Intensive Care Unit–Specific Virtual Reality for Critically Ill Patients With COVID-19: Multicenter Randomized Controlled Trial

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Abstract

Background: Although psychological sequelae after intensive care unit (ICU) treatment are considered quite intrusive, robustly effective interventions to treat or prevent these long-term sequelae are lacking. Recently, it was demonstrated that ICU-specific virtual reality (ICU-VR) is a feasible and acceptable intervention with potential mental health benefits. However, its effect on mental health and ICU aftercare in COVID-19 ICU survivors is unknown.

Objective: This study aimed to explore the effects of ICU-VR on mental health and on patients’ perceived quality of, satisfaction with, and rating of ICU aftercare among COVID-19 ICU survivors.

Methods: This was a multicenter randomized controlled trial. Patients were randomized to either the ICU-VR (intervention) or the control group. All patients were invited to an COVID-19 post-ICU follow-up clinic 3 months after hospital discharge, during which patients in the intervention group received ICU-VR. One month and 3 months later (4 and 6 months after hospital discharge), mental health, quality of life, perceived quality, satisfaction with, and rating of ICU aftercare were scored using questionnaires.

Results: Eighty-nine patients (median age 58 years; 63 males, 70%) were included. The prevalence and severity of psychological distress were limited throughout follow-up, and no differences in psychological distress or quality of life were observed between the groups. ICU-VR improved satisfaction with (mean score 8.7, SD 1.6 vs 7.6, SD 1.6 [ICU-VR vs control]; t64=–2.82, P=.006) and overall rating of ICU aftercare (mean overall rating of aftercare 8.9, SD 0.9 vs 7.8, SD 1.7 [ICU-VR vs control]; t64=–3.25; P=.002) compared to controls. ICU-VR added to the quality of ICU aftercare according to 81% of the patients, and all patients would recommend ICU-VR to other ICU survivors.

Conclusions: ICU-VR is a feasible and acceptable innovative method to improve satisfaction with and rating of ICU aftercare and adds to its perceived quality. We observed a low prevalence of psychological distress after ICU treatment for COVID-19, and ICU-VR did not improve psychological recovery or quality of life. Future research is needed to confirm our results in other critical illness survivors to potentially facilitate ICU-VR’s widespread availability and application during follow-up.

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Introduction

The increase in the survival of critically ill patients admitted to the intensive care unit (ICU) in the last few decades has revealed the effect of ICU treatment on quality of life [1-3]. Up to one-third of “general” ICU survivors experience a poor quality of life, predominantly owing to psychological sequelae such as anxiety, depression, and posttraumatic stress disorder (PTSD) [4-7]. These psychological impairments comprise the psychological component of the postintensive care syndrome (PICS); they are common and can last months to even years after patient ICU discharge [4,8,9]. Consequently, there is a need for post-ICU care.

As the demand for ICU beds and critical care services has skyrocketed during the current COVID-19 outbreak, so would that of ICU aftercare. As such, health care services will have to adapt rapidly to an anticipated surge of ICU patients and post-ICU care, and this will place an enormous strain on acute services [10]. The current pandemic is highlighting the urgency for a multimodal follow-up program and the need for patient-focused innovative solutions [11]. Difficulty accessing in-person clinics is a key barrier in the development of an ICU follow-up program and is probably hindered more owing to current COVID-19 regulations [12].

Recovery from COVID-19 could have the same multifaceted problems that occur after sepsis and other critical illnesses [17]. Recently, virtual reality (VR) was demonstrated to be a useful technique to improve post-ICU mental health in sepsis survivors and could also be safely used and implemented in post-ICU COVID-19 care [18-20]. As such, we hypothesized that an ICU-specific virtual reality (ICU-VR) intervention could improve the satisfaction with and rating of ICU aftercare and could contribute to psychological recovery. The aims of this study were therefore to explore the effects of ICU-VR on mental health and on patients’ perceived quality and satisfaction with and rating of ICU aftercare among COVID-19 ICU survivors.

Methods

Study Design

This multicenter, open-label, randomized controlled trial was conducted in a university teaching hospital and in 3 university-affiliated secondary care hospitals. Patients were included from June 2020 to February 2021 and were followed-up for 6 consecutive months. The study protocol was approved by the Medical Ethics Committee of the Erasmus Medical Centre, Rotterdam, and the participating centers’ institutional review boards (NL.73667.078.20, approved June 10, 2020) and has previously been published [21].

Participants

All consecutive adult (≥18 years) patients who were treated in an ICU of one of the participating hospitals and visited the COVID-19 post-ICU follow-up clinic were eligible for inclusion. COVID-19 was diagnosed on the basis of a positive finding on reverse transcription–polymerase chain reaction (RT–PCR) for SARS-CoV-2. Exclusion criteria were primary neurological impairments or documented active psychiatric diseases, an inability to understand the Dutch language, absence of a formal home address, and participation in other interventional trials that could confound the primary outcome. Patients and the public were not involved in the design, conduct, reporting, or dissemination plans of this study. A former ICU patient was involved in the development of the ICU-VR intervention.

Randomization and Masking

Patients were randomly assigned to either the ICU-VR (intervention) group or the control group at a 1:1 ratio, using a centralized internet-based randomization procedure by the study site’s principal investigator or a representative (Castor Electronic Data Capture [EDC]). Patients were randomized in a simple manner without stratification. The investigators were unaware of the assignment sequence. Owing to the nature of the intervention, blinding of patients and investigators was not possible.

Intervention

All patients were invited to a COVID-19 post-ICU follow-up clinic as part of regional standard care. During this visit, patients had a 60-minute-long consultation with an intensivist and an ICU nurse, during which the ICU treatment was reviewed, and patients were screened for PICS-related impairments and referred to an appropriate health care worker, if appropriate.

Patients in the ICU-VR group received the ICU-VR intervention once during this visit. ICU-VR is explained in depth elsewhere. In short, it was developed by an interdisciplinary team that included intensivists, ICU nurses, a psychologist, a psychiatrist, an investigator, and former ICU patients and was previously demonstrated to be safe and feasible [18,22]. ICU-VR consists of a 14-minute-long informational video that can be watched using VR, in which the patient is exposed to the ICU environment and receives voice-over explanations regarding different facets of the surrounding ICU and ICU treatment. ICU-VR consists of 6 scenes: (1) The ICU physician and nurse welcome the patient in front of the ICU. After being brought to and installed in the ICU, explanations are given (2) about the surveillance monitor, medication pumps, intubation (including tracheal tube suction), mechanical ventilation, and prone

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positioning; (3) about intravenous drips and lines and tracheotomy, including its procedures; (4) about the treatment team taking care of the patient; (5) about isolation measures and personal protection equipment; and (6) about COVID-19 [19,21]. The script and the YouTube version can be found elsewhere [21,23]. The ICU-VR intervention was watched using head-mounted display–VR glasses (Oculus Go) in combination with headphones.

**Study Procedures**

All COVID-19 ICU survivors were invited to the hospital’s COVID-19 post-ICU follow-up clinic 3 months after hospital discharge as part of regional standard care. One month prior to this visit, eligible patients were sent a study information brochure, and 2 weeks later, patients were contacted by telephone by a member of the study team to explain the study procedures. During their follow-up clinic visit, consent was obtained, and patients were randomized.

Patients randomized to the ICU-VR group received the ICU-VR intervention once during the concordant follow-up clinic visit, whereas patients randomized to the control group did not receive ICU-VR. Aside from the ICU-VR intervention, there were no differences between the study groups. These results are part of a larger study evaluating the long-term effects of ICU-VR after 6 months and the effect of VR crossover [21].

Prior to the follow-up clinic visit and 4 and 6 months after hospital discharge—that is, 1 and 3 months after the COVID-19 post-ICU follow-up clinic visit—psychological distress and quality of life were assessed. Six months after hospital discharge, all patients were asked about their satisfaction with and rating of ICU care and aftercare, and patients in the intervention group were asked about their perspectives on ICU-VR.

**Outcomes**

Primary outcomes were PICS-related psychological distress and quality of life up to 6 months after hospital discharge and were mandatory parts of the questionnaire.

Psychological distress was expressed as the prevalence and severity of PTSD, anxiety, and depression-related symptoms assessed using the Impact of Event Scale-Revised (IES-R; PTSD) and the Hospital Anxiety and Depression Scale (HADS; anxiety and depression), respectively [24,25]. The IES-R is a self-reported measurement consisting of 22 items that assesses subjective distress caused by a traumatic event and has previously been validated in ICU survivors [24,26,27]. It provides a total score ranging from 0 to 88, with higher scores indicating more severe symptoms. It also provides subscale scores to assess symptoms of intrusion, avoidance, and hyperarousal, which is the sum of all items in each section. An IES-R total score of ≥34 is considered the optimal cutoff for PTSD [28]. The HADS consists of 14 items and is commonly used to determine the levels of anxiety and depression that a patient is experiencing and has been validated in critical illness survivors [29-31]. Seven of the items relate to anxiety, 7 relate to depression, and each question is answered on a 4-point Likert scale. A sum score of ≥8 (ranging from 0 to 21, with higher scores indicating more severe symptoms) on either the depression or the anxiety subscale, is classified as clinically meaningful depression and anxiety, respectively [29].

Quality of life was assessed using the Short-Form 36 (SF-36) and the European Quality of Life, 5 Dimensions (EQ-5D) questionnaires [32,33]. The EQ-5D and SF-36 have been validated and tested in the ICU and have been recommended for use in critical care medicine [34-38]. The EQ-5D measures quality of life in 5 dimensions (mobility, self-care, usual activities, pain or discomfort, and anxiety or depression). In each domain, patients are asked if they experience no, slight, moderate, severe, or extreme problems, from which the weight of a health state can be computed, ranging from –0.446 (worst quality of life) to 1.000 (best quality of life) [39]. Additionally, SF-36 is a 36-item, patient-reported survey of health and health-related quality of life (HRQoL). It consists of 8 scaled scores, which are the weighted sums of the questions in their section, and a scale for health change. Each scale is directly transformed to a scale ranging from 0 (worst score) to 100 (best score) on the assumption that each question carries an equal weight. The 8 sections are physical functioning, social functioning, physical role functioning, emotional role functioning, emotional well-being, vitality, bodily pain, and general health perception [40]. In addition to these scales, mental and physical component scores can be calculated, which represent a patient’s mental and physical health state. These scores are computed so that the mean is 50 (SD 10) for the general population [41,42].

Patients’ perceived quality of life and patients’ satisfaction with and rating of ICU aftercare were assessed using a novel questionnaire, and the questions were nonmandatory to answer. The questionnaire was based on the Patient Satisfaction Questionnaire and Family Satisfaction with ICU Care tools, altered to the needs of this study [43-45]. This questionnaire consisted of 21 items and was categorized into five sections: perspectives on the added value of ICU-VR to ICU care and ICU aftercare (8 questions), perspectives on the timing and number of sessions (3 questions), overall perspective on the ICU-VR intervention (3 questions), perspectives on the effect of the ICU-VR intervention (4 questions; Multimedia Appendix 1). The first section was (partly) answered by all patients, irrespective of the randomization allocation, and the other sections were answered only by patients randomized to the ICU-VR group. All questions could be answered on a 10-point Likert scale, ranging from 1 (not at all) to 10 (very much), except for perspectives on the timing and number of sessions. The questionnaire was administered by telephone.

Baseline characteristics and survival were determined through patient record analysis. Additional demographics, such as educational level and preadmission employment status, were assessed using follow-up questionnaires.

**Statistical Analysis**

Based on a previous pilot study examining the feasibility, safety, and clinical relevance of sepsis ICU-VR (Cohen d effect size=0.77), we assumed the effect estimates of ICU-VR to be

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similar in this study [20]. Using a 2-sided \( \alpha \) value of .05, a power of .80, a 1:1 randomization, and an expected loss to follow-up of 20%, we aimed to include a minimum of 80 patients, with 40 patients in each study group.

Baseline demographics and treatment-related characteristics were quantified using descriptive statistics. Continuous variables are expressed as median (IQR) or mean (SD) values, depending on their distribution. Categorical variables are presented as absolute numbers and relative frequencies.

Differences between study groups in continuous variables, such as the IES-R sum score, the HADS anxiety and depression scores, the SF-36 subscales and the EQ-5D utility score, at several follow-up time points were analyzed using a mixed-effects linear regression model with a random intercept for each study site. Differences in continuous outcomes at the 3-month follow-up time point were adjusted by adding the 3-month outcome as an independent variable in the mixed-effects linear regression model. Patients were categorized on the basis of clinically meaningful cutoffs for the IES-R sum score and the HADS anxiety and depression scores. Differences in categorical variables between study groups at several follow-up time points were analyzed using a mixed-effects logistic regression model with a random effect for each site. Differences in categorical outcomes at the 3-month follow-up time point were adjusted by adding the 3-month outcome as an independent variable in the mixed-effects logistic regression model. Differences in continuous or categorical variables throughout follow-up were analyzed using a mixed-effects linear or logistic regression model with time, randomization, and a random intercept or slope for each individual and each study site as appropriate. Differences in linear or categorical outcomes at the 3-month follow-up time point were added to the mixed-effects linear or logistic model as independent variables to adjust for that difference.

Outcomes of the mixed-effects linear regression models are reported as coefficient (95% CI) values, which implies the estimated mean difference, and outcomes of the mixed-effects logistic regression models are reported as odds ratios (ORs) with corresponding 95% CI values.

All data were gathered using Castor EDC. All analyses were performed using SPSS (version 24.0; SPSS Inc) and R for Statistics (R Foundation for Statistical Computing). A \( P \) value of \( \leq .05 \) was considered statistically significant.

Data Sharing
All data sets created during this study are available upon reasonable request by the corresponding author.

Results
Results Overview
A total of 147 patients visited the COVID-19 post-ICU follow-up clinic, of whom 89 were enrolled (inclusion rate: 61%): 45 patients in the ICU-VR group and 44 patients in the control group (Figure 1). All patients in the ICU-VR group completed the ICU-VR intervention, and no adverse events were reported. Baseline demographics and treatment-related characteristics were well balanced between groups (Table 1). The mean age was 58 (SD 11) years, 63 patients (71%) were male, and the median ICU length of stay was 17 (IQR 9-29) days. Table 1 shows baseline demographics and treatment-related characteristics.
Figure 1. CONSORT (Consolidated Standards of Reporting Trials) flow diagram of the study. ICU-VR: intensive care unit–specific virtual reality.
Table 1. Baseline demographics and treatment-related characteristics.

| Characteristics | ICU-VR<sup>b</sup> group (intervention) (n=45) | Control group (n=44) |
|-----------------|---------------------------------------------|---------------------|
| **Baseline demographics** | | |
| Age (years), median (IQR) | 61 (54-65) | 59 (51-65) |
| Males, n (%) | 35 (78) | 28 (36) |
| **BMI, median (IQR)** | | |
| 27.6 (25.3-31.1) | 28.0 (25.3-31.2) |
| Participants with a BMI of >30, n (%) | 14 (31) | 18 (41) |
| **Educational level, n (%)** | | |
| Primary education | 14 (31) | 13 (30) |
| Intermediate vocational education | 15 (33) | 20 (46) |
| Higher vocational education | 13 (29) | 7 (16) |
| Academic education | 3 (7) | 4 (9) |
| Employment status, employed, n (%) | 23 (51) | 21 (48) |
| **Treatment-related characteristics** | | |
| Length of stay in the intensive care unit (days), median (IQR) | 14 (9-25) | 14 (7-28) |
| Length of hospital stay (days), median (IQR) | 22 (12-32) | 24 (13-40) |
| **Mechanical ventilation, n (%)** | | |
| Duration (hours), median (IQR) | 227 (169-343) | 383 (206-465) |
| Highest positive end-respiratory pressure (cm H<sub>2</sub>O), median (IQR) | 21 (17-28) | 20 (16-25) |
| Lowest fraction of inspired oxygen (%), median (IQR) | 28 (24-30) | 25 (22-30) |
| Lowest ratio of arterial oxygen (mm Hg), median (IQR) | 0.11 (0.09-0.23) | 0.11 (0.09-0.18) |
| Prone positioning, n (%) | 35 (77) | 36 (82) |
| **Medication** | | |
| Received noradrenaline, n (%) | 37 (82) | 35 (80) |
| Noradrenaline dose (µg/kg/minute), median (IQR) | 0.17 (0.10-0.30) | 0.14 (0.08-0.29) |
| Duration of noradrenaline use (hours), median (IQR) | 186 (32-249) | 167 (96-349) |
| Received midazolam, n (%) | 35 (78) | 33 (75) |
| Midazolam dose (mg/kg/hour), median (IQR) | 0.59 (0.43-0.71) | 0.51 (0.39-0.66) |
| Duration of midazolam use (hours), median (IQR) | 20 (13-93) | 20 (13-36) |
| Received remifentanil, n (%) | 32 (71) | 35 (80) |
| Remifentanil dose (µg/kg/hour), median (IQR) | 14 (10-16) | 14 (6-18) |
| Duration of remifentanil use (hours), median (IQR) | 33 (22-80) | 32 (23-72) |
| Received sufentanil, n (%) | 26 (58) | 28 (63) |
| Sufentanil dose (µg/kg/hour), median (IQR) | 0.55 (0.34-0.83) | 0.60 (0.38-0.70) |
| Duration of sufentanil use (hours), median (IQR) | 8 (1-13) | 10 (6-14) |
| Received rocuronium, n (%) | 22 (49) | 16 (36) |
| Rocuronium dose (mg/kg/hour), median (IQR) | 0.39 (0.05-0.77) | 0.32 (0.01-0.60) |
| Duration of rocuronium use (hours), median (IQR) | 22 (0-28) | 17 (0-22) |
| **Illness severity scores** | | |
| Simplified Acute Physiology Score (version 2), median (IQR) | 31 (26-36) | 31 (26-35) |
| Acute Physiology and Chronic Health Evaluation (version 4) score, median (IQR) | 49 (38-60) | 49 (42-59) |
| Admission Sequential Organ Failure Assessment score, median (IQR) | 2 (1-6) | 2 (1-4) |
Characteristics\textsuperscript{a} & ICU-VR\textsuperscript{b} group (intervention) (n=45) & Control group (n=44) \\
Highest Sequential Organ Failure Assessment score, median (IQR) & 8 (6-10) & 7 (6-9) \\

\textsuperscript{a}Baseline demographics and treatment-related characteristics were obtained at 3 months after hospital discharge via digital patient records.

\textsuperscript{b}ICU-VR: intensive care unit-specific virtual reality.

**Psychological Component of the PICS**

At the 3-month follow-up time point, a total of 31 of 89 patients (34\%) reported psychologic distress, with 10 patients (22\%) in the ICU-VR group and 21 patients (47\%) in the control group (OR 3.5, 95\% CI 1.4-8.9, \(P<.01\)). At 4 months, 38 patients (43\%) reported psychological distress, with 12 patients (27\%) in the ICU-VR group and 26 patients (59\%) in the control group (OR 3.0, 95\% CI 0.8-11.9, \(P=.11\)). At 6 months, 24 patients (31\%) reported psychological distress, with 9 patients (23\%) in the ICU-VR group and 15 patients (39\%) in the control group (OR 0.7, 95\% CI 0.2-2.9, \(P=.60\)).

At the 3-month follow-up time point, 7 patients (16\%) in the ICU-VR group and 16 patients (38\%) in the control group reported probable PTSD (OR 3.2, 95\% CI 1.4-8.9, \(P<.01\); Figure 2B). During follow-up, no differences were observed in PTSD scores or the proportion of patients who reported probable PTSD between randomization allocations (Figures 2A and 2B). Throughout follow-up, the PTSD score remained similar at 4 months (\(\beta=–.60\), 95\% CI –3.2 to 1.9, \(P=.63\)) after hospital discharge but improved at 6 months after hospital discharge (\(\beta=–3.1\), 95\% CI –5.8 to –0.4, \(P=.02\)). However, this improvement was independent of the randomization group (\(\beta=5.4\), 95\% CI –0.2 to 11.1, \(P=.06\)).

At the 3-month follow-up time point, 4 patients (9\%) in the ICU-VR group and 10 patients (22\%) in the control group reported probable anxiety (OR 3.3, 95\% CI 1.2-9.3, \(P=.02\); Figure 2D). Four months after hospital discharge, ICU-VR resulted in fewer patients with probable anxiety (n=8, 18\% vs 22, 50\%; OR 3.8, 95\% CI 1.1-12.7, \(P=.03\); Figure 2C) but not lower anxiety scores (median HADS anxiety score 3, IQR 1-5 vs 7, IQR 2-11; \(\beta=1.4\), 95\% CI –0.1 to 3.0; \(P=.07\)). There were no differences at 4 or 6 months after hospital discharge (Figures 2C and 2D). No natural decline in anxiety was observed, and the severity of anxiety and the prevalence of probable anxiety were lower in the ICU-VR group throughout the follow-up.

At the 3-month follow-up time point, 8 (18\%) patients in the ICU-VR group and 14 (33\%) patients in the control group reported probable depression (OR 2.3, 95\% CI 0.9-6.3, \(P=.10\); Figure 2F). Throughout the follow-up, no difference in the depression scores or the proportion of patients reporting probable depression was observed (Figures 2E and 2F). The severity of depression remained similar throughout the follow-up period.
Health-Related Quality of Life

The overall health-related quality of life, mental health–related quality of life, and physical health–related quality of life are depicted in Figure 3. Throughout the follow-up period, the overall, mental, and physical HRQoL remained similar until 4 months but improved at 6 months after hospital discharge, while overall quality of life, outcomes of individual EQ-5D domains, and subscales of the SF-36 score differed between groups during the follow-up period (Figures 3A-D, Multimedia Appendix 2 and Multimedia Appendix 3).
Figure 3. Quality of life outcomes. Boxplots of the overall quality of life (A), perceived health state (B), mental quality of life (C), and physical quality of life (D). Overall quality of life was expressed as the EQ-5D TTO score, the perceived health state as the EQ-5D VAS score, and the mental and physical quality of life as the mental and physical component scales of the SF-36, respectively. Differences between randomization groups at each follow-up time point and between follow-up time points (p, Time) and throughout the follow-up (p, Randomization) were analyzed using mixed-effects linear (severity) or logistic (prevalence) regression models. EQ-5D: European Quality of Life, 5 dimensions, ICU-VR: intensive care unit–virtual reality, MCS-36: Mental Component Summary, 36 items, PCS-36: Physical Component Summary, 36 items, TTO: trade time-off, VAS: visual analog scale.

Perspectives on ICU-VR

In total, 37 patients (84%) in the ICU-VR group and 32 patients (71%) in the control group gave their perspective about the intervention and the received care and aftercare (Figure 4 and Multimedia Appendix 4). Patients in the intervention group were more satisfied with the ICU aftercare (mean score: 8.7, SD 1.6 vs 7.6, SD 1.6 [ICU-VR vs control], \( t_{64} = -2.82, P = .006 \)) but not with the ICU care (mean score 8.9, SD 1.5 vs 8.5, SD 1.5 [ICU-VR vs control], \( t_{64} = -0.92; P = .36 \); Figure 4A). Additionally, patients in the intervention group rated the ICU aftercare higher (mean overall rating of aftercare 8.9, SD 0.9 vs 7.8, SD 1.7 [ICU-VR vs control], \( t_{64} = -3.25; P = .002 \)) but not the ICU care (mean score 8.9, SD 1.5 vs 8.7, SD 1.2 [ICU-VR vs control], \( t_{64} = -0.59; P = .56 \); Figure 4B). ICU-VR added to the satisfaction of ICU care according to 62% of patients (Figure 4C), satisfaction with ICU aftercare according to 65% of patients (Figure 4D), quality of ICU care according to 62% patients (Figure 4E), and quality of ICU aftercare according to 81% of patients in the ICU-VR group (Figure 4F).

Patients in the intervention group assigned a mean score of 8.7 (SD 1.0) out of 10 to the ICU-VR, on a Likert scale, and stated that ICU-VR improved their understanding of ICU treatment (score>5, 76%; mean score 7.2, SD 2.5) and decreased their frightening memories (score>5, n=24-37, 65%; mean score 6.6, SD 2.8; Multimedia Appendix 4).
**Discussion**

**Principal Findings**

We observed that ICU-VR improved patients’ perceived quality of, satisfaction with, and rating of ICU aftercare among COVID-19 ICU survivors. This method is feasible, acceptable, and innovative and could be implemented in regional ICU aftercare. Our results also demonstrate that approximately 31% of COVID-19 ICU survivors experienced decreased mental health in terms of psychological distress up to 6 months after.
hospital discharge and that ICU-VR did not improve psychological recovery or quality of life.

In contrast to our previous findings regarding patients with sepsis and a recent COVID-19 case report, we did not observe improved mental health or quality of life in the ICU-VR group [19,20]. In contrast to this study, we provided ICU-VR earlier post ICU admission (median 7-8 days) in our previous study. Notably, the patient with COVID-19 in our case report and those in the sepsis study had robust responses in terms of psychological distress symptoms, including PTSD. The COVID-19 critical illness survivors in this study received ICU-VR much later (3 months after hospital discharge). Therefore, the timing of the ICU-VR intervention could be important for its therapeutic effect. Although 3 months after hospital discharge is a clinically feasible time point, it can be argued that PTSD and anxiety, at that moment, have already fully developed, and treatment of fully established psychiatric disorders may require more complex treatment strategies. When ICU-VR is offered soon after ICU discharge; that is, in the initial few weeks patients are still processing what happened to them, and ICU-VR could be a valuable adjunct to improve factual recall and decrease frightening memories. In future studies, the timing of ICU-VR and the number of sessions needed should be further investigated.

The number of desired or needed VR sessions remains a matter of debate, and no study has determined the optimal number of sessions after ICU admission. Although an average of 8-14 sessions is used in nonhospital settings, we previously demonstrated that sepsis survivors desire a median of only one session [20,46,47]. In this study, more than half of the patients did not desire the ICU-VR intervention multiple times, although there was substantial interpatient variability. An important difference between the current study and the sepsis trial is that in the sepsis trial, patients could self-determine how many sessions they desired, and this could have potentially increased the effectiveness. Therefore, a more patient-centered approach instead of a prespecified number of times might be more suitable, though guidelines are currently lacking.

Additionally, we observed lower overall incidence rates of PTSD (22% vs 11%), anxiety (46% vs 21%), and depression (41% vs 18%) at 3 months compared to a recent nationwide study in the United Kingdom, which included all patients who received at least 24 hours of ICU treatment, and compared to previously observed studies involving patients with COVID-19, acute lung injury (and acute respiratory distress syndrome) survivors, and a Dutch cohort of critical illness survivors [8,9,16,48-50]. Importantly, our power calculation was based on the prevalence rates of psychological distress. This lower incidence might explain the lack of ICU-VR effectiveness for this item. The lack of predisposing factors (such as a pre-existing cognitive impairment) could possibly explain the low prevalence of PTSD and depression in the current population [51-53].

Satisfaction during and after ICU admission is increasingly becoming an issue of interest considering that low satisfaction negatively impacts psychological sequelae after critical care [54]. Evidence suggests that patients generally indicate that they are satisfied with ICU care [55,56]. We found similarly high levels of reported satisfaction and showed that despite these high numbers, ICU-VR improved satisfaction, ratings, and perceived quality of ICU aftercare. Moreover, 100% would recommend ICU-VR to other patients. This seems to suggest that satisfaction with patient care does not imply that there are no problems regarding some aspects of their inpatient experience or that they fully comprehended all ICU-related information. Our findings actually confirm the results from a recent review, which concluded that patients’ support after ICU admission is multifaceted and varies across several transition points after ICU discharge [57]. An analogy can be made to civil aviation, where satisfaction may be high, but customers still complain about specific aspects of the service [58]. ICU-VR could therefore serve as an additional modality to fulfill several individual patient needs during the transition from ICU to home. Additionally, despite the lack of a successful effect on “traditional” measurements, such as the psychological questionnaires in this study, more than half of the patients experienced a decrease in frightening memories. Therefore, ICU aftercare might be more complex than we thought and may require a more patient-centered approach for measuring the results of novel intervention methods.

Limitations

Several study limitations should be acknowledged. First, despite our randomization procedure, there were statistically significant differences in primary outcome measures between groups at the 3-month follow-up time point. To ensure that no effect was overestimated, we adjusted our outcomes for the 3-month follow-up time point outcomes by adding them as independent predictors to our regression models. Although this difference was unexpected considering the randomization procedure, we could have prevented these differences by stratifying the randomization procedure on the presence of psychological distress at the follow-up time point prior to randomization. In future studies, we should consider this when possible. Second, as both the ICU-VR intervention and the questionnaire were in Dutch, we could only include patients able to understand the Dutch language. This may have resulted in selection bias, and we do not know how ICU-VR performs in nonnative Dutch speakers or if a translated version has an effect in these patients. This is especially of interest as, owing to language restrictions, these patients are expected to understand less of their ICU treatment than native Dutch patients and may therefore benefit more from such an intervention. Third, we used a novel set of questionnaires to assess patient experiences. Although these were based on and altered from the Patient Satisfaction Questionnaire and Family Satisfaction with ICU Care tools, these questionnaires have not yet been validated [43-45].

Conclusions

In conclusion, ICU-VR is a feasible and acceptable innovative method to improve patient satisfaction with and rating of ICU aftercare and adds to its perceived quality. We observed a low prevalence of psychological distress after COVID-19 ICU treatment, and ICU-VR did not improve psychological recovery or quality of life. Future studies should explore ICU-VR’s widespread availability and application during ICU follow-up
and should determine whether the timing of ICU-VR impacts its effect on psychological PICS-related sequelae.

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Authors' Contributions
JHV, JvB, EW, DG, and MEvG conceived the study and initiated the study design. MEvG is the coordinating investigator and grant holder. DG is the principal investigator. TIMK provided statistical expertise in the clinical trial design, and JHV and TIMK devised the statistical analysis plan. JvB, EW, JAML, and AFCS are the local principal investigators at each study site. MEH initiated the regional post–COVID-19 follow-up clinic. JHV and MEH composed the questionnaires used in the study. MvB and LLHS assisted in participant inclusion and data collection. MEvG and TIMK independently verified the data. JHV, JB, and MEvG wrote the first manuscript draft, and all authors helped to further draft the manuscript. All authors read and approved the final version of the manuscript.

Conflicts of Interest
None declared.

Multimedia Appendix 1
Translation of the novel questionnaire assessing ICU care and aftercare satisfaction and global rating, and the perspectives on the ICU-specific virtual reality (ICU-VR) intervention.

[PDF File (Adobe PDF File), 401 KB - Multimedia Appendix 1]

Multimedia Appendix 2
Table S1. Outcomes of the individual European Quality of Life, 5 dimensions domains.

[PDF File (Adobe PDF File), 562 KB - Multimedia Appendix 2]

Multimedia Appendix 3
Table S2. Subscales of the Short-Form 36 throughout follow-up.

[PDF File (Adobe PDF File), 545 KB - Multimedia Appendix 3]

Multimedia Appendix 4
Table S3. Perspectives on the ICU-specific virtual reality intervention.

[PDF File (Adobe PDF File), 529 KB - Multimedia Appendix 4]

Multimedia Appendix 5
CONSORT (Consolidated Standards of Reporting Trials) checklist.

[PDF File (Adobe PDF File), 636 KB - Multimedia Appendix 5]

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Abbreviations

EDC: electronic data capture
EQ-5D: European Quality of Life, 5 dimensions
HADS: Hospital Anxiety and Depression Scale
HRQoL: health-related quality of life
ICU: intensive care unit
ICU-VR: intensive care unit–specific virtual reality
IES-R: Impact of Event Scale-Revised
OR: odds ratio
PICS: postintensive care syndrome
PTSD: posttraumatic stress disorder
RT–PCR: reverse transcription–polymerase chain reaction
SF-36: Short-Form 36
VR: virtual reality
