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**SYSTEMATIC REVIEWS AND META-ANALYSES**

**Effect of vitamin D supplementation on endothelial function — An updated systematic review with meta-analysis and meta-regression**

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**KEYWORDS**
Vitamin D; Atherosclerosis; Endothelial function; Endothelial dysfunction

**Abstract**

**Background and aims:** Atherogenesis and endothelial dysfunction contribute to cardiovascular risk and vitamin D has been implemented in endothelial repair. This systematic review, meta-analysis and meta-regression aims to establish the effect of vitamin D supplementation on endothelial function.

**Methods and Results:** To conduct the systematic review we searched the Cochrane Library of Controlled Trials, PubMed, ProQuest and EMBASE for randomized controlled trials that investigated the effects of vitamin D supplementation on flow-mediated dilation (FMD%), pulse wave velocity (PWV), and central augmentation index (AIx). Meta-analysis was based on a random effects model and inverse-variance methods to calculate either mean difference (MD) or standardized mean difference (SMD) as effects sizes. This was followed by meta-regression investigating the effect of baseline vitamin D concentrations, vitamin D dosing and study duration. Risk of bias was assessed using the Jadad scale and funnel plots.

We identified 1056 studies of which 26 studies met inclusion criteria for quantitative analysis. Forty-two percent of the 2808 participants had either deficient or insufficient levels of vitamin D. FMD% (MD 1.17% (95% CI 0.20, 2.54), p = 0.095), PWV (SMD −0.09 m/s (95% CI −0.24, 0.07), p = 0.275) and AIx (SMD 0.05% (95% CI −0.1, 0.19), p = 0.52) showed no improvement with vitamin D supplementation. Sub-analysis and meta-regression revealed a tendency for AIx and FMD% to increase as weekly vitamin doses increased; no other significant relationships were identified.

**Conclusions:** Vitamin D supplementation showed no improvement in endothelial function. More evidence is required before recommendations for management of endothelial dysfunction can be made.

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and the critical step in CVD commencement and progression with endothelial dysfunction (ED) and associated loss of endothelial integrity presenting as a precursor stage of atherosclerosis [3].

The vascular endothelium can be classified as a complex paracrine and endocrine organ that produces vasodilatory and vasoconstricting factors, most importantly, nitric oxide (NO) [4]. Endothelial dysfunction encompasses dysregulation of mechanistic processes facilitating vasodilatory function of the endothelium, and the actual functionality of the endothelium itself [5]. This broadens the assessments of ED beyond the simple scope of the physical functionality of the endothelial wall and expands the scope to include platelet aggregation and adhesion, vascular inflammation and thrombosis management [5]. Endothelial dysfunction is characterised by a decrease in bioavailable NO leading to a reduction in endothelial vasodilation in response to vasodilatory stimuli [6]. Endothelial dysfunction is the initial, and reversible step in atherogenesis development, and is used as a primary tool for clinical identification of the presence and extent of atherosclerosis by measuring endothelial vasoreactivity [5,7]. Measurement of ED is used as a predictor of CVD mortality risk [8]. Endothelial dysfunction is measured through both invasive and non-invasive methods, the most common being central augmentation index percentage (AIx), pulse wave velocity (PWV), and artery flow-mediated dilation (FMD%), the latter considered as the gold standard of ED assessment [5].

Vitamin D is a secosteroid hormone most commonly associated with calcium regulation regarding bone remodelling [9]. Lower serum concentrations of vitamin D have been associated with increased risk of CVD pathogenesis, including increased risk of hypertension, decreased myocyte contractility and increased arterial calcification [10]. Vitamin D receptors are expressed in numerous tissues, and due to their expression in cardiovascular tissues, focus has shifted to vitamin D effects on CVD risk factors, including endothelial function [9]. Speculation exists for the mechanisms for which vitamin D affects endothelial function through NO regulation. One study showed that calcitriol, the hormonally active metabolite of vitamin D, increases monocyte differentiation into myeloid angiogenic cells and augments their angiogenic capacity for endothelial repair, as well as increasing endothelial NO synthase expression, increasing endothelial function [11]. Several RCTs have reported equivocal results regarding the effects of vitamin D supplementation on arterial stiffness, an important measure of CVD risk, continuing the trend of conjecture regarding RCT results of vitamin D and CVD outcomes [9,10,12–16]. The positive effects of vitamin D on reducing CVD risk factors remain questionable due to RCT results being obtained from populations without hypovitaminosis D. This, and the short duration of the RCTs fails to confirm observational study results which show a positive correlation between vitamin D concentrations and CVD risk factor reduction [13]. Given the expression of vitamin D receptors throughout cardiac tissues, it is entirely conceivable that vitamin D supplementation may improve cardiovascular outcomes, with proposed mechanisms including decreasing the effects of chronic inflammation and decreased vascular smooth muscle cell proliferation [17,18].

This systematic review with meta-analysis and meta-regression aims to summarise the currently available evidence from RCTs investigating supplemental vitamin D effects on endothelial function in both healthy and clinical populations. The intent is to draw conclusions as to whether supplemental vitamin D can be recommended for prevention of establishment or progression of ED.

**Added value to previous meta-analyses on the same topic**

Previous systematic reviews and meta-analyses [9,12,19–23] have reported on the effects of vitamin D supplementation and markers of endothelial function; however, not all included FMD%, PWV and AIx as a primary outcome measure and only one provided analyses for all three [23]. In addition, a number of new trials have recently been published and are included in the present systematic review incorporating meta-analysis and meta-regression.

**Methods**

**Search strategy**

A systematic search was conducted to identify potential studies using the following electronic databases: the Cochrane Library of Controlled Trials, PubMed, ProQuest and EMBASE until 31st May 2019. Search criteria included numerous terms, both free text and MeSH regarding atherosclerosis, vitamin D2 and vitamin D3, endothelial function, endothelial dysfunction and terms titling methods for assessing endothelial function. Systematic reviews, meta-analyses, and study bibliographies were reviewed for additional studies. Three reviewers (NP, MP, GD) conducted the search and full article eligibility review.

**Study selection**

The following criteria were applied for study identification and selection (1) Randomized, double-blinded, placebo controlled trials; (2) studies conducted in all age groups with no additional exclusion criteria applied for sex, health status or CVD risk factors; (3) vitamin D2 or D3 as primary intervention (dose, administration method or route of vitamin D2 or D3 administration, or inclusion of other secondary interventions were not a basis for study exclusion). Studies using the vitamin D receptor activator Paricalcitol were excluded.

**Data extraction and outcome measures**

Data extraction was completed by three reviewers (NP, MP, GD). Primary outcomes measured were limited to endothelial function test measurements conducted via FMD%, PWV and AIx. Studies reporting FMD values were included
if the measurement was reported as relative FMD percentage (FMD%) or as absolute FMD (measured in cm, mm or μm) in the brachial artery. Where FMD was reported as both absolute FMD and FMD%, only data from the FMD% was extracted for analysis. Studies selected presented numerous PWV and AIx measurements and were included; studies that did not state PWV method were excluded. Units of measure reported for PWV outcomes were converted to m/s.

**Data synthesis**

Individual meta-analyses were completed for continuous data utilizing the change in mean and standard deviation (SD) values for the study populations. Where the change in mean and SD was not reported, the pre-intervention mean was subtracted from the post-intervention mean, and the change SD was calculated utilizing study group participant numbers in conjunction with group p-values or 95% CI. Where exact p-values or 95% CI were not available, the SD of the mean difference was imputed using the formula, \(SD = \text{square root}\left[\frac{(SD_{\text{pre-treatment}})^2 + (SD_{\text{post-treatment}})^2 - 2rSD_{\text{pre-treatment}} \times SD_{\text{post-treatment}}}{2}\right]\) [24], assuming a correlation coefficient (r) = 0.5, which is considered a conservative estimate. Where the standard error of the mean (SEM) was given instead of the SD, this value was converted to SD [24]. Data from studies that reported median and interquartile range (IQR 1st and 3rd quarter) was converted to mean and SD using the method described in Wan et al. [25]. Studies that reported data as median and 50% IQR had data converted to mean and estimated SD assuming the value given was approximately 1.35 standard deviations. Mean differences (MD) in conjunction with random effects inverse variance were applied to pooled data for FMD% analysis. Standardized Mean Differences (SMD) were utilized in conjunction with a random effects inverse variance to allow for comparison between differing effects sizes and methods of measurement of the outcome between selected studies, as was the case when conducting analyses for both PWV and AIx pooled data groups. Using current guidelines, SMD values of 0.2, 0.5 and 0.8 signify small, medium and large effects sizes, respectively [26]. Placebo group numbers were evenly divided amongst intervention study groups in selected studies that presented data for more than one intervention group. Forest plots were generated to provide visual representation of vitamin D effects on endothelial function. A 95% CI was selected to report changes in measured outcomes. Sensitivity analysis was conducted using the leave-one-out method for overall assessment of the intervention effect. Where studies did not report data suitable for pooling, a descriptive analysis was included. Meta-regression analysis was performed (NK) to investigate the heterogeneity of results using baseline vitamin D concentrations, weekly dose of vitamin D supplement and duration of study as covariates with a 5% level of significance and 95% confidence intervals [27]. All analyses were carried out and all figures were produced in Comprehensive Meta-Analysis (CMA) V3 (Biostat Inc., NJ, USA).

**Heterogeneity and publication bias**

Heterogeneity between included studies was calculated using CMA V3; the I² test was utilized for consistency appraisal between studies. Values < 25% show low risk of heterogeneity, whereas >75% shows high risk of heterogeneity. Values that fall between 25 and 75% show moderate risk of heterogeneity [24]. These values together with assessment of funnel plots and Egger’s regression test were used to evaluate overall heterogeneity and assess risk of publication bias [28].

**Study quality**

Study quality was assessed using the JADAD scale [29]. The maximum score possible is 5; with a score of ≤3 indicating high risk of bias, while a score of >3 indicated a low risk of bias. Study quality was performed by two assessors (NP, GD).

**Results**

Using allocated search criteria, the search of the four selected databases identified 1056 manuscripts. After removal of duplicates, 59 full-text articles were assessed for eligibility. After screening, a total of 32 full-text papers remained for inclusion in the qualitative analysis, of which 26 were eligible for inclusion in the meta-analysis as per PRISMA flow diagram (Fig. 1).

**Study characteristics**

Thirty-two studies were included in the qualitative analysis with 26 suitable for inclusion in the meta-analysis. Reasons for exclusion of the six studies from the meta-analysis are detailed in Supplementary Table 1. The total number of participants analyzed for the specified outcomes from the 26 studies [4,10,14–16,30–50] included in the meta-analysis was 2808. There was a diversity of participants in the studies, with the inclusion of both healthy and clinical populations. Of the 26 studies, six [4,14,30,42,45,48] provided a total of 327 participants diagnosed with Type 2 Diabetes Mellitus (T2DM) with 55 of these participants identified as being diagnosed with T2DM and non-alcoholic fatty liver disease (NAFLD) [30]. Four studies [15,38,40,50] provided a total of 273 participants diagnosed with various stages of Chronic Kidney Disease (CKD). Remaining studies of clinical populations included participants with increased risk of diabetes, polycystic ovarian syndrome (PCOS), human immunodeficiency virus (HIV), peripheral arterial disease (PAD), chronic fatigue, and CVD. Age group amongst participants ranged from 22.4 to 72.9 (average age 54.6 ± 14.8). Sex distribution was primarily male. Of the 26 included studies, 8 studies included participants with deficient (<12.5 ng/ml), 11 studies with inadequate (<20 ng/ml) and 7 studies with adequate (≥20 ng/ml) baseline vitamin D levels for optimal overall health as per National Institutes of Health recommendations – 58% of participants were
considered to have adequate vitamin D levels. Characteristics of all included studies are detailed in Table 1.

**Intervention details**

Intervention duration ranged from four to 57 weeks with most studies using a 16-week intervention period. Vitamin D supplementation dosing ranged from 892 IU/d to 7142 IU/d, with the higher doses due to singular large dosing either at the start of the study period [44,45,47], or large doses at interval periods longer than daily dosing [10,15,16,32,33,36–38,43,49,50]. Accounting for single dose effect over the intervention period most studies used between 2000 IU/d to 5000 IU/d and all studies supplemented vitamin D orally. All studies except for two [10,45,51] used vitamin D3, with one study using both vitamin D2 and D3 for the intervention [32]. In a number of studies, participants were using multiple pharmaceutical treatments in addition to either vitamin D or placebo. One study [33] included dosing of metformin with either 4000 IU of vitamin D plus 1500 mg of metformin, or placebo plus 1500 mg of metformin. One study included use of HIV medications [37] with either vitamin D or placebo. One study included 200 mg/d of calcium [42] with either vitamin D or placebo. Dosing intervals ranged from daily dosing to once every three months. To test endothelial function several methods were used across all studies, with the most common being FMD% [10,15,30,34,35,37,45,47–49], PWV [15,16,31–34,36,38,40–43,50] and AIx [4,14,31,33,34,36,38,40,42–44,46,47,49].
Table 1  Characteristics of studies included in the meta-analysis and -regression.

<table>
<thead>
<tr>
<th>First Author</th>
<th>Study Design</th>
<th>Health Status</th>
<th>n = analysed for specified outcomes</th>
<th>n = analysed Vitamin D group (VD)</th>
<th>Dose</th>
<th>n = analysed placebo group (C)</th>
<th>Duration In weeks</th>
<th>Baseline Vitamin D Concentration ng/ml ± SD or IQR</th>
<th>Outcome (Vitamin D versus Placebo)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Barchetta 2016</td>
<td>R, DB, placebo</td>
<td>T2DM with NAFLD</td>
<td>55</td>
<td>26</td>
<td>2000 IU/d (D₁)</td>
<td>29</td>
<td>24</td>
<td>16.06 ± 9.56 (C)</td>
<td>No effect on FMD</td>
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<td>Italy</td>
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<td>19.26 ± 9.48 (VD)</td>
<td>No effect on FMD</td>
</tr>
<tr>
<td>Borgi 2017</td>
<td>R, DB, placebo</td>
<td>Overweight/obese</td>
<td>84</td>
<td>43</td>
<td>50,000 IU/w (D₂)</td>
<td>41</td>
<td>8</td>
<td>15.4 (IQR 11.4–17.5)</td>
<td>No effect on FMD</td>
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<tr>
<td>United States</td>
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<tr>
<td>Breslasky 2013</td>
<td>R, DB, placebo</td>
<td>T2DM</td>
<td>32</td>
<td>19</td>
<td>1000 IU/d (D₃)</td>
<td>13</td>
<td>52</td>
<td>11.7 ± 6.5 (C)</td>
<td>Improved AIx</td>
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<td>Israel</td>
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<td>11.8 ± 10.9 (VD)</td>
<td>No effect on PWW</td>
</tr>
<tr>
<td>Bressendorff 2016</td>
<td>R, DB, placebo</td>
<td>Healthy</td>
<td>40</td>
<td>22</td>
<td>3000 IU/d (D₃)</td>
<td>18</td>
<td>16</td>
<td>12.8 ± 4 (C)</td>
<td>No effect on AIx</td>
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<tr>
<td>Denmark</td>
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<td>17 ± 11 (C)</td>
<td>No effect on AIx</td>
</tr>
<tr>
<td>Dalan 2016</td>
<td>PG, R, DB, placebo</td>
<td>T2DM</td>
<td>61</td>
<td>31</td>
<td>2000–4000 IU/d (D₃)</td>
<td>30</td>
<td>16</td>
<td>18 ± 7 (VD)</td>
<td>Improved PWW</td>
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<tr>
<td>Singapore</td>
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<tr>
<td>Forouhi 2016</td>
<td>R, DB, placebo</td>
<td>Increased risk T2DM</td>
<td>160</td>
<td>55</td>
<td>100,000 IU/m (D₃)</td>
<td>52</td>
<td>16</td>
<td>18.32 ± 10.52 (C)</td>
<td>Improved PWV</td>
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<tr>
<td>United Kingdom</td>
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<td>19.44 ± 9.88 (VD2)</td>
<td>No effect on PWW</td>
</tr>
<tr>
<td>Garg 2015</td>
<td>P, R, DB, placebo</td>
<td>PCOS</td>
<td>32</td>
<td>15</td>
<td>120,000 IU/m (D₃)</td>
<td>17</td>
<td>24</td>
<td>6.8 ± 2.46 (C)</td>
<td>No effect on PWW</td>
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<td>India</td>
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<td>7.7 ± 6.05 (VD)</td>
<td>No effect on AIx</td>
</tr>
<tr>
<td>Gepner 2012</td>
<td>P, R, DB, placebo</td>
<td>Healthy, post-menopausal</td>
<td>109 (FMD)</td>
<td>55 (FMD)</td>
<td>2500 IU/d (D₃)</td>
<td>54 (FMD)</td>
<td>16</td>
<td>32.3 ± 10.5 (C)</td>
<td>No effect on PWW</td>
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<tr>
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<td></td>
<td>30.3 ± 10.7 (VD)</td>
<td>No effect on PWW</td>
</tr>
<tr>
<td>Harris 2011</td>
<td>P, R, DB, placebo</td>
<td>Overweight</td>
<td>45</td>
<td>22</td>
<td>60,000 IU/m (D₃)</td>
<td>23</td>
<td>16</td>
<td>15.28 ± 1.2 (C)</td>
<td>Improved FMD</td>
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<td></td>
<td></td>
<td>13.72 ± 0.88 (VD)</td>
<td>Improved FMD</td>
</tr>
<tr>
<td>Kumar 2017</td>
<td>R, DB, placebo</td>
<td>CKD</td>
<td>117</td>
<td>58</td>
<td>300,000 IU at baseline and 8 w (D₃)</td>
<td>59</td>
<td>16</td>
<td>13.21 ± 4.78 (C)</td>
<td>Improved FMD</td>
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<td>India</td>
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<td>13.4 ± 4.42 (VD)</td>
<td>Improved PWW</td>
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<tr>
<td>Larsen 2012</td>
<td>R, DB, placebo</td>
<td>Hypertension</td>
<td>111 (PWV)</td>
<td>54</td>
<td>3000 IU/d (D₃)</td>
<td>57</td>
<td>20</td>
<td>23 ± 12 (C)</td>
<td>No effect on PWW</td>
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<td>Denmark</td>
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<td>23 ± 9 (VD)</td>
<td>No effect on AIx</td>
</tr>
<tr>
<td>Levin 2017</td>
<td>R, DB, placebo</td>
<td>CKD</td>
<td>102 (Alx)</td>
<td>52</td>
<td>15,000 IU/w (D₃)</td>
<td>30</td>
<td>21</td>
<td>29.4 ± 12.7 (C)</td>
<td>Improved PWW</td>
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<tr>
<td>Canada</td>
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<td>25.9 ± 10 (VD)</td>
<td>No effect on FMD</td>
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<tr>
<td>Longnecker 2012</td>
<td>P, R, DB, placebo</td>
<td>HIV</td>
<td>45</td>
<td>30</td>
<td>4000 IU/d (D₃)</td>
<td>15</td>
<td>12</td>
<td>6.2 (IQR 3.7–9.8)</td>
<td>No effect on FMD</td>
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<td>9 (IQR 7.1–13.1)</td>
<td>No effect on AIx</td>
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<td>Marksmann 2012</td>
<td>PG, R, DB, placebo</td>
<td>CKD</td>
<td>28 (PWV)</td>
<td>12 (PWV)</td>
<td>40,000 IU/w (D₃)</td>
<td>16 (PWV)</td>
<td>8</td>
<td>13.2 (IQR 9.4–17.1)</td>
<td>No effect on PWW</td>
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<td>Denmark</td>
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<td>9.5 (IQR 6.9–16.6)</td>
<td>No effect on AIx</td>
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<td>Martins 2014</td>
<td>R, DB, placebo</td>
<td>Overweight/obese</td>
<td>130</td>
<td>65</td>
<td>100,000 IU/m (D₃)</td>
<td>65</td>
<td>12</td>
<td>6.6 ± 2 (C)</td>
<td>No effect on AIx</td>
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<td>United States</td>
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<td>6.8 ± 2.08 (VD)</td>
<td>No effect on AIx</td>
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<tr>
<td>Mose 2014</td>
<td>R, DB, placebo</td>
<td>CKD stage 5 – chronic dialysis</td>
<td>41 (PWV)</td>
<td>18</td>
<td>3000 IU/d (D₃)</td>
<td>19</td>
<td>24</td>
<td>11.2 (IQR 8–27.6)</td>
<td>No effect on PWW</td>
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<td>Denmark</td>
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<td>11.2 (IQR 8–19.2)</td>
<td>Improved AIx</td>
</tr>
<tr>
<td>Pilz 2015</td>
<td>R, DB, placebo</td>
<td>Hypertension</td>
<td>153</td>
<td>81</td>
<td>2800 IU/d (D₃)</td>
<td>72</td>
<td>8</td>
<td>20.4 ± 5.7 (C)</td>
<td>No effect on PWW</td>
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<td>Austria</td>
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<td>22.0 ± 5.5 (VD)</td>
<td>No effect on AIx</td>
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<tr>
<td>Raed 2017</td>
<td>R, DB, placebo</td>
<td>Overweight</td>
<td>70</td>
<td>17</td>
<td>18,000 IU/m (D₃)</td>
<td>17</td>
<td>16</td>
<td>15.9 ± 3.9 (C)</td>
<td>Improved PWW</td>
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<td>United States</td>
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<td></td>
<td>14 ± 3.9 (VD-18,000)</td>
<td>No effect on PWW</td>
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<tr>
<td>Ryu 2014</td>
<td>P, R, DB, placebo</td>
<td>T2DM</td>
<td>45</td>
<td>24</td>
<td>2000 IU/d (D₃)</td>
<td>21</td>
<td>24</td>
<td>10.7 ± 2.6 (C)</td>
<td>No effect on PWW</td>
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<td>Korea</td>
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<td></td>
<td></td>
<td>12.3 ± 3.0 (VD)</td>
<td>No effect on AIx</td>
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<table>
<thead>
<tr>
<th>First Author</th>
<th>Study Design</th>
<th>Health Status</th>
<th>n = analysed for specified outcomes</th>
<th>n = analysed Vitamin D group (VD)</th>
<th>Dose</th>
<th>n = analysed placebo group (C)</th>
<th>Duration In weeks</th>
<th>Baseline Vitamin D Concentration ng/ml ± SD or IQR</th>
<th>Outcome (Vitamin D versus Placebo)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sluyter 2017</td>
<td>R, DB, placebo</td>
<td>Aged 50–84 yrs (&gt;60% HT)</td>
<td>517</td>
<td>256</td>
<td>200,000 (D₃) initially, then 100,000 IU/m</td>
<td>261</td>
<td>57</td>
<td>25.24 ± 9.84 (C) 24.84 ± 9.88 (VD)</td>
<td>No effect on PWV No effect on Alx</td>
</tr>
<tr>
<td>Stricker 2012</td>
<td>Pilot, R, DB, placebo</td>
<td>PAD</td>
<td>62</td>
<td>31</td>
<td>100,000 IU - single dose (D₃)</td>
<td>31</td>
<td>4</td>
<td>17 ± 5.5 (C) 16.3 ± 6.7 (VD)</td>
<td>No effect on Alx</td>
</tr>
<tr>
<td>Sugden 2008</td>
<td>P, PG, DB, R placebo</td>
<td>T2DM</td>
<td>34</td>
<td>17</td>
<td>100,000 IU - single dose (D₃)</td>
<td>17</td>
<td>8</td>
<td>14.56 ± 3.4 (C) 16.08 ± 4.12 (VD)</td>
<td>Improved FMD</td>
</tr>
<tr>
<td>Tomson 2017</td>
<td>PG, R, DB, placebo</td>
<td>Aged ≥65 yrs</td>
<td>305</td>
<td>102 102</td>
<td>2000 IU/d (D₃) 4000 IU/d (D₃)</td>
<td>101</td>
<td>52</td>
<td>20 – all groups</td>
<td>No effect on PWV No effect on Alx</td>
</tr>
<tr>
<td>Witham 2013b</td>
<td>PG, R, DB, placebo</td>
<td>Healthy</td>
<td>50</td>
<td>25</td>
<td>100,000 IU - single dose (D₃)</td>
<td>25</td>
<td>8</td>
<td>10.8 ± 6 (C) 10.8 ± 5.2 (VD)</td>
<td>No effect on FMD No effect on PWV No effect on Alx</td>
</tr>
<tr>
<td>Witham 2015</td>
<td>PG, R, DB, placebo</td>
<td>Chronic Fatigue</td>
<td>50</td>
<td>25</td>
<td>100,000 IU/2 months (D₃)</td>
<td>25</td>
<td>24</td>
<td>19.2 ± 8 (C) 17.6 ± 6 (VD)</td>
<td>No effect on FMD No effect on PWV</td>
</tr>
<tr>
<td>Yiu 2013</td>
<td>P, R, DB, placebo</td>
<td>T2DM</td>
<td>100</td>
<td>50</td>
<td>5000 IU/d (D₃)</td>
<td>50</td>
<td>12</td>
<td>21.9 ± 4.1 (C) 21.1 ± 4.4 (VD)</td>
<td>No effect on FMD No effect on PWV</td>
</tr>
</tbody>
</table>

Alx: augmentation index, C: control/placebo group; CKD: chronic kidney disease, d: day; DB: double blind, FMD: flow-mediated dilation, HIV: human immunodeficiency virus, HT: hypertension; IQR: interquartile range; m: month; NAFLD: non-alcoholic fatty liver disease, PAD: peripheral artery disease, PCOS: polycystic ovarian syndrome, PG: parallel group, P: prospective, PWV: pulse wave velocity, R: randomised, SD: standard deviation; T2DM: type 2 diabetes mellitus; VD: vitamin D supplement group; w: week.
Outcome measures

Flow-mediated dilation

Ten studies assessed FMD% as either a primary or secondary outcome and included a total of 689 participants [10,15,30,34,35,37,45,47–49]. Vitamin D supplementation did not improve FMD% with MD 1.17% (95% CI −0.20, 2.54), p = 0.095 (Fig. 2). Three additional studies [52–54] reported on FMD%, however, data did not allow for pooling. All three of these studies reported no change in FMD% with vitamin D supplementation. Sensitivity analysis for FMD% indicated that the studies of Kumar (2017) [15] and Harris (2011) [35] impacted the size and significance of the result. Upon removal of these studies, MD was reduced to 0.67% (95% CI −0.55, 1.88), p = 0.28 and 0.92% (95% CI −0.66, 2.49), p = 0.25, respectively (Supplementary Table 2).

Pulse wave velocity

Seventeen studies assessed PWV as either a primary or secondary outcome [15,16,31–34,36,38,40–43,50] and included a total of 1981 participants. Vitamin D supplementation did not improve PWV with SMD 0.09 m/s (95% CI −0.24, 0.07), p = 0.275 (Fig. 3). An additional four studies [51,52,55,56] reported on PWV (data did not allow for pooling) and showed no change in PWV with Vitamin D supplementation. Leave-one-out sensitivity analysis for PWV did not indicate that any study overly impacted the results (Supplementary Table 3).

Augmentation index percentage

Fifteen studies assessed AIx as either a primary or secondary outcome [4,14,31,33,34,36,38,40,42–44,46,47,49] and included a total of 1569 participants. Vitamin D supplementation did not improve AIx with SMD 0.05% (95% CI −0.1, 0.19), p = 0.52 (Fig. 4). An additional study [55] reported on AIx (data did not allow for pooling) and reported no change in AIx with vitamin D supplementation. Leave-one-out sensitivity analysis for AIx did not indicate that any study overly impacted the results (Supplementary Table 4).

Sub-analyses of vitamin D supplementation effect on FMD%, PWV and AIx

Sub-analyses were conducted for FMD%, PWV and AIx on the effect of baseline vitamin D levels (deficient, inadequate or adequate), vitamin D supplementation (daily, weekly, monthly or 2-monthly), the amount of vitamin D administered (<20,000 IU/week or >20,000 IU/week), and in populations with T2DM and CKD (Supplementary Figures 1 to 12).

Baseline vitamin D levels seemed to improve FMD% in those participants with inadequate vitamin D levels with MD 1.8% (95% CI −0.06, 3.68), p = 0.057. In participants with deficient or inadequate baseline vitamin D levels, PWV showed some improvement in vitamin D deficient participants with SMD 0.29 m/s (95% CI −0.01, 0.6), p = 0.06 and was significantly decreased in participants with inadequate vitamin D levels, favouring the placebo group, with −0.38 m/s (95% CI −0.09, −2.59), p = 0.01, respectively. AIx was not affected by baseline vitamin D levels (Supplementary Figures 1–3).

The timing of vitamin D supplementation did not significantly improve FMD%, PWV or AIx when supplemented daily, weekly, monthly or 2-monthly (Supplementary Figures 4–6).

Vitamin D dosage of <20,000 IU/week vs. >20,000 IU/week did not significantly affect FMD% or PWV; however, AIx showed a significant improvement when doses of >20,000 IU/week were administered with SMD 0.13% (95% CI 0.015, 0.25), p = 0.03 (Supplementary Figures 7–9).

Type 2 Diabetes – Pooled data for FMD%, PWV and Alx on Vitamin D supplementation in populations with T2DM did not demonstrate a significant improvement in any measure (Supplementary Figures 10 and 12).
additional study in a T2DM population [54], not included in the pooled analysis also failed to find any significant improvement.

Chronic Kidney Disease — Only one study [15] included in the review reported FMD% in a population with CKD; vitamin D supplementation improved FMD% with MD 5.49% (95% CI 4.35, 6.63), p = 0.00. Subgroup analyses for PWV (4 RCTs) and Alx (2 RCTs) failed to indicate any significant improvement (Supplementary Figures 10 and 12). Two additional studies [51,56] in CKD patients, not included in the pooled analysis, also failed to demonstrate any significant improvements.
Vitamin D supplementation on endothelial function

Meta-regression for covariates
The covariates baseline vitamin D concentration, weekly dose of vitamin D supplement and study duration were investigated for FMD%, PWV and Alx. There was a tendency for the mean difference of FMD% to increase as weekly dose of vitamin D supplementation increased ($p = 0.05$). No other significant relationships were found (Supplementary Figures 13–21).

Study quality assessment
The median JADAD score was 5 out of a maximum score of 5 (Supplementary Table 5). Most studies scored 5; however, a few studies lost points due to either not describing the method of blinding or randomization.

Heterogeneity and publication bias
Most analyses demonstrated moderate to high heterogeneity with $I^2 = 90.38\%$ for FMD%, $I^2 = 58.23\%$ for PWV and $I^2 = 38.07\%$ for Alx. Egger funnel plots showed little evidence of publication bias with an intercept of $-4.05$ (95% CI $-8.76, 0.67$), $p = 0.083$ for FMD%, $-0.16$ (95% CI $-1.87, 1.55$), $p = 0.85$ for PWV and $-0.93$ (95% CI $-2.47, 0.61$), $p = 0.217$ for Alx (Supplementary Figures 22–24).

Discussion
This systematic review with meta-analysis and meta-regression found little evidence to support the use of vitamin D supplementation to improve endothelial function. Pooled data failed to demonstrate any statistically significant improvement in FMD%, PWV and Alx, all of which are considered markers of endothelial function. These results are consistent with previous analyses on vitamin D supplementation and endothelial function [9,12,21,23] and another analysis that has identified a lack of positive effects for treatment of hypertension with vitamin D [57]. Our results and those of these previous analyses contrast with a recent analysis [19], which demonstrated a significant improvement in FMD%. This analysis differed from ours by conducting the assessment of vitamin D supplementation on FMD% with studies that did not meet our inclusion criteria and included studies that conducted assessment of supplemental vitamin D and vitamin D analogues [58]. Our analysis included the study by Sluyter (2017) [43] which recruited the largest number of participants to date investigating the effects of vitamin D on central and brachial blood pressure parameters and failed to find any significant effect of vitamin D supplementation on PWV and Alx. However, a subgroup analysis in participants with vitamin D deficiency indicated significant improvements in PWV and Alx [43].

Results amongst pooled study data are conflicting and possibly due to heterogeneity of vitamin D dosing, dosing schedules, baseline concentrations, duration of intervention, study power [44,59,60] and possible confounding factors which included sunlight exposure, diet, medications and unverified additional supplementation [16,42]. Choice of vitamin D type may also affect RCT results, as new evidence indicates that vitamin D3 is more biologically active than vitamin D2 [61,62]. However, only three out of 26 studies [10,32,45] included in this meta-analysis used vitamin D2 supplementation. Several studies included supplemental or pharmacological interventions as a part of the selected population group, including calcium [42], HIV medications [37] and metformin [33]. Vitamin D receptor genetic variation [63], body mass index and disease-related biases for vitamin D metabolism may have also influenced results that vitamin D supplementation may have a positive effect on populations with T2DM [14,45] and CKD [15,40,50,64]. This was not confirmed by recent systematic reviews [22,65]. One study [66] suggest that higher risk of CVD due to hypovitaminosis may be due to a phenotypical trait over one or several specific CVD risk factors, with additional sex-related differences such as hormonal factors, as indicated with women appearing to have an increased risk of a number of CVD risk factors in comparison to men. Standardization of populations for comparison whilst accounting for external factors including environmental, smoking and dietary intake variation increase the challenge for identification of actual long-term effects of vitamin D supplementation.

Individually, only eight trials included in this systematic review reported a significant improvement in FMD%, PWV or Alx [14–16,32,35,40,45,50], when the intervention groups were compared to the placebo groups. Two of the studies [16,35] were conducted in populations of overweight African-Americans, one of these studies only included participants who were vitamin D deficient [16]. Two of the studies were conducted in populations with T2DM or at risk of T2DM [14,45], one [32] in a population of participants at an increased risk of T2DM. Three studies were conducted in CKD participants [15,40,50].

Our sub-analyses investigated the effects of vitamin D supplementation considering participants’ baseline vitamin D levels, dosing schedules, dose of vitamin D administered, and chronic diseases. Participants with deficient or inadequate baseline vitamin D levels tended to show some improvements in PWV and FMD%, respectively; conflictingly, PWV was significantly decreased in those with inadequate levels of vitamin D at baseline. Dosing schedules did not improve FMD%, PWV and Alx; however, higher doses of vitamin D significantly improved the Alx especially in participants with deficient or inadequate baseline vitamin D levels. Sub-analysis of T2DM and CKD participants showed no improvements in endothelial function with only one study of CKD participants [15] finding a significant increase in FMD%. Populations with CKD are commonly vitamin D deficient [51,67] somewhat due to a reduction in vitamin D receptors and receptor resistance [68–71], impaired vitamin D tubular resorption [72] and reduced hepatic and renal synthesis of 1,25-dihydroxycholecalciferol due to decreased hepatic CYP450 enzyme expression and reduced renal 1α-hydroxylase [73]. The effectiveness of higher doses of vitamin D supplementation to improve FMD% may be due to a reduction in renal hyperparathyroidism through vitamin D...
induced suppression of elevated intact parathyroid hormone [15,74] by increasing catabolism of vitamin D to 1,25 hydroxyvitamin D [75]. Additionally, vitamin D supplementation could increase control of hemodynamic factors through reduction of renin-angiotensin-aldosterone system activity reducing the effects of hypertension on CKD [76–78]. Vitamin D supplementation may improve cardiovascular risk factors in non-diabetic patients in early stages of CKD.

Our meta-regression analyses found no significant relationships for the covariates baseline vitamin D levels, weekly dose of vitamin D or study duration.

Further research is still required to understand the mechanistic effects of vitamin D and other endogenous compounds for prevention of atherosclerosis. Future research in populations that commonly use several supplements, including vitamin D and calcium, may help to determine if combination supplementation can improve endothelial function and reduce the risk of atherosclerosis, such as in post-menopausal females supplementing with calcium to decrease osteoporosis risk. Additionally, analysis of whether nutritional or supplemental sources decrease risk of atherosclerosis would be of benefit.

There are a number of strengths to this paper. No study intervention dose fell below minimum Australian recommended dosing requirements for study participants, being 200 IU/day for populations aged 19–50 years old, 400 IU/day for populations aged 50–70 years old, and 600 IU/day for populations aged 70 years or older [79]. The meta-analysis utilized data only provided by trials that were designed as placebo-controlled, randomized, double-blind trials. Stringent selection criteria focused specifically on vitamin D2 and D3, excluding vitamin D analogues like the commonly prescribed vitamin D receptor activator Paricalcitol for CKD. The primary rationale for this was to investigate recommendations for over-the-counter options for the consumer to assist with improving endothelial function.

There were several limitations that need to be highlighted for this systematic review and the meta-analysis and meta-regression. The shortest intervention time was four weeks [44] which may be less than the estimated required time period for vitamin D repletion to occur due to the accepted half-life of vitamin D being eight weeks, and possible variations between half-lives of vitamin D2 and D3 [63]. Most studies used a 16-week intervention period which could be argued as still being too short to observe vitamin D facilitated changes on surrogate markers of arterial stiffness. Seven studies ran over 20–24 weeks, two studies had a study duration of 1 year, and only one study was conducted for longer than 12 months [43]. Our meta-regression analyses did not confirm any improvements in endothelial function with increased study duration. FMD%, PWV or AIx were not the primary outcome measures in some studies, hence statistical power may have affected the results of these studies. Only studies that measured FMD%, PWV and AIx were included; however, several other indices are markers of endothelial function and hence the results need to be interpreted with this context.

In conclusion, this meta-analysis and meta-regression suggests that vitamin D supplementation has no significant effect for improving endothelial dysfunction, although there is a tendency to improve FMD%. The current evidence is conclusive enough to suggest that supplementation or pharmacological prescription of vitamin D is not warranted for improving endothelial dysfunction. It is still unclear whether vitamin D supplementation is required for those population groups that require repletion. There is some evidence that vitamin D supplementation may be beneficial for populations with CKD, or overweight populations.

Conflicts of interest

All authors declare to having no actual or potential conflict of interest including any financial, personal or other relationships with other people or organizations within three years of beginning the submitted work that could inappropriately, or be perceived to influence, their work.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.numecd.2019.08.005.

References


Dreyer G, Tucker AT, Harwood SM, Pearse RM, Raftery MJ, Yaqoob MM. Ergocalciferol and microcirculatory function in chronic kidney disease and concomitant vitamin D deficiency: an


