Review

Skin Pharmacology and Physiology

Skin Pharmacol Physiol 2018;31:74–86 DOI: 10.1159/000485132 Received: March 10, 2017 Accepted after revision: November 7, 2017 Published online: January 6, 2018

Vitamin D and the Pathophysiology of Inflammatory Skin Diseases

Meenakshi Umar Konduru S. Sastry Fatima Al Ali Moza Al-Khulaifi

Ena Wang Aouatef I. Chouchane

Research Department, Division of Translational Medicine, Sidra Medicine, Doha, Qatar

Keywords

Vitamin D · Skin · Keratinocytes · Psoriasis · Atopic dermatitis

Abstract

Background: Vitamin D is a secosteroid, which was initially known for its skeletal role; however, in recent years, its functions in different organs have been increasingly recognized. In this review, we will provide an overview of vitamin D functions in the skin physiology with specific focus on its role in certain inflammatory skin conditions such as psoriasis and atopic dermatitis. *Methods:* A comprehensive literature search was carried out in PubMed and Google Scholar databases using keywords like "vitamin D," "skin," "atopic dermatitis," and "psoriasis." Only articles published in English and related to the study topic were included in this review. **Results:** Vitamin D is integrally connected to the skin for its synthesis, metabolism, and activity. It regulates many physiological processes in the skin ranging from cellular proliferation, differentiation, and apoptosis to barrier maintenance and immune functions. Vitamin D deficiency is asso-

KARGER

© 2018 S. Karger AG, Basel

E-Mail karger@karger.com www.karger.com/spp ciated with the risk of psoriasis and atopic dermatitis, and several clinical/observational studies have suggested the beneficial effect of vitamin D in the therapy of these 2 inflammatory skin disorders. **Conclusions:** Vitamin D exerts a pleiotropic effect in the skin and could be an important therapeutic option for psoriasis and atopic dermatitis.

© 2018 S. Karger AG, Basel

Introduction

The human skin acts as site of synthesis of vitamin D and also as target organ for the biologically active form of this vitamin. Vitamin D affects multiple functions in the skin ranging from keratinocyte proliferation, differentiation, and apoptosis to barrier maintenance and immunoregulatory processes [1]. Also, vitamin D is being considered as a therapeutic option for many skin pathologies [2]. In this review, we will discuss the nonclassical function of vitamin D in the skin and will evaluate its role in certain inflammatory skin conditions using atopic dermatitis (AD) and psoriasis as examples.

Aouatef I. Chouchane Research Department, Division of Translational Medicine, Sidra Medicine Al Luqta Street, PO Box 26999 Doha (Qatar) E-Mail achouchane@sidra.org

Main Structural and Functional Molecules in the Skin

The skin acts as first line of defense against infections. It consists of mainly 3 layers, the epidermis, dermis and hypodermis, and associated with it are several appendages like hair follicles, eccrine sweat glands, sebaceous glands, and apocrine glands. The epidermis consists of many cells like keratinocytes, melanocytes, Langerhans cells (a specialized subset of myeloid dendritic cells, DCs) and Merkel cells, among which keratinocytes account for 95% of the total epidermal cells. There are 4 distinct epidermal layers, each composed of keratinocytes at various differentiation stages [1]:

- 1. stratum basale: it consists of columnar, proliferating keratinocytes with an extensive network of keratins K5 and K14;
- 2. stratum spinosum: in this layer, keratinocytes initiate differentiation through synthesis of K1 and K10 keratins, involucrin, and enzyme transglutaminase;
- 3. granular layer: it is characterized by keratinocytes rich in electron-dense keratohyalin granules containing late differentiation markers like profilaggrin (precursor of filaggrin), and loricrin; it also consists of lipidfilled lamellar bodies that empty their contents into the intercellular spaces between the stratum granulare and stratum corneum and contribute to the permeability barrier;
- 4. stratum corneum (SC): the uppermost layer, consists of terminally differentiated dead cells known as corneocytes. The plasma membrane of corneocytes is replaced by an insoluble protein layer called "cornified envelope," made of structural proteins like involucrin, loricrin, filaggrin, and small proline-rich protein cross-linked by transglutaminase.

Filaggrin is a particularly important molecule in the SC, as it facilitates the aggregation of keratin filaments of the cytoskeleton into bundles, consequently collapsing corneocytes into flattened disks. Also, it contributes to the hydration of the SC by proteolysing into pyrrolidine carboxylic acid and transurocanic acid in conditions of low water content [3]. The constant thickness of the epidermis is maintained by the fine balance between basal cell proliferation and corneocyte desquamation. The desquamation process starts with the degradation of corneodesmosomes (modified desmosomes present in the SC) and is controlled by a number of proteases and their inhibitors. The human kallikrein (KLK)-related peptidases including the KLK5, KLK7, and KLK14 are the prominent proteases involved in desquamation. The lymphoepithelial Kazaltype 5 serine protease inhibitor is an important protease

inhibitor encoded by the *SPINK5* gene which has confirmed activity against the members of the KLK family [4].

Vitamin D: Synthesis and Functions

Vitamin D is a fat-soluble vitamin that occurs in 2 main forms: ergocalciferol (vitamin D₂) produced by plants and cholecalciferol (vitamin D₃) derived from animal-based foods. The major source of vitamin D in humans is the cutaneous synthesis in the presence of sunlight. The exposure of 7-dehydrocholesterol (7-DHC) to ultraviolet radiation B (UVB) of wavelength 290-315 nm results in the formation of previtamin D in the skin, which is thermally isomerized to the stabler vitamin D (cholecalciferol). The vitamin D, whether synthesized in the skin or obtained from diet, undergoes 2 hydroxylation reactions: first in the liver by vitamin D 25-hydroxylase (CYP2R1) enzyme to form 25-hydroxyvitamin D, 25(OH)D, also known as calcidiol and then in the kidney by 1a-hydroxylase (CYP27B1) to form an active metabolite, 1,25-dihydroxyvitamin D, 1,25(OH)₂D, also known as calcitriol. Both 25(OH)D and 1,25(OH)₂D may be metabolically inactivated through hydroxylation by 24-hydroxylase (CYP24A1) [5]. The levels of vitamin D in serum are tightly regulated by a feedback mechanism of calcium, phosphorus, parathyroid hormone, fibroblast growth factor and vitamin D itself [6, 7]. The vitamin D status is evaluated by measuring the serum 25(OH)D level, which is its major circulating form. According to the US Endocrine Society guidelines, vitamin D deficiency is defined as a serum level of 25(OH)D below 20 ng/mL (50 nmol/L) and vitamin D insufficiency as a serum 25(OH)D level between 21 and 29 ng/mL (52.5-72.5 nmol/L) [8].

The function of vitamin D was for a long time considered to be the maintenance of a normal skeletal architecture through calcium and phosphorus homeostasis, but in the last few decades, the extraskeletal effects of vitamin D became apparent, and its roles in the regulation of cell proliferation, differentiation, apoptosis, and in the immune modulation are increasingly recognized [9, 10]. These actions of vitamin D are mediated by the vitamin D receptor (VDR), which after activation interacts with retinoid X receptor (RXR) to form a heterodimeric complex. The VDR-RXR complex is recruited to the vitamin D response elements (VDREs) in the promoter of target genes to regulate their expression. This process is described as the genomic action of vitamin D, in contrast to the nongenomic action which is the direct effect that vitamin D has on several signaling pathways.

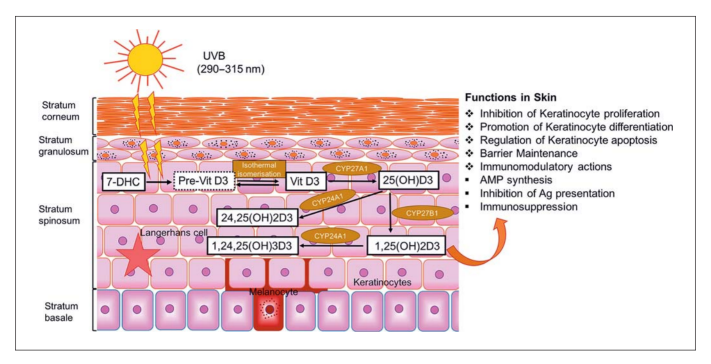


Fig. 1. Summary of vitamin D_3 pathway and functions in the human skin. Vitamin D_3 (Vit D_3) is synthesized in the skin from its precursor 7-DHC under the influence of UVB and metabolized to its active form, 1,25(OH)₂D₃ through 2 subsequent hydroxylation reactions by CYP27A1 and CYP27B1 enzymes. It is rendered inactive through the catabolic enzyme CYP24A1. 7-DHC, 7-dehydro-

Role of Vitamin D in Skin Physiology

Vitamin D plays a vital role in the skin: the keratinocytes are not only a source of vitamin D, but also a responder to its active form [1]. They are the only cells in the body that can synthesize vitamin D from its precursor 7-DHC, and which are equipped with the entire enzymatic machinery (CYP27A1 and CYP27B1) necessary to metabolize vitamin D into its active metabolite $1,25(OH)_2D$. Keratinocytes also express VDR, thus they respond in an autocrine and paracrine manner to the active form of vitamin D. The entire pathway of vitamin D₃ in human skin is shown in Figure 1.

Vitamin D and Epidermal Differentiation and Proliferation

Vitamin D affects the proliferation and differentiation of the skin either directly or through its interaction with calcium. Many in vitro studies have shown a dose-dependent effect of vitamin D on keratinocyte proliferation and differentiation. At low concentration $(10^{-9} \text{ M or less})$, 1,25(OH)₂D₃ was found to enhance keratinocyte prolif-

cholesterol; 25(OH)D₃, 25-hydroxyvitamin D₃; 1,25(OH)₂D₃, 1,25-dihydroxyvitamin D₃; 24,25(OH)₂D₃, 24,25-dihydroxychole-calciferol; 1,24,25(OH)₃D₃, 1,24,25-trihydroxycholecalciferol; CYP27A1, 25-hydroxylase; CYP27B1, 1 α -hydroxylase; CYP24A1, 24-hydroxylase; AMP, antimicrobial peptide; Ag, antigen; UVB, ultraviolet radiation B.

eration, while at high concentration (greater than 10^{-8} M), it inhibited the proliferation and promoted the differentiation [11, 12]. Several other factors like density of cells, calcium concentrations and presence or absence of serum influence the effect of vitamin D on in vitro keratinocyte proliferation [13]. The antiproliferative action of vitamin D on keratinocytes is mediated by the decreased expression of c-myc and cyclin D and by the increased expression of the cell cycle inhibitors p21^{cip} and p27^{kip} [1, 14]. 1,25(OH)₂D promotes keratinocyte differentiation through an increased synthesis of structural components (involucrin, transglutaminase, loricrin, and filaggrin) of the cornified envelope [14, 15]. The effect of vitamin D in the differentiation is also in part mediated by the (1) elevation of intracellular calcium levels caused by calcium receptor stimulation, (2) increased phospholipase $C-\gamma_1$ expression, and (3) enhanced formation of ceramides [15–17]. Vitamin D may also directly regulate the keratinocyte differentiation through interaction with VDR. This is evidenced by the fact that VDR knockout mice show reduced epidermal differentiation and exhibit low levels of involucrin, profilaggrin, and loricrin [18]. The process of vitamin D-mediated epidermal differentiation through VDR is sequential and requires selective binding of VDR to 2 major coactivators: vitamin D receptor-interacting protein (DRIP) and steroid receptor coactivator (SRC). It was observed that DRIP205 is predominantly expressed in proliferating keratinocytes and, as the cells differentiate, the expression of DRIP205 goes down, while the expression of SRC3 increases [19]. It was demonstrated that calcium also regulates the expression of these 2 coactivators and interacts with VDR for the differentiation of keratinocytes [20].

Vitamin D and Barrier Function

Another aspect of keratinocyte proliferation and differentiation is the maintenance of a proper epidermal barrier. Previous studies have shown that topical application of calcitriol (1,25[OH]₂D) restores the permeability barrier which was disrupted by application of corticosteroid or sodium lauryl sulfate [21, 22]. Vitamin D mediates its effect on the epidermal barrier by enhanced synthesis of structural proteins of the cornified envelope. Additionally, 1,25(OH)₂D regulates the processing of the longchain glycosylceramides essential for lipid barrier formation. Oda et al. [23] have shown that VDR knockout mice display a defective permeability barrier due to the reduced production of glucosylceramide and its decreased transport into the lamellar bodies, resulting in a lower lipid content in these bodies.

Vitamin D and Keratinocyte Apoptosis

The effect of vitamin D on the keratinocyte apoptosis is dose dependent, similar to its effect on cellular proliferation. At physiological concentrations, vitamin D prevents apoptosis triggered by various proapoptotic stimuli like ceramide, UV radiation, TNF- α , etc., while at high concentrations it induces apoptosis in keratinocytes [24]. The antiapoptotic or cytoprotective effect of vitamin D is shown to be mediated by sphinosine-1-phosphate. Other mechanisms are also reported to be responsible for the antiapoptotic effect of vitamin D like the activation of MEK/ERK and PI3K/Akt signaling pathways, and the increased ratio of antiapoptotic protein (Bcl-2) to proapoptotic protein (Bad and Bax) [25].

Vitamin D and Skin Immune Functions

The skin innate immune system comprises physical barrier structures like SC, immune cells (like neutrophils, monocytes, macrophages, DCs, natural killer [NK] cells, etc.) and antimicrobial peptides (AMPs). The cutaneous synthesis of AMPs is the primary protection mechanism of the skin against environmental insults or microbial invasion. Many resident cells of the skin (like keratinocytes, sebocytes, eccrine gland cells, and mast cells) and circulating cells recruited to the skin (like neutrophils and NK cells) contribute to the synthesis of AMPs in the skin [26, 27]. More than 20 proteins with antimicrobial function are recognized in the skin; however, β-defensin and cathelicidins are the 2 main groups of skin AMPs [26]. Defensins are classified in 3 subfamilies based on cysteinedisulfide pairing between β -sheet structure – α , β , and θ – of which only β -defensin is appreciably expressed in the skin. Humans have a single cathelicidin gene which encodes the inactive peptide hCAP18, which after cleavage generates the mature peptide LL-37. Cathelicidin and β-defensin mediate antimicrobial activity either directly by disrupting the bacterial cell membrane and viral envelope or indirectly by affecting various signaling pathways in the cells to initiate a host response. These 2 AMPs are also reported to promote keratinocyte proliferation and migration through EGFR signaling and STAT activation (necessary for skin wound healing), to stimulate cytokine or chemokine release through stimulation of G proteincoupled receptors and to induce IL-8 secretion through the ERK p38/MAPK pathway in mast cells and keratinocytes [28].

The level of AMPs is low in intact skin, and it increases following barrier disruption or infection. One of the possible ways it is done is through enhanced CYP27B1 expression, subsequent to skin insult, which increases the local synthesis of active vitamin D. Schauber et al. [29] have shown that following skin injury, TLR-2 is increased which in turn increases the level of cathelicidin through a vitamin D-dependent mechanism. Similarly, many studies have shown an increased expression of hCAP18/ LL-37 and defensin after 1,25(OH)₂D₃ treatment in keratinocytes and sebocytes [30–33]. Cathelicidin and β defensin are direct transcriptional targets of vitamin D, with cathelicidin being induced by binding of the 1,25(OH)₂D-VDR complex to the VDRE in the promoter region of the gene; however, β -defensin requires nuclear factor κ B along with the 1,25(OH)₂D-VDR complex for its transcription [34]. Vitamin D is also reported to regulate the AMP synthesis by mechanisms other than the direct transcriptional activation. The activity of cathelicidin and other AMPs in human skin is controlled through an enzymatic processing by serine proteases KLK5 and KLK7 [35]. Morizane et al. [36] showed that 1,25(OH)₂D₃ could affect the production of AMPs in the skin by regulating synthesis and protease activity of KLK5 and KLK7. In another study, Dai et al. [33] showed that the induc-

Downloaded by: 24.113.93.62 - 1/8/2018 6:38:06 PM tion of cathelicidin and β -defensin HBD-3 expression by 1,25(OH)₂D₃ is regulated by peroxisome proliferator-activated receptor- γ through AP-1 and p38 activity.

Besides regulating AMP synthesis in the skin, $1,25(OH)_2D_3$ and calcipotriol (an analog of vitamin D) mediate an immunosuppressive action in the skin through decreased antigen presentation either directly by affecting Langerhans cells or indirectly by modulating cytokine production by keratinocytes [37, 38]. Recently, many studies suggested that calcipotriol mediates tolerance or immunosuppression in the skin through induction of CD4+CD25+ T regulatory (T_{reg}) cells which prevents subsequent antigen-specific CD8+ T-cell proliferation and IFN-y production [39, 40]. Skin-homing cutaneous lymphocyte-associated antigen (CLA+) memory T cells preferentially home to cutaneous sites for host defense against pathogens. CCR10 is a chemokine receptor that is preferentially expressed by skin-homing CLA+ T cells which facilitate their entry into cutaneous sites by interacting with skin-associated CLC27 antigen. Studies investigating the effect of vitamin D on homing of memory T cells to the skin are contradictory. While some studies suggested that 1,25(OH)₂D₃ and its analogs prevent skin T-cell infiltration by downregulating the expression of CLA [41, 42], other studies showed that $1,25(OH)_2D_3$ induces CCR10 receptor expression on T cells promoting their homing to cutaneous sites [43-45]. Recently, a study has shown that seasonal variation in vitamin D level affects the skin-homing receptor expression with increased levels of CLA during the summer [46].

Role of Vitamin D in Certain Inflammatory Skin Diseases

Psoriasis

Psoriasis is a chronic multifactorial inflammatory disease where the immune dysregulation plays a major role by involving a crosstalk between the innate and adaptive immune system. There is an increased infiltration of innate immune system effectors like plasmacytoid dendritic cells (pDCs), myeloid dendritic cells (CD11c+ mDCs), neutrophils and NK cells, and abnormally high levels of AMPs (like β -defensins, S100 proteins or LL-37) in psoriatic lesions. It is suggested that a complex of host DNA and LL-37 acts as potent trigger for IFN- α production by pDCs and provides a mechanism of initiation of intolerance to self-DNA [47]. IFN- α derived from pDCs is supposed to drive the early inflammatory cascade in psoriasis by activating "quiescent" autoimmune T cells into pathogenic effectors through promoting activation or maturation of mDCs [48]. On activation, a subset of CD11c+ mDCs, known as TIP-DCs, expresses an increased level of TNF- α and inducible nitric oxide synthase enzyme (generates nitric oxide to induce vasodilation and inflammation) [49]. Additionally, another subset of mDCs produces IL-20 to enhance keratinocyte activation and proliferation, and IL-23 and IL-12 to activate a specific subset of T cells [50]. Neutrophils and NK cells recruited in psoriatic lesions further add to the inflammatory milieu of psoriasis through secretion of AMPs and proinflammatory cytokines [51, 52].

Psoriatic lesions are also characterized by an increased infiltration and activation of T cells particularly CD4+ T helper 1 (Th1) and CD8+ cytotoxic T cells, which predominantly secrete type 1 cytokines like TNF-a and IFN-γ. These lesions are also enriched in other types of T cells like IL-17-producing T cells and NK T cells. It was observed that IL-23 secreted by mDCs and other leukocytes induces the differentiation of naïve T cells into type 17 helper T cells (Th17) and type 17 cytotoxic T cells, both secrete IL-17, IL-17F, and IL-22 cytokines [53]. After activation, NK T cells also secrete Th1, Th2, and Th17 cytokines [54]. These type 17 cytokines together with IFN- γ and TNF-a result in activation and proliferation of keratinocytes. Thus, in response to cytokines secreted from DCs and T cells, keratinocytes become activated and produce AMPs, proinflammatory cytokines (IL-1, IL-6 and TNF- α), chemokines (CXCL8 through CXCL11 and CXCL20) and S100 proteins (S100A7-9) [55]. These soluble mediators act as chemoattractants for neutrophils and other immune cells. Therefore, a feedback loop exists between keratinocytes and infiltrating immune cells, which maintains a constant deregulated inflammatory process, characteristics of psoriatic disease. Unrestrained function of T cells in psoriasis may also be due to the dysfunction of T_{reg} cells. In fact, some studies have shown that T_{reg} cells isolated from psoriatic patients have a decreased suppressive function [56, 57], others demonstrated that they produce IFN-γ, TNF-α, and IL-17, suggesting a switch of their function from suppressive to proliferative [58].

Role of Vitamin D in Psoriasis

Vitamin D plays a critical role in psoriasis, and this is evidenced in many studies which reported either a deficiency or insufficiency of serum vitamin D in psoriatic patients [59–61]. Several case-control studies have shown significant lower levels of serum 25(OH)D in psoriatic patients compared to controls and reported an inverse correlation between serum 25(OH)D and the severity of the disease [62–66]. However, in a population-based screening, Wilson [67] showed that vitamin D deficiency is not common in psoriatic patients and that there is no significant difference in serum 25(OH)D levels in subjects with or without psoriasis. The 25(OH)D level varies with several factors, including race, dietary intake, and UV light exposure, therefore results of studies on vitamin D need cautious interpretation.

Vitamin D treatment may be effective in resolving psoriasis symptoms, and this is confirmed by many clinical studies. Finamor et al. [68] showed that psoriasis patients, who were receiving 35,000 IU of vitamin D₃ once daily for 6 months, had significant improvement in psoriasis area severity index score (PASI) with a marked increase in their serum 25(OH)D level. Several clinical trials have also demonstrated an excellent efficacy and safety profile of vitamin D analogs like calcipotriol, tacalcitol, and maxacalcitol in the treatment of psoriasis [69–71].

NB-UVB (narrow-band ultraviolet B light) and UVA/ UVB phototherapy, widely used in the treatment of psoriasis, are thought to mediate its beneficial effect in part by elevating the serum 25(OH)D level [72-74]. A clinical trial compared the efficacy and safety of various treatment regimens for psoriasis (calcipotriol monotherapy, NB-UVB phototherapy alone and combination of calcipotriol and NB-UVB) and demonstrated that the combination of calcipotriol and NB-UVB twice a week was superior to other treatment regimens in rapidly reducing the PASI score of patients [75]. The combination of vitamin D or its analogs and corticosteroid is also reported to be more effective than either of their monotherapy because of their complementary actions. In combination treatment, vitamin D may counteract the steroid-induced skin atrophy by restoring the epidermal barrier, while corticosteroid may reduce the perilesional skin irritation caused by vitamin D analogs [76-78].

Vitamin D exhibits an inhibitory effect in psoriasis through a multitude of ways. pDCs, which are supposed to initiate the inflammatory cascade in psoriasis, express transcriptionally active VDR and the vitamin D-metabolizing enzymes CYP27B1 and CYP24A1. It was shown that vitamin D treatment impairs the capacity of pDCs to induce T-cell proliferation and IFN- γ secretion [79]. Vitamin D is also supposed to affect the Th17 pathway: it was observed that application of vitamin D and its analogs on psoriatic lesions significantly decreased the infiltration of Th17 cells in the skin and inhibited their ex vivo expansion [80, 81]. In other studies, vitamin D was reported to suppress inflammatory cytokines like IL-12/23

Vitamin D not only modulates or suppresses inflammation in psoriasis; it also rectifies the abnormal epidermal function related to this condition. It was demonstrated that deletion in late cornified envelope genes, LCE3B and LCE3C, located within PSORS4 is a genetic risk factor of psoriasis. A study by Hoss et al. [86] has shown that 1,25(OH)₂D upregulated the LCE proteins (LC3A-E) in keratinocytes and provided a mechanism of ameliorating psoriasis in patients with LCE defects. The expression of tight junction proteins like claudin, ZO-1, and occludin, which are reduced in psoriatic skin, is correlated with VDR status, pointing out the role of vitamin D in the regulation of tight junction proteins in psoriasis [87]. Furthermore, vitamin D topical use normalized the expression and topography pattern of integrins and other activation markers like ICAM-1, CD26 and HLA-DR, which were altered on psoriatic skin [88].

The role of VDR polymorphisms in the risk of psoriasis was studied in several populations, with contradictory results. Richetta et al. [89] showed that among 5 common VDR polymorphisms (A-1012G, FokI, BsmI, ApaI, and TaqI), the A-1021G polymorphism is associated with the risk of psoriasis in an Italian population. In another study, ApaI and a specific haplotype of 5 VDR polymorphisms were associated with the risk of psoriasis in a Chinese population [90]. In contrast, studies in Croatian and Egyptian populations did not find any role of VDR polymorphisms in psoriasis [91-93]. The meta-analysis of studies investigating the role of VDR polymorphisms in psoriasis also suggests their ethnic specific association [94, 95]. The VDR polymorphisms, besides conferring a risk of psoriasis, are also reported to modulate the response of psoriasis patients to different treatment regimens. Ryan et al. [96] showed that psoriatic patients with the VDR TaqI polymorphism had a shorter remission pe-

79

p40, IL-1 α , IL-1 β , and TNF- α , which were present in abnormally high levels in psoriatic skin [82, 83]. Psoraisin (S100A7) and koebnerisin (S100A15), induced by Th17 cytokines, synergistically act as chemoattractants and "alarmins" to amplify inflammation in psoriasis. Calcipotriol was found to suppress Th17-induced psoriasin and koebnerisin in psoriatic skin [84]. In an epidermal reconstructed model of psoriasis, Datta Mitra et al. [85] showed that 1 α ,25-dihydroxyvitamin D₃-3-bromoacetate, a vitamin D analog, has a more potent antiproliferative action compared to 1,25(OH)₂D₃. He showed that bromoacetate reverses the psoriasiform changes induced by IL-22 in the reconstructed epidermal model by inhibiting the expression of AKT1, MTOR, chemokines (IL-8 and RANTES) and psoriasin (S100A7).

riod when treated with NB-UVB. Similarly, other studies suggested a positive association of wild-type alleles of A-1012G, FokI and TaqI VDR polymorphisms with topical calcipotriol response [97, 98].

Atopic Dermatitis

AD is a chronic or relapsing skin disorder caused by complex interactions between genetic, immunological, and environmental factors; it is characterized by chronic inflammation, disruption of the epithelial barrier, immunological abnormalities and increased serum IgE.

Skin Barrier Defect in AD. The epidermis of AD patients displays a significant barrier disruption and transepidermal water loss, which sensitizes AD skin to allergen penetration, bacterial, fungal, and virus invasion or colonization and inflammation. Various mechanisms are responsible for the barrier defect in AD: (1) deficiency or defects in structural proteins (like filaggrin, involucrin, loricrin, keratin K5 and K16, etc.), epidermal proteases, and protease inhibitors, (2) alteration in SC pH, and (3) decrease in skin ceramides, which supports lipid barrier and water retention [99, 100]. So far, loss of function mutation in the filaggrin gene represents the most significant genetic factor in the predisposition to AD, although only a fraction of patients (between a few and 50% depending on the populations studied) carry filaggrin mutations [101, 102].

Immune Dysregulation in AD. The immune dysregulation in AD is biphasic, with an initial Th2 phase in acute lesions, and Th0 and Th1 dominant inflammation in chronic lesions. Thus, there is an increased level of IL-4, IL-5, and IL-13 (Th2 cytokines) in the acute phase lesions, while Th1 cytokines like IFN-y, GM-CSF, and IL-12 are predominant in the chronic disease. Th0 cells are transitory and can differentiate into Th1 or Th2 cells [99]. Beside alteration in Th cytokines, the majority of AD cases (approx. 80%) displays high serum IgE levels with specific IgEs to food allergens or aeroallergens [103]. The outcome of Th cells in AD lesions is regulated by several factors. Thymic stromal lymphopoietin secreted by keratinocytes in atopic skin primes DCs, which drives naïve Th cells towards Th2 polarization and induces production of the proallergic cytokines IL-4, IL-5, IL-13, and TNF-a [104]. The DCs observed in atopic lesions are mainly of myeloid origin and comprise 2 populations: Langerhans cells and inflammatory DCs. It was observed that Langerhans cells are involved in Th2 polarization, while inflammatory DCs promote Th1 polarization in chronic lesions [105, 106]. T_{reg} cells play an important role in AD. Many studies have shown a high T_{reg} (CD4+CD25+Foxp3+) population with normal immunosuppressive activity in the peripheral blood of AD patients, which is also found to be positively correlated with the severity of the disease [107–110]. However, when stimulated with *Staphylococcus* enterotoxin B, T_{reg} cells lost their immunosuppressive activity suggesting a mechanism of T-cell activation by *Staphylococcus aureus* in AD lesions [107, 108]. There are few contradictory reports which suggested either low frequency or absence of T_{reg} cells in the peripheral blood and in skin lesions of AD patients [111, 112].

In addition to a defective adaptive immune system, AD patients have dysfunction in various components of the innate immune system like skin barrier disruption, diminished recruitment of innate immune cells (NK cells, pDCs, neutrophils) to the skin, TLR2 defects and reduced secretion of AMPs [113, 114].

Role of Vitamin D in AD

The effect of vitamin D levels on the prevalence and severity of AD was the subject of a large number of studies which yielded heterogeneous results. Epidemiological studies have shown an increased AD prevalence in populations living in higher geographic latitudes, with lower sun exposure and consequently less vitamin D production [115, 116]. Also, in large population-based studies, it was observed that there is an increased likelihood of developing AD in individuals with either deficient or insufficient vitamin D levels [117, 118]. Many observational studies including a meta-analysis have shown that the serum vitamin D level is lower in children and adults with AD compared to controls, and reported an association between vitamin D deficiency and risk of atopic eczema [119–121]. Also, the severity of AD was found to be negatively correlated with the vitamin D level, with moderate and severe AD groups having lower vitamin D levels compared to the mild AD group; this finding was supported by the use of objective tools, such as the SCORAD (Scoring Atopic Dermatitis) index, which was found to be inversely correlated with vitamin D levels in AD patients [120-123]. However, there are some contradictory reports, which suggest either no role of vitamin D or a positive association of vitamin D levels with the risk of developing AD [124, 125]. The maternal vitamin D level seems also to impact the risk of developing AD in infants: while 2 studies suggested that a higher maternal intake of vitamin D could increase the risk of infantile eczema [126, 127], others observed that a lower vitamin D level during pregnancy induced a risk of AD in infants during early years of their life [128]. Studies on the association of cord

80

serum 25(OH)D levels with infant AD are also contradictory [129–131]. The common polymorphisms in VDR and vitamin D-metabolizing genes have been investigated for their role in AD susceptibility. The VDR BsmI polymorphism increased the risk of AD in a Turkish population, and a specific haplotype of VDR BsmI, ApaI, and TaqI polymorphisms was overrepresented in severe AD patients in a German population [132, 133]. In another report, among 6 common polymorphisms in CYP24A1 and CYP27B1, CYP24A1rs2248359C allele and a specific haplotype were associated with an increased risk of severe AD [134].

As the majority of the literature suggests vitamin D deficiency as prominent risk factor of AD, studies have been carried out to examine the effect of vitamin D supplementation on phenotypes of AD. Many clinical trials including their meta-analysis have shown that vitamin D supplementation results in significant improvement in AD severity (measured by SCORAD and Eczema Area and Severity Index) [121, 135–138]. Di Filippo et al. [138] suggested that vitamin D supplementation exerts its positive effect on AD by normalizing the altered Th1 and Th2 cytokines like IL-2, IL-4, IL-6, and IFN-y in AD patients. In another study, Drozdenko et al. [139] showed that the oral intake of vitamin D increases the frequencies of CD38+ B cells to enhance the B-cell receptor-mediated response and decreases the IFN-y and IL-17 T-cell cytokine response in vitamin D-deficient individuals. Additionally UVA and UVB phototherapy is widely used in AD treatment because of its effects in the T cell-mediated immune response, and it is suggested that the beneficial effect of UVA/UVB phototherapy is also mediated by the correction of the vitamin D deficiency or insufficiency in AD patients [72]. An increased IgE response to common environmental and food allergens is a common feature in AD. It was found that vitamin D has an inhibitory effect on the allergic response: treatment of vitamin D suppressed the IgE production by human B cells and dampened IgE-mediated mast cell activation in both in vitro and in vivo settings [140]. Other than the effect on the adaptive immune system, vitamin D supplementation ameliorates the AD lesions by restoring the epidermal barrier defects and correcting the deregulated innate immune response. In fact, Kanda et al. [141] observed that a low serum vitamin D₃ level correlated with low serum LL-37 in AD patients. Also, the topical application and oral supplementation of vitamin D upregulated the expression of LL-37 in lesional and nonlesional skin in AD patients [114, 142]. More recently, a clinical improvement, assessed by a lower AD severity score, was noted in

AD patients, concomitantly to the increase in the LL-37 level, after vitamin D supplementation [143]. Büchau et al. [144] suggested that the positive effect of vitamin D on AMPs could be mediated by the inhibition of the expression of Bcl-3, which is upregulated in AD lesions causing a reduced expression of cathelicidin. AD patients are susceptible to the skin colonization and infection by S. aureus, which through the production of exotoxins with supra-antigenic properties aggravates the disease. Gilaberte et al. [145] have observed a significant association between low serum vitamin D levels and certain virulence genes of S. aureus in isolates of AD children suggesting some role of vitamin D deficiency in S. aureus colonization. Thus, vitamin D supplementation could be promising in reducing the cutaneous S. aureus burden in AD patients. In fact, a recent clinical trial showed a reduction in the skin colonization by S. aureus and an improvement in clinical symptoms of AD patients, who received an oral supplementation of 2,000 IUs of vitamin D daily for 4 weeks [146].

AD patients are also prone to skin infections caused by herpes virus; this complication, known as eczema herpeticum, is particularly common in AD children and could be life threatening. It has been demonstrated that LL-37 is by far less expressed in skin of AD complicated by eczema herpeticum compared to skin of patients with uncomplicated AD [147]. Treatment by vitamin D has a beneficial effect in children with eczema herpeticum, and this effect seems to be mediated by an increase in the LL-37 level in the skin [143].

Studies addressing the relationship between vitamin D and AD could be hampered by the geographic, seasonal and diet-related vitamin D variations in AD patients and healthy controls, and despite the myriad of studies advocating the important role of vitamin D in AD, the relationship could not be stated with certainty. Animal studies evaluating the effect of vitamin D in AD are not consistent either and yielded conflictual findings: some studies showed an induction of thymic stromal lymphopoietin with topical application of calcitriol or its low calcemic analog MC903, which resulted in AD- like syndrome in mice [148, 149]. However, in an allergen-induced animal model of AD, systemic administration of a low calcemic vitamin D agonist significantly improved the symptoms of atopic eczema by restoring the epidermal barrier and modulating the immune system [150]. It was also observed that administration of a VDR agonist to allergen-induced AD mice selectively increased the frequency of Foxp3+ T_{reg} cells, reduced the expression of IL-4 in a lesional skin model, and induced a significant improvement in barrier function through robust induction of several skin barrier genes (like loricrin, involucrin, filaggrin, and transglutaminase) and AMP β-defensin.

Conclusion

Beyond the classical phosphocalcic effect of vitamin D, its role in the proper functioning of several tissues/organs including the skin has been receiving a growing interest. Vitamin D exhibits a pleiotropic effect in the skin with its role as antiproliferative, prodifferentiative, antiapoptotic and immunomodulator. It is also intricately involved in many skin pathologies, and it positively influences the outcome of certain inflammatory dermopathologies. So far, therapeutic interventions (topical and systemic) based on vitamin D have been proved beneficial in psoriasis and AD. Future studies are needed to mechanistically and intensely explore the specific pathways affected by vitamin D using the latest advanced technologies and to assess the safety and efficacy of vitamin D-based treatment regimens in various inflammatory skin diseases.

Acknowledgments

This work was supported by a grant from Sidra Medicine, Qatar.

Disclosure Statement

The authors declare that there is no conflict of interest.

References

- 1 Bikle DD: Vitamin D and the skin: physiology and pathophysiology. Rev Endocr Metab Disord 2012;13:3–19.
- 2 Reichrath J, Zouboulis CC, Vogt T, Holick MF: Targeting the vitamin D endocrine system (VDES) for the management of inflammatory and malignant skin diseases: a historical view and outlook. Rev Endocr Metab Disord 2016;17:405–417.
- 3 Sandilands A, Sutherland C, Irvine AD, McLean WH: Filaggrin in the frontline: role in skin barrier function and disease. J Cell Sci 2009;122:1285–1294.
- 4 Deraison C, Bonnart C, Lopez F, Besson C, Robinson R, Jayakumar A, Wagberg F, Brattsand M, Hachem JP, Leonardsson G, Hovnanian A: LEKTI fragments specifically inhibit KLK5, KLK7, and KLK14 and control desquamation through a pH-dependent interaction. Mol Biol Cell 2007;18:3607–3619.
- 5 Bikle DD: Vitamin D metabolism, mechanism of action, and clinical applications. Chem Biol 2014;21:319–329.
- 6 Henry HL: Regulation of vitamin D metabolism. Best Pract Res Clin Endocrinol Metab 2011;25:531–541.
- 7 Christakos S, Ajibade DV, Dhawan P, Fechner AJ, Mady LJ: Vitamin D: metabolism. Endocrinol Metab Clin North Am 2010;39:243–253, table of contents.
- 8 Holick MF, Binkley NC, Bischoff-Ferrari HA, Gordon CM, Hanley DA, Heaney RP, Murad MH, Weaver CM: Evaluation, treatment, and prevention of vitamin D deficiency: an Endocrine Society clinical practice guideline. J Clin Endocrinol Metab 2011;96:1911–1930.
- 9 Zittermann A, Gummert JF: Nonclassical vitamin D action. Nutrients 2010;2:408–425.

- 10 Bikle D: Nonclassic actions of vitamin D. J Clin Endocrinol Metab 2009;94:26–34.
- 11 Itin PH, Pittelkow MR, Kumar R: Effects of vitamin D metabolites on proliferation and differentiation of cultured human epidermal keratinocytes grown in serum-free or defined culture medium. Endocrinology 1994;135: 1793–1798.
- 12 Gniadecki R: Stimulation versus inhibition of keratinocyte growth by 1,25-dihydroxyvitamin D₃: dependence on cell culture conditions. J Invest Dermatol 1996;106:510–516.
- 13 Svendsen ML, Daneels G, Geysen J, Binderup L, Kragballe K: Proliferation and differentiation of cultured human keratinocytes is modulated by $1,25(OH)_2D_3$ and synthetic vitamin D₃ analogues in a cell density-, calcium- and serum-dependent manner. Pharmacol Toxicol 1997;80:49–56.
- 14 Bikle DD: Vitamin D metabolism and function in the skin. Mol Cell Endocrinol 2011; 347:80–89.
- 15 Bikle DD, Ng D, Tu CL, Oda Y, Xie Z: Calcium- and vitamin D-regulated keratinocyte differentiation. Mol Cell Endocrinol 2001; 177:161–171.
- 16 Ratnam AV, Bikle DD, Cho JK: 1,25-Dihydroxyvitamin D_3 enhances the calcium response of keratinocytes. J Cell Physiol 1999; 178:188–196.
- 17 Pillai S, Mahajan M, Carlomusto M: Ceramide potentiates, but sphingomyelin inhibits, vitamin D-induced keratinocyte differentiation: comparison between keratinocytes and HL-60 cells. Arch Dermatol Res 1999; 291:284–289.

- 18 Xie Z, Komuves L, Yu QC, Elalieh H, Ng DC, Leary C, Chang S, Crumrine D, Yoshizawa T, Kato S, Bikle DD: Lack of the vitamin D receptor is associated with reduced epidermal differentiation and hair follicle growth. J Invest Dermatol 2002;118:11–16.
- 19 Oda Y, Sihlbom C, Chalkley RJ, Huang L, Rachez C, Chang CP, Burlingame AL, Freedman LP, Bikle DD: Two distinct coactivators, DRIP/mediator and SRC/p160, are differentially involved in VDR transactivation during keratinocyte differentiation. J Steroid Biochem Mol Biol 2004;89–90:273–276.
- 20 Bikle DD, Oda Y, Xie Z: Calcium and 1,25(OH)₂D: interacting drivers of epidermal differentiation. J Steroid Biochem Mol Biol 2004;89–90:355–360.
- 21 Effendy I, Kwangsukstith C, Chiappe M, Maibach HI: Effects of calcipotriol on stratum corneum barrier function, hydration and cell renewal in humans. Br J Dermatol 1996;135: 545–549.
- 22 Hong SP, Oh Y, Jung M, Lee S, Jeon H, Cho MY, Lee SH, Choi EH: Topical calcitriol restores the impairment of epidermal permeability and antimicrobial barriers induced by corticosteroids. Br J Dermatol 2010;162:1251–1260.
- 23 Oda Y, Uchida Y, Moradian S, Crumrine D, Elias PM, Bikle DD: Vitamin D receptor and coactivators SRC2 and 3 regulate epidermisspecific sphingolipid production and permeability barrier formation. J Invest Dermatol 2009;129:1367–1378.
- 24 Manggau M, Kim DS, Ruwisch L, Vogler R, Korting HC, Schafer-Korting M, Kleuser B: 1α,25-Dihydroxyvitamin D₃ protects human keratinocytes from apoptosis by the formation of sphingosine-1-phosphate. J Invest Dermatol 2001;117:1241–1249.

- 25 De Haes P, Garmyn M, Carmeliet G, Degreef H, Vantieghem K, Bouillon R, Segaert S: Molecular pathways involved in the anti-apoptotic effect of 1,25-dihydroxyvitamin D_3 in primary human keratinocytes. J Cell Biochem 2004;93:951–967.
- 26 Schauber J, Gallo RL: Antimicrobial peptides and the skin immune defense system. J Allergy Clin Immunol 2009;124:R13–R18.
- 27 Gallo RL, Hooper LV: Epithelial antimicrobial defence of the skin and intestine. Nat Rev Immunol 2012;12:503–516.
- 28 Schauber J, Gallo RL: Expanding the roles of antimicrobial peptides in skin: alarming and arming keratinocytes. J Invest Dermatol 2007; 127:510–512.
- 29 Schauber J, Dorschner RA, Coda AB, Büchau AS, Liu PT, Kiken D, Helfrich YR, Kang S, Elalieh HZ, Steinmeyer A, Zugel U, Bikle DD, Modlin RL, Gallo RL: Injury enhances TLR2 function and antimicrobial peptide expression through a vitamin D-dependent mechanism. J Clin Invest 2007;117:803–811.
- 30 Lee WJ, Cha HW, Sohn MY, Lee SJ, Kim DW: Vitamin D increases expression of cathelicidin in cultured sebocytes. Arch Dermatol Res 2012;304:627–632.
- 31 Heilborn JD, Weber G, Gronberg A, Dieterich C, Stahle M: Topical treatment with the vitamin D analogue calcipotriol enhances the upregulation of the antimicrobial protein hCAP18/LL-37 during wounding in human skin in vivo. Exp Dermatol 2010;19:332–338.
- 32 Weber G, Heilborn JD, Chamorro Jimenez CI, Hammarsjo A, Torma H, Stahle M: Vitamin D induces the antimicrobial protein hCAP18 in human skin. J Invest Dermatol 2005;124:1080-1082.
- 33 Dai X, Sayama K, Tohyama M, Shirakata Y, Hanakawa Y, Tokumaru S, Yang L, Hirakawa S, Hashimoto K: PPARgamma mediates innate immunity by regulating the 1alpha,25dihydroxyvitamin D_3 induced hBD-3 and cathelicidin in human keratinocytes. J Dermatol Sci 2010;60:179–186.
- 34 Liu PT, Schenk M, Walker VP, Dempsey PW, Kanchanapoomi M, Wheelwright M, Vazirnia A, Zhang X, Steinmeyer A, Zugel U, Hollis BW, Cheng G, Modlin RL: Convergence of IL-1beta and VDR activation pathways in human TLR2/1-induced antimicrobial responses. PLoS One 2009;4:e5810.
- 35 Yamasaki K, Schauber J, Coda A, Lin H, Dorschner RA, Schechter NM, Bonnart C, Descargues P, Hovnanian A, Gallo RL: Kallikrein-mediated proteolysis regulates the antimicrobial effects of cathelicidins in skin. FASEB J 2006;20:2068–2080.
- 36 Morizane S, Yamasaki K, Kabigting FD, Gallo RL: Kallikrein expression and cathelicidin processing are independently controlled in keratinocytes by calcium, vitamin D(3), and retinoic acid. J Invest Dermatol 2010;130: 1297–1306.

- 37 Bagot M, Charue D, Lescs MC, Pamphile RP, Revuz J: Immunosuppressive effects of 1,25-dihydroxyvitamin D₃ and its analogue calcipotriol on epidermal cells. Br J Dermatol 1994;130:424–431.
- 38 Dam TN, Moller B, Hindkjaer J, Kragballe K: The vitamin D₃ analog calcipotriol suppresses the number and antigen-presenting function of Langerhans cells in normal human skin. J Investig Dermatol Symp Proc 1996;1:72–77.
- 39 Ghoreishi M, Bach P, Obst J, Komba M, Fleet JC, Dutz JP: Expansion of antigen-specific regulatory T cells with the topical vitamin D analog calcipotriol. J Immunol 2009;182: 6071–6078.
- 40 Gorman S, Geldenhuys S, Judge M, Weeden CE, Waithman J, Hart PH: Dietary vitamin D increases percentages and function of regulatory T cells in the skin-draining lymph nodes and suppresses dermal inflammation. J Immunol Res 2016;2016:1426503.
- 41 Yamanaka K, Dimitroff CJ, Fuhlbrigge RC, Kakeda M, Kurokawa I, Mizutani H, Kupper TS: Vitamins A and D are potent inhibitors of cutaneous lymphocyte-associated antigen expression. J Allergy Clin Immunol 2008;121: 148–157. e143.
- 42 Yamanaka KI, Kakeda M, Kitagawa H, Tsuda K, Akeda T, Kurokawa I, Gabazza EC, Kupper TS, Mizutani H: 1,24-Dihydroxyvitamin D(3) (tacalcitol) prevents skin T-cell infiltration. Br J Dermatol 2010;162:1206–1215.
- 43 Sigmundsdottir H, Pan J, Debes GF, Alt C, Habtezion A, Soler D, Butcher EC: DCs metabolize sunlight-induced vitamin D_3 to "program" T cell attraction to the epidermal chemokine CCL27. Nat Immunol 2007;8:285– 293.
- 44 Baeke F, Korf H, Overbergh L, Verstuyf A, Thorrez L, Van Lommel L, Waer M, Schuit F, Gysemans C, Mathieu C: The vitamin D analog, TX527, promotes a human CD4+CD25highCD127low regulatory T cell profile and induces a migratory signature specific for homing to sites of inflammation. J Immunol 2011;186:132–142.
- 45 Khoo AL, Koenen HJ, Michels M, Ooms S, Bosch M, Netea MG, Joosten I, van der Ven AJ: High-dose vitamin D₃ supplementation is a requisite for modulation of skin-homing markers on regulatory T cells in HIV-infected patients. AIDS Res Hum Retroviruses 2013; 29:299–306.
- 46 Khoo AL, Koenen HJ, Chai LY, Sweep FC, Netea MG, van der Ven AJ, Joosten I: Seasonal variation in vitamin D(3) levels is paralleled by changes in the peripheral blood human T cell compartment. PLoS One 2012;7:e29250.
- 47 Lande R, Gregorio J, Facchinetti V, Chatterjee B, Wang YH, Homey B, Cao W, Su B, Nestle FO, Zal T, Mellman I, Schroder JM, Liu YJ, Gilliet M: Plasmacytoid dendritic cells sense self-DNA coupled with antimicrobial peptide. Nature 2007;449:564–569.

- 48 Nestle FO, Conrad C, Tun-Kyi A, Homey B, Gombert M, Boyman O, Burg G, Liu YJ, Gilliet M: Plasmacytoid predendritic cells initiate psoriasis through interferon-alpha production. J Exp Med 2005;202:135–143.
- 49 Lowes MA, Chamian F, Abello MV, Fuentes-Duculan J, Lin SL, Nussbaum R, Novitskaya I, Carbonaro H, Cardinale I, Kikuchi T, Gilleaudeau P, Sullivan-Whalen M, Wittkowski KM, Papp K, Garovoy M, Dummer W, Steinman RM, Krueger JG: Increase in TNF-alpha and inducible nitric oxide synthase-expressing dendritic cells in psoriasis and reduction with efalizumab (anti-CD11a). Proc Natl Acad Sci USA 2005;102:19057–19062.
- 50 Chu CC, Di Meglio P, Nestle FO: Harnessing dendritic cells in inflammatory skin diseases. Semin Immunol 2011;23:28–41.
- 51 Schön MP, Broekaert SM, Erpenbeck L: Sexy again: the renaissance of neutrophils in psoriasis. Exp Dermatol 2017;26:305–311.
- 52 Tobin AM, Lynch L, Kirby B, O'Farrelly C: Natural killer cells in psoriasis. J Innate Immun 2011;3:403–410.
- 53 Cai Y, Fleming C, Yan J: New insights of T cells in the pathogenesis of psoriasis. Cell Mol Immunol 2012;9:302–309.
- 54 Peternel S, Kastelan M: Immunopathogenesis of psoriasis: focus on natural killer T cells. J Eur Acad Dermatol Venereol 2009;23:1123– 1127.
- 55 Nestle FO, Kaplan DH, Barker J: Psoriasis. N Engl J Med 2009;361:496–509.
- 56 Soler DC, Sugiyama H, Young AB, Massari JV, McCormick TS, Cooper KD: Psoriasis patients exhibit impairment of the high potency CCR5(+) T regulatory cell subset. Clin Immunol 2013;149:111–118.
- 57 Mattozzi C, Salvi M, D'Epiro S, Giancristoforo S, Macaluso L, Luci C, Lal K, Calvieri S, Richetta AG: Importance of regulatory T cells in the pathogenesis of psoriasis: review of the literature. Dermatology 2013;227:134–145.
- 58 Yang L, Li B, Dang E, Jin L, Fan X, Wang G: Impaired function of regulatory T cells in patients with psoriasis is mediated by phosphorylation of STAT3. J Dermatol Sci 2016;81: 85–92.
- 59 Mattozzi C, Paolino G, Salvi M, Macaluso L, Luci C, Morrone S, Calvieri S, Richetta AG: Peripheral blood regulatory T cell measurements correlate with serum vitamin D level in patients with psoriasis. Eur Rev Med Pharmacol Sci 2016;20:1675–1679.
- 60 Maleki M, Nahidi Y, Azizahari S, Meibodi NT, Hadianfar A: Serum 25-OH vitamin D level in psoriatic patients and comparison with control subjects. J Cutan Med Surg 2016; 20:207–210.
- 61 Gisondi P, Rossini M, Di Cesare A, Idolazzi L, Farina S, Beltrami G, Peris K, Girolomoni G: Vitamin D status in patients with chronic plaque psoriasis. Br J Dermatol 2012;166: 505–510.

- 62 Chandrashekar L, Kumarit GR, Rajappa M, Revathy G, Munisamy M, Thappa DM: 25-Hydroxy vitamin D and ischaemia-modified albumin levels in psoriasis and their association with disease severity. Br J Biomed Sci 2015;72:56–60.
- 63 Orgaz-Molina J, Magro-Checa C, Rosales-Alexander JL, Arrabal-Polo MA, Castellote-Caballero L, Buendia-Eisman A, Raya-Alvarez E, Arias-Santiago S: Vitamin D insufficiency is associated with higher carotid intima-media thickness in psoriatic patients. Eur J Dermatol 2014;24:53–62.
- 64 Al-Mutairi N, Shaaban D: Effect of narrowband ultraviolet B therapy on serum vitamin D and cathelicidin (LL-37) in patients with chronic plaque psoriasis. J Cutan Med Surg 2014;18:43–48.
- 65 Atwa MA, Balata MG, Hussein AM, Abdelrahman NI, Elminshawy HH: Serum 25-hydroxyvitamin D concentration in patients with psoriasis and rheumatoid arthritis and its association with disease activity and serum tumor necrosis factor-alpha. Saudi Med J 2013;34:806–813.
- 66 Ricceri F, Pescitelli L, Tripo L, Prignano F: Deficiency of serum concentration of 25-hydroxyvitamin D correlates with severity of disease in chronic plaque psoriasis. J Am Acad Dermatol 2013;68:511–512.
- 67 Wilson PB: Serum 25-hydroxyvitamin D status in individuals with psoriasis in the general population. Endocrine 2013;44:537–539.
- 68 Finamor DC, Sinigaglia-Coimbra R, Neves LC, Gutierrez M, Silva JJ, Torres LD, Surano F, Neto DJ, Novo NF, Juliano Y, Lopes AC, Coimbra CG: A pilot study assessing the effect of prolonged administration of high daily doses of vitamin D on the clinical course of vitiligo and psoriasis. Dermatoendocrinology 2013;5:222–234.
- 69 Kragballe K, Beck HI, Sogaard H: Improvement of psoriasis by a topical vitamin D₃ analogue (MC 903) in a double-blind study. Br J Dermatol 1988;119:223–230.
- 70 Van de Kerkhof PC, Berth-Jones J, Griffiths CE, Harrison PV, Honigsmann H, Marks R, Roelandts R, Schopf E, Trompke C: Longterm efficacy and safety of tacalcitol ointment in patients with chronic plaque psoriasis. Br J Dermatol 2002;146:414–422.
- 71 Barker JN, Ashton RE, Marks R, Harris RI, Berth-Jones J: Topical maxacalcitol for the treatment of psoriasis vulgaris: a placebocontrolled, double-blind, dose-finding study with active comparator. Br J Dermatol 1999; 141:274–278.
- 72 Le P, Tu J, Gebauer K, Brown S: Serum 25-hydroxyvitamin D increases with NB-UVB and UVA/UVB phototherapy in patients with psoriasis and atopic dermatitis in Western Australia. Australas J Dermatol 2016;57:115–121.
- 73 Franken SM, Witte B, Pavel S, Rustemeyer T: Psoriasis and daily low-emission phototherapy: effects on disease and vitamin D level. Photodermatol Photoimmunol Photomed 2015;31:83–89.

- 74 Feldmeyer L, Shojaati G, Spanaus KS, Navarini A, Theler B, Donghi D, Urosevic-Maiwald M, Glatz M, Imhof L, Barysch MJ, Dummer R, Roos M, French LE, Surber C, Hofbauer GF: Phototherapy with UVB narrowband, UVA/UVBnb, and UVA1 differentially impacts serum 25-hydroxyvitamin-D₃. J Am Acad Dermatol 2013;69:530–536.
- 75 Takahashi H, Tsuji H, Ishida-Yamamoto A, Iizuka H: Comparison of clinical effects of psoriasis treatment regimens among calcipotriol alone, narrowband ultraviolet B phototherapy alone, combination of calcipotriol and narrowband ultraviolet B phototherapy once a week, and combination of calcipotriol and narrowband ultraviolet B phototherapy more than twice a week. J Dermatol 2013;40: 424–427.
- 76 Segaert S, Ropke M: The biological rationale for use of vitamin D analogs in combination with corticosteroids for the topical treatment of plaque psoriasis. J Drugs Dermatol 2013; 12:e129–e137.
- 77 Bagel J, Levi E, Tyring S, Knuckles ML: Reallife treatment profile of calcipotriene and betamethasone dipropionate topical suspension in patients with psoriasis vulgaris. J Drugs Dermatol 2014;13:1374–1379.
- 78 Eichenfield LF, Ganslandt C, Kurvits M, Schlessinger J: Safety and efficacy of calcipotriene plus betamethasone dipropionate topical suspension in the treatment of extensive scalp psoriasis in adolescents ages 12–17 years. Pediatr Dermatol 2015;32:28–35.
- 79 Karthaus N, van Spriel AB, Looman MW, Chen S, Spilgies LM, Lieben L, Carmeliet G, Ansems M, Adema GJ: Vitamin D controls murine and human plasmacytoid dendritic cell function. J Invest Dermatol 2014;134: 1255–1264.
- 80 Dyring-Andersen B, Bonefeld CM, Bzorek M, Lovendorf MB, Lauritsen JP, Skov L, Geisler C: The vitamin D analogue calcipotriol reduces the frequency of CD8+ IL-17+ T cells in psoriasis lesions. Scand J Immunol 2015;82: 84–91.
- 81 Fujiyama T, Ito T, Umayahara T, Ikeya S, Tatsuno K, Funakoshi A, Hashizume H, Tokura Y: Topical application of a vitamin D_3 analogue and corticosteroid to psoriasis plaques decreases skin infiltration of TH17 cells and their ex vivo expansion. J Allergy Clin Immunol 2016;138:517–528 e515.
- 82 Sato-Deguchi E, Imafuku S, Chou B, Ishii K, Hiromatsu K, Nakayama J: Topical vitamin D(3) analogues induce thymic stromal lymphopoietin and cathelicidin in psoriatic skin lesions. Br J Dermatol 2012;167:77–84.
- 83 Balato A, Schiattarella M, Lembo S, Mattii M, Prevete N, Balato N, Ayala F: Interleukin-1 family members are enhanced in psoriasis and suppressed by vitamin D and retinoic acid. Arch Dermatol Res 2013;305:255-262.

- 84 Hegyi Z, Zwicker S, Bureik D, Peric M, Koglin S, Batycka-Baran A, Prinz JC, Ruzicka T, Schauber J, Wolf R: Vitamin D analog calcipotriol suppresses the Th17 cytokine-induced proinflammatory S100 "alarmins" psoriasin (S100A7) and koebnerisin (S100A15) in psoriasis. J Invest Dermatol 2012;132:1416–1424.
- 85 Datta Mitra A, Raychaudhuri SP, Abria CJ, Mitra A, Wright R, Ray R, Kundu-Raychaudhuri S: 1alpha,25-Dihydroxyvitamin-D₃-3bromoacetate regulates AKT/mTOR signaling cascades: a therapeutic agent for psoriasis. J Invest Dermatol 2013;133:1556–1564.
- 86 Hoss E, Austin HR, Batie SF, Jurutka PW, Haussler MR, Whitfield GK: Control of late cornified envelope genes relevant to psoriasis risk: upregulation by 1,25-dihydroxyvitamin D₃ and plant-derived delphinidin. Arch Dermatol Res 2013;305:867–878.
- 87 Visconti B, Paolino G, Carotti S, Pendolino AL, Morini S, Richetta AG, Calvieri S: Immunohistochemical expression of VDR is associated with reduced integrity of tight junction complex in psoriatic skin. J Eur Acad Dermatol Venereol 2015;29:2038–2042.
- 88 Savoia P, Novelli M, De Matteis A, Verrone A, Bernengo MG: Effects of topical calcipotriol on the expression of adhesion molecules in psoriasis. J Cutan Pathol 1998;25:89–94.
- 89 Richetta AG, Silvestri V, Giancristoforo S, Rizzolo P, D'Epiro S, Graziano V, Mattozzi C, Navazio AS, Campoli M, D'Amico C, Scarno M, Calvieri S, Ottini L: A-1012G promoter polymorphism of vitamin D receptor gene is associated with psoriasis risk and lower allele-specific expression. DNA Cell Biol 2014;33:102–109.
- 90 Zhou X, Xu LD, Li YZ: The association of polymorphisms of the vitamin D receptor gene with psoriasis in the Han population of northeastern China. J Dermatol Sci 2014;73: 63–66.
- 91 Polic MV, Rucevic I, Barisic-Drusko V, Miskulin M, Glavas-Obrovac L, Stefanic M, Karner I, Lipozencic J, Bacun T, Mihaljevic I: Polymorphisms of vitamin D receptor gene in the population of eastern Croatia with psoriasis vulgaris and diabetes mellitus. Coll Antropol 2012;36:451–457.
- 92 Rucevic I, Stefanic M, Tokic S, Vuksic M, Glavas-Obrovac L, Barisic-Drusko V: Lack of association of vitamin D receptor gene 3'-haplotypes with psoriasis in Croatian patients. J Dermatol 2012;39:58–62.
- 93 Zuel-Fakkar NM, Kamel MM, Asaad MK, Mahran MZ, Shehab AA: A study of ApaI and TaqI genotypes of the vitamin D receptor in Egyptian patients with psoriasis. Clin Exp Dermatol 2011;36:355–359.
- 94 Liu JL, Zhang SQ, Zeng HM: ApaI, BsmI, FokI and TaqI polymorphisms in the vitamin D receptor (VDR) gene and the risk of psoriasis: a meta-analysis. J Eur Acad Dermatol Venereol 2013;27:739–746.
- 95 Lee YH, Choi SJ, Ji JD, Song GG: Vitamin D receptor ApaI, TaqI, BsmI, and FokI polymorphisms and psoriasis susceptibility: a meta-analysis. Mol Biol Rep 2012;39:6471–6478.

- 96 Ryan C, Renfro L, Collins P, Kirby B, Rogers S: Clinical and genetic predictors of response to narrowband ultraviolet B for the treatment of chronic plaque psoriasis. Br J Dermatol 2010;163:1056–1063.
- 97 Halsall JA, Osborne JE, Pringle JH, Hutchinson PE: Vitamin D receptor gene polymorphisms, particularly the novel A-1012G promoter polymorphism, are associated with vitamin D_3 responsiveness and non-familial susceptibility in psoriasis. Pharmacogenet Genomics 2005;15:349–355.
- 98 Saeki H, Asano N, Tsunemi Y, Takekoshi T, Kishimoto M, Mitsui H, Tada Y, Torii H, Komine M, Asahina A, Tamaki K: Polymorphisms of vitamin D receptor gene in Japanese patients with psoriasis vulgaris. J Dermatol Sci 2002;30:167–171.
- 99 Bieber T: Atopic dermatitis. N Engl J Med 2008;358:1483-1494.
- 100 Bieber T: Atopic dermatitis. Ann Dermatol 2010;22:125–137.
- 101 Le Lamer M, Pellerin L, Reynier M, Cau L, Pendaries V, Leprince C, Mechin MC, Serre G, Paul C, Simon M: Defects of corneocyte structural proteins and epidermal barrier in atopic dermatitis. Biol Chem 2015;396: 1163–1179.
- 102 Palmer CN, Irvine AD, Terron-Kwiatkowski A, Zhao Y, Liao H, Lee SP, Goudie DR, Sandilands A, Campbell LE, Smith FJ, O'Regan GM, Watson RM, Cecil JE, Bale SJ, Compton JG, DiGiovanna JJ, Fleckman P, Lewis-Jones S, Arseculeratne G, Sergeant A, Munro CS, El Houate B, McElreavey K, Halkjaer LB, Bisgaard H, Mukhopadhyay S, McLean WH: Common loss-of-function variants of the epidermal barrier protein filaggrin are a major predisposing factor for atopic dermatitis. Nat Genet 2006;38:441– 446.
- 103 Tokura Y: Extrinsic and intrinsic types of atopic dermatitis. J Dermatol Sci 2010;58: 1–7.
- 104 Soumelis V, Reche PA, Kanzler H, Yuan W, Edward G, Homey B, Gilliet M, Ho S, Antonenko S, Lauerma A, Smith K, Gorman D, Zurawski S, Abrams J, Menon S, McClanahan T, de Waal-Malefyt Rd R, Bazan F, Kastelein RA, Liu YJ: Human epithelial cells trigger dendritic cell mediated allergic inflammation by producing TSLP. Nat Immunol 2002;3:673–680.
- 105 Said A, Weindl G: Regulation of dendritic cell function in inflammation. J Immunol Res 2015;2015:743169.
- 106 Kerschenlohr K, Decard S, Przybilla B, Wollenberg A: Atopy patch test reactions show a rapid influx of inflammatory dendritic epidermal cells in patients with extrinsic atopic dermatitis and patients with intrinsic atopic dermatitis. J Allergy Clin Immunol 2003; 111:869–874.

- 107 Gaspar K, Barath S, Nagy G, Mocsai G, Gyimesi E, Szodoray P, Irinyi B, Zeher M, Remenyik E, Szegedi A: Regulatory T-cell subsets with acquired functional impairment: important indicators of disease severity in atopic dermatitis. Acta Derm Venereol 2015;95:151–155.
- 108 Ou LS, Goleva E, Hall C, Leung DY: T regulatory cells in atopic dermatitis and subversion of their activity by superantigens. J Allergy Clin Immunol 2004;113:756–763.
- 109 Lesiak A, Smolewski P, Sobolewska-Sztychny D, Sysa-Jedrzejowska A, Narbutt J: The role of T-regulatory cells and Toll-like receptors 2 and 4 in atopic dermatitis. Scand J Immunol 2012;76:405–410.
- 110 Roesner LM, Floess S, Witte T, Olek S, Huehn J, Werfel T: Foxp3(+) regulatory T cells are expanded in severe atopic dermatitis patients. Allergy 2015;70:1656–1660.
- 111 Ma L, Xue HB, Guan XH, Shu CM, Wang F, Zhang JH, An RZ: The Imbalance of Th17 cells and CD4(+) CD25(high) Foxp3(+) T_{reg} cells in patients with atopic dermatitis. J Eur Acad Dermatol Venereol 2014;28:1079–1086.
- 112 Verhagen J, Akdis M, Traidl-Hoffmann C, Schmid-Grendelmeier P, Hijnen D, Knol EF, Behrendt H, Blaser K, Akdis CA: Absence of T-regulatory cell expression and function in atopic dermatitis skin. J Allergy Clin Immunol 2006;117:176–183.
- 113 De Benedetto A, Agnihothri R, McGirt LY, Bankova LG, Beck LA: Atopic dermatitis: a disease caused by innate immune defects? J Invest Dermatol 2009;129:14–30.
- 114 Hata TR, Gallo RL: Antimicrobial peptides, skin infections, and atopic dermatitis. Semin Cutan Med Surg 2008;27:144–150.
- 115 Weiland SK, Husing A, Strachan DP, Rzehak P, Pearce N; ISAAC Phase One Study Group: Climate and the prevalence of symptoms of asthma, allergic rhinitis, and atopic eczema in children. Occup Environ Med 2004;61:609–615.
- 116 Schlichte MJ, Vandersall A, Katta R: Diet and eczema: a review of dietary supplements for the treatment of atopic dermatitis. Dermatol Pract Concept 2016;6:23–29.
- 117 Cheng HM, Kim S, Park GH, Chang SE, Bang S, Won CH, Lee MW, Choi JH, Moon KC: Low vitamin D levels are associated with atopic dermatitis, but not allergic rhinitis, asthma, or IgE sensitization, in the adult Korean population. J Allergy Clin Immunol 2014;133:1048–1055.
- 118 Kang JW, Kim JH, Kim HJ, Lee JG, Yoon JH, Kim CH: Association of serum 25-hydroxyvitamin D with serum IgE levels in Korean adults. Auris Nasus Larynx 2016;43: 84–88.
- 119 Yang AR, Kim YN, Lee BH: Dietary intakes and lifestyle patterns of Korean children and adolescents with atopic dermatitis: using the fourth and fifth Korean National Health and Nutrition Examination Survey (KNHANES IV,V), 2007–11. Ecol Food Nutr 2016;55: 50–64.

- 120 El Taieb MA, Fayed HM, Aly SS, Ibrahim AK: Assessment of serum 25-hydroxyvitamin D levels in children with atopic dermatitis: correlation with SCORAD index. Dermatitis 2013;24:296–301.
- 121 Kim MJ, Kim SN, Lee YW, Choe YB, Ahn KJ: Vitamin D status and efficacy of vitamin D supplementation in atopic dermatitis: a systematic review and meta-analysis. Nutrients 2016;8:E789.
- 122 Cheon BR, Shin JE, Kim YJ, Shim JW, Kim DS, Jung HL, Park MS, Shim JY: Relationship between serum 25-hydroxyvitamin D and interleukin-31 levels, and the severity of atopic dermatitis in children. Korean J Pediatr 2015;58:96–101.
- 123 Peroni DG, Piacentini GL, Cametti E, Chinellato I, Boner AL: Correlation between serum 25-hydroxyvitamin D levels and severity of atopic dermatitis in children. Br J Dermatol 2011;164:1078–1082.
- 124 Thuesen BH, Heede NG, Tang L, Skaaby T, Thyssen JP, Friedrich N, Linneberg A: No association between vitamin D and atopy, asthma, lung function or atopic dermatitis: a prospective study in adults. Allergy 2015;70: 1501–1504.
- 125 Back O, Blomquist HK, Hernell O, Stenberg B: Does vitamin D intake during infancy promote the development of atopic allergy? Acta Derm Venereol 2009;89:28–32.
- 126 Miyake Y, Tanaka K, Okubo H, Sasaki S, Arakawa M: Maternal consumption of dairy products, calcium, and vitamin D during pregnancy and infantile allergic disorders. Ann Allergy Asthma Immunol 2014;113: 82–87.
- 127 Gale CR, Robinson SM, Harvey NC, Javaid MK, Jiang B, Martyn CN, Godfrey KM, Cooper C: Maternal vitamin D status during pregnancy and child outcomes. Eur J Clin Nutr 2008;62:68–77.
- 128 Chiu CY, Huang SY, Peng YC, Tsai MH, Hua MC, Yao TC, Yeh KW, Huang JL: Maternal vitamin D levels are inversely related to allergic sensitization and atopic diseases in early childhood. Pediatr Allergy Immunol 2015;26:337–343.
- 129 Jones AP, Palmer D, Zhang G, Prescott SL: Cord blood 25-hydroxyvitamin D₃ and allergic disease during infancy. Pediatrics 2012;130:e1128-e1135.
- 130 Baiz N, Dargent-Molina P, Wark JD, Souberbielle JC, Annesi-Maesano I: Cord serum 25-hydroxyvitamin D and risk of early childhood transient wheezing and atopic dermatitis. J Allergy Clin Immunol 2014;133:147– 153.
- 131 Stelmach I, Majak P, Jerzynska J, Podlecka D, Stelmach W, Polanska K, Gromadzinska J, Wasowicz W, Hanke W: Cord serum 25-hydroxyvitamin D correlates with early childhood viral-induced wheezing. Respir Med 2015;109:38–43.

- 132 Heine G, Hoefer N, Franke A, Nothling U, Schumann RR, Hamann L, Worm M: Association of vitamin D receptor gene polymorphisms with severe atopic dermatitis in adults. Br J Dermatol 2013;168:855–858.
- 133 Kilic S, Silan F, Hiz MM, Isik S, Ogretmen Z, Ozdemir O: Vitamin D receptor gene BSMI, FOKI, APAI, and TAQI polymorphisms and the risk of atopic dermatitis. J Investig Allergol Clin Immunol 2016;26:106–110.
- 134 Hallau J, Hamann L, Schumann RR, Worm M, Heine G: A promoter polymorphism of the vitamin D metabolism gene Cyp24a1 is associated with severe atopic dermatitis in adults. Acta Derm Venereol 2016;96:169– 172.
- 135 Kim G, Bae JH: Vitamin D and atopic dermatitis: a systematic review and meta-analysis. Nutrition 2016;32:913–920.
- 136 Camargo CA Jr, Ganmaa D, Sidbury R, Erdenedelger K, Radnaakhand N, Khandsuren B: Randomized trial of vitamin D supplementation for winter-related atopic dermatitis in children. J Allergy Clin Immunol 2014;134:831–835 e831.
- 137 Amestejani M, Salehi BS, Vasigh M, Sobhkhiz A, Karami M, Alinia H, Kamrava SK, Shamspour N, Ghalehbaghi B, Behzadi AH: Vitamin D supplementation in the treatment of atopic dermatitis: a clinical trial study. J Drugs Dermatol 2012;11:327–330.
- 138 Di Filippo P, Scaparrotta A, Rapino D, Cingolani A, Attanasi M, Petrosino MI, Chuang K, Di Pillo S, Chiarelli F: Vitamin D supplementation modulates the immune system and improves atopic dermatitis in children. Int Arch Allergy Immunol 2015;166:91–96.

- 139 Drozdenko G, Heine G, Worm M: Oral vitamin D increases the frequencies of CD38+ human B cells and ameliorates IL-17-producing T cells. Exp Dermatol 2014;23:107– 112.
- 140 Yip KH, Kolesnikoff N, Yu C, Hauschild N, Taing H, Biggs L, Goltzman D, Gregory PA, Anderson PH, Samuel MS, Galli SJ, Lopez AF, Grimbaldeston MA: Mechanisms of vitamin D(3) metabolite repression of IgE-dependent mast cell activation. J Allergy Clin Immunol 2014; 133: 1356–1364, e1351– e1314.
- 141 Kanda N, Hau CS, Tada Y, Sato S, Watanabe
 S: Decreased serum LL-37 and vitamin D₃ levels in atopic dermatitis: relationship between IL-31 and oncostatin M. Allergy 2012; 67:804–812.
- 142 Mallbris L, Carlen L, Wei T, Heilborn J, Nilsson MF, Granath F, Stahle M: Injury downregulates the expression of the human cathelicidin protein hCAP18/LL-37 in atopic dermatitis. Exp Dermatol 2010;19:442–449.
- 143 Albenali LH, Danby S, Moustafa M, Brown K, Chittock J, Shackley F, Cork MJ: Vitamin D and antimicrobial peptide levels in patients with atopic dermatitis and atopic dermatitis complicated by eczema herpeticum: a pilot study. J Allergy Clin Immunol 2016; 138:1715–1719. e1714.

- 144 Büchau AS, MacLeod DT, Morizane S, Kotol PF, Hata T, Gallo RL: Bcl-3 acts as an innate immune modulator by controlling antimicrobial responses in keratinocytes. J Invest Dermatol 2009;129:2148–2155.
- 145 Gilaberte Y, Sanmartin R, Aspiroz C, Hernandez-Martin A, Benito D, Sanz-Puertolas P, Alonso M, Torrelo A, Torres C: Correlation between serum 25-hydroxyvitamin D and virulence genes of *Staphylococcus aureus* isolates colonizing children with atopic dermatitis. Pediatr Dermatol 2015;32:506– 513.
- 146 Udompataikul M, Huajai S, Chalermchai T, Taweechotipatr M, Kamanamool N: The effects of oral vitamin D supplement on atopic dermatitis: a clinical trial with *Staphylococcus aureus* colonization determination. J Med Assoc Thai 2015;98(suppl 9):S23–S30.
- 147 Bussmann C, Peng WM, Bieber T, Novak N: Molecular pathogenesis and clinical implications of eczema herpeticum. Expert Rev Mol Med 2008;10:e21.
- 148 Li M, Hener P, Zhang Z, Kato S, Metzger D, Chambon P: Topical vitamin D_3 and lowcalcemic analogs induce thymic stromal lymphopoietin in mouse keratinocytes and trigger an atopic dermatitis. Proc Natl Acad Sci USA 2006;103:11736–11741.
- 149 Leyva-Castillo JM, Hener P, Jiang H, Li M: TSLP produced by keratinocytes promotes allergen sensitization through skin and thereby triggers atopic march in mice. J Invest Dermatol 2013;133:154–163.
- 150 Hartmann B, Heine G, Babina M, Steinmeyer A, Zugel U, Radbruch A, Worm M: Targeting the vitamin D receptor inhibits the B cell-dependent allergic immune response. Allergy 2011;66:540–548.

Umar/Sastry/Al Ali/Al-Khulaifi/Wang/

Chouchane