CORRELATION BETWEEN PLASMA 25-HYDROXYVITAMIN D LEVEL WITH ANGIOGRAPHIC FINDINGS IN MALE STEMI PATIENTS

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ABSTRACT

BACKGROUND

Around 25% of deaths among Indians are attributable to CAD. Mortality from CAD in India has increased by 103% in males and 90% in females from 1985 to 2015.1 The prevalence of CAD in urban India is about double the rate of rural India and about 4-fold higher than in the USA.2 Vitamin D deficiency (VDD) has been associated with CVD risk factors such as hypertension3,4 and diabetes mellitus with markers of subclinical atherosclerosis such as intima-media thickness and coronary calcification as well as cardiovascular events such as myocardial infarction and stroke as well as CHF.5

Aims and Objectives of this study were to evaluate the 25-OH-D status in male patients with STEMI and to know the correlation between the 25-OH-D level and severity of angiography findings.

MATERIALS AND METHODS

Total 177 patients taken as per the inclusion and exclusion criteria mentioned below. We had taken 120 patients in our study group out of which 17 did not agree for angiographic interventions and 20 patients were detected to be having newly diagnosed case of type 2 DM, Hypertension, Alcoholic liver disease, smoking, etc. So they were excluded from the study. 83 patients were in study group. 57 patients were in control group. This study is a case control study. Approval from ethical committee was taken.

RESULTS

Mean age of presentation in the study group is 50.73 yrs. ± 11.73 yrs., range being 24 to 74 yrs as compared to the mean age group of 49.44 yrs. ± 13.2 yrs., range being 24-74 yrs. (p value > 0.05). Distribution patterns of angiographic findings in the study group are as described above i.e. 18 patients (21.7%) were having single vessel disease, 30 patients (36.1%) were having double vessel diseases and majority of the patients (35 patients) were having triple vessel diseases group i.e. 42.2%. Mean 25-OH-D level found in the TVD group was 9.66±3.60 ng/mL as compared to the SVD group where the level was 21.62±2.34 ng/mL i.e. significantly lower. One way ANOVA test was applied, f value found to be 103.59, p value<0.0001). Mean 25-OH-D level in the study group with complication was 10.72±2.77 ng/mL i.e. significantly lower as compared to the study group without complication where the mean 25-OH-D was 18.37±4.80 ng/mL (p value being <0.0001). It suggests that those patients with lower 25-OH-D were having more trend towards complications like heart failure, arrhythmia, periprocedural AMI, etc.

CONCLUSION

Our findings suggested low plasma 25-OH vitamin D in the STEMI patients when compared to the level in the age matched control group, and the lower the level of plasma 25-OH vitamin D the more severe were the incidences of severe angiography findings as described earlier specially in male patients and more were the incidence of complications. Hypovitaminosis D is another novel prognostic indicator for the severity of the CAD which can be treated aggressively to decrease the morbidity and mortality due to coronary artery disease.

KEYWORDS

Plasma 25-(OH) Vitamin D, STEMI


The vitamin D axis affects vascular SMC proliferation, inflammation, vascular calcification, the renin-angiotensin system (RAS), and blood pressure, all of which affect risk of CVD i.e. myocardial infarction (MI), a potentially life threatening manifestation. Vitamin D also impacts endothelial cell function, regulating endothelial cell-dependent vasodilation.5-8

Around 25% of deaths among Indians are attributable to CAD. Mortality from CAD in India has increased by 103% in males and 90% in females from 1985 to 2015. The prevalence of CAD in urban India is about double the rate of rural India and about 4-fold higher than in the USA.9

Vitamin D deficiency (VDD) has been associated with CVD risk factors such as hypertension10,11 and diabetes mellitus, with markers of subclinical atherosclerosis12 such as intima-
media thickness and coronary calcification as well as with cardiovascular events such as myocardial infarction and stroke as well as CHF. Low vitamin D levels have been associated with increased CVS morbidity & mortality in general population & in type 1 & 2 diabetes mellitus.13

Although VDD is frequently unrecognised clinically, laboratory measurement is easy to perform and treatment of VDD is relatively well tolerated and inexpensive.14

Because of its long half-life, 25-OH-D measurements are clinically useful for assessing vitamin D status in patients15 as it reflects the state of dietary supplement and endogenous production of vitamin D in the body. The level of 25-OH-D level also does not change like acute phase reactants in AMI, thus suggesting the constant level throughout the disease. Low level of vitamin D has been correlated with severity of CAD and the aim of this study is to correlate between the severity of angiographic findings with vitamin D level so that hypovitaminosis D can also be a marker of bad prognosis in coronary artery disease.

Despite these suggestive ecologic data and plausible mechanisms, data directly linking vitamin D levels to risk of MI are sparse. Because hypovitaminosis D is prevalent and easily correctable, establishing the relationship between vitamin D and risk of MI is important from treatment point of view.

**Epidemiology**

**Epidemiology in World**

Annual death rates in males and females caused by CVD are 37% and 41% respectively in the USA and 40% and 52%, respectively in Germany. In 1990, 63% of world mortality due to CVD were contributed by the developing countries16 which may rise to 76% of an estimated 25 million deaths in 2020 in economically developing countries.17

The French paradox indicates that mortality from CVD is relatively low in France despite a high intake of saturated fatty acids.18 The other paradoxes like The Italian paradox,19 The Northern Ireland paradox,20 The Albanian paradox21 indicate that some other cause might be contributing apart from the common aetiological risk factors for CVs disease. The Indian paradox indicates that CVD mortality rate in urban populations is higher compared with rural populations despite a very low fat intake.22 Hypertension is also less common or less severe at higher altitudes.23 Controlled clinical studies have demonstrated that regular exposure to UVA radiation but not to UVB radiation increases circulating 25(OH)D above a level of 100 nmol/L and also significantly reduces blood pressure by approximately 6 mmHg in hypertensive patients within an intervention period of 6 weeks.18

In Scotland, IHDM mortality rates showed a nadir in summer, approximately 30% lower compared with winter & winter peak and a summer nadir in male and female IHDM deaths.25 The prevalence of risk factors for CVD and also of disease rates is higher in urban than in rural communities in India26,27 & the same is true in African countries.28 In fact a South American study supported the same findings.6

**Epidemiology in India**

In view of being a tropical country, it was a disbelieve that VDD is uncommon in India.29 However, from the data available, VDD is very common in India in all the age groups and both sexes across the country.30-32 Vit D is a fat soluble vitamin and its synthesis in the body is dependent on multiple factors like latitude, atmospheric pollution, clothing, skin pigmentation and duration and time of exposure to sunlight. There is widespread prevalence of varying degrees (50-90%) of Vit D deficiency with low dietary calcium intake in Indian population according to various studies published earlier.30

CV Harinarayan et al have studied 25(OH)D and BMD in women of reproductive (WR) age group and postmenopausal women (PMW) in South India.33 They have reported VDD in 76% in WR, 70% in PMW, insufficiency in 16.5% in WR and 23% in PMW. VDD is considered to be present when serum 25(OH)D levels are <20 ng/mL; insufficiency between 20-30 ng/mL and sufficient when >30 ng/mL.34-36 In this study, there is VDD which coexists with low BMD.

**Vitamin D Physiology and Pathology**

There are 2 major forms of vitamin D, vitamin D2 (ergocalciferol) and vitamin D3 (cholecalciferol). Vitamin D2 is found in plants and can be consumed in fortified foods or as a supplement. Vitamin D3 is obtained from either dietary sources or through conversion of 7-dehydrocholesterol in the skin upon exposure to UVB radiation.1,4 Dietary sources of vitamin D are limited to fatty fish (wild or farm salmon, mackerel, tuna fish, sardines, and cod liver oil) and products fortified with vitamin D- depending on the country, these may include dairy products, cereals, margarine, flour, and orange juice.37,38

Both vitamin D3 obtained from exposure to the sun and from dietary sources are hydroxylated in the liver to 25-
hydroxyvitamin D (25(OH)D), which is then converted to the active form of vitamin D, 1,25-dihydroxyvitamin D (1,25(OH)2D) by 1-alpha-hydroxylase in the kidney.13,39,40 Though vitamin D is known as vitamin but basically it is a hormone with wide effects on body.41 Level of 25(OH)D is the marker that is more clinically relevant for VDD.37 Circulating hormone 1,25(OH)2D works at cellular level by affecting the genetic function and altering the protein synthesis which directly or indirectly regulates numerous genes at various organs.

**Definition of VDD**

VDD, a common problem in the USA and worldwide13,39 is defined as a level 75 nmol/L (30 ng/mL) approximately.42 Potential cutpoints for defining vitamin D status are listed in Table 13, including the IOM’s definitions of risk of deficiency, risk of inadequacy, adequacy, and risk of concern (including the fact that there may be a “U-shaped” relationship where very high levels may be detrimental.43 In the United States, the level of 25 (OH) D taken as low is 25 nmol/L (10 ng/mL).44 Indian definition of VDD has also been analysed as described below, 25(OH)D levels by RIA and PTH by IRMA were measured (made by BioSource Europe SA, Belgium) and its intensity categorised into three levels.45

- Mild deficiency- 15 ng/mL ≤ 25(OH)D < 20 ng/mL
- Moderate deficiency- 8 ng/mL ≤ 25(OH)D< 15 ng/mL
- Severe deficiency- 25(OH)D < 8 ng/mL

**Vitamin D and CV Risk Factors**

There is increasing evidence that vitamin D plays a role in determining risk for various cardiovascular outcomes, particularly metabolic syndrome (MetS) and type 2 diabetes mellitus (T2DM) and systemic hypertension.13,39,24

Epidemiological studies also suggest that the rate of hypertension, T2DM, and coronary heart disease (CHD) increase in proportion to increasing distance from the equator, suggesting a potential link to the vitamin D mechanism.

**Hypertension**

Vitamin D exerts a role in regulating the RAAS (a major contributor in CVD 1, 2, 10, 20, and VDD predisposes to upregulation of the RAAS and hypertrophy of both SMCs and left ventricle (LVH is a known risk factor or marker of CVD).41-26-27 In humans, 1, 25 (OH) 2D inhibits the RAAS and may lower blood pressure. VDD leads to hypertension as supported by Krause R et al study.28 Sugden J A et al study,6 NHANES III study. However, in a recent meta-analysis of 3 cohorts, lower 25(OH)D was associated with an 80% greater risk of incident HTN, whereas in a meta-analysis of 10 trials, supplementation with vitamin D only non-significantly reduced SBP by 2 mmHg and did not reduce DBP.24

Vitamin D also affects mechanisms related with MetS and T2DM pathophysiology, including impaired beta-cell function and insulin resistance, potentially by directly activating vitamin D receptors or by indirect effects by a calcium homeostasis regulation.44-47 Subsequently, this finding was confirmed by a meta-analysis of 5 observational studies in England,49 Pittas A. G et al study,50 Von Hurst P.R et al.51

Chronic VDD causes secondary hyperparathyroidism, which may mediate severe detrimental CV effects by various mechanisms. The threshold for elevation of PTH is a 25(OH)D level 30 ng/mL, with particular increases in Parathyroid levels at 25(OH)D levels 16 ng/mL. Elevated levels of PTH are associated with increases in arterial pressure and myocardial contractility, which can lead to apoptosis, fibrosis, and vascular SMC hypertrophy as well as LVH.10, 52 increase the risk of inflammation, as documented by elevated levels of C-reactive protein and interleukin-10.41 Administration of 1,25(OH)2D in the setting of VDD has been shown to down-regulate inflammatory biomarkers such as C-reactive protein.53 This may help to reduce the atherosclerosis.54-56

**Vitamin D, Cardiovacular Disease and Mortality**

In 1739 Framingham Offspring Study participants who were free of CV disease at baseline, the rate of major CV disease events was 53% to 80% higher among those with low vitamin D levels, with the increased risk magnified among those with hypertension.57 In an analysis of 13,331 adults from the NHANES III study followed up for a median of 8.7 years, mortality was inversely associated with vitamin D levels, with the lowest quartile of 25(OH)D ($17.8 ng/mL) having a 26% increased mortality compared with the highest quartile.2 In a prospective study of 3,258 consecutive patients scheduled for coronary angiography, low 25(OH)D and 1,25(OH)2D levels were independently associated with all-cause and CV mortality.4

**Vitamin D and Congestive Heart Failure**

In a recent study in 43 men and 17 women with left ventricular ejection fraction of 40% or less, longer 6-min. walk distance was correlated with higher 25(OH)D levels.58 The 6-min. walk distance is a frequently used test in heart failure patients to assess functional cardiac outcome. Data indicate that VDD is a causal factor of CHF and not just the result of disease-related alterations. Very low serum calcitriol levels (<37.5 pmol/L) have frequently been found in end-stage heart failure patients.59 In this study, patients in the highest calcitriol tertile had a hazard ratio for an event (Death or cardiac transplantation) of only 0.506 (95% CI 0.334–0.767) compared with patients in the lowest calcitriol tertile, after adjustment for potential confounders.

**Vitamin D and the Vasculature**

Classical risk factors in the pathogenesis of CHD are smoking, dyslipoproteinemia, hypertension, disturbed glucose metabolism, and proinflammatory processes.

Until recently, vascular calcification was considered to be a passive process that occurred as a nonspecific response to vascular damage without clinical significance. There is now accumulating evidence that vascular calcification is an active process.60 Almost all angiographically atherosclerotic lesions are calcified. Vascular calcification can cause thrombosis, arterial rupture, and myocardial infarction.

**Vitamin D and Cardiac Events**

During the last decade, deficiency of serum concentrations of vitamin D metabolites are found to be prevalent in the general population in western countries reason being an inadequate skin exposure to solar ultraviolet B radiation.

In a nonrandomised prospective study in 1739 Framingham Offspring Study participants, individuals with low 25(OH)D levels (<37.5 nmol/L) had a multivariable adjusted hazard ratio of 1.62 for incidents of cardiovascular disease such as myocardial infarction, coronary insufficiency, and heart failure compared with those with 25(OH)D levels of at least 37.5 nmol/L.41 In the Women's Health Initiative study; however, myocardial infarction, ischaemic attack, hospitalisation rate for heart failure, and cardiovascular death could not be prevented by supplementation with 1000 mg calcium and 10 mg vitamin D daily compared with the placebo group.62 But several limitations such as the low daily vitamin D dose and the lack of measurements of serum 25(OH)D levels or of calcitriol or of both of the vitamin D arm of this study makes data interpretation difficult. In a nested case-control study among male participants of the Health Professionals Follow-up study, men with low 25(OH)D levels (37.5 nmol/L) had a relative risk of 2.09 [95% confidence interval (CI) 1.24–3.54] of myocardial infarction compared with those considered to be sufficient (75 nmol/L), after adjustment for various lifestyle and other risk factors. Very recently, data on all-cause and cardiovascular mortality in association with vitamin D status have been published from a prospective cohort study of 3258 consecutive male and female patients scheduled for coronary angiography.53 During a median follow-up period of 7.7 years, 737 patients (22.6%) died, including 463 deaths from cardiovascular causes. Multivariate-adjusted hazard ratios for patients in the lower two 25(OH)D quartiles (median, 19.0 and 33.3 nmol/L) were higher for all-cause mortality (hazard ratio 2.08; 95% CI 1.60–2.70; and hazard ratio 1.53, 95% CI 1.17–2.01, respectively) and for cardiovascular mortality (Hazard ratio 2.22; 95% CI 1.57–3.13; and hazard ratio 1.82, 95% CI 1.29–2.58, respectively) compared with patients in the highest 25(OH)D quartile (median, 71.0 nmol/L). Similar results were obtained for patients in the lowest calcitriol quartile.

In line with this suggestion, a meta-analysis of controlled clinical trials came to the conclusion that vitamin D
supplementation reduced total mortality in middle aged to elderly adults by 7% during a trial size-adjusted mean of 5.7 years.64

Sunlight provides the most potent source of vitamin D, with approximately 3,000 IU vitamin D3 for 5 to 10 min. of mid-day, mid-year exposure of arms and legs for a light-skin Caucasian 4. Oral supplements of 50,000 IU of either D2 or D3 every 2 weeks.39,40 Among standard dietary sources of vitamin D, oily fish have the highest content of D3, ranging from 100 to 1,000 IU per 3.5 oz serving, whereas orange juice or milk fortified with vitamin D contains about 100 IU per serving.39,40,65 Each 100 IU of vitamin D ingested daily produces about 1 ng/mL increase in 25(OH)D levels.66,67

Generally, oral supplementation of either D2 or D3 increases levels of vitamin D reasonably well.68

Recent Reviews of Literature for Vit D & CAD

Serum 25(OH)D does not change after AMI and is likely to be a reliable marker of vitamin D status in patients with cardiovascular disease i.e. it does not behave like an acute phase reactant. Serum Vitamin D levels are independently associated with severity of CAD. VDD is associated with significant coronary stenoses in asymptomatic African American chronic cocaine users. Association of VDD with heart failure and sudden cardiac death noted in a large cross-sectional study of patients referred for coronary angiography.69 Independent association of low serum 25-hydroxyvitamin D and 1,25-dihydroxyvitamin D levels with all-cause and cardiovascular mortality. Low vitamin D levels predict stroke in patients referred to coronary angiography.70 The latest data supports the correlation of atherosclerosis and osteoporosis indicating the parallel progression of two tissue destruction processes with increased fatal and non-fatal coronary events, as well as higher fracture risk. Vitamin D inadequacy associated with low bone mineral density increases fall and fracture risk, leads to secondary hyperparathyroidism, calcifies coronary arteries and significantly increases cardiovascular disease.71

Two case-control studies and a small prospective study60 found that individuals with low 25(OH)D levels were at higher risk for ischaemic heart disease. The strongest test of the hypothesis that vitamin D lowers MI risk as seen by UK study of 2606 men and women where the participants were randomized to receive 830 IU of vitamin D daily (administered as 100000 IU of oral vitamin D3 every 4 months) or placebo for 5 years. The in-study 25(OH)D levels were 29.7 ng/mL in the vitamin D group and 21.4 ng/mL in the placebo group. There was a non-significant decrease in CVD incidence (RR,0.90; 95% CI, 0.77-1.06) and CVD mortality (RR, 0.84; 95% CI, 0.65-1.10) in the intervention group. Based on the present study, a difference of 3.3 ng/mL in 25(OH)D concentration would be associated with an RR of 0.92, which is compatible with the previous results. A recent meta-analysis of total mortality as a secondary end point of RCTs with varying levels of vitamin D vs placebo controls found a statistically significant 8% reduction in risk of total mortality in individuals who had received vitamin D. Although the authors could not evaluate cause-specific mortality, the relatively immediate effect of a large enough magnitude to affect total mortality would suggest a benefit on CVD risk. The largest RCT of vitamin D (and calcium) supplementation and CVD risk was from the Women’s Health Initiative, in which 282 postmenopausal women received either calcium (1000 mg/d) and vitamin D3 (cholecalciferol) (400 IU/d) or placebo. No reduction was observed in MI- or CHD-related deaths (hazard ratio, 1.04; 95% CI 0.92-1.18). These results seem to be in contrast to the present findings, suggesting 2 possible explanations. First, despite our efforts to exclude confounding, it is possible that uncontrolled or residual confounding explained these results. Alternatively, the range of vitamin D studied was much wider in the HPPS, which allowed us to detect an association. The difference between the medians of the top and bottom categories, for which we observed an RR of approximately 2, was 23.5 ng/mL (35.5-12.0 ng/mL), and the calculated reduction in MI risk per increment of 1 ng/mL of 25(OH)D was 2%. In the Women’s Health Initiative study, the effect of the treatment on 25(OH)D levels was not reported, but based on the dose and compliance, Lappe et al estimated it to be only 2 ng/mL.57 Based on the present data, such an increment would be expected to have only a 4% reduction in risk. To increase 25(OH)D levels from 12 to 35.5 ng/mL would require approximately 3000 IU of vitamin D daily.73 Although such intakes may seem high by current standards, increasing evidence demonstrates no toxic effects at intakes below 10000 IU/d.4 Because current sources of vitamin D provide much less (e.g. a glass of milk has approximately 100 IU), those who achieve high levels such as 35 ng/mL naturally do so largely through sun exposure.

MATERIALS AND METHODS

Aims and objectives of this study were to evaluate the 25-OH-D status in the male patients with STEMI and to know the correlation between the 25-OH-D level and severity of angiography findings. Total 177 patients taken as per the inclusion and exclusion criteria mentioned below. We had taken 120 patients in our study group out of which 17 did not agree for angiographic interventions and 20 patients were detected to be having newly diagnosed case of type 2 DM, Hypertension, alcoholic liver diseases, smoking, etc. So they were excluded from the study. 87 patients were in study group. 57 patients were in control group. This study is a case control study. Approval from ethical committee was taken.

All STEMI (As diagnosed by ACC/AHA criteria) (table 12)75 male patients in Cardiology Emergency Department who are considered for invasive strategy (either primary or pharmacoinvasive strategy as defined by PCI guidelines ACC/AHA 2011) were included. Patients excluded are the following as described by USA/ NSTEMACS/CSA patients, female patients with ACS, Poorly controlled DM, Dyslipidaemia, Smokers, Patients of old age more than 75 yrs., Morbidly obese patients, HIV positive individuals, chronic kidney disease, alcoholic liver diseases, patients who already received heparin or calcium or vitamin D, patients on glucocorticoid therapy or parathyroid hormone, chronic inflammatory diseases which affect the calcium level.

The cases were evaluated (ECG, echocardiography, angiography) and managed in the Department of Cardiology, S.C.B. Medical College and Hospital, Cuttack and the biochemical tests & investigations like CBC, RFT, FBS, PPBS, CK, CKMB, TROP T, LDH, vitamin D (25-OH-D), lipid Profile, were conducted in the Department of Biochemistry, S.C.B. Medical College and Hospital, Cuttack, The samples collected from volunteers without any clinical symptom or history of
any coronary heart disease or any chronic debilitating illness will be taken as control. Blood samples were collected in three 10-mL liquid EDTA blood tubes, placed on ice packs, stored in polystyrene foam containers, and returned to the blood storage department of biochemistry, and processing facility at the biochemistry department of S.C.B. Medical College, via overnight courier. Blood samples which were collected in the day time were analysed immediately. More than 95% of the samples arrived within 24 hours of collection. Plasma 25(OH)D levels were determined by means of ELISA method using vitamin D binding antibody.

Angiography findings will be categorised in the following groups as mentioned below (>50% of MLA or >70% MLA in all coronary vessel except LEFT main coronary artery where >50% of MLA).
1. SVD- single vessel disease.
2. DVD- double vessel disease (LAD+LCX, or LCX+ RCA or LAD +RCA proximal lesion excluded).
3. TVD group- triple vessel disease, Left main disease, Left main equivalent, Diffuse disease, Type Clesion.

The patients are divided into 2 groups according to complications like heart failure, arrhythmia, periprocedural AMI, contrast induced nephropathy, stent thrombosis. (Present or Absent).

Severity of angiographic findings were analysed by the SCAI classification and angiography classification. Data so collected was entered into the MS Excel Sheet. Statistical analysis was conducted using SPSS 16 software. A p value less than 0.05 was taken to be significant. Qualitative data was analysed by using Chi- square test. Quantitative data was analysed using student’s t test and one way ANOVA, as applicable.

Common causes of VDD are described in Table 11.

**Observations**

<table>
<thead>
<tr>
<th>Sl. No.</th>
<th>CAG Findings</th>
<th>Study Group N=83</th>
<th>Control Group N=57</th>
<th>P Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>SVD</td>
<td>18 (21.7%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>DVD</td>
<td>30 (36.1%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>TVD Group</td>
<td>35 (42.2%)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Table 4. Coronary Angiographic Profile of the Study Group (n = 83)**

<table>
<thead>
<tr>
<th>Age in Years</th>
<th>Mean 25-OH-D in Study Group (N=83)</th>
<th>Mean 25-OH-D in Control Group (N=57)</th>
<th>P Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>25-30 yrs</td>
<td>14.17±4.03</td>
<td>33.79±3.5</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>31-40</td>
<td>15.04±5.1</td>
<td>30.3±2.66</td>
<td></td>
</tr>
<tr>
<td>41-50</td>
<td>14.67±5.41</td>
<td>33.62±4.99</td>
<td></td>
</tr>
<tr>
<td>51-60</td>
<td>13.74±8.88</td>
<td>31.00±2.88</td>
<td></td>
</tr>
<tr>
<td>61-70</td>
<td>13.93±6.84</td>
<td>31.57±4.67</td>
<td></td>
</tr>
<tr>
<td>71-75</td>
<td>10.84±6.97</td>
<td>38.55±3.28</td>
<td></td>
</tr>
</tbody>
</table>

**Table 5. 25-OH-D Level Correlation with CAG Findings in Study Group (n = 83)**

<table>
<thead>
<tr>
<th>Sl. No.</th>
<th>Complication Present (N=47)</th>
<th>Complication Absent (N=36)</th>
<th>P Value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>10.72±2.77</td>
<td>18.37±4.80</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td></td>
<td>n 47 (56.6%)</td>
<td>36 (43.4%)</td>
<td></td>
</tr>
</tbody>
</table>

**Table 7. Distribution of 25-OH-D level in the Study Group with or Without Complication (n = 83)**

<table>
<thead>
<tr>
<th>CAG Findings</th>
<th>N</th>
<th>Age in years</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>DVD</td>
<td>14 (29.8%)</td>
<td>48.35±11.23</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>TVD group</td>
<td>33 (70.2%)</td>
<td>51.72±12.54</td>
<td></td>
</tr>
</tbody>
</table>

**Table 8. Correlation of CAG Findings in the Study Groups with Complications (n = 47)**

<table>
<thead>
<tr>
<th>Sl. No.</th>
<th>25-OH-D Quartiles</th>
<th>N</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>20.1-30.0</td>
<td>15 (18.1%)</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>10.1-20</td>
<td>47 (56.6%)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>3</td>
<td>0.1-10</td>
<td>21 (25.3%)</td>
<td></td>
</tr>
</tbody>
</table>

**Table 9. Three Quartile Distribution of 25-OH-D & CAG Findings in the Study Group (n = 83)**
### Table 10. Three Quartile Distribution of Vit-D & Severity of CAG Findings in the Study Group (n = 83)

<table>
<thead>
<tr>
<th>Sl. No.</th>
<th>25-OH-D Quartiles</th>
<th>SVD (N=18)</th>
<th>DVD (n=30)</th>
<th>TVD (n=35)</th>
<th>N</th>
<th>P Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>1st</td>
<td>20.1-30.0</td>
<td>13</td>
<td>1</td>
<td>1</td>
<td>15</td>
<td>&lt; 0.0001</td>
</tr>
<tr>
<td>2nd</td>
<td>10.1-20</td>
<td>5</td>
<td>29</td>
<td>13</td>
<td>47</td>
<td></td>
</tr>
<tr>
<td>3rd</td>
<td>0.1-10</td>
<td>0</td>
<td>0</td>
<td>21</td>
<td>21</td>
<td></td>
</tr>
</tbody>
</table>

### Table 11. Major Risk Factors for Vitamin D Deficiency

- Ageing
- Increased distance from the equator.
- Winter seasons.
- Darkly pigmented skin.
- Institutionalised/housebound.
- Sunscreens and cover-up clothing.
- Air pollution.
- Smoking.
- Obesity
- Physical inactivity.
- Genetic factors.
- Malabsorption.
- Renal disease.
- Liver disease.
- Certain medications.
- Glucocorticoids.
- Anti-rejection medications.
- Human immunodeficiency virus medications.
- Certain antiepileptic drugs.

### Either of the following Criteria Satisfies the Diagnosis for Acute, Evolving, or Recent AMI.

1. Typical rise and/or fall of biochemical markers of myocardial necrosis with at least one of the following.
   a) Ischaemic symptoms.
   b) Development of pathologic Q waves in the ECG.
   c) Electrocardiographic changes indicative of ischaemia (ST-segment elevation or depression).
   d) Imaging evidence of new loss of viable myocardium or new regional wall motion abnormality.
2. Pathologic findings of an acute myocardial infarction.

### Table 12. Revised Definition of Myocardial Infarction Criteria for Acute, Evolving, or Recent AMI
DISCUSSION

As depicted from the table 1, majority of patients in the study group are in the age group of 41-50 yrs. (28.9% vs. 24.6%) in the control group & 51-60 yrs. (32.5% vs. 21.1%) in the control group. (p value being more than 0.05, i.e. statistically insignificant). When we divided the age group in 2 groups according to age of 50 years, majority of the patients in study group were above the age of 50 yrs. i.e. 57.8% as compared to 49.12% in the control group. (p value being more than 0.05, i.e. statistically insignificant) as described in table 2.

As depicted in table 3, mean age of presentation in the study group is 50.73 yrs. ± 1.173 yrs., range being 24 to 74 yrs. as compared to the mean age group of 49.44 yrs. ± 13.2 yrs., range being 24-74 yrs. (p value being more than 0.05).

Distribution pattern of angiographic findings in the study group as per table 4 are as described i.e. 21.7% were having single vessel disease, 36.1% were having double vessel disease and majority of the patients were having triple vessel disease group i.e. 42.2%.

Table 5 shows that mean 25-OH-D level found in the control group was 9.66±3.60 ng/mL as compared to the SVD group where the level was 21.62±2.34 ng/mL, i.e. significantly lower. One way ANOVA test was applied, f value found to be 103.59, p value being less than 0.0001 i.e. highly significant.

Table 6 shows that average 25-OH-D level taken from the control group, prevalent in normal population was 32.47±4.35 ng/mL vs. 14.039± 5.39 ng/mL in the study group, (p value being < 0.0001), i.e. highly significant statistically. Age wise distribution of the 25-OH-D level was also analysed so as to know the range of the normal levels of vitamin D status in each group so as to prevent any age related difference in the Vit D level as compared to the study group. When we analysed the mean level of the Vit D in the age matched study group and control group, it was observed that the mean value of 25-OH-D was significantly lower in the study group as compared to the control group, the findings are more specific in older patients, p value being <0.0001, i.e. highly significant. The findings were more robust in the elderly patients i.e. 13.93±6.84 ng/mL in 61-70 yrs. group vs. 31.57±4.67 ng/mL in the control group with p value being <0.0001, i.e. highly significant, similarly in the 71-75 age group, the mean level was 10.8±6.97 ng/mL in the study group vs. mean level in the control group was 38.55±3.28 ng/mL (p < 0.0001).

Table 7 shows that mean 25-OH-D level in the study group with complication was 10.72±2.77 ng/mL i.e. significantly lower as compared to the study group without complication where the mean vitamin D level was 18.37±4.80 ng/mL (p value being <0.0001), i.e. highly significant statistically. It suggests that those patients with lower vitamin D level were having more trend towards complications like heart failure, arrhythmia, periprocedural AMI, etc.

As per table 8, also the complications like heart failure, arrhythmia or periprocedural AMI were noted in the TVD group as compared to the SVD group where not a single case of complication was noted & fewer complications noted in DVD group which has been depicted. On analysis, it was found that the lower the vitamin D level the more severe is the complication and the more is the severity of angiographic findings as depicted in the table. (p value being <0.0001).

Table 9 shows that after dividing the group of Vit D level in the 3 groups i.e. 1st quartile -insufficient Vit D level i.e. (20-30 ng/mL), 15 (18.1%) patients were in this group; 2nd quartile -mild to moderate insufficiency (10-20 ng/mL)- 47(56.6%) patients were in this group; and in 3rd quartile i.e. severe deficiency (<10 ng/mL), 21 (25.3%) were in this group.

Table 10 shows that after subgroup analysis of the study group in 1st quartile (n=15) it was observed that 13 out of 15 patients i.e. 86.6% were in SVD group as compared to 1 (6.7%) patient in DVD group & 1 (6.7%) patient in TVD group. Majority of patients were having relatively milder disease. In SVD group where 18 patients were there, we found the mean 25-OH-D level to be 21.62±2.34 ng/mL. Amongst these SVD group, we found that 13 out of 18 patients (72.23%) were in this quartile as compared to 5 patients (27.77%) who were in DVD group, p value being highly significant <0.0001. After subgroup analysis of the study group in 2nd quartile (n=47), it was observed that 5 out of 47 patients i.e. 10.64% were in SVD group vs 29 out of 47 patients i.e. 61.7% patients were in DVD group & 13 out of 47 patients i.e. 27.7% patients were in TVD group. Majority of patients were having DVD followed by TVD group. In DVD group, where 31 patients were there, we found the mean 25-OH-D level to be 14.60±2.10. Amongst DVD group, we found that 29 out of 30 patients (96.7%) were in this quartile as compared to 1 patient (3.3%) in 1st quartile. P value being highly significant <0.0001.

After subgroup analysis of the study group in 3rd quartile (n=21), it was observed that all patients i.e. 100% patients were in TVD group. In TVD group where 35 patients were there, we found the mean 25-OH-D level to be 9.66±3.60 ng/mL. Amongst the TVD group, we found that 21 out of 35 (60.7%) patients were in SVD group, 13 out of 35 patients (37.15%) were in 2nd quartile as compared to 1 patient (2.15%) in 1st quartile. P value being highly significant <0.0001.

Findings were in favour of lower level of 25-OH D level leading to increased severity of the angiographic findings.

A Danish study examined 25 hydroxyvitamin D (25[OH]D) levels measured in 128 patients admitted to the hospital with ischaemic heart disease (75 with angina pectoris and 53 with acute MI) and 409 control subjects and found that 25 (OH)D levels were significantly lower in those with angina (23.5 ng/mL [to convert to nanomoles per litre, multiply by 2.496]) or MI (24.0 ng/mL) than in controls (28.8 ng/mL). In a New Zealand case-control study, 3 of 179 patients with MI, cases had a lower mean 25(OH)D level (P=0.2), which was more pronounced in the winter-spring (P =0.03) than in the summer-autumn (P=2.1). The relative risk (RR) of MI decreased across increasing quartiles of 25(OH)D [10 ng/mL; RR, 1 [reference]; 10-13 ng/mL: RR, 0.56 [95% confidence interval (CI), 0.32-1.03]; 13.1-16.8 ng/mL: RR, 0.33 [95% CI, 0.17-0.64]; and 16.8 ng/mL: RR, 0.30 [95% CI, 0.150.61]). Multivariate analyses of major CVD risk factors did not appreciably alter the results. A small, nested, case-control study of MI based in the Tromso Heart Study (northern Norway) with only 30 cases and 60 matched controls found a slightly non-significant lower 25(OH)D level in cases (23.6 ng/mL) compared with controls (25.4 ng/mL).
CONCLUSION

The main finding of the study which was in favour of the above studies’ findings was that:

1. The level of vitamin D was low in the patients of coronary artery disease (established cases) when compared to the vitamin D level in the age matched control group.

2. The lower the level of vitamin D, the more severe was the incidence of severe angiography findings as described earlier and more is the incidence of complications.

Hypovitaminosis D is another novel prognostic indicator for the severity of the CAD.

Abbreviations

AMI - Acute myocardial infarction
25(OH)D - 25-hydroxyvitamin D
1,25(OH)2D - 1,25 Dihydroxyvitamin D
CHD - Coronary Heart Disease
CAD - Coronary Artery Disease
CKD - Chronic Kidney Disease
CRP - C-reactive Protein
CV - Cardiovascular
IOM - Institute of Medicine
MetS - Metabolic Syndrome
PTH - Parathyroid Hormone
RAAS - Renin Angiotensin Aldosterone System
T2DM - Type 2 Diabetes Mellitus
UVB - Ultraviolet B
VDD - Vitamin D Deficiency

REFERENCES


