Effect of intensive care unit-specific virtual reality (ICU-VR) to improve psychological well-being in ICU survivors: study protocol for an international, multicentre, randomised controlled trial—the HORIZON-IC study

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ABSTRACT
Introduction A substantial proportion of intensive care unit (ICU) survivors develop psychological impairments after ICU treatment, part of the postintensive care syndrome, resulting in a decreased quality of life. Recent data suggest that an ICU-specific virtual reality intervention (ICU-VR) for post-ICU patients is feasible and safe, improves satisfaction with ICU aftercare, and might improve psychological sequelae. In the present trial, we firstly aim to determine whether ICU-VR is effective in mitigating post-traumatic stress disorder (PTSD)-related symptoms and secondly to determine the optimal timing for initiation with ICU-VR.

Methods and analysis This international, multicentre, randomised controlled trial will be conducted in 10 hospitals. Between December 2021 and April 2023, we aim to include 300 patients who have been admitted to the ICU ≥72 hours and were mechanically ventilated ≥24 hours. Patients will be followed for 12 consecutive months. Patients will be randomised in a 1:1:1 ratio to the early ICU-VR group, the late ICU-VR group, or the usual care group. All patients will receive usual care, including a mandatory ICU follow-up clinic visit 3 months after ICU discharge. Patients in the early ICU-VR group will receive ICU-VR within 2 weeks after ICU discharge. Patients in the late ICU-VR group will receive ICU-VR during the post-ICU follow-up visit. The primary objective is to assess the effect of ICU-VR on PTSD-related symptoms. Secondary objectives are to determine optimal timing for ICU-VR, to assess the effects on anxiety-related and depression-related symptoms and health-related quality of life, and to assess patient satisfaction with ICU aftercare and perspectives on ICU-VR.

Ethics and dissemination The Medical Ethics Committee United, Nieuwegein, the Netherlands, approved this study and local approval was obtained from each participating centre (NL78555.100.21). Our findings will be disseminated by presentation of the results at (inter) national conferences and publication in scientific, peer-reviewed journals.

Trial registration number NL9812.

INTRODUCTION
Because of improved survival after intensive care unit (ICU) treatment, a new challenge arises.1 2 A substantial proportion of ICU survivors suffers from psychological impairments, such as post-traumatic stress disorder...
(PTSD), anxiety, and depression. Along with cognitive and physical impairments, these sequelae are referred to as the post-intensive care syndrome (PICS). PICS is common, can last for years after ICU discharge, and has a profound impact on daily functioning and quality of life.

Prevention and treatment of PICS have been recognized as a fundamental part of ICU care by the critical care community and recently it was demonstrated that the psychological component of PICS is the most important determinant of a decreased health-related quality of life (HRQoL) and impedes patients’ ability to rehabilitate. Although several interventions have been explored, such as keeping ICU diaries, organizing ICU follow-up clinics, and offering psychosocial support, studies on their effectiveness in terms of psychological distress or quality of life have yielded unsatisfactory and ambiguous results. As such, evidence-based interventions to improve psychological recovery and HRQoL are lacking.

Post-ICU psychological impairments may be caused by amnesia during the early period of critical illness in combination with sensory overload and sensory deprivation. Amnesia can lead to loss of factual recall of their ICU stay and patients can instead create delusional and frightening memories. Moreover, the typical ICU environment is characterized by unpatterned exposure and frequent sensory input such as light, noise and tracheal tube aspiration. The exposure to these extremes initiates the development of PTSD and anxiety. We hypothesised that exposure to the factual ICU environment, and additionally receiving ICU-related treatment information, could enhance ICU treatment understanding and subsequently could decrease delusional memories and psychological impairments.

Virtual reality (VR) allows users to fully immerse within a computer-generated three-dimensional environment. In psychiatry, exposure therapy using VR has been proven effective for the treatment of PTSD and anxiety, and thereby it addresses limitations of imaginal exposure. VR can effectively and easily be used to deliver structured and uniform information to patients. VR could, thus, be a valuable adjunct to safely inform and expose post-ICU patients to the environment traumatising them and could enhance psychological recovery. In the current study, our primary aim is to assess the effect of an ICU-specific VR intervention for post-ICU patients (ICU-VR) on PTSD-related symptoms. Second, we want to determine optimal timing for initiation with ICU-VR, to assess the effects of ICU-VR on anxiety-related and depression-related symptoms and HRQoL, and to assess patient satisfaction with ICU aftercare and their perspectives on ICU-VR.

METHODS AND ANALYSIS
Study design and setting
A multicentre, randomised controlled trial will be conducted in ICUs of 10 hospitals in the Netherlands (Erasmus Medical Centre (university hospital), Franciscus Gasthuis & Vlietland hospital, Maasstad hospital, Ikazia hospital, IJsseland hospital, Groene Hart hospital, Van Weel-Bethesda hospital, Haaglanden Medical Centre and the Albert Schweitzer hospital) and Belgium (Cliniques universitaires de Bruxelles—Hôpital Erasme, Bruxelles) (table 1). The Medical Ethics Committee United (MEC-U), Nieuwegein, the Netherlands, approved this study (NL78555.100.21, approved 25 October 2021), and local approval was obtained from the institutional ethics review boards of each participating hospital. Inclusion will be conducted from December 2021 to October 2022, and patients will be followed for 12 months after ICU discharge. Any modifications to the study protocol, which may affect the conduct of the study or patient safety, including changes of the study objectives, study design, study population, sample size, study procedures or significant administrative aspects, will be sent for approval to the MEC-U and the institutional ethics review boards. Health authorities will be informed in accordance with local regulations.

Study participants
We aim to include at least 300 patients. Patients admitted to the ICU for ≥72 hours, during which mechanically ventilated ≥24 hours, older than 17 years of age, and able to understand the Dutch language are eligible for inclusion. Patients admitted to the ICU with primary neurological impairment, a life expectancy <48 hours, or receiving palliative care, with documented active, established psychiatric disorders, a decreased cognitive function during inclusion (a telephone interview for cognitive status (TICS) score ≤27), with a new or active delirium during inclusion (defined as mentioning of a delirium in the daily status report of the treating physician or new administration of haloperidol), or without a formal home address will be excluded. Because the TICS is part of the study procedures, this will be assessed after inclusion and written informed consent. Patients with a TICS score ≤27 will be excluded after inclusion.

Randomisation and masking
Patients will be randomised in a 1:1:1 ratio to either the early ICU-VR group, the late ICU-VR group, or the usual care group. Randomisation will be according to a 1:1:1 ratio, stratified for study site, using a centralised internet-based randomisation procedure (Castor EDC, Amsterdam, The Netherlands). Due to the nature of the intervention, blinding of patients is not possible. Randomisation allocation will be coded in analysis with ‘0’ and ‘1’, and the analyst will as such be unaware of the randomisation allocation.

Intervention
ICU-VR for post-ICU patients is based on a uniform script that is designed by an interdisciplinary team and based on the several focus group meetings of this team. The content of the script is extensively described elsewhere and the
| Study site | Type of hospital | Type of ICU | Number of ICU beds |
|------------|-----------------|-------------|-------------------|
| Erasmus Medical Centre, Rotterdam, The Netherlands | Academic hospital | Mixed medical, surgical and cardiac ICU | 56 |
| Francisca Gasthuis & Vlietland, Rotterdam, The Netherlands | Community, teaching hospital | Mixed medical and surgical ICU | 19 |
| Massstad Hospital, Rotterdam, The Netherlands | Community hospital | Mixed medical and surgical ICU with an burn expertise centre | 25 |
| Ikazia Hospital, Rotterdam, The Netherlands | Community hospital | Mixed medical and surgical ICU | 12 |
| Lusseiland Hospital, Capelle a/d Ijssel, The Netherlands | Community, teaching hospital | Mixed medical and surgical ICU | 8 |
| Groene Hart Hospital, Gouda, The Netherlands | Community hospital | Mixed medical and surgical ICU | 12 |
| Van Weel-Bethesda Hospital, Dikrslad, The Netherlands | Community, teaching hospital | Mixed medical and surgical ICU | 6 |
| Haaglanden Medical Centre, The Hague, The Netherlands | Community, teaching hospital | Mixed medical and surgical ICU | 22 |
| Albert Schweitzer Hospital, Dordrecht, The Netherlands | Community, teaching hospital | Mixed medical and surgical ICU | 16 |
| Cliniques universitaires de Bruxelles – Hôpital Erasme, Bruxelles, Belgium | Academic hospital | Mixed medical and surgical ICU | 36 |

ICU, intensive care unit.
We also have written a movie directors script to produce an uniform ICU-VR film in each participating centre. The ICU-VR film was produced for each centre, that is, hospital specific, to optimise immersiveness and to deliver relevant and truthful information regarding ICU stay and ICU treatment. The point of view for the camera is the field of vision of the mock patient lying in an ICU bed. ICU-VR will be watched using head-mounted display VR (Pico G2 VR All-In-One Headset) and a headset.

**Study procedures**

An oversight of the study procedures is presented in [figure 1](#). Patients who are eligible for inclusion will be approached by an investigator of the research team or by a dedicated research nurse within 7 days after ICU discharge. A translation of the information for patients and the informed consent form are found in online supplemental data file 2.

After obtaining informed consent and completing the TICS assessment, patients will receive the first set of questionnaires (T0), consisting of a self-composed questionnaire regarding demographics and their history of mental health, the Impact of Event Scale-Revised (IES-R), the Hospital Anxiety and Depression Scale (HADS), the European Quality of life 5 dimensions (EQ-5D) and the short-form 36 (SF-36) (table 2). Patients are asked to fill in the HADS, EQ-5D, and SF-36 questionnaire both retrospectively and prospectively to obtain a baseline and over
time measure of patient anxiety and depression levels and quality of life. Patients randomised to the early ICU-VR group will receive ICU-VR between day 8 and day 15 after ICU discharge for a maximum of three times, unless the patient is discharged from the hospital ward sooner. The number of times ICU-VR is offered and accepted will be logged. Between 3 and 6 months after ICU discharge, all patients will visit the post-ICU follow-up clinic of the concurrent hospital. During this post-ICU follow-up visit, patients have a consultation with a dedicated ICU nurse and an intensivist. Patients randomised to the late ICU-VR group will receive ICU-VR once during their concurrent post-ICU follow-up clinic visit.

All patients will receive follow-up questionnaires at 1 month (T1), 3 months (T2), 6 months (T3) and 12 months (T4) after ICU discharge (Table 2).

**Outcomes and measurements**

The primary outcome is the effect of ICU-VR on the severity of PTSD-related symptoms at 6 months after ICU discharge.

The severity of PTSD-related symptoms will be expressed as the sum score of the IES-R and an IES-R sum score ≥24 will be considered as clinically relevant PTSD. The IES-R comprises 22 items, assessing subjective distress caused by traumatic events, and has been used commonly in survivors of critical illness. The IES-R yields a total score (ranging from 0 to 88; higher scores indicate more severe symptoms) and subscale scores can be calculated for symptoms of intrusion, avoidance and hyper arousal.

Secondary outcomes are the effects of ICU-VR on the severity and prevalence of PTSD-related, anxiety-related and depression-related symptoms and on HRQoL throughout follow-up, the patient satisfaction with ICU aftercare, and patient perspectives on ICU-VR.

The severity of anxiety-related and depression-related symptoms will be expressed as the HADS anxiety and depression scores, and a HADS anxiety or depression score ≥8 will be considered as clinically relevant anxiety and depression, respectively. The HADS comprises of 14 items and is commonly used to determine the levels of anxiety and depression. Seven of the items relate to anxiety and seven relate to depression.

HRQoL will be expressed as the overall HRQoL, which implies the time trade-off (TTO) score of the 5-level EQ-5D, and the mental HRQoL, which implies the mental component score of the SF-36.

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HRQoL will be expressed as the overall HRQoL, which implies the time trade-off (TTO) score of the 5-level EQ-5D, and the mental HRQoL, which implies the mental component score of the SF-36. The EQ-5D measures HRQoL in five dimensions, that is, mobility, self-care, usual activities, pain/discomfort and anxiety/depression. By giving a certain weight to each answer option, the country-specific TTO score can be calculated, ranging from −0.446 (worst quality of life) to 1.000 (best quality of life).

All patients will receive follow-up questionnaires at 1 month (T1), 3 months (T2), 6 months (T3) and 12 months (T4) after ICU discharge (Table 2).

### Table 2 Questionnaires per follow-up time point

| Questionnaire                | T0, inclusion | T1, 1 month after ICU discharge | T2, 3 months after ICU discharge | T3, 6 months after ICU discharge | T4, 12 months after ICU discharge |
|-----------------------------|---------------|---------------------------------|----------------------------------|---------------------------------|----------------------------------|
| Demographics                | X             | X                               | X                                | X                               | X                                |
| History of mental illness   | X             | X                               | X                                | X                               | X                                |
| IES-R (Post-Traumatic Stress Disorder) | X             | X                               | X                                | X                               | X                                |
| HADS (Anxiety and Depression) | X (retrospectively and prospectively) | X                               | X                                | X                               | X                                |
| SF-36 Quality of Life      | X (retrospectively and prospectively) | X                               | X                                | X                               | X                                |
| EQ-5D Quality of life      | X (retrospectively and prospectively) | X                               | X                                | X                               | X                                |
| Satisfaction with ICU care | X             | X                               | X                                | X                               | X                                |
| Perspectives on ICU-VR      | X (early ICU-VR) | X                               | X                                | X                               | X                                |
| Visit to healthcare professionals | X             | X                               | X                                | X                               | X                                |

EQ-5D, 5-Level European Quality of Life Questionnaire; HADS, Hospital Anxiety and Depression Scale; ICU, intensive care unit; ICU-VR, intensive care unit-specific virtual reality; IES-R, Impact of Event Scale Revised; SF-36, Short-Form 36.
equal weight. The eight sections are vitality, physical functioning, bodily pain, general health perception, physical role functioning, emotional role functioning, social role functioning and mental health.\textsuperscript{38–40} In addition, a mental and physical component scale, the MSC-36 and PCS-36, respectively, can be calculated as a reflection of physical and mental health.\textsuperscript{38–40}

Patient satisfaction with ICU aftercare will be assessed using a novel questionnaire, based on the Patient Satisfaction Questionnaire and Family Satisfaction with ICU Care tools, altered to the needs of this study.\textsuperscript{41, 42} Additional novel items were added to evaluate patient perspectives on the ICU-VR intervention.

We also explore feasibility and safety outcomes, and the cost-benefit ratio of ICU-VR. Feasibility will be expressed as the number of sessions patients in the early ICU-VR group will receive. Safety will be expressed as the number of ICU-VR sessions requiring interruption or termination due to side effects in terms of cybersickness, mainly experienced as nausea.\textsuperscript{36–37} For the cost-benefit ratio, costs will be expressed as, among others, development costs for ICU-VR, employment costs of ICU nurses offering the intervention, and the employment and organisational costs of the ICU follow-up clinic, and benefits will be expressed as the gain in quality-adjusted life years determined as the EQ-5D TTO score.

Demographics, such as age, gender, body weight, length, pre-existing comorbidities, previous ICU admissions and ICU readmissions, treatment-related characteristics, such as type of admission, ICU and hospital length of stay, mechanical ventilation-related characteristics, episodes of sedative coma and delirium during ICU treatment, assessed using the Richmond Agitation Sedation Scale and the Confusion Assessment Method for the ICU (CAM-ICU) scale, respectively, use of renal replacement therapy, infections and illness severity scores during ICU treatment, and 3-month, 6-month and 12-month mortality will be assessed using electronic patient records.\textsuperscript{30, 41} Additionally, patients will be asked about their educational level, employment status prior to and after ICU treatment, financial decrease after ICU treatment, consultations with healthcare professionals, and their history of mental health in follow-up questionnaires.

Data management
Data will be uploaded, stored, and maintained using the electronic data capture (EDC) system of Castor (Castor EDC, www.castoredc.com, Amsterdam, the Netherlands). The study team will be responsible for data entry and quality control activities. Data will be checked by at least two persons from the study team and will be stored for at least 15 years on either the Castor EDC server or as a hardcopy in the ICU of the participating hospitals. Questionnaires will be sent digitally using Castor EDC or via hardcopy via postal mail whenever requested.

To maintain anonymity, data will be coded with a number and this number will be the only reference to patient identification. The principal investigator is the only one in possession of the translation key, making it impossible to link data to the patient.

Sample size calculation
Based on two previous studies yielding an ICU-VR Cohen’s \(d\) effect estimate of 0.56 (late intervention) to 0.88 (early intervention), the power calculation of the current study is based on a Cohen’s \(d\) of 0.56.\textsuperscript{36–45} We performed a G*Power analysis based on the Wilcoxon Mann-Whitney test, with no expectation about the underlying distribution of the outcome (parental distribution: ‘min ARE’). Using a two-sided alpha of 0.05, a power of 0.80 and a 1:1 allocation ratio, this resulted in a required sample size per group of 60 patients.\textsuperscript{46} We will use this required sample size for all three groups resulting in a total sample size of 180 patients. We anticipated a loss to follow-up rate of 40\% for which we will anticipate in the current trial. We, therefore, aim to include a total of \((3\times60/0.60 =) 300\) patients, with 100 patients per group.

Statistical analysis
All continuous data will be presented as medians (95\% range). Categorical variables will be presented as absolute and relative frequency. Baseline demographics, treatment-related characteristics, and patient perspectives on ICU-VR will be summarised using descriptive statistics. Outcomes of mixed effects linear and logistic regression models will be presented as the coefficient of the model, which implies the estimated mean difference between groups, including its 95\% CI, as the log of coefficient of the model, that is, the OR, including its 95\% CI, respectively.

To analyse the effects of ICU-VR on the severity of PTSD-related, anxiety-related, and depression-related symptoms, on HRQoL, and on the prevalence of clinically relevant PTSD, anxiety, and depression at each follow-up time point, we will use mixed effects linear (for continuous outcomes) or logistic (for categorical outcomes) regression models. In these, the outcome at each follow-up time point will serve as dependent variable, the randomisation group, the retrospectively assessed pre-existent score/prevalence of the outcome (parental distribution: ‘min ARE’). In the outcome at each follow-up time point will serve as dependent variable, the randomisation group, the retrospectively assessed pre-existent score/prevalence of the outcome of interest and a random intercept and/or slope for each study site will be used. The effect of ICU-VR on the course of (1) the severity of PTSD, anxiety-related and depression-related symptoms, (2) HRQoL and (3) the prevalence of clinically relevant PTSD, anxiety and depression throughout follow-up will be analysed using mixed effects linear (for continuous outcomes) and logistic (for categorical outcomes) regression models, in which the outcome/prevalence of interest of all follow-up time points will be used as dependent variable, the randomisation allocation, time, the retrospectively assessed pre-existent score/prevalence of interest will serve as independent variables and a random intercept and/or slope for each study site will be used.

To determine when ICU-VR is most effective, that is, early versus late, differences in psychological distress and HRQoL between the early ICU-VR group and late
ICU-VR group at 6 and 12 months will be assessed. We will analyse these using mixed effects linear and logistic regression models. In these models, the score/prevalence of interest at either 6 months or 12 months after ICU discharge will be used as dependent variable, the randomisation allocation, the retrospectively assessed pre-existent score/prevalence of the outcome of interest will serve as independent variables and a random intercept and/or slope for each study site will be used. Differences in the course of the severity and prevalence of psychological distress and HRQoL between 6 and 12 months will be assessed using mixed effects linear and logistic regression models, in which the outcomes at 6 and 12 months will simultaneously be used as dependent variable, and time after discharge in months, randomisation allocation (early ICU-VR/late ICU-VR), the interaction between randomisation and time (randomisation×time), the pre-existent score of the outcome of interest will serve as independent variables and a random intercept and/or slope for each patient and each study site will be used.

We will analyse differences in the subscales of the SF-36, patient resumption to work, experienced financial decline and consultation with healthcare professionals using the abovementioned manners.

The main analysis will be an intention-to-treat analysis, in which all included patients will be included. Second, we will perform a per-protocol analysis, in which patients are included if (1) they are randomised to the control group, (2) they are randomised to the early ICU-VR group and received ICU-VR three times in the hospital ward and (3) they are randomised in the late ICU-VR group and received ICU-VR once during the ICU follow-up clinic visit. Thereafter, we will conduct a complete case analysis, in which all patients who have completed all assessment are included.

We will conduct the subanalyses in (1) patients who have been mechanically ventilated ≥72 hours, (2) patients who have been mechanically ventilated >7 days, (3) patients who have been treated in the ICU for >7 days, (4) patients who have been treated in the ICU for >14 days, (5) patients who had a delirium, as documented in the healthcare record, (6) per study site (study sites with <10 inclusions will be combined), (7) sepsis patients, to compare these results with our previously conducted pilot study.

If the loss to follow-up at 6 months after ICU discharge will be higher than anticipated, we will impute missing data using both the last observation carried forward method and multiple imputation according to the Markov-chain Monte-Carlo.25–27

All data will be gathered using Castor EDC (Castor EDC, Amsterdam, the Netherlands). All analyses will be performed using SPSS (IBM SPSS Statistics for Windows, V.27.0; IBM Corporation, Armonk, New York) and R for Statistics (R Foundation for Statistical Computing, Vienna, Austria, 2015). A p value of ≤0.05 will be considered statistically significant.

**Ethics and dissemination**

This study will be conducted in accordance with the principles of the declaration of Helsinki (version October 2013; www.wma.net) and in accordance with the Medical Research involving human subjects act (WMO) and other guidelines, regulations and acts. We received approval from the MEC-U, Nieuwegein, and local approval has been obtained from the institutional ethic review boards of each participating hospital. If deviation from the protocol is necessary, it will not be implemented without the prior review and approval of the MEC-U and each participating hospital’s institutional ethic review board. Signed informed consent will be obtained from all patients prior to any study procedure. Previous research demonstrated that (ICU-)VR is safe, feasible and well accepted.25–27 Informed consent forms will be kept in a locked cabinet in a limited access room in the ICU of the participating study sites. Data will be archived for 15 years. The handling of personal data complies with the Dutch Law. On completion of the study, its findings will be published in peer-reviewed journals and presented at national and international scientific conferences to publicise the research to healthcare professionals, health services authorities and the public. A summary of results will be made available to the study patients if requested.

**Patient and public involvement statement**

Former ICU patients were involved in the development of the ICU-VR intervention. Patients and/or the public were not involved in the design, or conduct, or reporting, or dissemination plans of the current research.

**Contributors**

JHV, JvB, E-JW, MVM, DG and MEvG conceived the study and initiated the study design. MEV is the coordinating investigator and grant holder. DG is the principal investigator. TK provided statistical expertise in the clinical trial design, and JHV and TK wrote the statistical analysis plan. MVM provided expertise in the field of psychology, and JV and MVM determined what questionnaires are used. JvB, E-JW, FT, AFCS, JHE, AMTJR, AD and SA are the local principal investigators at each study site. All authors contributed to the refinement of the study protocol and approved the study protocol. JHV and AMTJR wrote the first draft of the manuscript, JvB, E-JW, TK, EK, MV, DG, MEV helped to further draft the manuscript. JHV and ALJ will be responsible for data collection. All authors approved the final version of the manuscript.

**Funding**

This study was supported by DSW (for the HORIZON-IC project; no grant number available), Stichting Theia (grant number: 2020286), Stichting SGS (grant number: 2020355) and BeterKeten (for the HORIZON-IC project; no grant number available).
available. The funding sources had no role in the design of the study and collection, analysis, and interpretation of data nor in writing the manuscript.

Competing interests DG has received speaker fees and travel expenses from Dräger, GE Healthcare (medical advisory board 2009–12), Maquet, and Novalung (medical advisory board 2015–18). TK has received speaker fees from Guided, IBISA, Merck, Berlin Chemie, and Goodlife Healthcare. All other authors declare no competing interests.

Patient and public involvement Patients and/or the public were not involved in the design, conduct, or reporting, or dissemination plans of the current research.

Patient consent for publication Not applicable.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Not applicable.

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