The skin microbiome: potential for novel diagnostic and therapeutic approaches to cutaneous disease

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Abstract
A vast diversity of microorganisms, including bacteria, fungi, viruses, and arthropods, colonize the human skin. Culture-independent genomic approaches for identifying and characterizing microbial communities have provided glimpses into the topographical, temporal, and interpersonal complexity that defines the skin microbiome. Identification of changes associated with cutaneous disease, including acne, atopic dermatitis, rosacea, and psoriasis, are being established. In this review, our current knowledge of the skin microbiome in health and disease is discussed, with particular attention to potential opportunities to leverage the skin microbiome as a diagnostic, prognostic, and/or therapeutic tool.

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In and on our bodies, microbial cells outnumber human cells by a factor of 10. Microbial genes outnumber human genes by a factor of 100. It is increasingly apparent that this collective set of microorganisms and their genetic material, the “microbiome”, contributes genetic diversity, modulates disease, influences metabolic processes, and is essential for immunity. The human microbiome is also dynamic, and changes associated with health and disease have been described and mechanistically investigated. Being a tractable entity, the microbiome is also a prime target for manipulation to influence health and disease processes. In most cases, but especially on the skin, the microbiota is an easily accessible target for therapeutic intervention and/or diagnostic testing.

Our awareness and investigation into human-associated microbial communities was once limited by culture-based techniques. These methods that rely upon cultivation and isolation of microbial communities was once limited by culture-based techniques. Importantly, these methods do not rely upon cultivation of the microorganism, thus eliminating biases associated with culture-based techniques.

Multiple studies have employed 16S rRNA gene sequencing to characterize bacterial communities colonizing the skin, in health and disease. Immediately following birth, our skin is colonized with microbiota, and delivery mode may in part contribute to differences in colonization. Analysis of neonate skin microbiota indicates that those babies delivered via vaginal birth had skin microbiota similar to their mother’s vaginal microbiota, and babies delivered via Caesarean section had skin microbiota similar to their mother’s skin microbiota. It is unclear how long these colonization differences persist, as most vaginal bacteria are not typically found on the skin, and if the associated differences translate into functional consequences and/or disease risk. Other factors, including geography, lifestyle, and/or ethnicity may influence the skin microbiota. For example, Amerindians residing in the Venezuelan Amazon are colonized with distinctly different microbiota compared to subjects residing in the United States.

Generally, the 4 dominant phyla of bacteria residing on the skin are the Actinobacteria, Proteobacteria, Firmicutes, and Bacteroidetes. A feature of the skin is the topographical diversity of bacterial populations, with composition and diversity of bacteria colonizing the skin depending on the environment. Variability between individuals is high, similar to other sites of the human microbiome, as is temporal variability within the same individual. However, the dominant types of bacteria that reside on the skin appear to be relatively stable, with the rarer, less abundant types of bacteria accounting for the variability. These dominant types of bacteria, primarily Staphylococcus, Propionibacterium, and important insight into bacterial, fungal, and viral populations on the skin.

The advent of next-generation sequencing technologies has revolutionized our view of human-associated microbial communities. Using DNA sequencing methodology, we are now able to characterize and analyze microbiomes with greater precision and accuracy, and less bias compared to culture-based approaches. A common approach used to identify bacterial populations is based on sequencing of the small subunit bacterial 16S ribosomal RNA (rRNA) gene. Hypervariable regions within this gene contain species-specific sequences, that when compared to reference databases, allow identification of the bacteria of origin. A similar approach for fungal identification, based on the sequence lying between the fungal 18S and 5.8S rRNA genes, the internal transcribed spacer region (ITS), has been used to sequence and survey fungal populations of the skin. Increasing throughput and decreasing costs associated with DNA sequencing, along with the development of computational tools for the analysis of resultant sequence data, have made these approaches feasible to query microbial communities in health and disease states. Importantly, these methods

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More research is needed to identify probiotic strains and prebiotics are substrates that encourage the growth of beneficial bacteria. Increasing popularity for use in the gastrointestinal tract. Probiotics are live microorganisms with known health benefits, and are gaining strategies targeting the microbiota are on the horizon. Probiotics are a component of health and also many disease processes, strat- egies to perturbation, providing a sensitive readout of the environment in which they thrive. While these implementations of personalized and precision medicine are on the horizon, more research is needed to define the skin microbiota and its involvement in cutaneous disorders. Here we describe the healthy, skin microbiome as it is understood from recent genomic, culture-independent surveys (Figure), how it is perturbed in disease states (Table), and how microbiome research may one day inform precision and personalized medical approaches for the management and treatment of skin disease.

Sebaceous skin: acne and rosacea
Sebaceous areas, such as the back, face, behind the ears, tend to be colonized by high proportions of the lipophilic bacteria Propionibacterium (Figure). This finding is consistent with evidence that Propionibacterium hydrolyzes triglycerides found in sebum, thus releasing free fatty acids on the skin that then function to acidify and emolliate the skin. Further, sebaceous areas tend to be low in diversity and relatively stable temporally, compared to moist and dry sites of the skin, perhaps a reflection of the unique microen- vironment provided by sebaceous secretions that select for certain types of microbes.

The role of sebum in defining the skin microbiota may be reflected by age-associated changes in the composition and diversity of the skin microbiota. At birth, babies are colonized with higher relative abundance of Firmicutes than Actinobacteria. Generally, puberty-associated maturation of the sebaceous glands coincides with a shift in skin microbiota towards enrichment with Actinobacteria, including Corynebacterium and Propionibacterium. This age-related shift in skin microbiota may be a potential factor contributing to the decrease in incidence and severity of childhood skin diseases, for example atopic dermatitis.

The association between Propionibacterium acnes and acne vulgaris has been well-established. The effectiveness of antibiotics in acne treatment further supports a microbial role in disease pathogenesis. Based on 16S rRNA gene sequencing studies, it is now apparent that P. acnes is a member of the commensal micro- biota even in those without acne. These findings raise the question of when and how does a commensal become a pathogen? Insight into this question may require strain-level analysis; a recent study demonstrated that even though P. acnes relative abundance did not differ between individuals with acne compared to healthy indi- viduals, certain strains were highly associated with acne (Table). These strains carry unique genetic elements, not present in the strains associated with healthy individuals, which may contribute to virulence and pathogenicity.

Probiotic applications, where the skin microbiota is supple- mented with a beneficial microorganism, may offer promise for treatment of conditions such as acne. For example, succinic acid, a fatty acid fermentation product of Staphylococcus epidermidis, can inhibit the growth of P. acnes. Similarly, P. acnes can func-
tion as a probiotic that suppresses growth of *S. aureus*, in particular USA300 community acquired methicillin resistant *S. aureus* (MRSA), through fermentation of glycerol. Replacing antibiotics with therapies that do not place selective pressures on microorganisms is desirable, as antibiotics encourage evolution of antibacterial resistance mechanisms. Long-term antibiotic treatment for acne has been associated with acquisition of antibiotic resistance. Gram-negative folliculitis and pharyngitis are also associated with *rosacea*, which in part may contribute to dysbiotic cutaneous states. Because *Demodex* are present on healthy skin, and considered as part of the commensal microbiota, it is likely that host status contributes to their transition from commensal to pathogen. However, additional research is needed to provide insight into the mechanism in which microorganisms, including *Demodex*, influence the pathogenesis of rosacea.

### Moast skin: body odor and atopic dermatitis

Skin areas that are moist and/or occluded tend to be colonized by bacteria such as *Staphylococcus* and *Corynebacterium* (Figure), organisms that prefer high humidity in their habitat. Culture-dependent analyses of the axilla microbiota has revealed microorganisms, namely the *Corynebacterium*, that process odorless eccrine, apocrine, and sebaceous gland secretions, thus producing volatile organic fatty acids and thioalcohols that are responsible for malodor associated with sweating of the axilla. Sequencing of 16S rRNA genes has confirmed that *Corynebacterium* and *Staphylococcus* are the predominant axilla bacteria, with higher proportions of *Corynebacterium* in males, and those using deodorants harboring greater bacterial diversity. Genetic variation in ABCC11, specifically a 538G→A single nucleotide polymorphism (SNP) prevalent in East Asian populations, results in reduced concentration of axillary apocrine secretions and decreased production of axillary odor. This genetic variant also impacts the composition of the axillary skin microbiota, where relative abundance of *Staphylococcus* is decreased and *Corynebacterium* increased in individuals carrying A/A genotype, as compared to those with G/A and G/G genotype. Deodorants containing probiotics and/or prebiotics may in the future be a feasible strategy to combat body odor, where odor-producing bacteria would be competed away with recolonization by nonodor producing bacteria.

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<table>
<thead>
<tr>
<th>Disease</th>
<th>Study</th>
<th>Sample size</th>
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<tbody>
<tr>
<td>Acne vulgaris</td>
<td>Fitz-Gibbon et al 2013</td>
<td>49 acne + 52 controls</td>
<td>Association of <em>P. acnes</em> strain with acne</td>
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<tr>
<td>Rosacea</td>
<td>Casas C et al 2012</td>
<td>50 rosacea + 48 controls</td>
<td>Increased density and frequency of <em>Demodex follicularum</em></td>
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<tr>
<td>Atopic Dermatitis</td>
<td>Kong et al 2012</td>
<td>12 AD + 11 controls</td>
<td>Increased <em>Staphylococcus</em> and decreased diversity during flares</td>
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<td>Psoriasis vulgaris</td>
<td>Alekseyenko et al 2013</td>
<td>51 psoriasis + 51 controls</td>
<td>Combined increase in <em>Corynebacterium</em>, <em>Propionibacterium</em>, <em>Streptococcus</em>, and <em>Staphylococcus</em></td>
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<td></td>
<td>Gao et al 2008</td>
<td>6 psoriasis</td>
<td>Increased Firmicutes and Actinobacteria</td>
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<td></td>
<td>Fahlen et al 2012</td>
<td>10 psoriasis + 12 controls</td>
<td>Decreased <em>Staphylococcus</em> and <em>Propionibacterium</em></td>
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an environmental component. Further, colonization with \textit{S. aureus} is associated with AD and antibiotics and bleach baths are effective treatment strategies that target the microbiota,\textsuperscript{44} Profiling of the bacterial microbiota via 16S rRNA gene sequencing indeed confirmed increases in \textit{S. aureus} relative abundance during AD flares, and decrease in bacterial diversity (Table).\textsuperscript{13} Whether \textit{S. aureus} colonization is a cause or effect of changes in the host skin remains unclear, but could be addressed by fine-scale longitudinal study designs incorporating skin barrier testing and surveys of immune status and/or the use of animal models. Predicting the best treatment for AD can also be difficult, and information about the microbiome may be able to inform these decisions. Similar to fecal microbiota transplantation therapies to treat \textit{Clostridium difficile} infections,\textsuperscript{45} one could envision the implementation of skin microbiota transplantations to ameliorate \textit{S. aureus} colonization and AD symptoms without relying on antibiotic therapies that encourage acquisition of resistance.

Some evidence exists to suggest that gastrointestinal probiotics and prebiotics may offer benefit in preventing and reducing the severity of AD.\textsuperscript{46} In addition, low diversity in the gut microbiota during early infancy has been associated with development of atopic eczema later in life.\textsuperscript{47,48} These observations would suggest that there is a connection between the gut microbiota and the skin, potentially through stimulation and/or education of immune cell populations. Further exploration of this connection may provide insight into the mechanism of allergic and atopic disease, while providing an additional target, the gut microbiota, for manipulation (via probiotics, prebiotics, or microbiota transplant) in the treatment and/or prevention of skin disease.

\textbf{Dry skin: psoriasis}

Generally, dry areas of the skin harbor greater amounts of bacterial diversity and lesser bacterial load than sebaceous and moist sites, and seem to have less of a selective force, as those bacterial populations are not dominated by individual taxa.\textsuperscript{9} Instead, drier areas contain greater amounts of Proteobacteria and Bacteroidetes, in comparison to moist and sebaceous sites (Figure).\textsuperscript{8,9,49,50} Psoriasis vulgaris (plaque psoriasis) has a predilection for drier skin sites, such as the elbows, knees, and trunk. It has been hypothesized that similar to Crohn’s disease, psoriasis results from failure in immune tolerance to microbiota.\textsuperscript{51} Guttate psoriasis, also forming lesions on the trunk and limbs, is associated with beta-hemolytic streptococcal infection, where streptococcal superantigens drive T-cell stimulation and expansion in the skin.\textsuperscript{52} The bacterial microbiota colonizing plaque psoriasis has been investigated using culture-independent techniques, and some changes have been identified, including decreased relative abundance of \textit{Propionibacterium} in plaques (Table).\textsuperscript{15,16} In a larger study of 51 psoriasis patients and matched healthy controls, the psoriatic plaque microbiota was determined to contain an increased combined relative abundance of \textit{Corynebacterium}, \textit{Propionibacterium}, \textit{Staphylococcus}, and \textit{Streptococcus}, and clustered distinctly from controls.\textsuperscript{14} However, it is unclear if these changes are a cause or consequence of alteration of the skin barrier and the ecosystem. Longitudinal studies, profiling the dynamics of microbial populations during plaque resolution and relapsing could provide insight into the role of microbiota during the triggering, propagation, and maintenance of plaques. Should the previously mentioned hypothesis be correct regarding the similarity of psoriasis to Crohn’s disease, microbiome-based therapies employed in conjunction with other therapies to prevent and/or treat the disease may be feasible. Further, complement activation is a known feature of psoriasis,\textsuperscript{53,54} and given previous findings connecting complement to skin microbiota composition and diversity,\textsuperscript{55} taken together would also suggest a microbial component to psoriasis. Complement therapeutics, which are under development and already used for some conditions resulting from complement activation, could potentially be employed to modify the skin microbiota and/or treat psoriasis.

\textbf{Skin microbiome interactions with host immunity}

Emerging evidence now suggests that host immunity influences the skin microbiota, and conversely, the skin microbiota in part modulates cutaneous immunity. Skin microbiota in patients with primary immunodeficiency syndromes (STAT-3 deficient hyper-IgE, Wiskott-Aldrich, and dedicator of cytokinesis 8 syndromes) is altered from a healthy state, colonized with different bacterial and fungal species, suggestive of increased ecological permisiveness.\textsuperscript{56} Primary immunodeficiency patients suffer from atopic dermatitis-like eczema and increased risk of skin infections. A separate study of STAT1 (chronic mucocutaneous candidiasis) and STAT3 primary immunodeficiency patients documented increased colonization with Gram-negative bacteria, in particular \textit{Acinetobacter}, which also suppressed the cytokine response of primary leukocytes to \textit{S. aureus} and \textit{Candida albicans}, 2 pathogens that commonly cause skin infections in these patient populations.\textsuperscript{57}

A study by our group recently demonstrated that antagonism of complement signaling (through the C5a receptor) in mice resulted in a shift in skin microbial composition and diversity.\textsuperscript{55} Notably, microbial diversity and richness decreased significantly when complement was blocked. Innate immunity, such as through complement signaling, may in part maintain microbial diversity at the skin barrier. Diversity can be advantageous in an ecosystem, and low diversity has been associated with some dysbiotic skin disorders (ie, atopic dermatitis) and gastrointestinal disorders. Additionally, the commensal microbiota was found to induce expression of complement genes, suggesting that the skin microbiota in part regulates cutaneous immunity.\textsuperscript{55} Another study found that a molecule released by staphylococcal bacteria, lipoteichoic acid (LTA), inhibits inflammation following injury through a Toll-like receptor (TLR2) dependent pathway,\textsuperscript{58} suggesting that the microbiota modulates cutaneous inflammation and immunity. Regarding adaptive immunity, the commensal skin microbiota was found to autonomously direct T-cell function and the local inflammatory milieu, in addition to conferring protective immunity to infection by \textit{Leishmania}.\textsuperscript{59} These studies together support an emerging interactive relationship between the skin microbiota and cutaneous immunity.

\textbf{Additional applications of microbiome science to precision cutaneous medicine}

Another application of culture-independent profiling of microbial communities may lie in prognostic and diagnostic approaches to disease. For example, identifying variants of disease, based on the colonizing microbiota, may better inform the optimal treatment approach. In a study of 52 neuropathic diabetic foot ulcers, patients clustered into 3 groups based on their colonizing micro-
biota. While this study did not provide outcome information or association of specific microbiota with poor outcomes, it is possible that these clusters represent variants that may benefit from different management and treatment approaches. In this particular case, 1 cluster was characterized by high relative abundance of anaerobic bacteria, which were not identified by standard of care cultures the majority of the time, likely because anaerobes are notoriously difficult to culture. This information alone is useful in guiding decisions of what antibiotic regimens to implement. In a separate study of acute traumatic wounds, characteristic microbiota was identified that correlated with wound location, mechanism (blunt versus penetrating), and future complication. In this situation, microbiota may be useful in identifying those that will go on to future infectious complication. In both of these types of wounds, predicting those who will experience infectious complications is difficult and biomarkers to guide management and treatment are greatly needed.

The role of microbiota in skin aging, such as wrinkling and sagging, remains unclear and is an area where treatments based on the skin microbiota may offer promise. For example, its foreseeability that certain metabolites produced by skin microbiota may offer benefit by modulating cutaneous pro- and anti-inflammatory responses, similar to what has been shown in the gastrointestinal tract. Overall, more research is necessary to identify those microorganisms that may offer benefit, and their mechanism of action.

Conclusion

Advances in sequencing technology have enhanced our ability to identify and characterize microbial communities colonizing the skin. Our knowledge regarding the healthy cutaneous microbiota, and how it is altered in dermatological disease is steadily increasing. However, in order to apply this knowledge to diagnostic, prognostic, and therapeutic applications, further research is necessary to understand beneficial and harmful microorganisms and their mechanism. These applications are particularly promising in light of emerging antibiotic resistance across medically relevant bacterial strains. Further, microbiome therapeutics and diagnostics are highly applicable to precision and personalized medicine, and may in the future transform management and treatment of dermatological disease.

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References


