Psoriasis: Embarking a dynamic shift in the skin microbiota

Mrinal Gupta MD1 | Jeffrey M. Weinberg MD2 | Paul S. Yamauchi MD3,4 | Anant Patil MD5 | Stephan Grabbe MD6 | Mohamad Goldust MD7

1DNB Dermatology Consultant
Dermatologist, Treatwell Skin Centre,
Jammu, India
2Icahn School of Medicine at Mount Sinai,
New York, NY, USA
3Dermatology Institute and Skin Care
Center, Santa Monica, CA, USA
4Division of Dermatology, David Geffen
School of Medicine, University of
California, Los Angeles, CA, USA
5Department of Pharmacology, Dr. DY
Patil Medical College, Navi Mumbai, India
6Department of Dermatology, University
Medical Center of the Johannes
Gutenberg University, Mainz, Germany
7Department of Dermatology, University
Medical Center Mainz, Mainz, Germany

Correspondence
Mohamad Goldust, Department of
Dermatology, University Medical Center
Mainz, Mainz, Germany.
Email: mgoldust@uni-mainz.de

Abstract
Recent interest has arisen regarding the role of microbiome and its composition in the pathogenesis of psoriasis. Numerous studies have shown that there are alterations in skin flora arrangement between normal individuals and psoriatic patients. Psoriasis exacerbation could be interconnected with epidermal or mucosal colonization with streptococci, Malassezia, Staphylococcus aureus, or Candida albicans. The role of cutaneous and gut microbiome in psoriasis pathogenesis has recently been studied in both human and animal models. In this review, we try to evaluate various pathogenic mechanisms linking the microbiota and psoriasis. The literature research included peer-reviewed articles which included clinical trials, original reports, and scientific reviews. MEDLINE and PubMed databases were searched from January 1990 to March 2021, including the reference lists of articles meeting our criteria.

KEYWORDS
microbiome, microbiota, psoriasis

1 | INTRODUCTION

The overall prevalence of psoriasis has been significantly associated with several metabolic syndromes and systemic inflammatory disorders.1-3 In addition, stress, infections, diet, pain killers, and antibiotics are some factors that tend to elicit psoriatic lesions.4,5 Psoriasis has been shown to have a bimodal prevalence with a major onset between the age of 20 and 30 years and then again between 50 and 60 years. Recently, the etiopathogenesis of the disease has been linked to composition of microbiota of the host.6 Numerous researchers have observed variations in skin microflora between normal controls and psoriatic patients, suggesting a disease-associated alteration in the skin microbiome.7,8

The role of microflora in etiopathogenesis of psoriasis is also rendered plausible by the finding that infections caused by various microorganisms often lead to the initiation or flare up of lesions.9,10 Psoriasis has been found to be induced or exacerbated by certain pathogens, which include bacteria such as Staphylococcus aureus and Streptococcus pyogenes, viruses such as human papillomavirus and endogenous retroviruses, and fungi such as Malassezia and Candida albicans.11-14 Alekseyenko et al. observed that Corynebacterium, Staphylococcus, Propionibacterium, and Streptococcus are significantly increased in psoriatic lesions.15 Fahlén et al. observed that streptococci were significantly increased in psoriatic lesions, but Staphylococcus and Propionibacterium were significantly lower in psoriatic lesions in control skin.16 Gao et al. also found that Propionibacterium species were lower in psoriatic than in normal skin.17 Researchers have observed a decrease in Firmicutes and an increase in Proteobacteria in psoriatic skin.18 Researchers have found that Firmicutes are significantly increased in psoriatic lesions as compared with uninvolved skin in psoriatic patients and healthy skin controls. Actinobacteria and Propionibacterium have also been found to be reduced in the psoriatic lesion samples. Changes in intestinal microbiome may activate a systemic pro-inflammatory status, which
may contribute to disease pathogenesis.13 The role of gut-skin affiliation in etiopathogenesis of psoriasis has been acknowledged in both human and animal models of psoriasis.14–17 Recovery from intestinal dysbiosis may reduce the cutaneous symptoms of psoriasis in patients.18 Aggravation of plaque psoriasis has been linked to bacterial spread into the blood stream resulting from the enhanced intestinal permeability seen in the patients with psoriasis.19 Researchers have also observed that certain alterations in intestinal microbial composition, which are found in patients with inflammatory bowel disease and obesity, including decreased profusion of Akkermansia muciniphila, are also seen in psoriatic patients.20

A number of studies performed with mice and human hosts provided conclusive substantiation of the effect of intestinal bacteria on skin condition.21 It was observed that mice that were fed with Lactobacillus reuteri, a probiotic bacteria, had a thicker skin and regained better reproductive fitness.22,23

2 | DIVERSIFIED SKIN MICROBIOME

It has been observed that the changes in psoriatic skin microbiota are highly site-specific and have been shown to have greater heterogeneity as compared to the healthy skin. Changes in human microbiota content and diversity have been associated with various diseases, for example, decreased bacterial alpha diversity in gut microbiota is associated with obesity and inflammatory bowel disease,24,25 and alteration in vaginal microbiome is seen in bacterial vaginosis.26,27 The skin microbiome of psoriatic skin, non-lesional areas, and healthy skin consists of four prominent phyla, namely, Actinobacteria (53.8%–66.5%), Firmicutes (23.9%–28.3%), Proteobacteria (5.8%–12.0%), and Bacteroidetes (2.1%–2.9%).28 Skin microbiome is mainly constituted of Propionibacterium (22.8%–38.1%), Corynebacterium (21.4%–23.9%), and Staphylococcus (5.3%–9.2%) in various cutaneous diseases. Even though the predominant species are similar in different conditions, a steady change in microbiota composition is seen in healthy skin and non-lesional skin psoriatic microflora and lesional microflora, indicating that these microbiota alterations may precede the disease onset in predisposed patients and may contribute to disease pathogenesis. Actinobacteria and Proteobacteria have been identified as differentiating markers of the skin microbiota in psoriatic lesions and healthy skin. Conchiformibius, Lactococcus, Moraxella, and Acetobacter have been found to be associated with unaffected psoriatic skin, and these species may act as possible markers for differentiating normal skin from various disease conditions.26 16S rRNA sequencing and LEfSe analysis have helped in identifying that the healthy skin microbiome has predominance of Propionibacterium acnes and Propionibacterium granulosum, whereas Staphylococcus sciu is prominent in non-lesional psoriatic skin. It has also been observed that two Staphylococcus species—S. aureus and Staphylococcus pettenkoferi—are more common in lesions of psoriasis, whereas Staphylococcus is not significantly seen in any skin condition.27

3 | INTERRELATION AMONG DISTINCT BACTERIAL SPECIES

Like the case with any ecosystem, it has been seen that the symphony of skin microbiota is controlled by various environmental factors (eg, nutrient accessibility and immune response of the host) and interactions among various bacterial species. These interactions could be an important factor that affects the overall composition of the community, and thus, it is extremely important to understand these interactions which can provide important information regarding the composition and preservation of psoriasis-associated microbiome. In the psoriatic skin, three clusters of bacterial communities were identified at the genus level. Cluster A consisting of Corynebacterium, Prevotella, Porphyromonas, Finegoldia, Peptoniphilus, and Anaerococcus is the largest cluster. Cluster B consists of Paracoccus, Kocuria, Micrococcus, and Janibacter, and Cluster C consists mainly of Rothia and Streptococcus.28

Citing the earlier findings of impending significance of Streptococcus in the pathogenesis of psoriasis, the role of Rothia spp. in the pathogenesis of psoriasis also needs to be studied as it is extensively associated with Streptococcus. P. acnes, which is common in normal skin, negatively correlates with S. sciu and S. pettenkoferi, which are more common in the psoriatic skin microbiome. In accordance with this finding, it was also observed that P. acnes and Staphylococcus epidermidis were highly associated with S. aureus-induced mild psoriatic skin tissue and S. pettenkoferi was highly associated with S. aureus-induced severe psoriatic skin tissue, indicating that the antagonistic interactions between these bacteria may take part in the pathogenesis of psoriasis. Pseudocloclavibacter bifida has been found to be negatively associated with P. acnes and positively associated with S. sciu. A higher concentration of Pseudocloclavibacter bifida was seen in S. aureus-induced severe psoriatic skin samples. P. acnes and P. granulosum act as two major Propionibacterium species, and their levels positively correlate with each other. The correlation between P. acnes and P. granulosum and their interactions in normal skin indicate their role in maintaining cutaneous health.29

Psoriasis is characterized by indurated and inflamed plaques; hence, it is very easy to distinguish the psoriatic skin lesions from non-psoriatic and normal skin as each of them represents very distinctive microbial habitats that possibly will affect interactions between different microbes. In addition, it is also observed that a particular group of dissimilar species are present at different stages of the disease progression, confirming the assumption that diverse interactions occur in different disease conditions. Amazingly, species interactions observed in the microbiome linked with psoriatic skin lesions were more analogous to those observed in healthy skin than the non-lesional psoriatic skin. In normal skin, P. acnes correlates negatively with different bacterial colonization, signifying its inhibitory role in microbial growth. Lesser microbes were seen with P. acnes in non-lesional psoriatic skin, whereas only Pseudocloclavibacter bifida was found to be anticorrelated with P. acnes in lesional psoriatic skin. On the whole, P. acnes might possibly take part in altering the skin
microbiome by maintaining growth of various microorganisms under control, and the alteration of this balance in psoriatic skin can contribute to disease progression.\(^3\) Loesche et al observed that the rare bacterial species (<1%) comprised a larger proportion of psoriasis-associated microbiome than those present in non-lesional skin, which include species like Lactococcus, Paracoccus, Porphyromonas, Prevotella, Acinetobacter, Neisseria, and Fusobacterium.\(^3\)

4 | MALADAPTATION OF THE SKIN MICROBIOME

The sophisticated association between microbiome and the host begins at the time of birth as the pelvic floor microflora are transmitted to the newborn during childbirth and lactation through colostrum, which is rich in microbes. The juvenile newborn immune system, which lacks full immune response, has been found to tolerate a symbiotic relationship association between microbiome. It is demonstrated by the response of newborn innate immune cells to microbial exposure, which showed a mitigated inflammatory response in comparison to the adult immune cells.\(^3\) Microbe-associated molecular patterns (MAMPs) are the key factors responsible for interaction between the host and pathogen. An exception to the fundamental biological function of the immune system to recognize foreign molecules and subsequently initiate a directed response is that no immune response arises to host cutaneous and mucosal MAMPs, despite the constant interface of MAMPs with certain pattern recognition receptors (PRRs) that identify microbes and start an immunological response.\(^3\) Through these mechanisms, the commensal microbiota influence the hosts' postpartum immune responses.\(^3\) Other physiological obstacles, such as the epithelial cell layer which secretes IgA and goblet cells which produce mucus, not only separate the microbiota from having a direct interaction with mucosa but also allow it to impart localized and systemic effects.\(^3\) Dysbiosis has arisen as a fundamental study focus in the pathogenesis of microbiome-related inflammatory disorders such as inflammatory bowel disease.\(^3\) Dysbiosis, defined as a disparity between the microbiome and the host, can be regarded as a form of compromised homeostasis where the microbiome is altered toward a simpler and different pathological state.\(^3\) The mechanism that brings this change from symbiosis to dysbiosis is not well-explained, but a genetic predisposition has been hypothesized.\(^3\) Dysbiosis of microbiota in psoriatic patients enhances the translocation of bacteria from the gut and skin into systemic circulation, which, in turn, may predispose to chronic systemic inflammation in them.

5 | PORTRAYAL OF BACTERIA IN PSORIASIS

Based upon comprehensive understanding of T cells in the etiology of psoriasis, it is extremely important to establish a link between streptococcal infection and the unstable or guttate variants of psoriasis.\(^3\) M proteins, extracted from Group A, C, and G β-hemolytic streptococci, have been avowed as decisive factors as deterioration of psoriasis was tightly linked with the M-protein-yielding streptococci.\(^3\) It has been hypothesized that M proteins may simulate keratin factors with succeeding psoriatic T-cell activation.\(^3\) This hypothesis is authenticated by the finding that interactions between type IV collagen and α1β1 integrin, which is found entirely on the epidermal psoriatic T cells, leads to the growth of these cells, leading to appearance of psoriasis.\(^3\) T-cell triggering in guttate variants of psoriasis is altered by the effect of antigens such as streptococcal pyogenic toxins A and B and peptidoglycans.\(^3\) It can be postulated that the skin microbiome plays a part in etiopathogenesis of psoriasis. Bacteria such as Propionibacterium, Staphylococcus, Corynebacterium, and Streptococcus have been recognized as the main bacterial genera which may take part in psoriasis pathogenesis. Researchers have identified Firmicutes as the most common bacterium found in psoriatic lesional skin, and the Actinobacteria count was significantly less in lesional psoriatic skin than non-lesional and healthy skin.\(^3\)

Researchers have observed that S. aureus proteins increase Th17 differentiation in vitro, suggesting that S. aureus colonization can cause elevated Th17 activation and IL-17 secretion.\(^3\) Researchers have reported higher Th17 transcriptomic signals and IL-17A and IL-17F cytokine transcripts in Teff cells collected from the skin S. aureus-colonized mice.\(^3\) IL-17A is identified as one of the major agents in pathogenesis of psoriasis. Researchers also observed that other mediators of Th17 responses, such as IL23R and IL22, were also elevated upon S. aureus exposure in the keratinocytes, which may lead to IL-17-induced inflammation in psoriasis.\(^3\)

6 | ROLE OF FUNGI IN PSORIASIS

Several studies have established a possible etiopathogenic correlation between psoriasis and fungi. In 1980s, development of psoriatic plaques was observed in all 10 tested subjects as a result of application of Malassezia ovales suspension to their unaffected skin.\(^4\) Similar plaques were observed after secondary fungal deposition from the scalp of psoriatic patients. In a case series, a week-long course of oral antifungal therapy was reported to improve psoriatic lesions.\(^4\) Increased Malassezia yeast colonization in lesional skin has been found to cause psoriasis exacerbations.\(^4\)

Also, the anecdotal observation that topical miconazole regularly improves inverse psoriasis could be compatible with a role of fungi in psoriasis (authors’ own observation). The role of Malassezia is largely attributed to its action to upregulate tumor growth factor-β1, HSP70, and integrin chain expressions, leading to increased migration of immune cells and keratinocyte hyperproliferation in psoriatic patients.\(^4\) Malassezia furfur induces higher IgG and lower IgM production in psoriatic patients than in normal subjects.\(^4\) As seen with bacterial microbiome, nonculture techniques have also succeeded over culture techniques in detecting Malassezia in psoriatic and normal skin. M. restricta has been identified as the most common
Malassezia species in psoriatic skin. In spite of the perpetual findings, the mere identification of fungus itself is not an indicator to extricate psoriatic from non-psoriatic and healthy skin.52,53

7 | CONCLUSION

The interaction of the immune system and microbiota is of significance because it can lead to better understanding of psoriasis pathogenesis and, at the same time, give clearance for expansion of microbiome-targeted therapies. Although reports state a higher variation of the microbial population in psoriatic lesions and an intestinal dysbiosis in psoriatic patients, it is yet to be determined whether such changes in microbiota are the etiology or consequence of the disease. Further research is warranted to explore the precise role of microbiota in the management of psoriasis.

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ORCID

Anant Patil https://orcid.org/0000-0002-9455-4025
Mohamad Goldust https://orcid.org/0000-0002-8646-1179

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