The impact of vitamin D: From a fetus to an infant

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The focus of this paper is to review the data on the classic and non-classic role of VD with regards to pregnancy and the newborn period. Over the past decade, new evidence has shown that vitamin D (VD) deficiency, in addition to its classical role in calcium metabolism and bone homeostasis, may contribute to the risk of developing a wide range of chronic diseases. VD may produce a wide array of favourable biological effects via genomic, non-genomic or intracrine mechanisms, and therefore contributes to the improvement of human health. Some of these effects may be even more important during pregnancy. Data from animal and human studies implicate maternal VD deficiency as a significant risk factor for several adverse outcomes affecting maternal, foetal and child health. In the newborn period, these comprise bone health, growth and immune response. Conclusion – Recent evidence supports the fact that low maternal vitamin D status is associated with an increased risk of adverse pregnancy outcomes. This paper investigates the effects of vitamin D on the placento-foetal unit and the mother, in terms of calcium metabolism and non-calcium effects.

Introduction

From conception onward, a child develops under the influence of a series of biological events that enable the maturation of tissues, growth and adaptation. This multi-factorial process represents the interplay of genetic, environmental and dietary aspects. The biological background of various influences on the developmental process has been the subject of research in numerous studies on animals over the past few years. It has been proven that, within a critical period during intrauterine development, the so-called “window of vulnerability”, foetal development is sensitive to the changing environment, which enables phenotypic diversity (1). During this time, certain parameters associated with the mother, such as vitamin D (VD) deficiency, may affect the body composition, bone mass, and the physiology of the child (2). In the past decade, different studies have enabled researchers to establish a link between VD status in pregnant women and the outcome of pregnancy, the health of the foetus and the newborn infant, as well as the health of the offspring in their childhood and adulthood (2). The classic action of VD in maintaining calcium homeostasis and its impact on bone health has been known for a long time (3). However, recent findings suggest that VD also plays a role in the process of immunomodulation, cell proliferation and differentiation, as well as in many other physiological functions in various tissues and organs, including the brain, pancreas,
kidneys, placenta and heart (3). Lack of VD during pregnancy is very common in many parts of the world, and there is a link between low VD levels and the possibility of adverse outcomes (4, 5). Despite reports on the wide prevalence of VD deficiency and its potential consequences, the criteria for determining the optimal level in the body, and thus the quantity of VD intake needed to maintain these adequate levels, remain a matter of debate (6, 7, 8).

This paper presents data from the scientific literature, and discusses the role of VD during pregnancy, as well as the effects of maternal VD deficiency on the offspring’s health, regarding their bones and their entire organism (non-skeletal health).

**Physiology and the biological role of vitamin D**

VD, or calciferol, is regarded as a pro-hormone. The term VD defines a group of fat-soluble secosteroids, of which the main representatives are VD$_2$ (ergocalciferol) and VD$_3$ (cholecalciferol). Exposure to sunlight, to ultraviolet radiation B (UVB) in particular, triggers the conversion of pro-VD$_3$ in the skin (7-dehydrocholesterol) into VD$_3$, which binds to the binding protein for VD; it is then transported into the bloodstream and accumulated in adipose tissue or metabolised in the liver. Skin synthesis is the main source of VD, although certain foods, such as fish oil and cod liver oil, also provide direct intake of VD$_3$ (3, 9). Additionally, VD$_2$ is produced by UVB irradiation of the ergosterol in plants and fungi, but has lower bioavailability than VD$_3$. All forms of VD are biologically inactive until hydroxylated, which first takes place in the liver with the enzyme 25α-hydroxylase. This results in 25-hydroxy-vitamin D [25(OH)D]. Since this conversion is not controlled, the concentration of 25(OH)D increases in proportion to the synthesis and VD intake. The second hydroxylation occurs in the kidneys, with the assistance of 1α-hydroxylase, resulting in 1,25-dihydroxy-vitamin D [1,25 (OH)$_2$D], which is a biologically active hormone. Renal synthesis of 1,25 (OH)$_2$D is stimulated by the parathyroid hormone (PTH), and inhibited by the fibroblast growth factor. Serum calcium and phosphorus levels also stimulate the formation of active VD (3, 4) (Fig. 1).

The classic effect of VD is the homoeostasis of calcium and phosphorus, promotion of osteoid mineralization and thereby the maintenance of bone health (3). However, many tissues in the body (including the placenta, prostate, breast, colon, lung, bone, parathyroids, pancreas, vascular walls and the immune system) have receptors to bind the active form of VD, and many of them contain an enzyme to convert 25(OH)D to 1,25(OH)$_2$D. Locally produced 1,25(OH)$_2$D serves as an autocrine/paracrine factor that is essential for cell proliferation, differentiation, apoptosis, inhibition of angiogenesis and immunomodulation even early in development (3, 4, 10). For these reasons, it has been suggested that VD also has an extra-skeletal role – a non-classic action; for example, it act to affect the innate and acquired immune response, reduce cancer cell proliferation, influence the cardiovascular function and blood pressure, as well as hormone secretion, including the promotion of insulin secretion (3, 4, 10).

**Classification of vitamin D status**

The standard method for assessing the VD status in the body is to determine the 25(OH)D serum levels. 25(OH)D is a stable metabolite of serum VD, with a half-life of about 3 weeks, and it is the most appropriate indicator of VD status (10). Recently the criterion for the optimal VD status has been defined as the concentration of 25(OH)D
necessary to achieve not only adequate bone health, but also the health of the entire organism (11). Recent recommendations by the Endocrinological Association stressed that the optimal concentration of 25(OH)D is above 75 nmol/L, when the serum levels of VD and PTH are in equilibrium (12). Thus, a 25(OH)D concentration lower than

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Fig. 1. Vitamin D metabolism.
Vitamin D is either produced in the skin by exposure to UVB radiation or is ingested in the diet. Vitamin D (D represents vitamin D₂ or vitamin D₃) is converted by the vitamin D-25-hydroxylase (25-OHase) in the liver to 25(OH)D. 25(OH)D is converted in the kidneys by 1-OHase to 1,25(OH)₂D. Once formed, 1,25(OH)₂D enhances intestinal calcium and phosphorus absorption, and stimulates the expression of RANKL on the osteoblasts to interact with its receptor RANK on preosteoclasts to induce mature osteoclastic activity, which releases calcium and phosphorus (HPO₄²⁻). In addition, 1,25(OH)₂D inhibits the renal 1-OHase and stimulates the expression of the renal 25(OH)D-24-hydroxylase (24-OHase). The induction of the 24-OHase results in the destruction of 1,25(OH)₂D into a water-soluble inactive metabolite calcitroic acid, PreD₃, previtamin D. (By Holic MF J Clin Invest 2006). Adapted with permission from Holic MF.
50 nmol/l (<20 ng/ml or <50 nmol/l) is recognized as VD deficiency, suboptimal status (20–30 ng/ml (50–75 nmol/l)) and the target concentration for optimal VD effects [30–50 ng/ml (75–125 nmol/l)] (13). This level coincides with the outcome of pregnancy and the health of the foetus, indicating that this classification may also be appropriate during pregnancy and lactation (14). However data from the Hollis study suggest that a circulating 25(OH)D level of approximately 100 nmol/l (40 ng/ml) is required to optimize production of 1,25(OH)₂D during human pregnancy (15).

Prevalence of VD deficiency during pregnancy

VD deficiency is being increasingly recognized as a global epidemic that affects children, adolescents, adults and the elderly. The prevalence of VD deficiency is rising, mainly due to the modern lifestyle and malabsorption disorders (14, 16). The scientific literature offers no evidence supporting the benefits of population screening for determining VD status. According to the current recommendations, the level of 25-hydroxy VD is tested in individuals with diseases such as rickets, osteomalacia, osteopenia or osteoporosis, chronic kidney disease, liver failure, malabsorption, and obesity, in patients treated with anticonvulsant and/or antiretroviral drugs and glucocorticoids, and in pregnant and lactating women (14, 16). The prevalence of VD deficiency in pregnancy varies between 20 and 84% (17). A number of recent studies, which have estimated dietary intake of VD during pregnancy in different geographical locations, have reported significant differences in VD status, depending on the latitude, season, sun exposure and VD intake. In a Slovenian study, VD deficiency and insufficiency were found in 14 and 41% of pregnant women, respectively, and optimal levels were found in less than half of the study population (18).

Consequences of VD deficiency during pregnancy

There are a number of plausible biological pathways through which VD could influence maternal and foetal health during pregnancy. VD has important immune-modulating properties, helping to establish a proper maternal immune response to the placenta, reducing obstetrics risks associated with VD deficiency (17, 19) (Fig. 2).
**Preeclampsia/gestational hypertension**

One of the complications most closely associated with VD deficiency during pregnancy is preeclampsia (PE). Compared with normal pregnancies, PE is characterized by marked changes in VD and calcium metabolism, and already in the early 1990's, the role of VD in the pathogenesis of PE was hypothesized (18). Many observational studies have found an increased risk of preeclampsia in women with low VD status (20-23). Women with PE are known to have lower circulating 25(OH)D₃ levels than normotensive pregnant women (20-23).

In nested case–control studies, VD deficiency in pregnancy <50 nmol/l of 25(OH)D₃ was associated with almost fourfold odds of severe PE (20), and VD deficiency <37.5 nmol/l was even associated with a fivefold risk of developing PE (20). Bodnar et al. (21) showed that 25(OH)D₃ deficiency before 22 weeks of gestation is an independent risk factor for the manifestation of PE. Additionally, many randomized controlled trials, which have been undertaken to determine the impact of VD supplementation in PE patients, concluded that VD supplementation reduces the risk of PE (23). The mechanism suggested is the regulation of maternal and placental immunological and inflammatory responses, as has been shown in experimental models (24).

In syncytiotrophoblasts from preeclamptic pregnancies, the expression and activity of 1α-hydroxylase are restricted, suggesting an important role for VD at the placental site of the disease, and endorsing the role of VD in the regulation of the target genes associated with implantation, trophoblast invasion, and implantation tolerance (24, 25).

**Gestational diabetes mellitus**

VD plays a role in glucose homeostasis by multiple mechanisms: it regulates calcium levels, which in turn regulate insulin production and secretion by the endocrine pancreas (26, 27). It also improves the sensitivity to insulin of the target cells, such as adipose tissue, liver and skeletal muscles (26). Through its role in the regulation of immune cells, it protects β cells from detrimental immune attacks and even enhances their function (27). The role of VD in the development of pregnancy-related diabetes has been the subject of numerous studies, and it has been shown that VD deficiency affects the altered glucose homeostasis during pregnancy (28). Eight observational studies have recently been conducted to study the association of VD with gestational diabetes mellitus (GDM), and reported an inverse relationship between VD status in early pregnancy and risk of gestational diabetes (23). In a cohort of pregnant women whose VD values were measured before the gestational age of 16 weeks, an increase in the level of VD was associated with a reduced risk for the development of hyperglycaemia at the gestational age of 24-28 weeks (29). In contrast, some studies failed to establish a role of VD in the prevention of GDM (23). Although VD deficiency, or the dysfunction of VD receptors, relates to the pathogenesis of type 1 and type 2 diabetes, its role in GDM remains inconclusive.

**Preterm delivery**

One of the most common causes of preterm delivery is infection. Thus, through its role in anti-inflammatory pathways via nuclear factor-κB inhibition, VD could play a role in decreasing the incidence of infections and thereby preterm births (30). However, the exact role of VD in the pathogenesis of preterm birth has not yet been clearly defined. Many observational and interventional studies have investigated the association, with a lack of consistency in their results. Bodnar et al. (31) conducted a case control study with 1,126 cases and 2,327 controls, and found a protective association of VD sufficiency and preterm birth, after adjusting for the confounding factors. McDonnell et al. (32...
found a clear association between maternal 25(OH)D concentration and preterm birth risk in the general obstetrical population. Thus women with a 25(OH)D concentration of ≥40 ng/ml had a 62% lower risk of preterm birth compared to those with concentrations <20 ng/ml. These findings support the inverse association between maternal 25(OH)D and PTB risk found in the Hollis and Wagner et al. (33) randomized clinical trials, as well as epidemiological studies. However, other studies did not support these findings. A meta-analysis carried out by Pérez López et al. (34) showed conflicting reports on the role of vitamin D and the risk of preterm birth.

**Other impacts of VD status during pregnancy**

VD deficiency during pregnancy has been also significantly associated with an increased incidence of bacterial vaginosis (35). The incidence of bacterial vaginosis during pregnancy is important because it is associated with adverse outcomes, such as premature rupture of the membranes, premature delivery and the incidence of postpartum endometritis. In addition, a higher level of parturition by Caesarean section has also been reported in pregnant women with VD deficiency, although some other studies indicate that the method of delivery is independent of maternal VD status (36).

**The influence of maternal VD status on offspring health**

Numerous studies have shown that there is a positive correlation between the cord blood levels of 25(OH)D in the newborn and the maternal VD status, and that it represents approximately 60 to 89% of the maternal value (1). Therefore, maintaining optimal VD intake during pregnancy is essential to prevent insufficiency in foetal, neonatal and early childhood development. Physiologically active metabolite 1,25(OH)₂D does not cross the placenta (1, 37). Foetal kidneys and placenta provide the foetal circulation with 1,25(OH)₂D by expressing CYP27B1. Hormones involved in foetal growth also influence CYP27B1 activity, including insulin like growth factor 1, human placental lactogen, PTH-related protein (PTHrP), estradiol, and prolactin (38).

**Bone health in newborns**

Lack of VD during pregnancy may have an impact on foetal and neonatal bone mass and bone mineralization. In their study, Weiler et al. (39) detected a lower bone mineral content per kilogram of body weight in newborns with lower levels of 25(OH)D in the cord blood. Viljakainen (40) studied the relationship between the levels of 25(OH)D in the first trimester of pregnancy and the neonatal bone mass, measured by peripheral quantitative computed tomography in the postpartum period. He found that the newborns of mothers whose average concentration of 25(OH)D was below 43 nmol/l exhibited lower tibia bone mass than the newborns of mothers whose average concentration of 25(OH)D was higher than 43 nmol/l.

Studies have shown that chronic VD deficiency in the mother has a negative effect on the development of the skeleton of the foetus. Using three-dimensional ultrasound, Mahon et al. (41) measured the length and a cross section of the distal femur in foetuses between 19 and 34 weeks of pregnancy. They found that in the 34th week of pregnancy, the 25(OH)D levels were inversely proportional to the femoral splaying index (distal cross-section / femur length). These results suggest that the maternal VD status already affects the bone morphology in the 19th week of pregnancy (41).
Hypocalcaemia in newborns

In the view of what has been said above regarding the metabolism of VD and calcium during pregnancy, it is assumed that the concentration of calcium in a foetus and neonate depends on the maternal VD status (42). Studies evaluating how the supplementation of VD to mothers during pregnancy affects the homeostasis of calcium in the newborn have shown higher serum calcium levels within the first week after birth, while the decline in the incidence of hypocalcaemia was more prevalent with 25(OH)D cord blood levels below 10 nmol/l. Neonatal hypocalcaemia was only observed in the group receiving a placebo (43). The main clinical signs of hypocalcaemia in newborns are jitteriness and generalized convulsions, although asymptomatic hypocalcaemia has also been reported. Newborns may also be lethargic, eat poorly, vomit, and have abdominal distention. The degree of irritability does not appear to correlate with serum calcium values.

Anthropometry in newborns

The association between birth weight and maternal VD status or intake remains inconclusive. While some observational and interventional studies have found improvement in birth weight with maternal VD supplementation or improved VD status, several others did not confirm these findings (23). A recent longitudinal study (multi-ethnic cohort of 3,730 pregnant women) in The Netherlands, reported an increased risk for reduced foetal growth in women whose 25(OH)D concentration in early pregnancy was lower than 30 nmol/l (44). This study has also found a link between a genotype for the VD receptor and the risk of reduced foetal growth. Morley et al. (45) showed a connection between a FokI genotype for the VD receptor and the newborn’s size. FF or ff polymorphisms were associated with a lower birth weight if the mother had insufficient levels of VD, while neonates with ff polymorphism did not exhibit this association. Also, FokI polymorphism did not affect the size of the newborn if the maternal VD status was adequate (45). In addition, Robinson at al. reported lower maternal 25(OH)D₃ concentrations in 56 women with early onset PE and small for gestational age (SGA) infants, vs. infants with normal foetal growth, suggesting the impact of VD on foetal growth through placental mechanisms (24). On the other hand, both studies - the longitudinal AVON, as well as the Southampton study - failed to show a link between the level of 25(OH)D and the birth weight and length of children (46, 47). A recently published meta-analysis of RCTs by Pérez-López et al. (22) also established the positive outcomes of VD supplementation in pregnancy regarding offspring birth weight and length. In summary, the current studies suggest that the risk of SGA and smaller birth size is possible when pregnant women are VD deficient or insufficient. However, the associations of maternal VD deficiency with early postnatal growth outcomes and infant adiposity are less clear, with inconsistent directionality and a smaller number of studies.

Other impacts of maternal VD status on the offspring’s health

Consistent in vitro findings and observational clinical data have found an association between VD status in the cord blood and the lower incidence of respiratory tract infections in the first year of life. Cord blood 25(OH)D concentrations <75 nmol/l have also been linked to infantile wheezing and eczema, possibly due to adverse consequences on the early immune development of the foetus. However, there is still not enough evidence to show whether VD supplementation increases or decreases the risk of allergies, and therefore this requires further exploration (23, 48). Re-
One study found no association between maternal VD status during pregnancy and neurocognitive function, a recent larger-sized study linked maternal serum 25(OH)D levels during pregnancy with language development in the offspring (23, 48).

**Conclusion**

The awareness of the importance of adequate VD intake during pregnancy, to achieve the optimal health of the mother and her offspring, is growing rapidly. In addition to the traditional role of VD, which is associated with maintenance of calcium homeostasis and bone health, new findings suggest that the mother’s VD status plays an important role in ensuring the proper development of the placenta and foetus, and a proper immune response during pregnancy, as well as in achieving the optimal health of the offspring. The aetiology of the various maternal and foetal outcomes is complex and multifactorial, with many confounding factors. Most new insights on the importance of VD are based on observational studies, and require confirmation of the results by well-designed, randomized, placebo-controlled studies in which the studied group will receive additional VD during pregnancy (49). Determining the benefits of VD supplementation in pregnancy would require further evaluation through large, multicentre, double-blind randomized controlled clinical trials, with the focus on specific adverse pregnancy outcomes.

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