Vitamin D supplementation in pre-dialysis chronic kidney disease
A systematic review

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Abbreviations: 25(OH)D, 25-hydroxyvitamin D or calcidiol; 1,25(OH)2D, 1,25-dihydroxyvitamin D or calcitriol; [Ca2+]i, intracellular calcium concentration; BAP, bone-specific alkaline phosphatase; CKD, chronic kidney disease; CVD, cardiovascular disease; eGFR, estimated glomerular filtration rate; ESRD, end stage renal disease; FGF-23, fibroblast growth factor-23; GFR, glomerular filtration rate; KDOQI, Kidney Disease Outcomes Quality Initiative; PTH, parathyroid hormone; VDBP, vitamin D binding protein

Vitamin D deficiency is associated with a variety of skeletal, cardiometabolic, and immunologic co-morbidities that are present in chronic kidney disease (CKD). We performed a systematic review to investigate the effects of vitamin D supplementation, in the form of ergocalciferol or cholecalciferol, on various health outcomes in early CKD. Seventeen clinical trials were identified, only two of which were randomized, placebo controlled trials. The majority of studies supplementing with > 2,000 IU/day of cholecalciferol achieved optimal vitamin D status, whereas studies supplementing with ergocalciferol were less consistent. Studies varied widely in their effects on lowering serum parathyroid hormone concentrations. Few studies investigated effects of vitamin D treatment on other clinical health indicators in early CKD. Rigorous studies are necessary to investigate optimal vitamin D dosing strategies in early CKD for the maintenance of adequate vitamin D status, management of secondary hyperparathyroidism and improvement of non-skeletal related clinical outcomes.

Introduction

With the increasing prevalence of obesity, hypertension, and type 2 diabetes, there has also been an increase in the prevalence of chronic kidney disease (CKD).1 Complications of CKD include progression to end stage renal disease and need for costly dialysis or renal transplantation, bone disease and premature cardiovascular disease (CVD) morbidity and mortality.2 Strategies to manage CKD are thus a high priority, not only from a clinical, but also a public health perspective.3 Vitamin D may play a key role in the disease management of CKD. Accumulating evidence from epidemiological and experimental research suggests that vitamin D is integral not only for its classical effects on the skeletal system, but also for its extra-skeletal benefits such as those on cardiovascular health and immune function.4,5 In this regard, a recent meta-analysis indicated that higher circulating 25-hydroxyvitamin D (25(OH)D or calcidiol) concentrations were associated with lower all-cause mortality risk in all stages of CKD.6

Altered vitamin D metabolism and resultant changes in bone and mineral metabolism are key features of CKD. The kidneys are the primary site of 25(OH)D conversion to the hormonal form of vitamin D, calcitriol (1,25(OH)2D), by the enzyme 1α-hydroxylase.7 As kidney disease progresses and renal mass decreases, there is a decline in the availability of 1α-hydroxylase enzyme resulting in a decrease in calcitriol, followed by compensatory increases in parathyroid hormone (secondary hyperparathyroidism) to offset impaired intestinal calcium absorption and resultant hypocalcemia.8,9 Other abnormalities of CKD, such as elevated fibroblast growth factor (FGF)-23,10,11 accumulation of parathyroid hormone (PTH) fragments (N-terminally truncated or C-terminal)12 and uremic toxins,13 and reductions in glomerular filtration rate (GFR) and megalin contribute to calcitriol defects in CKD.8 Because of defects in renal calcitriol production, studies in the past have primarily focused on the use of calcitriol replacement (or other active vitamin D analogs) to control hyperparathyroidism in CKD.2,13 However, there is evidence that patients with end stage renal disease (ESRD) retain some capacity to produce calcitriol from 25(OH)D, either through residual renal functioning or extra-skeletal production of calcitriol.14,15 Extra-renal cells, such as parathyroid cells, smooth muscle cells, endothelial cells, pancreatic cells and immunomodulatory cells, contain the machinery to locally produce calcitriol.16 This may explain the associations of adequate vitamin D status with lower chronic disease risk. Thus, the ability to maintain sufficient serum concentrations of 25(OH)D, the substrate for extra-renal 1α-hydroxylase and local production of active vitamin D, is particularly important in CKD.

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Vitamin D players: vitamin D, cholecalciferol, ergocalciferol, renal disease, secondary hyperparathyroidism, chronic kidney disease

GFR, glomerular filtration rate; KDOQI, Kidney Disease Outcomes Quality Initiative; PTH, parathyroid hormone; VDBP, vitamin D binding protein

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Recent reports estimate up to 84% prevalence of vitamin D deficiency and insufficiency (defined in this review as 25(OH)D < 20 ng/ml and 25(OH)D < 30 ng/ml, respectively) in CKD populations. Reasons for high prevalence of vitamin D deficiency in the CKD population compared with the general population are not clear. Elevated proteinuria and thus loss of urinary vitamin D binding protein (VDBP), reduced recycling of 25(OH)D by megalin in proximal tubular cells, and differences in diet, reduced sun exposure, and other lifestyle factors such as limited outdoor activity may provide explanations. Given the dysregulated vitamin D metabolism and the high prevalence of vitamin D deficiency in patients with CKD, the increasingly recognized pleiotropic benefits of the vitamin D system, and the elevated risk of premature morbidity and mortality associated with both CKD and vitamin D deficiency, optimizing vitamin D status through supplementation may be of particular relevance in CKD patients, specifically in predialysis CKD wherein renal capabilities of 25(OH)D conversion to calcitriol have not been exhausted. To this end, the purpose of this review is to investigate the effects of vitamin D supplementation (cholecalciferol or ergocalciferol) on health outcomes and biomarkers related to mineral and bone metabolism, CKD progression, diabetes, and cardiovascular disease in predialysis stages of chronic kidney disease (Stages 2 to 4).

Materials and Methods

We searched the terms “cholecalciferol OR ergocalciferol AND chronic kidney disease” in PUBMED through the present date (January 30, 2011). Limits were pre-set to manuscripts published in the English language. Titles and abstracts were reviewed. Manuscripts were further excluded for review if (1) they did not use oral cholecalciferol or ergocalciferol; (2) they used analog compounds of vitamin D such as calcitriol, doxercalciferol or paricalcitol; (3) study participants received dialysis; or (4) studies involved renal transplant patients. Additional manuscripts were identified from the references of the selected manuscripts and selected recent reviews. Two independent reviewers (J.A., H.W.) reviewed the literature with these search criteria. When there was disagreement regarding inclusion of the manuscript for this systematic review, a third reviewer (V.T.) determined whether the manuscript was eligible.

Specific outcomes of interest included: (1) serum/plasma 25(OH)D, (2) serum/plasma PTH, (3) albuminuria/proteinuria, (4) glomerular filtration rate and progression across kidney stages and to dialysis, (5) measures and biomarkers of insulin resistance, glucose intolerance, and glycemic control, (6) blood pressure/hypertension, (7) measures and biomarkers of vascular function, (8) inflammation, oxidative stress, and measures of immunity, (9) mortality and (10) CVD events.

PubMed search results. One thousand and sixty-eight manuscripts were identified from the specified search terms (Fig. 1). After applying additional exclusion criteria, 22 manuscripts were identified as potential eligible studies based on titles and abstracts. After review of full-text, 1 was excluded because it was a publication of a protocol, 3 were excluded because they were in dialysis or end stage renal disease patients and 5 were excluded because of use of active vitamin D or analog or calcidiol. Two studies were identified from additional sources and two were
Vitamin D therapy on serum/plasma PTH. PTH significantly decreased in eight studies, with a variety of dosing protocols including both ergocalciferol and cholecalciferol. Ökka et al. reported similar decreases in PTH using either 5,000 IU or 20,000 IU cholecalciferol per week (~714 or 2857 IU/day). Hari et al. reported greater median reductions in PTH in Stage 2 (38.5%) compared with Stages 3 (25.2%) and 4 (15.2%). Both Al-Aly and Zisman et al. reported a significant decrease in PTH in Stage 3 (21.8 and 13.1% decrease) but not Stage 4 (11.3 and 2% decrease) CKD patients. Deville et al. reported a significant decrease in PTH in Stage 4 (17.9%) but not Stage 3 (12.7%) or five patients (7.0%), although the differences were attributed to sample size. Shroff et al. reported a significant effect of ergocalciferol treatment in preventing hyperparathyroidism compared with placebo in children.

Vitamin D therapy on other markers related to bone metabolism. Taner et al. reported an increase in serum calcium and phosphorus after 300,000 IU cholecalciferol per month for 12 wks, although the change was not significantly different than untreated controls. Qinibi et al. reported a statistically significant increase in serum phosphorus with no change in serum calcium in a retrospective study following KDOQI guidelines. Other studies reported no change in serum phosphorus or calcium.

Chandra et al. reported no changes in bone-specific alkaline phosphatase (BAP), C-telopeptide, or TRAP5b using 50,000 IU cholecalciferol per wk for 12 wks. Kovesdy et al. also did not find a significant change in BAP after ergocalciferol treatment for 16 wks, although there was a significant reduction in BAP using paricalcitol.

Vitamin D therapy on other markers related to CKD progression. Rucker et al. reported a decrease in estimated GFR (eGFR; 21 ± 8 to 20.5 ± 7.8 ml/min per 1.73m², p = 0.01) after cholecalciferol supplementation compared with controls; serum creatinine concentrations were not provided. Ökka et al. reported a decrease in eGFR in patients receiving 20,000 IU cholecalciferol/wk [mean eGFR (range): 51 (15–89) to 50 (15–82) ml/min/1.73m², p < 0.01] but not with 5,000 IU/wk after 1 y [mean eGFR (range): 43 (15–89) to 42 (7–80), p > 0.05]; serum creatinine increased in both groups. Four studies reported no change in eGFR.

Moe et al. reported a non-significant decrease in urine albumin/creatinine ratio over 12 wks that did not differ between treatment with cholecalciferol or doxercalciferol (pooled mean baseline and follow-up urine albumin/creatinine ratio: 853 ± 1153 to 471 ± 770 µg/mg). No studies reported progression to dialysis as an outcome.
Table 1. Summary of cholecalciferol or ergocalciferol trials in early CKD

| Author, yr   | Design, Intervention                      | Duration/ Follow-up | Participants                     | CKD stage | Baseline, Post 25(OH)D (ng/ml)* | Baseline, Post PTH (pg/ml)* | Other Outcomes                                      |
|--------------|--------------------------------------------|---------------------|----------------------------------|-----------|-------------------------------|----------------------------|-----------------------------------------------------|
| **Trials with daily cholecalciferol**                                                                                                           |
| Rucker 2009  | Prospective, randomized, controlled        | 12 wks              | 128 adults, age 69.0 ± 12.6 y    | 3–5       | 16 ± 5.6, 26.8 ± 10.4*        | 144.4 ± 81.7, 130 ± 72.2 | ↓ GFR                                               |
|              |                                             |                     |                                  |           |                               |                            | ↓ Serum Ca, P, albumin                               |
| Moe 2010     | Prospective, single-blind, randomized, active-controlled | 12 wks              | 47 adults, age 63.6 ± 10.2 y     | 3/4       | 14.0 ± 6.1, 37.2 ± 10.1*      | 109 ± 43, 97 ± 49       | ↔ Serum Ca, Serum P, urine ca/cr, urine alb/cr, blood pressure |
|              |                                             |                     |                                  |           |                               |                            | 4,000 IU/day × 1 mo then 2,000 IU/d vs Doxercalciferol (1 µg/d) |

| Hari 2010    | Prospective, observational                 | 6 weeks             | 42 children, age 7 (2–15) yr    | 2–4       | 17.9 (12.8–20.9)e, 48.2 (44–56.1)e* | 55.3 (47.1–77.8), 41.4 (31.2–56.8)e* | ↔ Serum Ca, P, alk phos, eGFR |
| Dogan 2008   | Prospective, randomized, controlled        | 4 wks               | 40 adults, age 49 ± 14 y         | 3/4       | 8.5 ± 3.6, 17.8 ± 21.4*         | 368 ± 274, 279 ± 179*  | ↔ Serum Ca, P, BUN, creatinine albumin               |
| Lajdova 2009 | Prospective, observational                 | 12 mo               | 87 adults, age 66 (19–88) yr     | 2–4       | 15 (5–60), 28 (14–72)*          | A: 63 (13–224), 48 (11–181)* | ↑1,25(OH)2D; ↑ Serum Cr (both groups), ↑ Urine Ca at month 8, ↑ GFR (group B) |
|              |                                             |                     |                                  |           | B: 16 (4–49), 37 (8–81)e*       | A: 50 (10–184), 40 (11–203)* | ↔ Serum Ca, P, alk phos, eGFR |
| Okša 2008    | Prospective, randomized, open-label        | 12 mo               | 44 adults, age 62 (19–77) yr     | 2/3       | Duplicate analysis of parent study | Duplicate analysis of parent study | ↓ [Ca2+], |
|              |                                             |                     |                                  |           |                               |                            | 5,000 IU/wk (714 IU/day) vs. B: 20,000 IU/wk (2857 IU/day) |
| Chandra 2008 | Prospective, double-blind, randomized, placebo-controlled | 12 wks              | 20 adults, age 60.9 ± 10.7 y     | 3/4       | 17.3 (11.8–25.2), 49.4 (33.9–72.0)* | 288.9 (178.7–467.2), 200.5 (114.0–352.8) | ↔ Serum Ca, 1,25(OH)2D, BAP, CTX, TRAPSb |
| Taner 2011   | Prospective, randomized, controlled        | 12 wks              | 48 adults, age 57.5 ± 10.2 y     | 3/4       | 13.4 ± 5.9, 82.9 ± 30.9*        | 134.3 ± 92.5, 93.8 ± 67.7* | ↑ serum Ca, P |
|              |                                             |                     |                                  |           |                               |                            | 300,000 IU/month (10,714 IU/day) vs. no treatment    |

| **Trials with weekly or monthly cholecalciferol**                                                                                               |
| Oka 2008     | Prospective, randomized, open-label        | 12 mo               | 85 adults, age 67.2 ± 11.8 y     | 3–5       | 17.0 ± 8.0, 42.1*              | 176.2 ± 112.2, 148.9* | ————                                               |
| Lajdova 2009 | Prospective, observational                 | 12 mo               | 22 children, age 10.7 ± 5.4 y    | 2–4       | Not reported                    | 122.1 ± 82.9, 80.1 ± 59.2* | ————                                               |
| Deville 2006 | Prospective, observational                 | 90 d                | 20,000 IU/day to 100,000/wk      | 3–5       | 17.0 ± 8.0, 42.1*              | 176.2 ± 112.2, 148.9* | ————                                               |
| Menon 2008   | Retrospective chart review                 | 12 wks              | 40 children, age 9.3 ± 3.5       | 2–4       | 20.0 ± 7.8, 33.3*              | 1,25(OH)2D            | ↔ Serum albumin                                  |
| Shroff 2012  | Prospective, double-blind, randomized, placebo-controlled | 6 mo-2 yrs          | 2,000–8,000 IU daily (KDOQI)     | 2–4       | 20.0 ± 7.8, 33.3*              | 1,25(OH)2D            | ↔ Serum albumin                                  |
Vitamin D therapy on CVD events and mortality. Lishmanov et al.\textsuperscript{41} reported fewer CVD events leading up to a mean follow-up of 27.2 mo in patients whose serum 25(OH)D concentrations increased by at least 25% [from baseline 25(OH)D < 30 ng/ml] after 6 mo of ergocalciferol compared with those who were not treated or did not respond to treatment [OR (95% CI): 0.37 (0.14–1.0); p = 0.05], after adjusting for age, baseline PTH, statin use, CVD history, diabetes status, and eGFR. All-cause survival and CVD-specific survival were higher in the treated group (p = 0.008 and 0.02, respectively). It should be noted that the treatment group had a lower history of diabetes compared with controls (53% vs. 75%, p = 0.02).

Vitamin D therapy on measures and biomarkers of cardiovascular risk. Only two studies reported outcomes specifically related to CVD risk,\textsuperscript{31,32} although they were not primary outcomes of the studies. Moe et al.\textsuperscript{32} reported a non-significant decrease in home-measured systolic and diastolic blood pressure (average of 1 week measurements; p = 0.17 and 0.11, respectively), yet no change in clinic-measured blood pressure (p > 0.40) after 2,000–4,000 IU cholecalciferol per day for 12 wks. Kovesdy et al.\textsuperscript{31} reported no change in cholesterol, B-type natriuretic peptide, pulse wave velocity, homocysteine, or body fat after ergocalciferol treatment, although there was a trend toward an increase in HDL cholesterol (p = 0.05).

Lajdova et al.\textsuperscript{45} reported a decrease in peripheral blood mononuclear cell intracellular calcium concentration ([Ca\textsuperscript{2+}]) after 5,000 IU/wk cholecalciferol treatment for 1 y, although [Ca\textsuperscript{2+}], was not the primary endpoint of the study. No studies reported outcomes related to measures and biomarkers of insulin resistance, glucose intolerance, or glycemic control.

Vitamin D therapy on inflammation, oxidative stress and measures of immunity. Kovesdy et al.\textsuperscript{31} measured C-reactive protein as a marker of inflammation, but did not find a significant change after ergocalciferol treatment. No studies have reported outcomes related to oxidative stress or immunity.

Adverse events. Hypercalcemia and hyperphosphatemia were reported in only one study,\textsuperscript{40} among stage 4 CKD patients receiving ergocalciferol based on KDOQI guidelines.\textsuperscript{2} This
was an observational trial, and incidences were resolved spontaneously.

Discussion

Several health outcomes and co-morbidities that present in the early stages of CKD, including those related to bone and mineral disturbances, as well as cardiometabolic risk factors, have been linked to vitamin D deficiency, which is highly prevalent in patients with CKD. Moreover, vitamin D deficiency has been shown to be a predictor of CKD progression and death in this population in clinical and epidemiologic studies. We have shown cholecalciferol to be more effective in raising and maintaining serum 25(OH)D concentrations in patients with CKD. However, a comparison of ergocalciferol in a head-to-head trial in raising 25(OH)D concentrations in patients with CKD. Additionally, vitamin D deficiency has been linked to vitamin D deficiency, which is highly prevalent in patients with CKD. Moreover, vitamin D deficiency has been shown to be a predictor of CKD progression and death in this population in clinical and epidemiologic studies.6,27,47-49

We conducted a systematic review of the published literature to investigate the effects of vitamin D supplementation (in the form of cholecalciferol or ergocalciferol) in patients with pre-dialysis stage CKD. The studies reviewed varied widely in the form of vitamin D, dosage and protocol. The reported outcomes primarily reflected bone and mineral outcomes, such as changes in serum/plasma 25(OH)D and PTH. There was limited data available with respect to extra-skeletal outcomes.

The National Kidney Foundation recommends an optimal serum/plasma 25(OH)D of greater than or equal to 30 ng/mL to define optimal vitamin D status in patients with CKD. Although all studies reviewed noted a significant improvement in 25(OH)D concentrations, a mean optimal vitamin D concentration was not achieved by all and may be due to several reasons including an insufficient dose of vitamin D and/or use of a less effective form of vitamin D (ergocalciferol, as opposed to cholecalciferol). In general the studies supplementing with the equivalent of 700 to 1,000 IU vitamin D per day34,36 did not achieve mean optimal vitamin D status. In a dosage comparison study, Okša et al. found that 20,000 IU cholecalciferol/wk (-2,857 IU/day), but not 5,000 IU/wk (-714 IU/day) was sufficient to achieve optimal status in CKD patients. Collectively, studies in this review suggest daily doses of vitamin D > 2,000 IU/day (or weekly/monthly equivalents) are required to achieve optimal vitamin D status.

The KDOQI guidelines recommend the use of ergocalciferol with a specific dosing strategy to treat vitamin D deficiency/insufficiency; however, several studies in healthy participants have shown cholecalciferol to be more effective in raising and maintaining 25(OH)D concentrations than ergocalciferol.50-52 No study has compared the effectiveness of cholecalciferol vs. ergocalciferol in a head-to-head trial in raising 25(OH)D concentrations in patients with CKD. However, a comparison of the placebo-controlled cholecalciferol trial in CKD by Chandra et al. to ergocalciferol trials with similar baseline serum 25(OH)D values and dosing strategies supports the hypothesis that cholecalciferol is more effective than ergocalciferol in CKD. Taken together, these data suggest KDOQI guidelines may need to be updated to recommend the use of cholecalciferol as opposed to ergocalciferol. Further study is needed to establish effective vitamin D dosing protocols specific to patients with CKD to achieve and maintain optimal serum 25(OH)D concentrations.

Alleviation of secondary hyperparathyroidism is a major target for CKD management and the most commonly investigated outcome for vitamin D supplementation in early CKD, as indicated by this review. Vitamin D is a known regulator of PTH secretion; however the effective amount and form needed to manage or prevent secondary hyperparathyroidism in early CKD are not known. The 2003 KDOQI guidelines made opinion-based recommendations to assess serum 25(OH)D concentrations and treat vitamin D insufficiency with ergocalciferol in patients with CKD Stages 3 and 4 only if PTH concentrations are elevated.2 Several studies identified in this review used modified vitamin D dosing strategies modeled after KDOQI and found varying effects on PTH. The sole study published following the exact 2003 KDOQI protocol did not find a PTH-lowering effect of ergocalciferol treatment.44 Kandula et al. performed a meta-analysis on the PTH lowering effects of any form of vitamin D (including calcitriol) or vitamin D analog and reported a significant decrease in PTH in early CKD patients. A 30% decrease in PTH has been cited as being clinically effective.42 In the only randomized, double-blind, placebo-controlled trial of vitamin D adults with early CKD, serum PTH decreased -30% after 12 weeks of cholecalciferol treatment, although the sample size may have not been large enough to detect a statistically significant change in PTH.29 Given the wide variability in the design of published vitamin D supplementation trials and the corresponding variability in PTH lowering effects, it is clear that further randomized, placebo-controlled trials are necessary to establish an optimal treatment strategy with vitamin D to manage secondary hyperparathyroidism.

CKD severity plays a role in the effectiveness of vitamin D in reducing serum PTH. Two studies reported a reduction in serum PTH in patients with Stage 3 but not Stage 4 CKD.40,42 These data suggest that vitamin D is most beneficial in lowering serum PTH when 1α-hydroxylase is more active.45 In this regard, it may be prudent to test and correct 25(OH)D concentrations prior to elevations in serum PTH among early stage CKD patients, despite current KDOQI guidelines.7 Indeed, ergocalciferol treatment in children with CKD stage 2–4 delayed the onset of secondary hyperparathyroidism compared with placebo, suggesting that optimization of vitamin D status may be especially important during early CKD. This remains to be examined in adult CKD patients.

Only two studies have investigated the use of active vitamin D analogs vs. supplemental vitamin D in CKD. One clinical trial showed paricalcitol more effectively suppressed PTH.31 The other suggested doxercalciferol may be better at reducing PTH but did not significantly differ from cholecalciferol.32 It is hypothesized that supplemental vitamin D may be a better choice given its lower cost, fewer side effects and potential extra-renal benefits that require adequate serum 25(OH)D as a substrate for conversion to 1,25(OH)₂D to act as a paracrine/autocrine hormone in local tissue.17 Studies have not investigated differences between vitamin D forms on other outcomes in this regard. Vitamin D analogs have been hypothesized to interfere with local paracrine/autocrine VDR activation and effects by inhibiting local 1α hydroxylase or promoting 24-hydroxylase expression thereby suppressing local calcitriol production.6,54 Furthermore, adequate serum 25(OH)D is able to suppress PTH synthesis and secretion in parathyroid tissue.17 Studies have not investigated differences between vitamin D forms on other outcomes in this regard. Vitamin D analogs have been hypothesized to interfere with local paracrine/autocrine VDR activation and effects by inhibiting local 1α-hydroxylase or promoting 24-hydroxylase expression thereby suppressing local calcitriol production.54
cells. These reasons underscore the importance of maintaining adequate serum 25(OH)D levels in CKD.

The presence of proteinuria in CKD is an independent predictor for disease progression to ESRD and mortality.56,57 Cross-sectional studies indicate inverse relationships between circulating 25(OH)D and albuminuria in patients with CKD.58,59 Experimental evidence suggests that vitamin D may protect against or reduce proteinuria via its inhibitory effects on renin gene transcription and subsequent angiotensin II production;60 via inhibition of renal TNFα converting enzyme expression;61,62 via direct upregulation of nephrin;63 and/or via upregulation of renal megalin expression.64 Paricalcitol has been shown to significantly decrease proteinuria in clinical trials of patients with CKD.65-68 and may have anabolic effects, as indicated by increases in serum creatinine and blood urea nitrogen.69 eGFR is highly influenced by serum creatinine which may explain the reported decreases in eGFR identified in the review after cholecalciferol supplementation.34,36 Only one clinical trial of vitamin D supplementation identified by this review reported albuminuria as an outcome, albeit secondary, and did not observe a significant reduction in albuminuria.52 Given the possibility that proteinuria may promote vitamin D deficiency through urinary loss of VDBP,21 future clinical trials aimed at raising serum 25(OH)D concentrations with cholecalciferol and using proteinuria as a primary outcome are warranted. As accelerated protein energy wasting occurs with progressive CKD,70 the implications for a vitamin D protective effect on muscle integrity in this population should also be further explored.

Low circulating 25(OH)D concentrations predict increased all-cause mortality risk in pre-dialysis CKD patients in large cohort studies.6,27,67,69 Mortality was reduced by 26% in early CKD patients who received calcitriol, and this was independent of PTH and other risk factors.71 The mechanisms mediating a reduced mortality risk with higher vitamin D status are yet unknown; however vitamin D may play a role in infection-associated morbidity and mortality, which is significantly greater among patients with CKD.72 Calcitriol promotes upregulation of antimicrobial defenses by the immune system,53,73-75 and a higher rate of infection-associated mortality has been linked to severe vitamin D deficiency in ESRD.76 No studies have investigated the relationship between vitamin D status and infection in early chronic kidney disease.

The cardioprotective effects of vitamin D may also mediate the associations between better vitamin D status and reduced mortality in CKD. Vitamin D deficiency in ESRD patients is associated with an increased risk of CVD events.77 In contrast, a meta-analysis did not find a benefit of vitamin D analogs on cardiovascular outcomes.78 We identified only one study to-date examining the effect of vitamin D supplementation on CVD events in CKD. In a retrospective data review, Lishmanov et al.71 reported a reduced incidence of CVD events (albeit not statistically significant), as well as lower all-cause and CVD-related mortality, in early CKD patients successfully treated with ergocalciferol. Findings were independent of changes in PTH, suggesting other mechanisms mediated the reduced cardiovascular risk. Long-term, prospective, randomized-controlled trials of vitamin D supplementation are required to confirm effects in patients with early CKD.

Adequate vitamin D status may be important for regulation of several predictors of CVD risk that are relevant in patients with CKD, such as hypertension and left ventricular hypertrophy,79-81 insulin resistance/glucose intolerance,82 systemic inflammation,83,84 and oxidative stress.85 A reduction in these CVD risk indicators with cholecalciferol and/or active vitamin D analog treatment has been shown in ESRD patients.79,85-93 Only two trials, as indicated in this review, have investigated effects of vitamin D supplementation on such measures of CVD risk with suggestive, but not significant, results, and not as primary outcomes.35,52 No study has specifically investigated if vitamin D supplementation improves glucose homeostasis in early CKD. Given the scarcity of data available, conclusions cannot be made regarding specific effects of vitamin D supplementation on CVD risk in early stage CKD. Prospective clinical trials are required to identify the best measure of cardioprotective effects of vitamin D supplementation in early stage CKD, as well as the optimal vitamin D formulation.

Conclusion

The current systematic review suggests that achievement of optimal vitamin D status (25(OH)D ≥ 30 ng/ml) in patients with early CKD may require greater than 2,000 IU vitamin D per day. Furthermore, cholecalciferol may be more effective than ergocalciferol in achieving and maintain optimal 25(OH)D concentrations, thus highlighting the need for a revision of the current KDOQI guidelines. Current recommendations suggest intervening with vitamin D after the recognition of hyperparathyroidism, however it is possible that treatment of vitamin D deficiency earlier in the progression of CKD could prevent the rise in secondary hyperparathyroidism. Although various studies reviewed found an improvement in serum/plasma 25(OH)D and a reduction in PTH, the clinical utility of such improvements in early CKD is not yet confirmed. Few studies examine tangible, relevant clinical outcomes such as CVD risk factors, progression to ESRD, and mortality. Future vitamin D supplementation trials in early CKD should be designed with these clinical indicators as the primary outcomes. Furthermore, future studies should investigate effective treatment strategies for maintaining optimal vitamin D status, as well as improving clinical outcomes.

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