A Calm, Dispassionate Look at Skin Microbiota in Atopic Dermatitis: An Integrative Literature Review

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ABSTRACT

Atopic dermatitis (AD) is a chronic common inflammatory skin disorder with clinical characteristics of pruritic, dry, and recurrent flares that involve the whole body. Recent studies have demonstrated that the skin microbiota, characterized by an overgrowth of Staphylococcus aureus (S. aureus), plays a critical role in the manifestation of AD. There is striking evidence that skin microbiota can modulate the development and progression of AD. Therefore, more and more therapeutic approaches are adopted for modifying skin microbiota. Here we discuss the role of skin microbiota in the etiology and maintenance of AD; furthermore, we summarize the effects of therapeutic treatments on skin microbiota in AD based on published literature. With the help of the theoretical guidance suggested by microbial metagenome analysis, the reconstitution of microbiota should be a promising way to harness the pathogens of AD and could be used as a brand-new therapeutic strategy in clinical trials. We believe that the targeted therapy of dysbiosis in AD may possibly become a unique approach to an integrated treatment program in the near future.

Keywords: Atopic dermatitis; Skin microbiome; Skin microbiota; Staphylococcus aureus; Therapy

Key Summary Points

Skin microbiota is a complex ecosystem composed of bacteria, fungi, and viruses. Integrity of the diverse microbes plays an essential role in maintaining homeostasis and preventing pathogens from invading skin. Recently, researchers found that increased colonization of Staphylococcus aureus (S. aureus) plays a critical role in the pathogenesis of atopic dermatitis (AD).

In this review, we summarize the clinical features of imbalanced skin microbiota associated with AD, and we emphasize the effects of each therapeutic treatment and their influences on skin microbiota. Besides traditional treatment approaches, such as emollient, antibacterial treatment, tacrolimus, narrowband UVB, coal tar, biological therapy, and contact with nature, we also discuss novel treatments that are targeting specific strains of human microbiota. Intriguingly, the targeted therapy of dysbiosis in AD has the potential to become an integrated treatment in the near future.

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INTRODUCTION

Skin microbiota is a complex ecosystem composed of bacteria, fungi, and viruses. Balance and integrity of the diverse microbes play an essential role in preventing pathogens from invading skin [1], while the skin microbiome refers to the composition of all microbial genes in the skin’s community [2]. Atopic dermatitis (AD), a worldwide heterogeneous, recurrent, chronic pruritic disease due to epithelial barrier dysfunction, immune dysregulation, and skin inflammation with dysbiosis of skin microbiota and colonization by the predominant pathogen *S. aureus*, shows an increasing tendency [3, 4]. Many studies have found alterations in the composition of the microbiome in patients with AD compared with that of healthy individuals, and microbiota differs between lesional skin and non-lesional skin [5, 6]. AD can also affect the skin of any part of the body, but it generally shows age-related morphological and distributional characteristics [7]. With the development of new technology, the application of whole genome sequencing (WGS) allows access to all genes of virtually all inflammation with dysbiosis of skin microbiota. Next-generation sequencing (NGS) technology in clinical bacteriology has been interrogating thousands of 16S ribosomal RNA gene amplicons from bacteria to archaea in floras from a patient with AD [8, 9]. Currently, exploration of the skin microbiome in AD is attracting more and more attention from researchers. Dysbiosis in microbiota has been universally considered an important factor in the pathogenesis of AD, and how to achieve and maintain a balance between skin microbiota and the host has become a hot research topic in the field of AD treatment [10]. Although traditional treatments including topical glucocorticoid and calcineurin inhibitors can inhibit inflammation, the recurrence of AD continues. Focusing on microbiota therapy is currently being developed to revise skin dysbiosis related to AD, with the expectation of obtaining long-term remission. In this review, we give a brief introduction on clinical features of AD, then discuss the pathogenesis of skin microbiota in AD, and finally discuss the treatment strategies based on the skin microbiome by reviewing literature published till October 2019. This article is based on previously conducted studies and does not contain any studies with human participants or animals performed by any of the authors.

CLINICAL FEATURES OF AD AND SKIN MICROBIOME

The consistent increment in the prevalence of AD in both developing and developed countries has depicted a global trend in the burden of AD, and the picture in the developing world may soon resemble that in developed countries [11], where 15–20% of children have been affected [12, 13]. Nearly 70% of children with AD show disappearance of symptoms or spontaneous remission during puberty even in individuals with genetic heterogeneity of filaggrin mutations [14]. However, about 2–5% of adults worldwide can be affected and present as adult-onset AD or infantile/childhood AD, and the dermatitis symptoms may persist or recur over many years [15, 16].

Newborns generally carry an adequately uniform microbiome. The microbial community composition of newborns is initially determined by the manner of delivery, i.e., newborns delivered naturally obtain bacteria in their mother’s vagina that are dominated by *Lactobacillus*, while those born by Caesarean section should inherit microbiota resembling those on their mothers’ skin [17]. Although the skin lesions of infants are rather diffuse, it should be noted that most of the lesions occur on their faces and extensor sides of extremities [18]. Skin lesions on adults vary from a small localized plaque to a widespread lichenification, and even erythroderma. Shi et al. in 2016 identified that the skin microbiome of AD was significantly different between young children and adults–teenagers [14]. The skin microbiome during AD flares was investigated at species level by Byrd et al. [9]; dynamics of the microbial community from the pediatric AD group was consecutively monitored throughout the course of the disease. Eventually, they found a great dominance of *S. aureus* in those with more...
severe AD; however, there was an obvious dominance of *Staphylococcus epidermidis* (*S. epidermidis*) in patients with mild symptoms.

**PATHOGENESIS OF AD REVEALED BY SKIN MICROBIOME**

*Skin Microbiome Profiling in Patients with AD*

The pathophysiology of AD is associated with genetic, immunological, environmental, and microbiota risk factors. Aberrations of the immune system have long been considered as having key roles, until recently; molecular epidemiology has tended to focus on the skin microbiome as a common precursor and pathological feature. At present, the pathology of AD includes three critical factors: impaired skin barrier function, microbial dysbiosis, and cutaneous immune abnormality predisposed to T helper 2 (Th2) immunity, which may aggravate one another [19, 20].

The microbiota dysbiosis in AD is generally thought to be strongly related to the colonization of *S. aureus* on the skin, and the colonized skin lesions are closely correlated with the relative level of *S. aureus* [21, 22]. *S. aureus* was abundant on the skin of patients with AD compared with normal controls [6, 14]. The affected skin sites, especially inflamed areas, were predominantly colonized by more *S. aureus* compared to unaffected sites in patients with AD [9, 14]. The abundance of *Staphylococcus* and *Corynebacterium* spp. increased dramatically in untreated patients during a flare, while bacterial diversity was decreased including the reduction of *Propionibacterium, Streptococcus, Acinetobacter*, and *Corynebacterium* spp. [4, 21]. Greater bacterial diversity with increased abundance of *Staphylococcus, Streptococcus, Corynebacterium*, and *Propionibacterium* spp. was observed after AD treatment with clinical remission, suggesting a positive correlation with disease severity [23]. It is reported in a meta-analysis of 95 observational studies that the pooled *S. aureus* colonized 70% of lesional skin, 39% of nonlesional skin, 62% of nasal colonized patients with AD, and 3% of nonatopic patients [6].

Kobayashi et al. in 2015 [24] demonstrated that natural occurrence of *S. aureus* and colonization of *Corynebacterium bovis* caused inflammation of eczematous dermatitis in a mice model that resembled that observed in patients with AD. Thus, the dysbiosis is regarded as a pathological factor that can drive eczematous inflammation in mice.

**S. aureus**

*Staphylococcus aureus* is a well-known gram-positive spherical bacterium that occurs in grape-like clusters when viewed under a microscope. It is a facultative aerobic coccus with positive coagulase reaction. This bacterium is a clinically unapparent “carrier” and could lead to infection and thus it has been depicted as a pathobiont.

The severity of AD is usually associated with biofilms formed by *S. aureus* [25]. A biofilm is a bacterial assemblage attached to the surface and enclosed in an extracellular matrix. Research has shown that biofilms induce immune evasion, leading to the recurrence of AD and hard-to-treat infections [26].

All *S. aureus* strains express superantigens including staphylococcal enterotoxin (SE) and toxic shock syndrome toxin-1 (TSST-1). The colonization of *S. aureus* could exacerbate AD and cause vicious cycles via inducing thymic stromal lymphopoietin (TSLP) and Th2/Th17-type inflammation [9].

Recently, it has been proven that *S. aureus* plays a crucial role in occurrence and development of Th2 skewing and skin inflammation by multiple ways including the induction of T cell-independent B cell expansion, the release of cytokine and proinflammatory lipoproteins, and furthermore the stimulation of mast cell degranulation [23, 27], leading to the release of characteristic interleukins (ILs), such as IL-31, a notable pruritus-related cytokine [27, 28] (Fig. 1). When *S. aureus* from patients with AD was transplanted to a mice model, the epidermides of mice were thickened and cutaneous Th2 and Th17 cells emerged and expanded [23].
The allergens, especially superantigens, can initiate immunoglobulin E (IgE) responses to histamine derived from the mast cells and basophils. The production of \textit{S. aureus} superantigen definitely facilitates the maturation of Th2 cells, and eventually leads to inflammatory response. Moreover, the colonization of \textit{S. aureus} on skin can produce multiple virulence factors such as toxins and enzymes contributing to the inflammation and skin barrier dysfunction, and consequently cause bacteremia and sepsis through the invasion of human skin infection by \textit{S. aureus}.

\textit{S. epidermidis}

\textit{Staphylococcus epidermidis} is considered a skin commensal, but its role in AD is unclear. Besides \textit{S. aureus}, it is increased from the reports on colonization of \textit{S. epidermidis} on lesional skin of pediatric AD. Interestingly, \textit{S. epidermidis} skin colonization in AD can produce antimicrobial peptides (AMP) that could selectively target \textit{S. aureus} [29]. It implies a compensatory mechanism of \textit{S. epidermidis} in controlling pathogens.

**TREATMENT TARGETING SKIN MICROBIOTA**

To extend a patient’s symptom-free period for as long as possible, multifactorial approaches are essential for the treatment of AD disease because of its multifactorial nature. The rationale of targeting the microbiota in AD derives from considerable benefits that related to the microflora diversity [28]. The abundance of \textit{S. aureus} is tightly associated with disease flares.

![Fig. 1 Pathogenic role of \textit{Staphylococcus aureus} involved in atopic dermatitis. \textit{S. aureus} expresses superantigens, such as SE and TSST-1, which activate basophils and mast cell and cause histamine release; \textit{S. aureus} activates LC releasing IL-4, IL-5, and IL-13, which induces AMP reduction and Th2 differentiation and inflammation; \textit{S. aureus} stimulates KC releasing proinflammatory cytokines including TSLP, IL-8, IL-18, and IL-1β and biofilm can induce KC apoptosis releasing inflammatory cytokines; \textit{S. aureus} can release δ-toxin inducing mast cell degranulation. SE staphylococcal enterotoxin, TSST-1 toxic shock syndrome toxin-1, LC Langerhans cell, TSLP thymic stromal lymphopoietin, IL interleukin, KC keratinocyte](image)
and therapeutic responses [30]. Microbial diversity during AD flares is strongly correlated with recent AD treatments. It is likely that treatment induces changes in skin bacterial diversity. It indicates that AD treatments diversify skin bacteria preceding clinical improvements in disease activity [21]. Here we describe all kinds of strategies used for treatment targeting skin microbiota in patients with AD.

**Emollient Use**

Considering that barrier dysfunction with epidermal water loss has an important relationship with AD, topical treatment using emollient is usually encouraged as a basic therapy [3, 31, 32]. Glatz et al. in 2018 conducted an experiment in 6-month-old infants with family history of AD [33]. The emollient group demonstrated higher bacterial diversity, decreased skin pH, and increased proportion of *S. salivarius*. It was a remarkable benefit of applying emollients in high-risk infants. Further studies are needed to confirm the preliminary results of emollients as a prophylactic therapy for AD [33]. Seite et al. in 2014 reported that when treated with emollients, the clinical symptoms of AD were alleviated in 72% of patients with AD, as indicated by an increased overall diversity and a decrease in the abundance of *Staphylococcus* species. Importantly, the microbial communities on the affected skin in the treatment group more closely resembled those on unaffected skin [5].

**Antibacterial Treatment**

It is essential to apply antibacterial agents such as antiseptics and antibiotics in clinically relevant bacterial infection in AD, and in patients with signs of bacterial superinfections, intermittent bleach baths and intranasal application of mupirocin ointment decreased the severity of AD [34]. However, it should be noted that the use of antibiotics in the treatment of systematic decontaminations of *S. aureus* is controversial as large-scale use of antibiotics would result in bacterial resistance and microbiota disorder [35]. Although bleach baths are widely accepted as an effective therapy for AD in some areas [36], they still remain highly controversial worldwide. Moreover, bleach baths were not more effective than water baths alone in reducing the severity of AD [37]. Moreover, sometimes the effects of bleach bath were contradictory in human studies. Gonzalez et al. in 2016 found that a 4-week duration period and/or twice-weekly baths may not be sufficient for additional impact on the cutaneous microbiome [38]. Another study evaluated the bleach bath by reviewing the effect of either topical corticosteroid alone or in combination with dilute bleach baths. The results showed improved clinical eczema representation and suppressed *Staphylococcus* species on lesional and non-lesional sites in both groups, suggesting an invalid effect of the additional dilute bleach baths [23]. Some researchers even expressed a word of caution based on the potential dangers in the application of such a strongly alkaline treatment [39].

**Tacrolimus Ointment for Treatment of AD**

Tacrolimus ointment, which is a very important anti-inflammatory topical drug, is recommended as the first-line treatment for AD with topical corticosteroids [40]. Wongpiyabovorn et al. [41] conducted an experiment with nine Thai patients with AD undergoing a 4-week monotherapy with tacrolimus. A lower prevalence of the phyla Actinobacteria and Proteobacteria and a higher prevalence of the phylum Firmicutes were identified in patients with AD. With the notable improvement of all clinical scores, there was a significant increase in some commensal genera including *Dermacoccus*, *Pseudomonas*, *Corynebacterium*, *Proteus*, *Micrococcus luteus*, and *Lactococcus* in the patients with AD after treatments with tacrolimus ointment, which should reflect the effect of tacrolimus on the skin microbiome. This study provides unequivocal proof that tacrolimus has a positive impact on the skin microbiome in patients with AD.
Narrowband Ultraviolet B (UVB) Phototherapy

UVB phototherapy is well established as a second-line treatment for adolescent and adult patients with AD. Kwon et al. carried out research on 18 patients with AD by using UVB phototherapy. A drastic increase in microbial diversity and decrease of *S. aureus* proportion were observed with clinical improvement of AD. The UVB treatment did not exert additive effects on the improvement of AD; however, it seemed to reduce AD recurrence [42].

Treatment with Coal Tar

Coal tar (CT), which is an ancient yet highly effective and safe therapeutic agent, was applied to seven patients with AD and ten healthy volunteers with vehicle treatment as control [22]. During CT treatment, the abundance of *Staphylococcus* decreased, whereas *Propionibacterium* abundance increased. The therapeutic effect of CT treatment was achieved by targeting the cutaneous bacterial composition through keratinocyte-derived AMP production via activation of the aryl hydrocarbon receptor (AHR).

Biological Therapy in Patients with AD

Dupilumab, a monoclonal antibody against interleukin-4 receptor alpha, has been used in numerous allergic diseases and works by blocking signaling of interleukin-4 and interleukin-13, type 2/Th2 cytokines. It has been confirmed to be effective in patients with moderate to severe AD in several double-blind, placebo-controlled studies and in clinical practice [43–45]. The result showed that clinical improvement is positively correlated with increased favorable microbial community and reduced abundance of *S. aureus* [46]. More novel systemic immunomodulatory agents for AD are under development, and it is promising that these therapies will not only improve skin immunity but also adjust *S. aureus* colonization by modulating their cell wall proteins [19].

Contact with Nature

A new theory is that regular contact with nature could enhance the diversity of microbiota [47]. For example, dog ownership confers a protective effect against AD development because dog owners spend more time outdoors and therefore have more exposure to natural environments, resulting in the alteration of their skin microbiota. Moreover, pet dogs also benefit owners’ health by reducing psychological stress and promoting associated immunomodulatory effects [48, 49].

Novel Treatments

Interestingly enough, novel treatment strategies seek to target specific aspects of dysbiosis in AD [29]. The commensal strains of coagulase-negative staphylococci (CoNS), collected from the skin of healthy individuals, protected against skin dysbiosis by reducing the colonization of *S. aureus* in patients with AD [29]. A prospective, double-blind, placebo-controlled clinical study in 75 volunteers with AD showed that cream containing 5% of the lysate of a nonpathogenic proteobacterium *Vitreoscilla filiformis* could significantly improve a clinical score and reduce transepidermal water loss (TEWL), itch, and insomnia compared with placebo [50]. A further study showed that topical transplantation with *Roseomonas mucosa* was effective in decreasing disease severity, topical steroid requirement, and *S. aureus* burden in patients with AD. There were no adverse effects or treatment complications in this study [51]. Overall, more research is to enable us to implement such skin microbiota as a therapeutic target in the pathogenesis and treatment of AD [30].

CONCLUSIONS

New insights into the pathophysiology of AD focus on the important role of microbial dysbiosis, which was a driving factor in the pathogenesis of AD according to animal experiments [4]. However, it is still unclear if causal relationships between the skin microbiota and
disease exist in patients with AD. *S. aureus* plays a critical role not only during AD flare but also in the treatment of skin dysbiosis. Treatment targeting the microbiota in patients with AD can effectively reduce the colonization of *S. aureus* and significantly increase the diversity of bacteria on the skin surface, suggesting an important role of microbial dysbiosis in the pathogenesis of AD. Thus, the establishment of the balance of skin microbiota in patients with AD is feasible for the treatment and prevention of AD disease. Pan-genome and comparative genome analyses of the specific bacterial species should reveal their genomic abundance in the healthy and diseased human skin microbiota, and all the skin microbiome of AD disease must be investigated in the context of the healthy skin microbiota. Probing the molecular mechanism of interaction between *S. aureus* from AD skin and the commensal strains will be a major focus for the future.

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**Data Availability.** The datasets generated during and analyzed during the current study are available from the corresponding author on reasonable request.

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