Skin Immunodysregulation Disorders - The T-Cell Mediated Dermatoses

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Introduction

The skin as an immunological organ

The skin harbors an immune organ, made up of keratinocytes, dendritic cells (also known as Langerhans cells), as well as immune-derived nomadic cells (e.g. lymphocytes, neutrophils and eosinophils), which can respond to a variety of insults including inflammation, wound repair and infection. In addition to playing a role in defense, the skin’s immune system also modulates the growth of commensal microbiota. Keratinocytes monitor colonizing bacteria, fungi and viruses using Pattern Recognition Receptors (PRRs) including Toll-Like Receptors (TLRs), mannose receptors and NOD-like receptors. These receptors recognize a variety of Pathogen-Associated Molecular Patterns (PAMPs), including flagellin, nucleic acids, lipopolysaccharide (from gram-negative bacteria), peptidoglycan and lipoteichoic acid (from gram-positive bacteria), as well as mannann and zymosin (from fungal cell walls), causing the activation the skin’s innate immune response [1-4].

The innate immune system

Upon stimulation from outside threats, a response occurs known as the innate immune system. “Innate” refers to the skin’s inherent ability to ward off invading microorganisms, as well as environmental threats such as allergens [1,3,5]. In the innate immune response keratinocytes and surrounding fibroblasts secrete Anti-Microbial Peptides (AMPs), as well as inflammatory cytokines and chemokines, to create a chemical milieu allowing for the direct interplay between keratinocytes, immune cells and microorganisms. AMPs bind to the cell surfaces of microorganisms, allowing immune cells to directly target and destroy bacteria, fungi and enveloped viruses [1,3-5].

It is important to note that the skin’s immune system can discriminate between harmless commensal versus harmful pathogenic microorganisms, a response reliant on TLRs. Desensitization of TLRs (ergo tolerance) occurs either through decreased receptor expression or increased inhibitory regulators [such as IL-1-receptor-associated kinase 3 (IRAK3/IRAK-M) or Suppressor of Cytokine Signaling 1 (SOCS1)] in response to prolonged exposure to commensal microorganisms. The activation of TLR signaling by commensal bacteria has also been shown to be necessary for cell survival and repair during infection. On the other hand, specific recognition by Pathogen-Associated Molecular Patterns (PAMPs) allows for immune system to target pathogenic species of bacteria, fungi and viruses [1-4,6,7].

Commensal skin flora also plays a role in skin defense. Staphylococcus epidermidis has recently been demonstrated to modulate the host innate immune system. For instance, S. epidermidis secretes modulins that inhibit pathogenic strains of bacteria including S. aureus and group A Streptococcus, lipoteichoic acid to inhibit skin inflammation through TLR-2/-3-mediated pathways, and increases keratinocyte expression of AMPs through a TLR2 dependent mechanism to enhance pathogenic microorganism destruction [8].

The Role of the Microbiome in Immunodermatoses

A microbiome is defined as a mini-ecosystem or collection of symbiotic, commensal and pathogenic microorganisms. *Corresponding author: Sharon E Jacob, MD, Department of Dermatology, Loma Linda University, VA Loma Linda, 11201 Benton Street, Loma Linda, CA, 92357, USA, Tel: (909)-558-2890, E-mail: sjacob@contactderm.net

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Microbes and microorganisms (bacteria, fungi and viruses) inhabiting an environment, in the skin’s case the stratum corneum [8]. The skin microbiome is influenced by a multitude of endogenous (genetic polymorphisms, AMPs, native immunity), as well as exogenous (hygiene hypothesis, hand washing, physical and chemical irritants) factors. Recent mouse models suggest that local skin flora balance is closely associated to effector and regulator T cell populations [8].

Disease states also affect the composition of the skin microbiome. For instance, patients with Atopic Dermatitis (AD) have increased susceptibility to S. aureus infection at sites of excoriation and bacterial inoculation. Upregulation in the expression of interleukins 4 and 13 (IL-4, IL-13) results in decreased epidermal barrier and thus susceptibility to microbial invasion. In addition, AD patients have excessive fungal colonization by Malassezia globosa and restricta leading to a distorted and non-diverse bacterial microbiome, with decreased colonization by organisms such as Clostridium. This lack of diversity allows for increased colonization by S. epidermidis, and infection by S. aureus [8,9].

Although less is known regarding the psoriasis microbiome, some researchers claim psoriasis patients have decreased Staphylococcus and propionibacteria species compared to healthy skin samples. During psoriasis exacerbations, AMPs such as cathelicidin and human β-defensin 2 are upregulated, suggesting an underlying disturbance in the skin microbiome. New studies have linked antibiotic use with increased incidence of psoriatic arthritis; although it is hypothesized that this mechanism might be related to alterations in the microbiome, no clear mechanism has been elucidated [8].

In addition to the localized skin microbiome, researchers have hypothesized that global microbiota may also play a role in the development of inflammatory skin diseases. The administration of Lactobacillus reuteri decreased the incidence of IgE-mediated AD, suggesting long-term alterations in the intestinal microbiome may decrease inflammatory efforts mediated by T helper-2 (Th2) cells [10]. Applying kefir, a fermented dairy product, to burn wounds has been shown to increase collagen formation and epithelialization through its anti-bacterial and anti-inflammatory properties [11]. Furthermore, the ingestion of probiotics (which include S. epidermidis) may help with neutrophil recruitment, disrupting biofilms, and increasing the release of anti-microbial peptides by keratinocytes and fibroblasts [12].

### Dysregulation of the Skin Immune Response and Skin Disorders

In inflammatory dermatitis and dermatoses states, the innate immune system may become dysregulated, for example, increased activity is seen in psoriasis and decreased activity is seen in AD [3,9,13] (Table 1). These changes lead to alterations in the cutaneous microbiome, and in some cases this may be a factor in disease activity. It is also possible that changes in the cutaneous microbiome, independent of the innate immune system, could also be a driving factor in some pathologic states.

#### Table 1: Comparing T-cell populations and cytokine production in select T-cell mediated dermatoses.

<table>
<thead>
<tr>
<th>Dermatoses</th>
<th>Predominant T cell population</th>
<th>Primary inflammatory cytokines</th>
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<tbody>
<tr>
<td>Psoriasis (Pso)</td>
<td>Th1 (initial)</td>
<td>CCL20</td>
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<td></td>
<td>Th17, Th22 (disease memory and recurrence)</td>
<td>IFN-γ</td>
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<td>TNF-α</td>
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<tr>
<td>Atopic Dermatitis (AD)</td>
<td>Th2 (initial)</td>
<td>IFN-γ</td>
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<td>Th1 (chronic)</td>
<td>IL-6</td>
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<td>TSLP</td>
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<tr>
<td>Allergic Contact Dermatitis (ACD)</td>
<td>NK cells</td>
<td>IL-17</td>
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<td>Th17</td>
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This table summarizes the differences in T-cell population and inflammatory cytokine expression pattern in psoriasis, AD and ACD. Psoriasis initially has Th1 cell predominance, whereas AD initially has Th2 predominance. ACD is propagated by dysregulation of NK and Th17 cells.
Psoriasis

Psoriasis affects approximately 2.5% of the global population. The most common form of psoriasis, “plaque” psoriasis, accounting for 90% of clinical psoriasis presentations, presents as sharply demarcated erythematous plaques with an overlying adherent silver scale [13,14]. These plaques are often located on extensor surfaces, such as the elbows and knees. In addition to the above “plaque” psoriasis, other subtypes exist including: “inverse” in which the axilla, groin, and gluteal cleft/folds are involved; “pustular” presenting with monomorphic pustules of the palms/soles which can then generalize throughout the body; “guttate” characterized by drop-shaped plaques after Streptococcus infection or a rapid systemic steroid taper; “sebo-” affecting the scalp, inner ear, face and skin folds; “oral” characterized by the fissured tongue; and “nail” with evidence of multiple pits or oil spots. Psoriatic Arthritis (PsA) affects about 30% of psoriasis patients and involves symmetric small joints, the sacroiliac joints and the spine [13,14].

Psoriatic lesions produce a large amount of AMPs and are characterized by a highly active innate immune response with a resulting shift in the resident T lymphocyte population favoring initially a Th1 subtype, and later, Th17. Antigens that have been linked to T cell activation in psoriatic patients include elements of the stratum corneum, squamous cell carcinoma antigen, nuclear ribonucleoprotein A, Heat Shock Protein (HSP), type I keratin (molecular mimicry due to similarity in structure to the Streptococcus M protein), keratin 17, human papilloma virus-5 (HPV-5) Deoxyribonucleic Acid (DNA), antimicrobial peptide cathelicidin (LL37; produced by keratinocytes), and ADAMTS-like protein 5 (produced by melanocytes). Increased amounts of TNF-α, IL-23 and IL-17 cause T cell activation, with overexpression of PAMs and TLR pathways, resulting in chronic inflammatory changes of the skin [13,14].

The early activation of T-helper 1 (Th1) cells by endocytosis of a TLR-9-antigen complex, induces the expression interferon-γ (IFN-γ) causing dendritic cells to produce Chemokine Ligand 20 (CCL20) and IL-23 leading to the expansion of the Th17 cell population. Increases in IL-17 production by Th17 cells leads to a chronic inflammatory cascade, including Th21/IL-21 mitogenic effects on keratinocytes and Th22/IL-22 keratinocyte activation and epidermal acanthosis. Once the chronic inflammatory pathway has been set into motion, Th22 and Th17 lymphocytes play a role in disease memory and allow for recurrence of psoriasis at previously affected sites (Figure 1) [13,14].
Atopic dermatitis

AD is a chronic, relapsing inflammatory dermatitis that affects approximately 2% of American adults and 15% of children [17,18]. AD presents as erythematous pruritic patches and plaques, most commonly found on the flexural surfaces, such as the axillae, antecubital fossae, and popliteal fossae. In addition to dysregulation of the innate immune system, AD is often also associated with changes in the skin microbiome affecting colonization and rates of infection [8,17,18].

Unlike psoriasis, AD lesions are associated with low levels of AMP. The loss of AMP expression may be a contributing factor to the decrease in cutaneous microbiota diversity [8,9]. It is estimated that 90% of AD patients are colonized with *S. aureus* on both lesional and non-lesional skin, versus less than 5% of healthy individuals. In comparison, normal, healthy skin flora is composed primarily of Actinobacteria, Firmicutes, Bacteroides and Proteobacteria species. This loss of microbial diversity is often amplified during acute AD flares; recent mouse models have shown that diversity can be restored using anti-inflammatory, antibiotic and Narrow-Band Ultraviolet B (NB-UVB) treatments [8,9,19]. In addition to changes in skin flora, researchers have also noted that the gut microbiome of AD patients differs in that there is an...
subsequently increases transepidermal water loss, allergen penetration with increasing levels of Immunoglobulin E (IgE), and colonization of the skin by *S. aureus* [9,19]. Other immune cell types such as eosinophils and basophils may contribute to host defense in the setting of decreased physical barrier and innate immunity. For instance, AD patients have been found to have tissue eosinophilia. These eosinophils deposit extracellular protein granules that may play a role in host defense. Basophils are recruited by IgE-activated mast cells and create anti-bacterial extracellular traps [9].

Traditional AD treatment options have focused on lipid repletion and heavy emollient use to help with loss of the physical barrier (emollients), shock treatment to acutely alter the pH of the skin (bleach baths and vinegar soaks), and anti-inflammatory agents (topical/systemic steroids, topical calcineurin inhibitors, and immunomodulators such as methotrexate, cyclosporine or mycophenolate mofetil) [20-22]. Early use of emollient has been shown to decrease risk of developing atopic dermatitis in children considered high risk given prior family history [23].

With an increase in the understanding of the patho-
Allergic contact dermatitis

Allergic Contact Dermatitis (ACD) most commonly presents as a relapsing dermatitis following exposure to specific allergens/haptens to which the person has been previously sensitized. Some of the most common reported allergens are metals (nickel, cobalt, chromium), flavor additives (balsam of Peru), plants (uroshiol found in the *Toxicodendron* genus of plants), topical antibiotics (neomycin, bacitracin), fragrance, colophony (rosin, fir), preservatives (formaldehyde, isothiazolinone), and rubber (mercaptobenzothiazole) [27,28]. Nickel ACD alone affects an estimated 2.5 million American adults and 250,000 children, costing about $5.7 million USD yearly for diagnosis and treatment [29].

ACD, purely an immunologic phenomenon, is classified as a type IV hypersensitivity reaction, in which there is an induction priming phase to the specific allergen, followed by the elicitation phase in which a response is triggered on re-exposure to the specific allergen. Water-soluble allergens/haptens cross the stratum corneum and, once engulfed, are combined with lipids from the cellular membranes of keratinocytes [30,31]. As part of the induction phase, antigenic epitopes are presented to naïve T cells by dendritic cells found in the skin. On re-exposure, these allergens cause activation of Natural Killer (NK) T cells initiating the elicitation phase [32].

New research suggests that there may be a further role in ACD for Th17 cells. Skin isolated from nickel ACD patients has shown to have increased expression of IL-17, which has downstream effects including neutrophil recruitment, as well as fibroblast and macrophage recruitment [33] (Figure 5).

The mainstay of treatment for ACD includes allergen/hapten avoidance, directed by confirmatory diagnostic patch testing. Topical steroids and calcineurin inhibitors may be used during the initial avoidance
phase for their anti-inflammatory effect. For severe cases, systemic immunomodulatory agents may be necessary. Currently there are no approved biologic therapies for ACD [27,34].

**Diagnosing T-cell mediated diseases**

The differential diagnosis for inflammatory skin diseases is wide and includes chronic inflammatory skin diseases to infectious diseases (e.g. tinea) to cutaneous lymphoma to medication reactions, and metabolic and congenital disorders. The diagnosis of immunologic dermatoses revolves around a thorough history and physical examination. It is important to query patient exposures and seasonal variations at time of symptom onset. Regional patterns and distributions can provide diagnostic clues, for example flexural surfaces are often affected by AD, extensor surfaces for psoriasis, while ACD often exhibits exposure pattern presentations (e.g. earlobes/neck/wrists for metal-associated jewelry reactions; exposed area dermatitis when there is aerosolization etiology; lips for flavoring compounds; and plantar food surface reactions when rubber components are the instigators).

If a history and physical does not reveal a clear source for the dermatitis, further tests may need to be performed. For instance, use a Potassium hydroxide (KOH) or mineral oil preparation to rule-out *Tinea* and scabies infections. An epicutaneous patch test is indicated when there is sustained or worsening chronic dermatitis [34]. A skin biopsy with stain for Hematoxylin and Eosin (H&E), can help with the diagnosis of specific pathologies such as psoriasis or malignancy, such as Cutaneous T Cell Lymphoma (CTCL) when combined with specific immunohistochemical stains. For instance, psoriasis is characterized by a hyperkeratotic epidermis with parakeratosis and neutrophils located in the stratum corneum [35]. On the other hand, CTCL may present with intraepidermal clusters of lymphocytes (Pautrier’s microabscesses) and perivascular lymphocytic infiltrate [36]. Although differences in histology may appear minute, they are important for the correct diagnosis, as well as appropriate and timely initiation of treatment.

**Conclusions**

The skin is a complex immunologic organ, able to protect the body from oncoming insults such as pathogenic microbiota and environmental insults. Increasingly, researchers are demonstrating the importance of the commensal skin microbiome in the immunologic response. Dysregulation of the innate immune system, causing changes in the resident T lymphocyte population with either a Th1 or Th2 predominance, creates fluctuations in inflammatory cytokine levels allowing for propagation of chronic inflammatory changes within the epidermis, subsequently causing the symptoms of psoriasis, AD, and ACD with which we are all familiar. Variations in the skin microbiome have been shown to play a role in disease processes such as psoriasis and AD. It is important to understand the underlying mechanisms of these T-cell mediated dermatoses when practicing diagnose these skin pathologies in a timely manner and determine the appropriate therapeutic option for patients.

**Figure 5:** Chronic inflammatory pathways of Allergic Contact Dermatitis (ACD).

Upon exposure to an allergen or hapten, dendritic cells present antigen to Th0 cells during the induction phase. On re-exposure, natural killer cells are activated and the elicitation phase begins. There is a possible role for Th17 cells in disease memory.
Conflicts of Interest

The authors have no conflicts of interest to disclose.

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References