TITLE PAGE

Title: The association between critical illness and changes in bone turnover in adults: A systematic review

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Competing Interests: Neil Orford, Claire Cattigan, Sharon L Brennan, Mark A Kotowicz, Julie Pasco, and D James Cooper declare no conflict of interest

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ABSTRACT
Purpose: Intensive care patients face health issues that extend beyond their critical illness and result in significant morbidity and mortality. Critical illness may result in altered bone turnover due to associated immobilisation, inflammation, exposure to medications that effect bone and calcium metabolism, and endocrine dysfunction. The aim of this study was to synthesise the existing evidence for altered bone turnover in adults admitted to Intensive Care.

Methods: A literature search using MEDLINE and EMBASE was performed from 1965 to March 2013. Reviewed studies investigated the relationship between critical illness and evidence of altered bone turnover (bone turnover markers, bone mineral density, or fracture). Studies were rated upon their methodological quality, and a best-evidence synthesis was used to summarise the results.

Results: Four cohort and seven case-control studies were identified for inclusion, of which five studies were rated as being of higher methodological quality. Ten of the studies measured bone turnover markers, and one study fracture rate. Findings were consistent across studies, and best-evidence analysis resulted in a conclusion that moderate evidence exists for an association between critical illness requiring admission to intensive care and altered bone turnover.

Conclusion: A positive association between critical illness requiring intensive care admission and bone turnover exists, although data are limited, and risk factors and the nature of the relationship are not yet understood. Prospective cohort studies that identify risk factors and extent of critical illness related bone turnover changes are required.

Key Words: Critical care, intensive care, bone turnover, mechanical ventilation, osteoporosis

Word count: 3890

Mini Abstract: Critical illness may lead to altered bone turnover and associated adverse health outcomes. This systematic review found moderate evidence for a positive association between critical illness and increased bone turnover. Prospective cohort studies that identify the extent and risk factors for critical illness related bone loss are required.
INTRODUCTION

Intensive care patients face health issues after their critical illness including increased mortality, reduced quality of life, reduced return to work, and ongoing economic and social costs to families and caregivers when compared to pre-illness and general population controls [1-7]. Despite an increasing awareness of long-term sequelae of critical illness, the identification of specific pathophysiology amenable to intervention remains elusive. Osteoporosis is a major public health problem, and is widely recognised as a chronic progressive disease with multifactorial aetiology [8]. The epidemiology and risk factors for primary and secondary osteoporosis are well described, including increasing age, female gender, low body mass index (BMI), smoking, excessive alcohol intake, positive family history, medications such as glucocorticoids, and predisposing disease or medical condition such as hyperthyroidism [8,9]. However as few as 13-27% of patients with osteoporosis are treated following a fragility fracture, suggesting it remains an under diagnosed disease [10,11]. Critical illness, with its associated immobilization, inflammation, and endocrine dysfunction, may lead to accelerated bone turnover. When combined with an ageing population with undiagnosed osteoporosis, this accelerated bone turnover may contribute to the burden of morbidity and mortality observed in survivors of intensive care [12,13]. In a review of metabolic bone disease in the intensive care unit (ICU), Hollander et al concluded an interventional study of bisphosphonates in survivors of ICU is needed [14]. However, there is no systematic review of the evidence for accelerated bone turnover following critical illness, or of the nature and risk factors for this relationship.

In this study we sought to systematically review and synthesise the current literature regarding the association between critical illness and changes to bone turnover. In addition we describe ongoing and planned research in this area.

METHODS

Search strategy and study selection
To identify studies that examined whether admission to an ICU was associated with changes in bone turnover in adults, we searched MEDLINE (1965 until 31st March 2013) and EMBASE (1974 until 31st March 2013) for citations of relevant articles. Our computerised search strategy employed the following medical subject headings (MeSH/EMTREE); (“mechanical ventilation” OR “critical illness” OR “chronic critical illness” OR “ventilator” OR “critical care” OR “intensive care”) AND (“bone turnover” OR “bone change” OR “fracture” OR “bone mineral density” OR “bone density” OR “bone loss” OR “CTX” OR “P1NP” OR “bone biomarkers” OR “bone markers”).
Studies were deemed eligible for inclusion in this review if they met the following criteria: full-text original articles; comprised either a cohort, case-control, or a cross-sectional study design; examined, in adults aged ≥18 years, associations between receiving mechanical ventilation in an ICU (with a length of stay 24 hours or greater), and de novo change in bone turnover (defined as loss of BMD, increase in bone turnover markers (BTMs), or incident fracture of at least one of the major osteoporotic sites of hip, wrist, humerus, or spine). Patients that were identified as osteoporotic at the time of ICU admission were included.

Studies were excluded if: published in languages other than English; utilized animal models; investigated patients with existing neurological illness that results in impaired weight bearing (including stroke with loss of weight-bearing, spinal cord injury, progressive neurological disease eg multiple sclerosis); were admitted to ICU for reasons of trauma-related fracture, or with existing metabolic bone disease; employed qualitative methodology; or were review articles, editorials, commentaries, dissertations, or were randomised control trials. Where interventional studies reported baseline data that fulfilled inclusion and exclusion criteria prior to intervention, the baseline data were included in the analysis.

We electronically restricted our search to identify articles that were: related to human subjects, published in English, and available in full-text. Reference lists of relevant studies deemed eligible for inclusion were manually searched, and citations were tracked for those publishing in the field of interest.

Two reviewers confirmed the search strategy (NRO and SLB) and one reviewer performed the computerized search and initial manual search (NRO). Complete details of the search strategy can be obtained from the corresponding author. For each eligible study, two reviewers (NRO and CEC) confirmed the selection of articles based on readings of the full text article. Where the eligibility of studies was ambiguous, two reviewers (NRO and CEC) held discussions to reach consensus. Where consensus could not be achieved, a third reviewer was consulted (SLB).

Methodological Quality Assessment
To assess the methodological quality (internal validity) of the included studies, two reviewers (NRO and CEC) undertook independent scoring using an adapted version of the scoring system published by Lievense et al [15] (Table 1); this methodological approach has previously been employed for reviews of observational studies in the field of musculoskeletal disorders [15,16]. Both reviewers independently scored each of the criteria as positive (1), negative (0), or not applicable (NA), with a maximum possible score for each study design of 100%. Where the score afforded to certain criteria differed between the reviewers, discussion
was held to achieve consensus; if disagreements were not resolved, a third reviewer (SLB)
was consulted to achieve a final judgment. Positive scores were summed to give an overall
internal validity score.

Data Analysis
As there were limited data and studies were heterogeneous, a “best-evidence” synthesis was
preferred rather than a meta-analysis. The studies were divided into subgroups according to
the type of study design. A cohort study was judged the most valid design, followed by case-
control study. Studies were then ranked according to their methodological quality score (Table
2). A study was considered to be of higher quality if the methodological quality score was
greater than the mean quality score of all studies [15-17]

RESULTS

Search Results and Study Characteristics
The results of our search are presented in Figure 1. Our electronic search strategy identified
13,185 studies, including 2,218 duplicates that were subsequently excluded. Of the remaining
10,967 studies, a total of 21 underwent full-text review to determine eligibility. A further six
studies were identified for full-text review from a manual search of citation lists. Of the 27 full-
text studies reviewed, 16 were excluded for not meeting the predetermined eligibility criteria,
resulting in 11 studies included in the final analysis [18-28]. In three of the case-control
studies, ICU patients were randomised to an intervention after comparison of baseline data
from the ICU cohort was compared to a control cohort [22,23,26]. The baseline data from
these studies was included in the analysis, whereas the data resulting from the randomised
intervention was excluded. One study performed in-vivo analysis of osteoclast number and
activity in ICU patients compared to controls, with further in-vitro analysis of osteoclast cells,
estoblastic cells, and serum activation factors [25]. As the in-vitro tests were not recognised
tests of bone turnover, they were excluded from the analysis.

Description of the studies
An overview of the included articles (n=11) is presented in Table 3. Four of the studies were
cohort study design [18-21] and the remaining seven were case-control study design [22-28].
Sample sizes ranged from nine [28], to 739 cases [18]. A group of seven patients were
shared by two studies, in which a total of 15 cases [23] and 33 cases [22] were enrolled. In
addition, these studies shared the same group of 50 controls.

The criteria for enrolment in the studies included requirement for mechanical ventilation in an
ICU [20,27,28], duration of mechanical ventilation greater then 48 hours [18] or 2 weeks
[22,23,24], chronic critical illness [25], admission to a respiratory care unit for prolonged


ventilatory support [19,21], ICU length of stay greater than 10 days [26]. Patient populations included mixed adults in nine of the studies [18-21,24-28], and males only in two studies [22,23]. A control population was present in the seven case-control studies, and in one of the cohort studies [18]. The controls were age and gender-matched healthy population-based participants in six studies [18,22-26], age but not gender-matched healthy controls in one study [28], and non-gender or age-matched participants with a history of rheumatism or mild osteoarthritis in one study [27]. Exclusion criteria included renal, metabolic, and liver disease in five studies [22,23,24,26,28], metabolic and neurological disease in four studies [22,23,24,26], and prior medications in four studies [22,23,24,26]. The follow-up time for patients ranged from 1 day [24,25], to 10 years [18].

Methodological quality assessment
The two reviewers scored 129 items and agreed on 117 items (90.70% agreement, $\kappa = 0.80$, standard error (SE) 0.05, 95% CI 0.70-0.91, strength of agreement considered “very good”). The 12 disagreements were resolved in a single consensus meeting. The range of methodological scores was 55% to 67% (Table 4), with the mean of quality scores 61%. Using the mean score as the cut off point, 5 of 11 were considered to be of higher methodological quality [18,19,22-24].

Results of all included studies
Assessment of bone turnover
Table 4 presents the findings of the reviewed studies. The BTM used in the studies included bone resorption and bone formation markers [29]. The bone resorption markers measured included urinary collagen type 1 cross-linked N-telopeptide (NTX) in two studies [19,21], urinary pyridinoline (Pyd) or deoxypyridinoline (Dpd) in seven studies [20,22,23,26-28], and urine or serum carboxy-terminal cross-linked telopeptide of type 1 collagen (ICTP/BCTX) in two studies [20,26]. One study reported serum osteoclast precursors (double-positive CD14+/CD11b+) and serum mature osteoclasts (triple-positive CD14+/CD11b+/VNR+) [25]. Bone formation markers were reported in four studies [22-24,26] and included serum skeletal alkaline phosphatase (SALP), serum osteocalcin (OC), serum procollagen type 1 C peptide (P1CP) and serum procollagen type 1 N peptide (P1NP). Incident fracture rate post ICU discharge was reported as the outcome in one study [18].

Results of bone turnover measurement
All studies that measured markers of bone resorption reported an increase in markers compared to controls or reference range, suggesting increased osteoclastic activity. The two studies observing urinary NTX levels in patients admitted to a respiratory care unit (prolonged ventilation unit) reported elevated NTX in 83% of patients [21], with baseline levels 4 to 6-fold greater than reference range [19,21] while urinary collagen cross-links (Pyd, Dpd) were
increased 4 to 14-fold compared to controls [20,22-24,26-28], and serum carboxy-terminal cross-linked telopeptide of type 1 collagen was increased 3 to 6-fold compared to controls or reference values [20,26]. The one study that measured osteoclast precursors and mature osteoclasts in serum described a significant increase in osteoclast precursors in critical illness compared to controls [25].

The studies that reported markers of bone formation described a varied increase in SALP compared to controls, a significant increase in P1CP and P1NP compared to controls, and a significant decrease in osteocalcin compared to controls [22-24,26]. These results suggest an increase in number and activity of immature osteoblasts, with low activity of mature osteoblasts.

The single study that reported the incidence of new fractures following ICU, described an increased incident fragility fracture risk in older female ICU survivors (rate 4.33/100 patient years, 95% CI 2.72-5.93) compared with age- and gender-matched population controls (rate 2.81/100 patient years, 95% CI 2.33-3.28) [18].

Exposure variables and increased bone turnover
Three studies reported a relationship between bone resorptive markers and either ICU or hospital length of stay, with a positive correlation between urinary NTX and ICU length of stay (r=0.42, p<0.01) [19], urinary NTX and both hospital (r=0.49,p<0.01) and ICU length of stay (r=0.42,p<0.01) [21], and increased urinary Pyd and Dpd in patients with an ICU length of stay of 5-days or greater compared to less than 5-days [28]. The relationship observed between vitamin D, parathyroid hormone, or calcium status was variable [19,20]. An association between bone formation markers and inflammatory markers was observed in three studies [22-24], an inverse correlation between bone resorptive markers and thyroid hormones in one study [24], and no correlation between nitric oxide breakdown products and bone resorption markers in one study [27]. The two studies that compared ICU patients with sepsis to other ICU cohort reported an increase in bone resorption markers in sepsis compared to trauma [27] and surgery [20].

Best-evidence analysis
As the reviewed studies employed different methodology, had recruited diverse populations (for example, differences in ages, gender and population sizes, among other factors), and examined varying follow up times, we performed a best-evidence analysis to cater for the high level heterogeneity. Our best evidence synthesis included studies that scored above the mean (>61%) for their methodological quality. Of the eligible studies, two cohort [18,19], and three case-control studies [22-24] were considered to be of higher methodological quality.
The higher-quality cohort studies described an increase in bone resorption markers, with a positive correlation between markers and duration of ICU stay [19], and an increase in fragility fracture in older women following ICU admission compared to age and gender matched healthy controls [18]. The higher-quality case-control studies described an increase in bone resorption, a pattern of increased bone formation consistent with an increased number and activity of immature osteoblasts and decreased activity of mature osteoblasts, and a correlation between inflammatory cytokines and bone resorptive activity [22-24]. These results are consistent with findings of the lower-quality studies.

In summary the result of our best evidence analysis is five of eleven studies (two cohort and three case-control) were considered of higher methodological quality, with consistent results. This is consistent with a conclusion that moderate evidence exists for an association between critical illness requiring intensive care admission and changes in bone turnover.

**DISCUSSION**

Overall, we identified limited but consistent data that examine the relationship between critical illness requiring ICU admission and bone turnover. A best evidence analysis of available literature provides moderate evidence for a positive association between critical illness requiring ICU admission and increased bone turnover, a finding in all the studies identified by this analysis. There are insufficient high-quality data available about the factors contributing to the relationship between ICU admission and bone turnover to allow interpretation of the nature of this association.

This review included studies with patients admitted to ICU for mechanical ventilatory support for greater than 24-hours, and assessed measures of bone turnover following ICU admission. We chose a relatively inclusive definition of critical illness for this review, as although chronic or prolonged critically ill are more likely to be at risk of increased bone turnover, the definition for chronic critical illness remains ambiguous [31], and the relationship between critical illness and bone metabolism relatively unexplored.

The criteria for new bone turnover used in this review included BMD assessment, BTMs, and incident fracture. Measurement of BMD remains the primary tool for fracture risk and osteoporosis treatment assessment, and is the central component of internationally agreed definitions of osteoporosis [32]. An important limitation of the studies identified by this analysis is that none report changes in BMD during or following critical illness. A single study reported an increased in fragility fracture risk in older females following ICU compared to population controls, and although this was a large study with a high methodological quality score, it was limited by its retrospective design [18]. There are no studies reporting the use of bone
histomorphometric analysis of bone biopsies or other methods for assessing bone microarchitecture (high resolution CT or MRI imaging) in critically ill patients. Bone histomorphometry could provide information regarding the effects of critical illness on microarchitectural deterioration, mineralisation and dynamic indices of bone resorption, and formation.

Bone turnover markers were the outcome measured in ten of the eleven studies identified in this review. BTMs are an important tool to assess progression of osteoporosis, fracture risk and treatment response [29,33,34]. Overall BTMs are separated into markers of bone resorption (PyD, DpD, B-CTX/ICTP) and bone formation (ALP, BALP, OC, P1CP, P1NP) [35]. However BTM levels are affected by a number of factors, requiring more complex interpretation. Osteocalcin is a marker of osteoblast function and bone formation, but smaller fragments are derivatives of bone resorption and included in assay. The bone formation markers P1NP and P1CP are both procollagen terminal extension peptides, but P1NP is more specific for bone formation. Also a number of BTMs are affected by biological factors including age, gender, co-existing disease, and medications [35]. Examples include decreased excretion of B-CTX in renal failure [35] and sensitivity of OC to glucocorticoid exposure [36].

The studies identified in this review consistently described changes in BTMs during critical illness suggestive of increased osteoclastic bone resorption (increased urinary DpD and PyD, serum B-CTX/ICTP), an increase in immature osteoblast number and activity (serum P1CP and P1NP), and reduced activity of mature osteoblasts (serum OC and ALP). The increase in bone resorption markers described in these studies is of the magnitude described in postmenopausal females, or metabolic bone disease [34,37,38], and has been likened to other metabolic bone disorders, such as Paget's disease, where uncoupling of bone osteoclast and osteoblast activity are described [26].

A limitation of the studies using BTMs to assess bone turnover in this analysis was the short duration of follow-up, ranging from 1-26 days, and a lack of premorbid assessment if bone turnover or skeletal health. When BTMs are used to assess treatment effect of anti-resorptive agents an interval of 3-6 months is normally recommended [36]. Although these studies are not designed to assess the effect of anti-resorptive agents on bone turnover, the short duration of follow-up decreases the ability to establish a causal relationship between critical illness and bone turnover.

An important limitation of the evidence identified by this review is the limited analysis of the effect of possible confounding variables on the association between critical illness and altered bone turnover. Although six of the studies provided an age and gender matched assessment
of a control group, the effect of other known causes of osteoporosis and variables known to affect the metabolism of BTMs (including menopausal status, renal failure, liver disease, diabetes, thyroid disease and medications) [37,38], were not consistently addressed. These variables are likely to occur in critically ill patients, leaving the possibility that altered metabolism of BTMs or known risk factors for osteoporosis are partly or wholly responsible for the observed increase in bone turnover.

The studies in this analysis do provide some information about the relationship between critical illness duration, inflammation, immunomodulation, endocrine dysfunction, and increased bone turnover. Higher levels of bone resorption markers were observed in ICU patients with a length of stay of greater than 5-days compared to less than 5-days [28]. This may indicate a relationship between duration of critical illness and bone resorption, although the lack of adjustment for confounders, including co-morbid illness such as renal failure, prevents the nature of this relationship being established.

Vitamin D deficiency with resultant secondary hyperparathyroidism and prolonged immobilisation may increase the risk of excessive bone resorption; however a range of metabolic abnormalities characterised as primary hyperparathyroidism, secondary hyperparathyroidism, and mixed disorder were described in critically ill patients with elevated bone resorption markers [19]. Two studies report the effects on bone turnover of treating vitamin D deficiency in critically ill patients. The interventional data from one study in this analysis described the effect of parenteral vitamin D 200 IU or 500 IU daily in long-term surgical ICU patients receiving parenteral nutrition. Higher dose vitamin D was associated with a relatively small increase in serum OC, and a decrease in serum B-CTX, but did not affect other BTMs. In addition the decrease in inflammatory markers interleukin-6 and C-reactive protein over time was more pronounced with the higher dose vitamin D [26]. However treating vitamin D deficiency with calcitriol did not lead to a reduction in bone resorption markers, suggesting that vitamin D deficiency was not the mechanism for accelerated bone turnover [21].

A positive relationship between inflammation and increased bone turnover was present in a number of studies [22-24,26], and was unrelated to severity of illness, type of illness, age or outcome [26]. Systemic inflammation has been identified as a marker for increased fracture risk in non-critically ill patients, with a 23% increase risk of fracture associated with each standard deviation increase in the inflammatory marker high-sensitivity C-reactive protein [39]. Inflammatory cytokines such as TNF-α and IL-6 are potent stimulators of RANK-ligand mediated activation of osteoclastogenesis and direct activation of osteoclast precursors [26]. Ongoing bone resorption did not correlate with inflammatory markers, which may reflect the influence of other mechanisms, a prolonged effect of cytokines through osteoclast activation
factors that increase maturation and lifespan of osteoclasts, or a direct effect of cytokines on osteoclast precursors. In one of the studies, concomitant treatment with glucocorticoids, thyroid hormones, or any other ICU medication did not significantly affect markers of bone turnover at any of the studied time points [26].

A series of studies included in this review by Van den Berghe et al [22-24], described changes to the somatotrophic, thyrotrophic, and gonadotrophic axes in prolonged critical illness, and included bone markers as a part of measures of target tissue effects. The studies describe a positive correlation between inflammatory cytokines and osteoclastic and osteoblastic activity, with variable effects of restoration of somatotrophic, thyrotrophic, and gonadotrophic axes on BTMs. The administration of growth hormone releasing peptide (GHRP) alone led to reactivation of pulsatile GH secretion in critically ill patients, but no changes in BTMs [22]. The addition of thyroid releasing hormone (TRH) led to increased osteocalcin, suggesting impaired maturation of osteoblasts may be explained by a suppressed thyroid axis [22]. Finally the addition of gonadotropin releasing hormone (GnRH) led to a further increase in osteocalcin [22,23]. This complex relationship between sex hormones and altered bone turnover markers in critical illness is not surprising given the increasingly complicated interaction between these regulators of osteoclast differentiation and activity [40].

In-vitro experiments have shown that compared to healthy controls, critically ill patients peripheral blood mononuclear cells (PBMCs) responded to the presence of osteoclastic activation factors with an increased number and activity of mature osteoclasts [25]. In addition, exposure of PBMCs to critically ill patient sera resulted in an increased formation of mature osteoclasts, whereas a model of bone formation showed a reduction in angiogenesis factor expression, and reduced vascularity and maturity of bone formation.

This systematic review provides moderate evidence of a relationship between critical illness and increased bone turnover. Increased bone turnover may lead to impaired fracture healing or post-ICU fragility fractures, with their associated morbidity and mortality. Increased bone turnover is associated with mortality in elderly patients [41] and patients with cardiovascular disease [42]. If future studies find that survivors of critical illness are at high risk of subsequent fragility fracture, target interventions to prevent or attenuate acute bone loss such as the early administration of antiresorptive therapies may be assessed as a broader fracture prevention strategy.

There is limited evidence examining the efficacy of bisphosphonates in this setting. A randomised controlled trial identified in the search strategy and excluded from this analysis, reported a transient decrease in serum B-CTX in chronic critically ill patients receiving a
single intravenous dose of ibandronate compared to placebo [43]. In addition the decrease in the bone turnover marker (serum OC) observed in postmenopausal women receiving ibandronate [44] was not observed in this study, supporting the theory that bone formation and resorption is uncoupled in critical illness. Although limited by small sample size, short follow-up, and limited extent and duration of effect, this study provides evidence that suppression of excessive bone resorption in critical illness is possible.

The higher-quality cohort and case-control studies, provide moderate evidence for an association between critical illness requiring intensive care admission, and increased bone turnover. A prospective observational study evaluating BMD changes in the year after critical illness, with comparisons to age and gender matched population controls, and adjustment for known risk factors and possible critical illness factors is now required.

ACKNOWLEDGMENTS

SLB supported by NHMRC Early Career Fellowship (1012472) SLB supported by NHMRC Early Career Fellowship (1012472)
REFERENCES


Figure 1: Summary of systematic search (QUOROM diagram)

Search Results n= 13,185
  - MEDLINE n=3717
  - EMBASE n=9468

Duplicates n=2218

Studies retrieved for more detailed evaluation n=10,967

Studies excluded based on title and abstract
  - Non English n=51
  - Non human n=20
  - Non adult =4747
  - Not ICU n= 1718
  - Not new bone turnover n=4092
  - Neurological study n=37
  - Not research paper=286

Potentially appropriate studies for full-text review n=21

Studies excluded based on full text eligibility n= 10
  - Not ICU 24hrs n= 1
  - Not human n=1
  - Not new bone turnover n=3
  - Not research paper=4
  - RCT=1

Additional studies identified through reference review of eligible full-text articles n=6

Studies excluded based on full text eligibility n= 6
  - Not new bone turnover n=3
  - Not research paper=3

Studies included in systematic review n = 11
Table 1: Criteria list for assessment of methodological quality for each eligible study, adapted from Lievense et al [15]

<table>
<thead>
<tr>
<th>Item</th>
<th>Criterion</th>
</tr>
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<tbody>
<tr>
<td><strong>Study Population</strong></td>
<td></td>
</tr>
<tr>
<td>1. Selection at uniform point</td>
<td>C/CC</td>
</tr>
<tr>
<td>2. Cases and controls drawn from same population</td>
<td>CC</td>
</tr>
<tr>
<td>3. Participation rate &gt;80% for cases/cohort</td>
<td>C/CC</td>
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<tr>
<td>4. Participation rate &gt;80% for controls</td>
<td>CC</td>
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<tr>
<td><strong>Assessment of risk factor</strong></td>
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<tr>
<td>5. Exposure assessment was blinded</td>
<td>C/CC</td>
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<tr>
<td>6. Exposure measured identically for cases and controls</td>
<td>CC</td>
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<tr>
<td>7. Exposure assessed prior to outcome</td>
<td>C/CC</td>
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<tr>
<td><strong>Assessment of bone turnover</strong></td>
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<tr>
<td>8. Bone turnover assessed identically in studied population</td>
<td>C/CC</td>
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<tr>
<td>9. Bone turnover reproducibly (coefficient of variation reported)</td>
<td>C/CC</td>
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<tr>
<td><strong>Study design</strong></td>
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<td>10. Prospective design used</td>
<td>C/CC</td>
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<tr>
<td>11. Follow-up time &gt;24 months</td>
<td>C</td>
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<tr>
<td>12. Withdrawals &lt;20%</td>
<td>C</td>
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<tr>
<td><strong>Analysis and data presentation</strong></td>
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<tr>
<td>13. Appropriate analysis techniques used</td>
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<tr>
<td>14. Adjusted for at least age and gender</td>
<td>C/CC</td>
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† C= applicable to cohort studies: CC= applicable to case-control studies.
Table 2: Criteria list for determining the level of evidence for best-evidence synthesis, adapted from Lievense et al. [15,16]

<table>
<thead>
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<td>Multiple high-quality cohort studies</td>
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<td>Moderate evidence</td>
<td>Generally consistent findings in:</td>
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<td>One high-quality cohort study and &gt;2 high-quality case-control studies</td>
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<td>&gt;Three high-quality case-control studies</td>
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<td>Limited evidence</td>
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<td>One or two case-control studies or</td>
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<td></td>
<td>Multiple cross-sectional studies</td>
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<td>Conflicting evidence</td>
<td>Inconsistent findings in &lt;75% of the trials</td>
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<td>Orford et al, Australia, 2011 [18]</td>
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<td>Lind et al, Sweden, 2000 [20]</td>
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<td>Nierman et al, USA, 2000 [21]</td>
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<td>Van den Berghe et al, Belgium, 2002 [22]</td>
<td>Prospective case-control</td>
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Table 3: continued

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<th>First author, country, year</th>
<th>Design</th>
<th>Subjects no. (% female)</th>
<th>Age</th>
<th>Patient description</th>
<th>Exclusions</th>
<th>Follow-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>Van den Berghe et al, Belgium, 2001 [23]</td>
<td>Prospective case-control</td>
<td>Cases 15 (0%)</td>
<td>67±12</td>
<td>Male adult patients with duration mechanical ventilation &gt;2 weeks, expected ICU LOS at least further 2 weeks</td>
<td>Metabolic, neurological, endocrine disease, liver and renal failure, medications</td>
<td>6 days</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Controls 50 (0%)</td>
<td>67±8</td>
<td>Age, BMI, gender matched healthy controls.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Van den Berghe et al, Belgium, 1999 [24]</td>
<td>Prospective case-control</td>
<td>Cases 14 (29%)</td>
<td>68±12</td>
<td>Adult patients with duration mechanical ventilation &gt;2 weeks, expected ICU LOS at least 2 further weeks</td>
<td>Metabolic, neurological, endocrine disease, liver and renal failure, medications</td>
<td>10 days</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Controls 65 (23%)</td>
<td>NA</td>
<td>Age, gender matched healthy controls</td>
<td></td>
<td>1 day</td>
</tr>
<tr>
<td>Owen et al, Belgium, 2012 [25]</td>
<td>Prospective case-control</td>
<td>Cases 12 (29%)</td>
<td>57 [26-80]</td>
<td>Adult patients with prolonged critical illness (ICU LOS=7-134 days)</td>
<td>Nil</td>
<td>1 day</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Controls 12 (29%)</td>
<td>57 [23-81]</td>
<td>Age, gender, BMI matched healthy controls</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Van den Berghe et al, Belgium, 2003 [26]</td>
<td>Prospective case-control</td>
<td>Cases 22 (46%)</td>
<td>62 [36-73]</td>
<td>Adult ICU patients with ICU LOS &gt;10 days</td>
<td>Chronic bone or kidney disease, prior glucocorticoids</td>
<td>10 days</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Controls 22/64 (46%)</td>
<td>62 [41-73]</td>
<td>Age, gender, BMI matched controls. Analysis of blood samples (n=22) and urine samples (n=64)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Smith et al UK, 2002 [27]</td>
<td>Prospective case-control</td>
<td>Cases 23 (56%)</td>
<td>65±2.9</td>
<td>Adult ICU patients admitted consecutively to ICU (sepsis n=20, trauma n=3)</td>
<td>Nil</td>
<td>1-6 days</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sepsis 20 (60%)</td>
<td>20±2.7</td>
<td>Adults with soft-tissue rheumatism or mild osteoarthritis (not gender and age matched)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Trauma 3 (33%)</td>
<td>40±1.8</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Controls 29 (55%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Shapses et al, USA, 1997 [28]</td>
<td>Prospective case-control</td>
<td>Cases 9 (33%)</td>
<td>73±7</td>
<td>Adult ICU patients requiring postoperative mechanical ventilation after gastrointestinal surgery</td>
<td>Diabetes, liver disease, renal disease</td>
<td>Up to 20 days</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Controls 17 (47%)</td>
<td>60±2</td>
<td>Age matched healthy controls</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

BMI = body mass index, GOS = Geelong Osteoporosis Study; a recruited random population-based sample, ICU= Intensive Care Unit, LOS = length of stay, NA = not available, NTX = collagen type 1 cross-linked N-telopeptide, RCU = respiratory care unit.
Table 4: Results of eligible studies included in review, presented by study design according to methodological quality assessment score

<table>
<thead>
<tr>
<th>Study First author</th>
<th>Bone turnover measure</th>
<th>Outcome</th>
<th>Result (presented as OR or HR with 95% CI ± p-value, mean±SD, or median [IQR])</th>
<th>Summary of significant findings</th>
<th>Quality Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cohort Orford et al. [18]</td>
<td>Fracture</td>
<td>Age-adjusted comparison of incident fragility fracture rate between female patients post-ICU discharge and community controls</td>
<td>Adjusted HR 1.65 (95% CI 1.08-2.52, p=0.02)</td>
<td>Increase in fragility fracture risk in older female ICU survivors compared with age- and gender-matched population control subjects.</td>
<td>64% (7/11)</td>
</tr>
<tr>
<td>Nierman et al. [19]</td>
<td>Biomarker - uNTX</td>
<td>1. Comparison to reference range 2. Metabolic categorisation into different states</td>
<td>Normal (n=4) NTX 43±23 Primary hyperresorption (n=4) NTX 176±88 Secondary hyperresorption (n=19) NTX 257±202 Mixed (n=22) NTX 200±122</td>
<td>Increased bone resorption in chronic critically ill patients</td>
<td>64% (7/11)</td>
</tr>
<tr>
<td>Lind et al. [20]</td>
<td>Biomarker - sICTP, uDpd</td>
<td>Comparison between sepsis and major surgery patients</td>
<td>Marker Sepsis Surgical D1 D3 D1 D3 uDpd 11±7.2 6.9±2.3 9.4±5.5 8.4±5.2 sICTP 25±16 25±15 12±13 16±12</td>
<td>Increased bone resorption in both septic and surgical patients in ICU</td>
<td>55% (6/11)</td>
</tr>
<tr>
<td>Nierman et al. [21]</td>
<td>Biomarker - uNTX</td>
<td>Comparison to reference range</td>
<td>uNTX 215±175, PTH 81±123</td>
<td>Increased bone resorption in chronic critical illness, with positive association with ICU and hospital LOS</td>
<td>55% (6/11)</td>
</tr>
<tr>
<td>Case-control</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Van den Berghe et al. [22]</td>
<td>Biomarker - sOC, sPICP, sSALP, uPyd, uDpd</td>
<td>Comparison between cases and control</td>
<td>sSALP 22.8±21.2 19.6±18.4 Controls 9.1±5.4 9.1±5.4 p-value 0.0008 0.02</td>
<td>Increased bone resorption and synthesis. Positive association between inflammatory cytokines and bone formation markers</td>
<td>67% (8/12)</td>
</tr>
<tr>
<td>Van den Berghe et al. [23]</td>
<td>Biomarker - sOC, sPICP, sSALP, uPyd, uDpd</td>
<td>Comparison between cases and controls</td>
<td>sSALP 296±175 325±219 Controls 19.6±14.8 22±10 p-value 0.0008 0.02</td>
<td>Increased bone resorption and synthesis. Positive association between inflammatory cytokines and bone formation markers.</td>
<td>67% (8/12)</td>
</tr>
<tr>
<td>Study First author</td>
<td>Bone turnover measure</td>
<td>Outcome</td>
<td>Result</td>
<td>Summary of significant findings</td>
<td>Score</td>
</tr>
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<td>--------------------</td>
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<td>---------------------------------</td>
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</tr>
<tr>
<td>Van den Berghe et al. [24]</td>
<td>Biomarker - sSOC, sPICP, sSALP, uPyd, uDpd</td>
<td>Comparison between cases and controls</td>
<td>sSALP: 13.5±11.4 vs 9.1±5.4, p = NS; sPICP: 256±108 vs 51.4±20.4, p &lt; 0.0001; sOC: 29.5±14.3 vs 34.5±29, p = 0.04; uPyd: 836±776 vs 61±27, p &lt; 0.0001; uDpd: 106±80 vs 12±5, p &lt; 0.0001</td>
<td>Increased bone resorption and synthesis in prolonged critically ill adults. Inverse association between bone resorption markers and thyroid hormones, and positive association between bone formation markers and inflammatory markers.</td>
<td>67% (8/12)</td>
</tr>
<tr>
<td>Owen et al. [25]</td>
<td>Biomarker - osteoclast precursor - mature osteoclast</td>
<td>Comparison between subjects and controls</td>
<td>Osteoclast precursor: 99.1% vs 83.9%, p &lt; 0.05; Mature osteoclast</td>
<td>Increased circulating osteoclast precursors in critical illness</td>
<td>58% (7/12)</td>
</tr>
<tr>
<td>Van den Berghe et al. [26]</td>
<td>Biomarker - sSOC, sPICP, sSALP, sBCTX, SPINP, uPyd, uDpd</td>
<td>Comparison between cases and control</td>
<td>sSALP: 9.4±3.9 vs 9.1±5.1, p = 0.7; sPICP: 388±360 vs 51±23, p = 0.0004; sPINP: 65[39,123] vs 38[30,46], p = 0.002; sOC: 21.8±10.8 vs 34.5±29, p &lt; 0.0001; BCTX: 0.7[0.46,1.03] vs 0.12[0.08,0.18], p &lt; 0.0001; uPyd: 328[215,503] vs 55[43,71], p &lt; 0.0001; uDpd: 55[36,76] vs 11[8,14], p &lt; 0.0001</td>
<td>Increased bone turnover, characterised by active immature osteoblasts, low activity of mature osteoblasts, and increased osteoclast activity.</td>
<td>58% (7/12)</td>
</tr>
<tr>
<td>Smith et al. [27]</td>
<td>Biomarker - uPyd, uDpd</td>
<td>- Comparison between cases and control - Multiple regression analysis</td>
<td>ICU sepsis: Age 65±2.9 vs 40±1.8, p = 0.001; uPyd: 417.0±155 vs 44.7±2.6, p = 0.001; uDpd: 67.4±17.2 vs 10.3±0.7, p = 0.001</td>
<td>Increased bone resorption in sepsis and trauma compared to controls. Nitric oxide (NO) not correlated with inhibition of bone resorption.</td>
<td>58% (7/12)</td>
</tr>
<tr>
<td>Shapses et al. [28]</td>
<td>Biomarker - uPyd, uDpd</td>
<td>Comparison between subjects and controls</td>
<td>Increased Pyd and Dpd (p&lt;0.001)</td>
<td>Increased bone resorption markers in subjects compared to controls, and increased bone resorption markers in longer stay ICU patients compared to shorter stay</td>
<td>58% (7/12)</td>
</tr>
</tbody>
</table>

uPyd= urinary deoxypyridinoline (Ref Range 1.3-8.4 nmol/L or 4.3-27.5 nmol/mmol Creat), sICTP/BCTX = serum carboxy-terminal cross-linked telopeptide of type 1 collagen (Ref Range 1.9-6.6 ug/L), IGFBP-3 (Insulin-like growth factor-binding protein 3), M-CSF=macrophage colony stimulating factor, uNOx = urinary Nitric Oxide breakdown products nitrate and nitrite (uNOx/creatinine, mmol/mmol), uNTX= urine collagen type 1 cross-linked N-telopeptide (Ref Range 12-80 BCE/mmol Cr), sOC= serum osteocalcin (Ref Range 13.2-254 ug/L), PBMC=Peripheral blood mononuclear cell, sPICP = serum carboxyl-terminal extension peptide of type 1 procollagen (Ref Range 75-254 ug/L), PTH = Parathyroid hormone (Ref Range 12-55 pg/ml), uPyd = urinary pyridinoline (Ref Range 27-182 nmol/mmol Creat), RANKL =receptor activator of NF-κB ligand, sSALP = serum skeletal alkaline phosphatase (Ref Range 9.0-15.0 ug/L).
Author/s:  
Orford, N; Cattigan, C; Brennan, SL; Kotowicz, M; Pasco, J; Cooper, DJ

Title:  
The association between critical illness and changes in bone turnover in adults: a systematic review

Date:  
2014-10-01

Citation:  
Orford, N; Cattigan, C; Brennan, SL; Kotowicz, M; Pasco, J; Cooper, DJ, The association between critical illness and changes in bone turnover in adults: a systematic review, OSTEOPOROSIS INTERNATIONAL, 2014, 25 (10), pp. 2335 - 2346

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File Description:  
Accepted version