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Virtual reality intervention alleviates dyspnea in patients recovering from COVID pneumonia

Betka Sophie*1, Kannape Oliver Alan*1,2,3, Fasola Jemina1, Lance Florian1, Cardin Sylvain2, Schmit Aline4, Similowski Thomas5,6, Soccal Paola Marina4, Herbelin Bruno**1, Adler Dan**4, Blanke Olaf**1

* Contributed equally to the publication
** Contributed equally to the publication

1Laboratory of Cognitive Neuroscience, Brain Mind Institute and Center for Neuroprosthetics, Faculty of Life Sciences, Ecole Polytechnique Federale de Lausanne, (EPFL), Geneva, 1202, Switzerland

2Mindmaze SA, Lausanne, Switzerland

3Virtual Medicine Center, University Hospital Geneva, Switzerland

4Division of Lung Diseases, University Hospital and Geneva Medical School, University of Geneva, Switzerland

5Sorbonne Université, INSERM, UMRS1158 Neurophysiologie Respiratoire Expérimentale et Clinique, F-75005 Paris, France

6AP-HP, Groupe Hospitalier Universitaire APHP-Sorbonne Université, site Pitié-Salpêtrière, Département R3S (Respiration, Réanimation, Réhabilitation respiratoire, Sommeil), F-75013 Paris, France

Corresponding author: Prof. Olaf Blanke; olaf.blanke@epfl.ch; +41 21 693 96 21

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Abstract (243w)

**Background**
Immersive virtual reality (iVR)-based digital therapeutics (DTx) are gaining clinical attention in the field of pain management. Based on known analogies between pain and dyspnea, we investigated the effects of visual-respiratory feedback, on persistent dyspnea in patients recovering from COVID-19 pneumonia.

**Methods**
We performed a controlled, randomized, single-blind, cross-over proof-of-concept study (feasibility and initial clinical-efficacy) to evaluate an iVR-based intervention to alleviate dyspnea in patients recovering from COVID-19 pneumonia. Included patients reported persistent dyspnea (≥5 on a 10-point scale) and preserved cognitive function (MoCA>24). Assignment was random and concealed. Patients received synchronous (intervention) or asynchronous (control) feedback of their breathing, embodied via a gender-matched virtual body. The virtual body flashed in a waxing and waning visual effect which could be synchronous or asynchronous to the patient’s respiratory movements. Outcomes were assessed using questionnaires and breathing recordings.

**Results**
Study enrollment was open between November 2020 and April 2021. Twenty-six patients were enrolled (27% women; age: median=55, interquartile range (IQR)=18). Data were available for 24 of 26 patients. The median (IQR) rating on a 7-point Likert-scale of breathing comfort improved from 1(2) at baseline, to 2(1) for synchronous feedback, but remained unchanged at 1(1.5) for asynchronous feedback (p<0.05) between iVR conditions). Moreover, 91.2% of all patients were satisfied with the intervention (p<0.0001) and 66.7% perceived it as beneficial for their breathing (p<0.05).

**Conclusion**
Our iVR-based DTx presents a feasible and safe respiratory rehabilitation tool that improves breathing comfort in patients recovering from COVID-19 infection presenting with persistent dyspnea. Future research should investigate the intervention’s generalizability to persistent dyspnea with other etiologies and its potential for preventing chronification.
Authors’ contributions

Every author named at the start of the article contributed to the study: Conceptualization (AD, BO, BS, HB, KO, ST), Data curation (BS, FJ), Formal Analysis (FJ, BS), Funding acquisition (AD, BO, BS), Investigation (FJ, SA), Methodology (AD, BO, BS, FJ, HB, KO), Project administration (AD, FJ, SA), Resources, Software (CS, HB, LF), Supervision (AD, BO, BS, KO), Visualization (BS, FJ), Writing – original draft (BS, FJ), Writing – review & editing (AD, BO, BS, CS, HB, KO, ST). All authors confirm that they had full access to all the data in the study and accept responsibility to submit for publication. FJ and BS verified the data.

Role of the funding source

The funders of the study had no role in study design, data collection, data analysis, data interpretation, or writing of the report. MindMaze SA involvement was limited to providing the devices for the study and in-kind contributions for the software development.
Introduction

Dyspnea is defined as "a subjective experience of breathing discomfort made of various sensations that can vary in intensity"\(^1\). In simpler words, dyspnea relates to the upsetting or distressing awareness of breathing activity. Beyond the symptom of cardiorespiratory dysfunction, dyspnea is a frightening and disabling experience. This is particularly true when it resists optimized treatment of the underlying condition, a situation termed "chronic breathlessness"\(^2\) or, more broadly, "persistent dyspnea"\(^3\). Persistent dyspnea deeply affects the lives of those afflicted. It profoundly deteriorates quality of life by impacting cognitive function, locomotion, and mental health\(^4\). Implicit to the definition of persistent dyspnea is the under-recognition of respiratory suffering by caregivers (e.g., during clinical consultations)\(^5\), as a major clinical burden. This invisibility, an important difference between dyspnea and pain impairs access to care \(^5\) and hinders the development of evidence-based targeted interventions\(^6\). The aim of the current study was therefore to develop and evaluate a non-invasive, non-pharmacological intervention that would use immersive Virtual Reality (iVR) to alleviate breathlessness in a clinical population, drawing on promising approaches that have also used embodied iVR for chronic pain.

Neuroscience evidence suggests that dyspnea occurs in conjunction with the recruitment of a neural network involving the insula, dorsal anterior cingulate cortex, amygdala, and medial thalamus, sharing important pathways with other brain functions such as pain processing\(^7\) and bodily self-consciousness\(^9\). This suggests that it is relevant to target the brain to relieve dyspnea when all cardiorespiratory approaches have been exhausted\(^13\).

In this regard, prior interventions using immersive VR-based Digital Therapeutics, also referred to as digiceuticals\(^11\), have demonstrated alleviation of chronic pain in patients with complex regional pain syndrome or spinal cord injury\(^12,13\). In the respiratory domain, visuo-respiratory stimulation has been associated with an increased feeling of breathing control (breathing agency)\(^14\), a reduced negative emotional state related to experimental dyspnea\(^15\), as well as changes in physiological measures of breathing\(^8,16\).
Persistent symptoms can occur beyond the initial period of COVID-19 infection recovery and affect patients who were managed in the community or in the acute care setting\textsuperscript{17}. Like general weakness, malaise, fatigue, and impaired concentration, dyspnea has consistently been reported in so-called long COVID cohorts with a high prevalence of around 25\% (CI95 18\% to 34\%)\textsuperscript{18}. In the case of persistent dyspnea, an extensive workup to identify respiratory sequelae or muscle deconditioning should be the foremost clinical preoccupation, mostly to guide the indications of pulmonary rehabilitation\textsuperscript{19}. Yet, dyspnea can be dissociated from physiological markers such as pulmonary function tests or lung imaging, in post-COVID situations\textsuperscript{18} as in a more general manner\textsuperscript{20}. This makes treatment and even diagnosis challenging. The importance of brain mechanisms in the pathogenesis of dyspnea justifies neuroscientific approaches for its management and implies that a cognitive intervention using a neuro-rehabilitation approach could be tested to understand and alleviate this debilitating symptom.

The present clinical study was performed to evaluate two primary outcomes: 1) the initial clinical efficacy of the COVID Virtual Reality (COVVR) intervention and 2) the overall feasibility of using iVR in patients recovering from COVID-19 pneumonia. This follows the guidelines for clinical trials using VR and corresponds to a VR phase 2 trial\textsuperscript{21}. The primary hypothesis with respect to efficacy was that our iVR intervention would alleviate dyspnea by improving breathing comfort in patients recovering from COVID-19 pneumonia presenting with persistent dyspnea. With respect to the feasibility of COVVR we hypothesized that the intervention could be used in an inpatient setting and would be accepted by the patients. We also had two research questions regarding potential perceptual changes in patients affected by persistent dyspnea. One, do patients maintain accurate awareness of their breathing movements, and two, do they maintain their sense of agency, or control, over their breathing\textsuperscript{17}?

Methods

\textit{Study Design}

A prospective controlled, randomized, single-blind, cross-over clinical study was conducted to evaluate both the efficacy and feasibility of a iVR biofeedback intervention to alleviate persistent dyspnea in patients recovering from COVID-19
pneumonia. This single-site study was carried out at the University Hospital (HUG) in Geneva, Switzerland and was approved by the Commission Cantonale d’Ethique de la Recherche de la République et Canton de Genève (2019-02360).

Patients
Thirty-nine patients were screened by a respiratory physician (AS). Patients that scored below 25 on the Montreal Cognitive Assessment (MoCA) were excluded (N=5); N=8 declined to participate. In total, N=26 patients were enrolled. Clinical inclusion criteria were that patients i) were recovering from COVID-19 pneumonia confirmed by Reverse Transcription Polymerase Chain Reaction (RT-PCR) for SARS-CoV-2, and ii) presented with persistent dyspnea with a self-rated intensity of five or higher (out of ten) on a visual analog dyspnea scale, at rest. The respiratory physician asked the dyspnea question as follows: "Do you have difficulty breathing?" then "On a scale of 0 to 10, with 0 being no difficulty to breathe and 10 being the worst difficulty to breathe that you can imagine, where do you rank?". This dyspnea rating was only used as an inclusion criterion and not as an outcome. The delay between the initial screening by the physician and the inclusion by the researcher varied between 1 hour to 2 days. As discussed below, a separate set of questions was used to evaluate the primary outcomes, as they could directly be compared to the prior studies on respiration-awareness. Patients had to be able to give consent and to understand and speak French or English. Patients who presented with unstable respiratory, neurological, or cardiac conditions, or psychiatric illness were excluded (see supplementary material for details on Screening). Patients underwent randomization in the respiratory ward during the recovering phase of COVID pneumonia only if they were in a stable respiratory, neurological or cardiac clinical condition.

Procedure

Figure 1

Setup
Eligible patients were installed in a semi-seated position in their hospital bed and wore a belt-mounted linear force sensor (Go Direct® Respiration belt, Vernier, Beaverton (OR), USA) fitted on the abdomen to allow proper recording of respiratory movements. They were also equipped with a head-mounted display (Zeiss VR
ONEPLUS, Oberkochen, Germany) holding a smartphone (Samsung Galaxy S8, Seoul, South Korea). The smartphone ran the VR application and connected via Bluetooth® to the respiration belt. MindMaze SA provided the hardware for the study, co-developed the application with the Laboratory of Cognitive Neuroscience at EPFL and deployed this on the smartphone. The application collects and processes respiratory data to render a computer-generated virtual environment in real-time.

Figure 2

**Intervention Conditions**

Patients were asked to look around in the VR environment and orient their gaze to a gender-matching virtual body lying on a bed next to them in a similar position as theirs (Figure 2.B). The virtual body flashed in a waxing and waning visual effect which could be synchronous or asynchronous to the patient’s respiratory movements. In the synchronous condition, the radiance of the visual flash was maximal at the end of inspiration and minimal at the end of the expiration. In the asynchronous condition, at the end of each visual flash, a duration between 2.5 and 33.3 seconds is randomly generated for the next visual stimulation, such that the feedback is both phase-shifted and frequency-modulated with respect to the actual respiration.

**Intervention Procedure**

Once the patient was ready, they were asked to close their eyes while their respiratory movements were recorded for two minutes. Participants were then asked to describe their current respiratory experience by answering two questions according to a classically used 7-point Likert-scale (from -3= Strongly disagree to +3= Strongly agree): 1) *I have difficulty breathing* (breathing difficulty) and 2) *My breathing is enjoyable* (breathing comfort). Questions were asked to patients in French language. This condition served as the baseline assessment for the breathing (dis)comfort items and the breathing rate. For comparison the two items were also included in the post-exposure questionnaires.

Table 1

After baseline assessment, participants were randomly assigned in a 1:1 ratio to receive either the sequence “synchronous/asynchronous” or “asynchronous/synchronous” (see supplementary material for details on
Randomization and masking. In each block, patients were first asked to look around in the virtual room, and then to orient their gaze towards the virtual body while relaxing for 5 minutes. They were not informed that the flashing of the virtual body was related to their breathing. Each block was followed by a subjective questionnaire of six items (7-point Likert scale) derived from previous visuo-respiratory studies\textsuperscript{14–16}, (see Table 1). Question 1 evaluated the awareness of the visuo-respiratory experimental manipulation (breathing awareness)\textsuperscript{14,16}, while question 3 pertained to the breathing agency\textsuperscript{14–16}. Questions 2 and 5 were included as control items. These should not differ between conditions but could flag if participants were influenced by baseline suggestibility. The breathing comfort items matched the baseline assessment (Q4 and Q6 respectively, Table 1). Finally, patients completed an \textit{ad hoc} questionnaire to assess the acceptance and feasibility of the iVR intervention (7-point Likert scale, see supplementary section Table S1). During the entire intervention (baseline included), oxygen therapy was administered through nasal cannulas to obtain a SpO2 level of 90-92%.

**Outcomes**

Following the recommendations for clinical trials in VR we conducted a VR2-type study with a focus on initial clinical efficacy and intervention feasibility as two primary outcomes\textsuperscript{21}. Efficacy was evaluated based on subjective feedback by the patients regarding their breathing comfort and difficulty (Q4 and Q6 of the above Table 1). Feasibility was evaluated using a feedback questionnaire. Agreement with the questionnaire items indicates better feasibility, acceptance, and perceived outcome.

Secondary outcome measures included respiratory parameters as well as the subjective reports of breathing awareness and agency. Both respiratory rate (breaths per minute) and respiratory rate variability (using inter-breath intervals) were measured using the respiration belt. Respiratory rate and variability were compared across the baseline and two intervention conditions. Breathing awareness and agency were evaluated using a 7-point Likert scale where agreement indicates stronger embodiment of the feedback.

The feedback questionnaires were administered by the researcher, directly after the intervention.
**Statistical Analysis**

All analyses were performed using R (version 4.1.0) and Matlab (version 2020a). The effect of synchrony on each measure was assessed using a linear mixed-effects model, with a random intercept for each patient. In addition to the experimental condition (synchronous vs asynchronous condition), each model also included the experimental sequence (starting the experiment with synchronous or asynchronous condition) and the interaction between the experimental sequence and the experimental condition as fixed effects. The statistical significance of the interactions was assessed using the likelihood ratio test. All p-values were two-sided and statistical significance was set at a p-value of 0.05.

Median, interquartile range (IQR), and rating frequency (in%) were computed for each feasibility item. To ensure clarity, observed percentages for ratings from 1 = Agree to 3 = Strongly agree were grouped, indicating overall agreement with the statement. A one-sided, one-sample t-test was used to determine if the mean of ratings was significantly greater than zero, indicating that, at least, the majority of patients were agreeing with the statement. For details regarding *Power calculation & Risk assessment* see supplementary material.
Results

Demographic and baseline information

Patient enrollment, randomization, and testing took place at the division of Pneumology at Geneva University Hospital between November 2020 and April 2021. Twenty-six patients were randomly assigned either to the “asynchronous/synchronous” sequence (N=12) or the “synchronous/asynchronous” sequence (N=14). At the time of database lock in May 2021, data was available for all except two (7.7 %) of 26 patients (Table 1).

Table 2

Subjective and physiological measures are reported in Table 2. At baseline, the median (and IQR) breathing comfort rating was 1(2) and the mean breathing difficulty rating was 1(3). Median values and interquartile ranges for each experimental condition, in function of the experimental sequence are provided in supplementary Table S2.

Primary outcomes

Efficacy of the intervention on breathing comfort and difficulty.

Table 3

Regarding the primary outcome breathing comfort, we observed that the median (and IQR) rating significantly improved from 1 (1.5) during the asynchronous condition to 2(1) during the synchronous condition, with an estimated difference between conditions of 0.54 (95% CI 0.05-1.04, p<0.05, Figure 3.A). Moreover, post-hoc paired one-sided t-tests, confirmed a significant difference between breathing comfort ratings during the intervention (synchronous) condition compared to baseline. No such difference was observed for the control (asynchronous) condition, excluding a mere effect of VR distraction (see supplementary section for statistical details). For the assessment of breathing difficulty, even though a similar trend was observed in the data, no significant main effect of experimental condition was observed (Figure 3.D). The experimental sequence had no significant effect on breathing comfort or breathing difficulty ratings.
Feasibility of the intervention

Feasibility ratings, the co-primary outcome, are depicted in Figure 4. The majority of the patients (91.2%) were satisfied with the intervention (Satisfaction: median (IQR)=2(2); t=5.20, p<0.0001, 95% CI 1.17 to inf). In addition, 66.7% rated the iVR intervention as beneficial for their breathing (Respiratory benefit: 1(2.25), t=1.81, p<0.05, 95% CI 0.04 to inf). Half of the patients reported that it made them feel better (Well-being benefit: 0.5(4), t=0.36, p>0.05, 95% CI -0.64 to inf), and a further 45.8% indicated that they would like to continue using the device during their recovery (Rehabilitation: 0(4), t=0.10, p>0.05, 95% CI -0.67 to inf) and at home (Home use: -1(4.25), t=-0.74, p>0.05, 95% CI -1.11 to inf). Finally, 37.5% would have liked to use the intervention earlier during their stay at the hospital (Hospital use: 0(2.25), t=-0.22, p>0.05, 95% CI -0.74 to inf). Descriptive statistics and statistical tests are described in the supplementary section (Table S3 and Figure S2).

Figure 3

Secondary outcomes

Effect of the intervention on subjective reports of breathing awareness and agency

The secondary outcome measures of this study included the subjective ratings for breathing awareness and agency and the physiological measures. The median (and IQR) breathing agency rating increased from -2(4) during the asynchronous condition to 1.5(4.25) during the synchronous condition, with an estimated difference of 1.58 (95% CI 0.34 to 2.83, p<0.05, Figure 3.B). The median breathing awareness rating increased from -1.5(4, asynchronous) to 2(2, synchronous), with an estimated difference of 2.17 (95% CI 1.07 to 3.27, p<0.0001, Figure 3.C). Neither control item differed between conditions (“It seemed as if I had three bodies”, Figure 3.E, “I felt as if the virtual body was drifting with the flashing”, Figure 3.F).

Effect of the intervention on respiratory parameters

Respiration rate and its variability did not differ between experimental conditions. The order of conditions did not significantly affect any of the secondary outcomes (see Figure S1).

Figure 4
Discussion

In this study, our COVID VR intervention (COVVR), improved breathing comfort in patients with persistent dyspnea recovering from COVID-19 pneumonia. Persistent dyspnea, a common but underreported condition, is defined as the breathlessness reported by patients despite receiving state-of-the-art treatment of their respiratory condition, and leads to major disabilities impacting cognition, locomotion, and mental health. COVVR may therefore provide an additional non-invasive and non-pharmacological tool for aiding patient recovery and satisfaction, with the potential of alleviating some of the burden of this debilitating symptom.

Patients reported a significant improvement in breathing comfort after a relatively short exposure (5 minutes) to synchronous visuo-respiratory COVVR stimulation compared to the asynchronous control condition and compared to their baseline breathing comfort. Our results extend recent observations in chronic pain studies to patients with persistent dyspnea. This previous iVR work indicated the value of personalized stimulation using cardio-visual, somatosensory-visual, and respiratory-visual feedback (see 14–16). These studies using multisensory bodily stimulations, including the present VR protocol, differ from previous interventions focused on using (1) immersive or non-immersive VR as a distraction tool or (2) the more recent efforts to digitizing patient education and cognitive behavioral therapy; they differ by being designed to impact the central processing of nociceptive and respiratory signals respectively. The specificity of these personalized iVR interventions, including COVVR, is highlighted by the cross-over randomized design of our study differing only in respiratory synchrony (synchronous versus asynchronous conditions), while being identical in all other aspects of VR exposure (i.e., presence of a virtual body animated by patient’s own breathing, total duration of breathing sequence, identical 3D virtual environment, etc.). This is markedly different from the more commonly applied, non-immersive VR-stimulations prevalent in medical research.

Next to the positive primary outcome, a similar beneficial effect of the intervention was observed for breathing agency, that is, the feeling of being in control of one’s breathing. Patients reported a stronger sense of control over their breathing for synchronous feedback as well as maintained awareness of their breathing movements. Patients further reported a global satisfaction regarding the VR
intervention and, more importantly, indicated that the iVR feedback improved their breathing. The COVVR study extends respiratory iVR studies in healthy individuals that have demonstrated increased breathing agency\textsuperscript{14-16}, changes in tidal volume variability\textsuperscript{16}, and translates this approach to the bedside. Monitoring these markers as well as the patient’s emotional state\textsuperscript{18}, may be instrumental to decreasing dyspnea-related anxiety and understanding its chronification.

While synchronous visuo-respiratory stimulation improved one item used for the assessment of dyspnea (Q6: my breathing was more enjoyable), synchronous stimulation was only associated with a non-significant improvement in the other item used for this assessment (Q4: I had difficulty breathing). The most probable reason for this finding is a lack of power related to our sample size. Another reason is that, although participants reported persistent dyspnea with a self-rated intensity of five or higher when screened by the respiratory physician, their agreement with item Q4 (I had difficulty breathing) was quite low just prior to the VR intervention, indicating a possible white coat effect. The semantics of the chosen items could also explain this finding. The item “My breathing is enjoyable” taps into affective processes whereas the item “I have difficulty breathing” explores sensory/perceptual processes.\textsuperscript{27} Further work should compare both breathing comfort and breathing difficulty items to validated respiratory questionnaires.

Mounting evidence using functional neuroimaging suggests that patients suffering from persistent dyspnea may exhibit “hypersensitivity” to afferent respiratory signals as a result of learned expectations\textsuperscript{28}. Perception and anticipatory processes of dyspnea are known to share breathing control mechanisms in the brainstem and the insular cortex.\textsuperscript{26} Consequently, once treatment of the underlying respiratory pathophysiology has been optimized, these neurorespiratory mechanisms should be considered as potential targets for pharmacological and non-pharmacological interventions\textsuperscript{10}. Pharmacological treatments have been shown to be useful: low dose oral sustained-release morphine administered for persistent dyspnea is associated with improved health status in COPD without affecting PaCO\textsubscript{2} or causing serious side-effects (especially in patients with mMRC stage 3-4).\textsuperscript{25} Pulmonary rehabilitation, an evidence-based multidisciplinary non-pharmacological intervention has also been shown to modify neural responses to learned breathlessness associations, likely due to central desensitization to dyspnea.\textsuperscript{26} While not directly
investigating neurorespiratory mechanisms, the present iVR paradigm, using carefully controlled visuo-respiratory conflicts, not only introduces a new complementary rehabilitation intervention but may help identify subjective (agency, awareness, dis/comfort) and physiological (breathing rate and variability) markers of “hypersensitivity”, based on perceptual and anticipatory brain processes of dyspnea.

Digital therapeutics are becoming popular in the field of chronic pain management. Dyspnea and pain share several similarities. They engage similar brain networks, are best characterized by multidimensional models, and both respond to opioid treatment. As the global COVID-19 pandemic has progressed, a significant proportion of patients experience prolonged symptoms beyond the initial period of acute infection, such as persistent dyspnea. The increasing number of patients isolated for prolonged periods has stressed an urgent need to develop multidisciplinary rehabilitation strategies that can be individualized and adapted to accommodate patients’ needs. Given our findings, we propose that our iVR intervention is a feasible and safe neuro-rehabilitation tool that could be considered to improve breathing comfort in patients experiencing persistent dyspnea after COVID-19 infection. As our intervention involves neurorespiratory processes its use could further be extended to persistent dyspnea with other etiologies. Offering a non-pharmacological, non-invasive intervention that can readily be adapted for home-use may be particularly relevant at a time when over 40% of adults are estimated to avoid medical care because of COVID-19-related concerns.

Our study comes with certain limitations. First, although it is based on an adequate power calculation for a proof-of-concept study, our results stem from a small sample. Eight participants (20%) refused to participate, unfortunately reasons for refusal were not recorded. It is therefore possible that selection biases have contributed to our results. Second, almost half of the patients who were selected as being severely dyspneic (i.e., visual analogic dyspnea scale ≥ 5) reported low agreement with the breathing difficulty item, at baseline. This may be due to the delay between the initial screening and the start of the intervention or the fact that the former was completed by the respiratory physician and the latter by a researcher. Further studies should also include validated multi-dimensional dyspnea scales, such as the Multidimensional Dyspnea Profile, as a multidimensional outcome would be ideal.

While we here focused on a specific population of patients recovering from COVID-
In conclusion, our study shows that a short exposure to an iVR-based digital therapeutic can improve breathing comfort and breathing control in patients recovering from COVID-19 pneumonia. Global satisfaction and respiratory benefit from the patients are reported, attesting to the feasibility of the present intervention. Although more clinical data are needed, immersive VR-based interventions may become a key factor of the multi-dimensional treatment of persistent dyspnea.
References


Figure legends

Figure 1: Study flow chart

Figure 2: Portable setup and virtual reality feedback. (A) A respiratory belt captures the respiratory movements of the chest and sends the signal to a smartphone via Bluetooth. A custom software generates the virtual environment. (B) A matched-gender virtual body is displayed and observed by the patients by slightly turning their head to the side. The virtual body is illuminated synchronously or asynchronously with respect to the patient’s chest movements. The top image represents the end of the expiration with a low flashing intensity, while the bottom image shows the end of the inspiration corresponding to the maximal luminosity in the synchronous condition. A video of the experiment can be found in the supplementary material.

Figure 3: Breathing comfort (A), Agency (B), Awareness (C), Breathing difficulty (D) and control items (E&F) test results. (A-C) Subjective measures for which the main effect of the experimental manipulation was significant. *p<.05; ***p<.001 (D-F) Subjective measures for which the main effect of the experimental manipulation was not significant.

The boxplots are depicting subjects’ ratings during asynchronous condition compared to the synchronous condition, independent of experimental sequence. The thick line within a box plot represents the median, the diamond represents the mean, the upper boundary of the box indicates the 25th percentile (Q1) and lower boundary the 75th percentile (Q3). The whiskers above and below the box indicate the minimal and maximal values (Q1 – 1.5*IQR and Q3 + 1.5*IQR respectively), while points above the upper or below the whiskers indicate outliers. Subjective ratings were measured using a 7-point Likert scale with -3 = Strongly disagree, -2 = Disagree; -1 = Somewhat disagree; 0 = Neither agree nor disagree; 1 = Somewhat agree; 2 = Agree; 3= Strongly agree.

Figure 4: Feasibility scores for all items. The boxplots are depicting subjects’ ratings for feasibility items. The thick line within a box plot represents the median, the diamond represents the mean, the upper boundary of the box indicates the 25th
percentile (Q1) and lower boundary the 75th percentile (Q3). The whiskers above and below the box indicate the minimal and maximal values (Q1 – 1.5*IQR and Q3 + 1.5*IQR respectively), while points above the upper or below the whiskers indicate outliers. Subjective ratings were measured using a 7-point Likert scale with -3 = Strongly disagree, -2 = Disagree; -1 = Somewhat disagree; 0 = Neither agree nor disagree; 1 = Somewhat agree; 2 = Agree; 3= Strongly agree.
Table

Table 1

<table>
<thead>
<tr>
<th>Items</th>
<th>Domain</th>
</tr>
</thead>
<tbody>
<tr>
<td>Q1 It seemed as if the flashing was my respiration</td>
<td>Breathing awareness</td>
</tr>
<tr>
<td>Q2 It seemed as if I had three bodies</td>
<td>Control</td>
</tr>
<tr>
<td>Q3 I felt as if the virtual body was breathing with me</td>
<td>Breathing agency</td>
</tr>
<tr>
<td>Q4 I had difficulty breathing</td>
<td>Breathing difficulty</td>
</tr>
<tr>
<td>Q5 I felt as if the virtual body was drifting with the flashing</td>
<td>Control</td>
</tr>
<tr>
<td>Q6 My breathing was enjoyable</td>
<td>Comfort</td>
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Table 1 Subjective questionnaire items
Table 2

<table>
<thead>
<tr>
<th>Patient Characteristics</th>
<th>Total (n=26)</th>
<th>Synchronous first (n=14)</th>
<th>Asynchronous first (n=12)</th>
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</thead>
<tbody>
<tr>
<td><strong>Gender</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male N (%)</td>
<td>19 (73%)</td>
<td>11 (79%)</td>
<td>8 (67%)</td>
</tr>
<tr>
<td>Female N (%)</td>
<td>7 (27%)</td>
<td>3 (21%)</td>
<td>4 (33%)</td>
</tr>
<tr>
<td><strong>Age (years)</strong></td>
<td>55/18 (35-81)</td>
<td>55/18 (38-81)</td>
<td>56.5/16.75 (35-73)</td>
</tr>
<tr>
<td><strong>MoCA</strong></td>
<td>27/3 (25-30)</td>
<td>27.5/1.75 (25-30)</td>
<td>27/4.24 (25-30)</td>
</tr>
<tr>
<td><strong>SpO2 on oxygen therapy</strong></td>
<td>94/4.3 (90-98)</td>
<td>95/5.5 (90-98)</td>
<td>92.5/4 (91-96)</td>
</tr>
<tr>
<td><strong>Oxygen flow (l/m)</strong></td>
<td>1/3 (0-8)</td>
<td>2/2 (0-8)</td>
<td>0.25/1.75 (0-4)</td>
</tr>
<tr>
<td><strong>Heart rate (bpm)</strong></td>
<td>74.5/22 (52-108)</td>
<td>79.5/24.5 (62-108)</td>
<td>70.5/11.5 (62-92)</td>
</tr>
<tr>
<td><strong>Days since first symptom onset</strong></td>
<td>17/22 (3-39)</td>
<td>14/21 (3-39)</td>
<td>18.5/16.25 (6-37)</td>
</tr>
<tr>
<td><strong>Contagious at time of testing N (%)</strong></td>
<td>16 (62%)</td>
<td>10 (71%)</td>
<td>7 (58%)</td>
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<tr>
<td><strong>Primary Outcomes</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Breathing comfort</strong></td>
<td>1/2 (-3-2)</td>
<td>0/2 (-3-2)</td>
<td>1/3.25 (-2-2)</td>
</tr>
<tr>
<td><strong>Breathing difficulty</strong></td>
<td>1/3 (-3-2)</td>
<td>1/3 (-3-2)</td>
<td>0/3.25 (-3-2)</td>
</tr>
<tr>
<td><strong>Secondary Outcomes</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Respiratory rate (bpm)</strong></td>
<td>21/10(7-35)</td>
<td>21/10(15-32)</td>
<td>22/10 (15-35)</td>
</tr>
<tr>
<td><strong>Respiratory rate variability (bpm)</strong></td>
<td>3/2(1-12)</td>
<td>3/2 (1-12)</td>
<td>3/2 (1-5)</td>
</tr>
</tbody>
</table>

Table 2 Characteristics of the patients at randomisation in the intent to treat population and primary, secondary outcomes. Data are presented as n (%) or median/IQR(Range). IQR: interquartile range. *Data were missing for some patients; the denominator in the asynchronous group was 10.
<table>
<thead>
<tr>
<th></th>
<th>beta</th>
<th>95% CI LB</th>
<th>95% CI UP</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Breathing comfort</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Main effect of Synchrony*</td>
<td>0.542</td>
<td>0.046</td>
<td>1.037</td>
<td>0.033</td>
</tr>
<tr>
<td>Asynch First*</td>
<td>0.25</td>
<td>-0.627</td>
<td>1.127</td>
<td>0.223</td>
</tr>
<tr>
<td>Synch First*</td>
<td>-0.583</td>
<td>-1.544</td>
<td>0.378</td>
<td></td>
</tr>
<tr>
<td>Breathing difficulty</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Main effect of Synchrony*</td>
<td>-0.5</td>
<td>-1.064</td>
<td>0.064</td>
<td>0.080</td>
</tr>
<tr>
<td>Asynch First*</td>
<td>-1.333</td>
<td>-2.260</td>
<td>-0.407</td>
<td>0.221</td>
</tr>
<tr>
<td>Synch First*</td>
<td>-0.667</td>
<td>-1.760</td>
<td>0.427</td>
<td></td>
</tr>
<tr>
<td>Agency</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Main effect of Synchrony*</td>
<td>1.583</td>
<td>0.335</td>
<td>2.832</td>
<td>0.014</td>
</tr>
<tr>
<td>Asynch First*</td>
<td>-0.667</td>
<td>-1.922</td>
<td>0.589</td>
<td>0.336</td>
</tr>
<tr>
<td>Synch First*</td>
<td>1.167</td>
<td>-1.285</td>
<td>3.618</td>
<td></td>
</tr>
<tr>
<td>Awareness</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Main effect of Synchrony*</td>
<td>2.167</td>
<td>1.068</td>
<td>3.266</td>
<td>&lt; 0.0001</td>
</tr>
<tr>
<td>Asynch First*</td>
<td>0.167</td>
<td>-0.894</td>
<td>1.227</td>
<td>0.064</td>
</tr>
<tr>
<td>Synch First*</td>
<td>2</td>
<td>-0.121</td>
<td>4.121</td>
<td></td>
</tr>
<tr>
<td>Control (Q2)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Main effect of Synchrony*</td>
<td>0.042</td>
<td>-0.042</td>
<td>0.125</td>
<td>0.312</td>
</tr>
<tr>
<td>Asynch First*</td>
<td>-3.000</td>
<td>-3.135</td>
<td>-2.865</td>
<td>0.302</td>
</tr>
<tr>
<td>Synch First*</td>
<td>0.083</td>
<td>-0.080</td>
<td>0.246</td>
<td></td>
</tr>
<tr>
<td>Control (Q5)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Main effect of Synchrony*</td>
<td></td>
<td></td>
<td></td>
<td>Did not converge as data are similar in both conditions</td>
</tr>
<tr>
<td>Asynch First*</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Synch First*</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Respiration Rate</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Main effect of Synchrony*</td>
<td>-0.275</td>
<td>-1.748</td>
<td>1.198</td>
<td>0.704</td>
</tr>
<tr>
<td>Asynch First*</td>
<td>23.346</td>
<td>19.460</td>
<td>27.231</td>
<td>0.053</td>
</tr>
<tr>
<td>Synch First*</td>
<td>2.685</td>
<td>-0.041</td>
<td>5.410</td>
<td></td>
</tr>
<tr>
<td>Respiration Rate Variability</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Main effect of Synchrony*</td>
<td>-0.295</td>
<td>-0.779</td>
<td>0.190</td>
<td>0.222</td>
</tr>
<tr>
<td>Asynch First*</td>
<td>4.594</td>
<td>3.344</td>
<td>5.843</td>
<td>0.810</td>
</tr>
<tr>
<td>Synch First*</td>
<td>-0.114</td>
<td>-1.082</td>
<td>0.854</td>
<td></td>
</tr>
</tbody>
</table>
Table 3 Results summary for the subjective and physiological measures. #: depicts the mean difference between synchronous and asynchronous conditions, regardless of the sequence, as well as its CI estimated by the linear mixed model (the p-value corresponds to the test of this difference being equal to zero); ¶: depicts the mean difference between synchronous and asynchronous conditions and its CI estimated by the linear mixed model for the experimental sequence “Asynchronous first” (the p-value corresponds to the result of the interaction test); +: depicts the mean difference between synchronous and asynchronous conditions and its CI estimated by the linear mixed model for the experimental sequence “Synchronous first” (the p-value corresponds to the result of the interaction test). CI: confidence interval, LB: Lower Bound, UB: Upper Bound. Subjective ratings were measured using a 7-point Likert scale with -3 = Strongly disagree, -2 = Disagree, -1 = Somewhat disagree; 0 = Neither agree nor disagree; 1 = Somewhat agree; 2 = Agree; 3 = Strongly agree.
**Conflict of interest statements**

CS and KOA were employees of MindMaze SA at the time of the study. No MindMaze SA products were used in this study. BO is member of the board and shareholder of Mindmaze SA. BO is founder and shareholder of Metaphysiks SA.

**Fundings information**

Dr Betka Sophie’s salary has been funded by a Marie Skłodowska-Curie Individual Fellowship (H2020-MSCA-IF-2019 894111/ RESPVR), awarded by the European Commission. Olaf Blanke is funded by the Bertarelli Foundation.

**Data sharing**

The anonymised patient data will be shared, while safeguarding the privacy of patients, via the LNCO data repository (https://gitlab.epfl.ch/LNCO-public).

**Acknowledgements**

We thank Dr Fosco Bernasconi for his statistical advice.

**Ethics committee approval**

This single-site study was carried out at the University Hospital (HUG) in Geneva, Switzerland and was approved by the Commission Cantonale d’Ethique de la Recherche de la République et Canton de Genève (2019-02360).
Figure 1

39 patients assessed for eligibility

- 5 ineligible (MoCA<25)
- 8 declined

26 enrolled

26 randomised

14 assigned to Synch-VR

- 2 discontinued
- 2 technical issues

12 assigned to Asynch-VR

Cross-Over

12 assigned to Synch-VR

12 assigned to Asynch-VR

24 included in analysis
Figure 2
Figure 3

(A) Breathing comfort
(B) Agency
(C) Awareness
(D) Breathing difficulty
(E) Control (Q2)
(F) Control (Q5)
Supplementary material

Virtual reality intervention alleviates dyspnea in patients recovering from COVID pneumonia
Betka Sophie*, Kannape Oliver Alan*, Fasola Jemina¹, Lance Florian³, Cardin Sylvain², Schmit Aline³, Similowski Thomas⁴, Soccal Paola Marina³, Herbelin Bruno**, Adler Dan**, Blanke Olaf**

* contributed equally to the publication
** contributed equally to the publication

¹Laboratory of Cognitive Neuroscience, Brain Mind Institute and Center for Neuroprosthetics, Faculty of Life Sciences, Ecole Polytechnique Federale de Lausanne, (EPFL), Geneva, 1202, Switzerland
²Mindmaze SA, Lausanne, Switzerland
³Division of Lung Diseases, University Hospital and Geneva Medical School, University of Geneva, Switzerland
⁴Sorbonne Université, INSERM, UMR1158 Neurophysiologie Respiratoire Expérimentale et Clinique, F-75005 Paris, France
⁵AP-HP, Groupe Hospitalier Universitaire APHP-Sorbonne Université, site Pitié-Salpêtrière, Département R3S (Respiration, Réanimation, Réhabilitation respiratoire, Sommeil), F-75013 Paris, France

Corresponding author: Prof. Olaf Blanke, Dr Dan Adler
Email: olaf.blanke@epfl.ch; Dan.Adler@hcuge.ch
Tel: +41 21 693 96 21
Supplementary material

Screening
Patients were screened by a respiratory physician during morning rounds on weekdays only, except for some leave days, in the division of lung diseases of HUG. Once referred, it was verified that the patients met the inclusion criteria by performing an anamnestic interview as well as the MoCA. The delay between the initial screening by the physician and the inclusion by the researcher varied between 1 hour to 2 days.

Randomization and masking
Patients were allocated to one of two starting conditions using a randomization script (i.e., randomizing the experimental sequence for each patient, using Matlab version 2020a), before data collection. Randomization was not restricted; no stratification or minimization factors were applied. Allocation was concealed to the clinicians screening patients.

Participant masking (blinding) was achieved by keeping both the procedure and the virtual environment identical for both tested conditions. Participants were naïve to the difference in the two conditions which consisted only of a change in feedback synchrony between respiratory movements and virtual body luminance. Experimenters were not blinded, however, the instructions were only given to the patient once and applied to both conditions.

Power calculation
Based on previous work on breathing agency¹, a sample size of 21 patients was estimated, using a two-sided paired t-test with an effect size of 0.65, alpha of 0.05 and a power of 0.8 to demonstrate a breathing agency difference of 0.5 point measured on a 7-point Likert-scale from -3 to 3- in subjective ratings between the two experimental conditions (i.e., synchronous and asynchronous).

Risk assessment
As head-mounted displays are widely used in clinical and healthy populations no specific safety analysis was performed within the scope of this study. There is no evidence that using HMDs carry risks beyond those of CRT screens (e.g., with respect to binocular vision or photosensitive epilepsy). As patients remained seated during the intervention there was no risk of falling or collisions.
<table>
<thead>
<tr>
<th>Items</th>
<th>Domain</th>
</tr>
</thead>
<tbody>
<tr>
<td>Q1 Did you enjoy the VR experience?</td>
<td>Satisfaction</td>
</tr>
<tr>
<td>Q2 Would you like to continue using the device during your recovery?</td>
<td>Rehabilitation</td>
</tr>
<tr>
<td>Q3 Would you have liked to use this earlier during your stay at the hospital?</td>
<td>Hospital Use</td>
</tr>
<tr>
<td>Q4 Would you like to continue using the device at home?</td>
<td>Home Use</td>
</tr>
<tr>
<td>Q5 Do you think the VR feedback improved your breathing?</td>
<td>Respiratory benefit</td>
</tr>
<tr>
<td>Q6 Did the VR feedback make you feel better?</td>
<td>Well-being benefit</td>
</tr>
</tbody>
</table>

Table S1 Feasibility questionnaire
Supplementary Results

Means and standard deviations in function of the experimental sequence

<table>
<thead>
<tr>
<th>Measures</th>
<th>Asynchronous Condition</th>
<th>Synchronous Condition</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Asynchronous first (n=12)</td>
<td>Synchronous first (n=12)</td>
</tr>
<tr>
<td>Comfort</td>
<td>Median</td>
<td>IQR</td>
</tr>
<tr>
<td>Difficult</td>
<td>-2</td>
<td>2</td>
</tr>
<tr>
<td>Agency</td>
<td>-2</td>
<td>3,5</td>
</tr>
<tr>
<td>Awareness</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td>Control (Q2)</td>
<td>-3</td>
<td>0</td>
</tr>
<tr>
<td>Control (Q5)</td>
<td>-3</td>
<td>0</td>
</tr>
<tr>
<td>Respiratory rate</td>
<td>20,33</td>
<td>6,34</td>
</tr>
<tr>
<td>Respiratory rate Variability</td>
<td>4,26</td>
<td>2,43</td>
</tr>
</tbody>
</table>

Table S2 Medians and interquartile ranges for the asynchronous and synchronous conditions, in function of the experimental sequence.

Breathing comfort - Tests against baseline

Using post-hoc paired one-sided t-tests, we found a significant difference between breathing comfort ratings during the synchronous condition compared to baseline (Difference: 1.25±0.431, t = 2.901, p < 0.01, 95% CI 0.511 to inf). This was not observed between breathing comfort ratings during the asynchronous condition compared to baseline (Difference: 0.708±0.547, t = 1.296, p > 0.05, 95% CI -0.229 to inf), excluding a mere effect of VR distraction.
Figure S1 Objective measures for which the main effect of the experimental manipulation was not significant. The boxplots depicting subjects’ physiology signal during asynchronous condition compared to the synchronous condition, independent of experimental sequence. The thick line within a box plot represents the median, the diamond represents the mean, the upper boundary of the box indicates the 25th percentile (Q1) and lower boundary the 75th percentile (Q3). The whiskers above and below the box indicate the minimal and maximal values (Q1 – 1.5*IQR and Q3 + 1.5*IQR respectively), while points above the upper or below the whiskers indicate outliers. Subjective ratings were measured using a 7-point Likert scale with -3 = Strongly disagree, -2 = Disagree; -1 = Somewhat disagree; 0 = Neither agree nor disagree; 1 = Somewhat agree; 2 = Agree; 3= Strongly agree.
Feasibility

<table>
<thead>
<tr>
<th>Items</th>
<th>Mean</th>
<th>SD</th>
<th>Median</th>
<th>IQR</th>
<th>t</th>
<th>df</th>
<th>p-value</th>
<th>CI LB</th>
<th>CI UB</th>
</tr>
</thead>
<tbody>
<tr>
<td>Satisfaction</td>
<td>1.75</td>
<td>1.649</td>
<td>2.00</td>
<td>2.00</td>
<td>5.201</td>
<td>23.000</td>
<td>0.000</td>
<td>1.173</td>
<td>inf</td>
</tr>
<tr>
<td>Rehabilitation</td>
<td>0.042</td>
<td>2.032</td>
<td>0.00</td>
<td>4.00</td>
<td>0.100</td>
<td>23.000</td>
<td>0.460</td>
<td>-0.669</td>
<td>inf</td>
</tr>
<tr>
<td>Usage at the hospital</td>
<td>-0.083</td>
<td>1.863</td>
<td>0.00</td>
<td>2.25</td>
<td>0.219</td>
<td>23.000</td>
<td>0.586</td>
<td>-0.735</td>
<td>inf</td>
</tr>
<tr>
<td>Usage at home</td>
<td>-0.333</td>
<td>2.22</td>
<td>-1.00</td>
<td>4.25</td>
<td>0.736</td>
<td>23.000</td>
<td>0.765</td>
<td>-1.110</td>
<td>inf</td>
</tr>
<tr>
<td>Respiratory benefit</td>
<td>0.708</td>
<td>1.922</td>
<td>1.00</td>
<td>2.25</td>
<td>1.806</td>
<td>23.000</td>
<td>0.042</td>
<td>0.036</td>
<td>inf</td>
</tr>
<tr>
<td>Well-being benefit</td>
<td>0.167</td>
<td>2.297</td>
<td>0.50</td>
<td>4.00</td>
<td>0.355</td>
<td>23.000</td>
<td>0.363</td>
<td>-0.637</td>
<td>inf</td>
</tr>
</tbody>
</table>

Table S3 Descriptive statistics and statistical tests of the feasibility items. Ratings were measured using a 7-point Likert scale with -3 = Strongly disagree, -2 = Disagree; -1 = Somewhat disagree; 0 = Neither agree nor disagree; 1 = Somewhat agree; 2 = Agree; 3 = Strongly agree. SD = standard deviation, IQR = Interquartile range, df = degree of freedom.
Figure S2 Percentage of feasibility scores for all items, rated on a 7-point Likert scale, with -3 = Strongly disagree, -2 = Disagree; -1 = Somewhat disagree; 0 = Neither agree nor disagree; 1 = Somewhat agree; 2 = Agree; 3= Strongly agree

Reference