Vitamin D supplementation in chronic spinal cord injury (VitD-SCI): study protocol for a randomised controlled trial

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ABSTRACT

Introduction Vitamin D insufficiency, a vitamin D status or serum 25(OH)D concentration of ≤75 nmol/L, is highly prevalent in individuals with a spinal cord injury (SCI). Vitamin D is important for the functioning of the musculoskeletal, immune and respiratory systems, which are relevant determinants of secondary health conditions in SCI. An insufficiency should be treated with vitamin D supplementation. However, there is a lack of evidence regarding the optimal dosage and duration of vitamin D supplementation for individualised and long-term management of the vitamin D status in the context of SCI. This paper presents the protocol for the vitamin D supplementation in chronic spinal cord injury (VitD-SCI) trial that aims to investigate the effect of a 12-month intake of vitamin D supplementation on vitamin D status as well as on secondary parameters among individuals with a chronic SCI.

Methods and analyses The VitD-SCI trial is a randomised, placebo-controlled, double-blinded, parallel-group superiority trial, conducted at the Swiss Paraplegic Centre. A total of 45 participants living with an SCI for at least 3 years (chronic SCI) and a vitamin D insufficiency at the first study visit, will be randomly assigned to one of three intervention groups. Participants receive either a monthly dosage of 24 000 IU or 48 000 IU vitamin D or a placebo for 12 months. Measurements taking place every 3 months include the assessment of vitamin D status (primary outcome) as well as bone mineral density, handgrip strength, fatigue, mood, pain and pressure injuries (secondary outcomes). Safety and tolerance of vitamin D supplementation will also be evaluated.

Ethics and dissemination The Swiss Ethics Committee for Northwest/Central Switzerland (EKNZ, 2020–01493) and the Swiss Agency for Therapeutic Products (Swissmedic, 2020DR3150) approved this study. Findings will be disseminated through peer-reviewed publications. Trial registration numbers NCT04652544 and SNCTP00004032.

INTRODUCTION

A suboptimal vitamin D status is prevalent in up to 93% of individuals with spinal cord injury (SCI) which is excessive compared with up to 40% among the general population. This excessive prevalence can be explained by the adverse impact of SCI on physiological functioning, including altered metabolism and gastrointestinal functioning, and lifestyle behaviours, such as reduced physical exercise, outdoor activity and sun exposure. Vitamin D is important for the optimal functioning of the musculoskeletal and respiratory system as well as the regulation of innate and adaptive immune responses. Particularly in the context of SCI, a lower vitamin D status has been associated with an elevated risk of respiratory illness, pressure injuries and depression, as well as poor physical function and bone mineral density. Therefore, the prevention of a vitamin D insufficiency, commonly defined as a vitamin D status of 75 nmol/L or less, is of great concern regarding secondary health conditions in SCI.
Vitamin D supplementation is a promising intervention to reduce and prevent vitamin D insufficiency as well as secondary complications, yet the provision of such clinical guidelines in the context of SCI is currently hampered by a lack of evidence regarding the efficacy, as well as the optimal dosage and duration of the supplementation. For the general non-SCI population (adults aged 19–70 years) a dosage of 600 IU – 800 IU/day has been recommended, although other experts consider this dosage as suboptimal. Among individuals with chronic SCI, a dosage of 800 IU/day for 12 months failed to raise vitamin D status to sufficient levels. Higher dosages of 2000 IU/day or 6000 IU/day during 3 months appeared more successful, but the respective studies did not investigate the long-term efficacy regarding vitamin D status or the lasting consequences for musculoskeletal and subjective health parameters. Though recommended, vitamin D status is not regularly monitored among the chronic SCI population.

To close this evidence gap, we present the protocol for the vitamin D supplementation in chronic spinal cord injury (VitD-SCI) trial: a placebo-controlled randomised controlled trial that investigates the efficacy of a moderate (24 000 IU/month) and a high (48 000 IU/month) dosage of vitamin D supplementation for 12 months among individuals living with an SCI for at least 3 years (chronic SCI). Besides vitamin D status (primary objective), further assessments include bone mineral density, fatigue, pain, mood, performance of daily activities and handgrip strength (secondary objectives). We hypothesise a dose-response effect of vitamin D supplementation on vitamin D status and anticipate that the evidence acquired during this study will effectively inform community-based policy regarding vitamin D supplementation in chronic SCI.

METHODS AND ANALYSES
Study design and setting
The VitD-SCI trial is a placebo-controlled randomised double-blinded superiority study, evaluating the effect of 12 months of vitamin D supplementation on vitamin D status and several secondary outcomes among individuals with a chronic SCI (figure 1). This protocol was designed according to the recommendations of the Standard Protocol Items: Recommendations for Interventional Trials 2013 Statement (online supplemental file 1). The study is planned to run from May 2021 to May 2023. Study visits take place at the Swiss Paraplegic Centre and participants will receive compensation for travel expenses. The VitD-SCI trial is a nested project of the population-based Swiss Spinal Cord Injury (SwiSCI) cohort study.

Patient and public involvement statement
Patient and public organisations (eg, SwiSCI), as well as clinicians from several specialised SCI centres, have been involved in the design of the study protocol. Study procedures, including the intervention and assessments, have been discussed and tested together with individuals with SCI.

Participants
The inception cohort of the SwiSCI cohort study (2013–present) provides the sampling frame for participant recruitment for the VitD-SCI trial. Eligible are individuals who provided consent to be contacted for future scientific studies and who fulfil the key VitD-SCI trial inclusion criteria, including a vertebral lesion level of C4 or below and time since SCI of at least 3 years. At present (May 2021), SwiSCI data indicate a sample of over 300 eligible individuals. Among individuals interested in participating, the remaining eligibility criteria (table 1) will be checked. Women of childbearing age must show a negative pregnancy test at the start of the experimental supplementation and must use adequate contraception during the 12-month treatment phase of the study. Participants are not allowed to take additional vitamin D supplements (>100 IU/day) or travel to countries with increased sun exposure (below the 37th parallel north) during the study. No further eligibility criteria, such as the completeness or the lesion level, were set, to allow a generalisation of the study results as well as to increase the feasibility. The recurrent assessment of contextual parameters for use as time-updated covariates in the data analysis will further support unbiased inference of the temporal efficacy of vitamin D supplementation. Informed consent will be obtained from all participants by qualified study employees (online supplemental file 2). A blood sample will be taken to determine vitamin D status at the first visit (Month 0).

Intervention
The trial consists of a three-armed intervention that applies a moderate or generally recommended dosage (24 000 IU vitamin D/month), and a high dosage (48 000 IU vitamin D/month) for comparison to placebo. Both dosages are prescribed in clinical practice and are within safe tolerable upper intake levels of 10 000 IU/day. Previous studies among individuals with SCI implementing even higher dosages of vitamin D supplementation (up to 50 000 IU/week) did not report any safety issues. The present study will further use a biweekly supplementation procedure, which showed superior adherence and accompanying efficacy over daily administration in achieving a sufficient vitamin D status. The biweekly supplementation schedule is supported by vitamin D pharmacokinetics, which indicates a half-life of serum 25(OH)D of up to 3 weeks.

As vitamin D supplement, the commercially available ViDe 3 Monthly Dose (Dr. Wild & Co. AG, Muttenz, Switzerland) will be used. One vial (5mL) contains an ethanol solution (65% alcohol by volume) with 600µg cholecalciferol, which corresponds to 24 000 IU vitamin D. The similar ethanol solution without cholecalciferol will be used as a placebo. Both the vitamin D supplement as well as the placebo will be filled in identical vials. An
independent and specialised pharmacy will label the vials with a number that is indicative of the order of intake but indiscriminate regarding content, making the vials indistinguishable in odour, taste and visual appearance. The first vial is to be taken within 7 days after the first study visit. Participants are instructed to ingest one single vial every 2 weeks by emptying the vial in a glass of water and immediately drinking the solution. Participants of the high dosage group will take one vial with 24 000 IU vitamin D₃ every 2 weeks. Similarly, participants of the placebo group will take one vial with placebo every 2 weeks. Participants of the moderate dosage group will alternately

Figure 1  Schematic overview of the study flow. The timeline of the study including the assessments is displayed starting from the screening (top of the figure) to the last study visit (bottom of the figure). (A) Study flow. (B) Legend. DXA, X-ray densitometry.
take one vial with 24 000IU vitamin D$_3$ or one vial with placebo every 2 weeks. Thus, an identical number of vials is assured across all intervention groups. The adherence of participants to the protocol cannot be objectively assessed, because the supplement intake occurs at home. To promote compliance with the study protocol, participants will receive a reminder every 2 weeks to take the vials. Participants will also be requested to keep track of the intake of each vial in a study diary and return all (un) used vials to the study centre.

Randomisation and blinding
Participants will be randomly allocated on a 1:1:1 ratio into one of the three intervention groups, according to a permuted block randomisation list with randomly varying block sizes (three or six) created using the module ‘ralloc’ (V.3.7.6) within Stata ((computer program), V.16, StataCorp, 2019). The allocation sequence will be securely stored and only accessible to an employee, who is not directly involved in the study. Participants and all study employees directly involved in the recruitment and measurements will be blinded to the assignment of the intervention until the end of the study (‘last participant out’). Parameters, such as blood and bone mineral density parameters, which may reveal the allocation across the intervention groups, will only be available to employees who have no direct involvement in the study.

Outcomes and assessments
Assessment periods and details are specified in table 2. Assessments are performed during screening or each of the 3 monthly visits, except for bone mineral density, which is measured at 0 and 12 months only to reduce radiation exposure. The primary outcome is the differential change in vitamin D status from 0 to 12 months. Transitional vitamin D status will also be evaluated every 3 months. The recurrent assessment of contributory parameters, including sun exposure and secondary health conditions, provide time-updated information to statistically control for within-person and between-person variation in investigating the efficacy of vitamin D supplementation on vitamin D dynamics. To evaluate the secondary effects of vitamin D supplementation on the musculoskeletal system, differential changes in the following secondary outcomes will be assessed: relevant blood parameters (calcium, parathyroid hormone, ionised calcium, phosphate, cystatin, estimated glomerular filtration rate, testosterone), bone mineral density, functional independence, pressure injuries, handgrip strength and falls. Differential changes in fatigue, mood and pain will provide insight into the effect of vitamin D supplementation on subjective parameters. Individual parameters, including demographics, SCI characteristics, skin phototype and sun exposure will also be collected. Furthermore, the occurrence of any new illness, injury or disease together with newly taken medication and supplements will be assessed at each visit. Since nutritional

Table 1  Eligibility criteria

| Inclusion criteria | Exclusion criteria |
|--------------------|--------------------|
| ► Informed Consent to the study as documented by a signature. | ► Contraindications to the investigational product, including known hypersensitivity or allergy to the cholecalciferol or alcohol solution. |
| ► Chronic (>3 years) traumatic or non-traumatic spinal cord injury a vertebral lesion level of C4 or below. | ► Clinically relevant disorders, including renal dysfunction, hepatic dysfunction, cardiovascular disease, lung disease, diabetes, blood disease, parathyroid disease, cancer, depression, alcohol abuse and/or the intake of significant concomitant medication (including osteoporosis treatment and benzothiadiazide derivatives). This will be assessed on an individual basis. |
| ► Between 18–60 years old. | ► Grade 3 or 4 pressure injuries. |
| ► Wheelchair dependency during activities of daily living, defined by a score of 0–2 in the Spinal Cord Independence Measure, subsection ‘Mobility in the house’. | ► Women who are pregnant or planning to become pregnant during the study period. |
| ► An insufficient vitamin D status (≤75 nmol/L) at the first visit. | ► Women who are breast feeding. |

Wheelchair dependency during activities of daily living, defined by a score of 0–2 in the Spinal Cord Independence Measure, subsection ‘Mobility in the house’.

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Table 2  Tools, parameters and timing of the study measurements

| Assessment tool | Parameter | Assessment periods |
|-----------------|-----------|--------------------|
| **Differential changes in blood parameters (sample taken from the antecubital vein)** | | |
| Electrochemiluminescence immunoassay | 25(OH)D (nmol/L). | 0M, 3M, 6M, 12M |
| Photometry—NM-BAPTA | Calcium (mmol/L). | 0M, 3M, 6M, 12M |
| Potentiometry with ion-selective electrodes | Ionised calcium (mmol/L). | 0M, 3M, 6M, 12M |
| Chemiluminescence immunoassay (CLIA) | Parathyroid hormone (ng/L). | 0M, 3M, 6M, 12M |
| Photometry—molybdate ultraviolet | Phosphate (mmol/L). | 0M, 3M, 6M, 12M |
| Photometry—particle-enhanced turbidimetric immunoassay | Cystatin C (mg/L). | 0M, 3M, 6M, 12M |
| Estimated from cystatin C following CKD-EPI and Grubb | Estimated glomerular filtration rate (mL/min). | 0M, 3M, 6M, 12M |
| CLIA | Testosterone (ng/dL). | 0M, 3M, 6M, 12M |
| **Differential changes in personal characteristics** | | |
| Body mass index | From weight and height (kg/m²). | 0M, 3M, 6M, 12M |
| Sun exposure based on Hanwell et al. | Time spent outdoors (min) with level of exposed skin.* | 0M, 3M, 6M, 12M |
| Physical activity | Sport per week (hours and frequency). | 0M, 3M, 6M, 12M |
| Medication and supplements | Sort and dosage (supplements only). | 0M, 3M, 6M, 12M |
| Further illnesses | Incidence since last measurement. | 0M, 3M, 6M, 12M |
| Skin phototype on the posterior forearm based on Fitzpatrick | Six categories ranging from light to very dark skin types. | Screening |
| SCI characteristics | Time since SCI (years), neurological level of injury, the degree of impairment following the American Spinal Injury Association Impairment Scale. | Screening |
| **Differential changes in bone mineral density** | | |
| X-ray densitometry, Lunar iDXA Serie with enCORE V.17 Software | T-scores for the forearm (radius), hip (femoral neck) and knee (distal femur and proximal tibia). | 0M, 12M |
| **Differential changes in functional independence** | | |
| Spinal Cord Independence Measure III | Self-care subscore (0–20), respiration and sphincter management subscore (0–40), indoor mobility subscore (0–10), outdoor mobility subscore (0–30), total score (0–100). | 0M, 3M, 6M, 12M |
| **Differential changes in urinary tract infections** | | |
| Exact questions of the first-wave questionnaire of the Swiss Spinal Cord Injury (SwiSCI) survey | Incidence since the last measurement, medical care required. | 0M, 3M, 6M, 12M |
| **Differential changes in pressure injuries** | | |
| Exact questions of the first-wave questionnaire of the SwiSCI survey | Localisation and grade following the European and United States National Pressure Ulcer Advisory panels classification. | 0M, 3M, 6M, 12M |
| **Differential changes in pain** | | |
| Exact questions of the first-wave and second-wave questionnaire of the SwiSCI survey, based on the International SCI Pain Basic Data Set and as well as the SCI Secondary Conditions Scale | The occurrence, kind, location and influence of pain during the last week as well as chronic pain (>3 months). | 0M, 3M, 6M, 12M |
| **Differential changes in strength** | | |
| Jamar Smart Hand Dynamometer (Patterson Medical, Warrenville, Illinois) | Three measurements of the left and right hand (mean kg). | 0M, 3M, 6M, 12M |
| **Differential changes in mood** | | |
| Hospital Anxiety and Depression Scale | Total score (14–56). | 0M, 3M, 6M, 12M |
| **Differential changes in fatigue** | | |
| Fatigue Severity Scale | Total score (9–63). | 0M, 3M, 6M, 12M |
| **Differential changes in falls** | | |
| Occurrence | Incidence since last measurement, reason daily activity, sport or accident) and medical care required. | 0M, 3M, 6M, 12M |

Continued
intake of vitamin D only accounts for a limited intake and vitamin D fortification of nutrition in Switzerland is rare, no dietary assessments are planned, which precludes scaling of vitamin D efficacy to between-person and within-person variation in nutrition intake. Due to feasibility reasons, no direct measurements of respiratory or immune function will be assessed.

Serious adverse events will be documented throughout the participant’s enrolment to evaluate the safety of the supplementation.

Sample size
We defined the minimal sample size as the sample size needed to detect a statistically meaningful difference in the time course of the primary outcome vitamin D status across the three intervention groups. For this, we estimated the sample size needed to detect an interaction between treatment and time across the three intervention groups using the power analysis package for repeated measures in Stata software, V.16.0 (StataCorp, 2019). The expected mean vitamin D status at each time point was calculated for each intervention group using the generalised pharmacokinetic curve equation of Heaney, et al. which we parametrised for the purpose of this study using the available data for vitamin D supplementation in chronic SCI.

Assuming a significance level of 5% and a power of 0.80, minimal sample sizes were calculated for a range of between-group variances (ie, 1000–3500, with 500 incremental steps) and correlations between repeated measures (ie, 0.1–0.9, with 0.1 incremental steps; online supplemental file 3). Based on this power analysis for sample size, we conservatively plan for 45 participants with a full data set, implying 15 participants in each of the three intervention groups. In the case of dropouts or a large amount of missing data for a given participant, recruitment will be continued until the minimal sample size of 15 participants per intervention group is effectively achieved. Sample size calculation did not take secondary outcome parameters into account. Since long-term studies among the chronic SCI population are rare, this study may inform the minimal sample size needed for future studies targeting the dynamics of secondary outcome parameters in response to vitamin D supplementation.

Data management and analyses
Data management
The web browser-based system secuTrial (interActive Systems, Berlin, Germany), which fulfils the Good Clinical Practice requirements, will be used for data capturing and management. This database allows for individualised accounts with predefined roles for each study employee and will be hosted by a study-independent data manager. Built-in quality control mechanisms and notification of missing data will reduce errors during data entry. Participants will be assigned a unique identification code and no directly identifiable information is stored in the database.

Analyses of primary and secondary outcomes
An intention-to-treat analysis will be performed to prevent potential bias resulting from the distortion of baseline equivalence after randomisation due to the withdrawal of participants, non-adherence to the study protocol, or an unwarranted level of missing data. Participants will only be included in the analyses for which they have data available. Protocol non-adherence, early withdrawals and loss to follow-up will be characterised and examined. Sensitivity analyses using different populations or participant groupings may be used to examine the robustness of the estimator generated by the intention-to-treat analysis. Any sensitivity analyses added after the breaking of the blind (post-hoc sensitivity analyses), including per-protocol analysis that will contain any participants recruited as a replacement for withdrawals or dropouts, will be clearly identified as such in the trial reporting.

Basic univariable statistical analysis techniques will be used to describe the study population as well as the primary and secondary outcomes at the different measurement points. To evaluate the longitudinal variation in the primary outcome and secondary outcomes, multilevel mixed-effects models that appropriately account for the within-individual and between-individual sources of variance in outcome variation will be used. In these models, participants are treated as random effects, while the treatment group is used as a fixed parameter. The choice of

Table 2

| Assessment tool Parameter Assessment periods |
|-------------------------------------------|------------------------------------------|
| Safety of the investigational medical products |                                |
| Serious adverse events The occurrence of life-threatening medical complications, the requirement of hospitalisation. | Any time |
| Side effects, tolerability and adherence Number of and reason for missed intake, side effects. | 0M, 3M, 6M, 12M |

Assessment periods are referenced to the date of the first visit (0M) and include screening (screening; max 30 days before 0M), 3-month follow-up (3M; 90±7 days), 6-month follow-up (6M; 180±7 days) and 12-month follow-up (12M; 365±7 days). Serious adverse events are continuously evaluated.

*Level of exposed skin: (1) hands and face, (2) hands, face and arms, (3) hands, face and legs, (4) entire body (bathing suit).
CKD-EPI, Chronic Kidney Disease Epidemiology Collaboration; NM-BAPTA, 5-nitro-5′-methyl-(1,2-bis(o-aminophenoxy)ethane-N,N,N′,N′-tetraacetic acid); SCI, spinal cord injury.
regression model will depend on the error distribution function of the respective outcome parameter. Time-updated covariates will also be used in regression modelling, for instance, to evaluate the impact of time-varying vitamin D status on the secondary outcomes. Time lags in time-updated covariates will also be evaluated to further detail temporal associations. Non-linear regression modelling will be used to evaluate pharmacokinetic curves for each of the intervention groups as described above. Thus, estimates are derived for the parameters $a$ (ie, change is constant at equilibrium), and $k$ (ie, the rate constant for the proportion of the total mass of 25(OH)D used or metabolised per day) for mutual comparison as well as with estimates from other populations (other SCI populations, general population or other health-condition groups).

No interim analyses are planned. If severe clinical deterioration is detected in more than one participant, or on the recommendation of the monitoring committee following a serious adverse event, the study will be suspended until a comprehensive safety review has been completed. If the trial is suspended or halted, an interim analysis will be performed.

**Handling of missing data and dropouts**

When appropriate, multiple imputation will be used to account for missing data. In case the number of dropouts is substantial and systematically related to baseline values of the primary outcome variable, logistic regression modelling will be used to derive inverse probability weights for use in regression modelling to statistically account for potential longitudinal selection bias. Participants who are withdrawn or dropout will be replaced. Sensitivity analysis will be used to address the robustness of the estimators to changes in the assumptions underlying the chosen imputation strategy.

**Quality assurance and safety provisions**

**Minimising bias and contamination**

Trained study employees will execute all of the measurements following a study manual. Measurement devices will be appropriately serviced and calibrated throughout the study. Questionnaires that have been validated among or adapted for the SCI population will be used. The collection of several possible confounders, including sun exposure, season and SCI level, allows for the correction of these factors. Participants will be asked to express their belief regarding the assigned intervention every 3 months, which will be used to compute a measure for the extent of successful blinding of the trial.

**Potential risks**

There are no guidelines regarding upper limits of serum 25(OH)D concentration among the SCI population, although vitamin D toxicity has been reported with levels exceeding 375–750 nmol/L among the general population. Side effects of vitamin D supplementation, resulting from an overdose, are not expected with the selected dosages in this study. Nevertheless, participants are asked about potential side effects and tolerance of the supplement, and relevant blood parameters will be monitored. Due to the 65% volume alcohol solution, the intake of the supplement and/or placebo could have a minor influence on the ability to drive or operate machines, but similar solutions have been proven harmless. The sum of the radiation exposure for the two bone mineral density measurements in this study is less than the average yearly radiation exposure due to radon in Switzerland and falls well within the dose guidance value for Swiss research projects. Serious adverse events will be documented and reported to the local authorities for the entire study duration, which encompasses the signing of the informed consent until the completion of the last protocol-specific procedure including a safety follow-up period. A risk-adapted monitoring will be applied and executed by the clinical trial unit of the study centre.

**Withdrawal criteria**

Participants can withdraw from the study at any time, without providing reasoning. Participants might be redrawn from the study in case of protocol non-adherence, the eligibility criteria are no longer fulfilled or health problems related to the intake of the vitamin D supplement or placebo, including a vitamin D intoxication (serum 25(OH)D concentration >375 nmol/L) or hypercalcaemia (ionised calcium concentration >3.5 mmol/L).

**Unblinding procedures**

The study physician together with the study employees will make study termination and unblinding decisions. Unblinding will only be allowed in the following circumstances; on study termination, in the case of premature study termination or in the case of emergencies or complications. An employee not directly involved in the measurements will perform the unblinding of individual participants during the study. Since each participant receives an individual batch of vials, revealing the blinding for one participant will not unblind the entire cohort.

**ETHICS AND DISSEMINATION**

**Ethics approval**

This study will be conducted in compliance with the current version of the Declaration of Helsinki and the International Conference on Harmonisation Good Clinical Practice guidelines, as well as all national legal and regulatory requirements. Both the Swiss Ethics Committee for Northwest/Central Switzerland (EKNZ, 2020–01493) and the Swiss Agency for Therapeutic Products (Swissmedic, 2020DR3150) have approved the study. The study has been registered at ClinicalTrials.gov (NCT04652544). After trial registration, we discovered not all of the secondary and further outcome parameters were mentioned in the registry, the missing outcome parameters were added subsequently. The regulatory
Dissemination policy
Results will be published in peer-reviewed journals and presented at scientific conferences. When desired, individual study results will be shared with the participants after the study end. Dissemination among persons with SCI in Switzerland will be achieved through the newsletters ‘Paraplegie’ and ‘Paracontact’ as well as through online media. Data and statistical code will be made available on request.

Trial status
This publication is based on version 3 of the VitD-SCI trial protocol dated 9 November 2020. The official start of recruitment was on 19 May 2021 and data collection is estimated to end on 31 May 2023. As of the time of submission, temporary interruptions due to the COVID-19 pandemic are a possibility and in case of extended disruption, a schedule adjustment may be required.

DISCUSSION
This is the first study to investigate a 12-month vitamin D supplementation among individuals with a chronic SCI. The existing studies lack the combination of long-term vitamin D supplementation with different dosages, while not only assessing vitamin D status but also secondary outcomes.

The VitD-SCI trial, for which the protocol is presented here, will provide valuable insights to optimise individual vitamin D supplementation. If a positive relation between a sufficient vitamin D status and secondary outcomes can be established, this could lead the way for standardised clinical recommendations for long-term management of vitamin D status and vitamin D supplementation among the chronic SCI population.

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