Vitamin D in Kidney Disease: Pathophysiology and the Utility of Treatment

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Vitamin D physiology has gained more importance and publicity than any of its counterparts in the water- and fat-soluble vitamin groups combined. This is partly because vitamin D deficiency is still widely prevalent in the developed world and the beneficial effects are thought to extend beyond the regulation of calcium and phosphorus homeostasis alone. Vitamin D deficiency becomes even more important in the various stages of chronic kidney disease (CKD); CKD itself is also on the increase. How vitamin D physiology is altered in CKD and how the various treatment modalities can alter the morbidity and mortality associated with CKD is the topic of discussion for this article. Chronic kidney disease, mineral and bone disorder (CKD-MBD) is the broad term used to describe the disease complex of hyperphosphatemia, secondary hyperparathyroidism, low levels of vitamin D, and their associated complications.1 The National Kidney Foundation, Kidney Disease Outcomes and Quality Initiative (KDOQI), and subsequently, Kidney Disease: Improving Global Outcomes (KDIGO), has endeavored to provide evidence-based clinical practice guidelines for various stages of CKD and has set certain target levels for the major factors involved in CKD-MBD.2,3 These factors include calcium, phosphorus, vitamin D, and parathyroid hormone (PTH). Whether achieving and maintaining these targets will provide any benefit for the overall survival of patients with kidney disease remains to be demonstrated.

CALCIUM

Calcium in present abundantly in the body. The skeleton acts as reservoir and buffer for calcium such that when a large oral load of calcium is ingested and absorbed, it gets buffered in the skeleton and when calcium is acutely needed, it is mobilized...
from the skeleton. Regulation of serum calcium is under complex control with short-term and long-term regulation. This regulatory control is governed by vitamin D metabolism, phosphorus metabolism, fibroblast growth factor 23 (FGF-23), and PTH. These regulators of calcium homeostasis are heavily dependent on each other, and there are continuous positive and negative feedback mechanisms. For various reasons that are discussed later, these regulatory mechanisms of calcium homeostasis become disrupted in CKD and result in the presentation of CKD-MBD.

**CKD-MBD**

Patients with kidney disease are at a many fold higher risk of bone fracture than their age-matched controls. Hyperphosphatemia and hyperparathyroidism are virtually universal in patients with advanced kidney disease. Vascular calcification can be debilitating and is strongly associated with the increased cardiovascular morbidity and mortality associated with CKD.\(^4\)–\(^6\) Many patients in CKD stage 4 (estimated glomerular filtration rate [eGFR] 15–29) never seem to progress to CKD stage 5 (eGFR<15), and mortality from cardiovascular disease may be a contributing factor. CKD-MBD starts early, usually by CKD stage 3 (eGFR 30–59) and, for the most part, is a silent problem that only becomes manifest as CKD advances. Vitamin D deficiency is widely prevalent in the general population and even more so in patients with kidney disease.

**NORMAL VITAMIN D PHYSIOLOGY**

Vitamin D is crucial for calcium and phosphorus homeostasis and the regulation of parathyroid function. In addition to obtaining vitamin D from diet, a significant amount is formed in the skin (Fig. 1). Ultraviolet rays of the correct wavelength (UVB) in sunlight convert 7-dehydrocholesterol in the skin to previtamin D. This is then transported to the liver where it is hydroxylated at carbon 25 to form 25-hydroxyvitamin D [25(OH)D]. This is the main storage form of vitamin D in the human body and is the

![Diagram of vitamin D metabolism](image)

**Fig. 1.** Normal vitamin D metabolism showing the capacity of the kidney and extrarenal sites to produce the active vitamin D sterol, 1,25-dihydroxyvitamin D.
vitamin D metabolite that reflects the state of vitamin D nutrition. Almost all 25(OH)D is bound to circulating vitamin D–binding protein (DBP) and is then filtered by the kidney and taken up by the proximal convoluted tubule by an endocytic receptor, megalin. The 25(OH)D-DBP complex is degraded in proximal tubule lysosomes, releasing 25(OH)D, which then translocates to the mitochondria. In the mitochondria, 25(OH)D is converted to 1,25-dihydroxyvitamin D by the enzyme 1α-hydroxylase and returned to the circulation as the active form of vitamin D. New research has shown the presence of this enzyme in organs other than the kidney, such as pancreas, brain, lymph nodes, heart, gastrointestinal tract, adrenal glands, and prostate gland, such that 1,25-dihydroxyvitamin D may be made locally in these tissues. The biologic role of the extrarenal 1α-hydroxylase and the local effects of 1,25-dihydroxyvitamin D in extrarenal sites is the subject of ongoing studies.

The actions of 1,25-dihydroxyvitamin D are mediated by binding to the vitamin D receptor (VDR) and result in the alteration of the transcription of many genes in the various target organs.

**VITAMIN D METABOLISM IN KIDNEY DISEASE**

Kidney disease seems to be a risk factor for vitamin D deficiency and as many as 70% to 85% of patients with CKD are found to have low levels of 25(OH)D. Many factors may contribute, including lack of sunlight, loss of 25(OH)D-DBP with heavy proteinuria, diabetes, chronic illness, decreased production of previtamin D in the skin, and other unknown factors. Thus, the concentration of substrate for conversion to 1,25-dihydroxyvitamin D is decreased. This is further complicated by the fact that with decreases in glomerular filtration rate (GFR) and decreased renal mass, there is decreased delivery of substrate to the renal 1α-hydroxylase, which limits the ability of the diseased kidney to produce the active 1,25-dihydroxyvitamin D. In addition, as CKD develops, phosphate retention occurs decreasing 1α-hydroxylase activity, directly and leading to increases in the levels of FGF-23, which in turn, can directly decrease the activity of 1α-hydroxylase. FGF-23 is a recently discovered phosphaturic hormone that is regulated by dietary phosphate, serum phosphate, and 1,25-dihydroxyvitamin D. FGF-23 levels increase early in CKD, presumably in response to phosphate retention, in an effort to increase phosphate excretion in conjunction with increases in PTH. Although PTH stimulates the activity of 1α-hydroxylase, the suppressive effect of FGF-23 on 1α-hydroxylase activity seems to dominate in this clinical situation. In addition, it has been suggested that accumulation of N-terminally truncated PTH peptides of C-terminal PTH fragments may decrease 1α-hydroxylase. Thus, because of these abnormalities (Fig. 2), it is not surprising that the levels of 1,25-dihydroxyvitamin D are reduced in CKD and progressively decline with advancing stages of CKD. In addition to decreased production of 1,25-dihydroxyvitamin D, there is also evidence for resistance to the actions of vitamin D as kidney disease progresses in that there may be decreased concentrations of the VDR, impaired binding of the 1,25-dihydroxyvitamin D binding to the VDR, and possibly impaired binding of the VDR complex to the vitamin D response elements in the nuclei. This altered vitamin D physiology in patients with renal disease contributes to subclinical or less commonly overt hypocalcemia and leads to the need for increased levels of PTH, which mobilizes minerals from the skeleton by stimulation of osteoclastic-mediated bone resorption. This process leads to the loss of lamellar bone and replacement with the woven and structurally weaker bone, increased propensity for fractures, and overall decreased quality of bone.
Abnormal phosphorus metabolism is a major factor in CKD-MBD. Phosphate retention, as a consequence of decreased GFR, is thought to be a major factor in the pathogenesis of CKD-MBD. Phosphorus retention can lead to increased PTH directly and indirectly by increasing the levels of FGF-23, which, in turn, decreases the activity of 1α-hydroxylase. High levels of phosphorus have been shown to promote vascular calcification in animal models and are associated with vascular calcification in experiments in vitro.

ROLE OF VITAMIN D TREATMENT IN CKD-MBD

Because of the major role of abnormal vitamin D metabolism in the disturbances of calcium and phosphorus homeostasis and in the pathogenesis of secondary hyperparathyroidism in the setting of CKD, use of vitamin D sterols is an important aspect of the therapy of patients with CKD. Current practice guidelines suggest evaluation for the presence of hyperparathyroidism early in the course of CKD, and if PTH values are elevated, vitamin D deficiency should be evaluated by measurement of 25(OH)D levels. If 25(OH)D levels are less than 30 ng/mL then vitamin D supplementation should be initiated. This is most often accomplished by the administration of ergocalciferol or cholecalciferol. Although the KDOQI guidelines suggest a dosage regimen for ergocalciferol, this is not always effective in achieving correction of the low levels of 25(OH)D in this patient group. The reasons for this are currently unclear. However, if the levels of 25(OH)D are increased then PTH values are seen to decrease in CKD stage 3. The decrease in PTH in patients with CKD stage 4 seems to be less, but again, there is marked heterogeneity in response to ergocalciferol. Accordingly, the use of active vitamin D sterols can be considered. The preparations available in North America are: calcitriol (1,25-dihydroxyvitamin D₃), doxercalciferol (1α-25-hydroxyvitamin D₂), and paricalcitol (19-nor-1,25-dihydroxyvitamin D₃). All are effective in reducing the secondary hyperparathyroidism associated with CKD, consistent with the finding that activation of the VDR in parathyroid glands results in decreased PTH gene transcription. Dosing should be monitored carefully to avoid toxicity, which is mainly manifested by hypercalcemia. In studies in animals, the analogue, paricalcitol, has been shown to be less calcemic and less phosphatemic than the native hormone. There are limited data on head-to-head studies of the active vitamin D sterols in patients, but paricalcitol seems to have the widest therapeutic window. It is likely that early
recognition of secondary hyperparathyroidism and initiation of therapy early in the course of CKD may lead to effective control of hyperparathyroidism and prevent parathyroid growth.

EFFECTS OF VITAMIN D ON THE CARDIOVASCULAR SYSTEM

Studies have shown that VDR knockout mice have an overactive renin-angiotensin-aldosterone system (RAAS) and develop hypertension and left ventricular hypertrophy (LVH). Similarly, Dahl salt-sensitive rats develop vitamin D deficiency and LVH with diastolic dysfunction, and the administration of paricalcitol has been shown to be associated with decreased LVH in these rats. Similarly, 1α-hydroxylase knockout mice develop hypertension, cardiac hypertrophy, and depressed cardiac function. These effects are not corrected by correction of calcium and phosphorus alone. Calcitriol, however, seems to ameliorate hypertension and improve cardiac function in this animal group. Therefore, it seems that the vitamin D system is a regulator of the renin-angiotensin system. These observations, coupled with other animal studies that have shown improved cardiac function and decreased LVH and the suggestion that 1,25-dihydroxyvitamin D may actually suppress the RAAS axis, have led to the initiation of 2 prospective trials in patients to evaluate this possibility. The Paricalcitol benefits in Renal Failure Induced Cardiac Morbidity (PRIMO) trials, PRIMO I and PRIMO II will evaluate the beneficial effects of paricalcitol in patients predialysis and those on dialysis, respectively (clinicaltrials.gov NCT00497146 and NCT00616902).

VITAMIN D AND OVERALL SURVIVAL

Some observational studies have shown decreased overall mortality in patients with end-stage renal disease (ESRD) who are being treated with an active vitamin D sterol. In 1 study there was a 26%, 2-year reduction in mortality in the patient groups who received some form of vitamin D versus those who received none. In another observational study by Teng and colleagues, in 60,000 patients undergoing hemodialysis there were lower rates of mortality in those treated with paricalcitol versus calcitriol. Other studies have also shown similar survival benefits with the use of paricalcitol. Melamed and colleagues (KI2006) noted a 26% reduction in mortality with injectable vitamin D as part of the Choices for Health Outcomes in Caring for ESRD (CHOICE) study. Tentori and colleagues showed a similar survival benefit irrespective of the formulation of the active vitamin D sterol (calcitriol, paricalcitol, or doxercalciferol). Similarly, Kalantar-Zadeh and colleagues showed paricalcitol administration to be associated with improved survival compared with those receiving no vitamin D in patients with ESRD. Similar apparent survival benefits associated with the use of active vitamin D sterols are now being presented in predialysis patients. Indeed, low levels of 25(OH)D have been associated with greater mortality in patients with cardiovascular disease as well as in patients with hypertension, and even in apparently normal people. These observational studies warrant the need for prospective, randomized trials to evaluate any survival benefit of vitamin D therapy in patients with CKD.

OTHER POTENTIALLY BENEFICIAL EFFECTS OF VITAMIN D TREATMENT IN CKD

Vitamin D metabolism has a role in immune function, which was elucidated by Liu and colleagues in studies evaluating the response of human macrophages to activation of toll-like receptors. These studies demonstrated that the antimicrobial protein cathelicidin was regulated by the generation of 1,25-dihydroxyvitamin D by the
macrophage 1α-hydroxylase. This pathway may be important in CKD, which has a high incidence of infections. Indeed, an association between low levels of the cathelicidin, hCAP18, and death from infectious causes in patients on hemodialysis has been observed.42

It is also possible that the vitamin D system may affect the progression of CKD. Several studies in animals show that administration of active vitamin D sterols can favorably affect the processes that lead to progression of kidney disease.43–50 Recently, studies have shown that paricalcitol seems to reduce proteinuria in patients with CKD,51 and this has been confirmed in a randomized controlled trial.52

SUMMARY

CKD is associated with decreased vitamin D metabolites, both the storage form 25(OH)D and the active form 1,25-dihydroxyvitamin D. This contributes to hyperparathyroidism, and increased levels of PTH mobilize minerals from the skeleton and increase the risk for fractures. Treatment with vitamin D sterols efficiently reduces secondary hyperparathyroidism of CKD. Observational studies suggest survival and other potential benefits of vitamin D treatment in the CKD population. These observations need to be verified with controlled prospective trials.

REFERENCES

