Vitamin D to prevent acute lung injury following oesophagectomy (VINDALOO): study protocol for a randomised placebo controlled trial

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Abstract

Background: Acute lung injury occurs in approximately 25% to 30% of subjects undergoing oesophagectomy. Experimental studies suggest that treatment with vitamin D may prevent the development of acute lung injury by decreasing inflammatory cytokine release, enhancing lung epithelial repair and protecting alveolar capillary barrier function.

Methods/Design: The ‘Vitamin D to prevent lung injury following oesophagectomy trial’ is a multi-centre, randomised, double-blind, placebo-controlled trial. The aim of the trial is to determine in patients undergoing elective transthoracic oesophagectomy, if pre-treatment with a single oral dose of vitamin D3 (300,000 IU (7.5 mg) cholecalciferol in oily solution administered seven days pre-operatively) compared to placebo affects biomarkers of early acute lung injury and other clinical outcomes. The primary outcome will be change in extravascular lung water index measured by PiCCO® transpulmonary thermodilution catheter at the end of the oesophagectomy. The trial secondary outcomes are clinical markers indicative of lung injury: PaO2:FiO2 ratio, oxygenation index; development of acute lung injury to day 28; duration of ventilation and organ failure; survival; safety and tolerability of vitamin D supplementation; plasma indices of endothelial and alveolar epithelial function/injury, plasma inflammatory response and plasma vitamin D status. The study aims to recruit 80 patients from three UK centres.

Discussion: This study will ascertain whether vitamin D replacement alters biomarkers of lung damage following oesophagectomy.

Trial registration: Current Controlled Trials ISRCTN27673620

Keywords: Acute lung injury, One lung ventilation, Oesophagectomy, Vitamin D

Background

Acute lung injury (ALI) and the more severe acute respiratory distress syndrome (ARDS) are common, devastating clinical syndromes of acute respiratory failure in the critically ill person. The incidence of ALI is 79 per 100,000 person years with a mortality rate of 30% to 65% [1]. Survivors of ARDS experience a significant reduction in health-related quality of life, with 46% reported to be unable to return to work within 12 months.

ALI is the final common pathway of response to a variety of direct pulmonary insults, such as bacterial/viral pneumonia and gastric aspiration, or indirect insults, such as abdominal sepsis or battlefield trauma. Only a relatively small proportion of patients develop ALI, with research suggesting that genetic, demographic (age), social (smoking, alcohol abuse) and other factors play a role in determining who develops ALI [1,2]. There are no current readily available tests that can clearly identify those who are at high risk of ALI and no therapeutic interventions proven to prevent its occurrence.
One lung ventilation (OLV) as a model for ALI/ARDS

To allow access to the oesophagus during surgery (using the transthoracic technique), one of the lungs is deflated and the subject is ventilated through the other lung. This is known as one-lung ventilation (OLV). There is a high postoperative incidence of ALI/ARDS [3-5] following OLV and unlike most insults leading to lung injury the delivery of OLV is predictably timed, thereby allowing serial studies to be carried out throughout the period of stimulus and development of the condition. Preoperative risk factors including age, respiratory function and cigarette smoking have been found to be related to the incidence of postoperative pulmonary complications [3,6-8]. It is unclear at present why only a percentage of patients undergoing OLV develop lung injury or why the lung injury typically occurs 24 to 48 hours after the cessation of OLV. Our data show that the development of lung injury is, however, associated with a doubling of in-hospital stay and elevated mortality.

We have extensively modeled the local and systemic inflammatory response to transthoracic oesophagectomy in 50 patients undergoing OLV. After OLV, patients have a neutrophilic alveolitis, with a significant alveolar and systemic inflammatory response. This is associated with the release of markers of both endothelial and alveolar epithelial dysfunction and an increase in the permeability of the alveolar barrier. This manifests clinically as increased extravascular lung water and a fall in oxygenation.

Alveolar levels of surfactant protein D and bronchoalveolar lavage fluid (BALF) protein permeability index are highest in those who develop ALI within 72 hours of OLV suggesting that peri-operative alveolar epithelial damage is a risk factor for the subsequent development of ALI. Immediate post-operative plasma markers of neutrophilic activation (myeloperoxidase, and matrix metalloproteinase-9 (MMP-9)) as well as the receptor for advanced glycation end-products (RAGE, a type I epithelial cell marker) are similarly raised in those who develop ALI within 72 hours of OLV. Proposed causative mechanisms for this injury include the ischaemic/reperfusion insult suffered by the collapsed lung, as well as oxidative stress and barotrauma causing epithelial injury to the ventilated lung [9]. These mechanisms are important in the pathogenesis of ALI making OLV a valid model for studying the pathogenesis of ALI in humans and exploring therapeutic strategies for preventing lung injury in a predefined subject population [10,11].

Vitamin D biology

Vitamin D₃, or cholecalciferol, is mainly formed in the skin after exposure to sunlight, then hydroxylated in the liver to 25-hydroxyvitamin D₃ (25(OH)D₃) and subsequently in the kidney to 1,25-dihydroxyvitamin D₃ (1,25(OH)₂D₃). When 25(OH)D₃ is sufficiently available, 24,25-dihydroxyvitamin D₃ (24,25(OH)₂D₃) is formed in the kidney which is further catabolised. Vitamin D metabolites are bound in the circulation to vitamin D binding protein (VDBP) which has a high affinity for 25(OH)D₃, 24,25(OH)₂D₃ and 1,25(OH)₂D₃ and, therefore, regulates free circulating concentrations of vitamin D metabolites. The biologically active metabolite 1,25(OH)₂D₃ can also be generated locally within tissues due to induction of extra-renal cyp27b1 (25(OH)D-1-alpha hydroxylase) and binds to the vitamin D receptor (VDR) resulting in modified gene expression.

Epidemiological studies have suggested a role for low vitamin D status in the risk of developing both viral and bacterial infection [12,13]. A recent study has further demonstrated the pleiotropic anti-inflammatory effects of vitamin D in patients with pulmonary tuberculosis [14]. Published data suggest that vitamin D deficiency is common in critically ill patients [15], and recent prospective studies suggest an association with increased morbidity and mortality [16-18]. Literature on acute vitamin D supplementation in critical illness is lacking but serious adverse events attributable to vitamin D supplementation are rare [18,19].

Is severe vitamin D deficiency a driver of post-OLV ALI?

Vitamin D has profound effects on human immunity acting as an immune system modulator, preventing excessive expression of inflammatory cytokines and increasing the ‘oxidative burst’ potential of macrophages, thereby enhancing bacterial killing. Vitamin D also stimulates the release of antimicrobial peptides such as LL-37 (cathelicidin) within the lung. LL-37 can also bind to and neutralize lipopolysaccharide (LPS), and functions as a chemoattractant for neutrophils, monocytes and T cells through a formyl peptide receptor-like molecule [20].

Respiratory epithelial cells convert 25(OH)D₂ to 1,25(OH)₂D₃ and activate VDR responsive genes increasing the production of hCAP18 from which LL-37 is cleaved within 24 hours [21]. In terms of the pathophysiology of ALI this could be important as LL-37 may drive epithelial repair responses as well as being an anti-microbial peptide [22]. Elevating local LL-37 may also be important as a downstream immunomodulator of vitamin D since it has recently been shown to reduce Toll-like receptor (TLR) agonist-mediated neutrophil-derived increases in IL-1β, IL-6, IL-8 and tumour necrosis factor-alpha (TNF-α) in addition to stimulating bacterial phagocytosis [23]. The ability of 1,25(OH)₂D₃ to directly inhibit nuclear factor-kappaB (NF-κb) signalling and suppress macrophage TLR expression suggests that vitamin D may also play a key role as a feedback regulator of macrophage responses [24,25].

The only study looking at vitamin D levels in patients with severe sepsis suggests that these patients have a lower serum vitamin D level than healthy control patients. This was associated with lower plasma levels of LL-37, 25-hydroxyvitamin D₃ and subsequently in the kidney to 1,25-dihydroxyvitamin D₃ (1,25(OH)₂D₃). When 25(OH)D₃ is sufficiently available, 24,25-dihydroxyvitamin D₃ (24,25(OH)₂D₃) is formed in the kidney which is further catabolised. Vitamin D metabolites are bound in the circulation to vitamin D binding protein (VDBP) which has a high affinity for 25(OH)D₃, 24,25(OH)₂D₃ and 1,25(OH)₂D₃ and, therefore, regulates free circulating concentrations of vitamin D metabolites. The biologically active metabolite 1,25(OH)₂D₃ can also be generated locally within tissues due to induction of extra-renal cyp27b1 (25(OH)D-1-alpha hydroxylase) and binds to the vitamin D receptor (VDR) resulting in modified gene expression.

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suggesting that this deficiency is of functional importance in vivo [26]. IL-1 and TNF production induced by TLR3 agonists from monocyte derived macrophages are inhibited to the same extent by 25(OH)D3 and 1,25(OH)2D3 after 24 - hours suggesting that the vitamin D metabolites may have a rapid anti-inflammatory action and that local intracrine activation of 25(OH)D3 can be anti-inflammatory [27].

Recent data have further implicated vitamin D in adaptive immunity because of its influence upon the differentiation of T cells between the regulatory T cell (Treg) and the pro-inflammatory T helper 17 (Th17) subsets [28-30]. Treg, Th17 cells are known to stimulate tissue inflammation and neutrophil chemotaxis, both of which are seen in ALI, predominantly by IL-17 production. It also appears that expression of markers of Treg cells (Foxp3) or Th17 cells (IL-17) by T cells may not be stable and that there is a greater degree of plasticity in their differentiation than previously appreciated [29]. Recent evidence has further suggested Treg cells are important in the resolution of experimental ALI. This suggests that local lung regulation of the balance between Treg and Th17 cells may be a determinant of resolution/persistence of neutrophilic inflammation which is known to be associated with a poor prognosis in human ALI.

Although the above suggest a potentially beneficial effect of vitamin D, we must exercise some caution as some studies have shown potentially adverse effects of vitamin D. In low dose nasal LPS challenge, cellular inflammation is actually lower in vitamin D receptor deficient (VDR KO) mice due to toll-receptor hypersensitiveness. In addition the chemotactic effects of LL-37 could in theory increase neutrophil recruitment to the lung; albeit in ALI, CXCL-8 and ENA-78 are the main chemokines driving neutrophil recruitment. More recently it has been shown that the plasma LL-37 concentration was decreased in vitamin D supplemented patients with tuberculosis, possibly representing a global suppressive effect of vitamin D supplementation on markers of acute phase response or an indirect response to enhanced microbial killing [14]. Despite these reservations, the predominant biological effects of vitamin D led us to hypothesise that vitamin D deficiency may be a risk factor for ALI, causing elevated inflammation which results in exaggerated epithelial damage in at risk, vitamin D deficient individuals.

The VINDALOO trial is a three centre randomised double blind, placebo-controlled trial aiming to define the safety and effectiveness of a single high dose of vitamin D in preventing ALI in a group of patients at high risk of developing the condition.

Methods/Design

Clinical trial registration

The trial is registered on the International Standard Randomised Controlled Trial Registry (ISRCTN27673620). The sponsor organisation for the trial is the University of Birmingham. The trial is funded by the Medical Research Council UK (MRC reference G1100196). The trial will be carried out in accordance with the Medical Research Council (MRC) Good Clinical Practice Guidelines, applicable UK legislation and Standard Operating Procedures of the Peri-operative and Critical Care Trials Group at the University of Birmingham. The trial will be reported in line with the Consolidated Standards of Reporting Trials (CONSORT) 2010 guidelines [31].

Outcome measures

Primary outcome

The primary outcome will be the extravascular lung water index (EVLWI) measured by PiCCO thermodilution catheter at the end of the oesophagectomy (measured within one hour post-operatively). In ALI the changes in lung compliance that are a cardinal feature of this disease occur due to the accumulation of extravascular lung water (EVLW). The PiCCO EVLWI has been shown to be an independent risk factor for mortality in ALI and has been used as the primary outcome in several clinical trials in ALI (BALTI-1, HARP) [32,33] as well as post-thoracotomy [34]. In BALTI-1 we demonstrated that the transpulmonary thermodilution technique (PiCCO) is able to detect a significant change in lung water with a treatment that might be expected to achieve this objective. We have tested the CV coefficient of variance of EVLWI measurements previously and found it to be 6.8% over six sequential assessments over two hours. Further, we have studied the perioperative changes in lung water following oesophagectomy and demonstrated that preoperative vitamin D status influences the level of accumulation of EVLW.

Secondary outcomes

The trial secondary outcomes are clinical markers indicative of lung injury: PaO2:FiO2 ratio, oxygenation index, development of lung injury/ARDS during the first 28 days, ventilator and organ failure free days, survival (28 and 90 day) and safety and tolerability of vitamin D supplementation.

Lung injury will be defined by the American European Consensus Conference definition [35] as the acute onset of: (1) bilateral infiltrates on the chest x-ray; (2) hypoxaemia with a PaO2:FiO2 ratio of <40 kPa; and (3) absence of clinical evidence of left atrial hypertension. Ventilator free days are defined in accordance with the ARDSnet criteria as the number of calendar days after initiating unassisted breathing to day 28 after randomisation, assuming a patient survives for at least 48 consecutive hours after initiating unassisted breathing [36]. Un-assisted breathing is defined as one of at least 48
consecutive hours of: (1) being extubated with face mask, nasal prong oxygen, or room air; (2) T-tube breathing; and (3) tracheostomy mask breathing, CPAP = 5 cm H2O with- out pressure support of intermittent mandatory ventilation assistance.

Organ failure free days are defined in a similar manner to ventilator free days with an organ failure free day being a day without evidence of non-respiratory organ failure. Organ failure will be defined as a sequential organ failure assessment (SOFA) score of greater than three [37].

Plasma indices of endothelial and alveolar epithelial function/injury, plasma inflammatory response will be measured by ELISA. Plasma vitamin D status (25(OH)D3, 1,25(OH)2D3, VDBP) and calcium will be assessed peri-operatively (pre-drug dose, pre-operative, post-operative, day 1 and day 3) as well as EVLWI day 1 post-operatively (measured at 9 am on day 1).

**Eligibility criteria**

Patients will be eligible for the trial if they fulfill the following criteria:

- Planned transthoracic oesophagectomy for oesophageal carcinoma at a participating centre.
- Men over 18 years old on the day of first dose of the investigational medicinal product (IMP).
- Women over the age of 55 or more than 2 years since menopause.
- Women of potential child bearing age (under 55 and less than two years since menopause) may be recruited provided they agree to use contraception during the pre-post-operative period (eight weeks).
- Ability to give written informed consent to participate in the study.

Patients fulfilling any of the criteria below will be excluded:

- Known intolerance of vitamin D.
- Known sarcoidosis, hyperparathyroidism, or nephrolithiasis.
- Known serum corrected calcium >2.65 mmol/L.
- Undergoing haemodialysis.
- Pregnant or breastfeeding.
- Patients with tuberculosis or lymphoma.
- Diagnosis of chronic obstructive pulmonary disease (COPD) with a forced expiratory volume in one second (FEV1) less than 50% predicted or resting oxygen saturations of less 92%.

**Power and sample size estimate**

There are no direct data to predict the effect size of vitamin D replacement upon EVLWI. In our preliminary work with 50 patients, the patients with 25(OH)D3 concentrations less than 15 nmol/L had the greatest increases in EVLWI (+3.2 ml/kg, +27%) compared to those patients with less severe deficiency pre–postoperatively (+1.0 ml/kg, +10% \( P = 0.013 \)) suggesting that severe vita- min D deficiency influences EVLWI. As a group, after undergoing oesophagectomy our patients have a mean post-operative EVLWI of 10.1 ml/kg and standard devia- tion of 2.9 ml/kg. For the study to be able to detect a change of 20% in EVLWI with a power of 80% we will require approximately 34 patients in each arm to reach the primary endpoint \( P = 0.05 \). An additional six patients will be needed to allow for dropouts, such as open/close cases, unexpected deaths and other difficulties with data collection. Thus, we intend to recruit 40 patients to each arm of the study.

With 34 patients completing each arm of the study, based upon preliminary unpublished data from 50 oesophagectomy cases, this study would detect a treatment effect of vitamin D upon the PO2:FiO2 ratio postoperative (PO) of ≤ 8.16 (± 20%), PO plasma soluble intercellular adhesion molecule-1 (sICAM-1) of ≤ 16 ng/ml (± 32%), plasma C-reactive protein (CRP) ≤ 53 ng/ml (± 33%), PO plasma von Willebrand factor (vWF) ≤ 55% relative to control plasma (± 24.7% change), PO plasma high-mobility group box 1 (HMGB-1) 5.42 ng/ml (± 54%), PO plasma myeloperoxidase (MPO) 105 pg/ml (± 57%), PO plasma MMP-9 51 ng/ml (± 57%), and PO plasma surfactant protein-D (SP-D) 691 ng/ml (± 61%). All calculations assume 80% power at a two-tailed significance level of 0.05.

**Trial conduct**

*Approach to patients and obtaining informed consent*

Patients will be identified through upper gastrointestinal cancer teams. Eligible patients will be invited to participate by their treating clinician, specialist clinical nurse or research nurse. If agreeable, written informed consent will be obtained, following a face to face discussion about the study.

**Randomisation and drug / placebo supply**

The trial drug manufacturer will produce the randomisation sequence using a block size of 10 with equal allocation between active and placebo groups. Drug (oral cholecalciferol oily solution Vigantol, 300,000 IU (7.5 mg)) and matching placebo (Miglyol 812 oil, the vehicle for vitamin D3 in Vigantol) will be supplied and packaged according to the randomisation sequence into numbered treatment boxes by Novalabs (Leicester, UK). Drug boxes will be supplied to centres in blocks of 10 thus ensuring an equal allocation between active and placebo groups to balance any differences in case mix, pre-operative, operative and post-operative care between centres. Patients will be randomised sequentially by allocating them to the next numbered treatment pack held at the centre.
Drug administration
Subjects will receive either 300,000 IU (7.5 mg) of vitamin D or placebo seven days prior to planned oesophagectomy. The drug will be administered by qualified medical or nursing staff.

Concomitant medications
Patients taking the following medications are not eligible for inclusion in the study:

- Taking more than 1,000 IU/day (25 mcg/day) vitamin D supplementation by month preceding enrolment.
- Taking cardiac glycoside, carbamazepine, phenobarbital, phenytoin, primidone or long-term immunosuppressant therapy.
- Patients taking benzothiadiazine derivatives at doses higher than that which is recommended in the British National Formulary (BNF).
- Patients taking a benzothiadiazine derivative in combination with a calcium supplement.

Post randomisation withdrawals and exclusions
Subjects may withdraw from the trial or the trial treatment at any time without prejudice. If a subject withdraws from the trial treatment, then they will be followed-up wherever possible and data collected as per protocol until the end of the trial. The only exception to this is where the subject also explicitly withdraws consent for follow-up.

Blinding/un-blinding
Patients, clinical and research / trial staff will be unaware of the arm of the study to which a patient is allocated. Active and placebo treatment packs and their contents will be identical in appearance. The protocol allows for emergency un-blinding in the event of significant concerns about patient safety. In the unlikely event that un-blinding is required the local investigator will discuss this with the Chief Investigator. All events will be logged.

Monitoring and reporting adverse events
VINDALOO is recruiting a population who are prone to recognized medical and surgical complications. It is expected that many of the patients will experience an event that might be seen as a serious adverse event but is a recognized complication following oesophagectomy. Adverse and serious adverse events which are recognised complications of surgery, for example, medical (pneumonia, sepsis) and surgical (chyle leak, anastomosis leak) will be recorded in the case report form.

Death and other serious adverse events thought to be related to the study drug or serious unexpected serious adverse reactions will be reported to the trial coordinating centre and Chief Investigator within 24 hours of becoming aware of their occurrence. The Chief Investigator will inform the sponsor and regulatory authorities.

Data collection
Data up until hospital discharge will be recorded in each subject’s Case Report Form (CRF) by a member of the trials team. Most of the data collected will be obtained from the patient’s hospital notes. In the unlikely event that a subject is transferred to another hospital, the study team will ensure that data collection is completed by the receiving hospital.

If the subject remains in hospital at 28 or 90 days, survival at these time points will be recorded by hospital staff. Mortality after hospital discharge will be obtained from the National Health Service (NHS) Statistical Tracing Service (NSTS).

Statistical analysis plan
Data will be analysed with the help of the trial statistician. Data will be analysed using SPSS for Windows 17.0. A detailed analysis plan will be developed during the trial prior to commencement of analysis. In brief, for continuously distributed data, differences between groups will be tested using independent samples t-tests with transformations of variables to Normality if appropriate, or non-parametric equivalents. Chi-squared tests (or Fisher’s Exact tests) will be used for categorical variables. A P value of 0.05 will be considered as significant. We will test for significant correlations between changes in the biological markers using standard methods. The treatment effect will be analysed on an intention-to-treat basis. For further examinations of relationships between a binary variable and known explanatory variables, the following tests will be applied as appropriate. Logistic regression will be used to provide the estimated risk ratios for the treatment effect with associated 95% confidence interval (CI). Time to event outcomes, such as duration of ventilation or duration of hospital stay, will be analysed by survival methods and reported as hazard ratios and 95% CI. A single final analysis is planned at the end of the trial.

Trial organisation/oversight
Trial oversight will be provided by a Trial Steering Committee (TSC) comprising investigators, clinicians and trialists. An independent data monitoring committee will monitor the safety of participants enrolled in the trial through regular review of adverse event reports. An interim analysis of efficacy is not planned.

Discussion
The preliminary data that this study is based upon suggests a significant role for vitamin D deficiency as a risk factor for the development of acute lung injury. This
trial will look at biomarkers of alveolar epithelial damage and inflammation to determine the proof of concept that vitamin D replacement may have efficacy as a preventative agent for the development of acute lung injury.

Trial status
Patient recruitment commenced in September 2012 and is expected to run for two years (last patient recruited in August 2014).

Abbreviations
ALL: Acute lymphoid malignancy; ARDS: Acute respiratory distress syndrome; BAL: Bronchoalveolar lavage; CI: Confidence interval; DBP: Vitamin D binding protein; ELISA: Enzyme-linked immunosorbent assay; EVLWI: Extravascular lung water index; IL: Interleukin; IMP: Investigational Medicinal Product; ISRCTN: International Standard Randomised; MMP-9: Matrix metalloproteinase-9; MRC: Medical Research Council; OLV: One lung ventilation; PICCO: Pulse Contour Cardiac Output Monitoring; RAGE: Receptor for advanced glycation end-products; REC: Research Ethics Committee; TNF-α: Tumour necrosis factor alpha; Th: T helper cells; Treg: Regulatory T cells; VDBP: Vitamin D binding protein; VDR: Vitamin D receptor; 25(CH)D: 25-hydroxyvitamin D.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
All authors made a substantial contribution to the protocol development. All authors have read and approved this manuscript.

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