Serum calcium and bone: effect of PTH, phosphate, vitamin D and uremia

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ABSTRACT

Hyperparathyroidism develops in chronic kidney disease (CKD). A decreased calcemic response to parathyroid hormone (PTH) contributes to the development of hyperparathyroidism and is presumed due to reduced calcium efflux from bone. Contributing factors to the decreased calcemic response to PTH in CKD include: 1) hyperphosphatemia; 2) decreased serum calcitriol; 3) downregulation of the PTH1 receptor; 4) large, truncated amino-terminal PTH fragments acting at the carboxy-PTH receptor; and 5) uremic toxins. Also, prolonged high dose calcitriol administration may decrease the exchangeable pool of bone calcium independent of PTH. The goal of the review is to provide a better understanding of how the above cited factors affect calcium efflux from bone in CKD. In conclusion, much remains to be learned about the role of bone in the regulation of serum calcium.

Keywords: Bone. Calcium. Parathyroid hormone. Phosphate. Uremia. Vitamin D.

The decreased calcemic response to parathyroid hormone (PTH) in chronic kidney disease (CKD)¹,² challenges us to evaluate how calcium is mobilized from bone. Hyperparathyroidism develops in CKD to prevent hypocalcemia. Factors contributing to the need for increased PTH include: 1) hyperphosphatemia or even inappropriately normal serum phosphorus for an elevated PTH; 2) decreased serum calcitriol; 3) downregulation of the PTH1 receptor; 4) accumulation of large, truncated amino-terminal PTH fragments such as 7-84 PTH; and 5) uremic toxins. A review of how these factors affect the calcemic response to PTH in CKD is instructive.

PARATHYROID HORMONE

The relationship between PTH and serum calcium is a sigmoidal curve with basal PTH approximately 25% of the maximal PTH response to hypocalcemia.³ During hypocalcemia, increased PTH secretion restores serum calcium to normal. Besides its bone effect, PTH increases renal calcium reabsorption and calcitriol production enhancing gut absorption of calcium. However, the instantaneous effect and that of greatest magnitude is attributed to calcium efflux from bone. Not often discussed is the role of a calcium gradient with greater mobilization of calcium from bone in hypocalcemia than in hypercalcemia. When increasing PTH doses were infused in parathyroidectomized rats, the serum calcium response was curvilinear with a steeper response between 5 and 10mg/dL than at higher values (Figure 1A) showing that the calcium gradient in hypercalcemia reduced the calcemic ac-
In the early 1990s, D’Amour and associates showed that during PTH stimulation from hypocalcemia, the ratio of intact PTH to carboxy (C)-PTH fragments increased and during PTH suppression from hypercalcemia, the ratio of intact PTH to C-PTH fragments decreased. Subsequently, the same authors showed that besides 1-84 PTH, the assay for intact PTH also detected non 1-84 PTH large truncated amino-terminal fragments of which 7-84 PTH was the prototype. In normal humans, non 1-84 PTH large truncated amino-terminal fragments represent about 20% of measured intact PTH, but in renal failure, these same fragments account for approximately 50%. Interest increased when the 7-84 PTH fragment was shown to inhibit the calcemic action of 1-84 and 1-34 PTH. The action of 7-84 PTH is most likely mediated by its binding to the C-PTH receptor present on osteocytes and other cells of osteoblast lineage including lining cells on the bone surface. Moreover, the osteocyte has important signaling functions to the bone surface. Finally, 7-84 PTH may internalize the PTH1 receptor in selected target cells in kidney and bone and also reduce calcitriol production. Thus, in advanced renal failure, greatly increased 7-84 PTH could decrease the calcemic response to PTH in absence of hypercalcemia.

An intriguing observation in azotemic rats is that parathyroidectomy completely restored the calcemic response to PTH even though hyperphosphatemia was present before the 1-34 PTH infusion (Figure 2 A). Before the PTH infusion, rats

**Figure 1. Serum calcium and phosphorus response to PTH dosing in parathyroidectomized rats.** Individual serum calcium (A) and phosphorus (B) values are shown for each dose of PTH infused for 48 hours with a subcutaneously implanted miniosmotic pump. During the study, rats were fed a normal calcium (0.6%) and phosphate (0.6%) diet. On the normal diet, a PTH dose of 0.022µg/100g-body weight/hr resulted in a normal serum calcium and phosphorus value.4
were maintained on a high calcium diet for more than two weeks after parathyroidectomy. It is possible that the total absence of 7-84 PTH or other PTH fragments resulted in the greater response to PTH. Other possibilities not mutually exclusive include that the total absence of PTH resulted in an upregulation of the PTH1 receptor, hypocalcemia lessened the gradient driving mineralization increasing the exchangeable bone pool of calcium, and the high calcium diet resulted in expansion of the exchangeable bone pool of calcium.

The size of the exchangeable pool of bone calcium differs among the different forms of renal osteodystrophy in CKD. In studies in hemodialysis patients performed in the 1980s and 1990s, we used low and high calcium dialysates during hemodialysis to induce hypo- and hypercalcemia to evaluate PTH stimulation and suppression. The capacity to defend against the development of hypocalcemia and hypercalcemia was much greater in hemodialysis patients with high (osteitis fibrosa) than with low bone turnover (adynamic bone or osteomalacia). A similarly expanded pool of bone calcium and increased calcium efflux from bone has been shown in dialysis patients with osteitis fibrosa studied with calcium isotopes. Moreover, in osteitis fibrosa, bone is deposited in a woven rather than the usual lamellar pattern. The abnormal woven bone may bind or incorporate calcium less well. Also, trabecular bone volume and thickness were greater in dialysis patients with high than low bone turnover suggesting that greater bone mass is present for calcium exchange in the former. A summary of the discussion on PTH and bone is provided in Table 1.

**Table 1. Parathyroid hormone and calcium efflux from bone**

<table>
<thead>
<tr>
<th>A. Classic Effects of PTH</th>
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<tr>
<td>1) Increase in calcium efflux from bone</td>
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<tr>
<td>2) Increase in urinary reabsorption of calcium</td>
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<td>3) Increase in urinary phosphate excretion and decrease in serum phosphorus</td>
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<td>4) Stimulation of calcitriol production</td>
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<th>B. Bone Effects of PTH</th>
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<td>1) Increase in osteoblasts</td>
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<td>2) Increase in osteoclasts</td>
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<td>3) Increase in bone formation rate</td>
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<td>4) Increase in calcium efflux</td>
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<th>C. Points of Discussion</th>
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<tr>
<td>1) Bone-blood calcium gradient</td>
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<td>2) Bone-blood phosphate gradient</td>
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<td>3) Exchangeable pool of bone calcium</td>
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<td>4) Resistance to the calcemic action of PTH in uremia</td>
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<td>5) Effect of large carboxy-terminal PTH fragments</td>
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<td>6) Parathyroidectomy</td>
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**PHOSPHATE**

Phosphate is an important factor in evaluating the calcemic response to PTH. Long ago Albright showed that the first action of infused parathyroid extract was phosphaturia followed by lowering serum phosphorus before a calcemic effect developed. In the 1960s, Raisz showed in an ex-vivo study that increasing the phosphate concentration in the medium reduced calcium efflux from bone both in the absence and presence of PTH. Phosphate also modifies calcium efflux from bone in animal and human studies. In parathyroidectomized rats given a normal phosphate diet and a fixed replacement dose of PTH sufficient to maintain normal serum calcium and phosphorus values, changing to a high phosphate diet resulted in hypocalcemia and hyperphosphatemia. In young, healthy subjects given continuous intravenous phosphate infusion for 7 days, hypocalcemia and hyperphosphatemia developed until a three-fold increase in PTH increased phosphate excretion by five-fold and normalized serum calcium and phosphorus. Conversely, phosphate depletion induces hypercalcemia despite marked suppression of PTH. Even high dose bisphosphonate in phosphate-depleted rats did not prevent hypercalcemia despite producing toxic effects in osteoclasts and a marked decrease in the osteoblast surface. These results support the concept that an osteoclast independent effect may be important for bone efflux of calcium. Finally, bisphosphonate treatment in phosphate-depleted rats increased urine calcium excretion more than in phosphate-depleted rats not receiving bisphosphonates suggesting that decreased calcium influx to bone may be a factor.

In renal failure, many studies have shown that phosphate loading or hyperphosphatemia decreases the calcemic response to PTH. When 5/6 nephrectomy was performed in parathyroidectomized rats, a replacement dose of 1-34 PTH sufficient to maintain normal serum calcium and phosphorus values in parathyroidectomized rats with normal renal function failed to maintain normal serum calcium and phosphorus levels in renal failure. Even though dietary phosphate was unchanged, the phosphate burden was presumably increased because of decreased renal excretion of phosphate. Conversely, dietary phosphate restriction in renal failure decreases PTH while maintaining the serum calcium value. In essence, less PTH is needed to maintain the serum calcium concentration. Even in patients with advanced CKD, dietary phosphate restriction decreases PTH while maintaining normal serum calcium values. A summary of the discussion of the effect of phosphate on serum calcium and bone is provided in Table 2.

**VITAMIN D**

The effect of vitamin D on the exchangeable pool of bone calcium is more difficult to define than that of phosphate. Under some circumstances, 1,25 vitamin D (1,25D) appears to
Table 2. Phosphorus and calcium efflux from bone

<table>
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<th>A. Classic Effects of Phosphorus</th>
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<tr>
<td>1) Calcitriol production</td>
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<td>2) Hyperphosphatemia/phosphate loading and FGF23 production</td>
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<td>3) Phosphate and PTH production</td>
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<th>B. Bone Effects of Phosphate</th>
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<tr>
<td>1) Hyperphosphatemia/phosphate loading decreases calcium efflux from bone</td>
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<tr>
<td>2) Hypophosphatemia/phosphate depletion increases calcium efflux from bone</td>
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<th>C. Points of Discussion</th>
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<tr>
<td>1) Phosphate loading increases PTH which in turn maintains normal serum calcium and phosphorus</td>
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<tr>
<td>2) Demand for PTH is increased in uremia for same phosphate load because of decreased capacity to excrete phosphate</td>
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<tr>
<td>3) Hyperphosphatemia/phosphate loading decreases calcemic response to PTH</td>
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<tr>
<td>4) Restricting dietary phosphate maintains a normal serum calcium despite reduced PTH values</td>
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enhance, but in others to reduce calcium efflux from bone. In vitamin D intoxication, studies have suggested that calcium efflux from bone is increased. In vitro and in vivo studies have shown that calcitriol increases osteoclast activity. In a case report of vitamin D intoxication, bone histomorphometry showed an increase in osteoclasts with a reduction in osteoblasts, an increased osteoid surface and volume, and decreased mineralization. Studies in the chick also found that vitamin D intoxication impaired bone formation. The effects on osteoid volume and bone formation suggest that calcium influx to bone may be decreased in vitamin D intoxication. Both corticosteroids and bisphosphonates effectivley treat hypercalcemia in vitamin D intoxication. However, in different settings, bisphosphonate treatment did not change the rate of recovery from EDTA-induced hypocalcemia in osteoporotic patients or prevent hypercalcemia in rats on a phosphate depleted diet perhaps suggesting different mechanisms.

In a recent study in intestinal-specific VDR-null mice in which calcium absorption is unresponsive to 1,25D, insights into the calcemic and bone effects of 1,25D during calcium deficiency were provided. In this mouse model, markedly elevated endogenous 1,25D levels maintained serum calcium normal by increasing calcium efflux from bone while bone mineralization was inhibited resulting in hyperosteoitis and a decrease in bone mass. The activation of bone resorption by 1,25D in calcium deficient intestinal-specific VDR-null mice was osteoblast mediated. Systemic VDR-receptor null mice, bone abnormalities were prevented when calcium repletion was achieved with a high calcium/lactose diet. In the intestinal-specific null model, although PTH values were elevated, the authors concluded that because markedly elevated PTH did not prevent hypocalcemia in systemic VDR null mice, 1,25D played a critical role in increasing calcium efflux from bone in intestinal-specific VDR-null mice. Finally, the calcemic bone actions of 1,25D were blocked by bisphosphonates. The authors concluded that high 1,25D levels in calcium deficiency: 1) increase calcium efflux from bone; 2) decrease calcium influx to bone by inhibiting bone mineralization; and 3) both 1) and 2) act to maintain serum calcium, but decrease bone mass.

In intestinal-specific VDR null mice, a marked elevation in 1,25D increased calcium efflux from bone. But a major question is whether such a mechanism is applicable in vitamin D deficiency? In dogs placed on a vitamin D and calcium deficient diet, 25D values decreased rapidly, but 1,25D values tripled by 24 weeks and remained greater than normal at 36 weeks before subsequently declining below normal. Ionized calcium values remained normal until 24 weeks before marginally decreasing at 36 weeks. Thus, despite vitamin D and calcium deficiency, serum calcium values were maintained normal for a prolonged time as 1,25D and PTH values tripled during the first 24 weeks. An elevated 1,25D and PTH together with hypophosphatemia contributed to the prolonged maintenance of serum calcium. The marked 1,25D increase during the first 24 weeks could have helped maintain serum calcium by enhancing calcium efflux from bone while also decreasing calcium influx to bone. In humans with established vitamin D deficiency, studies have shown that treatment with vitamin D or 25D increases 1,25D values to 3 to 4 times greater than normal. Thus, it is plausible in vitamin D deficiency that high 1,25D values have an important skeletal effect in maintaining serum calcium during its evolution and increasing serum calcium during initial treatment with vitamin D. Also, in evolving vitamin D deficiency, high 1,25D values perhaps together with PTH-induced decreases in serum phosphorus potentially contribute to the development of osteomalacia.

In renal failure, several studies have shown that acute calcitriol administration increases the calcemic response to PTH. In one study, high dose PTH was infused for 48 hours while calcitriol was administered to rats on a calcium free diet. Thus, neither calcitriol nor hypercalcemia could modify PTH levels. Calcitriol administration enhanced the calcemic response to PTH in azotemic rats on a low phosphate diet and in both normal and azotemic rats on a high phosphate diet (Figure 2 B).

Two studies in dialysis patients are of particular interest because long-term calcitriol treatment decreased the bone response to PTH. In one study, 14 children on peritoneal dialysis with biopsy-proven osteitis fibrosa were treated with high dose calcitriol for 12 months with dose adjustments for hypercalcemia and hyperphosphatemia. The average calcitriol dose was 1.25 µg thrice weekly. The dialysate calcium

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Adynamic bone and in 6 patients the bone formation rate decreased despite no change in PTH values. The latter result suggests that calcitriol treatment may have decreased the concentration was 3.5 meq/L and calcium carbonate was the primary phosphate binder. A repeat bone biopsy performed after one year showed that 12 patients developed normal or

Figure 2. Changes in the serum level of calcium and phosphorus during PTH infusion for 48 hours.
A) Effect of parathyroidectomy (PTX). During the PTH infusion rats with normal renal function (creatinine, mg/dL (Cr)<0.5), moderate renal failure (Cr=0.5-0.6) and advanced renal failure (Cr>0.6) were fed a low phosphate (0.2%) and calcium free diet. The interrupted line indicates rats with PTX. B) Effect of high phosphate diet. Rats with normal renal function (creatinine, mg/dL (Cr)<0.5), moderate renal failure (Cr=0.5-0.6) and advanced renal failure (Cr>0.6) were fed a high phosphate (1.0%) and calcium free diet. Interrupted line indicates daily calcitriol administration of 40ng/100g.20

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bone formation rate similar to results seen in intestinal-specific VDR null mice. A positive calcium balance from calcium carbonate and high dialysate calcium concentration could also have contributed.

In the other study, a hemodialysis patient with markedly elevated intact PTH (1912 pg/mL, Nichols assay) and serum alkaline phosphatase (1512 IU/L) was studied before calcitriol activated intact PTH (1912 pg/mL, Nichols assay) and serum alkaline phosphatase (1512 IU/L) was studied before calcitriol treatment, after 6 weeks of 2 μg intravenous calcitriol thrice weekly, and 6 weeks after calcitriol was stopped. Studies consisted of separate dialysis sessions with low (1 meq/L) and high (4 meq/L) calcium dialysate to determine maximal PTH secretion and suppression respectively. Despite 6 weeks of calcitriol treatment, predialysis serum calcium and PTH values were similar to pretreatment values (Figures 3 A and 3 B) as were serum phosphorus and alkaline phosphatase values. The maximal PTH value during induction of hypocalcemia was similar approaching 4000 pg/mL during both low calcium dialysis studies (Figure 3 B). However, during the low calcium dialysis in the first study, serum calcium, which started at 10.8 mg/dL, only decreased to 9.4 mg/dL at 180 minutes. After calcitriol treatment, despite a similar starting serum calcium concentration, serum calcium decreased to less than 8 mg/dL after only 15 minutes of dialysis (Figure 3A). Studies with the high calcium dialysate showed marked differences in bone buffering of calcium. Before calcitriol treatment, serum calcium slowly increased to 11.8 mg/dL at 180 minutes. However, after calcitriol treatment, serum calcium increased to 12.2 mg/dL after only 15 minutes (Figure 3 C). PTH suppression was similar (Figure 3 D). When dialysis studies were repeated after calcitriol had been stopped for 6 weeks, results were similar to those before calcitriol was started (Figure 3). Thus, in this hemodialysis patient with severe secondary hyperparathyroidism and an enhanced capacity to defend against the induction of hypocalcemia and hypercalcemia, calcitriol treatment for 6 weeks dramatically decreased calcium efflux from and influx to bone.

In both cited studies, a decreased bone/calcemic response to PTH was seen. In the second study, the exchangeable pool of bone calcium was seemingly greatly reduced by calcitriol treatment despite no change in PTH secretion. In the first study it could be argued that the decrease in osteoblasts and osteoclasts, which reflects bone remodeling, is different from an evaluation of bone efflux and influx of calcium. However, the marked reduction in bone formation rate together with the decreased cellular activity seen after calcitriol treatment is also a likely indicator of a decreased exchangeable pool of bone calcium.

How does the vitamin D/calcium biochemical profile differ between dialysis patients receiving high doses of calcitriol and patients with vitamin D intoxication? In dialysis patients, calcitriol treatment reduced calcium efflux from bone during a low calcium dialysis while in patients with vitamin D intoxication increased calcium efflux from bone is reported to be a major contributor to hypercalcemia. In vitamin D intoxication, hypercalcemia is persistent with suppressed PTH. In dialysis patients treated with calcitriol, if monitored diligently the serum calcium concentration is normal while the high PTH value generally decreases, but remains elevated. In the study in which calcitriol was given thrice weekly, measured serum calcium values were in the normal range before subsequent dosing but high 1,25D values would be expected immediately after calcitriol dosing. In dialysis patients, 25OHD values are often less than the desired goal, 30 ng/mL. Conversely, in vitamin D intoxication, 25OHD values are markedly elevated while serum calcitriol values are often but not always increased.

How does vitamin D affect calcium efflux from bone in vitamin D intoxication? In vitamin D intoxication, long-standing hypercalcemia leads to reduced renal function further exacerbating hypercalcemia, which may act to drive calcium into bone. Treatment with normal saline improves renal function by correcting the decreased extracellular volume, which in turn, increases renal excretion of calcium lowering serum calcium. As a result, the supersaturated bone releases more calcium for several reasons. These include an increasing calcium gradient between bone and blood, reduced serum phosphorus from improved renal function, and increased PTH secretion from a reduction in serum calcium. Such actions could account for the enhanced calcium release from bone reported in vitamin D intoxication. Finally, suggesting that increased bone calcium deposition may occur during the hypercalcemia of vitamin D intoxication is the report of increased bone mineral density in osteoporotic patients with vitamin D intoxication.

Why did high dose calcitriol treatment decrease the exchangeable pool of bone calcium in the dialysis patient with severe secondary hyperparathyroidism? In dialysis patients with osteitis fibrosa, there are abundant osteoblasts in which the VDR can be activated to generate osteoclasts and increase bone resorption. However, calcitriol administration decreased the exchangeable pool of bone calcium. In both cited studies in patients with osteitis fibrosa, a calcium-replete environment was present due to high dialysate calcium, the use of calcium-based phosphate binders, and intermittent or continuous hypercalcemia. In contrast, in studies in intestinal-specific VDR-null mice in which 1,25D-mediated increased bone resorption and decreased bone formation were seen, calcium depletion was present. However, when the VDR-null mouse was calcium replete, increased bone resorption and decreased bone formation were prevented. Other important differences are present between dialysis patients with osteitis fibrosa and intestinal-specific VDR null mice/patients with vitamin D deficiency. In the dialysis patients, calcitriol administration was intermittent in contrast to continuously elevated, endogenous 1,25D values. Although PTH values are increased in all the cited conditions, the calcium and phosphorus status are quite different. In the dialysis patients, hypercalcemia and
hyperphosphatemia are present while in intestinal specific null mice and vitamin D deficient patients, hypocalcemia and hypophosphatemia prevail. In the reported dialysis patient,\textsuperscript{64} increases in serum calcium were not seen with calcitriol treatment. However, whether increased bone buffering of calcium not only prevented hypercalcemia, but also drove the mineralization process decreasing the exchangeable pool of bone calcium is an interesting possibility. A summary of the discussion on 1,25D and bone is provided Table 3.

A final point of discussion is the effect of both 19-nor-1,25-dihydroxy-vitamin D\textsubscript{3} (paricalcitol), an analog of 1,25D, and the calcimimetic, cinacalcet, on bone resorption and serum calcium. Both agents have been used for the past 10 to 15 years for the treatment of hyperparathyroidism in dialysis patients. To achieve similar PTH suppression, the dose ratio of paricalcitol to calcitriol used has been 3 or 4:1. Most, but not all in vitro and animal studies have shown that the calcemic effect of paricalcitol is less than that of calcitriol.\textsuperscript{69-71} In studies in dialysis patients, paricalcitol suppresses PTH secretion at least as well if not better than calcitriol\textsuperscript{72} and may also have less calcemic effect.\textsuperscript{72,73} In studies in uremic rats, less vascular calcification has been seen with paricalcitol than with calcitriol\textsuperscript{74} and calcimimetics appear to be even more effective in preventing vascular calcification.\textsuperscript{75} A summary of the discussion on 1,25D and bone is provided Table 3.

Calcimimetics are an effective treatment for hyperparathyroidism in the uremic animals and dialysis patients. Because of its calcimimetic activity, hypocalcemia generally develops

\begin{figure}[h]
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\includegraphics[width=\textwidth]{Figure3}
\caption{Serum calcium and PTH response to low and high calcium dialysate before, during and after calcitriol (CTR) treatment. Shown is the effect on serum calcium and PTH during hemodialysis with low (1meq/L) and high (4meq/L) calcium dialysates. These studies were performed before the start of calcitriol (Pre-CTR), at the end of 6 weeks of calcitriol treatment (CTR) and 6 weeks after calcitriol treatment was stopped (Post-CTR).\textsuperscript{64}}
\end{figure}
Table 3. Calcitriol and calcium efflux from bone

A. Classic Effects of Calcitriol
1) Increase in gut calcium absorption
2) Increase in gut phosphate absorption
3) Increase in renal calcium reabsorption
4) Suppression of PTH mRNA synthesis
5) Stimulation of FGF23 and klotho

B. Bone Effects of Calcitriol
1) Increases calcemic action of PTH in azotemic rats
2) Increase in gut phosphate absorption
3) Increase in renal calcium reabsorption
4) Suppression of PTH mRNA synthesis
5) Stimulation of FGF23 and klotho

C. Points of Discussion
1) Calcitriol decreases bone formation in dialysis patients
2) Long-term calcitriol administration in dialysis patients may decrease the exchangeable pool of bone calcium
3) Bone effect of calcitriol may differ between calcium deplete and replete states
4) In hypercalcemia from vitamin D toxicity, improved renal function from volume expansion increases calcium excretion and may increase calcium efflux from bone due to gradient effect
5) Paricalcitol may have less calcemic effect than calcitriol

Indoxyl sulfate and p-cresol sulfate, a gut microbial product produced from metabolism of tyrosine, may alter PTH action on bone through the FGF23-klotho axis. Klotho blood values are low in CKD while FGF23 values are markedly elevated. In a mouse model of CKD and in isolated kidney cells, both indoxyl sulfate and p-cresol sulfate suppressed klotho production by epigenetic modification of the klotho gene. These toxins stimulate expression of DNA methyltransferase 1 which hypermethylates the klotho gene suppressing gene expression. A deficiency of klotho as exemplified by the klotho knockout mouse results in elevated values of serum calcium, phosphorus and 1,25D, ectopic calcification including extensive vascular calcification, and osteoporosis with decreases in osteoblasts and osteoclasts. FGF23 values are markedly elevated in the klotho knockout model similar to that in CKD. Thus, reduced levels of klotho in CKD, perhaps at least in part due to uremic toxins, could alter the bone response to PTH via hyperphosphatemia and/or extremely elevated FGF23 levels found in CKD. The potential role of uremic toxins on bone is summarized in Table 4.

A summary of the many factors affecting calcium efflux from bone discussed in our review is shown in Figure 4. Many unanswered questions exist about the exchangeable pool of bone calcium. These include 1) how PTH mobilizes calcium from bone, 2) what is the contribution of osteoclast-mediated calcium efflux in contrast to shifts across the quiescent bone surface, 3) how after induced hypocalcemia, serum calcium is restored to baseline values in parathyroidectomized animals, 4) how parathyroidectomy restores the calcemic response to PTH to normal in the azotemic rat, 5) how phosphate modifies calcium efflux from bone, 6) why bisphosphonate treatment during treatment. Several studies in rats have evaluated the effect of calcimimetics on bone. Two studies in rats used a 5/6 nephrectomy model and the other an adenine model of uremia. The latter model results in much more severe renal failure, hyperparathyroidism, hypocalcemia and hyperphosphatemia than the former model. In one of the 5/6 nephrectomy models, a high phosphate diet was used and in the other a relatively high calcium diet was used. Moreover, different calcimimetics were used in the three studies. In all the studies, serum calcium and PTH decreased, but the magnitude of the reduction in PTH differed. Changes in bone histology also differed considerably among the studies. In another study simulating primary hyperparathyroidism, the bone response to a two week continuous infusion of PTH was evaluated in mice with either a PTH gene knockout or PTH and calcium sensing receptor gene knockout (double knockout). The mice had normal renal function. The absence of the calcium sensing receptor reduced the bone effects of PTH in this model of primary hyperparathyroidism.

UREMIC TOXINS

A variety of factors which could potentially contribute to skeletal resistance to PTH are altered in CKD. Exposure of osteoblast-like cells to uremic serum affects the number and function of these cells including suppressed mitogenesis and IL-6 release and reduced PTHR expression. In CKD, levels of inhibitory and stimulatory insulin growth factor binding proteins 4 and 5 respectively, are altered and may contribute to abnormal osteoblast proliferation. Os
teoprotegerin, a decoy receptor of NF-kB ligand (RANKL), accumulates as renal failure progresses blocking osteoclast activation by PTH.

The concentration of absorbed products of intestinal microbial flora increase as renal function declines. These products, particularly indoxyl sulfate and p-cresol sulfate, may alter bone metabolism and contribute to bone resistance to PTH. Indole, formed by gastrointestinal bacteria during breakdown of tryptophan, is metabolized to indoxyl sulfate which is transported by an organic anion transporter into osteoblasts where it induces oxidative stress. Exposure of osteoblast cells in culture to indoxyl sulfate inhibits PTH-induced cyclic AMP generation and downregulates PTHR expression, an effect partially inhibited by the organic anion transporter inhibitor, probenicid. Elevated indoxyl sulfate values in hemodialysis patients inversely correlate with markers of bone formation. Furthermore, in a rat model of adynamic bone in which elevated indoxyl sulfate values were present, administration of the oral absorbent, AST-120, lowered indoxyl sulfate values and increased bone formation.
Table 4. Uremic toxins and calcium efflux from bone

A. Classic Effects of Uremia
1) Decrease in calcemic action of PTH
2) Secondary hyperparathyroidism
3) Decrease in calcitriol values
4) Hyperphosphatemia/phosphate retention
5) Increase in FGF 23 values

B. Bone Effects of Uremia
1) Renal osteodystrophy

C. Points of Discussion
1) Uremic serum inhibits mitogenesis of osteoblast-like cells
2) Exposure of osteoblasts to indoxyl sulfate inhibits PTH-induced cyclic AMP generation and downregulates PTH receptor expression
3) Lowering indoxyl sulfate in rat model of adynamic bone increases bone formation
4) Both indoxyl sulfate and p-cresol sulfate suppress klotho production

Figure 4. Schematic of factors affecting calcium efflux from bone are shown. The plus (+) sign indicates that the factor increases calcium efflux from bone. The negative (-) sign indicates that the factor decreases calcium efflux from bone. The following abbreviations are used: calcium (Ca); phosphorus (P); parathyroid hormone (PTH); calcitriol (CTR); fibroblast growth factor 23 (FGF23); and parathyroid hormone receptor (PTHR).

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DISCLOSURES

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**Conflicts of interest**

The authors declare that they have no conflicts of interest related to the contents of this article.

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