A pilot study assessing the effect of prolonged administration of high daily doses of vitamin D on the clinical course of vitiligo and psoriasis

Danilo C. Finamor,1 Rita Sinigaglia-Coimbra,1 Luiz C.M. Neves,2 Marcia Gutierrez,3 Jeferson J. Silva,1 Lucas D. Torres,1 Fernanda Surano,1 Domingos J. Neto,3 Neil F. Novo,6 Yara Juliano,6 Antonio C. Lopes4 and Cicero Gall Coimbra1,*

1Laboratório de Fisiopatologia Clínica e Experimental; Universidade Federal de São Paulo; São Paulo, Brazil; 2Instituto de Ciências da Saúde; Universidade Paulista; São Paulo, Brazil; 3Farmácia Sensitiva; São Paulo, Brazil; 4Disciplina de Clínica Médica; Universidade Federal de São Paulo; São Paulo, Brazil; 5Hospital Heliópolis; São Paulo, Brazil; 6Disciplina de Cirurgia Plástica; Universidade Federal de São Paulo; São Paulo, Brazil

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Abbreviations: BMI, body mass index; CIs, confidence intervals; CYP, cytochrome P450 superfamily; CYP27B1, 25-hydroxyvitamin D-1 alpha-hydroxylase; DBP, vitamin D-binding protein; VDR, vitamin D receptor; FGF23, fibroblast growth factor 23; HPLC, high performance liquid chromatography; IOM, Institute of Medicine; IS, internal standard; K m, Michaelis–Menten constant; PASI, psoriasis area and severity index; PTH, parathormone; RDA, recommended dietary allowance; Treg, regulatory T cells; VDR, vitamin D receptor; V max, maximal velocity of enzymatic reaction; UV, ultraviolet; UVA, ultraviolet A

Autoimmunity has been associated with vitamin D deficiency and resistance, with gene polymorphisms related to vitamin D metabolism frequently described in affected patients. High doses of vitamin D3 may conceivably compensate for inherited resistance to its biological effects. This study aimed to assess the efficacy and safety of prolonged high-dose vitamin D3 treatment of patients with psoriasis and vitiligo. Nine patients with psoriasis and 16 patients with vitiligo received vitamin D3 35,000 IU once daily for six months in association with a low-calcium diet (avoiding dairy products and calcium-enriched foods like oat, rice or soya “milk”) and hydration (minimum 2.5 L daily). All psoriasis patients were scored according to “Psoriasis Area and Severity Index” (PASI) at baseline and after treatment. Evaluation of clinical response of vitiligo patients required a quartile grading scale. All patients presented low vitamin D status (serum 25(OH)D3 ≤ 30 ng/mL) at baseline. After treatment 25(OH)D3 levels significantly increased (from 14.9 ± 7.4 to 106.3 ± 31.9 ng/mL and from 18.4 ± 8.9 to 132.5 ± 37.0 ng/mL) and PTH levels significantly decreased (from 57.8 ± 16.7 to 28.9 ± 8.2 pg/mL and from 55.3 ± 25.0 to 25.4 ± 10.7 pg/mL) in patients with psoriasis and vitiligo respectively. PTH and 25(OH)D3 serum concentrations correlated inversely. The PASI score significantly improved in all nine patients with psoriasis. Fourteen of 16 patients with vitiligo had 25–75% repigmentation. Serum urea, creatinine and calcium (total and ionized) did not change and urinary calcium excretion increased within the normal range. High-dose vitamin D3 therapy may be effective and safe for vitiligo and psoriasis patients.

Introduction

The steroid hormone known as the active form of vitamin D (“calcitriol,” “1,25-dihydroxy vitamin D” or “1,25(OH)2D”) generates a wide range of biological responses. The human genome possesses 2,776 positions occupied by the VDR; about 10% of the human genes are, therefore, directly and/or indirectly responsive to vitamin D.1 Similarly, diverse human cells (including bone, colon, breast, prostate, skin, muscle, blood vessel, brain and immune cells) express the enzyme 25-hydroxyvitamin D-1 α-hydroxylase (CYP27B1), indicating that the extra-renal intracrine and paracrine 1,25(OH)2D3 synthesis may critically affect the activities of many tissues and organs.2,4 Accordingly, cumulative data have associated low vitamin D status (as assessed by measuring “25-hydroxyvitamin D3” or “25(OH)D3”—the main circulating form of vitamin D) not only to rickets and osteoporosis, but also to an increasing number of prevalent health disorders including autoimmune, infectious and cardiovascular diseases, hypertension, diabetes and deadly cancers.2,5,7

Contrasting with the pathophysiological importance of 1,25(OH)2D3, vitamin D deficiency is a poorly recognized worldwide epidemic among both children and adults.2 Regular and moderate sun exposure (the only significant natural source of vitamin D for most subjects) is currently uncommon due to indoor lifestyle, sun avoidance and indiscriminate sunscreen use.7 Even though there is mounting evidence indicating that physiologic doses should be closer to those achieved through a few minutes of daily skin exposure to sunlight,2 the RDA upgraded (from
200 to 600 IU per day) by the Institute of Medicine in 2010 and are lower than the daily doses required to correct vitamin D deficiency in most subjects.

Higher daily doses may be particularly critical for controlling the activity of autoimmune disorders. Cumulative data over the past 30 years have established the regulatory effects of vitamin D on the activity of autoimmune disorders. Cumulative evidence also indicates that vitamin D promotes—by both direct and indirect mechanisms—regulatory T cells (Treg) that inhibit a variety of inappropriate (both innate and adaptive) immune responses underlying autoimmune disease.

Estimations of daily requirements of vitamin D3 for patients with autoimmune disorders should take account of the functional consequences of genetic polymorphisms related to vitamin D metabolism. For instance, the polymorphic changes of the enzyme CYP27B1 associated with autoimmunity predictably originate a relative resistance to vitamin D, i.e., a higher level of circulating 25(OH)D3 required to achieve normal 1,25(OH)2D3 concentrations within the immune cells. While the in vivo Km of the normal enzyme exceeds the physiologic range of 25(OH)D3 concentrations, a polymorphic variant may have a higher Km (decreased affinity for substrate) and/or a lower Vmax, requiring higher concentrations of substrate to achieve a physiologic rate of product formation (Fig. 1). Frequent genetic polymorphisms and wide (ten to a hundred fold) variability in enzyme expression of the cytochrome P450 superfamily of enzymes (CYPs) are common and may considerably modify enzyme activity, causing large interindividual variability in the rate of product formation. Therefore, a subject bearing a CYP27B1 polymorphism may be prone to develop autoimmunity for sustaining concentrations of 1,25(OH)2D3 within the immune cells that are insufficient for full intracrine and paracrine tolerogenic effects of this powerful steroid molecule. High doses of vitamin D3 leading to supraphysiologic range of circulating 25(OH)D3 may compensate for this genetic-related status of relative vitamin D deficiency, thereby establishing tolerance to auto-antigens.

Doses ranging up to 40,000 IU/day of vitamin D3 are probably safe for healthy individuals, and enzyme polymorphisms affecting vitamin D metabolism may conceivably increase tolerability in patients with autoimmune disorders. This study aimed to assess the therapeutic efficacy and safety of a daily dose of 35,000 IU of vitamin D3 administered with a low-calcium diet for 6 mo to patients with psoriasis and vitiligo.

**Results**

**Subjects.** The physical characteristics of the 25 patients who participated in this study are shown in Table 1.

**Laboratory parameters.** As shown in Table 2 and 3 and Figures 2 and 5, all patients presented low baseline levels of 25(OH)D3 (≤ 30 ng/mL or ≤ 75 nmol/L), with a mean of 14.9 ± 7.4 ng/mL in the psoriasis group and a mean of 18.4 ± 8.9 ng/mL in vitiligo patients. After 6 mo of treatment with vitamin D3 (35,000 IU per day), 25(OH)D3 levels increased significantly, to 106.3 ± 31.9 ng/mL in the psoriasis group (p < 0.0001, Wilcoxon signed rank test) and to 132.5 ± 37.0 ng/mL in vitiligo patients (p < 0.0005, Wilcoxon signed rank test).

Baseline concentrations of PTH were 57.8 ± 16.7 pg/mL in the psoriasis group and 55.3 ± 25.0 pg/mL in vitiligo patients. After 6 mo of treatment with vitamin D3 PTH levels decreased significantly to 28.9 ± 8.2 pg/mL in the psoriasis group (p = 0.0039, Wilcoxon signed rank test) and to 25.4 ± 10.7 pg/mL in vitiligo patients.
vitreous patients (p = 0.0005, Wilcoxon signed rank test) (Tables 2 and 3, Figs. 2 and 5).

Statistical analysis revealed a negative correlation between 25(OH)D3 and PTH levels (r = −0.6753, p = 0.0153 in the psoriasis group; r = −0.5091, p = 0.0015 in vitiligo patients, Pearson’s correlation). Linear regression also revealed negative correlation between these two parameters using data collected at baseline and 6-mo outcome (Figs. 2 and 5, r² and p values indicated).

The concentrations of (total and ionized) serum calcium, urea and creatinine did not differ significantly from baseline values after 6 mo of high-dose vitamin D3 treatment (Tables 2 and 3; Figs. 3 and 6: Wilcoxon signed rank test).

Urinary calcium excretion significantly increased at 6 mo of vitamin D3 treatment in both groups from 123.6 ± 60.0 mg/24 h to 226.8 ± 41.6 mg/24 h in psoriasis group (p = 0.0039, Wilcoxon signed rank test) and from 158.3 ± 73.6 mg/24 h to 230.1 ± 226.8 ± 41.6 mg/24 h in psoriasis group (p = 0.0039, Wilcoxon signed rank test), but remained within the normal range (Tables 2 and 3; Figs. 3 and 6). Statistical analysis revealed a positive correlation between urinary calcium excretion and 25(OH)D3 levels in both groups of patients (r = 0.5372, p = 0.0108 in the psoriasis group; r = −0.3972, p = 0.0122 in vitiligo patients, Pearson’s correlation). Linear regression also revealed a negative correlation between these parameters (Figs. 3 and 6, r² and p values indicated).

Clinical outcome. The clinical condition of all patients with psoriasis (n = 9) significantly improved during the treatment (Fig. 4 and 8; p = 0.0023, Wilcoxon signed rank test). Statistical analysis revealed a negative correlation between the PASI score and 25(OH)D3 levels in both groups of patients (r = −0.5614, p = 0.0011, Pearson’s correlation). Linear regression also demonstrated correlation between these parameters (Fig. 4, r² and p values indicated).

Two out of 16 vitiligo patients showed no repigmentation of the affected areas; four patients showed 1–25% repigmentation, five patients showed 26–50% repigmentation, five patients showed 51–75% repigmentation and none showed more than 75% repigmentation of the affected areas (Figs. 7 and 9).

Discussion

This is an open-label study where all patients with psoriasis or vitiligo received oral treatment with 35,000 IU of vitamin D per day for six months associated with preventive measures (partial dietary calcium restriction and a daily hydration of at least 2.5 L). The treatment provided benefit to 9 out of 9 patients with psoriasis and to 14 out of 16 patients vitiligo.

A placebo-controlled approach was avoided for ethical reasons. Cumulative data have implicated vitamin D deficiency in the pathophysiology of vitiligo and psoriasis. Vitamin D deficiency (deficient levels of a potent steroid hormone that modulates hundreds of human genes) also increases the risk of developing or aggravating a myriad of serious health disorders, including cancer or death from cardiovascular events. Administering placebo to vitamin D-deficient patients for the sake of randomization may not be ethically acceptable. Potential research participants would not take the chances of being assigned to the control group if informed of the association of vitamin D levels and disease activity. On the other hand, intentionally omitting essential information to facilitate consent to placebo treatment may imply disregard for patient autonomy and violation of the principle of beneficence on the part of the physician. Conversely, restoring the physiologic intracellular levels of a potent steroid hormone (by increasing the availability of substrate to its polymorphic activating enzyme) should be regarded as good medical practice.

Laboratory or clinical signs of toxicity (hypercalcemia, hypercalciuria or kidney dysfunction) were not observed in any of the 25 participants, including a patient with vitiligo who reached a serum concentration of 25(OH)D3 of...
and paracrine effects at sites of inflammation, where the local availability of vitamin D3 is also under strict control of other calcium- and phosphate-regulating hormones (PTH and FGF23). Conversely, the availability of calcium for bone metabolism and extra-renal control mechanisms for expression and activity of enzyme 1α-hydroxylase underlies the feedback downregulation (associated with 24-hydroxylase expression of VDR,48 as well as to body weight, body fat, age, skin color, season, latitude and sunning habits; optimal therapeutic doses of vitamin D in the current treatment with immunosuppressive drugs.

High-dose vitamin D3 supplementation to patients with autoimmune disorders is conceivably advantageous over 1,25(OH)2D3 treatment concerning lower calcemic effects and more efficient control of autoimmunity. Administration of 1,25(OH)2D3 or 1,25(OH)2D3 analogs overpasses critical regulatory mechanisms related to the calciotropic effects of vitamin D by directly stimulating intestinal VDR and calcium absorption. In contrast, administration of vitamin D3 increases circulating concentrations of 25(OH)D3, which then faces different renal and extra-renal control mechanisms for expression and activity of the enzyme 1α-hydroxylase. Renal 1α-hydroxylase undergoes feedback downregulation (associated with 24-hydroxylase upregulation) by 1,25(OH)2D3 and 1,25(OH)2D3 production is also under strict control of other calcium- and phosphate-regulating hormones (PTH and FGF23). Conversely, the availability of 25(OH)D3 to immune cells (the production of which is not tightly controlled by the liver) may be the primary determinant of the amount of 1,25(OH)2D3 produced for intracrine and paracrine effects at sites of inflammation, where the local expression of cytokines may instead facilitate the conversion of 25(OH)D3 by inducing the expression of 1α-hydroxylase.

Hypervitaminosis D is associated with upregulation of intestinal VDR and increased absorption of dietary calcium. A low calcium diet protects against vitamin D toxicity, not only by reducing the availability of calcium for gastrointestinal absorption, but also by facilitating vitamin D inactivation at sites related to calcium metabolism. Reduced intestinal calcium by dietary restriction of milk, dairy products and calcium-enriched foods (like oat, rice or soya “milk”) has contributed to minimize the calciotropic effects of high daily doses of vitamin D3 in the current study. Increased gastrointestinal absorption of calcium is partly responsible for the hypercalcemia in vitamin D intoxication and a low dietary calcium intake gradually reduces serum calcium in such patients. Preliminary data (not shown) obtained from patients treated with progressively higher doses of vitamin D3 up to 35,000 IU daily showed that the adoption of such easily understandable dietary recommendations reduced urinary calcium from borderline elevated levels (around 400 mg or 10 mmol per day, with serum calcium sustained at the upper normal range) to values within the normal range without changing vitamin D daily dose. Further restriction of dietary calcium (by also avoiding foods prepared with milk, such as smashed potatoes, bread, cakes and cookies) dropped urinary calcium to levels below the lower limit of the normal range adopted by the local laboratory (100 mg or 2.5 mmol per day) while serum calcium remained around the lower limit (8.6 mg/dL or 2.15 mmol/L).

Taken together, those data suggest that partial dietary calcium restriction efficiently prevents hypercalcemia and hypercalciuria by controlling the gastrointestinal availability of calcium under the calciotropic effects of the treatment paradigm employed in patients with psoriasis and vitiligo in this study. Potentiation of osteoclastic activity in bone by pharmacologically high levels of 25(OH)D3 competing for binding to VDR is a recognized mechanism of hypercalciemia in vitamin D intoxication. The high daily doses of vitamin D3 administered for 6 mo in this study, however, do not seem to have significantly enhanced osteoclastic activity in patients with psoriasis and vitiligo, since partial restriction of dietary calcium efficiently prevented hypercalcemia and hypercalciuria.

Compared with this study, longer-term follow up research should look for yet unknown potential side-effects related to high-dose vitamin D3 treatment of a larger sample of subjects with autoimmune disorders. Future studies should also aim to develop a method for setting the particular daily dose that compensates for genetic polymorphisms and other distinctive features, providing the highest therapeutic effect against autoimmunity without causing side effects like hypercalcemia/hypercalciuria due to enhanced osteoclastic activity. The individual dose needed to achieve optimal biological effects of vitamin D might be related not only to a single but also to multiple gene polymorphisms affecting vitamin D hydroxylases, DBP and/or VDR, as well as to body weight, body fat, age, skin color, season, latitude and sunning habits; optimal therapeutic effect in autoimmunity, in addition, should conceivably require pharmacological doses, much higher than those necessary for

**Table 3. Serum concentrations of 25(OH)D3, PTH, total calcium, ionized calcium, urea and creatinine and clinical status of 16 patients with vitiligo at baseline and after treatment with vitamin D (35,000 IU per day for 6 mo)**

| Parameter                  | Baseline (mean ± SD) | 6 mo (mean ± SD) | p value * |
|----------------------------|----------------------|------------------|-----------|
| 25(OH)D3 (30–100 ng/mL)    | 18.4 ± 8.9           | 132.5 ± 37.0     | 0.0005    |
| PTH (8–74 pg/mL)           | 55.3 ± 25.0          | 25.4 ± 10.7      | 0.0005    |
| Serum Calcium (8.4–11.0 mmol/L) | 9.2 ± 0.3           | 9.2 ± 0.3        | 0.7615    |
| Ionized Calcium (1.10–1.40 mmol/L) | 1.2 ± 0.2           | 1.2 ± 0.1        | 0.5417    |
| Urinary Calcium (100–300 mg/24h) | 158.3 ± 73.6       | 230.1 ± 81.4     | 0.0239    |
| Urea (< 50 mg/dL)          | 35.5 ± 7.2           | 33.9 ± 9.9       | 0.7148    |
| Creatinine (0.7 – 1.5 mg/dL) | 0.9 ± 0.2           | 0.9 ± 0.2        | 0.5016    |

* Wilcoxon signed rank test.
preventive measures. All interventions using high-dose vitamin D3 so far applied to treat autoimmune disorders, including the current one, have used arbitrarily selected doses, without taking into account the potentially large variability of resistance to the benefits and side-effects of vitamin D among the patients participating in the clinical trial. Such uniform research approach precludes the development of a rational protocol for establishing the optimal daily dose of vitamin D for each patient, which would improve both safety and therapeutic effectiveness in the clinical setting. Ultimately resulting from all these multiple variables interacting in the same organism, the serum concentration of PTH may be the best biological indicator for the individual setting of the optimal therapeutic dose of vitamin D3 for the treatment of autoimmune disorders. An isolated measurement of serum PTH concentration does not provide a reliable ancillary parameter to estimate the dose of vitamin D3 required for optimal control of autoimmunity. The enhancing effect of PTH on renal 1,25(OH)2D3 production declines with age, so that young patients may sustain efficient gastrointestinal absorption of calcium without raising PTH to the upper normal range or above in contrast to mature or old adults. Conversely, the magnitude of the decrease in serum PTH concentration, from the basal value to the level achieved after a period of treatment, may provide a reasonable estimation of how much the initial daily dose of vitamin D3 should be increased to reduce serum PTH levels to the lower reference range. A period of at least 2 mo should be allowed between the two serum PTH measurements, considering that 25(OH)D3 has a half-life of 15 d. Using the PTH level as an ancillary index of therapeutic response requires a diet only partially restricted in calcium (like the one described in this study) since excessive restriction of calcium intake would maintain increased bone resorption to preserve normocalcemia, thereby limiting or preventing vitamin D3-induced PTH drop. Avoiding excessively high doses of vitamin D3 capable of suppressing PTH and periodically measuring bone density, on the other hand, may conceivably indicate that 25(OH)D3 probably have not reached circulating concentrations capable of increasing osteoclastic activity. According to this view, the PTH reductions observed with sustained normal serum and urinary calcium after 6 mo of treatment either in patients with psoriasis (from 57.8 ± 16.7 pg/mL to 28.9 ± 8.2 pg/mL) or vitiligo (from 55.3 ± 25.0 pg/mL to 25.4 ± 10.7 pg/mL) did not reach the lower limit of the reference range (8–74 pg/mL), suggesting that they could obtain additional therapeutic benefit from even higher doses of vitamin D3 than those used in the current study without compromising the safety of treatment.

In summary, the present study suggests that, at least for patients with autoimmune disorders like vitiligo and psoriasis, a daily dose of vitamin D3 should be increased to reduce serum PTH levels to the lower reference range. A period of at least 2 mo should be allowed between the two serum PTH measurements, considering that 25(OH)D3 has a half-life of 15 d. Using the PTH level as an ancillary index of therapeutic response requires a diet only partially restricted in calcium (like the one described in this study) since excessive restriction of calcium intake would maintain increased bone resorption to preserve normocalcemia, thereby limiting or preventing vitamin D3-induced PTH drop. Avoiding excessively high doses of vitamin D3 capable of suppressing PTH and periodically measuring bone density, on the other hand, may conceivably indicate that 25(OH)D3 probably have not reached circulating concentrations capable of increasing osteoclastic activity. According to this view, the PTH reductions observed with sustained normal serum and urinary calcium after 6 mo of treatment either in patients with psoriasis (from 57.8 ± 16.7 pg/mL to 28.9 ± 8.2 pg/mL) or vitiligo (from 55.3 ± 25.0 pg/mL to 25.4 ± 10.7 pg/mL) did not reach the lower limit of the reference range (8–74 pg/mL), suggesting that they could obtain additional therapeutic benefit from even higher doses of vitamin D3 than those used in the current study without compromising the safety of treatment.

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![Figure 2](image-url)

**Figure 2.** Serum concentrations of 25(OH)D3 and PTH in patients with psoriasis before and after treatment with vitamin D (35,000 IU per day for 6 mo). (A) Box plot showing serum concentrations of 25(OH)D3 before and after treatment. (B) Same for the respective serum PTH concentrations. Significance level (Wilcoxon signed rank test) indicated in (A) and (B). (C) Serum concentrations of 25(OH)D3 and PTH respectively increased and decreased during treatment. (D) Linear regression of serum PTH on serum 25(OH)D3 concentrations is significant (significance level and r value are shown; dashed lines represent the 95% CIs for the linear regression line; baseline and 6-mo values are respectively shown as empty and filled circles).
under the CONEP number 0356/08). All patients gave spoken and written informed consent before inclusion in the study. This study included nine psoriasis patients and 16 vitiligo patients from two outpatient clinics in São Paulo, Brazil—the Dermatology Clinic of the Hospital Heliópolis and the Internal Medicine Clinic of Universidade Federal de São Paulo. Subjects were men or women over 18 y of age. The study protocol excluded (1) patients under treatment with thiazide diuretics or lithium to avoid drug-vitamin D interactions on calcium metabolism;58 (2) patients taking psoriasis- or vitiligo-inducing drugs; 59,60 (3) patients with cognitive impairment; (d) patients with renal insufficiency and (4) patients with elevated calcium/calciuria or primary hyperparathyroidism.

Materials and Methods

Subjects. The Ethics Committee of the Hospital Heliópolis (Sao Paulo Brazil) and the Ethics Committee of the Universidade Federal de São Paulo approved this study protocol (registered under the CONEP number 0356/08). All patients gave spoken and written informed consent before inclusion in the study.

This study included nine psoriasis patients and 16 vitiligo patients from two outpatient clinics in São Paulo, Brazil—the Dermatology Clinic of the Hospital Heliópolis and the Internal Medicine Clinic of Universidade Federal de São Paulo. Subjects were men or women over 18 y of age. The study protocol excluded (1) patients under treatment with thiazide diuretics or lithium to avoid drug-vitamin D interactions on calcium metabolism;58 (2) patients taking psoriasis- or vitiligo-inducing drugs; 59,60 (3) patients with cognitive impairment; (d) patients with renal insufficiency and (4) patients with elevated calcium/calciuria or primary hyperparathyroidism.
**Figure 4.** PASI scores in patients with psoriasis before and after treatment with vitamin D (35,000 IU per day for 6 mo). (A) Individual temporal profiles of the P.A.S.I. score during the treatment showing clinical improvement in all patients. (B) Linear regression of P.A.S.I. on 25(OH)D3 concentration is significant (significance level and r² value are shown; dashed lines represent the 95% CIs for the linear regression line; baseline and 6-mo values are respectively shown as empty and filled circles).

**Figure 5.** Serum concentrations of 25(OH)D3 and PTH in patients with vitiligo before and after treatment with vitamin D (35,000 IU per day for 6 mo). (A) Box plot showing serum concentrations of 25(OH)D3 before and after treatment. (B) Same for the respective serum PTH concentrations. Significance level (Wilcoxon signed rank test) indicated in (A) and (B). (C) Serum concentrations of 25(OH)D3 and PTH respectively increased and decreased during treatment. (D) Linear regression of serum PTH on serum 25(OH)D3 concentrations is significant (significance level and r² value are shown; dashed lines represent the 95% CIs for the linear regression line; baseline and 6-mo values are respectively shown as empty and filled circles).

**Design.** This is an open-label study where all patients received oral treatment with 35,000 IU of vitamin D (1.75 ml of a 20,000 IU/mL sunflower oil solution) per day. Whenever present, standard treatments were not changed. Following instructions, all patients excluded milk and dairy products (as well as calcium-fortified foods like soy, oat or rice milk) from their diet and
parathormone (PTH), serum urea and creatinine levels. By collecting all data from March to October 2011 and from March to October 2012, the authors attempted to minimize seasonal variations in 25(OH)D3 levels and related effects.

**Analytical methods.** 25(OH)D3 levels were determined by HPLC (LC 20AT) with UV detection at 265 nm (UV/VIS model SPD-20A). 25(OH)D3 standard, solvents and other reagents were of analytical grade (all purchased from Sigma-Aldrich). An internal standard (IS) of 25(OH)D3 compensated the reduced HPLC sensitivity to detect very low levels of 25(OH)D3. Stock solution of IS was prepared in methanol at 100 μg/mL and kept at −20°C in amber tube. The HPLC conditions

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**Figure 6.** (A), (B), (C), (D), (F) Box plots respectively showing concentrations of serum calcium, 24-h urinary calcium excretion, serum urea and serum creatinine in patients with vitiligo before and after treatment with vitamin D (35,000 IU per day for 6 mo), with a significant pre- and post-treatment difference only found for 24-h urinary calcium excretion (Wilcoxon signed rank test). (D) Linear regression of 24-h urinary calcium excretion on serum 25(OH)D3 concentrations is significant (significance level and r² value are shown; dashed lines represent the 95% CIs for the linear regression line; baseline and 6-mo values are respectively shown as empty and filled circles).

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ingested at least 2.5 L of fluid per day to prevent, respectively, excessive absorption of intestinal calcium and concentrated urinary calcium. Calcium supplementation was not allowed. The onset of symptoms suggestive of hypercalcemia (increased thirst, constipation, nausea, vomiting) would require performing extra laboratory tests. Regular medical follow-up appointments ensured compliance with dietary restriction and fluid intake.

After clinical diagnosis, all patients had clinical scoring, photo documentation of affected skin areas, laboratory tests performed before, at three and six mo of treatment. Laboratory tests included basal vitamin D status (25(OH)D3 levels), total and ionized serum calcium, total 24-h urinary calcium excretion,
were LiChrosorb RP-18 5 μm column (Merck), 40°C, acetonitrile-methanol-water (90:4:6, v/v) as the mobile phase, 3.2 min for retention time and flow rate of 1 ml/min. The calibration curve was linear over the concentration range of 9.8 ng/mL to 1250.0 ng/mL ($r^2 = 0.9994$).

Serum samples were immediately stored at −80°C until analysis. The following steps were performed under light protection. Each serum sample of 100 μL was thawed, added to 70 μL of methanol-2-propanol mixture (80:20) and then vortex-mixed for 30 sec. Subsequently, 400 μL of hexane was added, vortex-mixed again for 60 sec and centrifuged under 2500 g for 10 min at 10°C. A volume of 100 μL of the resulting organic phase was submitted to evaporation in a Concentrator model 5301 for 10 min at room temperature. The resulting dried residue was then resuspended in 50 μL of methanol plus 50 μL of IS (working solution of 200 ng/mL) and vortex-mixed for 2 min prior to injection into HPLC. The final result was obtained by deducting the amount corresponding to added IS from the ultimate figure released from HPLC. The IS method appropriately correlated with assessments of standard 25(OH)D3 concentrations and serum samples.

PTH was essayed by chemiluminescence, while other laboratory tests (total and ionized serum calcium, urea and creatinine) were analyzed by standard methods at the Central Laboratory of the University Hospital (Federal University of São Paulo).

Clinical parameters of vitiligo and psoriasis. The PASI score used for assessment of severity and extent of psoriasis enabled the evaluation of the response to treatment. Pre- and post-treatment photographic documentation of the affected skin under Wood’s light (UVA 351 nm) and a quartile grading scale enabled the evaluation of the clinical response to the treatment in patients with vitiligo. The percentage of the affected skin surface that achieved repigmentation at 6 mo of treatment yielded five categories of clinical response. These included “0” for no repigmentation; “1” for 1 to 25% repigmentation, “2” for 26 to 50% repigmentation, “3” for 51 to 75% repigmentation and “4” for repigmentation above 75% of the total baseline area of affected skin.

Statistics. Results are expressed in tables as mean ± standard deviation. The Wilcoxon assigned rank test was employed to assess the effect of treatment with vitamin D (50,000 IU daily for six months). The relationship between 25(OH)D3 and continuous variables (PTH and urinary calcium excretion) was evaluated using a Pearson correlation coefficient. Curves relating serum 25(OH)D3 and PTH or urinary calcium levels were fitted using linear regression models. Statistical analyses and graphical presentation were performed using the GraphPad Prism (Version 3.02, San Diego, USA).

Disclosure of Potential Conflicts of Interest
The authors declare no conflict of interest.

Acknowledgments
C.G.C., R.S.G., D.C.F. and A.C.L. designed the study; D.C.F. and R.S.C. conducted the research; M.G. provided and prepared vitamin D supplement, R.S.C., L.C.M.N., L.D.T., J.J.S. and F.S. performed the analytical method for 25(OH)D3 measurements; N.F.N. and Y.J. performed statistical analysis; C.G.C., R.S.C., D.C.F., L.C.M.N., D.J.N. and A.C.L. analyzed data; C.G.C., R.S.C. and D.C.F. wrote the paper. All authors have approved the manuscript. This project was supported by grants from CAPES, Institute for Investigation and Treatment of Autoimmune Disorders (I.I.T.A.) and UNIMED do ABC, Brazil. DCF was a CAPES fellow.

References
1. Ramagopalan SV, Heger A, Berlanga Al, Maugeri NJ, Lincoln MR, Burrell A, et al. A ChIP-seq defined genome-wide map of vitamin D receptor binding: associations with disease and evolution. Genome Res 2010; 20:1352-66; PMID:20736230; http://dx.doi.org/10.1101/gr.107920.110
2. Holick MF. The vitamin D deficiency pandemic and consequences for nonskeletal health: mechanisms of action. Mol Aspects Med 2008; 29:361-8; PMID:18801384; http://dx.doi.org/10.1016/j.mam.2008.08.008
3. Eyles DW, Smith S, Kinobe R, Hewison M, McGrath JJ. Distribution of the vitamin D receptor and 1 alpha-hydroxylase in human brain. J Chem Neuroanat 2005; 29:21-30; PMID:15586999; http://dx.doi.org/10.1016/j.jchemneu.2004.08.006
4. Adams JS, Hewison M. Extrarenal expression of the 25-hydroxvitamin D-1-hydroxylase. Arch Biochem Biophys 2012; 523:95-102; PMID:22446158; http://dx.doi.org/10.1016/j.abb.2012.02.016
5. Eyles DW, Burne TH, McGrath JJ. Vitamin D, effects on brain development, adult brain function and the links between low levels of vitamin D and neuropsychiatric disease. Front Neuroendocrinol 2013; 34:47-64.; PMID:23796576; http://dx.doi.org/10.1016/j. fnen.2012.07.001
6. Hewison M. An update on vitamin D and human immunity. Clin Endocrinol (Oxf) 2012; 76:315-25; PMID:21995874; http://dx.doi.org/10.1111/j.1365-2265.2011.04201.x
7. Holick MF. Evidence-based D-bate on health benefits of vitamin D revisited. Dermatoendocrinol 2012; 4:183-90; PMID:22928075; http://dx.doi.org/10.4161/derm.20015
8. IOM (Institute of Medicine). Dietary reference intakes for calcium and vitamin D. Committee to Review Dietary Reference Intakes for Calcium and Vitamin D. Washington, DC: The National Academies Press. 2011.
Figure 8. For figure legend, see page 11.
11. Figure 8. Photographs of two male patients with psoriasis before (A and C) and after (B and D) six months of treatment with vitamin D (35,000 IU per day). (A and D) A 59-year-old patient with BMI of 24.8 presenting a PASI score of 31 before treatment and achieving score of 18.2 after six months of treatment; his serum concentration of 25(OH)D was 22.8 ng/mL at baseline, reaching 127.5 ng/mL after 6 mo of treatment. (C and D) A 60-year-old patient with BMI of 33.6 presenting a PASI score of 40.4 at baseline, achieving score of 12.4 after six months; his serum concentration of 25(OH)D was 5.6 ng/mL, reaching 103.2 ng/mL after six months of treatment.
Figure 9. For figure legend, see page 13.
51. Henry HL. Regulation of vitamin D metabolism. Best Pract Res Clin Endocrinol Metab 2011; 25:531-41; PMID:21872796; http://dx.doi.org/10.1016/j. beem.2011.05.003

52. Bikle DD. Vitamin D and immune function: understanding common pathways. Curr Opin Pharmacol 2009; 9:58-65; PMID:19631030; http://dx.doi.org/10.1016/j.copha.2009.05.011

53. Beckman MJ, Horst RL, Reinhart TA, Beitz DC. Up-regulation of the intestinal 1,25-dihydroxyvitamin D receptor during hypervitaminosis D: a comparison between vitamin D2 and vitamin D3. Biochem Biophys Res Commun 1990; 169:910-5; PMID:2163637; http://dx.doi.org/10.1016/0006-291X(90)91979-3

54. Buckle RM, Gamlen TR, Pullen IM. Vitamin D intoxication treated with porcine calcitonin. Br Med J 1972; 3:205-7; PMID:4261142; http://dx.doi.org/10.1136/bmj.3.5820.205

55. Selby PL, Davies M, Marks J, Mawer EB. Vitamin D intoxication causes hypercalcaemia by increased bone resorption which responds to pamidronate. Clin Endocrinol (Oxf) 1995; 43:531-6; PMID:8548936; http://dx.doi.org/10.1111/j.1365-2265.1995. tb02016.x

56. Cannell JJ, Hollis BW. Use of vitamin D in clinical practice. Altern Med Rev 2008; 13:6-20; PMID:18377099

57. Tsai KS, Wahner HW, Offord KP, Melton LJ 3rd, Kumar R, Riggs BL. Effect of aging on vitamin D stores and bone density in women. Calcif Tissue Int 1987; 40:241-3; PMID:3107776; http://dx.doi.org/10.1007/BF02555255

58. Hariri A, Mount DB, Rastegar A. Disorders of calcium, phosphate, and magnesium metabolism. In: Mount DB, Sayegh MH, Singh AK, ed. Core Concepts in the Disorders of Fluid, Electrolytes and Acid-Base Balance. New York, NY: Springer US, 2013:103-146.

59. Duhra P, Foulds JS. Persistent vitiligo induced by diphencyprone. Br J Dermatol 1990; 123:415-6; PMID:2206982; http://dx.doi.org/10.1111/j.1365-2133.1990.tb06306.x

60. Basavaraj KH, Ashok NM, Rashmi R, Praveen TK. The role of drugs in the induction and/or exacerbation of psoriasis. Int J Dermatol 2010; 49:1351-61; PMID:21091671; http://dx.doi.org/10.1111/j.1365-4632.2010.04570.x

61. Fredriksson T, Pettersson U. Severe psoriasis - oral therapy with a new retinoid. Dermatologica 1978; 157:238-44; PMID:357213; http://dx.doi.org/10.1159/000250839

56. Fredriksson T, Pettersson U. Severe psoriasis - oral therapy with a new retinoid. Dermatologica 1978; 157:238-44; PMID:357213; http://dx.doi.org/10.1159/000250839

62. Shin J, Lee JS, Hann SK, Oh SH. Combination treatment by 1060 nm ablative fractional carbon dioxide laser and narrowband ultraviolet B in refractory non-segmental vitiligo: a prospective, randomized half-body comparative study. Br J Dermatol 2012; 166:658-61; PMID:22050270; http://dx.doi.org/10.1111/j.1365-2133.2011.10723.x