The Role of Vitamin D and Oxidative Stress in Chronic Kidney Disease

Keith C. Norris¹, Opeyemi Olabisi², M. Edwina Barnett³, Yuan-Xiang Meng⁴, David Martins⁵, Chamberlain Obialo⁴, Jae Eun Lee³, Susanne B. Nicholas¹

¹University of California Los Angeles, Los Angeles, CA. USA
²Harvard University, Boston, MA. USA
³Jackson State University, Jackson, MS. USA
⁴Morehouse School of Medicine, Atlanta, GA. USA
⁵Charles R. Drew University of Medicine and Science, Los Angeles, CA. USA

Number of Tables/Figures: none/none
Word counts: 1670 (excluding abstract, references and tables)
Running Head: Vitamin D, Oxidative Stress and Kidney Disease

Address for Correspondence:
Keith C. Norris, MD, PhD
911 Broxton Ave., Suite 103
Los Angeles, CA 90024
Phone 310-794-6973
Fax 310-794-0732
Email: kcnorris@mednet.ucla.edu
ABSTRACT

Chronic kidney disease (CKD) is a major non-communicable disease associated with high rates of premature morbidity and mortality. The prevalence of hypovitaminosis D (deficiency of 25(OH)D or 25D) is greater in racial/ethnic minorities and in patients with CKD than the general population. Low 25D is associated with bone and mineral disorders as well as immune, cardiometabolic and cardiovascular (CV) diseases. Thus, it has been suggested low 25D contributes to the poor outcomes in patients with CKD. The prevalence of hypovitaminosis D rises progressively with advancing severity of kidney disease with over 30% of patients with CKD stage 3 and 70% patients with CKD stage 5 estimated to have low levels of 25D. This report describes several of the abnormal physiologic and counter-regulatory actions related to low 25D in CKD such as those in oxidative stress and inflammatory systems, and some of the preclinical and clinical evidence or lack of thereof of normalizing serum 25D levels to improve outcomes in patients with CKD, and especially for the high risk subset of racial/ethnic minorities who suffer from higher rates of advanced CKD and hypovitaminosis D.

**Key Words:** Vitamin D; oxidative stress; kidney disease, disparities
Introduction

Chronic kidney disease (CKD) is a major non-communicable disease and is emerging as an important public health problem. Patients with CKD suffer from higher rates of premature morbidity and mortality due to a myriad of metabolic perturbations that arise as renal function declines. The prevalence of vitamin D deficiency (25(OH)D or 25D) is greater in patients with CKD than the general population. Over 30% of patients with CKD stage 3 [estimated glomerular filtration rate (eGFR) between 30 and 59 mL/min/1.73 m^2] have low levels of 25D, and the prevalence of 25D deficiency is as high as 60-70% in the later stages of CKD (stage 4 eGFR between 15 and 29 mL/min/1.73 m^2 and stage 5 eGFR <15 mL/min/1.73 m^2). Deficiency of 25D is associated with adverse clinical outcomes such as bone and mineral disorders (BMD) as well as immune, cardiometabolic and cardiovascular (CV) diseases. Thus, improving outcomes in patients with CKD requires normalization of many of the dysregulated physiologic and hormonal systems in CKD. While improving circulating levels of 25D are needed, optimizing improvement of immune and cardiometabolic health needs to take advantage of the unique interplay between 25D, oxidative stress and inflammation.

Vitamin D Deficiency/Insufficiency in Chronic Kidney Disease

25D is a pre-hormone that acts in concert with a variety of signaling pathways to influence a variety of cell actions throughout the body. Recommendations for adequate 25D levels vary, as the optimal concentrations for a given cell function and optimal clinical outcome varies. In addition, the level of 25D at which there is a clear adverse physiological manifestation due to the low value may vary by degree of impairment or...
activation of related counter-regulatory systems such as oxidative stress and inflammatory pathways.\textsuperscript{12} Reasons for 25D deficiency in the CKD population include, but are not limited to, phosphorus-restricted diets as well as decrease in dietary intake in general, and reduced endogenous synthesis of 25D due to limited outdoor activity and reduced sunlight exposure.\textsuperscript{2,13-16}

Hypovitaminosis D is commonly classified in terms of deficient and insufficient levels but these terms represent a spectrum of disease risk and not an explicit state of disease,\textsuperscript{9} leading to varying treatment recommendations. The Institute of Medicine (IOM) recommends serum 25D levels be maintained at 20-50 ng/ml.\textsuperscript{10} While the optimal serum 25D level for patients with CKD is not well defined, serum 25D concentrations below 12 ng/ml are associated with marked increased risk for BMD, cardiometabolic and CV diseases.\textsuperscript{10,11,17} Different organizations vary in their recommendations of the levels at which hypovitaminosis occur with 25D levels below 12-20 ng/ml considered deficient and levels below 20-30 ng/ml considered insufficient.\textsuperscript{10,11}

**Vitamin D, Oxidative Stress and Inflammation**

Several signaling pathways are charged with maintaining a healthy balance in the ongoing struggle between injurious oxidant and protective antioxidant events. These pathways include modification of proteins and DNA, and alteration in gene expression that may promote apoptosis, endothelial dysfunction and impairment of cellular immunity.\textsuperscript{18} A common metabolic pathway of stress-related cellular activation is an increase in reactive oxygen species (ROS) causing adverse cellular events termed oxidative stress, which is found in many chronic medical conditions such as
atherosclerosis, diabetes, immune-related disorders and CKD. Emerging evidence supports the role of 25D administration in attenuating oxidative stress via increased nuclear factor-erythroid-2-related factor 2 (Nrf2) and up-regulation of the expression of genes encoding antioxidant enzymes, as well as modulating levels of ROS through control of cellular antioxidants. In addition to oxidative stress, inflammation is a second major system implicated in pathogenesis of the premature CV diseases in patients with CKD. Nrf2 activates the antioxidant response element (ARE) and activation of the ARE downregulates redox-sensitive and inflammatory genes, including nuclear factor-kB (NF-kB). In patients with CKD, increasing oxidative stress leading to increased inflammation, and vice versa, are part of a deleterious cycle leading to over-production of each and adverse clinical sequelae. The ability of 25D repletion to attenuate this viscous cycle and reduce oxidative stress and inflammation through increasing Nrf2 and activating ARE represents a non-traditional regulatory role of the vitamin D pathway and a potential mechanism through which it may improve CKD-related cardiovascular disease, anemia, inflammation, and other clinical disorders.

Select Pre-Clinical and Clinical Studies

While its effects on attenuating oxidative stress in cell cultures and animal models have been robust, clinical results have been mixed, possibly due to factors such as differences in dosing, duration of treatment, and clinical setting (e.g., baseline 25D level, oxidative stress and inflammatory marker levels). Treatment with an analog of 1,25(OH)₂D, the active vitamin D hormone, reduced markers of systemic and intrarenal oxidative stress in mice with diabetic nephropathy. In addition, pretreatment with
1,25(OH)₂D administration reduced antioxidant activity and inflammation in a human cell culture model and animal models of oxidative stress. Despite promise in preclinical trials, its clinical effect in humans has been more variable. A Cochrane meta-analysis of seven randomized controlled trials of 25D administration to patients with polycystic ovary syndrome and normal kidney function found significant decreases in serum high-sensitivity (hs) C-reactive protein (CRP) and malondialdehyde (a marker of oxidative stress) with increased total antioxidant capacity, but no effect on nitric oxide or total glutathione levels. However, the administration of 1 mcg/day of paricalcitol [19-nor-1,25-(OH)₂D₂, an analog of the active form of vitamin D₂] or placebo for three months to 60 patients with diabetes and stage 3 or stage 4 CKD did not affect biomarkers of either oxidative stress or inflammation. By contrast one month of both 1 and 2 mcg/day of paricalcitol administration to 24 patients with CKD and a mean eGFR of 45 mL/min/1.73 m² significantly reduced hs-CRP and albuminuria compared to placebo. Coyne DW, et al. reported paricalcitol (1 mcg/day) and 1,25(OH)₂D (calcitriol; 0.25 mcg/day) were similarly effective in lowering intact parathyroid hormone (iPTH) and alkaline phosphatase in patients with secondary hyperparathyroidism in stages 3-4 CKD, with minimal elevations in serum calcium and phosphorus. However, two more recent outcome trials found no benefit from paricalcitol on clinical cardiovascular outcomes in patients with CKD. The PRIMO study (Paricalcitol Capsules Benefits in Renal Failure-Induced Cardiac Morbidity) found that forty-eight weeks of therapy with 2 mcg/day paricalcitol vs. placebo did not alter left ventricular mass index or improve certain measures of diastolic dysfunction in 227 patients with stage 3 and 4 CKD. Similarly, the OPERA trial (Oral Paricalcitol in Retarding Cardiac Hypertrophy, Reducing
Inflammation and Atherosclerosis in Stage 3 - 5 Chronic Kidney Disease) found a 52 week intervention with 1 mcg/day paricalcitol vs. placebo significantly improved secondary hyperparathyroidism but did not alter measures of left ventricular structure and function in 60 patients with stage 3-5 CKD. Thus, the promise of vitamin D to have a clinically significant impact on CV diseases in patients with CKD has not been shown in recent clinical trials. Whether or not there may be greater efficacy in select patients with higher levels of oxidative stress or inflammation and if these potential effects differ by dose and type of vitamin D used remains to be determined.

Possible Implications for Disparities

African Americans suffer from overall higher rates of hypovitaminosis D than other racial ethnic groups. Many cardiometabolic disorders that are also disproportionately high in African Americans are also associated with low levels of vitamin D. In addition, African Americans suffer from higher rates of the advanced stages of CKD and are 3 times more likely to develop end-stage kidney disease. For patients with CKD, African Americans typically have higher levels of intact parathyroid hormone. Interestingly, compared to White patients, African Americans on dialysis appear to have greater survival which has been linked, in part, to treatment with active vitamin D [1,25(OH)2D or analogs]. Whether this is an independent effect, related to a vitamin D - oxidative stress/inflammation interaction, or due to other causes is not known.

The role of hypovitaminosis D in African Americans is complex as emerging evidence has suggested certain vitamin D binding protein polymorphisms are associated with low measured levels of serum 25D but normal bioactive 25D, and these
polymorphisms are more prevalent in African Americans. Clinically this is highlighted by the findings of Gutierrez et al. who reported a strong correlation between serum 25D levels and bone mineral density among White and Hispanic patients, but no correlation for African Americans who had similar bone mineral density measures regardless of serum 25D level. Also, Robinson-Cohen reported increased CV events in Whites and Asians from over 6400 participants in the Multi-Ethnic Study of Atherosclerosis, but no relationship for Hispanics and African Americans, possibly related to ethnic differences in vitamin D binding protein polymorphisms or the vitamin D receptor. Berg and colleagues found both 24,25(OH)₂D levels and 25D levels were higher in White Americans compared to Black Americans, but the ratio of 24,25(OH)₂D to 25D was the same in both groups. Thus, the 24,25(OH)₂D to 25D ratio [vitamin D metabolite ratio (VMR)] may reflect free circulating 25D and could represent a more physiologically precise measure of bioactive 25D that should be independent of racial/ethnic differences in vitamin D binding protein levels and/or polymorphisms. Thus, the VMR may represent a new candidate biomarker for vitamin D status. Thus, modestly lower levels of serum 25D in African Americans may not require treatment and differing rates of these polymorphisms in different clinical studies may contribute to the conflicting outcomes following vitamin D administration.

**Summary/Conclusion**

Observational studies have linked low 25D levels in patients with CKD to progression to end-stage kidney disease, infections, fracture rates, hospitalizations, and all-cause mortality. A recent panel convened by the National Kidney Foundation recommended
patients with CKD and 25D levels less than 15 ng/mL should be treated, while those with serum 25D levels between 15 and 20 ng/mL may not require treatment unless there is evidence of counter-regulatory hormone activity. Whether evidence of increased oxidative stress and inflammation are other indicators of increased risk requiring treatment at modestly low serum levels is still to be determined. In addition, future studies are warranted to further assess the value of the 24,25(OH)₂D to 25D ratio and its correlation with clinical outcomes across racial/ethnic groups overall and in patients with CKD.
**Funding/Support:**

The authors are supported by the research grants from the NIH including: U54MD008149 (KN, MEB, YM, DM, JL), 3U54RR022762-03S4 (KN, YM, DM, GG), P20-MD000182 (KN, DM), UL1TR000124 (KN), and P30AG021684 (KN). The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health.

**Financial Disclosures:**

None of the authors declare any relevant conflicts of interest.

**Author Contributions:**

Keith C. Norris, Opeyemi Olabisi, Chamberlain Obialo, and Susanne B. Nicholas conceived and designed the concept and model. Keith C. Norris and M. Edwina Barnett drafted the paper. Opeyemi Olabisi, Yuan-Xiang Meng, David Martins, Jae Eun Lee and Susanne B. Nicholas made substantive edits and/or comments to the initial draft.

Acknowledgements: These data were presented in part as an oral presentation at the Research Centers in Minority Institutions (RCMI) Translational Science 2017 Conference.
References


