BMJ Open Improving physical function of patients following intensive care unit admission (EMPRESS): protocol of a randomised controlled feasibility trial

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ABSTRACT

Introduction Physical rehabilitation delivered early following admission to the intensive care unit (ICU) has the potential to improve short-term and long-term outcomes. The use of supine cycling together with other rehabilitation techniques has potential as a method of introducing rehabilitation earlier in the patient journey. The aim of the study is to determine the feasibility of delivering the designed protocol of a randomised clinical trial comparing a protocolised early rehabilitation programme including cycling with usual care. This feasibility study will inform a larger multicentre study.

Methods and analysis 90 acute care medical patients from two mixed medical-surgical ICUs will be recruited. We will include ventilated patients within 72 hours of initiation of mechanical ventilation and expected to be ventilated a further 48 hours or more. Patients will receive usual care or usual care plus two 30 min rehabilitation sessions 5 days/week.

Feasibility outcomes are (1) recruitment of one to two patients per month per site; (2) protocol fidelity with >75% of patients commencing interventions within 72 hours of mechanical ventilation, with >70% interventions delivered; and (3) blinded outcome measures recorded at three time points in >80% of patients. Secondary outcomes are (1) strength and function, the Physical Function ICU Test-scored measured on ICU discharge; (2) hospital length of stay; and (3) mental health and physical ability at 3 months using the WHO Disability Assessment Schedule 2. An economic analysis using hospital health services data reported with an embedded health economic study will collect and assess economic and quality of life data including the Hospital Anxiety and Depression Scales core, the Eurogol-5 Dimension-5 Level and the Impact of Event

Ethics and dissemination The study has ethical approval from the South Central Hampshire A Research Ethics Committee (19/SC/0016). All amendments will be approved by this committee. An independent trial monitoring committee is overseeing the study. Results will be made available to critical care survivors, their caregivers, the critical care societies and other researchers.

Strengths and limitations of this study

- ► Will investigate the implementation of a protocolised early rehabilitation intervention that is usual care in one NHS/university teaching institution, into other NHS institutions with different organisational
- The defined cohort has been demonstrated to benefit from this type of rehabilitation in alternative healthcare systems.
- Results will inform the design of a multicentre randomised controlled trial.
- This study is not designed to assess the effectiveness of the intervention.
- Inability to blind the intervention to patients, physiotherapist and clinicians involved in the delivery of the intervention.

Trial registration number NCT03771014.

INTRODUCTION

In 2018/2019, there were over 290000 admissions to adult intensive care units (ICUs) in the UK.1 Treatment advances have reduced mortality associated with critical illness²; however, survival does not represent the end of the story. ⁴ A complex interplay between baseline health status, acute disease and the traumatic effects of intensive care treatment is associated with long-term physical, psychological and social hardship.^{5–10} Patients discharged from the ICU have higher mortality, higher health service costs and a reduction in employment status compared with hospitalised patients not requiring ICU.811

ICU-acquired weakness is characterised by rapid muscle wasting, polyneuropathy and bone demineralisation, causing pain, weakness and impaired physical function. 12-14 Contributing factors are multifactorial,



although immobility due to the sedation required for tolerance of ventilation plays an important role. 15 16 Early mobilisation may mitigate these effects. 17-19 In 2009, Schweickert et al reported that patients who underwent early physical therapy (within 1.5 days of mechanical ventilation) had greater functional independence at hospital discharge than patients who had usual care physical therapy. A recent randomised controlled trial (RCT) on the impact of a progressive ICU mobility programme reported improved functional status at ICU discharge.²¹ Meta-analyses and systematic reviews report that early mobilisation of ICU patients may reduce duration of mechanical ventilation and improve short-term physical outcomes, 22-24 but mobilisation can be difficult to implement during a patient's stay in the ICU. Moreover, studies which used delayed rehabilitation, often more than a week after ICU admission, 25-27 have not replicated these outcomes. 28-34 Barriers to early mobilisation include heavy sedation, patient's illness, lack of resources and/or clinician buy-in. 35-38 In-bed cycle ergometry can provide passive activity in heavily sedated patients who are receiving vasopressors 39 40 with minimal physiological demand 40 41 and can be transitioned to active cycling as the patient's condition improves.²³ 42-44

We implemented cycle ergometry as part of an early protocolised rehabilitation quality improvement programme with physiotherapy technicians supporting the additional workload. Like other investigators, we reported reduced number of ventilator days and ICU length of stay. 21 46-49

The primary aim of this study was to evaluate the feasibility of an RCT investigating the effect of early protocolised rehabilitation versus usual physiotherapy care in ICU patients. Results will inform a prospective fully powered multicentre RCT. This protocol is reported according to Standard protocol items for clinical trials (SPIRIT 2013 Statement)⁵⁰ and Template for Intervention Description and Replication⁵¹ guidelines.

Aim

The aim of this study was to determine the feasibility of delivering study procedures comparing an early protocolised mobilisation programme that includes cycling with usual care.

Objectives

Feasibility will be determined by measures of the recruitment process, intervention fidelity and outcome measurement completeness, specifically, (1) study accrual rates: a minimum of 30% of eligible patients or one to two patients per site per month are enrolled; (2) protocol adherence: 75% of patients commencing intervention within 72 hours of ICU admission, with a minimum of 70% of planned interventions delivered; and (3) blinded outcome assessment: functional assessment performed at three time points in 80% of survivors. The results will inform a larger fully powered RCT.

METHODS AND ANALYSIS

Study design

This is a two-centre feasibility study using a two-arm RCT, randomised 1:1, with blinded outcome assessments at ICU discharge, hospital discharge and 3-month follow-up. Patients will be recruited from two general ICUs, located in the south of the UK. They will not be recruited from our ICU on account that the intervention is now standard practice at this site. Prior to each site opening to recruitment, an audit of current physiotherapy practice will be undertaken over a 4-week period to evaluate what constitutes 'usual care' in each institution.

Participants

Ninety patients will be recruited. Eligible patients will be over 42 years old and will have an acute/unplanned medical admission to the ICU. They will be functionally independent prior to ICU admission (Barthel Index>80), in the hospital for <5 days prior to intubation and

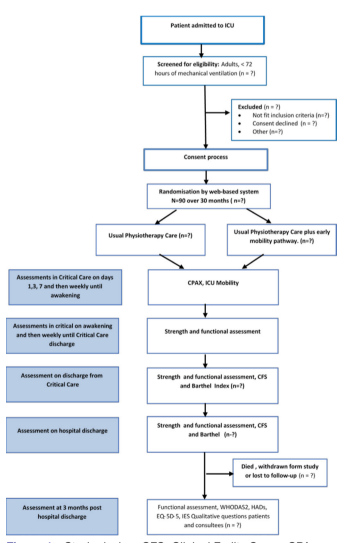


Figure 1 Study design. CFS, Clinical Frailty Score; CPAx, Chelsea Critical Care Physical Assessment Tool; EQ-5D-5L, Euroqol-5 Dimension-5 Level; HAD, ICU, intensive care unit; IES, Impact of Event Score; WHODAS 2.0, WHO Disability Assessment Schedule 2.

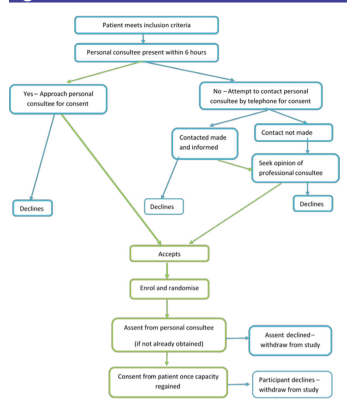


Figure 2 Consent pathway.

ventilation, intubated and ventilated for <72 hours, and expected to remain ventilated for a further 48 hours. Patients will be excluded if they are in the hospital for 5 days or more prior to ICU admission, have acute brain or spinal cord injury, known or suspected neurological/ muscular impairment, condition limiting use of cycle ergometry (eg, lower limb fracture/amputation), not expected to survive >48 hours decided by consulting an intensivist and persistent therapy exemptions in the first 3 days of mechanical ventilation. Figure 1 presents the planned flow of patients through the study.

Recruitment, consent and randomisation

The study team will screen all patients for eligibility. Recruitment began in June 2019 (and was temporarily suspended in March 2020 due to the COVID-19 pandemic). It is anticipated recruitment will continue until mid 2022. The majority of patients will have diminished capacity when first eligible; therefore, the consent process is multilayered and designed in accordance with the Mental Capacity Act (MCA) 2005⁵² (figure 2):

Patient informed consent: wherever possible, informed consent will be directly sought from the patient (see online supplemental files 1 and 2).

Personal consultee informed assent: if the patient is unable to provide consent, informed assent will be sought from the patient's personal consultee, within 6 hours of confirmation of eligibility. If the personal consultee is not available in person, attempts will be made to contact them by telephone. They will be asked to provide

written assent, at the earliest possible convenience (see online supplemental files 3 and 4).

Professional consultee informed assent: where both patient and personal consultee are not available to approve enrolment within 6 hours of confirmation of eligibility, assent will be sought from a professional consultee in accordance with the MCA. The professional consultee will be a consultant medical practitioner, independent from the study. The patient's personal consultee will be consulted at the earliest possible opportunity and assent will be requested to continue in the study.

In all cases, once the patient has regained capacity, they will be informed of the study and consent continuation will be sought. Following consent or assent, patients will be registered on a bespoke electronic data collection tool (ALEA) and randomly assigned to the protocolised early rehabilitation or usual care.

Staff training/site set-up

Participating sites will employ the equivalent of a fulltime therapy technician to deliver the study intervention, under the supervision of a senior critical care therapist. Both senior critical care therapists and therapy technicians will complete a training package delivered by the primary institution (University Hospital Southampton NHS Foundation Trust), where early rehabilitation with cycling is well established and embedded in usual care. This package includes seminars on the delivery of the protocolised early rehabilitation, use of the bespoke electronic database and 5 days of clinical shadowing.

Interventions

All patients will receive usual medical, nursing and physiotherapy care while in intensive care. Each bedside nurse will be asked at the start of the shift if they have been involved caring for a patient in the intervention arm of the study. The ICU physiotherapy team, who are not involved with the delivery of the study delivery, will deliver all usual physiotherapy interventions in both groups. The physiotherapist delivering usual care will be asked to verify if they have delivered any of the study interventions. In the intervention arm, the protocolised physiotherapy programme will commence within 72 hours of ICU admission or as soon as possible thereafter and will continue for 28 days or until ICU discharge, whichever occurs first. Patients' respiratory support can range from full mandatory ventilation through to oxygen supplementation with no mechanical support following extubation. Sedation is targeted throughout the time that the patient is intubated, and the ventilation mode is adjusted to patients' needs, compliance and comfort at discretion at the start of each physiotherapy intervention, the participants' level of sedation will be assessed using the Richmond Agitation-Sedation Scale (RASS)^{53 54} and the Confusion Assessment Method for ICU⁵⁵will be undertaken. RASS will be targeted to a RASS between -1 and +1 by the bedside nurse. After 28 days of ICU admission, all patients will receive usual care physiotherapy interventions.

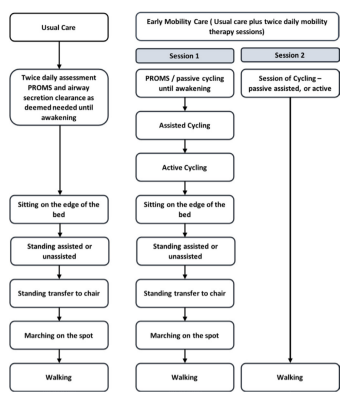


Figure 3 Study intervention pathway. (PROM = Passive Range of Movement)

Group 1: usual care control group

In this pragmatic study, physiotherapy interventions will be guided by individual assessment and start in accordance with the usual care pathway within each institution. The focus of each session may be respiratory support, mobilisation or a combination of both. Interventions delivered will be determined by the physiotherapist in conjunction with the attending physician. Interventions include, where appropriate, passive or active range of movement, positioning and respiratory physiotherapy, and when able, sitting on the edge of the bed, standing (assisted or unassisted), standing to transfer to chair, marching on the spot and walking (figure 3). Usual interventions may occur at any time of day.

Group 2: Protocolised rehabilitation pathway

Patients will have usual care physiotherapy, in addition to the two protocolised intervention within 72 hours of ICU admission or as soon as possible thereafter. Patients will be screened for safety criteria to withhold the intervention prior to each planned intervention session (table 1).

Those meeting criteria to withhold interventions will have issues addressed and reassessed for interventions 2 hours later. The two additional rehabilitation sessions will be delivered by the research physiotherapy staff including a therapy technician. This will comprise two mobility sessions the modality of the first, chosen at the discretion of the physiotherapist. The second session will be 30 min of supine cycling delivered in the afternoon.

The first rehabilitation intervention each day will be delivered in the morning. Planned interventions include passive or active range of movements, passive cycling, active cycling, in-bed exercises, sitting, mobilisation out of bed and walking. Daily assessment of the patient will be made to ensure the highest level of activity possible is provided for each individual patient given safety considerations and capability of the patient.

The second session will be cycling based. An in-bed supine cycle ergometer (MotoMed Letto 2) will be used

Table 1 Safety criteria for delivery of physical therapy interventions					
	Criteria to commence physiotherapy	Criteria to stop/withhold physiotherapy intervention			
Blood pressure	Mean Arterial Blood Pressure (MAP) 60–100 mm Hg, no change in vasopressor dose requirement for preceding 2 hours	Catecholamine-resistant hypotension with MAP <60 mm Hg			
Heart rate	Between 40 and 140 beats/min	<50 or >140 beats/min			
Respiratory rate	Sustained <40 breaths/min	Sustained >40 breaths/min			
Temperature		>40°C			
Oxygen requirement	If Fraction inspired oxygen (FiO ₂)>0.8 for passive exercise only				
	FiO ₂ <0.8 and (Positive End Expiratory Pressure) PEEP <15 cmH ₂ O				
Desaturation		Sats fall <85% for>1 min			
Other		 Fall. Unplanned extubation. Acute bleeding. New-onset arrhythmia. Signs/symptoms of acute myocardial ischaemia. Patient pain/distress. Clinical team decides therapy intervention not appropriate. Refusal by patient or representative. 			



to engage the participant in passive, assisted or active cycling, or a combination, depending on the degree of patient cooperation (figure 3). The aim was for the patient to have 30 min of cycling per day, following a standardised cycling programme. If cycling is in passive mode, patients will commence cycling at 5 revolutions per minute (RPM), building up to 20 RPM over 5 min and continue this for 20 min before 5 min 5 RPM cool down. In the assisted or active mode, after the 5 min warm-up, cycling will continue for 20 min at patient-selected RPM followed by a 5 min cool-down at 5 RPM. In-bed cycling sessions will stop when the patient is deemed to be able to stand and transfer from bed to chair for both mobility sessions for two consecutive days. If patients are considered unable to have concurrent mobility therapy and respiratory weaning, mobility therapy will take priority, in agreement with the senior clinical team. Individual participants will receive the trial intervention on 5 days/week (Monday-Friday) for the duration of their ICU stay or a maximum of 28 days, whichever comes first. Patients will be monitored for cardiovascular and respiratory stability and safety of indwelling lines, tubes and catheters with predetermined criteria for termination of any session (table 1). Deviations from the planned protocol will be reported to determine potential barriers to implementation. Patients will be able to decline any intervention or

outcome assessment at any time without compromise to their care.

Primary outcome: feasibility to deliver the protocol as

Feasibility will be determined by measures of the recruitment process, intervention fidelity and outcome measurement completeness, specifically,

- Study accrual rates: a minimum of 30% of eligible patients or one to two patients per site per month are enrolled.
- Protocol adherence: 75% of patients commencing intervention within 72 hours of ICU admission, minimum of 70% of planned interventions delivered.
- Blinded outcome assessment: functional assessment performed at three time points in 80% of survivors by physiotherapists working within the hospital but not within the ICU.

Secondary outcomes

The schedule of outcome assessments is detailed in table 2.

Strength and function

We will measure the Physical Function ICU Test-scored (PFITs) at awakening as described by De Jonghe et al⁶⁶

Table 2 Schedule of asses	sments								
	Randomisation	Day 1	Day 3	Day 7	Awakening	Weekly	ICU discharge	Hospital discharge	3 months posthospital discharge
Demographic data	Х								
Muscle assessment									
Medicial Research Council sum-score (MRCss) ^{60 61}					X	X	X	X	
Grip strength ⁶²					Χ	Χ	Х	Х	
Physical function									
CPAx ⁶³		Χ	Χ	Х	Χ	Χ	Χ		
ICU mobility ⁶⁴		Χ	Χ	Χ	Χ	Χ	Χ		
PFITs ⁵⁹					Χ	Х	Х		
Timed-Up and Go							Χ	Χ	Χ
Clinical Frailty Score ⁶⁹		(X)					Χ	Χ	Χ
Barthel Index		(X)					Χ	Χ	Χ
6 min Walk Test ⁷⁰								Χ	Χ
Health Related Quality of Life (H	IRQL)								
WHODAS 2.0 ⁷¹									Χ
HADS ^{72 73}									Χ
EQ-5D-5L ⁷⁴									Χ
Impact of Event Scale ⁷⁵									Χ
Health Economic Evaluation (CSRI)*									Χ

CPAx, Chelsea Critical Care Physical Assessment Tool; CSRI, Client Service Receipt Inventory; EQ-5D-5L, Eurogol-5 Dimension-5 Level; HADS, Hospital Anxiety and Depression Scale; ICU, intensive care unit; PFITs, Physical Function ICU Test-scored; WHODAS 2.0, WHO Disability Assessment Schedule 2.



then weekly within ICU and on ICU discharge.⁵⁷ PFITs is a reliable and valid four-item scale (arm strength, leg strength, ability to stand and step cadence), with a score range of 0-10 and is responsive to change and predictive of key outcomes.⁵⁸ Medical Research Council Manual Muscle Test Sum Score (MRC-ss)⁵⁹ 60 and handheld dynamometry⁶¹ will be measured on awakening, weekly, on ICU discharge and hospital discharge. Chelsea Critical Care Physical Assessment Tool (CPAx)⁶² and ICU Mobility Scale⁶³ will be assessed three times during the first week within ICU, on awakening, weekly thereafter within the ICU and at ICU discharge. Timed Up and Go, ^{64 65} Clinical Frailty Score (CFS) ^{66–68} and Barthel Index will be assessed at ICU discharge, hospital discharge and 3 months posthospital discharge. Preadmission Barthel Index and CFS will be assessed by proxy on admission from family member or next of kin. Six-minute walk test⁶⁹ will be performed, in accordance with American Thoracic Society guidelines, at hospital discharge and 3 months posthospital discharge.

Health-related quality of life (QOL) outcomes

The following will be measured at 3 months posthospital discharge: WHODAS 2.0, 70 Hospital Anxiety and Depression Scale score, 71 72 Euroqol-5 Dimension-5 Level (EQ-5D-5L), 73 Impact of Event Score 74 and Client Service Receipt Inventory questionnaire, designed for this study to evaluate costs that fall on patients and their carers. Resource use and costs including direct intervention costs of therapists and equipment and general hospital costs (per bed day) will be recorded for each patient.

Health economic substudy

We will also conduct an embedded health economic study to identify and define data collection for a future RCT where a full cost-effectiveness analysis (CEA) can be conducted. Within the feasibility study, we aim to address the following:

- ▶ What the quality of the data and what potential problems are there for reporting QoL (EQ-5D-5L), resource use and costs.
- ► The cost implications of the proposed intervention in terms of impact for the NHS (inpatient stay bed days) and identifying the main cost drivers.
- ► Is the EQ-5D-5L appropriate for use in the future RCT?

The economic outcomes will include secondary care resource use within hospitals during inpatient stay, primary care resource use following discharge up to 3 months and resource use related to providing the intervention. The results will be reported in the form of descriptive statistics and will be used to inform a future CEA within a definitive RCT.

Additional data collection

We will collect baseline data including demographic information, Functional Comorbidity Index, ICU diagnosis, APACHE II score, ventilation duration, ventilator-free

days, ICU and hospital length of stay, within ICU drug history and duration and type of usual care physiotherapy.

Implementation evaluation

We aim to investigate whether the protocolised early rehabilitation programme used in one NHS institution is transferable, as an RCT, into other similar NHS institutions. The design of a future multicentre study will be informed by identified facilitators and barriers to implementation. Implementation assessment will be based on the measures described by Proctor. 75 A cross section of ICU staff and patients will be interviewed and complete questionnaires at trial completion to identify barriers impacting delivery of the study. Understanding of the integration and sustainability of the intervention are necessary to inform the design of a powered RCT. Acceptability will be measured at the beginning and end of the study from investigators and clinical staff by direct discussions and questionnaire. Our experience informs us that the introduction of this intervention is dependent on a cultural change within any unit for a proactive focus on early mobilisation. We aim to explore measures to help optimise implementation. Adoption, feasibility and fidelity measures will be monitored during the study by regular meetings with the investigators. Patient screening logs will identify the number of patients eligible for the study and barriers to enrolment. We will assess the degree to which it is possible to separate the staff caring for the intervention group from those caring for the patients in the control group.

We will report whether trial participation has influenced usual care within the participating units by prestudy and poststudy audits. Participating sites will collect data regarding number and seniority of therapy staff with dedicated time to work within the ICU; delirium and sedation protocols used; time, type and frequency of rehabilitation interventions delivered, who delivers the interventions and reasons why usual care may not be delivered.

The feasibility outcomes described earlier will be used to power a larger RCT.

Data entry and checks

Data will be entered into the secure electronic case report form (ALEA) and data validation will take place according to the procedures set out in the data management plan and data validation plan, both developed apriori. Missing data will be assessed to identify any specific challenges with any items of data collected. Missing data level is expected to be less than 20%. Data loss and mortality will inform number of participants needed to design a larger RCT. As this is a feasibility study data imputation will not be undertaken. Prior to statistical analysis, variables will be checked for missing and impossible and improbable values as defined by clinical opinion. Questions regarding the data will be directed to the data manager.



Sample size calculation

This is a feasibility study, the results of which will be used to power a definitive study if appropriate; as such, no formal sample size calculation for effectiveness of the intervention has been undertaken. Ninety patients will be recruited, aiming for 30-45 participants at each site. We anticipate a 30% in hospital mortality /loss to follow-up with an estimate of 60 patients completing the study. This sample size of 90 will allow the estimate of recruitment rate to be made with a 95% CI of ±5.2% if the rate is observed to be around 30%, and with a CI of ±7.3% if the recruitment rate is observed to be around 50%. In addition, the sample of 90 recruited patients will allow the estimate of the mortality rate to be made with a 95% CI of $\pm 9.5\%$, assuming the mortality rate was around 30%. Finally, assuming the recruitment rate was around 30%, a sample of 300 patients approached to take part in the study, leading to 90 enrolled patients would allow for the recruitment rate to be estimated with a 95% CI of $\pm 5.2\%$. If the recruitment rate was nearer 50%, with 180 patients approached to recruit the 90 enrolled patients, the recruitment rate would be estimated with a 95% CI of $\pm 7.3\%$.

Statistical analysis

The analysis will be reported in line with the feasibility studies extension to the Consolidated Standards of Reporting Trials statement. ⁷⁶ The aims of the study were to estimate the recruitment, compliance and retention rates to inform the design of a future study and is not powered for hypothesis testing regarding the effectiveness of the intervention. Feasibility outcomes (recruitment, compliance and retention rates) will be presented with 95% CIs across the whole study population. Compliance and retention rates will also be presented by treatment arm to ensure balanced recruitment, but no formal statistical comparison tests will be made. Mortality and participant dropout rates will be presented with 95% CIs across the whole study population and within treatment arm. Clinical outcome data (secondary outcomes) will be presented as summary statistics using means and SDs or medians and ranges/IQRs, as applicable, across the whole study population and by treatment arm. These data will be used to inform the future trial but will not be used to draw conclusions about the effectiveness of the protocolised early rehabilitation intervention within this study.

Trial management

The chief investigator will ensure all study personnel are appropriately orientated and trained, oversee recruitment and report to the trial safety monitoring committee. Training will occur across sites using competency-based training developed at the primary site (University Hospital Southampton NHS Foundation Trust). A study steering group, consisting of an independent chair, expert members and two lay advisors will meet every 3 months. Fortnightly teleconferences with trial sites will be held to monitor conduct and progress. Timing and intervals of

visits and teleconferences will be reviewed at 3 months to ensure optimal time use.

The chief investigator and principal investigators will facilitate local monitoring by the research and development quality manager, research ethics committee (REC) review and provide access to source data as required. A monitoring report will be produced, summarising the visit, documents and findings. The chief investigator will ensure that all findings are addressed appropriately. The steering group will review all events in a timely manner. Additional monitoring will be scheduled where there is evidence or suspicion of non-compliance with the study protocol.

A data management and safety committee will be chaired by an independent expert. Quarterly reports will be given to the committee once recruitment has commenced.

Patient and public involvement

The study has been supported by patient advisory representatives. These representatives are members of the trial steering committee. Patient advisors partnered with us for the design of the study, the informational material to support the intervention, the burden of the intervention from the patient's perspective and contributed to the dissemination plan

Ethics and dissemination

Ethical approval has been granted by South Central Hampshire A Research Ethics Committee (REC reference 19/SC/0016). This study entitled: A feasibility study of Early Mobilisation Programmes in Critical Care (EMPRESS) was registered with Clinical Ttrials.gov on 10 December 2018.

Results of this proposed feasibility study will be disseminated for four key audiences: (1) patients and public; (2) intensive care staff, healthcare workers and potential future research delivery partners; (3) service delivery organisations; and (4) academic and potential future research collaborators. Dissemination activities will include feedback to patients and public involvement study focus group, feedback to study participants, presentations to local clinical teams and managers and commissioners and presentation at conferences attended by appropriate healthcare professionals. Where appropriate, results will be published in peer reviewed journals.

Safety and adverse events

Early mobility within ICUs is safe. In a review of physiotherapy in a critical care rehabilitation programme, 1110 patients underwent 5267 rehabilitation sessions; physiological abnormalities or potential adverse events occurred in only 6 per 1000 interventions. Mobilisation interventions will only be delivered if patients fit the safety criteria defined in table 1. Similar safety criteria have been used in other ICU rehabilitation studies. ^{78 79}

All adverse events will be documented. Any intervention will cease according to stopping criteria detailed in



table 1. Any such event will be recorded as an adverse event. The chief investigator will provide a monthly update to the safety monitoring committee. Serious adverse events are events that result in death, are lifethreatening or require prolonged hospitalisation. Any such event will be reported in accordance with the NHS Health Research Authority guidance.

DISCUSSION

EMPRESS is a feasibility study to assess if an RCT of protocolised rehabilitation with supine cycling can be delivered in ventilated patients in ICUs with differing organisational structures with blinded follow-up assessments. A recent meta-analysis indicated that protocolised rehabilitation significantly reduces duration of mechanical ventilation and ICU length of stay.²³ This is consistent with our findings when we introduced the early rehabilitation programme outlined here in our ICU. 45 Passive cycling commenced on ventilated patients may assist the recovery muscle strength in ICU patients, 43 although the overall benefits of leg cycle ergometry in the critically ill is inconclusive. 44 We describe a protocolised rehabilitation programme with supine cycling delivered as close to intubation as possible, at an intensity according to the patients' highest performance capability.

Both patient and organisational issues are recognised to the delivery of early rehabilitation of the critically ill patients.³⁵ A frequently reported challenge is the lack of appropriately qualified staff.⁸⁰ This study evaluates the safety, feasibility, effectiveness of delivery and cost efficiency of using therapy technicians to deliver protocolised rehabilitation interventions. In addition to the clinical benefits, early physical rehabilitation can also be cost saving.⁴⁹ Even with the cost of employment of additional therapy technicians specifically to assist in the delivery of we have found this early rehabilitation programme cost effective.⁸¹

This study will collect data on the dose of intervention delivered to all patients, reasons for non-delivery of protocol interventions, and the level of experience of therapists delivering the interventions. A qualitative process evaluation is designed to identify both patient and organisational challenges that have potential to be addressed in a potential future study. Findings will inform refinement of trial design and evaluation of the intervention, clarifying causal mechanisms behind study outcomes and providing additional context not adequately captured by the quantitative data. The process evaluation will be consistent with Medical Research Council guidance for conducting process evaluations of complex healthcare interventions. 82

Targeted sedation is embedded within this protocol as oversedation is one of the more commonly cited barriers to mobilisation of the ventilated patient.³⁵ This study opened to recruitment prior to the publication of the recommended core outcome set for critical care ventilation trials⁸³; however, three of the six outcomes listed

(duration of mechanical ventilation, duration of stay and health-related QOL) are secondary outcomes in this study and the other three outcomes are included in the data collected. This will be addressed should we proceed to a full RCT. Due to the nature of the intervention, it is not possible for this to be blinded; however, the follow-up assessments will be carried out by a blinded.

Results from EMPRESS will inform the design of a multicentred RCT, both identifying barriers to the implementation of the designed protocol and exploring how these may be addressed from feedback from the therapy and nursing teams in addition to the feedback from patients and their next of kin.

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Contributors RC and AB contributed equally to the preparation of the paper. RC and ZvW had the original idea for the study. RC, LD, IR, NH, AD, GS, ID and MG developed the trial protocol. IR devised the statistical analysis plan. MC developed the economic analysis. AB, GS, ID and RC prepared and submitted documents for research and development and ethical approval. RC, KM and AB wrote the manuscript. All authors reviewed the final version.

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Disclaimer The views expressed are those of the author(s) and not necessarily those of the National Institute for Health Research, NHS or the Department of Health and Social Care.

Competing interests None declared.

Patient and public involvement Patients and/or the public were involved in the design, conduct, reporting or dissemination plans of this research. Refer to the Methods and analysis section for further details.

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EMPRESS.

A feasibility study of early mobilisation in Critical Care. Patient Information Sheet

Version 2: 29th January 2019

Introduction

Study Title: EMPRESS: A study of very early mobilisation in Critical Care.

Invitation:

Your consultee has agreed on your behalf, to your participation on a research study. We would like to invite you to confirm whether you wish to continue or withdraw your participation from this research study.

This hospital is taking part in a national research study to investigate whether starting rehabilitation in the Intensive Care Unit, as soon as possible, will improve patient's long-term physical ability and quality of life.

When patients are sedated in Intensive Care, muscle wasting and weakness can occur very quickly and this can take a long time to recover from. Because we feel that it may be important to deliver rehabilitation physiotherapy as early as possible, it was agreed by your doctor and / or your relative/ friend that you could be involved in this study. This research has been approved by Hampshire Research Ethics Committee (IRAS number: 250165).

This patient information sheet provides information about the study to help you decide if you would like to continue to participate in it. It is important that you understand why the research is being done and what it involves.

Knowing what is involved will help you decide if you want to continue to take part in the research, so this Information Sheet explains the tests and treatments involved.

- Part 1 tells you about why we are doing this study and what will happen to you if you continue to take part.
- Part 2 gives you more detailed information about how we will run the study.

If you have no objection to continue taking part, we will ask you to read and sign a form that records your permission, called the consent declaration. We'll then give you a copy to keep. We will keep you fully informed during the study so you can let us know if you have any concerns or if you would prefer to be withdrawn. Taking part in this research is entirely voluntary. If you decide not to continue, you will still be offered the best possible standard of care.

Please ask us if there is anything that is not clear or if you would like more information. Take time to decide whether or not you wish to take part.

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PART ONE: Why are we doing this study and what will happen to you?

What is the purpose of this study?

We know that when patients are very unwell and need sedation in the Intensive Care Unit, they can lose muscle strength and size very quickly. It is normal to offer rehabilitation, but this often starts after the patient has woken up. By this time the muscles have already been affected. Previous studies have shown that this can take many months to recover from and may affect a patient's quality of life after leaving hospital.

In Southampton Hospital, researchers and physiotherapists started performing rehabilitation exercises much earlier than usual, even while the patient was sedated. They showed that this method reduced the patient's time on the ventilator and reduced the amount of time that they needed to be in Intensive Care.

We are now trying to discover whether this method will work in a number of different hospitals in the UK. We will also do some tests to see whether the patients who have this type of rehabilitation are stronger and able to engage in physical activity more easily, when they leave hospital and 3 months later.

Why have you been chosen?

You were enrolled in this study because during your admission to the Intensive Care you needed a ventilator (a machine to help you breathe) and sedation was needed to help keep you calm and comfortable. The treating doctor and physiotherapist thought that either very early rehabilitation or standard rehabilitation would be equally suitable. We may have given you very early physiotherapy already in the intensive care unit, because we are testing such early rehabilitation, but we would like to ask for your permission to continue.

Do I have to take part?

No. It is up to you to decide whether or not you would like to continue to take part. If you do, you will be given this information sheet to keep and be asked to sign the consent form. You are still free to withdraw at any time without giving a reason. The decision to withdraw at any time, or a decision not to take part, will not affect the standard of care you receive.

What will happen to me if I take part?

Participation in the study began in the Intensive Care Unit. The final tests will take place 3 months after you leave hospital.

In the Intensive Care Unit: Your treating doctor has assessed you to be eligible to take part in this study: EMPRESS. You were randomly allocated (like the flip of a coin) to receive either of the following:

 Standard physiotherapy: All patients on the trial will receive their normal physiotherapy. This will normally include activities to assist in keeping your airway

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clear and activities to maintain limb flexibility. These will not be affected by being on the trial.

OR

• Standard physiotherapy, as above, plus an extra 2 sessions of 30 minutes of rehabilitation from Monday to Friday. For patients receiving extra, early rehabilitation, in addition to your normal physiotherapy, you have been using a cycle machine that is designed to work, in the bed, even with sedated patients. As you wake up you have started to pedal for yourself, do some more bed-based exercises and finally get out of bed and start moving. All of these sessions have been and will continue to be run by a well-trained physiotherapist and the bedside nurses. We have already tested this method in University Hospital Southampton and it has reduced the length of time on the ventilator and ICU stay. During these sessions, you have been and will continue to be very carefully monitored for your own safety and the safety of lines, tubes and catheters.

These exercises will continue for a maximum of 28 days or less if you leave the Intensive care unit before then.

BOTH GROUPS

Additional assessments: So that we can test whether our new method works, patients
on the trial will undertake some extra assessments. These include a simple test of grip
strength by using a hand held pressure monitor; a test of arm and leg strength, ability
to stand and step and mobility and walking tests. There will also be quality of life and
health questionnaires.

There was a 50/50 chance of being allocated to either group. Neither you nor your doctor can decide which. No samples of blood are required for this research study.

In the hospital ward: When you have been discharged to a normal hospital ward, you won't receive any extra physiotherapy. Just before you go home, you will be tested again for muscle strength and mobility, including how far you can walk in 6 minutes. These tests will be supervised by a trained and experienced physiotherapist

Following discharge from hospital: Regardless of which group you were allocated to, after going home, you should follow the advice given to you by your doctors and physiotherapists. We have designed our study so that this will not affect our results.

You will be contacted by one of the critical care research team 3 months after you have been discharged home. We will arrange to see you for approximately one hour. During this visit we will test your walking speed, strength and agility. We will also ask for some questionnaires to be completed, which will assess how you feel about your quality of life and recovery.

The researchers would also like to have access to your medical record to obtain information relevant to the study. This information would be anonymised and kept confidential.

If you have any questions regarding the trial procedures, please don't hesitate to ask the intensive care or research doctors, physiotherapists and nurses.

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What do I have to do?

It is important to tell the doctor and the research staff about any treatments or medications you may have been taking, including over-the-counter medications, vitamins or herbal remedies, acupuncture or other alternative treatments. We would also like to know about any medical conditions which may affect the exercise.

Please let us know if you are involved in any other studies at this time.

What are the alternatives to participation?

Participation in this research is not the only option. You may decide to receive only standard or usual care. This is absolutely fine. Please feel free to discuss these options with your doctor before deciding whether or not to continue to take part in this research project.

What are the possible disadvantages of taking part?

Early mobility within ICU is safe. Potential risks may include, but not be limited to blood pressure or heart rate problems, breathing problems, problems with the tubes, lines and catheters.

In a review of physiotherapy within Intensive Care Units, involving over 1100 patients and 5267 episodes of physiotherapy in similar patients, there were 34 potential safety events (equivalent to 6 events in 1000 episodes of physiotherapy), Most of these were potentially related to changes in heart rate or blood pressure which settle quickly in stopping the physiotherapy.

In Southampton, over a four year period, we have treated over 500 patients in this way and had 2 non-serious adverse events.

The doctors, physiotherapists and nurses who will be caring for you while in the ICU, are trained to recognise the effects on the body associated with physical rehabilitation and will treat you accordingly. You will be continue to be monitored and assessed. Your safety is always our number one priority.

There may be side effects that the researchers do not expect or do not know about and that may be serious. Please tell the doctor immediately about any new or unusual symptoms that you get.

What are the possible benefits of taking part?

We cannot guarantee or promise that you will receive any benefits from this research. This study aims to further medical knowledge and may improve future treatment of patients who need to be on a ventilator, however it may not directly benefit you.

For how long will I be in the research study?

The final research assessment will take place 3 months after discharge from hospital. Once that is done, your participation in the study will end.

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What happens if there is a problem?

We will keep you fully informed of any problems which may be related to the study.

Will taking part in the study be kept confidential?

Yes. All of the information about participation and the data collected will be kept confidential.

Information held by the NHS and records maintained by the NHS Information Centre and the NHS Central Register may be used to help contact you and provide information about your health status. This information may be obtained and stored by the study research team to enable long term follow-up.

University Hospital Southampton is the sponsor for this study based in the United Kingdom. We will be using information from you and/or your medical records in order to undertake this study and will act as the data controller for this study. This means that we are responsible for looking after your information and using it properly. University Hospital Southampton will keep information about you for 10 years after the study has finished.

Your rights to access, change or move your information are limited, as we need to manage your information in specific ways in order for the research to be reliable and accurate. If you withdraw from the study, we will keep the information about you that we have already obtained. To safeguard your rights, we will use the minimum personally-identifiable information possible.

You can find out more about how we use your information at https://www.hra.nhs.uk/information-about-patients/

[Local NHS site name] will collect information from you and/or your medical records for this research study in accordance with our instructions.

(Local NHS site name) will keep your name, NHS number and contact details confidential and will not pass this information to University Hospital Southampton. [Local NHS site name] will use this information as needed, to contact you about the research study, and make sure that relevant information about the study is recorded for your care, and to oversee the quality of the study. Certain individuals from University Hospital Southampton and regulatory organisations may look at your medical and research records to check the accuracy of the research study. University Hospital Southampton will only receive information without any identifying information. The people who analyse the information will not be able to identify you and will not be able to find out your name, NHS number or contact details.

[Local NHS site name] will keep identifiable information about you, including the consent form from this study for 10 years after the study has finished.

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Contact Details:

Local PI

Consultant Critical Care,

Hospital address

Dr XXXX: 02381 XXXXXX

Research Nurse: 02381 XXXXXX

ICU: 02381 XXXXXX

PART 2: How we will run this study.

What if relevant new information becomes available?

During the research project, new information about the risks and benefits of the study may become known to the researchers. If this occurs, you will be told about this new information and the doctor will discuss whether this new information affects you.

If any information becomes available which could affect your participation in the study the research doctor will tell you about it and discuss whether you want to continue in the study. If you decide to not continue in the study, the research doctor will make arrangements for your care to continue as normal. If you decide to continue in the study you will be asked to sign an updated consent form.

Also, on receiving new information the research doctor might consider it to be in your best interests to withdraw you from the study. He/she will explain the reasons and arrange for your care to continue.

If the study is stopped for any other reason, you will be told why and your continuing care will be arranged.

What will happen if I don't want to carry on with the study?

You can withdraw from the study at any time without giving an explanation and be assured that it will not impact on any part of your further treatment.

If you decide to withdraw from the study, the researchers would like to keep your health information that has already been collected. This is to help them make sure that the results of the research can be measured properly. If you do not want them to do this, you can tell them when you withdraw from the research project.

What if there is a problem?

If you have any concerns regarding the study, please ask to speak to the ICU doctor in charge of your care or ask to speak to name of local PI, the consultant who is in charge of the study.

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Complaints:

If you have a concern about any aspect of this study, you should ask to speak with the researchers or the Intensive Care doctors and nurses, who will do their best to answer your questions. If you remain unhappy and wish to complain formally, you can do this through the NHS Complaints Procedure. Please localise with your hospital PALS contact details.

Harm:

In the event that something does go wrong and you are harmed during the research study there are no special compensation arrangements. If you are harmed and this is due to someone's negligence then you may have grounds for a legal action for compensation against University Hospital Southampton, but you may have to pay the legal costs. The normal National Health Service complaints mechanisms will still be available to you.

Involvement of the General Practitioner/Family doctor (GP)

If you are agreeable we would like to inform your GP of your participation in the study. If you do not wish for your GP to be informed, please let us know and indicate on the consent form that you do not wish your GP to be informed.

Will taking part in this study be kept confidential?

If you continue with the study, some parts of your medical records and the data collected for the study will be looked at by authorised researchers from University Hospital Southampton and University of Southampton who are sponsoring and organising the research. They may also be looked at by authorised people to check that the study is being carried out correctly. All will have a strict duty of confidentiality to you, as a research participant and we will do our best to meet this duty.

All information that is collected about you during the course of the research will be kept strictly confidential. Any information about you that leaves the hospital will have your name and address removed so that you cannot be recognised from it.

Anonymised data collected during the study may be sent to associated researchers in other countries, where the laws don't protect your privacy to the same extent as the law in the UK but the study team will take all reasonable steps to protect your privacy.

You have the right to check the accuracy of data held about you and correct any errors.

What will happen to the results of the research study?

They will be published in a medical journal, presented at conferences and lay press where possible.

Who is organising and funding the research?

Dr Rebecca Cusack from University Hospital Southampton is the lead researcher, who is organising the research.

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The research is funded by the NHS through the National Institute for Health Research, Research for Patient Benefit scheme.

Who has reviewed the study?

Hampshire Research Ethics Committee (IRAS number: 250165) have reviewed this study and given their approval.

Thank you very much for taking the time to read this information sheet at this very stressful time.

If you have any further questions please ask the doctors in Intensive Care, Dr (local PI) or one of the research team.

If you agree to continuing participation in this study, please keep this information sheet and you will be given a copy of the agreement form that you will be asked to sign.

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Version 2:

29th January 2019

FORM TO BE ON SITE SPECIFIC HEADED PAPER

University Hospital Southampton NHS Foundation Trust

Critical Care, Anaesthesia & Peri-operative Medicine Dept,
Research office CE 93. MP 24,
University Hospital Southampton,
Tremona Road,
Southampton
SO16 6YD

Tel: 023 8120 5308 Fax: 023 8120 5378

Consent form for patients participating in EMPRESS. A feasibility study of early mobilisation programmes in Critical Care.

Name of Researcher:	
Please initial box	
1. I confirm that I have read and understood the Patient Information Sheet (version Dated) for the EMPRESS study. I have had the opportunity to ask questions about the study and understand what is involved.	
2. I have no objection to taking part in the above study.	
3. I understand that my participation is voluntary and that I am free to withdraw from the study at any time, without giving any reason and without my care or legal rights being affected.	
4. I understand that relevant sections of my medical notes and data collected during the study, may be looked at by individuals from the research team, from regulatory authorities or from the NHS Trust, where it is relevant to my taking part in this research. I give permission for these individuals to have access to my records.	
5. I understand that information held by the NHS and records maintained by the NHS Information Centre and the NHS Central Register may be used to help contact me and provide information about my health status. I give permission for this information to be obtained and stored by the study research team to enable long term follow-up.	
6. I agree to my GP being informed of my participation in the study.	
7. I agree to take part in the above study	

Name of Participant:	Signature:	Date:
Person undertaking consultation (researcher):	Signature:	Date:

Original Informed Consent form to be filed in the Investigator Site File.

1 copy to be given to the patient

1 copy to be filed in the patients' hospital notes.

Version 2: 29th January 2019

Site specific header to be inserted here

EMPRESS.

A feasibility study of early mobilisation in Critical Care. Information for Consultee

Version 2: 29th January 2019

Introduction

Study Title: EMPRESS: A study of very early mobilisation in Critical Care.

Invitation: This hospital is taking part in a national research study to investigate whether starting rehabilitation in the Intensive Care Unit, as soon as possible, will improve patients' long-term physical ability and quality of life.

When patients are sedated in the Intensive Care unit, muscle wasting and weakness can occur very quickly and this can take a long time to recover from. Because we feel that it may be important to deliver rehabilitation physiotherapy as early as possible, we wish for your relative/friend to participate in the trial.

Because your relative/friend is unable to decide for himself/herself whether to participate in this research, we'd like to ask your opinion as to whether or not they would want to be involved. Please consider what you know about their wishes and feelings and what you think may be best for them.

If we have been unable to contact you, your relative/friend may have been enrolled as a participant in this research project with the approval of their treating doctor and the Hampshire Research Ethics Committee (IRAS number: 250165). If this is the case, then we seek to confirm that you are in agreement.

Knowing what is involved will help you decide if you want your relative/friend to continue to take part in the research, so this information sheet explains the tests and treatments involved.

- Part 1 tells you about why we are doing this study and what will happen to your relative if they take part.
- Part 2 gives you more detailed information about how we will run the study.

If you decide your relative/friend would have no objection to taking part, we will ask you to read and sign a form that records your permission, called the consultee declaration. We'll then give you a copy to keep. We will keep you fully informed during the study so you can let us know if you have any concerns or you think your relative/friend should be withdrawn. Taking part in this research is entirely voluntary. If you decide not to continue, they will still be offered the best possible standard of care.

Please ask us if there is anything that is not clear or if you would like more information. Take time to decide whether or not you wish your relative/friend to take part.

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PART ONE: Why are we doing this study and what will happen to my friend / relative?

What is the purpose of this study?

We know that when patients are very unwell and need sedation in the Intensive Care Unit, they can lose muscle strength and size very quickly. It is normal to offer rehabilitation, but this often starts after the patient has woken up. By this time the muscles have already been affected. Previous studies have shown that this muscle weakness may take many months to recover from and may affect a patient's quality of life after leaving hospital.

In Southampton Hospital, researchers and physiotherapists started performing rehabilitation exercises much earlier than usual, even while the patient was sedated. They showed that this method reduced the patient's time on the ventilator and reduced the amount of time that they needed to be in Intensive Care.

We are now trying to discover whether this method will work in a number of different hospitals in the UK. We will also do some tests to see whether the patients who have this type of rehabilitation are stronger and able to engage in physical activity more easily, when they leave hospital and 3 months later.

Why has my relative been chosen?

Your relative has been enrolled in this study because during their admission to the Intensive Care Unit he/she needed a ventilator (a machine to help them breathe) and sedation to help keep them calm and comfortable. The treating doctor and physiotherapist thought that either very early rehabilitation or standard rehabilitation would be equally suitable. We may have made a start already, because we are testing such very early rehabilitation, but we would like to ask for your permission to continue.

Does my relative/ friend have to take part?

No. It is up to you to decide whether or not you would like him/her to continue to take part. If you decide they can, you will be given this information sheet to keep and be asked to sign a permission form. You are still free to withdraw your relative at any time without giving a reason. The decision to withdraw at any time, or a decision not to take part, will not affect the standard of care your relative receives.

At an appropriate time, when we hope your relative has recovered sufficiently, we will ask their permission to use the data we have collected. If they do not agree we will not collect any new data and ask if we may use the data already collected.

What will happen to my relative if they take part?

Participation in the study will begin in the Intensive Care Unit. The final tests will take place 3 months after they leave hospital.

In the Intensive Care Unit: Your friend/ relative's treating doctor has assessed them to be eligible to take part in this study: EMPRESS. They were randomly allocated (like the flip of a coin) to receive either of the following:

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• **Standard physiotherapy:** All patients on the trial will receive their normal physiotherapy. This will normally include activities to assist in keeping their airway clear and activities to maintain limb flexibility. These will not be affected by being on the trial.

OR

• Standard physiotherapy, as above, plus an extra 2 sessions of 30 minutes of rehabilitation from Monday to Friday. For patients receiving extra, early rehabilitation, in addition to their normal physiotherapy, they will start using a cycle machine that is designed to work, in the bed, even with sedated patients. As your friend/ relative wakes up they will start to pedal for themselves, do some more bed-based exercises and finally get out of bed and start moving. All of these sessions will be run by a well-trained physiotherapist and the bedside nurses. We have already tested this method in University Hospital Southampton and it has reduced the length of time on the ventilator and ICU stay. During these sessions, they will be very carefully monitored for their own safety and the safety of their lines, tubes and catheters.

These exercises will continue for a maximum of 28 days or less if they leave the Intensive care unit before then.

BOTH GROUPS

Additional assessments: So that we can test whether our new method works, patients
on the trial will undertake some extra assessments. These include a simple test of grip
strength by using a hand held pressure monitor; a test of arm and leg strength, ability
to stand and step and mobility and walking tests. There will also be quality of life and
health questionnaires.

There was a 50/50 chance of being allocated to either group. Neither you nor their doctor can decide which. No samples of blood are required for this research study.

In the hospital ward: When your friend/ relative has been discharged to a normal hospital ward, they won't receive any extra physiotherapy. Just before they go home, they will be tested again for muscle strength and mobility, including how far they can walk in 6 minutes. These tests will be supervised by a trained and experienced physiotherapist

Following discharge from hospital: Regardless of which group your friend/ relative was allocated to, after going home, they should follow the advice given to them by their doctors and physiotherapists. We have designed our study so that this will not affect our results.

They will be contacted by one of the critical care research team 3 months after they have been discharged home. We will arrange to see them for approximately one hour. During this visit we will test their walking speed, strength and agility. We will also ask for some questionnaires to be completed, which will assess how they feel about their quality of life and recovery.

The researchers would also like to have access to your relative or friend's medical record to obtain information relevant to the study. This information would be anonymised and kept confidential.

EMPRESS: A feasibility study of early mobilisation programmes in Critical Care.

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If you have any questions regarding the trial procedures, please don't hesitate to ask the intensive care or research doctors, physiotherapists and nurses.

What do I have to do?

It is important to tell the doctor and the research staff about any treatments or medications you know your relative/friend may have been taking, including over-the-counter medications, vitamins or herbal remedies, acupuncture or other alternative treatments. We would also like to know about any medical conditions which may affect the exercise.

Please just let us know if your relative/friend is involved in any other studies at this time.

What are the alternatives to participation?

Participation in this research is not the only option. You may decide for your relative to receive only standard care physiotherapy. That is absolutely fine. Please feel free to discuss these options with your relative's doctor before deciding whether or not to continue to take part in this research project.

What are the possible disadvantages of taking part?

Early mobility within ICU is safe. Potential risks may include, but not be limited to blood pressure or heart rate problems, breathing problems, problems with the tubes, lines and catheters.

In a review of physiotherapy within Intensive Care Units, involving over 1100 patients and 5267 episodes of physiotherapy in similar patients, there were 34 potential safety events (equal to 6 events in 1000 episodes of physiotherapy). Most of these were related to changes in heart rate or blood pressure and settles quickly on stopping the physiotherapy.

In Southampton, over a 4 year period, we have treated over 500 patients in this way and had 2 events needing attention but neither resulted in harm to the patient..

The doctors, physiotherapists and nurses who will be caring for your relative or friend while in the ICU, are trained to recognise the effects on the body associated with physical rehabilitation and will treat accordingly. Your friend/ relative will be continually monitored and assessed. Their safety will always be our number one priority.

There may be side effects that the researchers do not expect or do not know about and that may be serious. Please tell the doctor immediately if you are worried about any new or unusual symptoms that your relative/friend gets.

What are the possible benefits of taking part?

We cannot guarantee or promise that your relative will receive any benefits from this research. This study aims to further medical knowledge and may improve future treatment of patients who need to be on a ventilator, however it may not directly benefit your relative/friend.

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Consultee Information Sheet Version 2: 29th January 2019

For how long will my relative/ friend be in the research study?

The final research assessment will take place 3 months after discharge from hospital. Once that is done, your friend/ relative's participation in the study will end.

What happens if there is a problem?

We will keep you and your friend/ relative, fully informed of any problems which may be related to the study.

Will taking part in the study be kept confidential?

Yes. All of the information about participation and the data collected will be kept confidential.

Information held by the NHS and records maintained by the NHS Information Centre and the NHS Central Register may be used to help contact your friend/ relative and provide information about their health status. This information may be obtained and stored by the study research team to enable long term follow-up.

University Hospital Southampton is the sponsor for this study based in the United Kingdom. We will be using information from their medical records in order to undertake this study and will act as the data controller for this study. This means that we are responsible for looking after their information and using it properly. University Hospital Southampton will keep information about them for 10 years after the study has finished.

Your friend/ relative's rights to access, change or move your information are limited, as we need to manage the information in specific ways in order for the research to be reliable and accurate. If they withdraw from the study, we will keep the information that we have already obtained. To safeguard their rights, we will use the minimum personally-identifiable information possible.

You can find out more about how we use information at https://www.hra.nhs.uk/information-about-patients/

[Local NHS site name] will collect information from their medical records for this research study in accordance with our instructions.

(Local NHS site name) will keep name, NHS number and contact details confidential and will not pass this information to University Hospital Southampton. [Local NHS site name] will use this information as needed, to contact your relative/ friend about the research study, and make sure that relevant information about the study is recorded for their care, and to oversee the quality of the study. Certain individuals from University Hospital Southampton and regulatory organisations may look at medical and research records to check the accuracy of the research study. University Hospital Southampton will only receive information without any identifying information. The people who analyse the information will not be able to identify patients and will not be able to find out their name, NHS number or contact details.

[Local NHS site name] will keep identifiable information, including the consent form from this study, for 10 years after the study has finished.

EMPRESS: A feasibility study of early mobilisation programmes in Critical Care.

Consultee Information Sheet Version 2: 29th January 2019

Contact Details:

Local PI details Address

Dr local PI: 02381 XXXXXX

Research Nurse: 02381 XXXXXX

ICU: 02381 XXXXXX

PART 2: How we will run this study.

What if relevant new information becomes available?

During the research project, new information about the risks and benefits of the study may become known to the researchers. If this occurs, you will be told about this new information and the doctor will discuss whether this new information affects your relative.

If any information becomes available which could affect participation in the study the research doctor will tell you about it and discuss whether you want your relative to continue in the study. If you decide your relative should not continue in the study, the research doctor will make arrangements for your relative's care to continue as normal. If you decide to allow your relative to continue in the study you will be asked to sign an updated agreement form.

Also, on receiving new information the research doctor might consider it to be in your relative's best interests to withdraw them from the study. He/she will explain the reasons and arrange for their care to continue.

If the study is stopped for any other reason, you will be told why and your relative's continuing care will be arranged.

What will happen if I don't want my relative to carry on with the study?

You can withdraw your relative from the study at any time without giving an explanation and be assured that it will not impact on any part of your relative's further treatment.

If you decide to withdraw your relative from the study, the researchers would like to keep your relative's health information that has been collected. This is to help them make sure that the results of the research can be measured properly. If you do not want them to do this, you can tell them when you withdraw your relative from the research project.

What if there is a problem?

If you have any concerns regarding the study, please ask to speak to the ICU doctor in charge of your friend/ relative's care or ask to speak to (name of local PI), the consultant who is in charge of the study.

EMPRESS: A feasibility study of early mobilisation programmes in Critical Care.

Consultee Information Sheet Version 2: 29th January 2019

Complaints:

If you have a concern about any aspect of this study, you should ask to speak with the researchers or the Intensive Care doctors and nurses, who will do their best to answer your questions. If you remain unhappy and wish to complain formally, you can do this through the NHS Complaints Procedure. (Please localise with your hospital PALS contact details)

Harm:

In the event that something does go wrong and your relative is harmed during the research study there are no special compensation arrangements. If your relative is harmed and this is due to someone's negligence then your relative may have grounds for a legal action for compensation against University Hospital Southampton, but they may have to pay the legal costs. The normal National Health Service complaints mechanisms will still be available to them.

Involvement of the General Practitioner/Family doctor (GP)

If you are agreeable we would like to inform your friend/ relative's GP of their participation in the study. If you do not wish for their GP to be informed, please let us know and indicate on the consent form that you do not wish for their GP to be informed.

Will allowing my relative to take part in this study be kept confidential?

If your relative joins the study, some parts of their medical records and the data collected for the study will be looked at by authorised persons from University Hospital Southampton and University of Southampton who are sponsoring and organising the research. They may also be looked at by authorised people to check that the study is being carried out correctly. All will have a strict duty of confidentiality to your relative as a research participant and we will do our best to meet this duty.

All information that is collected about your relative during the course of the research will be kept strictly confidential. Any information about your relative that leaves the hospital will have their name and address removed so that they cannot be recognised from it.

Anonymised data collected during the study may be sent to associated researchers in other countries, where the laws don't protect your privacy to the same extent as the law in the UK but the study team will take all reasonable steps to protect your privacy.

Your relative has the right to check the accuracy of data held about them and correct any errors.

What will happen to the results of the research study?

They will be published in a medical journal, presented at conferences and lay press where possible.

EMPRESS: A feasibility study of early mobilisation programmes in Critical Care.

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Who is organising and funding the research?

Dr Rebecca Cusack from University Hospital Southampton is the lead researcher, who is organising the research.

The research is funded by the NHS through the National Institute for Health Research, Research for Patient Benefit scheme.

Who has reviewed the study?

Hampshire Research Ethics Committee (IRAS number: 250165) have reviewed this study and given their approval.

Thank you very much for taking the time to read this information sheet at this very stressful time.

If you have any further questions please ask the doctors in Intensive Care, Dr (local PI) or one of the research team.

If you agree to your relative participating in this study please keep this information sheet and you will be given a copy of the agreement form that you will be asked to sign.

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Southampton
SO16 6YD

Tel: 023 8120 5308 Fax: 023 8120 5378

Consultee declaration form for patients participating in EMPRESS. A feasibility study of early mobilisation programmes in Critical Care.

Name of Researcher: Please initial box	
1. I [name of consultee] have been consulted about [name of potential participant]'s participation in this research project. I have read and understood the Consultee Information Sheet (version; Dated). I have had the opportunity to ask questions about the study and understand what is involved.	
2. In my opinion he/she would have no objection to taking part in the above study.	
3. I understand that I can request he/she is withdrawn from the study at any time, without giving any reason and without his/her care or legal rights being affected.	
4. I understand that relevant sections of his/her medical notes and data collected during the study, may be looked at by individuals from the research team, from regulatory authorities or from the NHS Trust, where it is relevant to their taking part in this research. I give permission for these individuals to have access to their records.	
5. I understand that information held by the NHS and records maintained by the NHS Information Centre and the NHS Central Register may be used to help contact me or my friend / relative and provide information about their health status. I give permission for this information to be obtained and stored by the study research team to enable long term follow-up.	
6. I agree to their GP being informed of their participation in the study.	

When completed: 1 for participant; 1 (original) for researcher site file; 1 to be kept in medical notes.

Version 2: 29th January 2019 IRAS ID:250165

<u>Please confirm either:</u>		
I confirm that I will act as the personal con-	sultee for:	
Relationship to participant:		
Name of consultee:	Signature:	Date:
Person undertaking consultation (researcher):	Signature:	Date:

Original Informed Consent form to be filed in the Investigator Site File.

1 copy to be given to the patient

1 copy to be filed in the patients' hospital notes.

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