The Effects of Vitamin D Therapy on Left Ventricular Structure and Function – Are These the Underlying Explanations for Improved CKD Patient Survival?

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Abstract
Cardiovascular disease is a major cause of death among patients with chronic kidney disease and vitamin D deficiency is a common problem also among these patients. Abnormalities in left ventricular size and function are frequent, as they are encountered in 70–80\% of incident dialysis patients. These alterations develop early in the course of renal disease and their prevalence progresses in parallel with the decline in renal function. This process of left ventricular dilatation with compensatory hypertrophy continues after the institution of dialysis therapy, especially in the first year. The main factors responsible for the progression of left ventricular hypertrophy (LVH) are considered to be blood pressure and anemia, and in patients receiving hemodialysis, the arteriovenous fistula, volume overload and abnormalities in mineral metabolism. This additional potential set of factors related to LVH – mineral and bone metabolism – is intriguing and begs an immediate question: by what possible mechanism can these factors be linked to cardiac morphology? Recent observational studies have indeed indicated that vitamin D treatment was associated with a significant reduction of cardiovascular death among dialysis patients, and a reduction in LVH; in contrast, other studies suggested that excess vitamin D contributes to risk of hypercalcemia and vascular calcification, which is associated with reduced survival and morbidity. This review examines the evidence linking vitamin D with cardiac structure and function.

Introduction
Cardiovascular disease is a major cause of death among patients with chronic kidney disease (CKD) and vitamin D deficiency is a common problem also among these patients. Abnormalities in left ventricle (LV) size and function are frequent, as they are encountered in 70–80\% of incident dialysis patients [1]. These alterations develop early in the course of renal disease and their prevalence progresses in parallel with the decline in renal function [2, 3]. This process of LV dilatation with compensatory hypertrophy continues after the institution of dialysis therapy, especially in the first year [4]. The main factors responsible for the progression of left ventricular hypertrophy (LVH) are considered to be blood pressure and...
anemia [5], and in patients receiving hemodialysis (HD), the arteriovenous fistula, volume overload and abnormalities in mineral metabolism.

In a very recent cross-sectional analysis of prevalent Chinese patients, a higher prevalence of left ventricular hypertrophy (LVH) was found in HD than in peritoneal dialysis (PD) patients. As left ventricular mass index (LVMI) was associated with high blood pressure and volume overload, it was suggested that PD may preserve a more physiological hemodynamic status even [6]. At the same time, few reports from the literature demonstrate that single interventions (blood pressure control or anemia control) may decrease LV dysfunction. London et al. [7] showed that correction of anemia and strict BP control determined a decrease in LVM – associated with improvement in patient survival. Contradictory data were reported by Foley et al. [8], who found that complete anemia correction (hemoglobin = 13–14 g/dl) did not lead to regression of LVM in established LVH, but might prevent further LV dilatation. In all these studies, the enrolment period occurred before the emergence of the K/DOQI and European Best Practice Guidelines (EBPG) for HD practice. Covic et al. [9] found that strict implementation of EBPG guidelines was associated with an improvement in LV structural and functional parameters, in a significant percentage (61.2%) of dialysis patients. Anemia, mineral metabolism and blood pressure emerged as the most important determinants for changes in LVMI [9].

This additional potential set of factors related to LVH – mineral and bone metabolism – is intriguing and begs an immediate question: by what possible mechanism can these factors be linked to cardiac morphology?

Recent observational studies have indeed indicated that vitamin D treatment was associated with a significant reduction of cardiovascular death among dialysis patients, and a reduction in LVH [10]; in contrast, other studies suggested that excess vitamin D contributes to risk of hypercalcemia and vascular calcification, which is associated with reduced survival and morbidity [11].

This review summarizes the available literature regarding the relationship between vitamin D and cardiovascular disease, especially LVH.

**Vitamin D and Vitamin D Receptor in General**

The main source of vitamin D is de novo synthesis in the skin; under the action of ultraviolet light from sunlight, 7-dehydrocholesterol is converted to vitamin D3. Vitamin D is hydroxylated in the liver by 25-hydroxylase to 25-(OH) vitamin D. A second, subsequent hydroxylation in the proximal renal tubule by 1-hydroxylase is required to form the active metabolite 1,25(OH)2D3. The kidney is the major but not the only site for calcitriol production.

The majority of the molecular action of 1,25(OH)2D3 is mediated by the vitamin D receptor (VDR). The binding of 1,25(OH)2D3 to its nuclear VDR leads to VDR activation and heterodimerization with the retinoid X receptor (RXR) and other cofactors, to form the VDR-RXR complex, which binds to the vitamin D response element of target genes to regulate gene transcription. The VDR has been found in many tissues, not only in the classical target organs – intestine, kidney, and bone – but also in different cell types, including the immune system. This opens up the possibility of pleiotropic roles and actions for vitamin D [12]. Additionally, 1,25(OH)2D3 appears to bind to one or more cell surface receptors which, through second messenger pathways, mediate certain non-genomic effects [13]. These 2 observations, the widespread expression of the VDR, and the non-classical actions of vitamin D, make this one of the most complex and fast-moving topics in nephrology.

The predominant physiological function of the vitamin D is to maintain the calcium and the phosphate homeostasis, and this is accomplished by close coordination with parathyroid hormone (PTH). The wide distribution of VDR in almost every tissue provides a molecular basis to explain the beneficial effect of vitamin D on immune function [14], neuroprotection [15], and cardiac function [16].

Currently, vitamin D deficiency is recognized as endemic in old age and CKD [17]. Several recent studies have identified a high prevalence of vitamin D deficiency and insufficiency in otherwise healthy adults and children living in North America, Europe, and even sun-drenched countries. Thomas et al. [18] examined a cohort of 290 consecutively hospitalized patients; they discovered that 57% had a serum 25(OH)D below 40 nmol/l (the lower limit of normal) and 27% were classed as 'severe' vitamin D deficiency. In the MORE trial, Lips et al. [19] found that serum 25(OH)D was below 50 nmol/l in 28.4% of the postmenopausal women participating in this trial. In a multicenter study of 43 osteoporosis centers from all regions of Italy, including 700 women, values of 25(OH)D lower than 12 ng/ml were found in 76% of study subjects [20].

The major general cause of vitamin D deficiency is the lack of sufficient sun exposure. Vitamin D deficiency causes poor mineralization of the collagen matrix in
young children’s bones leading to growth retardation and bone deformities known as rickets and will precipitate and exacerbate osteopenia, osteoporosis, and fractures in adults.

Vitamin D deficiency is associated with colon and prostate cancer, type 1 diabetes, hypertension, coronary heart disease, and with alterations in the immune and inflammatory system.

The Relationship between the Vitamin D System and Cardiovascular Abnormalities

The association between vitamin D and cardiovascular disease events is widely debated and analyzed in the recent literature. In a very recent cross-sectional study, Pilz et al. [21] measured 25-hydroxyvitamin D [25(OH)D] levels in 3,299 Caucasian patients who were routinely referred for coronary angiography. They found that vitamin D deficiency is associated with prevalent myocardial dysfunction, heart failure, and sudden cardiac death. Poole et al. [22] compared serum vitamin D levels of 44 patients admitted to an acute stroke unit with those of 96 healthy, ambulant elderly subjects; they found that 77 percent of the stroke patients were deficient in vitamin D. In an elegant prospective study, Giovannucci et al. [23], using 18,000 men, showed that low levels of 25(OH)D are associated with higher risk of myocardial infarction in a graded manner, even after controlling for factors known to be associated with coronary artery disease.

On the other hand, Hsia et al. [24] recently reported that use of calcium and vitamin D supplements was not associated with a reduction in cardiovascular events in the Women’s Health Initiative. These apparently discrepant findings could be explained by several factors: the Women’s Health Initiative was a fracture-prevention trial and was not designed to evaluate cardiovascular risk, the investigators may have used an inadequate dose of vitamin D and the baseline 25(OH)D levels were not measured in the Women’s Health Initiative. A more recent meta-analysis of 18 randomized clinical trials, including the Women’s Health Initiative, did show that participants randomized to vitamin D supplementation experienced fewer deaths compared to those randomized to placebo [25].

Although the link between vitamin D deficiency and cardiovascular disease may be, in part, mediated through elevated PTH and calcium-phosphate metabolism, recent scientific evidence showed that vitamin D has 3 major potential protective mechanisms. First, experimental studies indicate that 1,25-OHD could determine the regulation of the renin-angiotensin axis by directly suppressing renin gene expression. Second, the presence in the cardiac muscle cells of a VDR, a calcitriol-dependent Ca\(^{2+}\) binding protein and a calcitriol-mediated rapid activation of voltage-dependent Ca\(^{2+}\) channels. Consequently, calcitriol administration can normalize the impaired contractility of the myocardium that is observed under experimental vitamin D deficiency. Third, vitamin D deficiency triggers secondary hyperparathyroidism, which then directly promotes cardiac hypertrophy (the direct PTH toxicity hypothesis).

Modulation of the Renin-Angiotensin System by Vitamin D

Vitamin D may play an important role in reduction of LVH through modulation of the renin-angiotensin system (RAS). The RAS plays a key role in the regulation of volume and blood pressure homeostasis, and over-activation of the RAS is a major pathogenic factor for hypertension, cardiac hypertrophy and atherosclerosis.

In observational and prospective cohort studies, low VDR activation has been associated with increased risk of hypertension, and is inversely correlated with plasma renin activity in patients with essential hypertension. Seasonal changes in blood pressure were described recently, higher blood pressure being associated with winter climate [26]. Vitamin D status, increasing after sun exposure, was associated inversely with blood pressure in a recent epidemiological study of a large normotensive population in the United States [27].

In HD patients, in a small, non-controlled study, intravenous calcitriol led a significant reduction in renin and angiotensin-II levels, and a significant regression of LVH [28]. These findings were later supported by data from animal and in vitro studies. Li et al. [29], using the vitamin D receptor knockout mouse model, showed that activated 1,25(OH)\(_2\)D\(_3\) inhibits renin secretion from the juxtaglomerular apparatus; they found that plasma angiotensin-II levels, as well as both renin mRNA and protein levels in the kidney, are elevated in vitamin D receptor knockout mice. More recently, Yuan et al. [30] found that 1,25(OH)\(_2\)D\(_3\) in vitro suppresses renin gene transcription by blocking the activity of the cyclic AMP-response element in the renin gene promoter.

Xiang et al. [31] completed the pathogenic link, showing that 1,25(OH)\(_2\)D\(_3\) functions as an endocrine suppres-
sor of renin biosynthesis and the subsequent genetic disrup-
tion of the VDR results in over-stimulation of the RAS, leading
to high blood pressure and cardiac hypertrophy. The cardiac
hypertrophy seen in vitamin D receptor knockout mice is a
consequence of activation not only of the systemic, but also of
the cardiac RAS [31]. These findings are confirmed by a recent
paper published by Zhou et al. [29] who demonstrated that
mutant mice deficient in 1,25(OH)2D3 biosynthesis develop
hypertension, cardiac hypertrophy, and impaired cardiac systolic
function due to the over-stimulation of the renal and car-
diac RAS.

Not all authors agree with this concept (see also be-
low). Simpson et al. [33] found that vitamin D deficiency
indeed induces myocardial hypertrophy and extracellular
matrix production and deposition in the myocardial
tissue of the rats; trichrome staining of heart tissue
showed marked increase in fibrotic lesions in vitamin D
receptor knockout (VDR-KO) mice. However, analysis of
plasma renin activity, angiotensin-II and aldosterone lev-
els showed elevated but not significantly different renin
activity in knockout mice versus normal mice and no sig-
nificant differences in angiotensin-II or aldosterone lev-
els [33, 34]. These finding suggest therefore that there
may be renin-independent associations between vitamin
D and cardiac structure.

**Other Effects of Vitamin D Therapy on Cardiac
Structure and Function**

Vitamin D also has several direct effects on cardiac
tissue growth and development including increased col-
lagen deposition, hypertrophy, and hyperplasia.

Zitterman et al. [35] described that an increased serum
level of atrial natriuretic peptide (ANP) gene expression
is one of the earliest and most reliable markers of cardiac
hypertrophy in neonatal rat cardiac myocytes. In patients
with LVH, virtually every biochemical or physical per-
turbation that results in ‘hypertrophy’ (increased cell
size, increased protein synthesis, and reorganization of
sarcomeric structure) also leads to activation of ANP
gene expression and a reduced circulating level of cal-
citriol, indicating an inverse relationship between vita-
m D level and cardiac hypertrophy [35].

These clinical findings are also supported by previous
animal and in vitro studies that provide alternative po-
tential explanations. In rats, Weishaar et al. [36] showed
that vitamin D deficiency induced myocardial hypertro-
phy (increased heart weight/body weight ratio) and ex-
tracellular matrix production via increased c-myc pro-
tein levels. Conversely, Gardner’s group found that cal-
citriol inhibited myocyte hypertrophy and the expression
of actin and ANP, markers of myocardial hypertrophy;
these later findings were confirmed in Wu et al. [37]
where it was found in vitro that 1,25(OH)2D3 inhibited
endothelin-induced myocyte hypertrophy and the ex-
pression of skeletal actin and ANP genes. More recently,
Nibbelink et al. [38] demonstrated that 1,25(OH)2D3 in-
creases expression of myotrophin (myotrophin is a myo-
cardial hypertrophy-inducing factor initially document-
ed in cardiomyopathic hearts).

Extracellular matrix remodeling, mediated by ma-
trix metalloproteinase, may be involved in progressive
LV remodeling, dilatation, and heart failure. Rahman
et al. [39] demonstrated that tissue inhibitors of metallo-
proteinase were significantly underexpressed in VDL-
KO mice; in the same animal model, the heart myofi-
brils showed highly significant cellular hypertrophy and
the fibrotic lesions were increased [39]. In humans,
in an Indo-Asian population, apparently healthy, but
with endemic vitamin D deficiency, plasma metallopro-
teinase levels were inversely related to vitamin D status
[40]. One year later, after treatment with vitamin D,
mean plasma metalloproteinase levels were decreased
significantly. Overproduction of transforming growth
factor-β (TGF-β) is almost universally observed in
models of experimental renal failure and it is recognized
to play a crucial role in renal fibrogenesis; conversely,
a blockade of its activity is associated with attenuation of
progressive renal failure. There is ample evidence that
TGF-β stimulates the synthesis of many extracellular
matrix components while reducing collagenase produc-
tion, as VDR activation is associated with reduced ex-
pression of vascular endothelial growth factor (VEGF)
and TGF-β. Angiotensin-II, the major effector molecule
of the RAS, has also been strongly implicated in renal
fibrosis. Numerous experimental studies, as well as clin-
cal studies, have shown that angiotensin-II blockade re-
duces renal injury and fibrosis in part by reducing TGF-
β. Cell culture studies have convincingly demonstrated
that angiotensin-II directly stimulates transcription and
bioactivation of TGF-β [42] as well as the synthesis
of VEGF in podocytes, which is mediated, in part,
through the activation of the p38 mitogen-activated
protein kinase pathway [42]. However, angiotensin-II
blockade does not normalize TGF-β levels, and disease
progression is slowed, but not halted.
Inflammation and Vitamin D

Inflammation is strongly associated with increased morbidity and mortality in CKD. Pro-inflammatory cytokines, tumor necrosis factor (TNF-α) and interleukin (IL)-1 or IL-6 (the central regulator of the inflammatory process) are elevated in CKD. Activated vitamin D has potent anti-inflammatory effects, by reducing production of the T-helper type 1 cytokines, IL-2, interferon-γ, and TNF-α and by suppression of inflammatory macrophage reactions [43]. Cytokine production may be modulated by calcitriol either as a result of its direct interaction with monocytes or indirectly through its effect on calcium and PTH metabolism. Panichi et al. [44] indicated a dose-dependent in vitro inhibition of the IL-1β and TNF-α production in uremic subjects incubated with physiologic doses of calcitriol. Therapeutic doses of calcitriol are also able to induce a marked decrease of cytokine production in vivo. Vitamin D deficiency is associated with elevated levels of C-reactive protein (CRP), and supplementation of this vitamin led to a 23% reduction in CRP levels at 1 year in a healthy cohort [45]. Alborzi et al. found the same relationship in their recent study [46]. All of their study subjects were vitamin D deficient (<30 ng/ml) and 60% also had an elevated baseline high-sensitivity CRP level, indicating subclinical inflammation. This study showed a reduction in CRP levels within 1 month after treatment with paricalcitol in patients with CKD. In contrast, in another recent paper, Ewers et al. [47] examined the connection between vitamin D status and low-grade systemic inflammation (serum concentrations of CRP) in an adult population of kidney-transplant patients. No impact of vitamin D status on low-grade systemic inflammation was found [47].

Many studies have demonstrated that vitamin D is a potent suppressor of cell proliferation and an inducer of the differentiation in numerous cells types [48]. The anti-proliferative action is mediated predominantly through a G1/S block of the cell cycle. However, there are comparatively fewer studies which have analyzed the effects of vitamin D on vascular smooth muscle cell (VSMC) proliferation, and the results of these are rather different. Cardus et al. [49] found that calcitriol induced a dose-dependent increase in VSMC proliferation (achieved by shortening the G1 phase). This effect is mediated by VEGF. This study was performed in vitro, and the concentration of 1,25(OH)2-vitamin D was significantly higher than the normal serum level, suggesting that 1,25(OH)2-vitamin D overdosing may cause stimulation of VSMC proliferation [37]. In contrast, O’Connell et al. [50] showed that 1,25(OH)2D3 deficiency produced hyperplasia and increased c-myc protein levels in the hearts of vitamin D3-deficient rats. The mechanism by which 1,25(OH)2D3 regulates myocyte proliferation involves blocking entry into the S phase of the cell cycle [49].

Vitamin D plays also an important role in cardiac function. Cardiac muscle cells possess a VDR and calcitriol-dependent Ca2+ binding protein. In a recent observational study in patients with cardiac dysfunction, both vitamin D deficiency and hyperparathyroidism were common findings. Moreover, a calcitriol-mediated rapid activation of voltage-dependent Ca2+ channels exists in cardiac muscle cells, and the uptake of calcium by cardiac muscle cells is in part regulated by vitamin D3. In experimental studies, vitamin D administration can normalize the impaired contractility of the myocardium. Baksi et al. [51] observed an increased contractility in response to increasing concentration of the extracellular calcium bath in atria in rats maintained on a vitamin D-deficient diet. The direct effects of vitamin D deficiency were suggested by Weishaar et al. [36]: rats fed a vitamin D-deficient diet have increased amounts of collagen (quantified by measuring hydroxyproline per gram of heart tissue and collagen deposition in the extracellular space of the myocardium). This effect could not be suppressed by normocalcemia, suggesting that indeed this is a direct action/consequence of vitamin D deficiency [51].

Vitamin D and Arterial Stiffness

Arterial stiffness in uremic patients is increased and arterial/coronary calcifications are frequent. Increased arterial stiffness is a consequence of several factors, including chronic fluid overload, the state of chronic microinflammation, sympathetic overactivity, activation of the renin-angiotensin-aldosterone axis, advanced glycation end products, lipid peroxidation, abnormalities of the nitric oxide system, and not the least, the calcification of the vessel wall. Arterial calcification is closely related to arterial stiffness: more calcified arteries are obviously losing their elastic properties [52]. Increased arterial calcification and stiffness may at least in part explain the very high morbidity and mortality in end-stage renal disease patients.

A passive process is implicated in vascular calcification, i.e. calcium-phosphate precipitation in the vessel walls, associated with an active biological process, ‘ossification’ of the vascular wall structure. Important con-
Contributors to these calcifications are hyperphosphatemia and an increased calcium-phosphate product (>55 mg²/dl²). At the same time, high levels of phosphate and/or calcium directly activate genes associated with osteoblastic functions in the smooth muscle cells (e.g., bone matrix protein 7, α2-HS glycoprotein, and matrix GLA protein) [53].

The first logical approach for reducing vascular calcification is the appropriate handling of the calcium-phosphate metabolism. Currently used calcium-containing phosphate binders (like calcium carbonate or acetate) may promote vascular calcification. Non-calcium-containing phosphate binders (like sevelamer hydrochloride or lanthanum carbonate) may reduce the calcium-load in dialysis patients [54]. In some in vitro studies and in animal models, calcitriol has induced vascular calcification, by oversuppression of PTH and induction of a low-turnover bone disease state, or by increased calcium-phosphorus (Ca × P) product [55]. Vitamin D analogues (paricalcitol, doxercalciferol, maxacalcitol) are capable of effective parathyroid suppression; they are structurally modified for fewer calcemic and phosphatemic effects. In a very recent study, Cardus et al. [56] demonstrated that paricalcitol has different effects on vascular calcification compared with calcitriol. Both compounds raised the serum calcium and Ca × P product compared to control, but only calcitriol caused an increase in the calcification of the abdominal aorta. Similar results were observed by Hirata et al. [57]. These results support the fact that different VDR activators exert differential effects on vascular calcification independent of serum calcium, phosphorus and Ca × P product.

Recently, Noohan et al. [58] demonstrated that paricalcitol and doxercalciferol display differential effects on aortic calcification, independent of serum Ca, P and Ca × P. These findings suggest a different mechanism of action between these 2 VDRA activators. The mechanism for the differential effects remains unclear [58].

**Vitamin D, PTH and LVH**

The function of vitamin D is accomplished in close coordination with the PTH. Vitamin D deficiency triggers hyperparathyroidism, which itself tends to promote cardiac hypertrophy.

In vitro, there are several possible explanations for increased LVM in primary/secondary hyperparathyroidism. PTH acts on adult cardiomyocytes by binding to the PTH/PTHrP receptor, inducing a rise in calcium intracellular levels [60], activating protein kinase C, which activates hypertrophic pathways inside the cell [61]. Expression of some cardiac proto-oncogenes may be enhanced, which may lead to altered expression of several genes involved in cardiac structure and function, and may stimulate the translation of contractile and non-contractile cardiac muscle proteins, all leading to LVH [62]. Additionally, Amann et al. [63] showed that elevated PTH levels cause irreversible interstitial fibrosis with collagen deposition. Regarding the relationship between PTH and systolic function, a direct effect of PTH on myocardial contractility has not been demonstrated in adult human myocytes, but the intracellular influx of calcium induced by PTH has been shown to increase contractility in animal cells.

In patients with primary hyperparathyroidism, there are conflicting reports on the relation between PTH and LVH, as well as on the cardiac effects of parathyroidectomy in these patients. Stefenelli et al. [64] reported that 81.6% of patients with primary hyperparathyroidism (PHPT) had hypertrophy of the intraventricular septum. Längle et al. [38] found that 50% of the patients with PHPT had LVH. Piovesan et al. [65] found a correlation between PTH values and LVM; this correlation suggests an action of the hormone in the pathogenesis of LV hypertrophy, confirmed also by the decrease of LVM after the reduction of PTH levels. Some studies have not detected any cardiac hypertrophy in PHPT patients [66], but in general these included fewer patients with a shorter duration of hyperparathyroidism.

The increased prevalence of LVH seems to be independent of blood pressure, since cardiac hypertrophy is also a common finding in normotensive PHPT patients. Furthermore, Stefenelli et al. [64] found that only 55% of the recruited PHPT patients with LVH had a history of hypertension. In the subgroup of patients who were receiving antihypertensive treatment, parathyroidectomy was not followed by a reduction of LVH; whilst in patients without evidence of hypertension there was a significant reduction of LVH [64].

In dialysis patients with secondary hyperparathyroidism, several investigators have shown a relationship between PTH and LVH. Stróżecki et al. [67] showed that the LVM was significantly greater in patients with remarkably high plasma PTH levels. Some studies have reported an improvement of cardiac function after parathyroid treatment in dialysis patients. For example, parathyroidectomy in 12 patients on maintenance HD resulted in a 22% reduction in LVMI [68]. However, Fujii et al. [73] consider that in dialysis patients many factors (such as
hypertension, anemia and volume overload, calcium and phosphate levels) may influence the cardiac abnormalities observed in these patients, so that it is very difficult to evaluate the relative importance and relationship between all these competing factors. None of the studies account for all these potentially relevant factors. The only study that performed load- and rate-independent measurement of cardiac function before and after parathyroid treatment under identical conditions was made by Fellner et al. [69] and included only 7 stable hemodialysis patients; it did not find a direct effect of PTH on cardiac function. Further interventional studies are clearly necessary to clarify the contribution of these factors to LVH in dialysis patients.

Calcitriol or Newer Vitamin D Analogues?

Vitamin D therapy has intrinsic dose-dependent calcemic and phosphatemic properties. This has heightened interest in structurally modified derivatives, with the hope that these unwanted effects will be reduced.

Although early reports indicated less increase in serum calcium and phosphorus at 12 months, and a significantly higher survival over the 36 months with vitamin D analogues (e.g. paricalcitol) compared to calcitriol [70], more recent trials are less positive. The Current Caring for Australians with Renal Impairment guidelines for CKD stage 5, published in 2006, indicated that there is insufficient evidence to support the superiority of newer vitamin D analogues over calcitriol [71]. Palmer et al. [72] using the 76 available randomized trials, found no evidence for the superiority of newer vitamin D analogues over established vitamin D compounds, for any major outcome.

Although promising, the survival benefit of the newer vitamin D analogues needs more evaluation, and must be subjected to the full rigor of an independently-monitored hard end-point outcome trial.

Conclusion

Effects of vitamin D compounds are necessarily pleiotropic as VDRs exist in almost every tissue and are incompletely understood. Vitamin D is an important influencing factor in the development and physiology of cardiac tissue. In dialysis patients, clinical studies suggest improvement in survival after treatment with vitamin D and a beneficial effect on LVH. These can be achieved by negatively regulating the RAS, modulating the inflammatory response to blood vessel injury, as well as inhibiting cardiomyocyte hypertrophy and proliferation. Nevertheless, definitive proof is still lacking, and in an era of mostly negative clinical trials, and ever-increasing pressure on health-care budgets, evidence derived from a solid, prospective, randomized controlled trial is mandated. However, the growth of basic scientific, observational, and interventional research in this field is explosive and is likely to lead to therapeutic advances in the next decade.

References


The minireview by Covic and colleagues from Guy’s hospital in London, UK, draws the reader’s attention to the recent literature suggestive of a survival advantage for vitamin D in patients with end-stage renal disease (ESRD) on renal replacement therapy as well as in those with cardiovascular disease. It focuses on the cardiovascular benefits of vitamin D including its effect on vascular and cardiac remodelling. Whilst there is a considerable and rapidly expanding literature on the subject of vitamin D and survival advantage, readers should be critical in their appraisal of the literature as such survival benefit may be confounded by indication and case mix. The healthier patients may be given vitamin D supplementation. Readers should also be aware that a recent meta-analysis failed to show an advantage and that the DOPPS review showed that, whilst vitamin D was associated with a survival benefit in models prone to bias, no difference in mortality was observed in instrumental variable models that tend to be more independent of unmeasured confounding [1]. Clearly, a randomised control study of vitamin D supplementation in ESRD patients would be valuable. Whether such a clinical trial is feasible in such a patient group is doubtful in view of the established advantage of vitamin D supplementation on the management of renal osteodystrophy in these patients.

Reference