Immune Alterations in IgE and Non IgE-Associated Atopic Dermatitis

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Abstract: Atopic dermatitis is a complex disease in which a strong interaction between alterations of skin barrier and the adaptive immune system coexists. In the recent years, new findings have underlined the importance of skin proteins, especially filagrin, which participate to the outmost layers of the skin. To strengthen this physical barrier, many factors are available, such as antimicrobial peptides, chemokines and cytokines produced by keratinocytes. Skin disruption can easily allow the allergen penetration and the local keratinocytes can promote the adaptive immune response toward a Th2 phenotype. On the other side, allergic Th2 cytokines may downregulate the production of skin barrier proteins, facilitating the penetration of allergens. Moreover, data on murine models show the absolute relevance of the systemic immune system to develop clinical skin reaction. Since the clinical aspect of patients with AD does not show different patterns whatever is the prevalent underlying mechanism, in clinical practice it is difficult to translate the different endotypes beside the IgE and non IgE associated forms. The aim of this review is to point out to the most recent knowledge in this field, which makes AD more difficult to frame in a unique clinical entity.

Keywords: Atopic dermatitis, endotype, filaggrin, IgE, immune system, skin barrier, Th2.

INTRODUCTION

In 1970, few years after the discovery of IgE (1966), Johansson et al. published a paper where, describing the spectrum of diseases with high IgE levels, indicated among these atopic eczema and classified it into two subgroups: atopic eczema (frequently with high IgE) and non-atopic eczema (a skin disease with the same features - skin lesions and their distribution pattern- but without evidence of IgE sensitization to aero- or food allergens) [1]. This was the first observation in which a difference in atopic eczema was underlined after the discovery of IgE. Indeed, in 1933, Wise and Sulzberger had already proposed the definition of “atopic dermatitis” (AD) to emphasize its close association with other atopic diseases, especially allergic rhino-conjunctivitis and asthma [2]. The possible link with the immune system and allergic diseases stimulated numerous studies to investigate the underlying pathogenic mechanisms.

THE FIRST IMMUNOLOGICAL STUDIES

In the course of the years, the immunological studies on AD followed the progressive update knowledge: among the earlier ones is paradigmatic the work published in 1975 by the group guided by Rebecca Buckley, that described an alteration of cell-mediated immunity in inverse relationship with the level of total IgE, detected analyzing peripheral white cells of subjects with atopic eczema [3]. Thus far, in these initial studies lymphocytes were differentiated into two main sub-groups through the formation of rosettes E. However, these researches strongly reflected on the clinical level: in 1978 the Lancet published a double-blind trial conducted by the Institute of Child Health in London [4]. In this study, authors highlighted the advantages of an exclusion diet in children suffering from AD with more severe clinical features; these data have been later confirmed by further observations. These studies lead to profound changes in clinical practice: from this time the elimination diet had a stronger impact on the therapeutic strategies of paediatricians.

In order to better define the disease, clinician tried to separate the two distinct forms of AD: one characterized by the presence of high IgE and associated allergic manifestations (also named “extrinsic AD”) and the other one with normal IgE without allergic symptoms (or “intrinsic AD”), with a high prevalence of the former type but different percentage in relation to age and severity of the disease (Table 1) [5-16].

In the meantime, the immunological network incredibly complicated: with the advent of monoclonal antibodies an important step in the knowledge of the immune system had been made. It was possible to distinguish different subtypes of lymphocytes with different functions; this allowed much more accurate results to be obtained, and moved the target organ from the blood to the skin.

In 1981 the first data on T cell subsets using monoclonal antibodies in AD patients appeared: Leung et al. reported data about in vitro cellular reactivity from 22 patients with AD by using monoclonal antibodies to recognize different peripheral lymphocytes [17]. Patients with AD showed a lower rate of T3+ cells (now named CD3+) and T8+ cells (now named CD8+) but not of T4+ (now named CD4+)
especially and 5 IFN INF eosinophils with increase and of vivo inverse numbers.

Immune- 

At the Laske Hochreutener Walker Akdis Cabon 13 13 model: of skin patients showed the 

1. IFN mRNA. as the list of secreting higher of IL-10 higher of IL-4, IL-5 and IL-13 in response to specific allergens, but decreased number of T cells producing IFN-γ in peripheral blood samples of patients with AD [19, 20].

At the same time the immunological studies were also directed to the comprehension of skin function; it was found that the skin cells were able to recognize antigens and to elicit a systemic immune response. Spargel et al. [21] described the mechanism of epicutaneous sensitization in a murine model: the induction of allergic sensitization by ovalbumin, a well known allergenic protein, leads to an increase in levels of total and ovalbumin-specific IgE and to the development of the atopic dermatitis skin lesions, with a local appreciable infiltrate of CD3+ T cells, eosinophils and neutrophils, and expression of IL-4, IL-5 and IFN-γ mRNA. These cutaneous immunohistological features corresponded to the Th2 response (increased IL-4, IL-5, and IL-13 levels) of the acute phase of AD, while chronic lesions showed meanly a prevalence of Th1-cytokines (IL-12 and IFN-γ). The two types of AD did not show dramatic differences in the expression of inflammatory cytokines: IL-5 and IFN-γ were detectable in similar amounts, while IL-4 and IL-13 showed a lower expression in non IgE AD, especially in the lesional skin.

### Table 1. Prevalence of non IgE-associated type atopic dermatitis (AD) in different studies.

<table>
<thead>
<tr>
<th>Reference</th>
<th>No. of Patients</th>
<th>Pts with Non IgE-Associated AD, n (%)</th>
<th>Age (Years)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kägi et al. 1994 [8]</td>
<td>33</td>
<td>14 (42%)</td>
<td>19-55</td>
</tr>
<tr>
<td>Cabon et al. 1996 [9]</td>
<td>59</td>
<td>27 (45%)</td>
<td>0-12</td>
</tr>
<tr>
<td>Wüthrich 1999 [10]</td>
<td>93</td>
<td>17 (18%)</td>
<td>37</td>
</tr>
<tr>
<td>Fabrizi et al. 1999 [12]</td>
<td>72</td>
<td>8 (11%)</td>
<td>36</td>
</tr>
<tr>
<td>Akdis et al. 1999 [13]</td>
<td>1151</td>
<td>117 (10%)</td>
<td>11-51</td>
</tr>
<tr>
<td>Laske &amp; Niggemann 2004 [14]</td>
<td>345</td>
<td>93 (27%)</td>
<td>1-24</td>
</tr>
<tr>
<td>de Benedictis et al. [15]</td>
<td>2184</td>
<td>764 (35%)</td>
<td>1-2</td>
</tr>
<tr>
<td>Ricci et al. 2014 [16]</td>
<td>184</td>
<td>15 (8%)</td>
<td>8-18</td>
</tr>
</tbody>
</table>

comparing to controls. An interesting finding was the higher T4+/T8+ ratio observed in 17 of 22 patients with atopic dermatitis but not in the control subjects.

The main characteristics that suggested an immune pathogenesis in AD were summarized by Donald Leung in 1995 as listed below: increased IgE levels, skin test sensitization to multiple allergens, higher spontaneous histamine release by basophils, lower number of CD8 suppressor/cytotoxic lymphocytes with less effective function, higher expression on mononuclear cells surface of the low-affinity receptor for IgE CD23, increased rate of Th2-like cells secreting IL-4 and IL-5, and decreased rate of numbers of those ones secreting IFN-γ with a significant inverse correlation between in vitro IFN-γ production and in vivo IgE serum concentrations in AD patients [18]. Furthermore, other studies confirmed the increased number of T cells producing Th2-like cytokines such as IL-4, IL-5 and IL-13 in response to specific allergens, but decreased number of T cells producing IFN-γ in peripheral blood samples of patients with AD [19, 20].

At the same time the immunological studies were also directed to the comprehension of skin function; it was found that the skin cells were able to recognize antigens and to elicit a systemic immune response. Spargel et al. [21] described the mechanism of epicutaneous sensitization in a murine model: the induction of allergic sensitization by ovalbumin, a well known allergenic protein, leads to an increase in levels of total and ovalbumin-specific serum IgE and to the development of the atopic dermatitis skin lesions, with a local appreciable infiltrate of CD3+ T cells, eosinophils and neutrophils, and expression of IL-4, IL-5 and IFN-γ mRNA. These cutaneous immunohistological features corresponded to the Th2 response (increased IL-4, IL-5, and IL-13 levels) of the acute phase of AD, while chronic lesions showed meanly a prevalence of Th1-cytokines (IL-12 and IFN-γ). The two types of AD did not show dramatic differences in the expression of inflammatory cytokines: IL-5 and IFN-γ were detectable in similar amounts, while IL-4 and IL-13 showed a lower expression in non IgE AD, especially in the lesional skin.

### THE ADVENT OF GENETIC STUDIES

The new advance on basic science is promptly reflected on clinical research: with the advent of the new technologies of gene analysis, the skin barrier became the protagonist. In 2002, Coxson and Moffatt [22] understood that a gene or a cluster of genes, coding for proteins involved in the formation of the deep layers of the skin, may play a key role in the pathogenetic mechanisms of atopic dermatitis, including allergic sensitization. But only in 2006 data from a cohort of Irish patients with AD were published in Nature Genetics [23]: authors found alterations in the nucleotide sequence of the gene encoding for filaggrin. This protein is the most important among those of the so called “epidermal differentiation complex” and is fundamental to preserve the skin barrier integrity. Moreover, filaggrin is able to promote the aggregation of the keratin filaments and its functional defects have described as a risk factor for the onset of AD. Furthermore, in subjects with AD two variants of this gene (R510X and 2282del4) were associated with the occurrence of allergic asthma. Otherwise, these alterations of the gene of filaggrin were observed in not more than 30% of patients, so that was insufficient to explain the pathogenesis of AD. Later, further studies on additional proteins that might contribute to the integrity of the skin barrier had been performed, and other skin proteins (e.g. loricrin, claudine 1) have been associated to the development of AD [24, 25].

### THE IMMUNE RESPONSE: WHAT WE KNOW NOW

The contribution of multiple cell types, and the existence of multiple cytokine patterns at different evolution stages give an idea of the high complexity of the mechanisms involved in the immune response in patients with AD.

#### The Skin Immune Response

A model to study the immune response is to analyze the cellular skin infiltrate after atopy patch test (APT) to house dust mites (HDM). Epicutaneous application of HDM frequently induces eczema in the nonlesional skin of 40-50% of patients with AD. By using immunohistochemistry and molecular analysis, it has been shown that also cytotoxic T
cells are implicated in the pathogenesis of these lesions; the analyses from a murin model showed that very few CD8+ T cells infiltrating the skin (about 5% among the CD45+ cells) are sufficient to trigger the HDM-induced AD lesions and their presence is even indispensable, because mice depletion with anti-CD8 monoclonal Abs completely suppressed the inflammatory process [26]. In a recent study, patients with mild to moderate AD were enrolled on the basis of the positivity of APT to HDM; in nine patients the skin biopsy showed that CD8+ T cells are involved in the early phase of the response to allergen exposure. A hypothesis is that the apoptosis of keratinocytes and the epidermal spongiosis, which are both pathological hallmarks of AD, are due to the cytotoxic activity of CD8+ cells [26].

Recently, in vivo studies evidenced new relevant details in the skin immune response. In certain allergic tissue reactions potent agents with vasodilator and permeability functions were recognized to be expressed by the inflammatory cells in the lesional skin, such as the calcitonin gene-related peptide (CGRP) and the vascular endothelial growth factor (VEGF). Skin biopsy specimens from atopic dermatitis lesions were collected after various times from the cutaneous allergen exposure and analysed by using single and double immunohistochemistry and in situ hybridization: neutrophils and CD3+ T lymphocytes were the main CGRP+ cells detected at the late-phase of the reaction (i.e., 6 hours) [27]. In the setting of allergic inflammation, the wide CGRP production by neutrophils may clarify the characteristic vasodilatation that can be observed in the late-phase of the skin reaction: thus it could be at least in part a neutrophil-dependent phenomenon [27]. Otherwise, in patients with chronic AD and in those with psoriasis, the lesional skin biopsy specimens show that dendritic cells (DC) did not increase a preferential T-cell subset in a disease-specific manner. The capacity of each DC subset to increase Th1, Th2, Th17 and Th22 subsets was the same in the two diseases, but an upregulation of specific chemokine expression such as CCL17, CCL18, and CCL22 was observed only in patients with AD [28]. Moreover, in patients with AD, cutaneous biopsy showed an impaired IFN-γ-mediated signaling pathway and a decreased IFN-γ production both in DCs and in their precursor cells; this condition might contribute to the Th2 bias [29]. On the other side, the increased IFN-γ responses suggest the role of multiple new factors involved in the mechanisms of apoptosis and inflammation in the development of AD [30].

Also other mediators, such as TNF-α and the TNF-like weak inducer of apoptosis (TWEAK), cooperate in the induction of keratinocyte apoptosis and in the lesional production in AD patients [31]. Indeed, during disseminated viral infections such as eczema herpeticum, which is caused by herpes simplex virus, patients suffering from AD showed a mixture of defects both in the skin barrier function and in the innate and adaptive immune responses. In particular, an impaired IFN-γ response was observed in human AD complicated by eczema herpeticum: indeed genetic variants as single nucleotide polymorphisms were found in IFN-γ and IFNGR1 genes and they were significantly related with eczema herpeticum and abnormal IFN-γ production [32].

Also mutations of STAT6 (signal transducer and activator of transcription 6) gene increase make patients with AD more prone towards disseminated viral skin infections [33]. Also the cellular transcription factor Specificity protein (Sp)-1 is involved in diverse cellular functions and represents a critical player during the antiviral responses of skin keratinocytes. Sp1 deficiency in AD patients with viral infection may contribute to increase the risk to develop a disseminated infection of the skin [34].

Recently, the molecular and cellular pathogenetic mechanisms of lesional and nonlesional AD (intrinsic and extrinsic) have been studied in 51 patients with severe AD by using the gene expression assay (real-time PCR). While a prominent infiltrate of T cells and DCs was observed in lesional skin of both types of AD, patients with the intrinsic form (n=9) showed a Th17 immune response more increased than those with extrinsic AD (n=42) [35]. Moreover, higher activation of all inflammatory axes, with a particular involvement of Th17 and Th22 cytokines, including the Th2 products, was detected in patients with intrinsic AD. A positive association between Th17-related molecules and severity of AD, assessed by SCORAD index, was found only in patients with intrinsic AD, whereas patients with extrinsic AD showed a characteristic positive association between the SCORAD index and Th2 cytokine levels (IL-4 and IL-5) and inverse association with proteins involved in the skin differentiation (e.g. loricrin and perilipin) [35].

**AGAINST THE BIPHASIC MODEL**

The current model to describe the skin inflammation in AD largely descends from experimental studies performed by using APTs with environmental allergens to induce acute lesions and simulate the acute phase of the disease [36, 37].

As previously described, AD pathogenesis is characterized as a biphasic T cell-mediated disease: an early Th2 pattern which predominates in the acute phase and a late Th1 pattern which prevails in the chronic phase [36, 37]. On the basis of an experimental model comparing spontaneous acute AD lesions with chronic lesions from the same patient should permit the develop of a new viewpoint [38]. This new perspective to explore this mechanism was recently performed: biopsy specimens from acute lesion, chronic lesion (>72 hours duration), and nonlesional skin and blood samples from the same patient were collected from 17 patients with moderate-to-severe AD. A significant increases in gene expression levels of Th2 (i.e. IL-4, IL-13, and IL-31) and Th22 (i.e. IL-22) cytokines was associated with the onset of acute lesions; the quantitative gene expression showed also a raise of some other inflammation products as IL-31, IL-22, S100A7, S100A8, and S100A9 with a positive correlation between the SCORAD index and IL-22 mRNA expression [38]. Th1 products induced by interferon, in particular IFN-γ, was also decreased in acute skin lesions. A small increase of Th17-cytokines were observed in acute disease associated with a higher increase in IL17-regulated products (CCL20, peptide inhibitor 3 elafin, and lipocalin 2). Instead IL-22 mRNA and its associated products (S100A7, S100A8, S100A9, and IL-32) showed a progressive increase and were detected both in acute and in chronic lesions [39]. The chronic phase is also characterized by an intensified release of Th2-related cytokines with the exception of IL-4; indeed a decrease of IL-4 levels has been observed from acute to chronic lesions [38]. The result of skin inflammation is a marked activation of the gene cluster codifying for proteins of the Epidermal differentiation
immune and underlying inoculation, essential elevated eosinophils in inflammatory hyperexpression Immune expression synthesis a other action nonatopic in healthy Immune receptors is number and activity of microbial proteins: immunological and observations of healthy the epidermis of skin infections. This observation suggests that the mechanism underlying cathelicidin deficiency is insufficient alone to explain the pathobiology of disseminated skin infections and other immunological alterations should be involved in patients with eczema herpeticum. The action of cathelicidin seems to be different but complementary to the mechanism of action of IFN: indeed while cathelicidin acts extracellularly by damaging the viral envelope, IFN inhibits the intracellular viral transcription and translation. Therefore in patients with AD both intracellular and extracellular defects in the antiviral response of the host are involved in increasing the susceptibility to viral infections. Keratinocytes can modulate the release of inflammatory mediators (e.g., cytokines, chemokines, and AMPs) through the expression of specific receptors for the effectors of the innate immune response overall defined pattern recognition receptors (PRRs). The relation between AD and the innate immune receptors system has been clearly reviewed by Kuo et al., who remarked their synergic action in the production of pro-Th2 cytokines, as the Thymic Stromal Lymphopoietin (TSLP), IL-25 and IL-33, which drive the immune response towards a Th2 pattern.

TSLP is preferentially produced by epithelial cells such as keratinocytes and mucosal cells of the airway; the TSLP expression level is increased in the epidermal layers of AD patients if compared with healthy subjects and it shows a correlation with SCORAD index, in particular with the dry skin score.

In 10 adult patients with mild to moderate AD, it was observed that the epicutaneous application of HDM using the APTs promotes the induction of TSLP and of CCL17/Thybus and activation-regulated chemokine (TARC), as a potential indicators of TSLP bioactivity.

ADAPTIVE IMMUNE SYSTEM

After the review article by Leung et al. issued in 1995 [18], an intriguing paper by Akdis et al. published in 1999 [13]. The authors investigated the immunologic mechanisms among 1151 chronic AD patients (10% of them with non-IgE associated AD) and they found that skin T cells were always implicated in both subtypes of AD, responding to staphylococcal enterotoxin B, superantigens and cytokines (IL-2, IL-5, IL-13 and IFNγ). Interestingly, skin T cells from non-IgE associated form expressed lower IL-13 and IL-5, while IL-4 was not found in any of the two types. Moreover, authors found a relevant expression of CD23 in the activated B cells of patients with IgE associated AD, while the non-IgE associated form was mainly characterized by the absence IL-13-induced-B cells and subsequent IgE secretion.

Recently, new lymphocytes' phenotypes and cytokines have been identified. Th22 cells, originally identified as circulating T-cell clones with skin-homing properties, express receptors for chemokines (CLA, CCR4, CCR6 and CCR10.7) and do not coproduce IL-17, IL-4, or IFNγ. In thirteen patients with severe chronic AD a raised rate of T cells characterized by skin-homing capability and expressing both IL-13 and IL-22 was found. Indeed, this peculiar subtype of T-cells named “IL-13/IL-22-coproducing T cells” might act a potential key role in the pathogenesis of the disease.

Furthermore, not only Th2 cells have been identified as producers of IL-31, since also DCs, monocytes and mast cells are implicated in IL-31 synthesis: these cells have been isolated in the skin of subjects affected by AD and the expression of IL-31 was found to be raised also in the sera of these patients with a strong correlation with the grade of severity of the disease [59]. In addition, IL-31 plays a relevant role also in the process of skin differentiation: in a human 3-dimensional skin model filaggrin was observed to be downstream regulated by IL-31 [59]. Also the role of IL-10 is crucial in modulating the adaptive immune system: IL-10 owns suppressive properties both on DCs’ maturation and subsequent cytokines expression, and on Th1 cell differentiation. Simultaneously, it contains properties of effector T-cells and enhances the function of Tregs, suggesting that the interaction between the different cell subsets (DCs, Tregs and effector T cells) is crucial in down-regulating
an unbalanced activation of the immune system [60]. Studies on IL-10-deficient (IL-10−/−) mice have shown that DC-specific IL-10R−/− rodents have an intensified hypersensitivity reactions towards contact hapten, arguing that the pathway of IL-10 signalling is crucial in limiting the contact hypersensitivity reaction, while it is not essential in T-cell priming [61]. In the end, IL-10, suppressing the expression of proinflammatory cytokines from monocytes and macrophages, plays also a critical role in modulating the responses of both the adaptive and innate immune systems [60, 61].

Also the interactions between the endocrine and the immune systems play an influential role in AD [62]: levels of corticotropin-releasing hormone (CRH) receptor expressed by T cells were significantly lower in subjects with AD compared to controls. In contrast, in the healthy population, CRH induces the upregulation of IL-4 by Th2 cells and the inhibition of Th1 cells-induced IFN-γ secretion. In subjects with AD, T cells were not shown to secrete IL-4 and IFN-γ after CRH treatment, while CRH significantly inhibits the synthesis of IL-10 from Tregs, giving an explanation to the stress-induced reoccurrences of AD skin lesions [62].

The relation between immune system and the development of AD skin lesions has been distinctly observed in a murine model showing the importance of the adaptive immunity in the spontaneous appearance of skin lesions. Filaggrin-deficient flaky tail (ft) mice (Flgft/ft) show spontaneous skin inflammation that imitates human AD. The breeding of Flgft/ft with Balb/c Rag2-deficient (Rag2−/−) generated mice with a lack of function of filaggrin as well as of T and B lymphocytes (Rag2−/−/Flgft/ft) [63]: these Rag2−/−/Flgft/ft mice didn’t develop skin lesions during the observation period of 32 weeks, confirming that T cells and the adaptive immune system give an essential contribution to the development of skin inflammation [63].

A link between cytokines and AD has been shown by recent results from genome-wide association studies [64]. Data were obtained from a public repository about independent populations from two birth cohorts from the United Kingdom (ALSPAC, 12 set 7) and Germany (Multicenter Allergy Study [MAS]): a significant association was detected for 318 genetic markers related to the expression of pro-inflammatory genes, identifying novel genetic risk factors for AD. A mutation in the amino acid sequence of the IL-6 receptor (IL-6R Asp358Ala; rs2228145) resulting in a functional change of the protein, has been significantly associated with AD, in particular with the persistent forms [64].

THE INTERACTIONS AMONG SKIN IMMUNE RESPONSE AND THE INNATE AND ADAPTIVE IMMUNE SYSTEM

The interplay among skin barrier and the innate and adaptive immune system in AD has been clearly summarized in recent review articles [65, 66]. When the barrier function is compromised, hapten and protein antigens easily penetrate in the skin, promoting the switch toward a Th2-cytokine pattern. Th2 (IL-4, IL-13, and IL-31) and Th22 (IL-22) cytokines act as suppressors of the major final differentiation of skin proteins in keratinocytes (i.e., filaggrin and loricrin), enhancing the barrier disruption [35, 38, 63]. Moreover, IL-4 and IL-13 have inhibitors capacity on the production of AMPs, directly inhibiting the expression of HBD-2 and HBD-3 and, indirectly, of hBD-3 and LL-37 (via inhibition of STAT6 activation) and subsequently suppression of TNF-α/IFN-γ and IFN-γ production. On the other hand, IL-17 and IL-22 cytokines, derived from the recently identified T-cell subsets, have many effects on epidermal keratinocytes, enhancing the production of proinflammatory mediators such as S100A7, S100A8, and S100A9 proteins. In addition, keratinocytes by themselves release cytokines, chemokines and AMPs, which enhance the production of pro-Th2 cytokines, including TSLP, IL-25, and IL-33, and drive the immune response to surface antigens toward a Th2 profile. Finally TSLP stimulate DCs and induces the expression of cell-surface activation markers to promote Th2-skewed inflammatory responses (Table 2).

Table 2. Major interactions between skin barrier/innate immunity and adaptive immunity in AD patients.

<table>
<thead>
<tr>
<th>Main Links Between Innate and Adaptive Immunity in AD</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Eosinophils act as immunoregulatory cells, playing a role in the switch from a Th2-cytokine pattern in acute lesions of AD to a more Th1-like pattern in the chronic stage</td>
</tr>
<tr>
<td>2. TSLP (secreted by keratinocytes) stimulate DCs and induces the expression of cell-surface activation markers to promote Th2-skewed inflammatory responses</td>
</tr>
<tr>
<td>3. Keratinocytes release cytokines, chemokines and AMPs, which acts producing pro-Th2 cytokines, including TSLP, IL-25, and IL-33, and drive the immune response to surface antigens toward a Th2 profile.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Main Links Between Adaptive Immunity and Skin Barrier/Innate Immunity in AD</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. IL-4, IL-13 cytokines have inhibitors capacity on epidermal differentiation and production of antimicrobial peptides; they could inhibit the expression of hBD-2 and hBD-3 and might indirectly inhibit hBD-3 and LL-37 via inhibition of STAT6 activation and subsequently inhibit TNF-α/IFN-γ production</td>
</tr>
<tr>
<td>2. IL-17 and IL-22 cytokines derived from the recently identified T-cell subsets have many effects on epidermal keratinocytes, S100A7, S100A8, and S100A9 mRNAs (human keratinocytes).</td>
</tr>
<tr>
<td>3. Tα2 (IL-4, IL-13, and IL-31) and Tα22 (IL-22) cytokines were shown to suppress the major final differentiation proteins (i.e., filaggrin and loricrin)</td>
</tr>
</tbody>
</table>

DCs: dendritic cells; TSLP: thymic stromal lymphopoietin.
FROM THE OLD NOMENCLATURE TO THE “UNIFYING HYPOTHESIS”

In the last years the nomenclature and classification of AD has been revised: initially AD was classified in two subtypes named “extrinsic” (associated with atopic sensitization) and “intrinsic”. The intrinsic type was not associated with specific IgE sensitization and with any of the clinical manifestation of other atopic diseases and was characterized by normal levels of total serum IgE. Later, the nomenclature changed and the current classification called the extrinsic form as “IgE-associated” and the intrinsic one as “non-IgE-associated AD” [67, 68]. However, a recently proposed model unified these two subtypes, considering them as different stages of the same pathogenetic process: indeed, while in the early infancy the non-IgE associated form is predominant, at a later stage the allergic sensitization occurs in most of the patients determining the switch to the IgE-associated phase [69-71].

CONCLUSION

AD is a complex disease in which many actors contribute to develop the clinical phenotypes. Nevertheless, many of these are not directly bound to the pathogenetic mechanism implicated: the two main actors are the primitive skin barrier defects (in particular the alterations of the filaggrin gene) and the immune system Th2-directed (i.e. cells, cytokines and chemokines allergic-oriented). The primitive barrier defect facilitates the passage of allergens and antigens; this contact activates innate immunity to develop a Th2-oriented response. On the other hand, allergic Th2 cytokines may downregulate the production of barrier proteins facilitating the income of allergens. These two main alterations may interact with different proportion each other (Fig. 1) determining many potential distinct endotypes, even if the phenotypic appearance of the skin shows similar characteristics in both pathways.

It is difficult to translate in clinical practice such complex pathogenesis: the majority of patients (about 80%) have an “IgE-associated AD”, as recently suggested to be defined by Thomas Bieber [71]. Some patients, with the “non-IgE associated” form, seem to show an immune alteration profile only at the skin level, where together with the increase of IL5 and IL13, a wider production of IL17 appears to be stimulated, supporting the hypothesis of a link with an autoimmune course. Since skin disruption can easily allow allergen penetration and the local keratinocytes can promote an allergic immune response, the clinical intervention should be addressed to the strict control of the skin inflammation by the application of emollients with antiinflammatory and proactive properties in order to interrupt this mechanism. On the other hand, a wider effort should be addressed to evaluate which is the allergic sensitization pattern of the patient, in particular during the paediatric age, since at this stage preventive strategies are still possible and effective. Meanwhile, how much the sensitization directly influences the course of AD should be determined by the physician on the basis of the clinical severity of the diseases and on the results of the allergic response.

ABBREVIATIONS

AD = Atopic dermatitis
AMPS = Antimicrobial peptides
APT = Atopy patch test
CGRP = Calcitonin gene-related peptide
CRH = Corticotropin-releasing hormone
DC = Dendritic cell
HBD = β-defensin
HDM = House dust mites
PRRs = Pattern recognition receptors
Sp-1 = Specificity protein-1
STAT6 = Signal transducer and activator of transcription 6 gene
TSLP = Thymic stromal lymphopoietin
TWEAK = TNF-like weak inducer of apoptosis
VEGF = Vascular endothelial growth factor

CONFLICT OF INTEREST

The authors confirm that they have no conflict of interest.

ACKNOWLEDGEMENTS

Declared none.

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