High-dose vitamin D supplementation in multiple sclerosis – results from the randomized EVIDIMS (efficacy of vitamin D supplementation in multiple sclerosis) trial

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Abstract

Background: Epidemiological, preclinical, and non-interventional studies link vitamin D (VD) serum levels and disease activity in multiple sclerosis (MS). It is unclear whether high-dose VD supplementation can be used as an intervention to reduce disease activity.

Objectives: The study aimed to compare the effects of every other day high- (20,400 IU) versus low-dose (400 IU) cholecalciferol supplementation on clinical and imaging markers of disease activity in patients with relapsing–remitting MS or clinically isolated syndrome.

Methods: The EVIDIMS (efficacy of vitamin D supplementation in multiple sclerosis) trial was a multicentre randomized/stratified actively controlled explorative phase 2a pilot trial with a double-blind intervention period of 18 months, add on to interferon-β1b.

Results: Fifty-three patients were randomized, and 41 patients completed the study. Cholecalciferol supplementation was well tolerated and safe in both arms. After 18 months, clinical (relapse rates, disability progression) and radiographical (T2-weighted lesion development, contrast-enhancing lesion development, brain atrophy) did not differ between both treatment arms. Post-study power calculations suggested that the sample size was too low to prove the hypothesis.

Conclusions: The results neither support nor disprove a therapeutic benefit of high-dose VD supplementation but provide a basis for sound sample size estimations in future confirmatory studies. www.clinicaltrials.gov/NCT01440062

Keywords: Multiple sclerosis, clinical trial, vitamin D, efficacy, treatment, supplementation

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Introduction

Preclinical and epidemiological studies link 25-hydroxy (OH) vitamin D, hereafter referred to as vitamin D, serum levels to disease activity of multiple sclerosis (MS). In particular, associations of higher vitamin D levels in patients with MS with favourable clinical and magnetic resonance imaging (MRI) parameters of disease activity have been reported.1–3 Based on these data, MS clinicians commonly boost vitamin D levels in patients by oral cholecalciferol or ergocalciferol supplementation.

While safety data are rather unequivocal and suggest that vitamin D supplementation up to 280,000 IU/week is fairly safe,4 prospective data from interventional treatment trials of vitamin D supplementation on clinical efficacy, doses needed, and optimal serum levels in MS are sparse, inconsistent, and in part conflicting.5,6 A phase I/II randomized controlled open-label study comparing continuous intake of up to 40,000 IU/day or 4000 IU/day for 52 weeks in 49 patients with MS showed a significant reduction of annualized relapse rate in the...
The EVIDIMS clinical trial was a multicentre, randomized, stratified, active controlled, double-blind, explorative phase II trial. Two doses of vitamin D3 (cholecalciferol) were compared: in the high-dose arm participants received 20,400 IU (1 ml cholecalciferol oil (20,000 IU) +1 tablet (400 IU)) every other day for 18 months, while in the low-dose arm 400 IU (1 ml placebo oil (0 IU) +1 tablet (400 IU)) were given every other day. To rule out confounding effects of variable disease-modifying treatments, all participants were required to be on stable treatment with interferon-beta-1b, started at least 3 months prior to inclusion. Interferon-beta treatment was continued throughout the study.

Patients with relapsing–remitting (RR)MS or clinically isolated syndrome (CIS) were recruited at seven German study centres (Berlin (four centres), Potsdam, Halle, Teupitz; for details see supplementary information). After a short screening period eligible patients were stratified according to gender and vitamin D level (< or ≥20 ng/ml (50 nmol/l)) and then randomized 1:1 to either high-dose or low-dose treatment as an add-on to continuous treatment with IFN β1b. Clinical evaluations including Expanded Disability Status Scale (EDSS)12 and MS Functional Composite (MSFC)13 were performed at screening and then every 6 months. Standardized MRI was performed at screening and after 12 and 18 months using identical hardware for all patients (at the Charité study centre, Berlin). Pharmacovigilance laboratory testing was done after 4 weeks and thereafter every 3 months.

The primary endpoint was the number of new T2-weighted (T2w) hyperintense lesions on brain MRI at month 18 compared with baseline. Major secondary endpoints were relapse rate and disability progression on the clinical side, T2w lesion volume, number of gadolinium (gd) lesions, and brain atrophy on the MRI side, and number of adverse events on the safety side. More detailed information on the design and conduct of the EVIDIMS trial has been published previously,11 and additional information is provided as supplementary information. The study protocol (in German) is available from the corresponding author upon qualified request.

The study was approved by the German Federal Institute for Drugs and Medical Devices (BfArM, 4037578), as well as by the local ethics committees (11/0386-ZS Ek 13) and is registered at European Clinical Trials Database (EudraCT 2011-002785-20) and ClinicalTrials.gov (NCT01440062). It was conducted strictly adhering to the study protocol and to applicable national laws (Arzneimittelgesetz, 14. Novelle, 2005), the Harmonized Tripartite Guideline for Good Clinical Practice (ICH GCP), and the principles of the Declaration of Helsinki in its applicable version. All participants gave written informed consent at screening prior to any study-related procedures.

Study population
The main inclusion criteria were: male and female patients with either CIS or a definite diagnosis of MS according to the 2005 revised McDonald criteria14 and a relapsing–remitting disease course; age at randomization 18–65 years; EDSS score 0–6; continuous treatment with IFN β1b for at least 3 months; and no relapse within 30 days prior to randomization. Important exclusion criteria were: relevant hepatopathy or renal dysfunction; history of sarcoidosis, nephrolithiasis, pseudo hypoparathyroidism; vitamin D supplementation >500 IU/day within 6 months prior to randomization; pregnancy; hypercalcemia or urine calcium/creatinine ratio >1; concomitant medication with hydrochlorothiazide, digoxin, digoxin, phenytoin, barbiturates; inability to provide informed consent; incompatibility with MRI procedures. A comprehensive listing of inclusion
Outcome measures

Clinical evaluations and safety laboratory tests. All clinical evaluations and scorings were performed by trained study personnel blinded to the patients’ treatment allocations according to standardized operating procedures. MS relapses were defined as: (re)occurrence of new or previous central nervous system dysfunction in the absence of infections or hyperthermia, duration ≥24 h, time-lag from the onset of previous relapse ≥30 days. Disability was assessed by EDSS and MSFC.12,13 EDSS rating was performed by trained (Neurostatus) and board-certified physicians otherwise not involved in the management of study participants. MSFC scoring was done by trained study personnel. All study-related safety laboratory tests were done by Bioscientia laboratories (Bioscientia Institut für Medizinische Diagnostik GmbH, Ingelheim, Germany).

MRI. Standardized MRI was performed in all participants in one study centre at Charité - Universitätsmedizin Berlin, using a Magnetom TIM TRIO 3 Tesla MRI (Siemens Healthineers, Erlangen, Germany). A sagittal 3D SPACE T2w sequence (TR 5,000 ms, TE 502 ms, TI 900 ms, flip angle 9 degree, isotropic resolution 1 mm3, GRAPPA 2) was used to obtain T2w images. In addition, we applied a 3D fluid-attenuated inversion recovery sequence (SPACE-FLAIR, TR 6,000 ms, TE 388 ms, TI 2,100) with identical spatial parameters, and a high-resolution 3-dimensional T1w sequence (MPRAGE, TR 1,900 ms, TE 3.03 ms, TI 900 ms, flip angle 9 degree, isotropic resolution 1 mm3 GRAPPA 2) with identical spatial parameters. A 3D T1w sequence (VIBE, TR 4.8 ms, TE 2.2 ms, flip angle 9 degree, isotropic resolution 1 mm3 GRAPPA 2) with identical spatial parameters was applied 5 min after injection of 0.1 mmol/kg gd-DTPA (Gadovist, Bayer, Berlin, Germany). Raw data were transferred to a Linux workstation and processed following a semi-automated procedure, including an image coregistration (FMRIB’s Linear Image Registration Tool, FMRIB Analysis Group, University of Oxford, Oxford, UK) and inhomogeneity correction routine embedded into the MedX v.3.4.3 software package. Brain atrophy (percentage brain volume change (PBVC) and normalized brain volume (NVB)) was estimated with SIENA.15 part of FSL.16 Thalamic volumetry was performed using FIRST (FMRIB Analysis Group, University of Oxford, Oxford, UK).

Statistical analyses. Due to lack of data on the effect of vitamin D on T2w lesions, no statistical sample size calculation was performed in advance using a given error of the 1st kind and stipulated power. A sample size of 80 patients (40 in each arm) was planned, based mainly on feasibility.

Due to the explorative character of the study, statistical testing has to be understood as explorative, and data analyses were mainly descriptive for all endpoints. For univariate independent group comparisons exact Mann–Whitney U tests and exact Chi-Square tests were used. Survival data such as MS relapses were analysed by Kaplan–Meier curves with appropriate log-rank tests and multiple Cox-regressions. For time series data, a nonparametric multivariate analysis of longitudinal data (MANOVA) in a two-factorial design was applied (1st factor (independent): treatment groups, 2nd factor (dependent): study visits). When appropriate, multivariate nonparametric analysis of covariance (MANCOVA) using baseline values and additional parameters including age, gender, and disease duration as covariates was complemented. The PBVC was adjusted using baseline NBV. Missing values were replaced by multiple imputations. Statistical significance was assumed at the p = 0.05 level. Because of the explorative nature of analyses, no adjustments for multiple comparisons were performed. Both intent-to-treat (ITT) and per-protocol (PP) analyses were carried out. The ITT group comprised 53 patients. The PP definition was regular study termination and protocol-conform study drug intake. The PP group comprised 38 patients. All calculations were performed using SAS Version 9.4 [TSIM3] Copyright 2002–2012 by SAS Institute Inc., Cary, NC, USA, IBM SPSS Statistics, Version 25, Copyright 1989, 2010 SPSS Inc., an IBM Company, Chicago, IL, USA and The R Project for Statistical Computing, Version 3.0.2 (2017-04-21).

Results

Study population

After starting recruitment, the study was terminated early. Fifty-three patients were randomized and...
Baseline characteristics of patients (ITT).

Table 1. Baseline characteristics of patients (ITT).

<table>
<thead>
<tr>
<th></th>
<th>High dose</th>
<th>Low dose</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total (n) [within group (%)]</td>
<td>28 [53]</td>
<td>25 [47]</td>
<td>&gt;0.99a</td>
</tr>
<tr>
<td>Disease course (n)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RRMS</td>
<td>26</td>
<td>25</td>
<td>0.49a</td>
</tr>
<tr>
<td>CIS</td>
<td>2</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Female [within group (%)]</td>
<td>20 [71]</td>
<td>17 [68]</td>
<td>&gt;0.99a</td>
</tr>
<tr>
<td>Mean disease duration onset to screening (months) [SE]</td>
<td>41 [2.1]</td>
<td>45 [1.8]</td>
<td>0.26b</td>
</tr>
<tr>
<td>Mean BMI at screening [SE]</td>
<td>27.2 [1.3]</td>
<td>25.5 [0.9]</td>
<td>0.61b</td>
</tr>
<tr>
<td>Median EDSS [range]</td>
<td>2.0 [5.0]</td>
<td>2.5 [6.0]</td>
<td>0.18b</td>
</tr>
<tr>
<td>Mean 25(OH)vitamin D serum level (ng/ml) [SE]</td>
<td>18.8 [1.9]</td>
<td>17.8 [1.7]</td>
<td>0.99b</td>
</tr>
<tr>
<td>Mean T2w lesion count (n) [SE]</td>
<td>52.6 [6.7]</td>
<td>76.1 [10.7]</td>
<td>0.08b</td>
</tr>
<tr>
<td>Mean T2w lesion volume (ml) [SE]</td>
<td>4.6 [0.9]</td>
<td>10.4 [1.9]</td>
<td>0.01b</td>
</tr>
<tr>
<td>Mean brain parenchymal fraction (ml) [SE]</td>
<td>1163.1 [25.8]</td>
<td>1121.3 [18.1]</td>
<td>0.4b</td>
</tr>
<tr>
<td>Mean thalamus volume (ml) [SE]</td>
<td>15.5 [0.4]</td>
<td>14.4 [0.4]</td>
<td>0.05b</td>
</tr>
<tr>
<td>Total gd+ lesions (n)</td>
<td>4</td>
<td>2</td>
<td>&gt;0.99b</td>
</tr>
</tbody>
</table>

BMI: body mass index; CIS: clinically isolated syndrome; EDSS: Expanded Disability Status Scale; gd+: gadolinium enhancing lesions; ITT: intention to treat; MSFC: Multiple Sclerosis Functional Composite; n: number; RRMS: relapsing remitting MS; SE: standard error.

aexact Chi-Square tests; bexact Mann–Whitney U test

Vitamin D levels, MRI and clinical endpoints

Important explorative clinical and paraclinical outcome parameters (univariate, ITT population) are presented in Table 2. While on study medication, 25(OH)vitamin D levels increased from 19 (7–53) ng/ml at baseline to 62 (52–80) ng/ml at month 18 in the low-dose group and from 17 (4–35) to 23 (18–27) ng/ml in the low-dose group (p < 0.001).

The primary endpoint, number of new T2w hyperintense lesions on brain MRI at month 18 compared with baseline, was not different between arms, as were T2w lesion counts (p = 0.15) when including baseline, age, gender, and disease duration as covariates in a MANCOVA for longitudinal data. Similarly, T2 lesion volume at month 18 compared with baseline was not different between arms (p = 0.98). A numerical but not significant (p = 0.09) higher number of cumulative new gd+ lesions in the low-dose arm was driven by only four patients. MANCOVA with baseline as covariate showed no differences in the thalamic volume (p = 0.95), PBVC (p = 0.84), and the brain parenchymal fraction (p = 0.43) (Table 2). Disease progression measured by median EDSS (p = 0.64) and mean MSFC z-score (p = 0.31) in a MANCOVA with baseline as covariate was not different in both arms (Table 2).

Safety endpoints

The numbers and severity of adverse events did not differ between groups (p = 0.5, exact Chi-Square test). In total, 41 adverse events were recorded; of these 23 were in the high-dose and 18 in the low-dose arm. The most often recorded adverse events were respiratory infections (n = 25) and musculoskeletal complaints (n = 15), but without differences between study arms. The majority of adverse events were mild or moderate and not considered related to the study medication. No serious adverse events and no vitamin D-related toxicity or safety issues were recorded.
**Figure 1.** The CONSORT flow diagram.
Figure 1 shows the numbers of screened, randomized, and analysed patients in both treatment arms. The reasons for exclusion from randomization, drop out or exclusion from analysis are displayed.

ITT: intention to treat; PP: per protocol; n: number.

**Table 2.** Main clinical and MRI outcome parameters after 18 months and the changes from baseline (ITT).

<table>
<thead>
<tr>
<th></th>
<th>High dose</th>
<th>Low dose</th>
<th>p-value</th>
<th>Change from baseline</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean 25OH vitamin D serum level (ng/ml) [SE]</td>
<td>65.0 [5.5]</td>
<td>22.3 [1.4]</td>
<td>&lt;0.001&lt;sup&gt;c&lt;/sup&gt;</td>
<td>45.9 [5.4]</td>
<td>5.9 [2.3]</td>
</tr>
<tr>
<td>Cumulative number of relapses (n)</td>
<td>5</td>
<td>7</td>
<td>0.6&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median EDSS [range]</td>
<td>2.0 [3.5]</td>
<td>2.0 [5.5]</td>
<td>0.26&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0 [4]</td>
<td>0 [2.5]</td>
</tr>
<tr>
<td>Mean MSFC z-score change [SE]</td>
<td>-0.24 [0.14]</td>
<td>-0.44 [0.28]</td>
<td>0.13&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.12 [0.07]</td>
<td>-0.09 [0.16]</td>
</tr>
<tr>
<td>Mean T2w lesion count (n) [SE]</td>
<td>53.4 [7.3]</td>
<td>84.1 [13.5]</td>
<td>0.05&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.3 [0.1]</td>
<td>2.1 [1.4]</td>
</tr>
<tr>
<td>Mean T2w lesion volume (ml) [SE]</td>
<td>3.6 [0.5]</td>
<td>11.9 [2.2]</td>
<td>&lt;0.001&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.1 [0.1]</td>
<td>0.2 [0.3]</td>
</tr>
<tr>
<td>Mean brain parenchymal fraction (ml) [SE]</td>
<td>1167.7 [25.9]</td>
<td>1126.6 [20.7]</td>
<td>0.97&lt;sup&gt;a&lt;/sup&gt;</td>
<td>-9.3 [3.7]</td>
<td>-7.3 [2.6]</td>
</tr>
<tr>
<td>Mean brain volume changes (%) [SE]</td>
<td>-0.61 [0.12]</td>
<td>-0.52 [0.10]</td>
<td>0.98&lt;sup&gt;a&lt;/sup&gt;</td>
<td>na</td>
<td>na</td>
</tr>
<tr>
<td>Mean thalamus volume (ml) [SE]</td>
<td>15.7 [0.3]</td>
<td>14.4 [0.5]</td>
<td>0.048&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.41 [0.6]</td>
<td>-0.12 [0.1]</td>
</tr>
<tr>
<td>Cumulative number of new gd+ lesions (V0–V6) (-)</td>
<td>2</td>
<td>14</td>
<td>0.09&lt;sup&gt;a&lt;/sup&gt;</td>
<td>na</td>
<td>na</td>
</tr>
</tbody>
</table>

EDSS: Expanded Disability Status Scale; gd+: gadolinium enhancing lesions; ITT: intention to treat; MSFC: Multiple Sclerosis Functional Composite; n: number; SE: standard error; na: not applicable.
<sup>a</sup>exact Mann–Whitney U tests; <sup>b</sup>Kaplan–Meier analysis, log-rank tests and multiple Cox regression; <sup>c</sup>multivariate analysis of covariance (MANCOVA) when including baseline values, age, gender and disease duration as covariates in a longitudinal analysis.
Discussion

Upon 18 months of continuous vitamin D intake we did not detect a different effect of high-dose versus low-dose supplementation of MS patients on important clinical and MRI parameters. We also did not record any relevant vitamin D-related adverse events. The study was carefully planned to account for heterogeneity of patients and variability of data (i.e. stratification, monocentric standardized MRI for all patients, standardization of evaluation, separation of treating and evaluating personnel). We chose an actively controlled paradigm because we expected a high incidence of vitamin D deficiency, and it was considered ethically not justifiable to deprive participants of any vitamin D supplementation. In fact, at baseline, the mean serum vitamin D levels were < 20 ng/ml in both arms (Table 1). The higher dose (corresponding to 10,200 IU per day) was chosen empirically under consideration of safety aspects and was higher than the doses given in the majority of previous trials, but was lower than the treatment arm dose (14,007 IU) in the SOLAR trial. The lower dose was derived from recommendations of the German Nutrition Society at the time when the study was designed, which were 200 IU per day. One might speculate that the difference between both doses was too low, or that already the low dose was immunologically active. But the observation of a rather small increase in the mean vitamin D serum level in the low-dose arm (from 18 to 22 ng/ml) in contrast to the pronounced rise in the high-dose arm argues against an insufficient difference between both doses. The rather mild disease activity in the entire cohort, which might be explained by the disease-modifying pretreatment with IFN β and the add-on treatment paradigm to an approved disease-modifying drug, may also have contributed to the negative results. But from an ethical point of view it was considered unacceptable to ‘treat’ MS patients with active disease solely by vitamin D and to withhold an approved treatment.

Finally, the rather small sample size may well account for the lack of differences in both study arms during therapy with a drug that significantly reduces relapses and MRI activity. The targeted sample size of 80 participants was mainly based on feasibility and did not result from a statistically supported sample size calculation. Nevertheless, unforeseen recruitment difficulties such as the contemporaneous approval of oral MS drugs and a highly comparative environment with a large number of recruiting clinical trials may explain why, despite a fairly long recruitment period of 45 months, only 53 patients were randomized. A substantial dropout rate further reduced the power to detect primary endpoint changes with \( \alpha = 0.05 \) (two-sided) to 11% (PP population) and 4% (ITT population), respectively. However, these figures provide important information for future trials on vitamin D supplementation in MS. To detect a difference in the T2w lesion count with a power of 80% and a type 1 error (\( \alpha \)) of 0.05 (two-sided) 252 participants (PP population) or 613 (ITT population) per arm would be necessary. The corresponding numbers for the hypothetical endpoint T2w lesion volume would be 3147 or 33,202 participants per arm. These power considerations suggest the in principle feasibility to demonstrate (or disprove) a disease-modifying effect of vitamin D supplementation in MS patients, at least for the endpoint T2w lesion count.

Tolerability and safety even of the high-dose were excellent in this study. The number of adverse events was similar in both arms, and adverse events recorded were not considered related to vitamin D intake. Most importantly, we did not observe hypercalcaemia. Thus, the EVIDIMS trial suggests that a mean daily intake of about 10,000 IU cholecalciferol is safe, at least under the premises of the cohort studied, i.e. in patients aged 25 to 62 without any relevant kidney dysfunction and exclusion of certain co-medications.

In conclusion, the results of the EVIDIMS study neither support nor disprove a beneficial effect of vitamin D supplementation in MS. However, the explorative trial results provide valuable information for sample size calculation and feasibility of future confirmatory trials on this topic. The data further support the safety and tolerability of a mean daily dose of 10,200 IU cholecalciferol in patients with MS.

Author Contributions

All authors approved the final version of the manuscript. JD designed the trial, drafted the study protocol, treated patients, generated data, and drafted the manuscript. PBK collected and processed data, performed statistical analyses, and drafted the manuscript. KDW was the responsible biometrician, performed the statistical analyses, and drafted the manuscript. EB, FH, JF, BB, OH and KA were principal investigators of a study site, treated patients and generated data. JW was in charge of acquisition and processing of MRI data. SKP supported the statistical analyses. JBS treated patients, generated data and coordinated as well as supervised the process of data collection. AB coordinated and supervised the process of data collection,
compilation and evaluation and drafted the manuscript. FP designed the trial, drafted the study protocol, generated data, and drafted the manuscript.

Conflict of Interests
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Supplemental Material
Supplemental material for this article is available online.

References

