# VITDALIZE UK: Effect of high-dose vitamin D3 on 28-day mortality in adult critically ill patients with severe vitamin D deficiency

Submission date	Recruitment status	[X] Prospectively registered
17/09/2020	Recruiting	[X] Protocol
Registration date	Overall study status	Statistical analysis plan
24/09/2020	Ongoing  Condition category	Results
Last Edited		Individual participant data
17/11/2025	Nutritional, Metabolic, Endocrine	[X] Record updated in last year

#### Plain English summary of protocol

Background and study aims

Vitamin D deficiency (low vitamin D levels) is common in patients who are unwell (around 70%). This has been found to be related to an increased risk of infection and death. There are many reasons why patients are poorly and those who do survive can suffer long-term health problems in the future. It is not known whether being vitamin D deficient is a cause or effect of being unwell, and research into whether vitamin D is useful is not clear. There are no guidelines to measure and treat patients admitted to intensive care who are critically ill and are vitamin D deficient. Vitamin D is cheap and easily available, and if using vitamin D is found to help, can be quickly put into standard practice in hospitals. VITDALIZE is an international trial that aims to recruit 2400 patients from across Europe. Countries that are participating include the UK, Austria, Germany and Belgium. The UK part of VITDALIZE aims to recruit 600 patients into the trial. The aim of this trial is to see if giving a high dose of vitamin D in critically ill patients can improve survival, length of hospital stay, and quality of life.

#### Who can participate?

Males and females aged 18 years and above admitted to ICU who are severely vitamin D deficient

#### What does the study involve?

Participants will be visited or contacted up to six times (days 0, 5, 12, 28, 90 and 1 year). At the beginning of the trial, participants will be given either a high dose of vitamin D or placebo (dummy supplement) on day 0 and a daily dose of either vitamin D or placebo from day 1 to day 90. On day 5 (if still in the hospital), the research team will take some more blood to see how participants are responding to treatment. On days 0, 28, 90 and after 1 year the research team will contact the participant (either in person if still in hospital or by telephone) to ask a few questions about their health. Participants will be asked to consent to provide some optional blood samples at up to three additional timepoints for future approved research. This would mean providing blood samples (25-30 ml; equivalent to 2 tablespoons) on days 0, 5 and 12.

What are the possible benefits and risks of participating?

There may be no direct benefits of taking part, but the results will lead to the best treatment being offered to patients who are unwell and vitamin D deficient.

Where is the study run from?

Birmingham Clinical Trials Unit at the University of Birmingham (UK)

When is the study starting and how long is it expected to run for? October 2019 to July 2027

Who is funding the study?

National Institute for Health Research Health Technology Assessment Programme (NIHR HTA) (UK)

Who is the main contact?

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#### Contact information

#### Type(s)

Scientific

#### Contact name

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#### Type(s)

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#### Contact name

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#### Additional identifiers

Clinical Trials Information System (CTIS)

2016-002460-13

Integrated Research Application System (IRAS)

274476

ClinicalTrials.gov (NCT)

NCT03188796

**Grant Code** 

DRKS00016940, HTA 17/147/33

Central Portfolio Management System (CPMS)

46276

#### Study information

#### Scientific Title

Effect of high-dose vitamin D3 on 28-day mortality in adult critically ill patients with severe vitamin D deficiency: the UK arm of an international multi-centre, placebo-controlled, Phase III double-blind trial

#### Acronym

VITDALIZE UK

#### Study objectives

- 1. The primary hypothesis is that in critically ill patients with severe vitamin D deficiency as defined by 25(OH)D concentration  $\leq 12 \text{ng/ml}$  (30nmol/L), a high-dose vitamin D replacement strategy, compared to placebo, leads to 28-day survival.
- 2. Further hypotheses are that high-dose vitamin D supplementation reduces hospital and ICU mortality, 90-day and 1-year mortality, reduces the length of stay in ICU and hospital, and improves health-related quality of life of patients and is cost-effective.

#### Ethics approval required

Old ethics approval format

#### Ethics approval(s)

Approved 03/11/2020, South Central – Oxford C Research Ethics Committee (Level 3, Block B, Whitefriars, Lewins Mead, Bristol, BS1 2NT, UK; +44 (0)207 104 8041; oxfordc.rec@hra.nhs.uk), REC ref: 20/SC/0300

#### Study design

Randomized; Interventional; Design type: Treatment, Drug

#### Primary study design

Interventional

#### Study type(s)

Treatment

#### Health condition(s) or problem(s) studied

Vitamin D deficiency

#### **Interventions**

Once randomised into the trial, the patient will receive either:

- 1. Intervention: A single loading high-dose oral/enteral vitamin D3 (540,000IU cholecalciferol, Oleovit™, Fresenius Kabi, Austria, dissolved in 37.5 ml of medium-chain triglycerides MCT) followed by 4000 IU daily (10 drops) for 90 days.
- 2. Control: Placebo, identical regime of loading dose of 37.5 ml MCT (Fresenius Kabi, Austria) followed by MCT (10 drops) daily for 90 days.

#### Intervention Type

Supplement

#### Primary outcome(s)

All-cause mortality, measured face-to-face (if an inpatient), by telephone, medical records, NHS digital/ONS data, at 28 days after randomisation

#### Key secondary outcome(s))

- 1. Mortality measured using telephone call, medical records, NHS digital/ONS data at 90 days and 1 year
- 2. ICU and hospital mortality measured face-to-face (if an inpatient), by telephone, medical records, NHS digital/ONS data at day 5, 28, 90 and 1 year
- 3. Hospital and ICU length of stay measured face-to-face (if an inpatient), by telephone, medical records, NHS digital/ONS data, starting at day 0, ending at discharge from the trial site or day 90 or mortality, whichever occurs first
- 4. Organ dysfunction measured by Sequential Organ Function Assessment score (SOFA), number of organ failures (0-6; defined as > 2 SOFA points in each of the 6 categories) on day 5
- 5. Hospital and ICU readmission measured using telephone call, medical records, NHS digital /ONS data until day 90
- 6. Discharge destination (home, rehabilitation, other hospital) measured using telephone, medical records, NHS digital/ONS data at discharge
- 7. Assessment of psychosocial functions measured using Katz Activities of Daily Life at day 90
- 8. Self-reported infections requiring antibiotics measured face-to-face (if an inpatient), by telephone, medical records, NHS digital/ONS data until day 90
- 9. Health-related quality of life measured using EQ-5D-5L at 90 days and 1 year
- 10. Disability assessment measured using WHODAS 2.0 at 90 days and 1 year
- 11. Secondary healthcare utilisation (ICU and hospital length of stay, readmissions and utilisation of hospital and community care resources after hospital discharge 1 year after randomisation), from Hospital Episode Statistics, civil registry data held by NHS Digital and patient questionnaires in the first year after randomisation
- 12. Health economics analysis measured face-to-face (if an inpatient), by telephone, medical records, NHS digital/ONS data at day 28, 90 and 1 year
- 13. Cost-effectiveness of screening for and treating VDD in critical illness measured using

telephone call, medical records, NHS digital/ONS data at day 28, 90 and 1 year 14. Cost per quality-adjusted life-year gained measured using telephone call, medical records, NHS digital/ONS data at 1 year after randomisation and at end of life

#### Exploratory outcome:

Health-related quality of life measured using proxy EQ-5D-5L and proxy WHODAS 2.0 at randomisation (day 0)

#### Safety outcomes:

- 1. Hypercalcaemia measured using medical records on day 5
- 2. Self-reported falls, fractures measured face-to-face (if an inpatient), by telephone, medical records, NHS digital/ONS data until day 90
- 3. New episodes of kidney stones measured using medical records until day 90

#### Completion date

31/07/2027

#### Eligibility

#### Key inclusion criteria

- 1. Patients ≥18 years
- 2. Anticipated ICU stay ≥48 hours
- 3. Admission to ICU ≤72 hours before screening for VDD
- 4. Severe VDD (25(OH)D ≤12 ng/ml [30 nmol/l]) using either the hospital's clinical laboratory or rapid bedside testing after ICU admission

#### Participant type(s)

**Patient** 

#### Healthy volunteers allowed

No

#### Age group

Mixed

#### Lower age limit

18 years

#### Upper age limit

100 years

#### Sex

All

#### Total final enrolment

0

#### Key exclusion criteria

1. Severe gastrointestinal dysfunction (>400 ml nasogastric tube residual volume)/unable to receive trial medication

- 2. Not expected to survive initial 48 hours of admission or treatment withdrawal imminent within 24 hours
- 3. Patient with DNAR (Do Not Attempt Resuscitation) orders in place
- 4. Hypercalcemia (>2.65 mmol/l corrected calcium and/or >1.35 mmol/l ionized calcium at screening)
- 5. Known kidney stones within the last 12 months
- 6. Known active tuberculosis within the last 12 months
- 7. Known sarcoidosis within the last 12 months
- 8. Women of childbearing age who have tested positive for pregnancy or who are lactating
- 9. Known hypersensitivity to the trial drug or excipient
- 10. Medical team deem it not suitable to include patient
- 11. Known prisoners in the custody of HM Prison and Probation services

#### Date of first enrolment

12/04/2021

#### Date of final enrolment

31/07/2026

#### Locations

#### Countries of recruitment

United Kingdom

**England** 

Northern Ireland

Wales

#### Study participating centre

**University Hospitals Birmingham NHS Foundation Trust** Trust HQ, PO Box 9551

Queen Elizabeth Medical Centre Edgbaston

Birmingham

England

B15 2TH

#### Study participating centre South Tees Hospitals NHS Foundation Trust

James Cook University Hospital Marton Road Middlesbrough Cleveland England TS4 3BW

#### Study participating centre Bolton NHS Foundation Trust

The Royal Bolton Hospital Minerva Road Farnworth Bolton England BL4 0JR

#### Study participating centre Guy's and St Thomas' NHS Foundation Trust

St Thomas' Hospital Westminster Bridge Road London England SE1 7EH

### Study participating centre Belfast Health & Social Care Trust

Knockbracken Healthcare Park Saintfield Road Belfast Northern Ireland BT8 8BH

#### Study participating centre Mid Yorkshire Hospitals NHS Trust

Pinderfields Hospital Aberford Road Wakefield England WF1 4DG

#### Study participating centre East Lancashire Hospitals NHS Trust

Royal Blackburn Hospital Haslingden Road Blackburn England BB2 3HH

#### Study participating centre Great Western Hospitals NHS Foundation Trust

Great Western Hospital Marlborough Road Swindon England SN3 6BB

Study participating centre
Diana, Princess of Wales Hospital
Scartho Road
Grimsby
England

# **Study participating centre Derriford Hospital**Derriford Road

Derriford Road
Derriford
Plymouth
England
PL6 8DH

**DN33 2BA** 

#### Study participating centre Royal Bournemouth Hospital Bcsc

Royal Bournemouth Hospital Castle Lane East Bournemouth England BH7 7DW

#### Study participating centre Sandwell and West Birmingham Hospitals NHS Trust

City Hospital Dudley Road Birmingham England B18 7QH

#### Study participating centre Musgrove Park Hospital

Musgrove Park Taunton England TA1 5DA

#### Study participating centre Scunthorpe General Hospital

Cliff Gardens Scunthorpe England DN15 7BH

#### Study participating centre The Royal Oldham Hospital

Rochdale Road Oldham England OL1 2JH

#### Study participating centre

Nottingham University Hospitals NHS Trust - Queen's Medical Centre Campus

Nottingham University Hospital Derby Road Nottingham England NG7 2UH

#### Study participating centre

Royal Liverpool University Hospital NHS Trust

Royal Liverpool University Hospital Prescot Street Liverpool England L7 8XP

#### Study participating centre Barnsley Hospital NHS Foundation Trust

Gawber Road Barnsley England S75 2EP

#### Study participating centre Hampshire Hospitals NHS Foundation Trust

Basingstoke and North Hampshire Hos Aldermaston Road Basingstoke England RG24 9NA

#### Study participating centre Kings College Hospital

Denmark Hill London England SE5 8AB

#### Study participating centre Kingston Hospital

Galsworthy Road Kingston upon Thames England KT2 7QB

#### Study participating centre Leeds Teaching Hospitals NHS Trust

St. James's University Hospital Beckett Street Leeds England LS9 7TF

#### Study participating centre Northampton General Hospital NHS Trust

Cliftonville Northampton England NN1 5BD

# Study participating centre The Dudley Group NHS Foundation Trust

Russells Hall Hospital Pensnett Road Dudley England DY1 2HQ

#### Study participating centre Mersey and West Lancashire Teaching Hospitals NHS Trust

Whiston Hospital Warrington Road Prescot England L35 5DR

# Study participating centre East Suffolk and North Essex NHS Foundation Trust

Colchester Dist General Hospital Turner Road Colchester England CO4 5JL

#### Study participating centre Whittington Health NHS Trust

The Whittington Hospital Magdala Avenue London England N19 5NF

#### Study participating centre

East Kent Hospitals University NHS Foundation Trust

Kent & Canterbury Hospital Ethelbert Road Canterbury England CT1 3NG

#### Sponsor information

#### Organisation

Medical University of Graz

#### **ROR**

https://ror.org/02n0bts35

#### Funder(s)

#### Funder type

Government

#### **Funder Name**

Health Technology Assessment Programme; Grant Codes: 17/147/33

#### Alternative Name(s)

NIHR Health Technology Assessment Programme, Health Technology Assessment (HTA), HTA

#### **Funding Body Type**

Government organisation

#### **Funding Body Subtype**

National government

#### Location

**United Kingdom** 

#### **Results and Publications**

#### Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request

(Updated 20/11/2020, previously: The datasets generated and/or analysed during the current study will be included in the subsequent results publication)

#### IPD sharing plan summary

Available on request

#### **Study outputs**

Output type
<u>Protocol article</u>

Details

Date created Date added Peer reviewed? Patient-facing? 12/11/2019 15/01/2025 Yes No

HRA research summary			28/06/2023 No	No
Participant information sheet	Participant information sheet	11/11/2025	11/11/2025 No	Yes
Study website	Study website	11/11/2025	11/11/2025 No	Yes