In this mini-review of hypocalcemia in the critically ill patient historical and new insights in the function of the PTH-vitamin D axis are discussed. Special attention is given to the role of locally produced calcitriol in the immune system and in the inflammatory cytokine response of the vitamin D deficient critically ill patient. Future directions in this research area and in the forthcoming research on the use of the anabolic recombinant PTH analogues in critical care medicine are discussed.

**Keywords:** Hypocalcemia; PTH-vitamin D; PTH, 25(OH)D3, 1,25 di-(OH)D3

**Abbreviations:** PTH: Parathyroid Hormone; cAMP: Cyclic Adenosine 3,5, Monophosphate; ECF: Extracellular Fluid Compartment; VDRs: Vitamin D Receptors; AMP: Antimicrobial Peptide; BMD: Bone Mineral Density; CRP: C - Reactive Protein

**Introduction**

In a balanced diet, roughly 1000 mg of calcium is ingested each day and about 200 mg is secreted in the bile and other gastrointestinal secretions. Depending on the concentrations of circulating parathyroid hormone (PTH) and active i.25.dl-(OH)D3 about 200-400 mg is absorbed from the intestine each day. The remaining calcium is excreted in the stool. Maintenance of body calcium stores depends on dietary calcium intake, absorption of the gastrointestinal tract and renal calcium excretion. PTH enhances distal tubular calcium reabsorption, independently of sodium, while proximal tubule calcium excretion is sodium dependent and coupled to phosphaturia, urinary cAMP (Cyclic Adenosine 3,5, Monophosphate) and nephrogenous cAMP excretion.

About 99% of body calcium is stored in bone mainly as hydroxyapatite crystals. About 1% of calcium is freely exchangeable with the extracellular fluid compartment (ECF) and therefore is available to buffering changes in calcium balance. Normal total serum calcium concentrations range from 8,8 to 10,4 mg/dl or 2,20-2,60 mmol/l. About 40% of the total blood calcium is bound to plasma proteins, primarily albumin. The remaining 60% includes ionized calcium and calcium complexed with phosphate and citrate. Ionized calcium is the physiologically active form and acts as an intracellular second messenger involved in skeletal muscle contraction, excitation-contraction coupling in cardiac and smooth muscle and activation of protein kinases and phosphorylation leading to the formation of other second messengers as cAMP and inositol 1,4,5-triphosphate mediating the cellular response to numerous hormones including epinephrine, glucagon, secretin and cholera toxin.

Ionized calcium is assumed to be 50% of the total serum calcium and can be estimated based on total serum calcium levels and albumin levels. In hypoalbuminemia measured total serum calcium is often low reflecting a low concentration of protein-bound calcium while ionized calcium may be normal. Measured total serum calcium decreases or increases by about 0,8 mg/dl or 0,2 mmol/l for every g/dl decrease or increase in albumin. Thus an albumin concentration of 2,0 g/dl (normal 4,0 g/dl) should itself reduce measured serum calcium by 1,6 mg/dl or 0,4 mmol/l, Even in the presence of a normal serum albumin changes in blood pH can alter the equilibrium constant of the albumin-calcium complex with acidosis reducing the binding and alkalosis enhancing it. In critically ill or post-surgical patients with hypoalbuminemia correcting total serum calcium for albumin is not necessarily accurate because of changes in pH and affinity of calcium binding [1]. A total serum calcium level less than 2,1 mmol/l and an ionized calcium level of less than 1,0 mmol/l is defined as hypocalcaemia [2]. Ionized calcium measurements offer additional benefits in the diagnosis of primary hyperparathyroidism because they are correlated with PTH levels and adenoma size and may be more sensitive marker of disease severity than total serum calcium, which is an appropriate first-line biochemical test [3].

Analysis for the ionized calcium level must be performed rapidly with whole blood to avoid changes in pH and anion chelation. Blood should be drawn in a heparinized syringe or an EDTA blood bottle which is transported to the gas analyzer and handled immediately for the best results. Falsely elevated ionized calcium levels may be seen with elevated acetaminophen levels, alcohol, hydralazine and hemolysis although this effect is modest. Falsely depressed levels can be seen with heparin, oxalate or pyridoxine.

In cases of hypocalcemia serum magnesium should always be checked. Occasionally inadequate dietary magnesium as in malnutrition and chronic alcoholism can lead to hypomagnesemia. Excessive use of thiazide diuretics and long term proton pump inhibitor use are other causes of hypomagnesemia. Hypomagnesemia impairs parathormone (PTH) secretion resulting in hypocalcemia and hypophosphatemia [7,9].

The differential diagnosis ranges from (pseudo) hypoparathyroidism, renal failure, post-thyroidectomy surgery, acute pancreatitis to widespread osteoblastic metastasis.
in prostate and breast cancer [10]. This mini-review will focus on hypocalcemia in the critically ill patient.

### Hypocalcemia in the critically ill

The first clinical observations and studies in hypocalcemic critically ill patients date back to the early 70’s and 80’s of the last century [11,12]. The incidence of hypocalcemia in critically ill patients varies widely depending on the different underlying diseases and comorbidity. In an analysis of 12 studies performed between 1988 and 2014 Aberegg believes the incidence of hypocalcemia in critically ill patients ranges from 50-88% [13].

However some critical annotations must be made. Some studies used corrected calcium values while others measured ionized calcium. It is also not clear if the calcium measurements were made for diagnostic, screening, homeostatic or daily routine purposes. The only thing that is evident at present is that some 50% of critically ill patients in an ICU will have hypocalcemia at some moment at some day during their stay [13].

Does that mean half of critically ill ICU patients has some latent disorder of calcium metabolism? Probably not but it is understandable that early studies focused on possible alterations in the PTH-Vitamin D axis in the hypocalcemic critically ill.

### The PTH-Vitamin D axis

Zalago reviewed in 1992 earlier human and animal investigations and in vivo and in vitro studies. He concluded that hypocalcemia in critically ill patients usually results from impaired PTH secretion or action (end-organ resistance), impaired vitamin D synthesis or action or calcium chelation and precipitation [14].

In a small 1999 study Lepage et al. [15] showed significant decreases in ionized calcium during abdominal surgery with compensatory significant rises in PTH levels depending on the severity of the surgery. Hypomagnesemia was absent in this study thus PTH secretion seemed intact under these circumstances. Alterations in blood pH during anaesthesia were not present and there was also no relationship between the decrease in ionized calcium and blood transfusions.

In 2000 Lind et al. [16] studied the dynamics of PTH secretion by a calcium chloride infusion on day 1 and 3 in 13 septic ICU patients with hypocalcemia and 13 surgical patients with normocalcemia. Serum PTH levels were increased in both groups. In both study groups an increased PTH secretion was found.

Is there end-organ resistance to PTH in hypocalcemic critically ill patients as Zalago suggested? End-organ resistance is tested in pseudohypoparathyroidism and in other disease states by PTH infusion followed by measuring the renal response in terms of phosphaturia and nephrogenous cAMP excretion [17]. These studies were not done in hypocalcemic critically ill patients with normal serum magnesium levels.

In hypomagnesemic, hypocalcemic critically ill patients Rude et al showed already in 1976 that functional hyperparathyroidism and end-organ resistance to PTH was caused by magnesium deficiency only by administering PTH extract in these patients.

The renal response was evaluated by measuring urinary cAMP excretion. Within minutes of i.v. magnesium administration serum PTH levels rose from undetectable levels to sky high. Restoring of the PTH end-organ resistance took at least 4 days [18].

So the suggestions of Zalago for the existence of impaired PTH secretion and PTH end-organ resistance in hypocalcemic critically ill patients with normal serum magnesium levels cannot be confirmed by evidence. Nevertheless these suggestions are widely cited in nearly all articles about this subject.

### Vitamin D and metabolites

Vitamin D deficiency is common in ICU patients and prevalence rates range from 38%-100% in observational studies [19]. Critically ill patients with a prolonged stay at an ICU may develop vitamin D deficiency for a number of reasons including the lack of sunlight, malnutrition, decreased renal 1-alpha-hydroxylation in acute renal failure and increased tissue conversion of 25(OH)D3 to 1.25 di-(OH)D3 during acute stress and inflammatory response [20,21].

Vitamin D is usually supplemented in critical care along with nutritional support. Approximately 100-300 IU of vitamin D2 or vitamin D3 daily is provided in standard and enteral parenteral regimen [22]. It is questionable whether this relatively small amount is sufficient.

Vitamin D deficiency could contribute certainly to hypocalcemia in the critically ill by diminishing gastro-intestinal absorption of calcium and decreasing renal tubular reabsorption of calcium.

Lee has postulated the theoretical concept of functional vitamin D deficiency in critical illness [19,21]. This means that the clinical consequences of vitamin D deficiency are not only dependent on the severity of vitamin D depletion but are also related to tissue requirements. In such a model the circulating 25(OH) D3 pool represents a substrate reservoir for conversion to active metabolites at the tissue level during acute stress.

It is conceivable that the physiological needs of appropriate conversion of the inactive 25(OH)D3 to the biologically active prohormone 1.25 di-(OH)D3 is generated through a regulatory system of local and renal 1-alpha hydroxylases to satisfy the demand of tissue requirements. When circulating vitamin D levels are sufficient and tissue requirement is relatively low organ function is normal and physical health maintained.

In acute stress and critical illness however, multifactorial hypocalcemia is very common and may lead to a compensatory rise in PTH levels which would enhance the conversion of 25(OH)D3 to 1.25 di-(OH)D3 to maintain calcium homeostasis as a result of increased bone resorption and intestinal calcium absorption respectively. Due to secondary hyperparathyroidism the consumption of 25(OH)D3 would further exacerbate vitamin D deficiency. This could be also an explanation for the high prevalence of vitamin D deficiency in the ICU.

Evidence in support of this hypothesis came from studies that showed secondary hyperparathyroidism was linked to hypocalcemia and low 25(OH)D3 levels [23-25].

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Furthermore in contrast to the general population in which calcitriol levels are maintained within the normal range despite low 25(OH)D3 levels calcitriol levels are up to 50% lower in ICU patients. Calcitriol levels correlated positively with 25(OH)D3 in this setting. This suggests substrate deficiency may be more profound in critically ill patients [20,25-28]. Therefore vitamin D deficiency in the ICU is a mismatch between substrate supply and tissue requirements. Despite stimulation of renal 1-alpha hydroxylase local mitochondrial tissue 1-a hydroxylases are unable to generate adequate 1.25 di-(OH)D3 levels. PTH resistance associated with hypomagnesemia, acute renal failure and functional hypoparathyroidism may further compromise 1.25 di-(OH)D3 production.

It is clear that in this vision the widely held concept of vitamin D end-organ resistance in the absence of hypomagnesemia, as suggested by Zalago in 1992, has been abandoned completely.

In a 2014 systematic review and meta-analysis de Haan et al showed Vitamin D deficiency to be a risk factor for infection, sepsis and mortality in the critically ill [29].

The cytokines

Proinflammatory cytokines such as interleukin-1(IL-1), interleukin-6 (IL-6) and tumor necrosis factor-alpha (TNF-alpha) are important mediators of the acute response to critical illness. An interplay between serum calcium, the PTH-vitamin D axis and the cellular immune system has been proposed [30]. Calcitriol has a central role in this interplay by its paracrine and autocrine non-skeletal effects.

Nearly 35 years ago the presence of vitamin D receptors (VDRs) in activated human immune cells was discovered as well as the ability of 1.25 di-(OH)D3 to inhibit T cell proliferation and the ability of activated macrophages to produce 1.25 di-(OH)D3 in patients with sarcoidosis [31-33]. This conversion has not been found in the tuberculosis granuloma and is specific for the sarcoid granuloma for unknown reasons. Nevertheless patients with a vitamin D deficiency are more susceptible to TB and recover faster with vitamin D supplementation to their tuberculostatic regimen [34]. It now appears that 1.25 di-(OH)D3 modulates the immune response in macrophages by upregulating the expression of the antimicrobial peptides cathelicidin and defensin which inhibit the intracellular growth of M.tuberculosis. Apart from the antimicrobial effects 1.25 di(OH)D3 also modulates the antigen presentation and secretion of chemokines and cytokines enhancing macrophage innate immune function by a cooperating system of toll-like receptors and vitamin D receptors as described below [34-36].

The significance of these observations concerning innate immunity was clarified with the discovery of toll-like receptors (TLRs) in monocytes and macrophages nearly 25 years later: TLRs are transmembrane receptors and recognize microbial ligands including bacterial lipopolysaccharides and are responsible for the transcriptional induction of genes like CYP27B1 and the vitamin D receptor (VDR) [37].

In this way circulating serum 25(OH)D3 enters monocytes, is converted by mitochondrial 1-alpha hydroxylase to active 1.25 di-(OH)D3 and then binds to the VDR. The complex of 1.25 di-(OH)D3 VDR bound leads to the transcription of cathelicidin expression, which is a potent antimicrobial peptide (AMP) that promotes autophagy [38]. Cathelicidin expression can be activated by 1.25 di-(OH)D3 alone. Another important AMP, defensin, is also produced by TLR induced expression [35,38-40]. These AMPs are now in phase 1 and phase 2 clinical trial stage [35].

Markers of systemic inflammation including C-reactive protein (CRP), TNF-alpha and interleukin-6 (IL-6) are typically increased during ICU stay. Prolonged hypercytokinemia leads to multi-organ failure [20].

Vanden Berghe et al. [20] showed that these circulating levels of inflammation biomarkers were 5-400 fold higher upon intensive care admission. In prolonged critically ill patients parenteral vitamin D3 supplementation of 500 IU per day reduced CRP and IL-6 levels by approximately 40% and 60% respectively and the fall of the inflammatory markers was significant compared to the control group receiving 200 IU vitamin D3 daily [20,21].

The multiple functions of vitamin D in the immune system’s response to infection lead to the suggestion that vitamin D could be useful in the treatment of sepsis. Gram-negative sepsis in particular is known to be associated with hypocalcemia and vitamin D deficiency in the critically ill. The VDR recognizes the lipopolysaccharides in the Gram-negative bacterial outer membrane [41].

Jeng et al. [37] showed that vitamin D binding protein levels are significantly lower in critically ill subjects with sepsis compared to critically ill patients without sepsis and healthy controls. When they examined plasma levels of the endogenous AMP LL-37 (cathelicidin) they found that lower levels of LL-37 were associated with lower levels of 25(OH)D3. This association supports in vivo data that vitamin D plays some roles in regulating the production of AMPs such as LL-37 in cultured macrophages.

Since many cells of the immune system have VDRs vitamin D status may prove to be an important factor in the management of the sepsis syndrome and critical illness. In this promising research area a lot has to be elucidated yet.

Recently De Fillipis et al. showed that vitamin D reduces the inflammatory response in a gingivitis model by modulating the human AMP beta-defensin-3 [42].

Results in healthy volunteers exposed to experimental human endotoxinemia didn’t show an association between vitamin D and cytokine levels [43]. May be that decreased vitamin D binding protein levels which are normal in healthy volunteers play a crucial yet unknown role in bacterial susceptibility [41].

Treatment

Correcting hypocalcemia

The treatment of hypocalcemia depends on the cause, the severity, the presence of symptoms and the speed of developing hypocalcemia [2]. Mostly hypocalcemia is mild and requires only
supportive therapy. Sometimes it is severe with seizures, tetany, refractory hypotension and cardiac arrhythmias that require an aggressive approach.

Studies in critically ill patients have yielded conflicting results regarding calcium supplementation. Some studies suggest that serum calcium normalizes mostly within 4 days after admission to the ICU and that low levels of calcium are protective and attempted correction may be harmful [13,44]. In contrast other studies have concluded that both mild and moderate hypocalcemia are associated with increased mortality. One large retrospective study showed that calcium supplementation improved 28 day survival in the ICU in critically ill patients [45,46].

In the MIMIC -2 study the only large retrospective study of sufficient statistical power to date both hypercalcemia and hypocalcemia were associated with altered mortality in a complex form. Interestingly mild hypercalcemia on ICU admission was found to be protective and was associated with reduction in 28- day mortality while hypocalcemia showed the opposite effect. This curve showed an U-shape, implicating endless calcium supplementation may be harmful. However many confounders were present as e.g the lack of standardized calcium supplementation, hypovitaminosis D3, absence of dialysis patients and differences in timing of blood samples. A sub analysis showed later also a beneficial effect of calcium supplementation on 90-day mortality. Follow up data studying morbidity and the overall clinical outcome are not available because too many patients were lost in the follow-up [46].

Are there evidence based guidelines for correcting hypocalcemia? No, the Endocrine Society released an Emergency Guidance for the management of acute hypocalcemia that is so very brief it has no additional value for daily clinical practice [47].

**A useful scheme would be**

**i. Mild:** (Ionized Calcium; 1, 0-1,2 mmol/l): Calciumgluconate i.v; 1-2 g over 2 hours.

**ii. Severe:** Without seizure or tetany; 0,5mg/kg/hr i.v; may be increased to 2mg/kg/hr not to exceed 3-4 g i.v over 4 hours.

**iii. Hypocalcemic tetany:** 3 g calciumgluconate i.v over 5-10 minutes, followed by continuous i.v infusion of 0,5 mg/kg/hr that may be increased to 2 mg/kg/hr. Monitor serum calcium q4-6 hours to maintain serum calcium levels.

**Vitamin D supplementation**

In general there are two U.S guidelines issued for vitamin D supplementation in the general population at risk for vitamin D deficiency. These are released by the IOM (Institute of Medicine) and by a Task Force of the Endocrine Society. They differ in their definitions of vitamin deficiency by using different cut-offs of 25(OH)D3 levels. To complicate matters further the Endocrine Society Task Force focuses on diseased people and advises higher supplemental doses while the IOM focuses on the normal population [48]. U.K NICE guidelines are directed at population groups at risk but do not refer in their recommendations to the ICU group of patients. An initial i.v dose of 50,000-60,000 IU should be considered. In fact it is not known if vitamin D should be given as loading i.v bolus, weekly i.v doses or only by daily TPN or feeding tube when oral supplementation is not possible [22,49,50].

In studies of vitamin D supplementation in critically ill patients different vitamin D formulas were used. The Spanish Mata-Granados Group demonstrated that 60,000 IU cholecalciferol administered orally at day 0 and day 4 evoked a significant rise in 25(OH)D3 and 1.25-di-(OH)D3 levels whereas the application of 2ug calcitriol intravenously on alternate days had no impact on these levels in septic patients [28].

In another small study 540,000 IU cholecalciferol or placebo was delivered once by feedtube or orally to 25 critically ill patients at an intensive care unit Already on day 1 the 25(OH)D3 levels were significantly higher in the intervention group. On day 1 the 1.25-di-(OH)D3 levels were significantly higher but this was only very transiently. Eighty percent of the vitamin D treated patients achieved 25(OH)D3 levels above 30 ng/ml.

Of interest is the transient and short-lived rise of 1.25 di-(OH) D3 levels. Probably this was due to a substrate boost for local tissue mitochondrial 1-alpha-hydroxylase activity. There were large individual differences in 25(OH)D3 levels probably due to large variations in gastrointestinal absorption [51]. Intravenous formulations could overcome this problem but i.v administration of calcitriol had no impact on 25(OH)D3 and 1.25-di-(OH)D3 levels in septic patients as shown in the Mata-Granados study [28].

It seems that the local tissue conversion of 25(OH)D3 to locally produced 1.25 di-(OH) D3 is essential in vitamin D function as e.g stimulation of AMP3 (37,38,41). This supports Lee’s hypothesis that Vitamin D deficiency in the critically ill is a mismatch between substrate supply (25(OH) D3, cholecalciferol) and tissue requirements further [19]. So a “one fits it all” vitamin D supplementation regimen can’t be given.

**Parathormone**

Human recombinant 1-84, full length, PTH (Natpara) proved to be efficacious and safe in the REPLACE study for the treatment of hypoparathyroidism [52]. In the RACE study long-term treatment of hypoparathyroidism improved hypercalciuria as the 3-year analysis showed [53]. Another recombinant PTH analogue is the 1-34 teriparatide (Forteo) which showed to be superior to alendronate for increasing lumbar spine bone mineral density (BMD) in postmenopausal osteoporosis at 12 months of treatment. However teriparatide was not superior to alendronate in reducing fracture risk [54]. Both these analogues have anabolic properties and effects [52-57] Studies about their efficacy in the hypocalcemia of the critically ill patient are awaiting as they could reverse or improve bone hyperresorption in the critically ill patient [27].

**Conclusion**

Hypocalcemia occurs in about 50 % of critically ill patients in an intensive care setting. Hypocalcemia should be measured by ionized calcium with a concomittant serum magnesium.
measurement. In cases of hypocalcemia and hypomagnesemia the serum magnesium deficiency should be corrected always at first. When serum magnesium is normal there is no impaired PTH secretion or PTH end-organ resistance in the hypocalcemic critically ill patient neither there is vitamin D end-organ resistance in the vitamin D deficient critically ill. Local tissue conversion of cholecalciferol by 1-alpha mitochondrial tissue hydroxylase seems to play an essential role in the critically ill patient’s immune system function as eg the production of antimicrobial peptides (AMPs) and in the acute inflammatory cytokine response. These functions of locally produced calcitriol can only be performed when the circulating pool of substrate of cholecalciferol is sufficient. In septic critically ill patients a mismatch between substrate supply and tissue requirements frequently occurs and contributes to the high prevalence of vitamin D deficiency in critically ill patients. Evidence based guidelines for calcium-and vitamin D supplementation in critically ill patients are hardly present. Future research in the immune and other possible functions of locally produced calcitriol in vitamin D deficient critically ill patients might resolve these knowledge gaps. Tha anabolic effects of the recombinant PTH analogues in the critically ill patient will be another forthcoming research topic in critical care medicine.

References


