

1 *Running title: Long-term olfactory dysfunction due to COVID-19*

2
3 ORIGINAL CONTRIBUTION

4
5 **The COVANOS trial – insight into post-Covid olfactory dysfunction**
6 **and the role of smell training**

7
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SUMMARY (200 words)

Background: *Olfactory dysfunction is a cardinal symptom of COVID-19 infection, however, studies assessing long-term olfactory dysfunction are limited and no randomised-controlled trials (RCTs) of early olfactory training have been conducted.*

Methodology/Principal: *We conducted a prospective, multi-centre study consisting of baseline psychophysical measurements of smell and taste function. Eligible participants were further recruited into a 12-week RCT of olfactory training versus control (safety information). Patient-reported outcomes were measured using an electronic survey and BSIT at baseline and 12 weeks. An additional 1-year follow-up was open to all participants.*

Results: *218 individuals with a sudden loss of sense of smell of at least 4-weeks were recruited. Psychophysical smell loss was observed in only 32.1%; 63 participants were recruited into the RCT. The absolute difference in BSIT improvement after 12 weeks was 0.45 (95%CI: -0.69 to 1.59, $p=0.43$) higher in the intervention arm. 76 participants completed 1-year follow-up; 10/19 (52.6%) of participants with an abnormal baseline BSIT test scored below the normal threshold at 1-year, and 24/29 (82.8%) had persistent parosmia.*

Conclusions: *Early olfactory training may be helpful, although our findings are inconclusive. Notably, a number of individuals who completed the 1-year assessment had persistent smell loss and parosmia at 1-year. As such, both should be considered important entities of long-Covid and further studies to improve management are highly warranted.*

Key words: COVID-19, anosmia, parosmia, quality of life, olfactory training

105

106 INTRODUCTION

107

108 Shortly after the emergence of SARS-CoV-2, it became evident that sudden loss
109 of sense of smell is a cardinal symptom of Covid-19 and early recognition is key in
110 affected patients and healthcare workers in particular.⁽¹⁻⁴⁾ It is typically more common
111 in those with mild disease or who are otherwise asymptomatic⁽⁵⁾. To date, nearly 277
112 million cases of COVID-19 have been reported (22 December 2021), with 11.7 million
113 in the UK and 51.4 million in the USA.⁽⁶⁾ With an incidence of roughly two-thirds, over
114 150 million individuals, globally, will have lost their sense of smell during this
115 pandemic, including roughly 5 and 29 million in the UK and the USA, respectively.⁽⁷⁾

116 Encouragingly, the vast majority of patients will recover their sense of smell
117 within the first two months, on average; however, olfactory dysfunction has been
118 reported in patients even six-months after initial infection.⁽⁸⁻¹⁰⁾ In their assessment of
119 51 patients with acute smell loss beyond 7 days at 8 months, Renaud *et al* demonstrated
120 persistent hyposmia in 2 patients (3.9%).⁽¹¹⁾ Comparatively, another study has
121 demonstrated olfactory dysfunction in 46% of patients followed up beyond 1-year, with
122 functional anosmia in 7%.⁽¹²⁾ Altogether, the precise burden of long-term olfactory
123 dysfunction remains unknown but is likely substantial.

124 In the COVID-19 context, both the British Rhinological Society (BRS) and
125 Clinical Olfactory Working Group (COWoG) recommend olfactory training based on
126 existing evidence of its efficacy, particularly for post-viral olfactory dysfunction.⁽¹³⁻¹⁷⁾
127 While the use of oral and topical steroids was very controversial at the beginning of the
128 pandemic, and at the time of the planning of the trial, recent evidence indicates a
129 potential benefit. However, the evidence is not robust.^(18,19) In line with this, the BRS
130 further recommend oral steroids, steroid rinses, and omega-3 supplements whilst the
131 COWG acknowledge a potential role for oral and topical steroids and vitamin A
132 drops.^(16,17) Both emphasize the need to examine the use of further medical treatment
133 on a case-by-case basis with careful risk assessments undertaken.

134 Here, we aim to obtain long-term follow-up data of individuals with olfactory
135 dysfunction for at least four weeks prior to enrollment during the COVID-19 pandemic
136 and evaluate the efficacy of early olfactory training in a parallel, 2-arm, randomised
137 controlled trial.

138

139 MATERIALS AND METHODS

140

141 *Trial Design and Recruitment*

142 This study, entitled ‘COVID-19 and Anosmia’ (acronym: ‘COVANOS’) was
143 sponsored by University College London and conducted across four NHS trusts: Barts
144 Health NHS Trust, Guy’s and St. Thomas’, James Paget University Hospitals/Norfolk
145 and Norwich University Hospitals, and Wrightington, Wigan and Leigh NHS
146 Foundation Trusts. Ethical approval was obtained through the UK Health Research
147 Authority Research Ethics Committee (ref. 20/WM/0147). Participants were recruited
148 through trust-wide email and poster advertisements directed primarily toward
149 healthcare workers (HCWs), who were identified via surveys which were conducted
150 across all these NHS Trusts and results published separately.^(20,21)

151 Individuals with persistent and sudden loss of sense of smell (at least 4 weeks)
152 were invited to participate in the study. A positive COVID-19 test was not a
153 requirement for participation, as availability of testing was extremely limited at the
154 beginning of the pandemic when the trial was launched. However, information
155 regarding COVID-19 antigen and antibody testing were collected *post-hoc* from those
156 for who data were readily available. All participants underwent psychophysical smell
157 testing using the Brief Smell Identification Test (Brief Smell Identification Tests™ -
158 Cross-Cultural Smell ID Test, Sensonics Inc., US) A subgroup of participants also
159 underwent gustatory testing using Taste Strips (Burghart Messtechnik GmbH,
160 Germany). Participants also completed a validated electronic survey [*submitted for*
161 *publication*], which collected relevant demographic data, details of symptoms
162 experienced, co-morbidities and other Covid-19 related symptoms including olfactory
163 function assessment. This included self-rating of smell and taste function with the
164 corresponding prompts: ‘How would you rate your sense of smell today (0 being really
165 bad, 10 being completely normal)?’ and ‘How would you rate your sense of taste
166 (salt/sweet/sour/bitter/savoury) today (0 being really bad, 10 being completely
167 normal)?’ As well, participants were asked a series of quality of life (QoL)-related items,
168 which were scored on a 7-point Likert scale. These items were separated into 4
169 categories: the impact of their smell dysfunction 1) on their social and professional life,
170 2) with regards to eating habits, 3) on their sense of anxiety and 4) the extent to which
171 it was annoying.

172 Recruitment took place either in-person at designated clinics across the NHS
173 trusts or remotely through email and post, the latter due to lockdown measures. Where

174 relevant, all study materials were posted to the participants with additional
175 correspondence by email. Informed consent was obtained for all participants.

176 Those with a BSIT score of 8 or less (considered abnormal smell, as published
177 previously)⁽²²⁾ were further invited to participate in the smell training trial (RCT), which
178 consisted of randomisation to either undergo 12 weeks of olfactory training using
179 Sniffin' Sticks (Duft-Quartett, Burghart Messtechnik GmbH, Germany; treatment
180 group) or receive safety information only (control group). Eligible participants were
181 randomised 1:1. Both arms were followed up at 12 weeks with regular correspondence
182 by email throughout the duration of the trial to ensure compliance and safety. At the
183 end of the 12-week periods, participants completed a follow-up BSIT and electronic
184 'End of Study' survey.

185 All participants enrolled at baseline within the eligible timeframe, for whom a
186 valid email address was available, were invited to participate in 1 year follow-up
187 assessments. This included all participants irrespective of baseline BSIT result and RCT
188 participation. The follow-up included a final electronic survey and BSIT. In addition to
189 questions related to their sense of smell, which were identical to those in the baseline
190 and 12-week follow-up surveys, participants were also asked about any symptoms of
191 long-Covid, including fatigue, brain fog, chest pain, joint pain, amongst others.

192

193 *Statistical Methods*

194 The primary outcome was the absolute difference between the intervention and
195 control arms in BSIT score smell improvement, measured as a change from baseline at
196 12-weeks. Secondary outcome measures were quality of life in relation to anosmia and
197 COVID-19 infection, compliance, and safety of olfactory training in the intervention
198 arm and the identification of predictive biomarkers for clinical outcome. A total sample
199 size of 200 patients, 100 per arm, was calculated to detect the target standardised effect
200 size of 0.5 at the two-sided 5% significance level with 90% power, after allowing for
201 up to 15% dropout.

202 Descriptive statistical analysis was conducted on participant characteristics and
203 associations were evaluated using Chi-Square and Fisher's Exact tests, where
204 appropriate. Trial arms were compared using linear and logistic regression adjusted for
205 baseline score where absolute as well as standardised effect sizes and odds ratios (with
206 95% confidence intervals and *P*-values) are presented, respectively. Smell and quality

207 of life scores were compared at different time-points using the paired samples t-test and
208 differences between groups were assessed using the Mann-Whitney U test. All
209 statistical tests were performed on SPSS version 27.

210

211 RESULTS

212

213 *Recruitment and Enrolment*

214 A total of 227 participants were recruited into the study and completed the
215 baseline BSIT between 4th May 2020 and 4th January 2021. One participant withdrew
216 at this time. Eight participants were further excluded due to a lack of evidence of
217 persistent smell loss ascertained through the baseline questionnaire. A final cohort of
218 218 participants was included in subsequent analyses.

219 Seventy participants scored 8 or below at 4 weeks following onset of the loss of
220 sense of smell and were subsequently invited to participate in the smell training trial.
221 At this point, most participants (67.9%, 148/218) scored within the normal range of the
222 BSIT test at the required 4 weeks and were thus ineligible for the RCT. Of the 70
223 participants who were eligible, 63 were enrolled into the smell training trial with 7
224 declining participation. 12-week follow-up data was available from 51 participants: 26
225 intervention and 25 controls, respectively. Four participants in the treatment arm had
226 withdrawn their participation or were removed from the study due to non-compliance
227 with the olfactory training regimen; a further 3 participants were lost to follow-up. In
228 the control arm, there were no withdrawals nor removals whilst 5 participants were lost
229 to follow-up.

230 In addition, 169 of the 218 participants in the overall cohort were re-contacted
231 for further assessments after approximately 1-year (8-13 months depending on the time
232 of recruitment). Of these, 76 participants completed the electronic survey and 56
233 completed an additional BSIT. **Figure 1** presents the flow of participants through the
234 study.

235

236 *Baseline characteristics and potential predictors of baseline BSIT score*

237 Of the 218 participants recruited with a persistent loss of sense of smell and
238 eligible for analysis (self-reported, at least 4 weeks), 190 completed the baseline
239 questionnaire. The median age was 44.0 years (range 22–78), and 85.0% (163/189)
240 were female (see **Table 1**). 72.1% (137/190) were never-smokers with 22.1% (42/190)

241 having smoked previously and 5.8% (11/190) being current smokers. 73.2% (139/190)
242 of participants consume 1–14 units of alcohol per week, 6.3% (12/190) consuming 15–
243 21 units per week and 2.1% (4/190) consuming over 21 units per week and 18.4%
244 (35/190) having never consumed alcohol.

245 24.7% (47/190), 12.6% (24/190) and 8.9% (17/190) had a history of sinonasal
246 disease, asthma, and high blood pressure, respectively (**Table 2**). Of those with a history
247 of high blood pressure, 58.8% (10/17) had been treated with either angiotensin-
248 converting enzyme (ACE) inhibitors or angiotensin-II receptor blockers (ARBs). There
249 was no evidence of an association between demographic factors nor medical history
250 with an abnormal BSIT test at baseline.

251 With regards to COVID-19 status, 50.5% (96/190) had tested positive by PCR
252 test before recruitment at one month post-initial infection, and the remaining 49.5%
253 (94/190) were recruited upon experiencing a sudden-onset smell loss within the last 1-
254 2 months with a suspected COVID-19 infection (PCR testing was not readily available
255 at the beginning of the pandemic, when the isolated symptom of smell loss was not an
256 indication for testing). *Post-hoc* COVID antibody and antigen testing results were
257 obtained for a subgroup of participants. Of the sixty-five participants for whom
258 antibody testing results were readily available, fifty-three (81.5%) tested positive. For
259 those who had reported a positive COVID-19 antigen result at the time of recruitment,
260 87.5% (28/32) also had a positive antibody result. For those who had not undergone
261 COVID-19 antigen testing at the time of recruitment, 76.0% (19/25) had a positive
262 antibody result in the time thereafter.

263 All eligible participants had one or more symptoms in addition to the loss of
264 sense of smell at the time of onset, with 75.8 (144/190) of participants reporting a loss
265 of sense of taste. Other common symptoms were fatigue (70.5%, 134/190), aches and
266 pains (53.2%, 101/190), fever (38.9%, 74/190), shortness of breath (36.8%, 70/190),
267 persistent cough (32.6%, 62/190) and sore throat (30.0%, 57/190). Moreover, 26.8%
268 (51/190) reported nasal congestion, 14.2% (27/190) reported having experienced
269 metallic taste and 13.7% (26/190) reported a burning sensation in the nose or mouth
270 (**Table 2**). Whilst most symptoms were more common in those with abnormal BSIT
271 test at 4 weeks, there was strong evidence in terms of reporting of shortness of breath
272 ($p=0.011$), difficulty breathing ($p=0.037$), aches and pains ($p=0.015$) and chest pain
273 ($p=0.009$).

274 Regarding self-reported qualitative smell dysfunction (**Supplemental Table 1**),
275 41.0% (73/178) reported distorted smell, 25.3% (45/178) reported having experienced
276 phantom smells, and 6.7% (12/178) reported a heightened sense of smell. For most
277 participants, the change in smell occurred suddenly (69.7%, 106/172), whilst 23.0%
278 (35/172) reported the change occurring over days. For those who had smell issues,
279 67.1% (102/152) reported that the issue was consistent throughout the day, 18.4%
280 (28/152) reported that the issue fluctuates, occurring more often than not, and 14.5%
281 (22/152) reported that the issue occurs occasionally throughout the day with the
282 majority of the time being normal.

283 Regarding taste function, 30 participants from our first participating centre
284 underwent taste testing. Most participants had normal taste function with regards to
285 sweet (93.3%, 28/30), salty (96.6%, 28/29), sour (86.7%, 26/30) and bitter (96.7%,
286 29/30). We did not pursue taste testing for the remainder of the cohort due to the remote
287 nature of the study and logistical constraints, and due to the fact that these initial results
288 demonstrated that the underlying impairment was not due to an impaired taste function
289 (sweet, etc.) but rather to do with the perception of flavours, as a result of smell
290 dysfunction, which would not be appropriately captured with this measure.

291 Regarding smell function, the mean BSIT score at baseline was 9.1 (Std. Dev.
292 = 2.12) (**Table 3a**). 67.9% (148/218) had normal smell (BSIT 9-12), 24.4% (53/218)
293 had mild anosmia (BSIT 6-8), 7.8% (17/218) had moderate anosmia (BSIT 3-5). No
294 participants scored within the severe anosmia range (BSIT 0-2).

295

296 *Primary and secondary outcomes for early smell training at 12-weeks and at 1 year*

297 The mean BSIT score for both trial arms at 12-weeks was 7.9 (Std. Dev. 2.23)
298 (**Table 3a**). Considering the change in BSIT score from baseline to 12-week follow-up,
299 the absolute difference between the trial arms is 0.45 points (95% CI: -0.69 to 1.59,
300 $p=0.43$), which corresponds to a standardised effect size of 0.22 (95% CI: -0.34 to 0.77),
301 after adjusting for baseline BSIT score. This was a smaller observed effect than the
302 target standardised difference of 0.5, and in a smaller sample than planned (i.e. more
303 uncertainty). Although not significant, the odds were higher in the treatment arm,
304 compared to the control arm, of having normal smell following early olfactory training
305 after 12-weeks (OR=2.38, 95% CI: 0.73 to 7.76, $p=0.15$), after adjusting for baseline
306 BSIT score (**Table 3b**).

307 Of the participants who responded to the invitation for a 1-year follow-up, 19
308 participants, who completed the 12-week RCT, responded. At this time-point
309 (control=8, intervention=11), the absolute difference in the change in BSIT score
310 between the trial arms is 0.65 (95% CI: -1.01-2.31, $p=0.42$), which corresponds to a
311 standardised effect size of 0.31 (95% CI: -0.38-1.01), after adjusting for baseline BSIT
312 score. Similar to at 12-weeks, we observed increased odds of having normal smell at 1
313 year with olfactory training (OR=2.3, 95% CI: 0.37-14.61, $p=0.37$), after adjusting for
314 baseline BSIT score, however this was not statistically significant (**Table 3b**).

315 *Long Covid and Proportion of Patients with Persistent Anosmia and/or Parosmia at 1-* 316 *Year*

317 The median number of months between the 1-year follow-up and baseline
318 enrolment was 10 months (range 8-13). For all participants, who participated in the 1-
319 year follow-up, the mean BSIT score was 9.5 (Std. Dev. 1.71). 75.0% (42/56) scored
320 with the normal range, while 23.2 (13/56) and 1.8% (1/56) had mild and moderate
321 anosmia, respectively (Table 4a). The change in BSIT score from baseline was 0.2 (Std.
322 Dev. 1.77). When considering the RCT participants only (n=19), there were slight
323 improvements in BSIT scores in both the treatment (n=11) and control arms (n=8) at 1-
324 year compared to baseline (Table 4a). However, for both arms combined, only 47.4%
325 (9/19) scored within the normal smell range at 1-year.

326 In an exploratory analysis of potential predictors of psychophysical long-term
327 smell loss, neither gender, smoking/alcohol history, nor medical history were
328 associated with an abnormal BSIT test at 1-year in the responding cohort
329 (**Supplemental Tables 2 and 3**). However, evidence of associations between the
330 experience of certain COVID-19 symptoms at baseline and an abnormal BSIT result at
331 1-year was observed: aches and pains ($p=0.030$) and/or diarrhoea ($p=0.011$)
332 (**Supplemental Table 4**).

333 Regarding subjective measures of olfactory dysfunction, the mean change in
334 participants' sense of smell self-rating, from baseline to 1-year, was 1.39 (Std. Dev.
335 2.29). This did not correlate with the change in BSIT result (Spearman's correlation
336 coefficient = 0.11, $p=0.465$).

337 The overall rate of parosmia in the responding cohort at 1-year was 43.4%
338 (33/76). 24 of the 29 participants, who reported parosmia at baseline, continued to
339 experience this symptom at 1 year (**Table 4**). In addition, experience of parosmia at 1-
340 year was more likely in those with abnormal BSIT scores at baseline (OR=3.56, 95%

341 CI: 1.30-9.69, $p=0.013$) Additionally, we observed a correlation between parosmia at
342 1-year and an abnormal BSIT score at 1-year, which approached significance ($p =$
343 0.055).

344 Regarding phantosmia, 9 of the 18 participants in the responding cohort, who
345 reported the symptom at baseline continued to experience it at 1-year. Experience of
346 phantosmia at 1-year was more likely in those with abnormal BSIT scores at baseline
347 (OR=5.18, 95% CI: 1.51-17.7, $p=0.009$) and significantly correlated with an abnormal
348 BSIT score at 1-year ($p = 0.011$).

349 Considering all participants, who completed the 1-year survey irrespective of
350 RCT enrolment, 65.8% (50/76) reported experiencing at least one symptom of long
351 Covid, with extreme tiredness/fatigue (39.6%, 30/76) brain fog (25.0%, 19/76), joint
352 pain (21.1%, 16/76), insomnia (17.1%, 13/76) and heart palpitations (14.5%, 11/76)
353 being the most common. For the participants for whom a 1-year BSIT and survey result
354 were available ($n=56$), brain fog significantly correlated with an abnormal BSIT result
355 at 1-year ($p = 0.037$) (**Supplemental Table 5**).

356

357 *Changes in Quality-of-Life Measures at baseline and after 1-Year*

358 When comparing QoL scores at 1-year and at baseline, improvements (i.e.,
359 negative change) were seen for most items (**Table 5**). The evidence for these
360 improvements was most robust for items 1 (mean difference -1.0, 95% CI: -1.60 to -
361 0.49, $p=0.001$), 1a (mean difference -0.8, 95% CI: -1.62 to -0.05, $p=0.038$), 2b (mean
362 difference -1.4, 95% CI: -2.05 to -0.78, $p<0.001$), 2c (mean difference -1.0, 95% CI: -
363 0.25 to -1.67, $p=0.010$), 4 (mean difference -1.0, 95% CI: -1.69 to -0.31, $p=0.008$), 4a
364 (mean difference -0.9, 95% CI: -1.51 to -0.32, $p=0.004$) and 4b (mean difference -0.9,
365 95% CI: -1.53 to -0.22, $p=0.011$).

366 In an exploratory analysis of differences in the mean scores between those who
367 experience both anosmia and parosmia at baseline, compared to anosmia only, only two
368 items were significantly different: item 2b, “Because of the changes in my smell, I don’t
369 enjoy food or drinks as much as I used to” ($p=0.045$) and 4b, “The changes in my sense
370 of smell annoy me when I am eating” ($p=0.023$) (**Supplemental Table 6**).

371

372 DISCUSSION

373

374 Crucially, our study confirms that most individuals who experience olfactory
375 dysfunction secondary to proven and/or presumed COVID-19 infection will recover

376 their sense of smell within the first four weeks. Indeed, two-thirds of our participants
377 scored within the ‘normal’ range of the BSIT at enrollment. As such, while the target
378 for the study was to recruit 200 participants, we found that this would be infeasible
379 within the timeframe of the study due to the high recovery rate within the first four
380 weeks. This is in line with previous studies, which have reported 60-70% of COVID-
381 19 patients recovering their sense of smell within the first month.⁽²³⁻²⁵⁾ However, there
382 remains a subset of individuals who will experience persistent anosmia, as
383 demonstrated in our study. In those who responded to the 1-year follow-up, most of
384 those with persistent anosmia at baseline, i.e. at least 4 weeks, exhibited some degree
385 of hyposmia even after 1 year. Furthermore, 52.6% of the 19 RCT participants (both
386 arms combined), who responded at 1-year, saw no improvement in their sense of smell.

387 Regarding early olfactory training without steroids, valid conclusions cannot be
388 drawn regarding a potential benefit after 12 weeks due to the small number of
389 participants who were ultimately eligible and enrolled in the RCT. Although some
390 benefit may be gained, observed effect sizes were lower than those targeted in the study
391 design and respective power analysis. With regards to the minimal clinically important
392 difference (MCID) between the two groups, there has been no formal study assessing
393 this in the context of anosmia/parosmia. Whilst a previously reported MCID of at least
394 1.0 for the BSIT appeared to be useful in evaluating chronic rhinosinusitis before and
395 after endoscopic sinus surgery, it is unclear whether this is applicable for the current
396 study.⁽²⁶⁾ Altogether, further investigation is needed to determine the efficacy of this
397 treatment.

398 Evaluating 10-weeks of olfactory training either on its own or in conjunction
399 with oral corticosteroids, others have reported that only those in the latter group saw a
400 clinically significant improvement in their olfactory score.⁽²⁷⁾ This finding suggests that
401 the addition of steroids to early olfactory training may significantly improve the sense
402 of smell, as shown with 6-month olfactory training at 1 year.⁽²⁸⁾ At the start of the
403 pandemic when this trial was planned and registered there was significant concern
404 regarding the use of both oral and intranasal steroids in SARS-CoV-2 infection.
405 However, evidence now suggests that olfactory training together with topical
406 corticosteroids, including nasal lavage may be the best approach. Further prospective
407 trials are warranted to determine the efficacy of these approaches and re-evaluate some
408 of the consensus guidelines, as corticosteroids appear to be effective for other types of
409 post-viral olfactory loss.^(16,29,30)

410 A large proportion of the participants who responded at one year reported
411 experiencing parosmia and, to a lesser extent, phantosmia. This is in line, albeit higher
412 than a previous report, which observed a 43.1% prevalence of parosmia after 6
413 months.⁽³¹⁾ Importantly, parosmia is emerging as a key symptom of long-Covid and our
414 study suggests its increasing prevalence at one year which we further show correlates
415 significantly with an abnormal baseline and 1-year BSIT test which in itself correlates
416 with the long-term Covid symptom of brain fog. This underscores the neurological
417 insult that occurs in a subgroup of patients which then causes a persistent central
418 nervous symptom complex.

419 It is apparent that there is a significant number of individuals who may suffer
420 from persistent symptoms of parosmia which can be debilitating. Whilst certain
421 strategies are currently used in standard practice, such as sodium valproate or similar,
422 these largely rely on anecdotal evidence with a lack of randomised, controlled trials.
423 This poses as a crucial gap in the management of long-term olfactory dysfunction.
424 Furthermore, the mechanism of parosmia has yet to be elucidated in the context of
425 COVID-19 and why late-onset parosmia occurs is unknown. While some researchers
426 have explored the neuroinvasive capacity of the virus, other research indicates that the
427 infection of sustentacular cells or the presence of viral products in the
428 microenvironment may cause the observed neurological sequelae.⁽³²⁻³⁵⁾ It is likely that
429 the cause for the symptoms is multifactorial and further investigations are highly
430 warranted.

431 Regarding quality of life, there were some improvements over the 1-year period
432 for all participant assessed, however, the scores for several items were similar, which
433 may be due to the persistent negative impact of smell dysfunction on these aspects of
434 life, particularly regarding feelings of anxiety as well as the impact on eating. Crucially,
435 considering the proportion of our participants who reported experiencing parosmia at
436 the 1-year assessment, it is important to note the specific way this condition impacts
437 quality of life in comparison to anosmia/hyposmia. A number of our participants have
438 reported, anecdotally [free text option, Supplemental Table 7], the challenges they have
439 faced psychologically and emotionally due to parosmia, which can be seen in the
440 differences in QoL scores between those who experienced parosmia and anosmia,
441 compared to anosmia only. However, these findings may be confounded by other
442 aspects of COVID-19 infection and the ongoing pandemic that we were unable to
443 account for in this study. Indeed, items related to the impact of smell dysfunction

444 socially may be confounded by the changes in societal restrictions as part of the
445 COVID-19 pandemic response and less to do with objective and/or qualitative smell
446 loss.

447 Our RCT is limited by its sample size, as we were unable to recruit our intended
448 target due to the extremely high rates of smell recovery prior to 4-weeks post-onset.
449 Furthermore, due to the fact that COVID-19 testing was not readily available at the start
450 of the pandemic, not all subjects had formal proof of having had COVID-19 infection.
451 As well, relatively high drop-out rates were observed (13 participants did not complete
452 the RCT or were lost to follow-up). Potential non-compliance is also a limitation, which
453 was largely due to the need to conduct the study remotely, in general, to comply with
454 local safety guidelines. Most of the participants completed the BSIT remotely and
455 unsupervised; this may be influenced by external factors, such as a family member
456 providing help. As such, careful instructions were provided to the participants to
457 mitigate these and results should be considered within the study context. Non-
458 compliance may also be an issue with regards to the olfactory training RCT. Regarding
459 the control group, there may be a chance that these participants conducted ‘at-home’
460 olfactory training in any case, with this information so readily available on the internet
461 and through support organisations. We attempted to mitigate these by providing clear
462 instructions, communication with the participant during the study and the subsequent
463 exclusion of those determined to be non-compliant, making these potential biases less
464 likely. Another potential limitation was the use of the BSIT, itself, as our primary
465 measure of olfactory function. While it is easy to use for the participant and suitable for
466 the remote nature of the study, we acknowledge that this tool is not as sensitive as other
467 more extensive tests (e.g. the University of Pennsylvania Smell Identification Test).
468 Furthermore, a major component of the study was the electronic survey, which was
469 completed by participants at baseline, 12 weeks and at one year. Findings from these
470 data may be subject to recall and response bias although this is likely limited as
471 participants were mainly asked to report their condition at the time of the survey. Lastly,
472 while the majority of participants were invited to complete the 1-year follow-up (169
473 of 218) a smaller-than-expected proportion responded. The remaining 50 participants
474 were recruited at a stage that was too late to be included within the timeframe of the 1-
475 year follow-up analysis. Hence, a response bias cannot be excluded. However, when
476 comparing the demographic details between responders and non-responders and

477 baseline BSIT scores, we did not observe a substantial difference (Supplemental Table
478 8). Therefore, response bias is likely minimal.

479 In summary, early olfactory training may be helpful, although the findings of
480 this trial are inconclusive. For those who responded to the 1-year follow-up, we
481 observed that those with persistent smell loss beyond 4 weeks are unlikely to recover
482 at 1 year with a high proportion of these participants also experiencing long-term
483 parosmia in addition to other symptoms of long Covid-19. As such, both anosmia and
484 parosmia should be considered important entities of long-Covid and further studies to
485 improve on their long-term management are highly warranted.

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487

488 ACKNOWLEDGEMENTS

489 Sisters Gill and Jackie at Barts Health who supported the recruitment at Barts Health
490 NHS Trust. We are also grateful for the support of the UCL/UCLH Joint Research
491 Office and NHS Health Research Authority, including Helen Penistone, Joe Wakeham,
492 Corcyra Burley, Susanne Emerton, Eyoanwan Simon-Modebe, Nowin Zahedi Fard,
493 and Pushpsen Joshi; the Research and Development teams at the local sites, including
494 Jack Biddle, Christina Armoogum, and Diane Heaton; Professor Chrissy Roberts and
495 the Open Data Toolkit team at the London School of Hygiene and Tropical Medicine
496 for facilitating electronic survey collection; Professor David Collier, Dr. Alastair Noyce
497 and Professor Sir Mark Caulfield from Barts Health NHS Trust for their support. We
498 would also like to thank Torsten Burghart and the team from Burghart Messtechnik for
499 kindly providing us with the olfactory training kits (Duft-Quartett) needed for the trial.

500

501 AUTHORSHIP CONTRIBUTION

502 ML, VJL (Co-CIs) and SJ, CP, CH, NK (local PIs) planned the study and led on the
503 study. All the above authors and CY, SP, NEL, TT, JM and JL and NC were also
504 involved in the planning/conduction of the study and/or analysis and interpretation of
505 the data. ML, JL, DC, RG, JR, CW, AT, JW, RB supported the acquisition of the data.
506 ML, JL, and VJL wrote up the manuscript draft with the help and input of all authors.

507

508 CONFLICT OF INTEREST

509 CP is a trustee of the charity Fifth Sense. All other authors declare no relevant conflict
510 of interest.

511

512 FUNDING

513 The study was funded by the Rhinology and Laryngology Research Fund, GSST charity
514 and WWL charity. TAT is funded by a BHF Intermediate Research Fellowship
515 (FS/19/35/34374). JCM, TAT and ML are directly and indirectly supported by the
516 UCLH and Barts NIHR Biomedical Research Units.

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TABLES

Table 1. Demographic characteristics of overall cohort by baseline BSIT result.

		Normal Smell (BSIT > 8) n=148		Abnormal Smell (BSIT 8 or less) n=70		Total n=218		p- value
		n	%	n	%	n	%	
Age (median, range)		44.0 (22 – 68) N=128		42.0 (23 – 78) N=61		44.0 (22 – 78) N=190		
Gender	Female	108	85.0	55	88.7	163	86.2	0.654
	Male	19	15.0	7	11.3	26	13.8	
	<i>Missing*</i>	21	NA	8	NA	29	NA	
Education	GSEs or eq.	15	11.7	3	4.8	18	9.5	0.391
	A-Levels or eq.	6	4.7	4	6.5	10	5.3	
	Degree	36	28.1	25	40.3	61	32.1	
	Higher Ed.	12	9.4	6	9.7	18	9.5	
	Post-Grad	51	39.8	22	35.5	73	38.4	
	Vocational	8	6.3	2	3.2	10	5.3	
	<i>Missing*</i>	20	NA	8	NA	28	NA	
Ethnicity	White	111	86.7	54	87.1	165	86.8	0.193
	Mixed	3	2.3	2	3.2	5	2.6	

	Indian	9	7.0	1	1.6	10	5.3	
	Pakistani	0	0.0	0	0.0	0	0.0	
	Bangladeshi	0	0.0	1	1.6	1	0.5	
	Chinese	0	0.0	0	0.0	0	0.0	
	Black	3	2.3	4	6.5	7	3.7	
	Other	2	1.6	0	0.0	2	1.1	
	<i>Missing*</i>	20	NA	8	NA	28	NA	
Smoking History	Never	93	72.7	44	71.0	137	72.1	0.843
	Former	27	21.1	15	24.2	42	22.1	
	Current	8	6.3	3	4.8	11	5.8	
	<i>Missing*</i>	20	NA	8	NA	28	NA	
Alcohol History	Never	24	18.8	11	17.7	35	18.4	0.986
	1-14 units/week	93	72.7	46	74.2	139	73.2	
	15-21 units/week	8	6.3	4	6.5	12	6.3	
	Over 21 units/week	3	2.3	1	1.6	4	2.1	
	<i>Missing*</i>	20	NA	8	NA	28	NA	

653 *baseline questionnaires were not available from 28 participants (either incomplete or not returned); as
654 such, only information regarding objective smell testing were available for these.
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661 **Table 2.** Participant medical history and COVID-19 symptomology and associations with baseline
662 BSIT result for overall cohort.
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	Normal Smell (BSIT > 8) n=148		Abnormal Smell (BSIT 8 or less) n=70		Total n=218		p-value
	n	%	n	%	n	%	
Medicinal History							
Sinonasal Disease	29	22.7	18	29.0	47	24.7	0.372
Diabetes	1	0.8	1	1.6	2	1.1	0.547
COPD	0	0.0	0	0.0	0	0.0	NA
Asthma	16	12.5	8	12.9	24	12.6	1.000
Bronchitis	1	0.8	1	1.6	2	1.1	0.547
Other Chronic Lung Disease	0	0.0	0	0.0	0	0.0	NA
Cancer	2	1.6	2	3.2	4	2.1	0.598
Stroke	0	0.0	0	0.0	0	0.0	NA
Heart Disease	0	0.0	0	0.0	0	0.0	NA
Arthritis	6	4.7	3	4.9	9	3.8	1.000
SLE	1	0.8	0	0.0	1	0.5	1.000
Other Autoimmune disease	4	3.1	2	3.2	6	3.2	1.000

High Blood Pressure	12	9.4	5	8.1	17	8.9	1.000
If high BP, treatment with ACEi/ARBs (n=17)	6	50.0	4	80.0	10	58.8	0.338
Any	53	41.4	25	40.3	78	41.1	1.000
<i>Missing*</i>	20	NA	8	NA	28	NA	
COVID-19 Symptoms							
Persistent Cough	38	29.7	24	38.7	62	32.6	0.249
Shortness of Breath	39	30.5	31	50.0	70	36.8	0.011
Sore Throat	33	25.8	24	38.7	57	30.0	0.091
Loss of Smell	128	100.0	62	100.0	190	100.0	NA
Loss of Taste	93	72.7	51	82.3	144	75.8	0.206
Hoarse Voice	7	5.5	9	14.5	16	8.4	0.050
Fever	4	35.2	29	46.8	74	38.9	0.153
Fatigue	86	67.2	48	77.4	134	70.5	0.176
Difficulty Breathing	16	12.5	16	25.8	32	16.8	0.037
Nasal Congestion	32	25.0	19	30.6	51	26.8	0.485
Burning in Nose/Mouth	17	13.3	9	14.5	26	13.7	0.824
Aches/Pains	60	46.9	41	66.1	101	53.2	0.015
Diarrhoea	29	22.7	10	16.1	39	20.5	0.342
Delirium	2	1.6	2	3.2	4	2.1	0.598
Chest Pain	13	10.2	16	25.8	29	15.3	0.009
Abdominal Pain	12	9.4	8	12.9	20	10.5	0.459
Metallic Taste	16	12.5	11	17.7	27	14.2	0.377
Skipped Meals	34	26.8	15	24.2	49	25.9	0.860
<i>Missing</i>	20	NA	8	NA	28	NA	

*baseline questionnaires were not available from 28 participants (either incomplete or not returned); as such, only information regarding objective smell testing were available for these.

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Table 3a. Summary of BSIT scores at baseline, 12-weeks and 1-year.

		All	RCT	
			Treatment	Control
Baseline score		N=218	N=33	N=30
	Normal, n(%)	148 (67.9)	0 (0.0)	0 (0.0)
	Mild, n(%)	53 (24.4)	25 (75.8)	22 (73.3)
	Moderate, n(%)	17 (7.8)	8 (24.2)	8 (26.7)
	Severe, n(%)	0 (0.0)	0 (0.0)	0 (0.0)
	mean (std. dev.)	9.1 (2.12)	6.5 (1.70)	6.7 (1.51)
12-week score		N=51	N=26	N=25
	Normal, n(%)	21 (41.2)	13 (50.0)	8 (32.0)
	Mild, n(%)	25 (49.0)	11 (42.3)	14 (56.0)
	Moderate, n(%)	4 (7.8)	1 (3.8)	3 (12.0)
	Severe, n(%)	1 (2.0)	1 (3.8)	0 (0.0)
	mean (std. dev.)	7.9 (2.23)	8.0 (2.52)	7.8 (1.92)
<i>Change from baseline</i>	mean (std. dev.)	1.3 (2.07)	1.5 (2.49)	1.0 (1.53)
1-year score		N=56	N=11	N=8
	Normal, n(%)	42 (75.0)	6 (54.5)	3 (37.5)

	Mild, n(%)	13 (23.2)	5 (45.5)	4 (50.0)
	Moderate, n(%)	1 (1.8)	0 (0.0)	1 (12.5)
	Severe, n(%)	0 (0.0)	0 (0.0)	0 (0.0)
	mean (std. dev.)	9.5 (1.71)	8.6 (1.29)	8.0 (2.33)
<i>Change from baseline</i>	mean (std. dev.)	0.2 (1.77)	1.6 (1.97)	0.9 (1.81)

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Table 3b. Primary and secondary outcomes for early olfactory training at 12-weeks and at 1-year.

	Treatment vs. Control	
12-weeks (n=51)	Difference in BSIT change between arms	0.45 (95% CI: -0.69 to 1.59) $p = 0.43$
	Standardized effect size	0.22 (95% CI: -0.34 to 0.77)
	Odds of having normal smell	OR=2.38 (95% CI: 0.73 to 7.76) $p = 0.15$
1-Year	Difference in BSIT change between arms	0.65 (95% CI: -1.01 to 2.31) $p=0.42$
	Standardized effect size	0.31 (95% CI: -0.38 to 1.01) $p = 0.36$
	Odds of having normal smell	OR=2.33 (95% CI: 0.37 to 14.61) $p=0.37$

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Table 4. Prevalence of parosmia and phantosmia at baseline and at 1-year.

	Normal	Abnormal	Total
Parosmia at baseline	<i>(n=117)</i>	<i>(n=61)</i>	<i>(n=178)</i>
Present	43 (36.8%)	30 (49.2%)	73 (41.0%)
Absent	74 (63.2%)	31 (50.8%)	105 (59.0%)
Parosmia at 1-year	<i>(n=51)</i>	<i>(n=25)</i>	<i>(n=76)</i>
Present	17 (33.3%)	16 (64.0%)	33 (43.4%)
Absent	34 (66.7%)	9 (36.0%)	43 (56.6%)
Parosmia for paired samples (1-year/Baseline)	<i>(n=48)</i>	<i>(n=25)</i>	<i>(n=73)</i>
Present / Present	13 (27.1%)	11 (44.0%)	24 (32.9%)
Present / Absent	4 (8.3%)	5 (20.0%)	9 (12.3%)
Absent / Present	4 (8.3%)	1 (4.0%)	5 (6.8%)
Absent / Absent	27 (56.3%)	8 (32.0%)	35 (47.9%)
	Normal	Abnormal	Total
Phantosmia at baseline	<i>(n=117)</i>	<i>(n=61)</i>	<i>(n=178)</i>
Present	25 (21.4%)	20 (32.8%)	45 (25.3%)
Absent	92 (78.6%)	51 (67.2%)	133 (74.7%)
Phantosmia at 1-year	<i>(n=51)</i>	<i>(n=25)</i>	<i>(n=76)</i>

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Present	5 (9.8%)	9 (36.0%)	14 (18.4%)
Absent	46 (90.2%)	16 (64.0%)	62 (81.6%)
Phantomia for paired samples (1-year/Baseline)	<i>(n=48)</i>	<i>(n=25)</i>	<i>(n=73)</i>
Present / Present	2 (4.2%)	7 (28.0%)	9 (12.3%)
Present / Absent	2 (4.2%)	2 (8.0%)	4 (5.5%)
Absent / Present	6 (12.5%)	3 (12.0%)	9 (12.3%)
Absent / Absent	38 (79.2%)	13 (52.0%)	51 (69.9%)

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Table 5. Mean QoL scores at baseline and at 1-year.

		Baseline, All Cases		Paired Analysis		
		Baseline		Baseline		1-Year
		N	Mean (Std. Dev.)	N	Mean (Std. Dev.)	Mean (Std. Dev.)
1	Has the loss of smell affected you socially? (i.e. in your work and personal life)	84	4.3 (1.58)	22	4.9 (1.13)	3.8 (1.47)
1a	The changes in my sense of smell make me feel isolated.	88	3.1 (1.77)	24	3.8 (1.62)	3.0 (1.49)
1b	Because of the changes in my sense of smell, I have problems with taking part in activities of daily life.	88	2.6 (1.68)	24	3.2 (1.89)	2.7 (1.49)
1c	The changes in my sense of smell make me feel angry.	88	4.0 (1.88)	24	4.8 (1.69)	4.4 (1.53)
2	Has the loss of smell affected your eating habits?	66	4.4 (1.67)	16	4.8 (1.33)	4.7 (1.25)
2a	Because of the changes in my sense of smell, I cook less often than I used to (or visit restaurants less often than I used to).	88	3.7 (2.07)	24	4.4 (1.98)	4.0 (1.94)
2b	Because of the changes in my smell, I don't enjoy food or drinks as much as I used to.	88	5.4 (1.86)	24	6.2 (0.88)	4.8 (1.77)
2c	Because of the changes in my sense of smell, I eat less than I used to or more than I used to.	87	3.8 (2.00)	24	4.1 (1.83)	3.2 (1.66)
3	Has the loss of smell affected your anxiety levels?	64	3.2 (1.67)	15	3.5 (1.46)	3.5 (1.85)
3a	Because of the changes in my sense of smell, I feel more anxious than I used to feel.	87	3.3 (1.78)	24	3.9 (1.82)	3.5 (1.72)
3b	Because of the changes in my sense of smell, I feel more socially isolated.	88	2.7 (1.67)	24	3.5 (1.64)	3.0 (1.57)
3c	Because of the changes in my sense of smell, I have to try harder to relax.	88	2.7 (1.72)	24	3.7 (1.76)	3.0 (1.52)
4	To what degree is the loss of smell annoying to you?	59	5.7 (1.72)	15	6.3 (0.72)	5.3 (1.23)

4a	I am worried that I will never get used to the changes in my sense of smell.	87	5.3 (1.96)	24	6.3 (1.00)	5.4 (1.17)
4b	The changes in my sense of smell annoy me when I am eating.	87	5.3 (2.06)	24	6.1 (0.90)	5.3 (1.33)

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790

791 FIGURE LEGEND

792

793 Figure 1. Flow-chart

794

795 TABLE LEGEND

796

797 **Table 1.** Demographic characteristics of overall cohort by baseline BSIT result.

798

799 **Table 2.** Participant medical history and COVID-19 symptomology and associations with baseline
800 BSIT result for overall cohort.

801

802 **Table 3a.** Summary of BSIT scores at baseline, 12-weeks and 1-year.

803

804 **Table 3b.** Primary and secondary outcomes for early olfactory training at 12-weeks and at 1-year.

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806 **Table 4.** Prevalence of parosmia and phantosmia at baseline and at 1-year.

807

808 **Table 5.** Mean QoL scores at baseline and at 1-year.

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