL ARTICLE**

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### Pre-registered controlled comparison of auditory function reveals no difference between hospitalised adults with and without COVID-19

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### ABSTRACT

**Objective:** Several viruses are known to have a negative impact on hearing health. The global prevalence of COVID-19 means that it is crucial to understand whether and how SARS-CoV2 affects hearing. Evidence to date is mixed, with studies frequently exhibiting limitations in the methodological approaches used or the populations sampled, leading to a substantial risk of bias. This study addressed many of these limitations.

**Design:** A comprehensive battery of measures was administered, including lab-based behavioural and physiological measures, as well as self-report instruments. Performance was thoroughly assessed across the auditory system, including measures of cochlear function, neural function and auditory perception. Hypotheses and analyses were pre-registered.

**Study samples:** Participants who were hospitalised as a result of COVID-19 (n = 57) were compared with a well-matched control group (n = 40) who had also been hospitalised but had never had COVID-19. **Results:** We find no evidence to support the hypothesis that COVID-19 is associated with deficits in auditory function on any auditory test measure. Of all the confirmatory analyses, only the self-report measure of hearing decline indicated any difference between groups.

**Conclusion:** Results do not support the hypothesis that COVID-19 infection has a significant long-term impact on the auditory system.

### Introduction

While several viruses are known to negatively impact the auditory-vestibular system (Cohen, Durstenfeld, and Roehm 2014), and direct SARS-CoV-2 infection of the inner ear has been observed (Jeong et al. 2021), the extent to which COVID-19 is related to audio-vestibular sequelae remains unclear. Recent systematic reviews estimate the prevalence of post-COVID-19 hearing loss symptoms at around 3-4%, and post-COVID-19 tinnitus symptoms at around 5-10% (Almufarrij and Munro 2021; Beukes et al. 2021; Jafari, Kolb, and Mohajerani 2022; Lough et al. 2022). Lough et al. (2022) estimated the prevalence of post-COVID-19 rotatory vertigo to be 2.4%. Most of the studies included in the reviews used self-report metrics, and the quality of these studies, where judged, was mostly considered "fair" (i.e. results deemed to be unbiased despite missing details). A systematic review from Meng et al. (2022) concluded that it is still unclear whether COVID-19 increases the risk of sudden sensorineural hearing loss. The global, and ongoing, prevalence of COVID-19 (WHO 2022), means that it is crucial to increase our understanding of whether and how COVID-19 affects hearing.

The considerable challenges associated with conducting research during the COVID-19 pandemic, alongside the need for rapid publication of pandemic-related research, has meant that studies to date often feature understandable but significant limitations (Ioannidis et al. 2022; Kapp et al. 2022). Case-control studies investigating COVID-19 and hearing often show bias in selection of the control group or lack of details about the groups' characteristics or selection. Small sample sizes are also common, as is incomplete reporting of methodology or results and lack of long-term follow-up. Within the bounds of these limitations, results from case-control studies have been mixed. Some report auditory deficits in COVID-19 patients, such as reduced otoacoustic emissions (Daikhes et al. 2020; Kokten et al. 2022; Mustafa 2020) and increased hearing thresholds (Gedik et al. 2021; Kokten et al. 2022; Mustafa 2020). Others find no significant impact of COVID-19 on auditory symptoms or hearing thresholds (Dror et al. 2021; Taitelbaum-Swead et al. 2022). While Dorobisz et al. (2023) found reduced auditory function on a range of measures in a large group of patients with long-COVID versus healthy controls, the long-COVID group were

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#### **KEYWORDS**

COVID-19; SARS-Cov2; hearing loss; auditory function



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selected on the basis of reporting post-COVID-19 hearing impairment, significantly limiting any conclusions that can be drawn from the comparison.

Other studies have focussed on differences between self-report measures in COVID-19 participants and controls, again with mixed results. Saunders et al. (2022) found that those who had had COVID-19 were more likely to report new or worse auditory symptoms compared to controls. However, AlJasser et al. (2022) found no significant difference in self-report of hearing or tinnitus symptoms between their COVID-19 and control groups, though their COVID-19 group were more likely to report rotatory vertigo (which is consistent with vestibular dysfunction). While the Saunders et al. (2022) data are compelling due to a large group size, inclusion of control group and inclusion of both pre- and post-COVID-19 data, the authors themselves highlight the potential for bias, inconsistency and inaccuracy in selfreport data, and hence the danger of drawing conclusions about causality (see also Saunders et al. 2023).

The present study overcomes many of the limitations present elsewhere. A relatively large sample of participants who were hospitalised as a result of COVID-19 infection was compared with a well-matched control group who had also been hospitalised but had never had COVID-19. Care was taken to recruit well-described, unbiased samples, and these groups were tested well beyond the typical COVID-19 recovery window. The use of a mobile research van for testing helped to remove barriers to participation, with the goal of increasing the diversity of participants. A comprehensive battery of auditory test measures was undertaken to thoroughly assess auditory ability and to isolate the specific loci of any COVID-19-related disorder. The combination of objective, behavioural and self-report measures recorded within the same set of participants represents the most comprehensive and thorough contribution from a single auditory study to date.

The protocol and hypotheses for the study were pre-registered (Guest et al. 2021). For each outcome measure, the prediction was that COVID-19 participants would show a deficit relative to control participants.

### **Materials and methods**

### Participants

Ninety-seven participants took part in the study; 57 in the COVID-19 group and 40 in the control group. Groups were matched for age, gender, body mass index (BMI) and time since hospital admission (see Table 1 for summary).

Information about specific COVID-19 variants was not available, and information about vaccine status was not sought. Extensive details of participant health and demographic characteristics can be found in theSupplementary Materials in the online repository for the project (https://osf.io/rc5fu/).

	Control group	COVID-19 group
N	40	57
Median age (and IQR), in years	57.5 (20.5)	58 (21)
Gender (female/male/other)	18/22/0	20/37/0
Mean BMI (and SD)	29.3 (6.8)	31.6 (6.3)
Majority ethnic group	White (95%)	White (81%)
Mean time since hospital admission (and SD), in months	9.1 (6.0)	10.7 (3.0)

Participants were recruited primarily via the Cross Speciality Research Nursing team at the Manchester NHS Foundation Trust using inpatient and outpatient clinic hospital records. Additional participants were recruited via word of mouth and advertising. Advertisements for the study referred only to experience of hospitalisation and omitted mention of hearing health to avoid biasing responses. Inclusion criteria for participation were: aged between 18 and 70 years old; admitted to hospital at least once (but no more than twice) in 2020-2021; and no self-report of profound hearing loss. For inclusion in the COVID-19 group, participants must have been hospitalised for COVID-19. For inclusion in the control group, participants must have been hospitalised with any other (i.e. non-COVID) illness, and must not knowingly have had COVID-19 at any time. Control participants were admitted for a range of illnesses, predominantly for respiratory conditions (25 out of 40 participants). Details of reasons for hospitalisation can be found in the Supplementary Materials. The study was approved by the London Central NHS Research Ethics Committee (ref: 21/PR/0137).

### Measures

Health and demographic data were collected by experimenters at the beginning of test sessions. Otoscopic examination and tympanometry were performed prior to all testing. Tympanometry was recorded with an Interacoustics Titan device, using a 226 Hz probe tone.

Outcome measures were categorised into three broad domains: cochlear function, neural function (peripheral and central) and auditory perception. Measures are described in detail below and are summarised in Table 2. Each measure was conducted in both ears where possible.

### **Cochlear function**

- i. Pure-tone audiometry (PTA). Testing took place in a sound-treated booth. Data collection was performed according to British Society of Audiology recommended procedures (British Society of Audiology 2018) at air conduction frequencies of 0.25, 0.5, 1, 2, 4, 8, 12.5 and 16 kHz and bone conduction frequencies of 0.5, 1 and 2 kHz, with appropriate masking applied to the non-test ear according to the recommended procedures. Testing took place using either an Interacoustics Callisto or Maico audiometer, with appropriately calibrated circumaural headphones (DD450 or HDA 300 respectively).
- ii. Distortion product otoacoustic emissions (DPOAEs). Measured using primary tones labelled f1 and f2, with a ratio (f2/f1) equal to 1.22. The following f2 frequencies were measured; 0.5, 1, 2, 4, 8 and 10 kHz (primary tone intensity levels used for f1 = 65 dB SPL and f2 = 55 dB SPL). This was recorded using the Interacoustics Titan device. For each frequency, a total recording time of 35 s was used, with frequencies tested in a descending order.

### Neural function

i. Acoustic reflex thresholds (ARTs). Recorded ipsilaterally with the Interacoustics Titan device, in automatic screening mode, with threshold criterion set to sensitive (0.03 ml). Measured using wideband evoking stimulus (spectral properties: "As per 'Broadband noise' specified in IEC 60645-5, but with 500 Hz as lower cut-off frequency"), with a 226 Hz

Table 2. Summary of outcome measures and their basic characteristics.

Hypothesis ID	Measure	Basic characteristics
Cochlear function		
1	Standard-frequency pure-tone audiometry (PTA) thresholds	Mean of thresholds at 0.25 to 8 kHz
2	Extended high-frequency (EHF) audiometry thresholds	Mean of 12.5 and 16 kHz thresholds
3	Standard-frequency distortion product otoacoustic emission (DPOAE) amplitudes	Mean of amplitudes at 0.5 to 8 kHz
4	EHF DPOAE amplitude	Amplitude at 10 kHz
Neural function		
5	Acoustic reflex threshold (ART)	Threshold for broadband (BB) noise elicitor using 226 Hz probe tone
6	Auditory brainstem response (ABR) wave I amplitude	Peak-trough amplitude
7	ABR wave I-V inter-peak interval	Interval between wave I peak and wave V peak
Auditory perception	on	
8	Digits-in-noise (DiN) signal-to-noise ratio for 71% correct (SNR71%)	Monaural threshold for identification of digits in speech-shaped noise
9	Speech, Spatial and Qualities of Hearing scale (SSQ12) change score	Sum of the 12 change scores
10	Tinnitus change score	Binary outcome: tinnitus onset/worsened vs tinnitus stable/absent

probe tone. Presentation started at 60 dB HL automatically increasing in 5 dB steps until two responses meeting the 0.03 ml criterion were observed at a single presentation level. Presentation stopped automatically once threshold was found or a maximum 100 dB HL presentation level was reached. The procedure was repeated twice and an additional, third time if there was  $\geq$ 10 dB difference between the first two threshold measurements.

Auditory Brainstem Response (ABR). Testing took place in ii. a sound-treated booth. ABRs were recorded using the Interacoustics Eclipse with ER3A insert phones. Appropriate correction for the sound wave delay due to the length of the insert tubing was included in the clinical interface. Stimuli were monaural 80 dB nHL broadband clicks presented at a rate of 11.1/sec. A two-channel recording was performed between the high forehead and both mastoids, using the ipsilateral mastoid recording when reporting results for a given ear. The ground electrode was on the low forehead. Online band-pass filtering of the EEG signal was applied between 0.1 and 2 kHz. A recording window of 0-15 ms was applied. The procedure was stopped after 5000 accepted epochs were recorded (with online artefact rejection of ±40 µV). Participants were in a reclined armchair for the duration of testing and instructed to keep their eyes closed, stay relaxed and to sleep if possible.

### Auditory perception

i. Digits-in-noise (DiN) signal-to-noise ratio (SNR) for criterion performance of 71% (SNR71%) correct responses (Smits, Kapteyn, and Houtgast 2004). Testing took place in a sound-treated booth. Digit-triplet stimuli were presented monaurally via TDH 39 headphones driven by a Cakewalk UA25 EX sound card, with presentation controlled by custom MATLAB (The MathWorks Inc. 2021) code and listener responses delivered via mouse and screen. In each trial, three consecutive digits (excluding the digits with two syllables, zero and seven) were spoken by a female British-English talker. A speech-shaped-noise masker was fixed at a level of 70 dB SPL while the level of the digit-triplet targets varied adaptively. Two digits out of three had to be entered correctly, in the correct order, for a trial to be scored as correct and a two-down one-up stepping rule applied (therefore tracking the 71% correct point on the psychometric function). The adaptive track had four initial turnpoints (6 dB step size) and six threshold turn-points (2 dB step size), with a starting SNR of 6 dB. SNR71% was calculated as the average of the SNRs at the final six turnpoints. The ear to be tested first was randomly selected per participant. Participants were provided with a short practice run before data collection began.

- ii. The short form of the Speech, Spatial and Qualities of Hearing scale, the SSQ12 (Noble et al. 2013), consists of 12 items requiring participants to indicate how easily they are able to perform or experience a range of everyday listening scenarios, using a scale of 0 to 10. Additionally, participants were asked to indicate whether their ability to perform or experience each scenario was worse, the same, or better compared to one month prior to their hospitalisation. (See the questionnaire section below for full details of the scoring of questionnaire responses.)
- iii. Tinnitus change score. A binary change score was assigned to each participant to indicate whether or not tinnitus had worsened following hospitalisation. (Tinnitus was defined as prolonged spontaneous tinnitus, i.e. tinnitus that occurs spontaneously and lasts for longer than 5 min). Participants' tinnitus was coded as having worsened (a tinnitus change score of 1) in any instance where (a) it was not present before hospitalisation but had occurred since, (b) it was occurring more frequently currently than before hospitalisation or (c) it was now present in both ears where previously it had only been in one. In all other cases participants were assigned a tinnitus change score of 0. Information about participants' experiences of tinnitus was collected at the beginning of test sessions, as detailed in the sections below.

### Questionnaires and other self-report measures

In addition to the SSQ12, all participants also completed the following questionnaires:

- (i) Fatigue Assessment Scale (FAS, Michielsen et al. 2004). Participants completed this with reference to their present experiences at the time of taking part in the study.
- (ii) Impacts of Illness and Hospitalisation (IIH). A custom, non-standardised questionnaire to assess impacts of illness and hospitalisation on social contact, loneliness, sleep, irritability, exercise, financial anxiety and depression/anxiety and depression (see Supplementary Materials for full details).

Participants also completed each of the following questionnaires if they met criteria for having experienced relevant symptoms, as defined in the section below:

- (iii) Dizziness Handicap Inventory (DHI; Jacobson and Newman 1990)
- (iv) Hearing Handicap Inventory for Adults (HHIA; Newman et al. 1990)
- (v) Tinnitus Handicap Inventory (THI; Newman, Jacobson, and Spitzer 1996)

Before testing began, all participants provided information about their health and their experiences of illness and hospitalisation. For experiences of dizziness, hearing difficulties and tinnitus, participants provided information for both their current experience and that in the period of time before getting ill and going into hospital. For tinnitus, participants were provided with a definition of tinnitus and asked whether they had ever experienced it, whether the experience was for longer than 5 min at a time, and whether it occurred spontaneously (i.e. not only due to infection or noise exposure). Participants who reported prolonged, spontaneous tinnitus were additionally asked how often it occurred (with response options of "Most or all of the time", "A lot of the time" and "Some of the time"), if it affected one or both ears and if the tinnitus pulsed. For hearing, participants were asked if they had any difficulty with their hearing, if they found it very difficult to follow a conversation in the presence of background noise and whether the difficulty affected one or both ears. For dizziness, participants were asked whether they suffered from attacks of dizziness in which things seemed to spin around them and whether they suffered from attacks of dizziness in which they seemed to move.

Which additional questionnaires participants were subsequently presented with was dependent on the responses given to the previous sets of questions. Participants were presented with the DHI if they had experienced attacks of dizziness, with the HHIA if they reported having experienced difficulty with their hearing and with the THI if they reported having experienced prolonged spontaneous tinnitus.

Each of the SSQ12, DHI, HHIA and the THI questionnaires were modified to include an additional metric for each item, to identify recent changes in experience. Directly following each standard questionnaire item, respondents were asked to indicate whether their current experience of the phenomenon in that item was "worse", "the same" or "better" than it was one month prior to hospitalisation. For analyses, these responses were assigned a value of 1, 0, -1, respectively, and summed to provide an overall "change score".

Measures are listed above to correspond with their order of appearance in the hypotheses listed in Table 1. The order in which tests were completed during test sessions was typically: Tympanometry, ARTs, DPOAEs, PTA, ABRs and DiN. Participants then completed questionnaires at the end of the session.

### Procedure

Test sessions were completed either in a bespoke auditory mobile research van or in a lab on site at the University of Manchester, depending on participants' availability and preference. When testing in the van, the tester would typically drive to, park and test outside the participants' homes. The van included a singlewalled sound-treated booth, and measurements of background noise at each location never exceeded 30 dB A. Background noise measurements were taken at the start of the test session using a type 2 sound level meter located where the centre of the participant's head would be located. During the test, the experimenter (in the non-sound-treated control booth) would subjectively monitor noise levels for any aberrations (e.g. the rare occurrence of a large vehicle driving past) and would wait for the noise to cease before recommencing testing. The on-campus lab contained a double-walled sound-treated booth. Sixty-six participants were tested in the van (40 COVID-19; 26 controls) and 31 participants were tested in the lab on campus (17 COVID-19; 14 controls). All testing was conducted by two experimenters (authors AV and IJ).

All participants completed the same procedures, regardless of experimental group or testing environment (the range of questionnaires completed varied according to participants' experiences, as detailed above). Testing was completed in a single session, typically lasting around 2 h. Participants were compensated for their time at a rate of £10 per hour.

### Pre-processing

For all analyses, data points were averaged across ears per participant. Where data were missing for one ear, data from the single ear were used in place of the average across ears for that participant. The number of participants contributing data from both ears or from only one ear for each outcome measure can be seen in Table 3. Analysis of ABR data was performed in two steps, firstly using an algorithm to automatically detect peaks and troughs, followed by visual inspection and manual correction of misidentified peaks. Where no peak was observable in the waveform, an amplitude value of 0 was assigned, and no latency value was assigned. For ART/PTA/DPOAE measurements that exceeded the limits of the equipment an appropriate floor or ceiling value was used. In the questionnaire data, for cases where participants were not required to complete a questionnaire (if a participant did not report any experience of dizziness, for example, they would not have been given the DHI to complete) they were assigned a change score of "0" in analyses to reflect the fact that hospitalisation had not had any impact on their experience of problems or symptoms. Further information about pre-processing of data can be found in the Supplementary Materials.

### Analyses

All processing and analyses were performed in R (R Core Team 2022), except for the processing of ABR data and automated peak-detection, which was conducted in MATLAB (The MathWorks Inc. 2021). Analyses are fully reproducible using the openly available code and de-identified data in the online repository for the project, which can be found at https://osf.io/rc5fu. Confirmatory analyses were pre-registered (Guest et al. 2021).

### **Confirmatory** analyses

For our continuous outcome measures, ANCOVA was performed with participant group as a between subject factor, and with age, gender and number of nights spent in hospital as covariates.

For our single outcome measure with a binary outcome, change in tinnitus (Hypothesis 10), logistic regression was performed with participant group as a between subject factor, and with age, gender and number of nights spent in hospital as covariates.

 Table 3. Summary of the number of participants contributing data in each test and the number of participants who completed each questionnaire.

Measure	Participant group	Data from both ears	Data from one ear only	Total N
PTA (standard frequencies)	Control	40	0	40
• •	COVID-19	57	0	57
PTA (EHF)	Control	40	0	40
	COVID-19	57	0	57
DPOAE (standard frequencies)	Control	39	1	40
	COVID-19	52	4	56
DPOAE (EHF)	Control	39	1	40
	COVID-19	52	4	56
ART	Control	36	3	39
	COVID-19	53	3	56
ABR, wave I amplitude	Control	28	7	35
	COVID-19	36	18	54
ABR, wave I–V latency	Control	28	7	35
	COVID-19	35	19	54
Digits-in-noise	Control	40	0	40
	COVID-19	57	0	57
Questionnaires				
DHI	Control			15
	COVID-19			23
FAS	Control			40
	COVID-19			57
HHIA	Control			18
	COVID-19			26
IIH	Control			40
	COVID-19			57
SSQ12	Control			40
	COVID-19			57
THI	Control			9
	COVID-19			8

### N per test

Table 3 summarises the number of participants included in statistical analyses for each test, and whether they contributed data from one or both ears. With one exception, missing test data for DPOAE and ART was due to either the presence of cerumen prohibiting testing and/or inability to obtain an adequate seal. The exception was one participant who requested to stop the test during data collection for ART. For the ABR wave I amplitude data, total missing ears consisted of 22 ears not tested due to cerumen, 15 due to an equipment fault (described fully in the Supplementary Materials), three which were excluded following manual inspection of the waveform revealing excessive noise, and one from a participant who found the experience uncomfortable and requested to stop before data were collected. For the ABR wave I-to-V interval data, missing ears were the same as for the amplitude data, plus an additional one ear which was not included in the analysis due to there being no identifiable wave I peak.

### Results

Summaries for the models used for each hypothesis can be found in Tables 4 and 5.

### Hypotheses 1 & 2: PTA thresholds at standard frequencies (0.25 to 8 kHz) and EHF (12.5 kHz)

Pure-tone audiograms and average thresholds are shown in Figure 1. A similar pattern of mild, high-frequency loss is present in both experimental groups. No statistically significant differences were found between groups at either standard or extended high frequencies. Age was significantly associated with higher thresholds at both standard (F(1, 92) = 39.66, p < 0.001; Eta2 (partial) = 0.30) and extended high frequencies (F(1, 92) = 156.57, p < 0.001; Eta2 (partial) = 0.63). All other *ps* were > 0.05 and can be found in Table 4.

## Hypotheses 3 & 4: DPOAE amplitudes at standard frequencies (0.5 to 8 kHz) and EHF (10 kHz)

Mean DPOAE amplitudes for standard and extended high frequencies are shown in Figure 2. No statistically significant differences between COVID-19 participants and controls were observed for DPOAE amplitudes, at either standard or extended high frequencies. Age was significantly related to lower amplitudes at both standard (F(1, 91) = 53.63, p < 0.001; Eta2 (partial) = 0.37) and extended high frequencies (F(1, 91) = 38.54, p < 0.001; Eta2 (partial) = 0.30). All other *ps* were > 0.05, full details are shown in Table 4.

### Hypothesis 5: ARTs

Mean ARTs for both experimental groups are shown in Figure 3. Means and distributions of thresholds are similar across groups. No statistically significant differences were found between groups for ARTs. Greater age was associated with a significant increase in thresholds (F(1, 90) = 8.18, p = 0.005; Eta2 (partial) = 0.08). All other *ps* were > 0.05 and can be found in Table 4.

### Hypothesis 6: ABR wave I amplitude

Peak-to-trough amplitudes for wave I and intervals for wave I to wave V peaks are shown in Figure 4, as are waveforms for the grand means for each experimental group.

Amplitudes for wave I did not exhibit a statistically significant difference between the COVID-19 group and the control group. Wave I amplitudes were significantly larger for women than for men (F(1, 84) = 9.93, p = 0.002; Eta2 (partial) = 0.11) and significantly reduced with age (F(1, 84) = 45.03, p < 0.001, Eta2 (partial) = 0.35). All other *ps* were > 0.05, full details are shown in Table 4.

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Table 4.	Model	summaries	for	Hypotheses	1	to	9.
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Hypothesis	Parameter	Sum of squares	df	Mean square	F	р	$\eta_p^2$
1. PTA (standard)	(Intercept)	257.05	1	257.05	2.61	0.110	0.03
	Gender	8.61	1	8.61	0.09	0.768	0.00
	Age (years)	3911.49	1	3911.49	39.66	< 0.001	0.30
	Length of stay in hospital	3.06	1	3.06	0.03	0.860	0.00
	Participant group	151.22	1	151.22	1.53	0.219	0.02
	Residuals	9074.03	92	98.63			
2. PTA (EHF)	(Intercept)	9478.43	1	9478.43	27.90	<0.001	0.23
	Gender	1217.99	1	1217.99	3.58	0.061	0.04
	Age (years)	53,194.40	1	53,194.40	156.57	< 0.001	0.63
	Length of stay in hospital	931.72	1	931.72	2.74	0.101	0.03
	Participant group	96.39	1	96.39	0.28	0.596	0.00
	Residuals	31,257.49	92	339.76			
3. DPOAE (Standard)	(Intercept)	321.25	1	321.25	19.00	<0.001	0.17
	Gender	2.05	1	2.05	0.12	0.729	0.00
	Age (years)	906.90	1	906.90	53.63	< 0.001	0.37
	Length of stay in hospital	18.16	1	18.16	1.07	0.303	0.01
	Participant group	7.31	1	7.31	0.43	0.512	0.00
	Residuals	1538.95	91	16.91			
4. DPOAE (EHF)	(Intercept)	5.63	1	5.63	0.23	0.635	0.00
	Gender	28.32	1	28.32	1.14	0.288	0.01
	Age (years)	954.35	1	954.35	38.54	< 0.001	0.30
	Length of stay in hospital	1.40	1	1.40	0.06	0.812	0.00
	Participant group	6.59	1	6.59	0.27	0.607	0.00
	Residuals	2253.64	91	24.77			
5. ART	(Intercept)	27,404.71	1	27,404.71	236.49	< 0.001	
	Gender	0.02	1	0.02	0	0.989	0
	Age (years)	947.79	1	947.79	8.18	0.005	0.08
	Length of stay in hospital	81.94	1	81.94	0.71	0.403	0.01
	Participant group	24.80	1	24.80	0.21	0.645	0
	Residuals	10,429.37	90	115.88			
6. ABR, wave I amplitude	(Intercept)	887,135.37	1	887,135.37	137.88	< 0.001	0.62
	Gender	63,888.93	1	63,888.93	9.93	0.002	0.11
	Age (years)	289,719.07	1	289,719.07	45.03	< 0.001	0.35
	Length of stay in hospital	7670.80	1	7670.80	1.19	0.278	0.01
	Participant group	1291.07	1	1291.07	0.20	0.655	0.00
	Residuals	540,452.08	84	6433.95			
7. ABR, wave I–V interval	(Intercept)	58.15	1	58.15	963.38	< 0.001	0.92
	Gender	0.17	1	0.17	2.81	0.097	0.03
	Age (years)	0.52	1	0.52	8.64	0.004	0.09
	Length of stay in hospital	0.07	1	0.07	1.10	0.297	0.01
	Participant group	0.07	1	0.07	1.12	0.293	0.01
	Residuals	5.07	84	0.06			
8. DiN	(Intercept)	1408.43	1	1408.43	567.35	< 0.001	0.86
	Gender	5.26	1	5.26	2.12	0.149	0.02
	Age (years)	37.54	1	37.54	15.12	< 0.001	0.14
	Length of stay in hospital	0.09	1	0.09	0.04	0.848	0
	Participant group	1.88	1	1.88	0.76	0.387	0.01
	Residuals	228.39	92	2.48			
9. SSQ12	(Intercept)	28.89	1	28.89	2.34	0.13	0.03
	Gender	48.44	1	48.44	3.92	0.051	0.04
	Age (years)	5.57	1	5.57	0.45	0.504	0.04
	Length of stay in hospital	7.34	1	7.34	0.59	0.443	0.01
	Participant group	59.25	1	59.25	4.79	0.031	0.05
	Residuals	1125.56	91	12.37	7.7 2	0.051	0.05

Table 5.	Model	summary	for	Hypothesis	10,	change	in	tinnitus.
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Hypothesis	Parameter	Fit	В	Ζ	р	β
10. Change in tinnitus	(Intercept)		-39.12	0	0.995	-39.37
-	Gender		18.6	0	0.997	18.6
	Age (years)	-0.01	0	0.893	-0.07	
	Length of stay in hospital		0.01	0	0.705	0.12
	Participant group		18.62	0	0.996	18.62
	AIC	35.18				
	BIC	48.05				
	Tjur's R2	0.07				
	Sigma	0.52				
	Log loss	0.13				

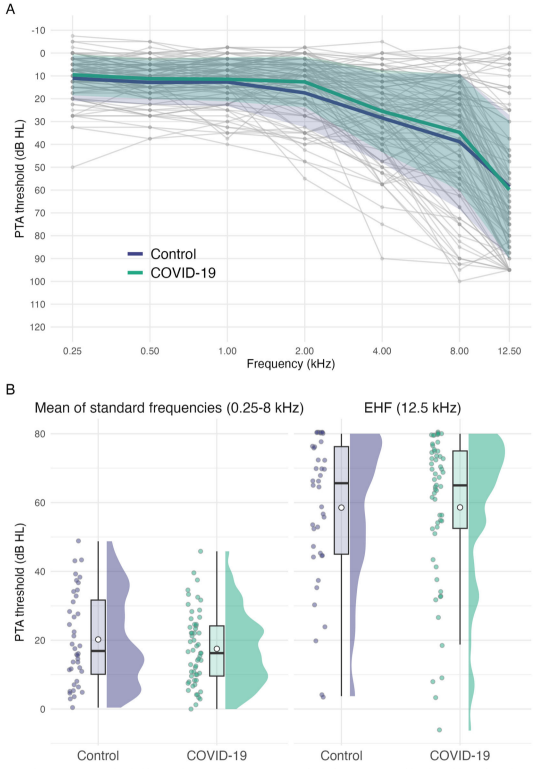


Figure 1. Panel A: Air conduction pure-tone thresholds. Grey lines and points represent individual participants. Bold, coloured lines show the means for each group at each frequency. Shaded ribbons around the bold lines show 1 SD from the mean. Panel B: Mean air conduction thresholds. Mean of standard (0.25–8 kHz) and extended high (12.5 kHz) frequencies. Jittered, coloured points show the raw data. Boxplot whiskers show 1.5 times the interquartile range. The hollow point inside boxplots shows the mean. Distribution curves show the probability density.

### Hypothesis 7: ABR wave I-V inter-peak interval

### Hypothesis 8: DiN SNR71%

The wave I–V inter-peak interval was not statistically significantly different between COVID-19 participants and controls. Age was significantly associated with a shortening of this interpeak interval (F(1, 84) = 8.64, p = 0.004; Eta2 (partial) = 0.09). All other *ps* were > 0.05, full details are shown in Table 4. DiN SNR71% SNRs are shown in Figure 5. No statistically significant difference in thresholds was observed between COVID-19 participants and controls. Age was significantly related to increased thresholds (F(1, 92) = 15.12, p < 0.001; Eta2 (partial) = 0.14). All other *ps* were > 0.05, full details are shown in Table 4.

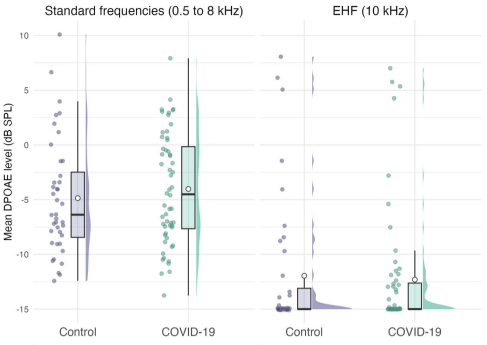


Figure 2. DPOAE levels: Means of standard (0.5 to 8 kHz) and extended high (10 kHz) frequencies. Jittered, coloured points show the raw data. Boxplot whiskers show 1.5 times the interquartile range. The hollow point inside boxplots shows the mean. Distribution curves show the probability density.

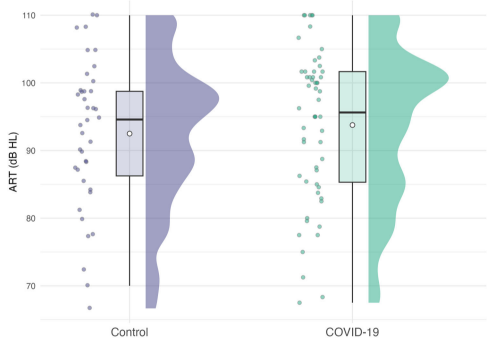


Figure 3. Mean ART. Jittered, coloured points show the raw data. Boxplot whiskers show 1.5 times the interquartile range. The hollow point inside boxplots shows the mean. Distribution curves show the probability density.

### Hypothesis 9: SSQ12 change score

Change scores for the SSQ12 are shown in Figure 6. Distributions for both groups are concentrated around 0, indicating that the majority of participants did not report any overall change in experience (the range of the scale shows the maximum and minimum scores possible; a total score of +12 would show a participant reported worsening of experience on every item).

SSQ12 change scores differed between the COVID-19 and control groups. On average, COVID-19 participants reported that their hearing abilities and experiences had worsened on about two to three items (M = 2.35) out of 12, compared to only around one item (M = 0.74) out of 12 in the control group. This difference is statistically significant (F(1, 91) = 4.79, p = 0.031; Eta2 (partial) = 0.05), but would not survive adjustment for multiple comparisons when considered collectively with the other outcomes measured. All other *ps* were > 0.05, full details are shown in Table 4.

The SSQ12 contains nine "pragmatic" subscales which categorise the area of difficulty each item is associated with (e.g. speech in noise, multiple speech streams, etc., with some items referring to more than one subscale). Exploratory analysis of these subscales

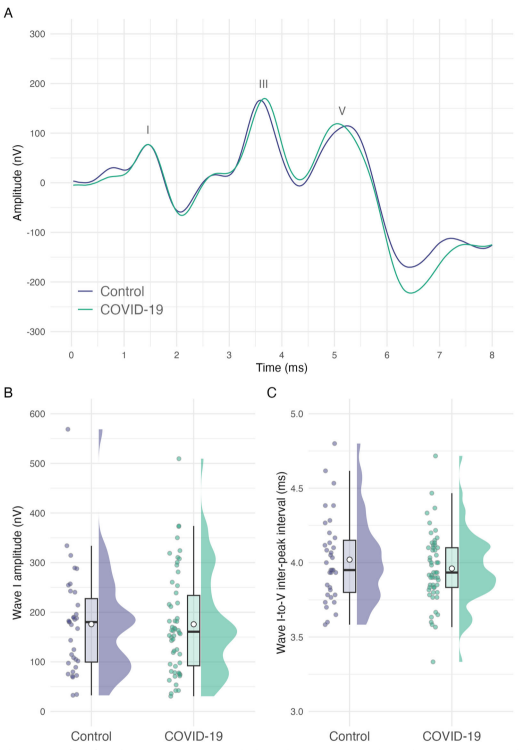


Figure 4. Panel A: Grand average waveforms for each group. Panel B: Wave I amplitude. Panel C: Wave I–V inter-peak interval. For panels B & C jittered, coloured points show the raw data. Boxplot whiskers show 1.5 times the interquartile range. The hollow point inside boxplots shows the mean. Distribution curves show the probability density.

showed that the largest difference between groups was for the item associated with listening effort. For this item/subscale, approximately three in 10 participants in the COVID-19 group reported an increase in effort since hospitalisation, compared to only one in 10 participants in the control group.

hospitalisation. Consequently, attempts to fit a logistic model to this data resulted in weak explanatory power (Tjur's  $R^2 = 0.07$ ). No statistically significant effects were observed (all ps > 0.05, full details can be found in Table 5).

### Hypothesis 10: change in tinnitus

Across the sample, only four participants, all from the COVID-19 group, reported that their tinnitus had become worse since

### **Exploratory** analyses

T-tests for questionnaire scores were performed to compare differences in responses between the COVID-19 group and control

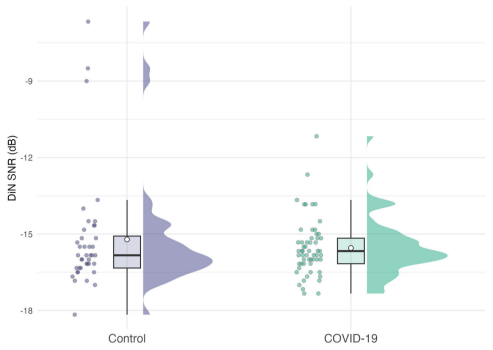


Figure 5. SNR threshold for digits-in-noise test. Jittered, coloured points show the raw data. Boxplot whiskers show 1.5 times the interquartile range. The hollow point inside boxplots shows the mean. Distribution curves show the probability density.

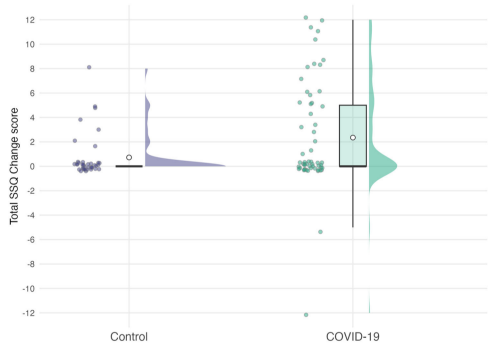


Figure 6. Total change scores on the SSQ12 questionnaire. Positive scores indicate a worsening of experience since hospitalisation. The range on the y-axis represents the minimum and maximum total scores possible. Jittered, coloured points show the raw data. Boxplot whiskers show 1.5 times the interquartile range. The hollow point inside boxplots shows the mean. Distribution curves show the probability density.

participant group. COVID-19 participants reported that their illness had had a greater overall impact on their lifestyle and mental state than control participants did, as assessed by the IIH questionnaire (t(94.18) = -3.58, p < 0.001; Cohen's d = -0.74). In the HHIA, COVID-19 participants reported that their hearing problems had worsened after hospitalisation to a greater extent than control participants did (t(80.09) = -2.93, p = 0.004; Cohen's d = -0.65). The COVID-19 group had a mean change score of 3.25 (out of a maximum of 25), compared to the control group mean of 0.82. For the remaining questionnaires (DHI, FAS, SSQ12 and the THI), comparisons of scores between COVID-19 and control groups produced p values > 0.05.

Participants provided ratings of their current general health and also for their general health as it was before being hospitalised. Ratings of pre-hospitalisation health did not significantly differ between groups. Both groups reported that their health was worse since hospitalisation than it was before, and the degree of change was significantly higher for COVID-19 participants than it was for controls (t(94.98) = 2.39, p = 0.019; Cohen's d = 0.49), mirroring the finding in the IIH that illness and hospitalisation had had a greater impact.

To assess any potential impact of the test environment (i.e. research van or university lab), all analyses performed for the confirmatory hypotheses were repeated with the inclusion of test environment as an additional covariate. No statistically significant impact of test environment was observed for any of the test outcomes.

All data collected were included in analyses regardless of tympanometry outcomes for individual ears. The proportion of ears categorised as non-normal (e.g. negative pressure, low compliance, etc.) was the same in each group (15% of total ears). To assess any potential impact of including non-normal tympanometry outcomes, all confirmatory analyses were repeated on a subset of the data containing only ears categorised as normal during tympanometry. No statistically significant differences between participant groups were observed on any test or questionnaire outcome. This pattern of results is identical to that reported above, other than for the SSQ change score, for which a marginally significant difference between groups was observed in the main analyses above. All exploratory analyses can be found in the Supplementary Materials.

### Discussion

The current study addressed a number of limitations found in existing studies of the effect of COVID-19 on hearing. Auditory measurements from COVID-19 participants were compared with those of tightly matched controls, following rigorous, pre-registered protocols and hypotheses. Bias was minimised at all stages, from advertising and recruitment of participants, through to the use of blinding where feasible in analyses of data. A comprehensive battery of auditory tests and questionnaires was undertaken, to probe the integrity of the auditory system at all levels. All outcome measures are reported and all findings are fully reproducible (de-identified data and code for analyses are publicly available, as detailed previously). We find no evidence that COVID-19 infection is associated with large-scale, long-term changes in auditory function.

This key finding is consistent with a recent comparison of hearing thresholds using PTA. Taitelbaum-Swead et al. (2022) controlled for age and duration of time between before-and-after tests and reported no significant impact of COVID-19 on hearing thresholds in PTA. While some studies have found differences in auditory function (hearing thresholds or otoacoustic emissions) associated with COVID-19, these have had multiple limitations such as bias in group selection (Dorobisz et al. 2023; Mustafa 2020), absence of control group (Kokten et al. 2022) and incomplete reporting of methods or results (Daikhes et al. 2020; Gedik et al. 2021), which make it difficult to draw firm conclusions from the data.

In the current work, no statistically significant differences were observed between groups across any of the confirmatory analyses of auditory tests. A statistically significant difference (for the raw p value, 0.031, uncorrected for multiple comparisons) was found for the self-reported change score associated with the SSQ12. That is, COVID-19 participants tended to report greater declines in perceived hearing ability than control participants following hospitalisation, as measured by how many of the listening experience items on the questionnaire they reported had got worse since hospitalisation. In terms of the statistical significance of the difference between groups, this was a moderately sized

effect (partial eta squared of 0.05). In absolute terms, a mean change score of 2.35 in the COVID-19 group and 0.73 in the control group is equivalent to participants reporting worsening, on average, on around 2 SSQ12 items out of 12 in the COVID-19 group and around 1 item out of 12 in the control group. Exploratory analysis of the pragmatic subscales in the SSQ12 showed the largest difference between COVID-19 and control groups to be in the category of "listening effort".

While the mean difference between groups is small for SSQ12 change scores, the discrepancy between lab-based and self-report measures is an intriguing one. Findings elsewhere suggest that self-report of post-COVID symptoms and experience is a complex issue, in which disentangling the influence of psychosocial factors and recall bias is a substantial challenge (Saunders et al. 2022, 2023). Nonetheless, an experience of increased listening effort would tie in with a model of post-COVID auditory symptoms relating to wider post-viral effects, such as fatigue and cognitive impairment (National Institute for Health and Care Excellence 2020), rather than a specific pathology of the auditory system. The mean FAS score for both groups met that scale's criterion for the presence of fatigue (threshold for the presence of fatigue is a total score of  $\geq$  22; the mean COVID-19 group score was 25.11, and the mean control score was 22.11). Ten of the COVID-19 group (18%) and 2 of the control group (5%) met the criterion for extreme fatigue (a total score of  $\geq$  35). In a similar pattern to the SSQ12 change scores, exploratory analysis of the HHIA change scores also revealed the COVID-19 group reported that their hearing problems had worsened to a greater degree than controls did, further indicating a greater perceived hearing deficit post-hospitalisation compared to the control group.

### **Deviations from protocol**

One deviation from the registered protocol is noted. The sample recruited for the study was smaller than the registered target size (n = 96 per group), meaning our analyses are not as highly powered as originally planned. This point is discussed further in the limitations section below.

### Limitations and future research

One potential limitation of the study is that the recruitment target of 96 people per group was not achieved. By the latter stages of the study, COVID-19 infection in the UK was so widespread that recruiting control participants who had never had the virus became a substantial challenge. Achieving the target sample would have increased statistical power, allowing for a greater degree of confidence in the outcomes of analyses and more accurate estimates of the size of significant effects. However, despite this limitation, distributions of data for each outcome show no obvious trends towards differences between groups, other than for self-reported SSQ12 and HHIA change scores. There is no indication in the data collected that larger group sizes would have led to statistically or clinically significant differences between groups on any other outcome measures.

The study aimed to achieve minimal bias between the two groups by ensuring that each had similar durations of recent hospitalisation, matching for age and gender and by imposing few other restrictions on inclusion. This resulted in unbiased but highly heterogeneous groups. Efforts were made to minimise bias in recruitment of the sample. Suitable candidates for the COVID-19 group were identified from lists of patients who had been admitted to COVID-19 and intensive care unit (ICU) wards. To obtain as close a match as possible for the control group, suitable candidates were identified primarily from lists of patients with non-COVID-19 respiratory illnesses and ICU admissions. Despite efforts to match characteristics across groups, differences in the experiences of the two groups remain a potential source of bias. Whether or not the participant had spent time in ICU was not systematically recorded, for example, and so any potential effects of this experience could not be confidently assessed (though no clear difference in ICU admission was apparent during collection of health and background information prior to test sessions). Similarly, further factors such as noise exposure, medical history and medications could also impact auditory function. While these factors were not routinely recorded in the current study, we have no reason to expect systematic differences between groups.

With increased prevalence of COVID-19 there is increased opportunity for studies to adopt within-participant designs. Direct assessment of individuals' hearing before and after COVID-19 infection would be a more sensitive measure than the between-groups design used in the current work.

Information about specific COVID-19 variants was not available, and information about vaccine status was not asked. However, examination of participants' hospitalisation dates shows that all of the COVID-19 group had already been hospitalised prior to the emergence of the Omicron (B.1.1.529) variant in the UK. Even the most recent participant to be hospitalised was admitted several weeks before the first reported case of the Omicron (B.1.1.529) variant in the UK. It also seems likely that a majority of the COVID-19 group were not vaccinated at the time of infection. Twenty-five percent of the group were hospitalised before the date of the first person to be vaccinated in the UK, and a further 25% were hospitalised within three weeks of this date, during which time only the very elderly and vulnerable were eligible to receive a vaccine.

### Conclusions

The global prevalence of COVID-19, and the importance of hearing for human functioning, means it is crucial to understand whether and how the virus might affect hearing. The existing literature for the effects of COVID-19 paints a mixed and inconsistent picture, likely due to significant limitations in the methodological approaches used, the populations studied and substantial risk of bias. The current work is a rigorous examination of the potential auditory impacts of COVID-19, in which bias was minimised at all stages. The range of outcomes measured is the most comprehensive to date. All hypotheses, as well as testing and analysis procedures, were pre-registered, and data and analyses are accessible and reproducible.

Results do not support the hypothesis that COVID-19 infection has a significant long-term impact on the auditory system. This is important and welcome public health information. Selfreport measures suggest it is not uncommon for patients to perceive changes in their hearing following COVID-19 infection, nor for them to attribute changes to the illness. Knowledge that self-perceived listening difficulties may have a basis beyond discernible physical changes in the auditory system can help health care professionals to provide appropriate counselling and management plans to support patients experiencing these difficulties.

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### **Authors' contributions**

The original study idea was conceived by authors KJM, CJP and HG. All authors contributed to further development of the protocols, procedures and pre-registration. Authors ASV and IRJ completed data collection. Author IRJ performed the data analysis. Authors IRJ and ASV prepared the manuscript. Author IRJ prepared the materials for open sharing of data and analyses. All authors provided critical intellectual feedback to successive versions of the manuscript. All authors read and approved the final version of the manuscript. Joint first authorship is shared between ASV and IRJ.

### **Disclosure statement**

No potential conflict of interest was reported by the author(s).

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### Data availability statement

Supplementary analyses, materials, code for analysis and de-identified data are available in the online repository for the project, which can be found at https://osf.io/rc5fu/, DOI: 10.17605/OSF.IO/RC5FU.

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