REVIEW

Vitamin D in critically ill patients - from molecular damage interactions to clinical outcomes benefits. When, why, how?

Alida Moise
Anesthesia and Intensive Care Unit, „Prof. Dr. D. Gerota” Hospital, Bucharest, Romania

Correspondence to:
Alida Moise, PhD, Chief of Anesthesia and Intensive Care Unit, „Prof. Dr. D. Gerota” Hospital
Profesional address: „Prof. Dr. D. Gerota” Hospital, no 29-31 V.V. Stroiescu Street, Bucharest 21392, Bucharest, Romania
E-mail: alidamoise@gmail.com

Conflicts of interests
Nothing to declare

Acknowledgment
None

Funding: This research did not receive any specific grant from funding agencies in the public, commercial or not-for profit sectors.

Keywords: Vitamin D; effect; cognition; delirium; obesity.

These authors take responsibility for all aspects of the reliability and freedom from bias of the data presented and their discussed interpretation.

Abstract
Vitamin D - „the sunshine vitamin” is essential for the good functioning of the human body. The most important forms of the vitamin D are the vitamin D2 and the vitamin D3, both biologically inactivated. Vitamin D can come from: diet or nutritiv suplimentts and skin. The activation of vitamin D is effect in two steps to the physiologically active 1,25-dihydroxyvitamin D [1,25(OH)2D].

The biological actions of Vitamin D involve regulation of gene expression at the transcriptional level and are mediated through binding to a Vitamin D Receptor (VDR). Vitamin D has multiple roles: hormonale/nonhormonale, skeletale/nonscheletale, genomece/nongenomece. Interesting is inversely corelation between Vitamin D and total body fat (BMI) and correalation between Vitamin D and cognitive impair-ment, especially Alzheimer Disease or delirium during hospitalisation.

The current recomendations regarding the supplying with Vitamin D are different for re- gions of the globe, also differ depending on the baseline serum Vitamin D and on the desired effect. So, potential nonskeletal effects occur at levels >30ng/ml, above 50-75ng/ml, serum level who should become the target of the supple-mentation. The loading dose should be considered perioperatetely for rapid effects.

In conclusion, Vitamin D is more than just a vitamin. It is a substance with multiple roles in body’s economy, and in recent years there has been an interest in the relation be-
between vitamin D deficiency and obesity or cognitive impairment. The majority of the data supports association, not causation, of low vitamin D levels. In other words, much of data does not clearly support the idea that vitamin D supplementation in a patient with low vitamin D levels reduces the risk of these diseases. But, the supplementation is very easy and no harm might be done.

**Introduction**

Vitamin D is also called „the sunshine vitamin“. It is essential for the good functioning of the human body. In the history of Vitamin D there also is a Nobel Premium. It was won by Adolf Windause for chemistry in 1928 “for his studies on the constitution of the sterols and their connection with vitamins”. Where does this name come from, why was this vitamin called the vitamin D in 1922? Simple, because there already existed vitamins with the names: A, B, C so the only letter that was not taken was D [1].

**Forms**

The vitamin D is actually a group of some vitamins, that are soluble in fat (Vitamins D1-D5), also called seco-steroisis. The most important forms of the vitamin D are the vitamin D2, discovered in 1932 and the vitamin D3, discovered in 1936. Both forms function as prohormones and are considered biologically inactived [2-4].

![Ergocalciferol (Vitamin D2)](image1.png)  ![Cholecalciferol (Vitamin D3)](image2.png)

**Figure 1. The most important forms of Vitamin D**

**Sources of Vitamine D**

Vitamin D can come from:

1. **diet** (fatty fish like salmon, sardines, tuna and eel; fish oils like cod liver oil; mushrooms and eggs, milk and products of milk, cereals, bread, pastele, butter, margarine and oil) or nutritiv suplimentts.

Some countries, like the USA, add vitamin D to foods like milk, soy milk, yogurt, butter, cheese, cereal grains and beer [3].

2. **skin**, through photochemical synthesis - the major natural source of the vitamin. It is dependent on moderate sun exposure to the face, arms and legs, averaging 5–30 minutes twice per week (UVB light between 270 and 300 nm). These wavelengths are present when the UV index is greater than 3. At angles greater than 45° above the horizon (at sea level), vitamin D production will be occurring, although some recent research suggests that vitamin D production may occur at angles as low as 30°.

The production of vitamin D in the skin depends on:

- the season of the year,
- the latitude,
- use of sunscreen,
- clothing,
- amount of skin exposed,
- age - synthesis of vitamin D declines with increasing age, due in part to a fall in 7-dehydrocholesterol levels and due in part to alterations in skin morphology [5],
- the amount of melanin in their skin – darker skin requires longer than fair skin.

It is thought that as little as five minutes per day of incidental sun exposure is sufficient for the production of Vitamin D, for someone who burns easily, and up to 20 minutes is sufficient for a person with darker skin [4,15].

In the skin (derm), photochemically, non-enzimatic, into thermal isomerization of 7-dehydrocholesterol to previtamin D3, inactive biological which is then rapidly converted to vitamin D3 (cholecalciferol) by temperature- and membrane-dependent processes, but excessive exposure to sunlight degrades previtamin D2 and vitamin D3 into inactive photoproducts [12,15].

**Metabolism of Vitamin D**

Dietary vitamin D (either D2 or D3) is absorbed with other dietary fats in the small intestine via chylomicrons. Its absorption is dependent on the presence of fat in the lumen, which triggers the release of bile acids and pancreatic lipase with the formation of lipid-containing micelles, which diffuse into enterocytes.

Vitamin D enters the circulation from the lymph or from the skin, it is prelucrated by the liver or storage tissues within a few hours. The activation of vitamin D is effect in two steps:

- **the first step** occurs in the liver and converts previtamin D to 25-hydroxyvitamin D [25(OH)D], calcidiol, also biologically inactive form. This etape is mediated by the 25-hydroxylase (cytochrome P450 enzyme 2R1 - CYP2R1). 25OH-D is the major circulating form of
vitamin D [15]; it circulates bound to a specific plasma carrier protein, Vitamin D Binding Protein (DBP). DBP also transports vitamin D and its metabolites [6-8,14,15].

The second step occurs primarily in the kidney and forms the physiologically active 1,25-dihydroxyvitamin D [1,25(OH)₂D], calcitriol, mediated by 1α-hydroxylase (CYP27B1) [4,8,10,15] and regulated by serum phosphorus, calcium and itself, as by fibroblast growth factor 23 (FGF-23), parathormone (PTH) with up-regulation and down-regulation via fibroblast-like growth factor-23. The 1α-hydroxylase gene is also expressed in several extra-renal tissues, but its contribution to calcitriol formation in these tissues is unknown. CYP27B1 is also expressed extrarenally in bone, placenta, prostate, keratinocytes, macrophages, T-lymphocytes, dendritic cells, several cancer cells, and the parathyroid gland and enables the production of 1,25(OH)₂D. This active form of vitamin D is locally active and exerts auto- or paracrine effects [8-12].

Serum albumin and serum DBP (D-binding protein) are the principal carriers of vitamin D and its metabolites. The affinity of albumin for 25 (OH)Vitamin D is substantially lower than that of DBP. DBP have the important role by actively facilitating the acquisition of 25(OH)D/1,25(OH)₂D or by defining the amount of ‘free vitamin D’ that is available for passive cellular uptake [14,18].

Excess fat-soluble vitamins D2, D3 after 24 hours are stored in the liver and fatty tissue for approximately two months. Adipose tissue as the primary site of vitamin D accumulation and there was enhanced uptake and clearance of vitamin D by adipose tissue in obese subjects compared with those of normal weights. Adipose tissue stores of vitamin D probably represent “non-specific” stores sequestered because of the hydrophobic nature of vitamin D, but the extent to which the processes of accumulation or mobilization are regulated by normal physiological mechanisms remains unknown at this time [4].

When the body needs more vitamins, vitamin D-2 and vitamin D-3 are converted to their active 25-hydroxyvitamin D. The released 25-hydroxyvitamin D can circulate in the body for approximately three weeks. After the body is replenished, the biologically active form is stored in fat tissues for months; 25-hydroxyvitamin D is released from the storage irregularly depending on the body’s need. By the time the physician detects vitamin D deficiency, serum concentration of 25-hydroxyvitamin D is less than 20ng/ml. At this point, vitamin D stores have been depleted [4].

Normal serum level and deficiency

Laboratory measures 25(OH)D concentration, that is consideret the functional indicator of vitamin D status [16,20].

The interpretation of vitamin D levels varies in literature: some consider the deficiency is <25-30ng/ml, others think that Vitamin D deficiency is diagnosed when 25(OH)D <20 ng/mL and vitamin D insufficiency is defined as 25(OH)D of 21–29 ng/mL [20].

For endocrinologists the value for 25(OH)Vitamin D more 30 ng/mL is considered sufficient, with 40–60 ng/mL being the preferred range.

Vitamin D intoxication usually does not occur until 25(OH)D >150 ng/mL and results only from the excessive consumption of supplements with hypercalcaemia [15,20].

The clinical practice corellates the level of Vitamin D with its effects. The threshold for normality should probably be the serum 25OHD required for maximum suppression of parathyroid hormone, greatest calcium absorption and highest bone mineral density [41].

Effects of Vitamin D

The biological actions of calcitriol involve regulation of gene expression at the transcriptional level and are mediated through binding to a Vitamin D Receptor (VDR), a specific zinc-finger nuclear receptor superfamily of steroid/thyroid hormone receptors, located primarily in the nuclei of target cells, and regulates expression of up to 2000 genes, directly or indirectly [15].

VDR is present in most tissues and cells in the body, allowing it to do her roles:
- jejenum and ileum (increases absorption of calcium and magnesium),
- bone (role in the mineralization),
- cardiovascular system (endothelium, vascular smooth muscle - may protect against atherosclerosis; cardiomyocytes –inhibition of cell proliferation without apoptosis; promotes cardiomyotube formation, induces cardiac differentiation) [37],
- pancreatic beta cells (mediated by upregulation of insulin receptors and modulates inflammation);
- hepatocytes (down-regulates fibrogenic TGF-β signaling, anti-inflammatory effects by inhibiting monocyte activation and TNF-α and IL-1 expression),
- cells of immune system like (acts as an
immunomodulator: local macrophage synthesis of 1,25(OH)2D is implicated in response to TLR signaling is also a key feature of innate immunity (13), inhibition of production of interleukin by activated T-lymphocytes and of immunoglobulin by activated B-lymphocytes, differentiation of monocyte precursor cells, modulation of cell proliferation, influences several growth factors (IGF-1, TGFβ, MAPK5, NF-Kb [12,13,15,22], -muscle -increases protein synthesis, protects against insulin resistance [37], - skin (antiproliferative, immunosuppressive and prodifferentiating effects), - adipocytes – inhibits intracellular fat accumulation, enhances basal lipolysis without cell toxicity, upregulation of β-oxidation-related genes, lipolytic enzymes and vitamin D-responsive genes [37].
- selected cancer cells - antiproliferative, prodifferentiating, increases apoptosis, decreases angiogenesis: date for renal carcinoma, thyroid, breast, colon, neural cells and prostat - the nonhormonal, intracrine, and paracrine actions of 1-hydroxylated vitamin D metabolites in man [8, 9,13,21,29,37].

The classification of the roles:
- hormonale/ nonhormonale,
- skeletale/nonscheletale,
- genomice/nongenomice (mediated through the VDR transcriptional effects inside the cell nucleus / VDR induces rapid signaling, situated on the cell membrane and/or cytoplasm, via calcium) [15,21,22].

A large number of cellular processes are regulated by calcium - as a primary signaling pathway or it can operate as a modulator signal.
- in non-excitable cells - fertilization, proliferation, metabolism, secretion and smooth muscle contraction.
- in excitable cells- contraction in the heart or memory formation in neurons.
- a modulator signal that can induce subtle changes in the generation and function of this primary Ca2+ signal: Alzheimer’s disease (AD) and cardiovascular disease.

Beside its role in homeostasis of the calcium, Vitamin D has a role in regulating cell proliferation and differentiation, in insulin resistance, obesity, metabolic syndrom and various cancers. Also vitamin D possess antiproliferative, pro-apoptotic and immunomodulatory effects in cancer [8,15]. For non-hormonal roles of vitamin D, the regulation is a lot more sensitive. There is the association between the deficiency of vitamin D and many acute and chronic illnesses including disorders of calcium metabolism, autoimmune diseases (multiple sclerosis, rheumatoid arthritis, etc), some cancers (breast, colon, prostat), type 2 diabetes mellitus, respiratory, cardiovascular disease and infectious diseases (tuberculosis, influenza, viral), but not it is not proven enough yet. [15]. Also, Vitamin D supplementation seemed to decrease general mortality in elderly people [23]. The majority of the data supports association, not causation, of low vitamin D levels.

Obesity and Vitamin D

Interesting is inversely correlation between total body fat (BMI) and 25-OH-D levels. More often in individuals who are not obese, body size and adiposity are inversely associated with blood 25(OH)D concentrations. BMI is inversely associated with the increase in serum 25(OH)D levels in response to vitamin D supplementation [24-28].

Hypotheses for correlation BMI- level of vitamin D:
- may be due to the increased metabolic clearance of vitamin D through enhanced uptake in fat tissue and/or the decreased bioavailability of vitamin D once it is deposited in the fat tissue;
- to storage of vitamin D in fat - The concept of storage of vitamin D in the adipose tissue has gained recent popularity and some studies recommend a higher intake of vitamin D in obese women, however there are no longitudinal studies to supports that suggestion.
- sedentary lifestyle, low sunlight exposure,
- true vitamin D deficiency - genetic changes in vitamin D metabolism
- diet
- other unknown factors [15,16,19,24-28] But, more trials did not show significant change in total body fat mass after treatment with vitamin D [28].

Cognition and Vitamin D

The most of important correlation is between Vitamin D and cognitive impairment, because the impact on human society and because of validation of the information available on this subject.

Boston, hosted in July 2013 the meeting of the international experts at the invitation summit on „Vitamin D and cognition in older adults“. Based upon literature and expert opinion, the task force focused on key questions on the role of vitamin D in Alzheimer disease (AD) and related disorder. Each question was discussed and voted using a Delphi-like approach [29].
Conclusions were:
- hypovitaminosis D increases the risk of cognitive decline and dementia in older adults,
- not be used thus far as a diagnostic or prognostic biomarker of Alzheimer disease due to lack of specificity and insufficient evidence,
- hypovitaminosis D should be screened in this population because of its high prevalence and supplemented, if necessary, but this advice was not specific to cognition.

There is a site dedicated to Vitamin D with information about this vitamin:
Vitamin D. There are a lot of articles for Vitamin D and cognition on the site.

A significant association between AD and low levels of vitamin D has been demonstrated. The role of vitamin D in AD are:
- regulating calcium-sensing receptor expression,
- enhancing amyloid-β peptides clearance,
- downregulating matrix metalloproteinases,
- upregulating heme oxygenase 1,
- suppressing the reduced form of nicotinamide adenine dinucleotide phosphate expression.

Furthermore, vitamin D supplements appear to have a beneficial clinical effect on AD by:
- regulating micro-RNA,
- enhancing toll-like receptors,
- modulating vascular endothelial factor expression,
- modulating angiogenesis,
- regulating calcium-sensing receptor expression,
- enhancing amyloid-β peptides clearance,
- down regulating matrix metalloproteinases,
- up regulating heme oxygenase 1,
- suppressing the reduced form of nicotinamide adenine dinucleotide phosphate expression.

The results of a meta-analysis revealed that subjects with deficient vitamin D status (25(OH)D level < 50 nmol/L) were at increased risk of developing AD by 21% compared with those possessing 25(OH)D level > 50 nmol/L [31].

Time of progression to severe stage of Alzheimer’s disease was slower in patients under treatment with vitamin D compared with those without treatment (5.4 ± 0.4 years vs. 4.4 ± 0.16 years respectively, p=0.003) [32].

Similarly, the risk of cognitive impairment was up to four times bigger in the severely deficient elders (25(OH)D <25 nmol/L) in comparison with individuals with adequate levels (75 nmol/L) [33].

There are functional explications: neurosteroid properties of vitamin D. The effects of Vitamin D are exercised through its nuclear hormone receptor the VDR, which is expressed in neuronal and glial cells in almost all regions of the CNS, and in special in the hippocampus, hypothalamus, cortex and subcortex, the areas essential for cognition. The active form of vitamin D has a trophic function of neuronal differentiation and maturation via control of the synthesis of neurotrophic agents such as nerve growth factor (NGF) and glial cell line-derived neurotrophic factor (GDNF). Moreover, 1,25OHD regulates the genetic expression of numerous neurotransmitters in the brain, including acetylcholine, dopamine, serotonin and c-aminobutyric acid, notably in the hippocampus [34,40].

The vitamin D is unregulator of neuroprotection, antiepileptic and anticalcification effects, neuro-immunomodulation, interplay with neurotransmitters and hormones, modulation of behaviors, brain ageing, and some other, less-explored, brain processes [35].

The activities of vitamin D for consolidation of cognitive may represent primary effects in the brain rather than, secondary systemic effects given the expression of vitamin D receptors in the brain, and Vitamin D can prevent neuronal damage through the neurotrophic factor and through the process of detoxification. It also contributes to regulating behaviour and protecting the brain through an antiinflammatory anti-oxidant buffer [36].

Moreover, for clinical, there are the speculation that vitamin D may thus play a role in protecting the brain from delirium. So, Vitamin D promotes neuroprotection by modulating the production of choline acetyltransferase. Choline acetyltransferase is the key enzyme in biosynthesis of acetylcholine and cholinergic deficits are implicated in delirium. In addition, Vitamin D may contribute to neuroprotection by modulating the production of glial cell-derived neurotrophic factor, NO synthase [38-40].

Vitamin D status before hospitalisation is associated with the risk of delirium, a problem of great interest in this moment. The level of pre-hospital 25(OH)D < 10 ng/ml is indeed associated with a significant increase in the odds of delirium in patients during hospitalisation but future studies need to complete this information [40].

In a recent study, the low level of vitamin D has been associated with an increased risk of elderly patients developing cognitive impairment in postoperative [43].
Vitamin D supplementation

The current recommendations regarding the supplying with Vitamin D are different for regions of the globe. These recommendations also differ depending on the baseline serum Vitamin D and on the desired effect [16,42].

So, potential nonskeletal effects occur at levels >30ng/ml, above 50-75ng/ml, serum level which should become the target of the supplementation.

There are on the website of The Endocrine Society the following schemes:
- 400-600ui/day for children (0-18y) with skeletal effects, but not known clear at this time what are potential nonskeletal health benefits. For a level of Vitamin D>30ng/ml are required at least 1000 IU/d.
- 600 IU/d for 19-50y. For a level of Vitamin D >30ng/ml are required at least 1500–2000 IU/d.
- 800IU/d for >50y, also for a level of Vitamin D >30ng/ml are required at least 1500–2000 IU/d [20, 27].

There are special situations when if vitamin D is given at least two to three times more for their age group:
- obese children and adults
- children and adults on anticonvulsant medications, glucocorticoids, antifungals such as ketoconazole, and medications for AIDS.

The administration of high doses suggest that the maintenance tolerable upper limits of vitamin D, which is not to be exceeded without medical supervision is 4000 IU/d for anyone over 8 yr [20].

Also, there are high variability in response to a given dose of vitamin D supplementation [16,17,18]. The higher doses vitamin D3 supplementation can not always increases the serum level upper 50ng/ml, the baseline serum being very important. Therefore, for all adults who are vitamin D sever deficient and for rapid effects is recommended be treated with loading doses (50,000 IU of vitamin D2 or vitamin D3 once a week for 8 wk or its equivalent of 7,000 IU of vitamin D2 or vitamin D3 daily) to achieve a blood level of 25(OH)D above 30 ng/ml, followed by maintenance therapy of 1500–2000 IU/d [20,42].

We strongly believe that the loading dose should be considered for rapid correction of the deficiency of Vitamin D, especially perioperatively in all oxidative stress situations. More like a concentration below 50 nmol/L for 25(OH)D was an independent and significant risk factor for delirium (odds ratio = 2.7; 95% confidence interval: 1.04–7.2, P = 0.04) in a multivariate regression analysis adjusted for all registered confounders at hip fracture patients in postoperative [44]. Also, Vitamin D is „essential for both optimal nerve function and recovery following stress” [40]. But, although the loading doses administered perioperatively seem attractive, association between rapid improvement of vitamin D status and the incidence of the delirium should be verified in future studies.

At this time, for prevention of cognitive diseases the recommendation is 4000-5000 IU/d and for Alzheimer’s disease 4,000 IU/d of vitamin D [45].

Conclusion

Vitamin D is more than just a vitamin. It is a substance with multiple roles in body’s economy, and in recent years there has been an interest in the relation between vitamin D deficiency and obesity or cognitive impairment.

The majority of the data supports association, not causation, of low vitamin D levels. In other words, much of data does not clearly support the idea that vitamin D supplementation in a patient with low vitamin D levels reduces the risk of these diseases.

But, the supplementation is very easy and no harm might be done.

References

6. Strushkevich N, Usanov SA, Plotnikov AN, Jones G, Park HW. Structural analysis of CYP2R1 in complex with vitamin D3. Jour-


31. Shen L, Ji HF. Vitamin D deficiency is associated with increased risk of Alzheimer’s disease and dementia: evidence from me-


37. Al-Tarrah K, Hewison M, Moiemen N, Lord JM. Vitamin D status and its influence on outcomes following major burn injury and critical illness. Burns Trauma 2018;6:11


Abbreviations

1,25(OH)₂D - 1,25-dihydroxyvitamin D
25(OH)D - 25-hydroxyvitamin D
AD - Alzheimer’s disease
BMI - body mass index
CYP27B1 - cytochrome P450 enzyme 27B1
DBP - Vitamin D Binding Protein
FGF-23 - fibroblast growth factor 23
GDNF - glial cell line-derived neurotrophic factor
IL-1 – interleukine 1
NGF - nerve growth factor
NO - nitrogen
PTH - parathormone
RNA – ribonucleic acid
TGF-β – Tumor growth factor β
TNF-α – tumor necrosis factor α
VDR - Vitamin D Receptor
UVB – ultraviolet radiation

Vitamin D