Protocol

BMJ Open Nutrition and Exercise in Critical Illness Trial (NEXIS Trial): a protocol of a multicentred, randomised controlled trial of combined cycle ergometry and amino acid supplementation commenced early during critical illness

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ABSTRACT

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Introduction Survivors of critical illness often experience significant morbidities, including muscle weakness and impairments in physical functioning. This muscle weakness is associated with longer duration mechanical ventilation, greater hospital costs and increased postdischarge impairments in physical function, quality of life and survival. Compared with standard of care, the benefits of greater protein intake combined with structured exercise started early after the onset of critical illness remain uncertain. However, the combination of protein supplementation and exercise in other populations has demonstrated positive effects on strength and function. In the present study, we will evaluate the effects of a combination of early implementation of intravenous amino acid supplementation and in-bed cycle ergometry exercise versus a 'usual care' control group in patients with acute respiratory failure requiring mechanical ventilation in an intensive care unit (ICU).

Methods and analysis In this multicentre, assessorblinded, randomised controlled trial, we will randomise 142 patients in a 1:1 ratio to usual care (which commonly consists of minimal exercise and under-achievement of guideline-recommended caloric and protein intake goals) versus a combined intravenous amino acid supplementation and in-bed cycle ergometery exercise intervention. We hypothesise that this novel combined intervention will (1) improve physical functioning at hospital discharge; (2) reduce muscle wasting with improved amino acid metabolism and protein synthesis in-hospital and (3) improve patient-reported outcomes and healthcare resource utilisation at 6 months after enrolment. Key cointerventions will be standardised. In-hospital outcome assessments will be conducted at baseline, ICU discharge and hospital discharge. An intentto-treat analysis will be used to analyse all data with additional per-protocol analyses.

Ethics and dissemination The trial received ethics approval at each institution and enrolment has begun. These results will inform both clinical practice and future

Strengths and limitations of this study

- This is the first randomised controlled trial (RCT) evaluating the combination of exercise and protein supplementation started in the early phase of critical illness.
- We have developed a rigorous framework to evaluate the effect of the study intervention on the patient's functional recovery and outcomes.
- As a phase II RCT, the study has a relatively small sample size recruited from four participating centres.
- This study evaluates a combined intervention and will not be able to independently evaluate the effect of the nutrition versus exercise on study outcomes.

research in the area. We plan to disseminate trial results in peer-reviewed journals, at national and international conferences, and via nutritional and rehabilitation-focused electronic education and knowledge translation platforms. **Trial registration number** NCT03021902; Pre-results.

INTRODUCTION

Up to 20 million people worldwide receive life support in intensive care units (ICUs) each year.¹ More than 750 000 Americans require mechanical ventilation annually,^{2 3} and 300 000 receive ventilation for >5 days.³⁻⁶ More critically ill patients are surviving hospitalisation due to recent medical advances.⁷ However, this survival comes at a cost. ICU survivors frequently experience significant post-ICU morbidities, commonly physical morbidities, including muscle weakness and impairments in physical functioning that can persist for years.⁸⁻¹⁴ Muscle weakness in the ICU is associated with delayed liberation from ventilation, extended ICU and hospital stays, worse long-term survival, physical functioning and quality of life (QOL).⁸⁹¹⁴⁻¹⁸ A recent review stated that the¹⁹ highest ranking research priority in the critical care nutrition/ metabolism field was to evaluate the effect of protein dose with active and passive mobilisation in the acute phase of critical illness.

Recent randomised controlled trials (RCTs) have shown that providing increased total calories to ICU patients may not improve outcomes.²⁰⁻²⁸ However, only two of these RCTs delivered >50% of recommended protein, and these two delivered only ~33% of the protein dose targeted in our RCT.^{20 28} Moreover, observational studies report that optimising daily protein intake, rather than total caloric intake, decreases infections, mechanical ventilation duration, time to discharge and mortality.²⁹⁻³³ Although some studies report that increased protein intake is associated with greater muscle wasting,³⁴ later ICU discharge³⁵ and increased mortality,³⁶ these studies did not adjust for a key confounding variables: the total time that protein was received or total caloric energy received.³⁷ Moreover, a small RCT demonstrated that greater protein intake is associated with improved pulmonary function in ICU patients with chronic obstructive pulmonary disease (COPD),³⁸ and attenuation of the muscle loss observed in the context of critical illness.^{39 40} A recent study found that infusing intravenous amino acids for 3 hours in ICU patients improved protein balance and stimulated an anabolic response.⁴¹ Many small trials have found that 2.0-2.8 g/kg/day of protein intake is safe and improves nitrogen balance.^{42–51} A recent RCT of IV amino acids of up to 2.0g/kg/day in 474 ICU patients demonstrated safety, and although amino acid therapy did not preserve kidney function (the primary outcome), this study did not measure any performance-based measures and did not combine the amino acids intervention with exercise.⁵² A posthoc analysis from the trial examining the effects of baseline kidney function on mortality found those with normal kidney function had a reduction in 90-day mortality.⁵³ Furthermore, those with baseline kidney dysfunction and/or those with a risk of progression of acute kidney injury (AKI) found no significant effect on 90-day mortality. Recent guidelines and systematic reviews recommend up to 2.0-2.5 g/kg/day and suggest that these doses are safe; thus, we propose to evaluate a similar dose.^{54 55}

With increasing recognition of physical complications after critical illness, recent studies have evaluated exercise interventions started early after ICU admission. These studies have demonstrated safety and feasibility, and provide some evidence of reduced myofibrillar proteolysis, less muscle atrophy and consequently, improved strength and physical functioning, and decreased durations of mechanical ventilation and ICU stay.^{56–59} However, these studies are mainly single site, with modest sample sizes, and have grossly under-delivered protein to patients. Moreover, a number of recent trials of early exercise interventions have largely been negative.^{60–62}

Hence, further evaluation of early ICU exercise interventions is needed, particularly in combination with amino acid supplementation, to reduce muscle weakness and physical complications after critical illness.

In various patient populations, combining protein and exercise has the greatest benefits compared with either nutrition or exercise alone. In older people, combined exercise and protein supplements improve protein synthesis and strength versus either intervention alone.^{63–67} In one study, combined exercise and supplementation in elderly patients increased muscle strength by 40% over exercise alone and by 130% over supplementation alone.⁶⁷ In patients with obesity,⁶⁸ HIV/AIDs⁶⁹ and chronic obstructive pulmonary disease,⁷⁰ as well as healthy volunteers undergoing bed rest,^{71 72} the combination of exercise and nutritional intervention (vs nutrition alone) yields the greatest benefit on muscle mass and strength. In a meta-analysis, protein supplementation and exercise (vs exercise alone) enhanced strength and muscle mass in non-critically ill adults.⁷³ Moreover, a recent RCT of 93 ICU survivors, conducted in the out-patient setting, demonstrated that combined oral amino acid supplementation and exercise improved 6 min walk distance (6MWD, the primary outcome for our proposed trial) measured 3 months after enrolment, compared with either amino acids or exercise alone.⁷⁴ We chose to use in-bed cycling because loss of lean body mass with bed rest is most pronounced in the legs.^{75 76} Our research finding also demonstrates greater weakness in legs versus arms in ICU patients.⁷⁷ Moreover, leg strength is critical to ambulation, and thus key to functional independence and living at home.⁷⁸⁻⁸³ Although the generalisability of these findings to patients in the early phase of critical illness is unclear, these data provide biologic plausibility that this combined intervention may reduce physical impairments.⁸⁴

Our multicentred, assessor-blinded, phase II RCT delivers a combination of intravenous amino acid supplementation and in-bed cycle ergometry exercise early after the onset of critical illness, versus usual care, in patients requiring mechanical ventilation in the ICU. The hypothesis is this novel RCT is that the combined intervention will (1) improve physical functioning at hospital discharge; (2) reduce muscle wasting with improved amino acid metabolism and protein synthesis in-hospital and (3) improve patient-reported outcomes and healthcare resource utilisation at 6 months after enrolment.

METHODS AND ANALYSIS Study design

This is a phase II multicentre, assessor-blinded, RCT conducted at four academic medical centres in the USA: the University of Vermont Medical Center (UVMMC), Johns Hopkins University (JHU), Harborview Medical Center (HMC) and Wake Forest University Baptist Medical Center (WFUBMC). The data co-ordinating centre is the Clinical Evaluation Research Unit (CERU)

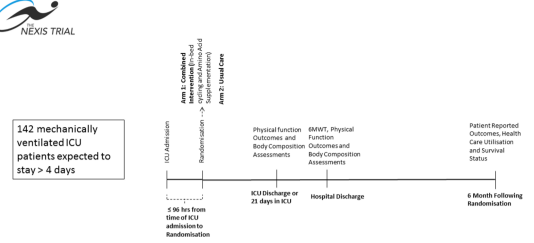


Figure 1 Study design and timeline. ICU, intensive care unit; 6MWT, 6 min walk time.

at the Kingston General Hospital in Kingston Canada (see www.ceru.ca). See figure 1 for study design. This trial is funded by the National Institutes of Health (NIH) and the notice of award was received in April 2017 with the duration of the trial is expected to be 5 years.

Eligibility criteria

Eligibility criteria are listed in table 1. We expect our intervention to be most effective when delivered early⁸⁵; thus, we will limit enrolment to the first 96 hours after intubation. Patients hospitalised for a longer time prior to their ICU admission may be less responsive to our intervention.³⁴ As such, we have excluded patients if they have spent greater than 5 days admitted to hospital in the 14 days leading up to the current ICU admission.

Participant selection and recruitment

This combined intervention is targeted to critically ill patients with acute respiratory failure. Eligibility is determined by daily screening at the study site ICUs by trained members of the research team. Final approval of patient eligibility must be given by a trained site investigator. The planned flow of patients through the study is depicted in figure 2. All data will be captured in an electronic database to monitor recruitment at each study site.

Consent

Once patients have been screened and confirmed by the site investigator or a subinvestigator, the participant or Legally Authorised Representative is approached for informed consent (online supplementary file). The research staff engages the Legally Authorized Representative (LAR) in a conversation to discuss the trial and ensure that they have understood the material. They are given ample time to review the materials and ask any relevant questions.

Randomisation

After consent, patients are randomised 1:1 to receive either (1) the combined intravenous amino acid and cycle ergometry exercise intervention or (2) usual care. Randomisation will be stratified by site and hospital length of stay prior to randomisation (<48 hours vs \geq 48 hours). Randomisation will further be restricted by using permuted blocks of random size within strata. The randomisation list was commuter generated by the senior biostatistician at the data coordinating centre who is uninvolved with site enrolment and unaware of which site codes map to which sites. The randomisation is implemented using the data coordinating centre's secure central web-based randomisation system which maintains concealment of future allocations and has been used successfully for several large international RCTs.

Blinding

Due to the nature of in-bed cycle ergometry, it is not possible to blind this study to patients, families or ICU clinicians. To minimise bias, blinded assessors are completing all outcome assessments and we are collecting data on key cointerventions to evaluate balance between groups in cointerventions. To assist with blinding outcome assessments, patients are prompted (using standardised language) *not* to disclose their perception of treatment allocation during the outcome assessments.⁸⁶ Moreover, to evaluate effectiveness of blinding, outcome assessors document their 'best guess' regarding allocation (intervention vs control) at each assessment.^{86 87}

Study intervention

Description of the amino acid intervention

The amino acid intervention is provided in addition to 'usual care' enteral and/or parenteral nutrition via an intravenous infusion (Clinisol 15% by Baxter) to target a *total* protein delivery of 2.0–2.5 g/kg/day. Consistent with a prior RCT demonstrating safety in the ICU,⁵² amino acids are delivered based on ideal body weight.⁸⁸ Given a typical daily enteral/parenteral protein intake for ICU patients of 1.2 g/kg/day,⁸⁹ the expectation is to administer approximately 1.2 g/kg/day of amino acid infusion (which yields 1.0 g/kg/day protein⁹⁰) to reach a total of 2.2 g/kg/day total protein. Amino acids are infused via an

Inclusion criteria

1. >18 years old

intervention)

2. Requiring mechanical ventilation

total duration of mechanical

3. Expected ICU stay ≥4 days after

exposure to the proposed

enrolment (to permit adequate

with actual or expected

ventilation ≥48 hours

Inclusion and exclusion criteria for stud

Table 1

	6		
	-		
a for study entry			
Exclusion criteria	Rationale for exclusion		
1. >96 continuous hours of mechanical ventilation before enrolment	Intervention most effective delivered early ⁸⁵		
2. Expected death or withdrawal of life-sustaining treatments within this hospitalisation	Patients unlikely to receive benefit		
3. No expectation for any nutritional intake within the subsequent 72 hours	Intervention intended to occur in addition to standard clinical nutritional intake		
4. Severe chronic liver disease (MELD score ≥20) or acute hepatic failure	Amino acid supplementation may be harmful in patients with severe liver disease		
5. Documented allergy to the amino acid intervention	Unable to receive proposed intervention		
6. Metabolic disorders involving impaired nitrogen utilisation	Unable to receive amino acid infusion		
7. Not ambulating independently prior to illness that leads to ICU admission (use of gait aid permitted)	Unable to perform outcome assessments		
8. Pre-existing primary systemic	May not benefit from proposed		

8. Pre-exis neuromuscular disease (eq. Guillain Barre)

9. Neuromuscular blocker infusion (eligible once infusion discontinued if other inclusion criteria met)

10. Intracranial or spinal process affecting motor function

11. Pre-existing cognitive impairment or language barrier that prohibits outcomes assessment

12. Patients in hospital >5 days prior to ICU Muscle weakness likely already established admission 13. Lower extremity impairments that prevent cycling (eg, amputation, knee/hip

injury) 14. Remaining intubated for airway

protection only 15. Weight ≥150 kg

- 16. Physician declines patient enrolment
- 17. Insufficient intravenous access

18. Pregnant 19. Incarcerated Unable to receive proposed cycling intervention Less likely to have muscle weakness and benefit from the interventions

intervention (ie, different mechanism of

Do not meet safety criteria for cycling

intervention (ie, different mechanism of

Unable to perform outcome assessments

May not benefit from proposed

muscle weakness)

muscle weakness)

intervention

Exceeds maximum weight permitted for use of the cycle device

Not appropriate to conduct trial Need dedicated access for nutrition intervention for several hours a day

Unknown effects in fetus

Vulnerable population

ICU, intensive care unit.

indwelling central venous catheter, when available. The solution also can be diluted to a 7.5% solution and delivered via peripheral venous catheter if needed. Clinicians caring for patients in both groups are encouraged to maintain euvolemia with diuresis, when clinically appropriate. Intravenous amino acids beginas close to the time of randomisation as feasible. The target dose, delivered as

a continuous infusion, will be started during or immediately after the exercise session. Usually, a participant will receive the amino acid infusion every day that the participant receives the cycling intervention. However, on a day when the cycling intervention is withheld (eg, safety reason), the amino acid infusion will still be given. Amino acid infusion will continue until ICU discharge or 21 days,

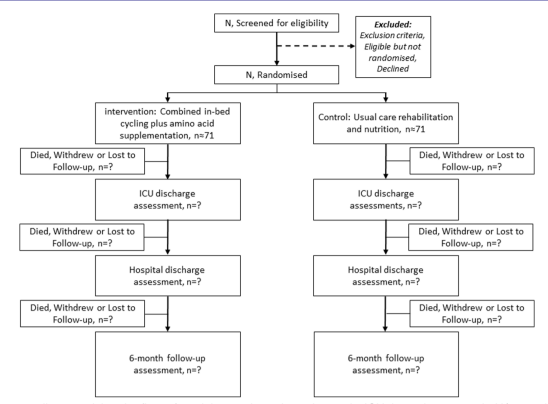


Figure 2 Consort diagram giving the flow of participants throughout the study. ICU, intensive care unit; N/n, number.

whichever occurs earlier. If the patient is discharged from ICU, then readmitted, amino acid administration will continue until the 21st calendar day after randomisation.

Guidelines recommend that ICU patients with AKI receive standard protein intake as it appears to be utilised, improves nitrogen balance and does not lead to increased urea generation.⁵⁵ Additionally, guidelines suggest that ICU patients receiving renal replacement therapy (RRT) should receive increased protein, up to a maximum of 2.5 g/kg/day.⁵⁵ Therefore, patients with moderate to severe AKI or chronic kidney disease who are not yet receiving RRT will be allowed to participate in the study. However, because these patients are at risk of developing azotemia with increased amino acid intake, the urea will be monitored daily as part of standard of care. If urea is >100 mg/dL and there is no plan for RRT that day, the amino acid infusion can be decreased by half of current rate in discussion with the ICU clinical team. It is important to note that azotemia, alone, has not been shown to be harmful; nevertheless, this approach will be taken.54

Description of in-bed cycle ergometry exercise

The cycling intervention will be delivered by trained staff, and started as close to the time of randomisation as feasible (ie, within 24 hours of randomisation). All sites will use a MotoMed Letto II cycle ergometer. The cycling sessions will occur according to a detailed protocol beginning with a safety assessment and continue for the first at least 5 days per week. The safety guidelines for cycling and the cessation of a cycling session can be found in box 1.

The intervention group will receive cycling sessions, for up to 45 min duration (as tolerated by patient),⁸³ with vigorous verbal encouragement to promote active cycling. This goal of a 45 min cycling duration was chosen for several reasons: (1) to help prevent under-dosing of the exercise, with it being >2x the dose delivered in the prior positive RCT of cycle ergometry (20 min/ day in ICU),⁸³ (2) our own experience suggests feasibility/tolerability of sessions >20 min in duration⁹¹ and (3) even longer cycling sessions may not be beneficial, with a recent small RCT reporting no difference in muscle loss between cycling 1 hour versus 2 hours per day.⁵⁶ The implementation of the cycling intervention is protocolised to provide graduated resistance during each session and between daily sessions. Cycling will continue through ICU discharge or 21 calendar days after randomisation, whichever occurs earlier (same as amino acids). The intervention specifically occurs during the ICU stay (and not post-ICU) since this represents the portion of hospitalisation in which patients are most exposed to bed rest/immobility (ie, due to ventilation and severity of illness). If the patient is discharged from ICU, then readmitted, the cycling intervention will continue until the 21st calendar day after randomisation. Proper implementation of the cycling will be overseen locally by site investigators and research staff. The data coordinating centre will run periodic data reports to review the implementation of the combined intervention. The protocol that is used in this RCT is adapted from an existing protocol that was developed and extensively used at one of the study sites.⁹¹

Box 1 Cycling safety guidelines

- A. Criteria to not commence cycling session
 - (Cycling should not occur if any of the following conditions are present for greater than 15 min within the 2 hours prior to cycling.)
 - Heart rate <50 or >140 bpm, or new arrhythmia.
 - New onset of chest pain of potential cardiac origin.
 - Presence of femoral extracorporeal membrane oxygenation (ECMO) or intra-aortic balloon pump (IABP).
 - Mean arterial pressure <65 mm Hg or below target or >120 mm Hg or above target.
 - A single vasopressor as outlined below:
 - Dopamine >12.5 mcg/kg/min.
 - Phenylephrine >2 mcg/kg/min.
 - $\quad Norepine phrine > 1 \ mcg/kg/min.$
 - ≥ 2 vasopressors at same time, as outlined below:
 - Vasopressin at ≥ 0.04 units/min.
 - Dopamine >10.0 mcg/kg/min.
 - Phenylephrine >1.6 mcg/kg/min.
 - Norepinephrine >0.8 mcg/kg/min.
 - Participant is pale/sweaty and requests not to start due to feeling unwell.
 - Fi0, >0.8.
 - PEÉP >15 cm H₂0.
 - Sp0₂ falls >10% of resting level or <85% for more than 60 s, or below target level for more than 60 s in participants with abnormal baseline Sp0₂
 - Recent receipt of neuromuscular blocker medication.
 - Clinical team's opinion that participant should not receive cycling despite the absence of above criteria.
- B. Criteria to terminate cycling session
 - Heart rate <50 or >140 bpm or new arrhythmia.
 - New onset of chest pain of potential cardiac origin.
 - Presence of femoral ECMO or IABP.
 - Mean arterial pressure <65 mm Hg or below target or >120 mm Hg or above target.
 - A single vasopressor as outlined below:
 - Dopamine >12.5 mcg/kg/min.
 - Phenylephrine >2 mcg/kg/min.
 - Norepinephrine >1 mcg/kg/min.
 - ≥2 vasopressors at same time, as outlined below:
 - Vasopressin at ≥ 0.04 units/min.
 - Dopamine >10.0 mcg/kg/min.
 - Phenylephrine >1.6 mcg/kg/min.
 - Norepinephrine >0.8 mcg/kg/min.
 - Participant is pale/sweaty and requests to stop due to feeling unwell.
 - Fi0₂>0.8.
 - $PEEP > 15 cm H_20$.
 - Sp0₂ falls>10% of resting level or <85% for more than 60 s, or below target level for more than 60 s in participants with abnormal baseline Sp0₂.
 - Fatigue.
 - Patient declines to continue.
 - Clinical team's opinion that participant should not stop cycling despite the absence of above criteria.

Standard ICU cointerventions for all patients

Important aspects of routine care will be standardised for both intervention and usual care group patients based

Table 2 Patient variables—collected a	at enrolment
Patient variables – collected at enrolment	Collection method
Age/sex/ethnicity/race demographic data	Chart review
Body mass index	Chart review
Comorbidities: Charlson* and functional indices**	Chart review
Baseline function: Functional Status Score for ICU ⁷⁷	Proxy interview (see Outcomes below)
Baseline function: Katz Activities of Daily Living Scale ¹⁵⁴ and Lawton IADL	Proxy interview (see Outcomes below)
Baseline function: SF-36 Physical Function domain and walk impairment question	Proxy interview
Clinical Frailty Scale	Proxy interview
ICU admission diagnosis (eg, sepsis, renal failure)	Chart review
Severity of illness: APACHE II†	Chart review
Patient location immediately prior to ICU and to hospital admission	Chart review

*Charlson Index: a score for in- patients derived from 19 comorbidities; an increased score reflecting increased 1 -year mortality. ¹⁵⁵

**Functional Index: an 18-diagnosis scale for ICU *patients* predicting 1 - year SF-36 Physical Function score; increased score reflecting worse function.¹⁵⁶⁻¹⁵⁸

†APACHE II: a severity of illness index using age, medical conditions and acute physiology, with higher scores reflecting increased short-term mortality. ¹⁵⁹

IADL, instrumental activities of daily living; ICU, intensive care unit.

on pre-existing protocols at all study sites, including (1) turning by nurse every 2 hours while bedbound; (2) frequency, duration and intervention type of all physical therapy and occupational therapy sessions in the ICU; (3) dietician consultation and prescription of a standard enteral nutrition formula (at approximately 20–25 kcal/kg/day and 1.2 g/kg/day) within 48 hours after intubation; (4) blood glucose control with insulin; (5) electrolyte replacement protocols; (6) daily sedation interruption or minimisation and (7) daily spontaneous breathing trials as part of ventilator weaning protocols.

On-study data collection

Demographic, laboratory, physiological, nutritional and rehabilitation data will be collected at study enrolment, then daily in the ICU as outlined in tables 2 and 3. The database will be under password protection in an institutional computer located at the CERU and only the research staff will have access.

Primary outcome

In developing our evaluation framework, we followed the recommendations of a recent expert consensus statement.⁹² The primary aim of this study is to measure in-patient muscle strength and physical functioning. This will

 Table 3
 ICU-related variables—collected at enrolment and

 DAILY during ICU stay*

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ICU variables—collected at enrolment and daily during ICU stay	Collection method
Sedation medications and dose, with sedation status—RASS score ¹⁶⁰ and CAM-ICU	Chart review
Neuromuscular blocker, corticosteroid drug use and dose	Chart review
Insulin dose and blood glucose level	Chart review
SOFA§ organ failure score (including vasopressor data)	Chart review
Creatinine, creatine phosphokinase, blood urea nitrogen	Chart review
Nutrition received (calories/protein, type and route of feeding)	Chart review
Mobility/rehabilitation received	Chart review
Compliance with proposed intervention regimen	CRF review
SOFA: a validated composite score of 6 organ sy	istems used to

§SOFA: a validated composite score of 6 organ systems used to assess the severity of ICU organ dysfunction.¹⁶¹

CRF, case report form; RASS, Richmond Agitation-Sedation Scale.

be evaluated using one primary and multiple secondary outcomes. The timing of these are provided in table 4. The primary outcome will be the walking distance achieved during a 6min walk test (6MWT) measured at hospital discharge. Implementation of the test will be based on the 2014 ATS standards, with adaptation, as needed, for the in-patient setting and ICU survivor population.⁹³ The test will only be performed once, rather than twice as recommended due to the feasibility of asking critically ill patients to perform the test multiple times. The 6MWT is a reliable, valid, responsive measure of physical function⁹⁴ for survivors of acute respiratory failure. Hospital discharge was chosen as the endpoint to assess the primary outcome, in congruence with other cycling studies.^{60 95 96}

Secondary outcomes

Secondary strength-related outcome measures will include the following.

- 1. Overall strength using Medical Research Council (MRC) sum-score evaluated via standardised 'manual muscle testing' with each of 12 muscle groups assessed using a 6-point MRC scale⁹⁷ and summed to a total score (range: 0–60).^{8 9 14 77 85 98-100}
- 2. Quadriceps force, via hand-held dynamometry^{83 101} for of both lower extremities. Each will be scored by averaging the results of three trials.^{102 103}
- 3. Distal strength measured via isometric hand grip strength via a hydraulic hand dynamometer performed bilaterally as per American Society of Hand Therapist guidelines¹⁰⁴ and evaluated using normal values.¹⁰⁵

Secondary Physical Functioning outcomes will include the following.

- 1. Short Physical Performance Battery which measures balance, walking speed and rising from a chair^{7880106–110}
- 2. Functional Status Score for ICU, which is a 5-item, 35-point assessment of bed mobility, transfers and ambulation, designed for ICU patients^{77 86 111 112} and was designed and validated specifically in ICU patients evaluated 8-point Functional Independence Measure response scale used throughout rehabilitation assessments^{113–116} and is responsive to change during recovery for ICU patients.^{77 86 112 117 118}

Body composition

Body composition will be assessed using additional secondary outcomes. Ultrasound (US) will be used to measure rectus femoris and vastus intermedius cross-sectional area and thickness using a published protocol.³⁴ ⁷¹ ^{119–123} Changes in muscle echodensity will be measured using quantitative greyscale analysis.³⁴ ¹²³–¹²⁵ We will do US at baseline (shortly after randomisation), ICU discharge and hospital discharge. Chest CT and abdominal CT scans will be obtained when clinically available, with chest CT used to measure pectoralis muscle area as it correlates with clinical outcomes in patients with COPD, lung cancer and critical illness,126-128 and abdominal CT scan used to measure abdominal and visceral adipose tissue at the level of the third lumbar vertebra. We will obtain all CT data, when performed for clinical purposes at any point during hospital stay (including during admission process prerandomisation), for comparative analyses between participants in the intervention and control groups. Due to expense, radiation exposure, and required transport out of the ICU for CT, we will not obtain specific research CT scans.

Hospital Acquired Infections

An additional secondary outcome is hospital-acquired infections. Data suggest that increased protein intake reduces infections.^{20 30 33} Recording culture results and antibiotics administered, along with pertinent clinical data will enable adjudication of infectious complications using pre-existing methodology.^{129 130}

Patient-reported outcomes

Finally, outcomes after hospital discharge will be assessed via 6-month phone-based follow-up. Health-related QOL will be measured using SF-36 version 2 (SF-36 v2) and EQ-5D-5L. The SF-36 is valid and reliable across a variety of patient groups, including ICU survivors.¹³¹¹³² The EQ-5D-5L is included, in addition to SF-36 v2, because it is suitable for patients with inattention and fatigue,133 134 recommended for use in ICU survivors.¹³⁵¹³⁶ Physical functional status will be measured using Katz activities of daily living (ADL)¹³⁷ and Lawton's Instrumental ADL (IADL)¹³⁸ scales, as well as return to baseline work/activity and living location. Mental and cognitive function will be measured, in addition, using the HADS, IES-R and MoCA-BLIND screening questionnaires as part the recommended Core Outcome Measurement Set for evaluating postdischarge outcomes in acute respiratory failure survivors.¹³⁶ Healthcare resource utilisation will be

	Instrument	Assessment timing
Primary outcome		
Physical functioning	6 min walk distance (6MWD)	Hospital discharge
Secondary outcomes		
Overall strength-upper and lower extremity	MRC sum-score	Hospital discharge
Quadriceps force-lower extremity strength	Hand-held dynamometry	Hospital discharge
Distal strength-hand grip strength	Hand grip dynamometry	ICU and hospital discharge
Overall Physical Functional status	SPPB and FSS-ICU	ICU and hospital discharge
Physical functioning (ADL)	Katz ADL	Hospital discharge
Mortality	Chart review	ICU and hospital discharge
Length of ventilation, ICU and hospital stay	Chart review	ICU and hospital discharge
ICU readmission and reintubation	Chart review	Hospital discharge
Hospital-acquired infections	Chart review	Hospital discharge
Discharge location (eg, home vs rehab)	Chart review	Hospital discharge
Body composition	Ultrasound of rectus femoris, vastus intermedius, tibialis anterior	Enrolment, ICU and hospital discharge
Body composition (when clinically available)	Chest CT scan (above the aortic arch)	Only when clinically available
Body composition (when clinically available)	Abdominal CT scan at third lumbar vertebra	Only when clinically available
Health-related quality of life	SF-36 and EQ-5D-5L	Telephone survey at 6 months
Physical functioning	Katz ADL; Lawton IADL	Telephone survey at 6 months
Physical functioning	Return to baseline work/activity	Telephone survey at 6 months
Physical functioning	Living location	Telephone survey at 6 months
Mental and cognitive functioning	MoCA-BLIND, HADS and IES-R	Telephone survey at 6 months
Healthcare resource utilisation	Admission to ICU, hospital, rehabilitation and nursing facility	Telephone survey at 6 months

MRC, Medical Research Council; SPPB, Short Physical Performance Battery.

Table 4 Primary and secondary outcomes – all performed by blinded assessors*

assessed through a structured interview regarding admissions to hospital, skilled nursing and rehabilitation facilities as done in prior research.^{11 139–142} In order to improve retention, a call will be made to participants at 3 months to update contact information and act as a reminder of upcoming follow-up assessments to be completed at the 6-month time point.

Adverse events

Patients will be monitored daily, their medical records examined and their care providers queried for adverse events that are serious and unexpected in nature. Unexpected SAEs will be recorded from the time of randomisation until ICU discharge or day 21, whichever comes first. These unexpected SAEs which are inconsistent with underlying pathophysiology or progression of underlying disease will be documented in study source documents and reported unexpected SAEs that are related or possibly related to participation in the study will be reported to the participating site IRBs, DSMB and NHLBI on an expedited basis. Deidentified reporting will occur within 7 calendar days of receipt of the initial report for fatal and life-threatening events. All other events (ie, non-fatal and non-life-threatening) will be reported within 15 calendar days of receipt of the initial report. In addition, we will capture events that may not be serious but may be related to the amino acid infusion (uremia>100 mg/dL, peripheral phlebitis) or cycle ergometry exercise (times when the safety criteria thresholds for blood pressure, heart rate are reached and cycling stopped).

Statistical analysis

Power calculation

Sample size was determined based on the primary outcome, 6MWT at hospital discharge. We plan to enrol 64 evaluable patients per arm (total n=128 evaluable patients) which achieves >80% power using the Wilcoxon Rank-sum test at a two-sided alpha=0.05 to detect differences in 6MWD distribution between the two groups with discrimination (ie, ROC AUC or concordance index)>64%. Figure 3 depicts a sample scenario yielding a concordance index of 64% and 80% power.

Using a traditional t-test approach, this sample size would have 80% power with α =0.05 (two sided) to detect a 50-m difference in 6MWD (with SD=100 based on our

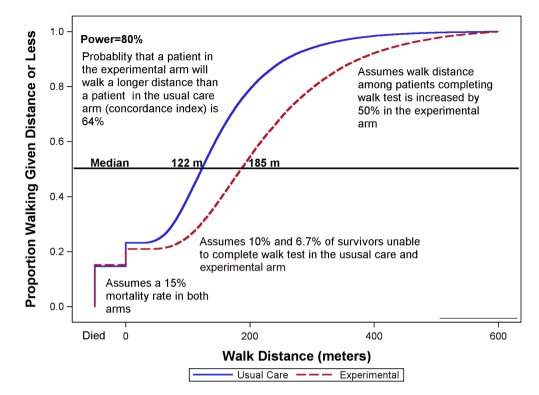


Figure 3 Sample empirical distribution function of walk distance.

prior RCT data in ICU patients (NCT01206166)). We expect <10% non-participation rate for the primary outcome (ie, patients capable of 6MWD but declining testing); thus, we will enrol a total of 142 patients.

Primary analysis

All analyses will follow the intention-to-treat principle. A two-tailed p-value<0.05 will be considered statistically significant. The 6MWD will be compared between the two groups using the rank-based Mann-Whitney U test.¹⁴³ This approach allows inclusion of all randomised patients, per the intent-to-treat principle, by assigning decedents a lower value than all survivors (eg, -1) and assigning patients incapable of performing the 6MWD test a value of 0, including the rare patients discharged while still receiving mechanical ventilation.¹⁴³ ¹⁴⁴ In addition to this rank-based test, we will describe the differences in 6MWD between the two groups graphically as shown in figure 3. Time from randomisation to 6MWD testing will be described by arm, and a sensitivity analysis will adjust for the time to testing before comparing the adjusted ranked 6MWD between arms. A per-protocol analysis will also be performed by excluding patients randomised to the intervention arm who do not receive at least 3 days of the combined intervention and patients in the usual care group who stay less than 3 days in the ICU.

Secondary analysis

Secondary continuous outcomes will be analysed similar to the primary outcome. Categorical secondary outcomes will include death and unable-to-perform as potential categories and will be analysed using Fisher's exact tests. As secondary outcomes will be considered hypothesis-generating, we will not formally correct for multiple comparisons but will consider the number of secondary comparisons when interpreting our results.

The number of missing values will be described for all outcomes. Outcomes will not be considered missing due to death since the proposed statistical methods include decedents in the between arm comparisons. Patient characteristics will be compared between those with versus without missing outcomes.^{145–147} If >5% of outcome data are missing, multiple imputation will be used,¹⁴⁸ and for the primary analysis a 'missing not at random' sensitivity analysis will be performed using the tipping point approach of the pattern mixture model with multiple imputation, as per the SAS MI procedure.^{149 150}

Patient and public involvement

This research question was the highest priority topic arising from an expert panel on research priorities for intensive care nutrition and metabolism.¹⁹ The outcomes and associated measurement instruments being utilised have been informed by a robust international Delphi consensus process with a panel that included almost 25% patient/family representatives,^{151 152} and is also informed by many foundational studies (leading up to the Delphi) that included patient/family input, as summarised in a recent publication.¹⁵³ The intervention that formed the foundation for this RCT was extensively tested with critically ill patients for its feasibility, including any need to

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stop the intervention due to patient request, agitation, pain or physiological issues.⁹¹ We did not include patients and members of public in the design process, recruitment or conducting of the study. The results will be published in peer-reviewed journals and reference to these works will be posted on our websites (eg, www.criticalcarenutr ition.com), which are in the public space.

ETHICS

The study sponsor is the NIH/NHBLI. The University of Vermont is the lead site. The NIH/NHBLI will take no part in design, conduct of the study, collection, management, analysis, interpretation of the data, preparation, review and approval of the manuscript. We have constituted a data monitoring committee to provide a thirdparty assessment of interim analyses and review of the scientific literature as it evolves over the duration of the trial.

DISCUSSION

We are conducting the first randomised trial of combined exercise and nutrition applied early in the context of critically illness. We will test whether this intervention improves the functional recovery and QOL of survivors of critical illness. If proven effective, this combined intervention has potential to improve care of ICU patients and have an important public health impact on the growing number of ICU survivors. After completion of this phase II RCT, a decision regarding progression to a phase III RCT will be based on the study findings of feasibility, safety and benefit, in addition to funding considerations. The limitations of work include a lack of blinding of the study interventions, which is impossible to do. Accordingly, we have blinded outcome assessors and standardised key cointerventions, and will report on their use in each group. We are utilising novel rank-based statistical analyses to account for patients who die before assessment of the primary outcome, as well as patients who are unable to walk at hospital discharge. The small sample size and the limited number of sites limits the generalisability of our findings. Nevertheless, our results will surely inform both clinical practice and future research in this area.

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Contributors DKH, RS, DN, DCF, CH, AD, ND and MM designed the study and the protocol submission. DKH and GJC wrote the manuscript. RS, DN, DCF, CH, AD, ND and MM revised the manuscript and approved the final version.

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Competing interests For purposes of conducting this NIH/NHLBI-funded clinical trial, Dr Heyland reports a grant and donated amino acid product from Baxter Healthcare Corporation and an equipment loan from Reck Medical Devices. For purposes of conducting this NIH/NHLBI-funded clinical trial, Dr Needham reports a grant and donated amino acid product from Baxter Healthcare Corporation and an equipment loan from Reck Medical Devices. For purposes of conducting this NIH/ NHLBI-funded clinical trial, Dr Stapleton reports a grant and donated amino acid product from Baxter Healthcare Corporation and an equipment loan from Reck Medical Devices. Dr Mourtzakis, John Clarke and Andrew Day have nothing to disclose. Dr Files reports grants from National Institute of Health during the conduct of the study. Dr Files also reports a grant and donated amino acid product from Baxter Healthcare Corporation and an equipment loan from Reck Medical Devices. Dr Hough reports grants from NIH during the conduct of the study. Dr Hough also reports a grant and donated amino acid product from Baxter Healthcare Corporation and an equipment loan from Reck Medical Devices. Nicolaas E Deutz declares no conflicts of interests, but discloses that he is a coinventor of several patents, owned by others, has served on scientific advisory boards for Novartis and Baxter and has been a consultant for Abbott Nutrition, Ajinomoto, OCERA and VitaNext. Texas A&M CTRAL receives funding from NIH, NSF, Abbott Nutrition, ICAAS, ESPEN fellowship, George Abramson Donation, Yani Mizubuti Donation and Internal Grants.

Patient consent for publication Not required.

Ethics approval The trial received approval of the ethics committee at each institution. It will be conducted in accordance with International Conference on Harmonisation Good Clinical Practice guidelines and the approved protocol. We plan to disseminate the results in peer-reviewed journals, at national and international conferences and via content specific web-based knowledge translation platforms (see and .

Provenance and peer review Not commissioned; externally peer reviewed.

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