1	TITLE PAGE
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3	Manuscript title:
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5	The effect of vitamin D supplementation on knee osteoarthritis, the VIDEO study: a randomised
6	controlled trial
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61	Word count: 3285
62	Running title: Vitamin D in knee osteoarthritis

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Objective: Epidemiological data suggest that low serum 25-hydroxyvitamin D_3 (25-OH- D_3) levels are associated with radiological progression of knee osteoarthritis (OA). This study aimed to assess whether vitamin D supplementation can slow the rate of progression.

Method: A 3 year, double-blind, randomised, placebo-controlled trial of 474 patients aged over 50 with radiographically evident knee OA comparing 800 IU cholecalciferol daily with placebo. The primary outcome was rate of medial joint space narrowing (JSN) over three years. Secondary outcomes included lateral JSN, Kellgren and Lawrence grade, WOMAC pain, function, stiffness and the Get up and Go test.

Results: Vitamin D supplementation increased 25-OH-D₃ from an average of 20.7 (SD 8.9) μ g/L to 30.4 (SD 7.7) μ g/L, compared to 20.7 (SD 8.1) μ g/L and 20.3 (SD 8.1) μ g/L in the placebo group. There was no significant difference in the rate of JSN over three years in the medial compartment of the index knee between the treatment group (average -0.01 mm/year) and placebo group (-0.08 mm/year), average difference 0.08 mm/year, (95% CI [-0.14 to 0.29], p=0.49). No significant interaction was found between baseline vitamin D levels and treatment effect. There were no significant differences for any of the secondary outcome measures.

Conclusion: There is no clear evidence that vitamin D supplementation slowed the rate of JSN or led to reduced pain, stiffness or functional loss over a three year period. On the basis of these findings we consider that vitamin D supplementation has no role in the management of knee OA.

Abstract word count: 248/250

Key words: Vitamin D, knee, randomized placebo controlled trial, osteoarthritis

Introduction

Knee Osteoarthritis (OA) is a chronic, painful disease associated with considerable morbidity, costs and disability ¹. In the U.S., it is estimated that over a third of people aged over 60 have radiographic knee OA² and over 50% of these with knee OA will go on to have a total knee replacement in their lifetime³. At present there are no licensed treatments that alter disease progress and management is primarily concerned with symptom control to retain or improve joint function.

Vitamin D deficiency (defined as 25-hydroxyvitamin D₃(25-OH-D₃) serum levels below 20μg/mL ^{4 5}) is common in the UK with estimates of over 12% for people living in private households and 30% of care home residents in the over 65s. There has been considerable interest in the association between vitamin D deficiency and OA incidence and progression. Vitamin D has a number of important biological functions in bone, cartilage and muscle⁶ which has led to the hypothesis that vitamin D supplementation may prevent the progression of OA. There is evidence from a number of, but not all, epidemiological studies suggesting that low dietary intake of vitamin D and low serum 25-OH-D₃ levels are associated with increased radiological progression of knee OA ⁷⁻¹³. Epidemiological data from the Framingham Study demonstrated that low vitamin D intake was associated with a three to four-fold increased risk of radiographic progression at two skeletal sites over 8-10 years.⁷ Further analysis of a separate cohort of patients in the Framingham study, along with another cohort from the Boston Osteoarthritis of the Knee Study (BOKS) found no association between vitamin D status and joint space or cartilage loss in knee OA ¹².

Findings from RCTs have thus far not conclusively settled this debate ¹⁴⁻¹⁷. A 12 month trial of vitamin D in 107 vitamin D insufficient subjects with knee OA found a small but statistically significant improvement in pain ¹⁴. A trial of 146 subjects with symptomatic knee OA found that vitamin D

supplementation for two years had no effect on the structural progression of OA using MRI as the primary outcome ¹⁶. A further post hoc analysis of a RCT concluded that calcium plus vitamin D supplementation for two years in post-menopausal women had no effect on self-reported frequency or severity of joint symptoms ¹⁷. As these trials were heterogeneous in terms of patients recruited, sample sizes and some also used calcium in addition to vitamin D supplements, it is important to have a large RCT with a prolonged follow up to provide further clarity on the role of vitamin supplementation in patients with knee OA.

Aim

The primary aim of this trial was to determine whether vitamin D supplementation can reduce the rate of structural progression of knee OA as measured by change in medial joint space assessed on a weight-bearing radiograph over a 3-year period. Secondary outcomes included changes in pain and function.

Methods

Study design

The VIDEO study was a double-blind, randomised, placebo-controlled trial performed at five UK NHS hospitals. Participants were randomly assigned to receive either 800IU of oral cholecalciferol or matched placebo daily. Data from clinical trials indicated that 800IU/day of cholecalciferol can produce significant increases in serum 25-hydroxyvitamin D₃ levels and that these increases are evident within one month of starting treatment¹⁸. The protocol was approved by the Scotland A Research Ethics Committee and the trial was registered with EudraCT: ref. 2004-000169-37,

ISRCTN94818153, CTA No. 11287/0001/001. The trial was conducted in accordance with Good Clinical Practice guidelines and the Declaration of Helsinki.

Participants were identified from GP lists, patient referrals to hospitals and via radio advertisements. Patients were eligible if they: were aged >50 years, ambulatory, had radiological evidence of knee OA at medial tibio-femoral knee compartment (Modified Kellgren & Lawrence (K&L) score 2/3, JSW >1mm) and knee pain for most days of the previous month. Reasons for exclusion were: secondary OA, inflammatory arthritis, early morning knee stiffness for >30 minutes, cod liver oil or vitamin supplementation containing vitamin D >200 IU, glucosamine or chondroitin use for <three months, osteoporotic fracture, previous knee surgery or arthroscopy within six months, use of bisphosphonates within two years. Eligible participants were invited to a screening appointment. Informed consent was taken along with knee radiographs, which were assessed by the local clinician to determine eligibility.

Randomisation and blinding

Eligible participants were randomised centrally by the UK Medical Research Council Clinical Trials

Unit (MRC CTU) via telephone to receive either oral vitamin D or matching placebo tablets (1:1) by

computer-generated randomisation with stratification by recruitment centre. Treatment allocation

was concealed from the patients, clinicians, outcome assessors and investigators. Both the active

treatment and placebo were manufactured by Thompson and Capper Ltd, and packed by Bilcare

Global Clinical Supplies (Europe) Ltd.

Trial procedures

At the baseline visit knee bilateral radiographs and blood samples were taken, and the assigned drug dispensed in six month packs. Radiographs and blood sampling were repeated at 12 months and 36 months. Questionnaires (WOMAC) were completed at 6-monthly intervals until the final visit. Blood was drawn to measure serum 25-OH-D $_3$ at baseline and 12 months to assess baseline vitamin D status and response to supplementation. Serum vitamin D $_2$ and D $_3$ concentrations were assayed at King's College Hospitals NHS Foundation Trust via mass spectrophotometry using the MassChrom reagent kit (Chromsystems Instruments & Chemicals GmbH).

Outcome measures

compartment of the index knee (knee with the smallest joint space width (JSW) at baseline in the case of bilateral disease), as measured by the rate of JSN (mm/year) over the three years. Knee X-rays were taken using the MTP technique ¹⁹ using a foot map to improve accurate re-positioning at follow up visits.

All joint space measurements were performed by a single reader. Reproducibility was excellent, and comparable to previous results using the same software package ^{20, 21}; intra-rater intra-class correlation coefficients (ICCs) were: 0-96 medial 95% CI [0-88-0-98], 0-98 lateral 95% CI [0-94 0-99].

Secondary outcomes measures included: rates of change in minimum JSW of the lateral compartment, and of the medial and lateral compartments of the contralateral knee, Kellgren and Lawrence (K&L)^{22, 23} grade, WOMAC VAS scores (0-100 pain, stiffness, function and total) in the index

The primary outcome measure was radiological progression of knee OA in the medial joint

knee, and Get up and Go test. Baseline and follow-up X-rays were graded for K&L grade by a Clinical Orthopaedic Fellow, with an intra-reader Kappa of 0.68.

Sample size

The study was designed to detect a clinically important mean difference of 0·22mm/year in the rate of JSN between treatment groups over three years, assuming a standard deviation of 0·7 mm ^{24, 25}, with 80% power at the 5% significance level. Allowing for 32% drop-out rate, the total sample size required was 470.

Statistics

specified analysis plan which was finalised prior to database lock and breaking the blind.

To assess JSN a longitudinal analysis was performed using a linear mixed regression model with fixed effects for treatment, time, treatment by time and adjustment for: baseline JSW, centre, gender, glucosamine or chondroitin use, age and BMI. To allow for between patient differences the model included a random patient intercept. The central parameter of interest was the treatment by time interaction, which represents the average difference in the rate of JSN/year between the treatment groups. Continuous secondary outcomes were analysed similarly. Changes in ordinal outcomes over time were analysed using ordinal logistic regression models with robust Huber-White sandwich estimators of standard errors. The effect of treatment on the proportion of patients with clinically

Analysis was conducted following the intention-to-treat principle and in accordance with a pre-

significant progression (JSN>0.5mm in the index knee) at three years was obtained using a Poisson

regression model with robust error estimates. For patients who had a total knee replacement (TKR) in the index knee during the trial, clinically significant progression was assumed.

Mean imputation was used to deal with missing covariate values ²⁶. For patients who had TKR during the trial, data before surgery was included and data after surgery assumed to be missing. All missing outcome values were assumed to be missing at random and multiple imputation by chained equations was used ^{27, 28}. Sensitivity analyses, including analysis of the complete cases and a range of missing not at random mechanisms, were performed to assess the robustness of the primary results to the effect of missing data (for full details see supplementary file eTable 2 and eFigure 1). All statistical analyses were performed using Stata/IC version 12·1 (StataCorp, College Station, TC, USA).

Results

In total, 474 participants were recruited between 19/01/2005 and 13/06/2008. Table 1 shows baseline clinical data and baseline radiographic characteristics. Additional baseline variables can be found in the supplementary file, eTable 1. The treatment and placebo groups were well matched for clinical characteristics and showed a similar distribution of radiographic characteristics. The distribution of serum 25-OH-D₃, divided into tertiles (table 3), was almost identical in the two groups, with 50% of both groups vitamin D₃ deficient ($<20\mu g/L$).

As shown in Figure 1, 198 of participants in the placebo group (84%) and 188 of those in the treatment group (79%) attended the 3-year follow-up visit. Six patients in the placebo group and seven in the vitamin D group received a TKR of the index knee during the follow up period. Due to a combination of technical and logistic reasons, including poor positioning and quality a number of radiographs from attending patients, including baseline, could not be evaluated for JSW accurately. JSW in the medial compartment of the index knee was missing for a total of 37/474 patients (8%) at

baseline (18/237 placebo, versus 19/237 active), 110/474 patients (23%) at year one (58/237 placebo versus 52/237 active) and 183/474 (39%) at year three (87/237 placebo versus, 96/237 active). 38% of the missingness at year one (42/110) was due to unreadable X-rays (23 placebo and 19 active). 30% of the missingness at year three (55/183) was due to unreadable X-rays (27 placebo versus 28 vitamin D). The remaining missingness at year three occurred due to withdrawal 54% (99/183, 49 placebo (3 with TKR of index knee at one year) and 50 active (1 with TKR of index knee at one year)), loss to follow-up 10% (18/183, 7 placebo and 11 active), TKR of the index knee 5% (9/183, 3 placebo and 6 active) or death 1% (2/183, 1 placebo and 1 active). Missingness of X-ray data did not vary by treatment arm. 380/474 patients (189/237 placebo, 191/237 active) had baseline and at least one follow up JSW reading available and were analysed separately as a sensitivity analysis. A separate analysis of the 242/474 patients (125/237 placebo, 117/237 active) with complete follow-up was also performed along with additional sensitivity analysis to assess the impact of missing data (supplementary file eTable 2 and eFigure 1).

Vitamin D analysis

At 12 months, serum vitamin D_3 levels had increased from an average of 20.7 (8.9) μ g/L at baseline to 30.4 (7.7) μ g/L in the vitamin D group. Levels decreased for those receiving placebo from 20.7 (8.1) μ g/L at baseline to 20.3 (8.1) μ g/L at 12 months (table 3). The number of patients with vitamin D deficiency (<20 μ g/L) fell to 7% in the vitamin D group but rose to 54% in the placebo group.

Radiographic results

There was no significant difference in the rate of JSN over three years in the medial compartment of the index knee between treatment groups (-0.01mm/year versus -0.08mm/year for vitamin D and placebo respectively), between group difference 0·08 mm/year, 95% CI [-0·14 to 0·29], p=0·49 (figure 2, table 2). Sensitivity analyses conducted to assess the effect of missing values on the estimated treatment effect produced results no different from the primary analysis (supplementary file eTable 2 and eFigure 1). No interaction between baseline vitamin D status and treatment effect (Δ) was found (<20 µg/L, Δ 0·06, 95% CI [-0·20 to 0·32]; 20 µg/L to 30 µg/L, Δ 0·05, 95% CI [-0·20 to 0·29]; >30 µg/L, Δ 0·05, 95% CI [-0·30 to 0·40]) (Figure 3).

There was no difference in the proportion of patients with clinically significant progression of JSN (JSN>0.5mm in the index knee) at three years between the vitamin D group (39%) and placebo group (37%). The absolute risk difference was 2% (95% CI [-10% to 14%], p = 0.76) (eTable 4).

We explored the hypothesis that there may be an interaction between treatment effect and baseline JSN. The interaction did not reach significance (p=0.86, N=474).

Secondary outcomes

The placebo group showed an increase in WOMAC pain whereas the vitamin D group showed a small decrease (0.71 versus -0.08 per year, between group difference -0.79, 95% CI [-2.31 to 0.74], table 2, eFigure 2). WOMAC stiffness decreased in both groups (-2·02 versus -0.50 per year for vitamin D and placebo groups respectively, between group difference -1.52, 95% CI [-3.24 to 0.21]). WOMAC

function increased for both groups (0.42 versus 1.07 per year for vitamin D and placebo, between group difference -0.65, 95% CI [-2.09 to 0.79]) . None of the above differences achieved statistical significance.

Odds ratios of a higher K&L grade per year were calculated as 1.32 (Vitamin D) and 1.23 (placebo) for the index knee and 1.19 (Vitamin D) and 1.18 (placebo) for the contralateral knee. This gave a treatment by time odds ratio, which represents the increase in odds of a higher K&L grade per year for vitamin D patients relative to placebo, of 1.07 (95% CI [0.88 to 1.31]) for the index knee and 1.01 (95% CI [0.80 to 1.27]) for the contralateral knee (table 2). The odds of a higher get up and go test grade per year for Vitamin D patients was 1.00 and 1.04 for placebo patients. There was no significant difference in the odds of a higher get up and go test grade over time between the treatment groups (OR = 0.96, 95% CI [0.73 to 1.27]). Additional secondary outcomes were assessed and treatment effect estimates can be found in the supplementary file eTable 4. All outcomes at three years are summarised in eTable 5.

Adverse events

There was no difference in the proportion of patients experiencing SAE's between the vitamin D (59/237, 25%) and placebo group (64/237, 27%), p = 0.67 or in the rates of occurrence of hypercalcaemia (five placebo, three vitamin D) or hypercalciuria (34 placebo, 46 vitamin D).

Discussion

There is no clear evidence that vitamin D supplementation, at a dose of 800 IU cholecalciferol daily, had an effect on the progression of knee OA over the three year period, as measured by changes in JSW, or on knee pain, function or stiffness. This is despite the fact that participants had high rates of vitamin D deficiency at trial entry, and the level of supplementation was sufficient to increase serum vitamin D levels by $10~\mu g/L$ on average in the first year of treatment, reducing the proportion of participants with deficiency by over 80%.

Previous research has not provided a consensus on the effect of vitamin D on the progression of knee OA, with observational studies and RCTs generating conflicting findings. Several high quality epidemiologic studies have demonstrated an association between low serum vitamin D and /or vitamin D intake and the risk of either OA incidence or progression ⁸⁻¹¹, however others have shown no association ^{12, 13, 15, 29-31}. These studies vary in methodology and were also subject to a number of important biases.

McAlindon performed a two year RCT of 2000 IU/day oral cholecalciferol for patients with symptomatic knee OA. The primary outcomes were cartilage volume loss measured by MRI and knee pain by WOMAC. The population studied had similar baseline concentrations of vitamin D but greater baseline JSW (approximately 5mm vs. 3.5mm). The results demonstrated that despite 61.3% of patients achieving target concentrations of vitamin D, there were no significant improvements over placebo in any of the outcomes. Sanghi *et al* performed a 12 month RCT of vitamin D

supplementation in patients with knee OA and vitamin D deficiency ¹⁵. They demonstrated a statistically significant reduction in pain and increase in physical function in a group taking vitamin D compared with placebo, however the difference between the two groups was not deemed to be clinically important ³².

The results from our study, which is substantially larger than the previous studies, are consistent with the above results. The VIDEO trial contributes several new findings. Firstly, we measured JSN and K&L grade in the contra-lateral knee. This is important as pathogenic mechanisms may be different in the contra-lateral joint compared with the index knee which exhibits later stage disease in patients with bilateral OA, as suggested in the Doxycycline trial by Brandt *et al* ²⁵. In addition, we measured JSN in the medial and lateral compartments individually. Although medial compartment disease is far more prevalent, and the majority of previous studies focus only on joint space changes in the medial compartment ^{4, 25}, it is important to measure JSN in the lateral compartment to ensure disease progression is not missed ³³. We looked at the association of the treatment effect with baseline [25-OH-D₃] concentration and the change in vitamin D concentration after 12 months of treatment. This study has a longer follow-up period than previous trials, with three year JSN having been shown in a previous study to be predictive of the incidence of osteoarthritis related knee surgery ³⁴.

Strengths and potential limitations

A key strength of VIDEO was the inclusion of patients who were not biochemically vitamin D deficient. Laslett *et al* found that vitamin D deficiency was associated with incident or worsening of knee pain over a five year period ³⁵, suggesting that vitamin D supplementation would be effective in attenuating the progression of knee pain only in those who already show moderate deficiency.

However, 50% of VIDEO participants had vitamin D insufficiency ($<20 \,\mu g/L$) at baseline. When analysis of treatment effect on JSN was broken down by baseline vitamin D status, no significant interactions with the treatment effect were found. Vitamin D supplementation had no effect on the change in joint space width even in subjects who were vitamin D deficient.

We acknowledge limitations. The radiographs from the screening visits were read by the local PI at each centre to establish eligibility into the trial. A clinical orthopaedic fellow re-read all the baseline x-rays for the final analysis. This explains why a proportion of the baseline radiographs were determined to be K&L grade 1, while the inclusion criteria specified K&L ≥2. The difference between the definitions of the two grades relates to a possible vs. definite osteophyte, this boundary being particularly subjective. The distribution however was similar between the two groups and would be unlikely to bias the results of the trial. Of interest, it allowed us to assess the effect of vitamin D in very early OA.

The proportion of participants lost to follow-up by the three year visit (16% placebo group, 21% treatment group) could be considered a limiting factor. This rate of loss is consistent with other OA trials ^{4, 17, 25, 36} and the sample size calculation allowed for 32% loss to follow up. An additional number of x-rays were unevaluable for JSW due to technical and logistic reasons. However, there was no evidence of a differential loss to follow up or unevaluable X-rays between treatment arms and detailed sensitivity analyses to assess the impact of missing data (described in supplementary file) were consistent with the primary analysis.

Conclusions

There is no clear evidence that vitamin D supplementation, at a dose sufficient to elevate serum vitamin D_3 levels by 10 μ g/L in one year, slowed the rate of JSN or led to reduced pain, stiffness or

functional loss over a three year period, when compared with placebo. On the basis of these findings we consider that vitamin D supplementation has no role in the management of knee OA.

Acknowledgements

We would like to thank the support of the staff of the VIDEO Osteoarthritis Study including the staff at MRC CTU at UCL who managed/conducted the VIDEO trial, the VIDEO research nurses at Southampton Centre for Biomedical Research, the Royal National Orthopaedic Hospital, Royal Victoria Infirmary Newcastle, Salford Royal Hospital, and Norfolk and Norwich University Hospital. Additional thanks to Dr Iva Hauptmannova, PhD, Royal National Orthopaedic Hospital, for her support and assistance throughout the study on behalf of the sponsor site.

We would like to acknowledge Dr Kirsten Leyland, PhD, University of Oxford, for support and assistance with all aspects of the radiographic measurements, and Charlotte Arden for performing joint space width measurements

We would also like to thank the participants of the VIDEO study who made this work possible.

Author Contributions

RK, NKA, FB, TWON, AM, CC, CJD contributed to the design of the work and acquisition of the data.

AB and SAT contributed to the acquisition of the data. SC, CJD, SS, DJH, SJ contributed to the analysis

of the data.

All authors contributed to drafting the work or revising the content critically and all authors have approved the final version.

NKA had full access to all of the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

389 Role of the funding source 390 391 Funding was received from Arthritis Research Campaign (now Arthritis Research UK, grant number 392 K0576). Additional support was received from the NIHR Musculoskeletal Biomedical Research Unit, 393 University of Oxford. Researchers were independent from funders and sponsors. The funder of the 394 study had no role in study design, data collection, data analysis, data interpretation, or writing of the 395 report. The corresponding author had full access to all the data in the study and had final 396 responsibility for the decision to submit for publication. 397 398 **Conflict of interests** 399 400 All authors have completed the Unified Competing Interest form at 401 www.icmje.org/coi_disclosure.pdf and declare the following interests: 402 NA reports consultancy work for Merck, Roche, Smith & Nephew, Q-Med, Nicox, Flexion, payment 403 for lectures from Bioiberica and Servier, outside of the submitted work. 404 CC reports personal fees from Servier, personal fees from Amgen, personal fees from Eli Lilly, 405 personal fees from Merck, personal fees from Medtronic, personal fees from Novartis, outside the 406 submitted work. 407 408 409 **Ethics statement** 410

411	The trial was registered with EudraCT: ref. 2004-000169-37, ISRCTN94818153, CTA No.
412	11287/0001/001, and the protocol received full approval from the Scotland A Research Ethics
413	Committee (NHS REC Application Reference: 04/MRE10/30). The full protocol can be accessed at
414	http://www.ctu.mrc.ac.uk/our_research/research_areas/other_conditions/studies/video/.
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416	Data sharing statement
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418	Anonymised patient level data and statistical code available from the corresponding author at
419	nigel.arden@ndorms.ox.ac.uk.
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554	Figure Legends
555	
556	Figure 1. Consort flow diagram for the VIDEO study
557	Figure 2. Mean Joint Space Width in the medial compartment of the index knee with 95% CI's by
558	treatment group (N = 474). All available readings were included in primary analysis and multiple
559	imputation was used to impute missing values, assuming all missing outcome values were missing at
560	random, conditional on treatment and the covariates included in the imputation model. Both centre
561	and baseline BMI were included in the imputation model.
562	Figure 3. Scatterplot of baseline Vitamin D₃ against three year change in Joint Space Width by
563	treatment group with linear fit imposed (N = 463).
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Table 1 Baseline Clinical and radiographic Characteristics as mean (sd) or number (%).

	N vitamin D /	Vitamin D	Placebo
	N Placebo		
Age (yrs)	237/237	64 (8)	64 (8)
Sex: (% Female)	237/237	144 (61%)	145 (61%)
Index knee: % Right	237/237	136 (57%)	146 (62%)
BMI (kg/m²)	236/237	30 (5)	29 (5)
Family history of knee or hip OA	236/235	113 (48%)	109 (46%)
Heberdens nodes	237/237	145 (61%)	165 (70%)
Bouchards nodes	237/237	71 (30%)	83 (35%)
CMC joint OA	237/237	105 (44%)	101 (43%)
% Bilateral knee OA	237/237	169 (71%)	166 (70%)
% Taking analgesics	237/237	104 (44%)	98 (41%)
% Taking glucosamine or	237/237	109 (46%)	104 (44%)
chondroitin			
% Taking cod liver oil	236/236	73 (31%)	78 (33%)
WOMAC pain score	236/232	33 (18)	31 (19)
WOMAC function score	236/232	36 (21)	35 (20)
WOMAC stiffness score	236/231	47 (24)	43 (24)
WOMAC total score	236/232	36 (19)	35 (19)
Worst K&L grade+ (of	234/236		
medial/lateral)			
Index knee:			
0		3 (1%)	3 (1%)
1		62 (26%)	59 (25%)
2		86 (37%)	92 (39%)

3		70 (30%)	66 (28%)
4		13 (6%)	16 (7%)
Worst K&L grade ⁺ (of			
medial/lateral)			
Contra-lateral knee:			
0	234/236	2 (1%)	2 (1%)
1		77 (33%)	87 (37%)
2		65 (28%)	70 (30%)
3		54 (23%)	43 (18%)
4		29 (12%)	26 (11%)
TKR Contra-lateral knee		7 (3%)	8 (3%)
Medial JSW index knee (mm) $^{\scriptscriptstyle \dagger}$	218/219	3.49 (1.48)	3.58 (1.47)
Lateral JSW index knee (mm) †	222/219	5.27 (1.95)	5.42 (1.87)
Medial JSW Contra-lateral knee [†]	214/213	3.40 (1.69)	3.62 (1.60)
(mm)			
Lateral JSW Contra-lateral knee [†]	216/212	5.38 (2.07)	5.22 (1.90)
(mm)			
Baseline Vitamin D ₃ (in μg/L)		20.7 (8.9)	20.7 (8.1)

^{*}Baseline X-rays were missing for 3 individuals in the vitamin D group. 1 placebo patients X-ray disc was corrupt therefore could not be read. Due to X-ray quality issues, including poor positioning, the numbers of readable JSW measures vary by region and by knee.

Table 2 Treatment effect estimates for primary and secondary outcomes

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Rate of change of Joint Space width	Vitamin D	Placebo	Difference [95% CI]
(mm/year)			
Primary Outcome:			
Medial compartment index knee	-0.01	-0.08	0.08 [-0.14 to 0.29]
Secondary Outcomes:			
Lateral compartment index knee	-0.11	-0.18	0.07 [-0.19 to 0.33]
Medial compartment contra-lateral	-0.03	0.03	-0.06 [-0.26 to 0.13]
knee			
Lateral compartment contra-lateral	-0.10	-0.07	-0.03 [-0.27 to 0.21]
knee			
	Vitamin D	Placebo	Difference [95% CI]
Clinically significant progression	39%(N=92)	37%(N=88)	2% [-10% to 14%] ¹
(Medial index JSN>0.5mm)			
Rate of change per year	Vitamin D	Placebo	Difference [95% CI]
WOMAC pain	-0.08	0.71	-0.79 [-2.31 to 0.74]
WOMAC stiffness	-2.02	-0.50	-1.52 [-3.24 to 0.21]
WOMAC function	0.42	1.07	-0.65 [-2.09 to 0.79]
WOMAC total	0.11	0.84	-0.72 [-1.92 to 0.48]
	Vitamin D	Placebo	Treatment x Time
			OR [95% CI]
Odds of a higher K&L grade per year	1.32	1.23	1.07 [0.88 to 1.31]
index knee			
Odds of a higher K&L grade per year	1.19	1.18	1.01 [0.80 to 1.27]
contra-lateral knee			
Odds of higher grade in Get up and	1.00	1.04	0.96 [0.73 to 1.27]

go test per year

584	N=474 (N=237 Vitamin D, N = 237 Placebo). WOMAC scores range from 0 to 100, 0 = no
585	pain/disability, 100 = extreme pain/disability. Get up and Go test graded 1 - normal to 6 – abnormal.
586	¹ Corresponds to a relative risk of 1.05 [0.77 to 1.44].
587	

	N vitamin D /	Vitamin D	Placebo
	N Placebo		
Baseline Vitamin D₃:	232/231		
<20 μg/L		117 (50%)	115 (50%)
20 μg/L to 30 μg/L		79 (34%)	87 (38%)
>30 μg/L		36 (16%)	29 (12%)
Baseline Vitamin D ₃ (in μg/L)		20.7 (8.9)	20.7 (8.1)
Baseline Vitamin D ₂ :	232/231		
<2.2 μg/L		228 (98%)	218 (94%)
≥2.2 μg/L		4 (2%)	13 (6%)
Baseline Vitamin D ₂ (in μg/L)*	4/13	5.0 (2.7)	3.8 (1.7)
12 month Vitamin D _{3:}	206/206		
<20 μg/L		14 (7%)	111 (54%)
20 μg/L to 30 in μg/L		97 (47%)	67 (32%)
>30 μg/L		95 (46%)	28 (14%)
12 month Vitamin D ₃ (in μg/L)		30.4 (7.7)	20.3 (8.1)
12 month Vitamin D ₂ :	206/206		
<2.2 μg/L		203 (99%)	193 (94%)
≥2.2 µg/L		3 (1%)	13 (6%)
12 month Vitamin D ₂ (in μg/L)*	3/11	3.3 (0.76)	4.2 (2.3)

12 month change Vitamin D_3 (µg/L) 201/201 9.4 (8.3) -0.8 (5.7) *Vitamin D_2 reported in μ g/L for patients with Vitamin $D_2 \ge 2.2 \mu$ g/L only. Data presented as mean(sd) or number (%) for categorical variables. Vitamin D₃ and Vitamin D₂ were not available at baseline for 5 vitamin D and 6 placebo patients and at 12 months for 31 vitamin D and 31 placebo patients, for reasons unknown.

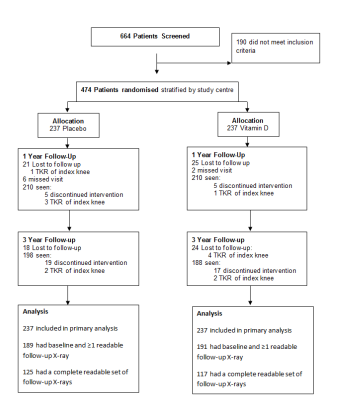


Figure 1. Consort flow diagram for the VIDEO study

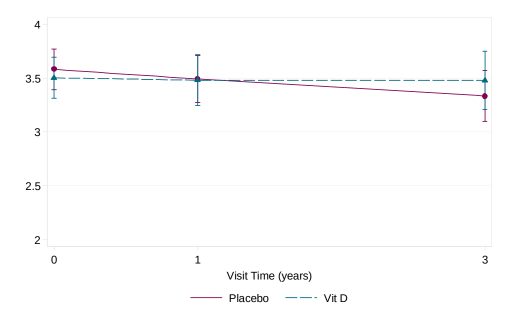


Figure 2. Mean Joint Space Width in the medial compartment of the index knee with 95% Cl's by treatment group (N = 474 All available readings were included in primary analysis and multiple imputation was used to impute missing values, assuming all missing outcome values were missing at random, conditional on treatment and the covariates included in the imputation model. Both centre

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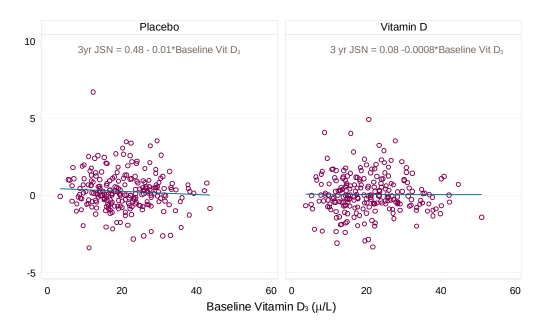


Figure 3. Scatterplot of baseline Vitamin D_3 against estimated three year change in Joint Space Width by treatment group with linear fit imposed (N = 463).