Microbiome Modulation as a Therapeutic Approach in Chronic Skin Diseases

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Abstract: There is a growing quantity of evidence on how skin and gut microbiome composition impacts the course of various dermatological diseases. The strategies involving the modulation of bacterial composition are increasingly in the focus of research attention. The aim of the present review was to analyze the literature available in PubMed (MEDLINE) and EMBASE databases on the topic of microbiome modulation in skin diseases. The effects and possible mechanisms of action of probiotics, prebiotics and synbiotics in dermatological conditions including atopic dermatitis (AD), psoriasis, chronic ulcers, seborrheic dermatitis, burns and acne were analyzed. Due to the very limited number of studies available regarding the topic of microbiome modulation in all skin diseases except for AD, the authors decided to also include case reports and original studies concerning oral administration and topical application of the pro-, pre- and synbiotics in the final analysis. The evaluated studies mostly reported significant health benefits to the patients or show promising results in animal or ex vivo studies. However, due to a limited amount of research and unambiguous results, the topic of microbiome modulation as a therapeutic approach in skin diseases still warrants further investigation.

Keywords: microbiome; probiotics; prebiotics; synbiotics; atopic dermatitis; psoriasis; chronic ulcers; seborrheic dermatitis; burns; acne

1. Introduction

In 1683, Antoni van Leuwenhoek made the first microscopic observation of bacteria colonizing the surface of the human skin [1]. Joshua Lederberg first suggested the term microbiome in 2000, meaning the collective genome of commensal, symbiotic and pathogenic bacteria, archaea and eukaryote of the human body [2].

The skin microbiome includes bacteria, fungi, viruses, micro-eukaryotes (mites), archaea, and phages [3]. They can be found not only on the surface of the epidermis, but also in sweat, sebaceous glands and associated hair follicles [4]. The composition of microbiome differs among different regions affected by numerous factors including age, gender, genetics, immunity, hormonal balance, sleep routine, stress, metabolic factors, hygiene and skin care routine, chemical or ultraviolet radiation exposure, physical activity, climate, environmental pollution and availability of nutrients [5]. The initial colonization of the skin depends on the delivery mode, with neonates delivered vaginally acquiring the species (spp.) present in the vaginal tract (e.g., Lactobacillus, Prevotella, Sneathia) in contrast to children delivered by Cesarean section, that acquire microbiome associated with the...
The skin microbiome of newborns is less complex than for adults [8]. In adults, the longest assessment of the skin microbiome composition lasted for 2 years, indicating that the skin microbiome remains rather stable despite changes in the environment [9]. The most dominant group in the skin microbiome are bacteria [10]. The most dominant species are *Staphylococcus epidermidis*, *Cutibacterium acnes* and *Corynebacterium*, which overall are estimated to constitute 45–80% of the skin microbiome [5]. Considering bacteria, sebaceous and dry areas are dominated by *Cutibacterium spp*. Moist environments, with a greater humidity level, harbor mostly *Staphylococcus* and *Corynebacterium* [11]. Regarding fungi, *Malassezia spp.* are present on the whole body surface, predominant in oily sites (face, back); however, in the foot site the fungal diversity is greater [11]. The viruses identified on the human skin include *Papillomaviridae*, *Polyomaviridae* and *Circoviridae* families [4]. The dust mites, found in 23–100% of the population, are considered commensals; however, it is known that *Demodex* mites may be associated with blepharitis and rosacea [3]. The data on the type and role of the phages are limited, yet it was found that they can modulate the skin microbiome [12]. Skin microbiome alterations were found in the background of numerous dermatological diseases, including acne, atopic dermatitis (AD) and psoriasis, among others [13,14].

Similarly to the skin, the gut microbiome starts to shape after the delivery. Depending on dominant genera, three robust clusters of intestinal microbiome, referred to enterotypes, may be distinguished: enterotype 1 with dominant *Bacteroides*, enterotype 2 with dominant *Prevotella* and enterotype 3 with dominant *Ruminococcus* [15]. The interactions between the gut microbiome and the skin are complex and not yet fully elucidated; however, several pathways bringing light to this topic may be found [16]. Gut bacterial dysbiosis may lead to reduced short-chain fatty acid (SCFAs) production, as well as disruption of the gut barrier integrity and increased permeability, that results in bacterial translocation, activation of immune cells to produce pro-inflammatory cytokines and promotes chronic, low-grade systemic inflammation [14]. Probiotics are defined by the World Health Organisation as living microorganisms that confer a health benefit when administered in adequate amounts [17]. Administering probiotics results in the stabilization of the gut bacterial community, restoration of the bacterial microbiota “signature” in the gut, producing bacteriocins, altering microRNA (miRNAs), competing with pathogens for certain nutrients and improving the gut barrier function [18].

Antibiotics have a diverse effects on the skin and gut microbiome’s ecological balance, depending on the antibiotic class, dosage and duration [19]. Antibiotic therapies are essential in the treatment of chronic dermatological diseases including acne and rosacea [20]. They possibly reduce the proportion of pathogenic bacteria and promote the growth of potentially beneficial microorganisms [21]. However, dysbiosis of the healthy gut microbiome’s composition, induced by antibiotics, can cause and aggravate disease [22]. Moreover, antibiotic therapies have a high rate of adverse reactions and their overutilization increases the probability of developing resistance [21].

Prebiotics are substances, such as carbohydrates or fibres, that can promote the growth of beneficial bacteria. They can be defined as selectively fermented ingredients that allow specific changes in the composition as well as in the activity of the gastrointestinal microflora. Similarly to probiotics, they confer benefits upon host health. As prebiotics are typically fibers that cannot be digested by the host, they are metabolized by the gut microbiome in the colon, which results in an abundance of certain bacterial species and metabolites production, including SCFAs [4]. The combination of probiotics and prebiotics, administered simultaneously, is referred to as ‘synbiotics’, where the two agents show synergism [23].

Pre- and probiotics, modifying the gut microbiome, may be used for targeting skin health [24]. In the present paper, it was aimed to review the current knowledge concerning skin diseases, in which the supplementation of pre- or probiotics is beneficial via the modulation of the skin or gut microbiome.
2. Materials and Methods

A search of PubMed (MEDLINE) and EMBASE databases was conducted, using a combination of keywords such as: “microbiome”; “modulation”; “prebiotic”; “probiotic”; “skin”; “skin disease” using MeSH and Emtree methods. The majority of results concerned the following diseases: AD, psoriasis, chronic wounds, SD, burns and acne. A second search in PubMed (MEDLINE) and EMBASE databases was conducted, using a combination of keywords such as: “probiotic” or “prebiotic” and the name of each of the mentioned diseases. The literature review was based on the PRISMA principles (Figure 1). Works in English published until June 2021 were included with inclusion criteria as follows: full text articles available, use of probiotics or prebiotics to treat the skin diseases. The total number of records considered into analysis was 563 on AD, 188 records on psoriasis, 628 records on ulcers, 20 records on SD, 126 records on burns and 165 records on acne. After the duplicates were removed, 470 records on AD, 101 on psoriasis, 78 on ulcers, 17 on SD, 30 on burns and 165 on acne were further analyzed. As a very limited number of studies were available on the topic of microbiome modulation with probiotics and prebiotics supplementation in all skin diseases except for AD, the authors decided to include case reports, original studies on animal or in vitro model in cases of other skin diseases. 113 articles were included in the final analysis concerning the effects of probiotics, prebiotics and synbiotics supplementation in AD, SD, psoriasis, burns, chronic ulcers and acne. We analyzed the number of patients, race, country of origin of the study population, study and control group size, study type, type of intervention and outcomes.

![Figure 1. PRISMA Flow Diagram for the identification and screening of the included studies.](image-url)
3. Results

**Atopic dermatitis.** We identified 21 original studies which investigate the influence of probiotic supplementation in pregnant women and newborns with family history of AD or allergic diseases (Table 1). Additionally, 11 studies assessed the prevention of AD with prebiotics (Table 2). 37 original studies present the treatment of AD with probiotics and prebiotics in infants (Table 3), children (Table 4) and adults (Table 5). No published data are available on the use of prebiotics in the treatment of adult patients with AD.

**Psoriasis.** Three publications concerning the administration of probiotics were found—one case report and two original studies. Four original studies investigating animal models were identified. No studies concerning prebiotics administration in psoriatic patients were found. The results are presented in Table 6.

**Chronic ulcers.** One case report and two clinical trials were found. Two studies concerning probiotics supplementation in the animal model were also identified; the authors also included two in vitro studies on probiotics application (Table 7). No published data on the topic of prebiotics supplementation in chronic ulcers was found.

**Seborrheic dermatitis.** Two clinical trials addressing oral administration of probiotics and one concerning topical administration were identified. One of the studies reported SD as a side effect of probiotic administration (Table 8).

**Burns.** 16 studies reporting the influence of pre- or probiotics on the healing of burns, the permeability of gut barrier in patients suffering from burns or the complications including sepsis were found, including seven original studies and two case reports. Both in clinical trials and in the animal model, the oral and topical administration of probiotics was investigated. One animal study also investigated the effects of local administration with daily sub-eschar injections. A single study on the prebiotics influence on the gut barrier permeability was also reported (Table 9).

**Acne.** 11 studies on the effects of probiotics and prebiotics on acne were included in the final analysis: seven clinical trials, divided into two groups with oral supplementation or topical application, and four in vitro studies. Studies concerning the use of bacterial strains not considered as probiotics (C. acnes, S. epidermidis) or probiotics modulating gut microbiome during antibiotic therapy were excluded from the analysis. The results are presented in Table 10.
Table 1. Prebiotics in the prevention of AD.

| Number (No.) of Study | Author | Patient Population (Number) | Type of Study | Intervention | Results |
|-----------------------|--------|-----------------------------|---------------|--------------|---------|
| 1                     | Kalliomaki et al.—2007 [25] | Pregnant women (n = 159) with a family history of AD, continuing after delivery and their children (n = 132) | Double-blind, randomised placebo-controlled trial | Participants received two capsules of placebo (n = 95) or LGG (n = 64) daily for 2–4 weeks before expected delivery | The frequency of AD was significantly reduced |
| 2                     | Rautava et al.—2006 [26] | Infants with 6 months of exclusive breast-feeding (n = 38) | Double-blind placebo-controlled study | Infant formula supplemented with either LGG and Bb-12 (n = 38) or placebo (microcrystalline cellulose) (n = 43) daily until the age of 12 months | Supplementation of probiotics increased protective cow’s milk-specific IgA responses. 13% of the infants receiving probiotics and 20% of those receiving placebo manifested with AD, cow’s milk allergy was confirmed in none of the infants receiving probiotics and in 8% of the infants receiving placebo |
| 3                     | Abrahamsson et al.—2007 [27] | Pregnant women with a family history of at least 1 allergic disease (n = 188) and then their infants (n = 188) | Prospective, double-blind, placebo-controlled, multicenter trial | The mothers were taking *L. reuteri* (n = 95) or placebo (n = 93) 4 weeks before term and continued daily until delivery, after birth, the baby continued with the same product up to 12 months of age | The cumulative incidence of AD was similar in the probiotic and the placebo groups (36% vs. 34%) |
| 4                     | Taylor et al.—2007 [28] | Infants with atopic mother (n = 178) | Randomized, double-blind, placebo-controlled | Newborns of women with allergy received either *L. acidophilus* (n = 89) or placebo (n = 89) daily for the first 6 months of life | Not reduction in the risk of AD and increased allergen sensitization |
| 5                     | Wickens et al.—2008 [29] | Pregnant women (n = 474) and their infants (n = 474) | Double-blind randomized placebo-controlled trial | Daily supplementation with either HN001 (n = 157) or HN019 (n = 158) or placebo (n = 159) from 35 weeks gestation until birth, continuing to 6 months after birth in mothers if breastfeeding, and from birth till 2 years in all infants | Prevention of the development of AD |
| Number (No.) of Study | Author | Patient Population (Number) | Type of Study | Intervention | Results |
|----------------------|--------|-----------------------------|---------------|--------------|---------|
| 6                    | Huurre et al.—2008 [30] | Pregnant women (n = 140) and infants (n = 138) | Placebo-controlled prospective intervention study | Oral administration of LGG and Bb-12 each day (n = 72/70), or placebo (microcrystalline cellulose and dextrose anhydrate) (n = 68). Atopic sensitization was at the age of 6 and 12 months and in mothers at third trimester of pregnancy | There was no difference between infant sensitization in the probiotic and the placebo group |
| 7                    | Kopp et al.—2008 [31] | Pregnant women (n = 105) with a family history of at least one allergic disease and their children (n = 96) | Double-blind, placebo-controlled prospective trial | Administration of either the probiotic LGG (n = 54) twice daily or placebo (n = 51) 4-6 weeks before expected delivery, followed by a postnatal period of 6 months | After a 2 year follow-up, administration of probiotic did not reduce the incidence nor altered the severity of AD |
| 8                    | West et al.—2009 [32] | Healthy infants with birth weight >2500 g who were vaginally delivered (n = 89) | Double-blind, placebo-controlled randomized intervention trial | Daily intake of cereals supplemented with LF19 (n = 89) or identical cereals without LF19 supplementation (n = 90) from 4 to 13 months of age | Decreased cumulative incidence of AD |
| 9                    | Niers et al.—2009 [33] | Pregnant women (156) and then their infants with a positive family history of allergic disease (n = 156) | Double-blind, randomized, placebo-controlled trial | Probiotic bacteria were prenatally administered to pregnant mothers (n = 78) during the last 6 weeks of pregnancy and postnatally for 12 months to their infants (n = 78); the intervention group received once daily B. bifidum W23, B. lactis W52, and L. lactis W58 in a freeze dried powder | Decreased incidence of AD |
| 10                   | Soh et al.—2009 [34] | Infants with a positive family history of allergic disease (n = 253) | Double-blind, placebo-controlled randomized clinical trial | Infants (n = 127) received commercially available cow’s milk formula with probiotic supplementation of BL999 and L. rhamnosus daily for the first 6 months. Infants in the control group (n = 126) received milk without probiotics | No effect on the prevention of AD or allergen sensitization |
| Number (No.) of Study | Author | Patient Population (Number) | Type of Study | Intervention | Results |
|-----------------------|--------|-----------------------------|---------------|--------------|---------|
| 11                    | Kim et al.—2010 [35] | Pregnant women with a family history of allergic diseases (n = 112), continuing after delivery and their infants (n = 68) | Double-blind, randomized, placebo-controlled trial | Pregnant woman received supplement of *B. bifidum* BGN4, *B. lactis* AD011 and *L. acidophilus* AD03 (n = 33) or placebo (n = 35), starting at 4–8 weeks before delivery and continuing until 6 months after delivery. Infants were exclusively breastfed during the first 3 months, and were fed with breastmilk or cow’s milk formula from 4 to 6 months of age | The prevalence of AD in the first year of life was significantly lower in the probiotic group. |
| 12                    | Dotterud et al.—2010 [36] | Pregnant women (n = 415) and their infants (n = 278) | Randomized, double-blind trial | Pregnant women received probiotic milk (n = 138) or placebo (n = 140) from 36 weeks of gestation to 3 months postnataally during breastfeeding | Decreased cumulative incidence of AD. |
| 13                    | Boyle et al.—2011 [37] | Pregnant women (n = 250), their partner or a previous child was affected by allergic disease including asthma, eczema, food allergy or allergic rhinitis | Randomized controlled trial | Participants were allocated to take probiotic treatment with LGG (n = 125) or maltodextrin placebo (n = 125) each morning from 36 weeks gestation until delivery. Infants were assessed during their first year for eczema or allergic sensitization | Prenatal treatment was not associated with reduced risk of eczema or IgE-associated eczema but decreased breast milk soluble CD14 and IgA levels |
| 14                    | Rautava et al.—2012 [38] | Pregnant women with atopic sensitization (n = 241) and their infants (n = 205) | Double-blind, randomized, placebo-controlled trial | Pregnant women received a dietary food supplement with the combination of LPR and BL999 (n = 81) or the combination of ST11 and BL999 (n = 82) or placebo (78) | Administration of specific probiotics is a safe and effective way in reducing the risk of AD |
| 15                    | Ou et al.—2012 [39] | Pregnant women with atopic diseases determined by history, total immunoglobulin (Ig)E > 100 kU/L, and/or positive specific IgE (n = 191) | Prospective, double-blind, placebo-controlled clinical trial | Pregnant woman received either LGG ATCC 53103 (n = 95) or placebo (n = 96) from the second trimester of pregnancy | Reduced severity of maternal allergic disease |
### Table 1. Cont.

| Number (No.) of Study | Author | Patient Population (Number) | Type of Study | Intervention | Results |
|-----------------------|--------|-----------------------------|---------------|--------------|---------|
| 16                    | Lau et al.—2012 [40] | Infants with at least single heredity for atopy (n = 606) | Randomized, placebo-controlled trial | From week 5 until the end of month 7, infants were treated orally with bacterial lysate containing heat-killed gram-negative *E. coli* and gram-positive *E. faecalis* (n = 303) or placebo (n = 303) | Prevention of the development of AD |
| 17                    | Allen et al.—2014 [41] | Pregnant women (n = 454) and then their infants with a positive family history of allergic disease (n = 454) | Randomised, double-blind, placebo-controlled, parallel group trial | Women from 36 weeks gestation and their infants to age 6 months received daily either the probiotic (*L. salivarius* CUL61, *L. paracasei* CUL08, *B. animalis* subsp. *lactis* CUL34 and *B. bifidum* CUL20) (n = 220) or placebo (n = 234) | Cumulative frequency of AD at 2 years of age was similar between the two groups |
| 18                    | Cabana et al.—2017 [42] | Infants (n = 184) | Randomized, double-blind controlled trial | The intervention group received a daily capsule of LGG and inulin for the first 6 months of life (n = 92); the control group received inulin (n = 92) | At 5 years of age, the cumulative incidence of asthma was significantly higher in the control group (17.4%) than in the intervention group (9.7%) |
| 19                    | Wickens et al.—2018 [43] | Pregnant women, continuing after giving birth. The patient or her partner had a history of atopic disease (n = 473) | 2-centre, parallel double-blind, randomized placebo-controlled trial | *HN001* (n = 157), *HN019* (n = 158) or placebo (n = 159) was taken daily by mothers from 14-16 weeks of gestation until 6 months post-partum. Their infants were also given the same capsule daily from birth until the age of 2 years | Prevention of the development of AD and atopic sensitization |
| 20                    | Plummer et al.—2019 [44] | Preterm infants, born <32 gestational week and weighing <1500 g (n = 281) | Multi-center, double-blind, placebo-controlled randomized trial | Infants in the intervention group (n = 127) received a probiotic combination *B. infantis*, *Str. thermophilus*, and *B. lactis* once daily (in a maltodextrin base powder) and the placebo group (n = 154) received maltodextrin | No effect on the incidence of allergic diseases or atopic sensitization |
| 21                    | Schmidt et al.—2019 [45] | Infants with birthweight >2500 g, gestational age >36 weeks (n = 144) | Double-blind, placebo-controlled intervention trial | The intervention group (n = 144) received sachets of maltodextrin supplemented with LGG and Bb-12, and the placebo group (146) received maltodextrin only | A significantly lower incidence of AD in the probiotic group |

**Abbreviations:** AD, atopic dermatitis; *B.*, *Bifidobacterium*; Bb-12, *Bifidobacterium lactis* Bb-12; BL999, *Bifidobacterium longum* BL999; *E. coli*, *Escherichia coli*; *E. faecalis*, *Enterococcus faecalis*; IgE, immunoglobulin E; IgA, immunoglobulin A; *L.*, *Lactobacillus acidophilus*; LF19, *Lactobacillus paracasei* F19; LGG, *Lactobacillus rhamnosus* GG; *HN001*, *Lactobacillus rhamnosus* HN001; *HN019*, *Bifidobacterium animalis* subsp. *lactis* strain HN019; *Str.*, *Streptococcus*; ST11, *Lactobacillus paracasei* ST11; No, number.
### Table 2. Prebiotics in the prevention of AD.

| No. of Study | Author             | Patient Population (Number)                                      | Type of Study                           | Intervention                                                                                                                                  | Results                                                                                           |
|-------------|--------------------|-----------------------------------------------------------------|----------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------|
| 1           | Moro et al.—2006  | Infants at risk for atopy (n = 259)                              | Prospective, double-blind, randomised, placebo controlled trial | Participants received either hydrolysed cows’ milk with GOS/FOS in the prebiotic group (n = 102) or maltodextrin in the control group (n = 104) | Development of AD was significantly more frequent in the control group                             |
| 2           | Ziegler et al.—2007 | Healthy infants (n = 226)                                        | Double-blind, randomized, controlled, parallel-group, prospective trial | Participants were divided into 3 different formula groups: control group-PDX (n = 76), PG4 group-PDX+GOS (n = 74), PDL8 group-PDX, GOS, LOS (76). Formula was fed for 120 days | No differences among the groups in growth rate                                                   |
| 3           | Arslanoglu et al.—2008, 2012 | Healthy infants with a parental history of atopy (n = 134)  | Prospective, double-blind, randomised, placebo controlled trial | Participants received either GOS/FOS prebiotic supplement in the intervention group (n = 66) or maltodextrin supplementation in the control group (n = 68) | Cumulative incidences for AD, recurrent wheezing, and allergic urticaria were higher in the placebo group after 2 years |
| 4           | Grüber et al.—2010, 2016 | Healthy infants with low risk of atopy (n = 1130)               | Double-blind, controlled, randomized, prospective intervention study | Participants were divided into three groups: prebiotic group (n = 414) - mixture of GOS, FOS, pAOS, breastfed group (n = 300), control group (n = 416) | After 1 year, AD occurred in significantly fewer infants from the prebiotic group                |
| 5           | Niele et al.—2012  | Preterm infants (n = 94)                                         | Prospective, double-blind, randomised, placebo controlled trial | Volunteers received either enteral GOS, FOS and pAOS supplementation (n = 48) or placebo (n = 46) during first month of life | No decrease in the incidence of allergic and infectious diseases during first year of life      |
| 6           | Pontes et al.—2016 | Healthy children (1–4 years of age) (n = 256)                   | Double-blind, randomized, controlled trial | The intervention group (n = 125) received cow’s milk-based beverage containing DHA, PDX, GOS, β-glucan and the control group (n = 131) were fed cow’s milk three servings/day up to 28 weeks | Participants in the intervention group were associated with fewer episodes of allergic manifestations |
| 7           | Boyle et al.—2016  | Infants with an atopic parent (n = 1047)                        | Parallel-group, multicentre, randomized double-blind controlled trial | Three groups: prebiotic group (n = 432) - mixture of GOS, FOS, pAOS, breastfed group (n = 184), control group (n = 431) | Prebiotics did not prevent AD in the first year of life                                          |
| 8           | Ranucci et al.—2018 | Infants (n = 400) with an atopic parent                          | Randomised, double-blind, placebo-controlled trial | Participants received either prebiotic formula containing GOS/PDX (n = 201) or standard formula (n = 199) in the first 48 weeks of life | No significant differences in the cumulative incidence of AD and its intensity and duration between groups |
### Table 2. Cont.

| No. of Study | Author                          | Patient Population (Number) | Type of Study                                           | Intervention                                                                                     | Results                                                                                                                                 |
|-------------|--------------------------------|-----------------------------|---------------------------------------------------------|-------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------|
| 9           | Wopereis et al.—2018 [56]      | Healthy infants (n = 138)   | Double-blind, randomized, controlled parallel-group nutritional intervention trial | Participants were divided into three groups: prebiotic group (n = 51) -mixture of GOS, FOS, pAOS, breastfed group (n = 30), control group (n = 57) | Metabolites and pH of infants receiving GOS/FOS/pAOS was closer to breastfed infants than to infants receiving standard cow’s milk formula. After 18 months, AD occurred in significantly fewer infants in the prebiotic group |

Abbreviations: AD, atopic dermatitis; FOS, fructo-oligosaccharides; GOS, galacto-oligosaccharides; LOS, lactulose; pAOS, acidic oligosaccharides; PDX, polydextrose.

### Table 3. Pre- and probiotics in the treatment of AD in infants.

| No. of Study | Author                          | Patient Population (Number) | Type of Study                                           | Intervention                                                                                                                                                                                                 | Results                                                                                                                                 |
|-------------|--------------------------------|-----------------------------|---------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------|
| 1           | Majamaa et al.—1997 [57]        | Infants aged 2.5 to 15.7 months with AD (n = 27), nursing mothers of infants with AD (n = 10) | Randomized controlled trial                           | Infants with AD and cow’s milk allergy received cow’s milk without (n = 14) and with (n = 13) the addition of LGG; the second part of the study involved 10 breast-fed infants who had AD and cow’s milk allergy. In this group LGG was given to nursing mothers | Probiotic bacteria downregulated hypersensitivity reactions and intestinal inflammation                                               |
| 2           | Isolauri et al.—2000 [58]       | Infants with AD, mean age of 4.6 months (n = 27) | Randomized double-blind placebo-controlled study       | Probiotic-supplemented, Bb-12 (n = 9) or LGG ATCC 53103 (n = 9), extensively hydrolysed whey formulas or to the same formula without probiotics (n = 9)                                                                 | First clinical demonstration of specific probiotic strains modifying AD                                                                 |
| 3           | Kirjavainen et al.—2003 [59]    | Infants with AD, mean age was 5.5 months (n = 43) | Randomized double-blind manner                          | Infants were randomly assigned into placebo (n = 10), viable LGG (n = 17), or heat-inactivated LGG groups (n = 16) and extensively hydrolyzed whey formula or the same formula supplemented with viable or heat-inactivated LGG               | Supplementation of infant formulas with viable but not heat-inactivated probiotic was effective for the management of AD and cow’s milk allergy |
| No. | Study | Author | Patient Population (Number) | Type of Study | Intervention | Results |
|-----|-------|--------|-----------------------------|---------------|-------------|---------|
| 4   | Viljanen et al.—2005 [60] | Infants with AD under the age of 12 months (n = 230) | Randomized double-blinded study | First group (n = 80) received capsules containing LGG ATCC 53103; the second group (n = 76) a mixture of probiotics: LGG, *L. rhamnosus* LC705, *B. breve* BB9, and *Propionibacterium* JS; and there was a placebo group (n = 74) | Treatment with *L. rhamnosus* alleviated AD symptoms in IgE-sensitized infants |
| 5   | Weston et al.—2005 [61] | Children aged 6-18 months with moderate or severe AD (n = 56) | Randomized double blind placebo controlled trial | The children were given a *L. fermentum* VRI-033 PCC (n = 28) or placebo (n = 28), twice daily for 8 weeks | Supplementation with probiotic bacteria is beneficial in improving the extent and severity of the symptoms |
| 6   | Taniuchi et al.—2005 [62] | Infants with cow milk hypersensitivity and AD (n = 10) | Randomised placebo controlled trial | Orally given lyophilized bifidobacteria *B. breve* M-16V (n = 10) strain or placebo (n = 7) | Significantly increased proportion of bifidobacteria in the fecal microflora |
| 7   | Folster-Holst et al.—2006 [63] | Infants (n = 54) aged 1-55 months with moderate-to-severe AD | Randomized, double-blind, placebo-controlled study | Participants received a hydrolysed whey-based formula as placebo (n = 17), or supplemented with either *L. rhamnosus* (n = 17) or LGG (n = 16) for 3 months | No significant differences between the groups in the clinical symptoms |
| 8   | Brouwer et al.—2006 [64] | Infants less than 5 months old with AD (n = 50) | Randomized, double-blind, placebo-controlled study | LGG (n = 26) or placebo (n = 27) was received during an 8-week period | No clinical or immunological effect of *L. rhamnosus* |
| 9   | Grüber et al.—2007 [65] | Infants with AD aged 3–12 months (n = 54) | Randomized trial | LGG (54) or placebo (48) as a food supplement for 12 weeks | No therapeutic effect of probiotic against mild to moderate AD |
| 10  | Flintermann et al.—2007 [66] | Children aged 0.5–2.8 years with AD (n = 13) | Randomized trial | Probiotics (n = 7) or placebo (n = 6) was randomly assigned to the patients. The probiotics contained a mixture of *L. acidophilus* W55, *L. casei* W56, *L. salivarius* W57, *L. lactis* W58, *B. infantis* W52, *B. lactis* W18 and *B. longum* W51 | Probiotics enhanced the production of Th1 and regulatory cytokines *in vitro* |
| 11  | van der Aa LB et al.—2010 [67] | Infants with AD SCORAD > or =15, aged < 7 months and exclusively formula fed (n = 90) | Double-blind, placebo-controlled multi-centre trial | Extensively hydrolysed formula with *B. breve* M-16V and a galacto-/fructo-oligosaccharide mixture (n = 46) or the same formula without synbiotics (n = 44) for 12 weeks | Symbiotic mixture does not have a beneficial effect on the severity of AD, but it modulates the intestinal microbiota |
| No. of Study | Author                        | Patient Population (Number)                                                                 | Type of Study                                                                 | Intervention                                                                                                                                                                                                 | Results                                                                                          |
|-------------|-------------------------------|-------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------|
| 12          | Gøbel et al.—2010 [68]        | Children from 7 to 24 months of age with AD (n = 50)                                        | Randomised double-blind placebo-controlled trial                             | First group: *L. acidophilus* NCFM and other supplements in a capsule given (n = 17), Second group: *B. lactis* Bi-07 and supplements in a capsule given (n = 17). Third group received placebo (n = 16). Treatment was given daily for 8 weeks | No overall beneficial effects on the degree of SCORAD index.                                    |
| 13          | Nermes et al.—2011 [69]       | Infants with AD (n = 39)                                                                   | Double-blind study                                                           | Extensively hydrolysed casein formula supplemented with (n = 19) or without (n = 20) LGG (ATCC 53103) was given to the two different groups for three months                                                             | Probiotics may enhance gut barrier function                                                      |
| 14          | Farid et al.—2011 [70]        | Infants and children aged 3 months to 6 years with AD (n = 40)                             | Randomized, double-blind, placebo-controlled study                          | Patients in the probiotic group (n = 19) received synbiotic containing a mixture of *L. casei*, *L. rhamnosus*, *Str. thermophilus*, *B. breve*, *L. acidophilus*, *B. infantis*, *Lactobacillus bulgaricus* and FOS twice daily for 8 weeks | Mixture of probiotics and FOS improved the severity of symptoms                                 |
| 15          | Gore et al.—2012 [71]         | Infants with AD (n = 208)                                                                  | Randomized-controlled trial                                                  | Infants were randomized to daily supplements containing *L. paracasei* or *B. lactis* (n = 137) or placebo (n = 71) for a 3-month period, while receiving extensively hydrolysed whey-formula (dairy-free diet) | No benefit in the treatment of eczema and no effect on the progression of allergic disease       |
| 16          | Shafiei et al.—2011 [72]      | Infants aged 1-36 months with moderate-to-severe AD (n = 41)                               | Randomized double-blind placebo controlled trial                           | Mixture of seven strain probiotics plus FOS (n = 20) or placebo (n = 21), administered daily as a powder for two months                                                                                 | No improvement of AD                                                                             |
| 17          | Ivakhnenko et al.—2013 [73]   | Infants aged of 3-12 months with the diagnosis of AD and allergy to cow’s milk protein (n = 60) | Open randomized prospective clinical study                                   | Bb-12 and *Str. thermophilus* TH-4 intake for half of volunteers (n = 30). The other half of the volunteers (n = 30) received placebo for 4 weeks                                                                 | Improved clinical symptoms                                                                      |
| 18          | Lin et al.—2015 [74]          | Infants with AD (n = 40)                                                                   | Randomized controled study                                                  | The intervention group (n = 20) received *B. bifidum* triple viable capsules for 4 weeks with a dosage of one capsule three times a day. The control group (n = 20) were not given a placebo drug | Positive effect on the prevention and treatment                                                 |
Table 3. Cont.

| No. of Study | Author                    | Patient Population (Number) | Type of Study                      | Intervention                                                                 | Results                                                                 |
|--------------|---------------------------|----------------------------|-----------------------------------|-----------------------------------------------------------------------------|-------------------------------------------------------------------------|
| 19           | Guo et al.—2015 [75]      | Adult AD patients (n = 180) | Randomized trial                  | Participants were divided into two groups. Participants received routine symptomatic treatment and combination of probiotics (microecologics) (n = 90) or symptomatic treatment (n = 90) orally twice a day for one month | Application of microecologics as an adjuvant therapy was effective     |
| 20           | Wu et al.—2017 [76]       | Children aged 4-48 months with AD and with SCORAD ≥ 15 at enrollment. (n = 66) | Two-center, double-blinded, randomized and placebo-controlled study | Treatment group (n = 33)—one capsule containing L. rhamnosus a day, control group (n = 33)—one capsule of placebo a day for 8 weeks | Probiotic was effective in decreasing AD symptoms                      |

Abbreviations: AD, atopic dermatitis; B., Bifidobacterium; Bb-12, Bifidobacterium lactis Bb-12; FOS, fructo-oligosaccharides; IFNy, Interferon gamma; IL-10, Interleukin 10; L., Lactobacillus; LGG, Lactobacillus rhamnosus GG; Propionibacterium JS, Propionibacterium freudenreichii sup. shermanii JS; SCORAD, Scoring Atopic Dermatitis; Str.: Streptococcus, Th1, T helper type 1 cells.

Table 4. Pre- and probiotics in the treatment of AD in children.

| No. of Study | Author                    | Patient Population (Number) | Type of Study                      | Intervention                                                                 | Results                                                                 |
|--------------|---------------------------|----------------------------|-----------------------------------|-----------------------------------------------------------------------------|-------------------------------------------------------------------------|
| 1            | Rosenfeldt et al.—2003 [77]| Children aged 1 to 13 years with AD (n = 43) | Double-blind, placebo-controlled, crossover study | The patients were randomized in two groups to receive either placebo followed by active treatment or active treatment followed by placebo. 2 probiotic lyophilized L. rhamnosus 19070-2 and L. reuteri DSM 122460 were given in combination for 6 weeks | Combination of probiotics was significantly effective in the management of AD |
| 2            | Sistek et al.—2006 [78]   | Children aged between 1 and 10 years with AD (n = 59) | Randomized controlled trial       | L. rhamnosus and B. lactis (n = 29) or placebo (n = 30) were given daily as a powder for 12 weeks | Combination of probiotic bacteria improved AD only in food sensitized children |
| 3            | Passeron et al.—2006 [79] | Children aged at least 2 years with AD (n = 48) | Double-blind prospective randomized study | L. rhamnosus Lcr35 plus prebiotic preparation (n = 24) or prebiotic preparation alone (n = 24) was given three times a day for 3 months | Both synbiotics and prebiotics used alone seem able to significantly improve the manifestations of AD |
Table 4. Cont.

| No. of Study | Author                      | Patient Population (Number)                  | Type of Study                          | Intervention                                                                 | Results                                                                 |
|--------------|-----------------------------|----------------------------------------------|----------------------------------------|------------------------------------------------------------------------------|--------------------------------------------------------------------------|
| 4            | Gerasimov et al.—2010 [80]  | Children aged 1–3 years with moderate-to-severe AD (n = 90) | Randomized, double-blind, placebo-controlled, prospective trial | Infants were randomly assigned into placebo (n = 47), and intervention group (n = 43). Mixture of *L. acidophilus* DDS-1, *B. lactis* UABLA-12 with fructo-oligosaccharide or placebo twice daily for 8 weeks | Significant clinical improvement                                           |
| 5            | Woo et al. —2010 [81]       | Children aged 2 to 10 years with AD (n = 45)  | Double-blind, placebo-controlled trial | Volunteers received either *L. sakei* KCTC 1075SBP (n = 45) or placebo (n = 33) daily for 12 weeks | Substantial clinical improvement and a significant decrease in chemokine levels |
| 6            | Han et al.—2012 [82]        | Children aged 1–13 years presenting with AD (n = 83) | Randomized, double-blind, placebo-controlled study | *L. plantarum* CJLP133 (n = 44) or placebo (n = 39) was given to children twice a day for 12 weeks. SCORAD scores, eosinophil counts, serum total IgE, IFN-γ and IL-4 were evaluated | SCORAD score at week 14 was significantly lower in the probiotic group |
| 7            | Wu et al.—2012 [83]         | Children aged 2-14 years with moderate-to-severe AD (n = 54) | Double-blind, randomized, clinical trial | One capsule twice daily for 8 weeks containing either *L. salivarius* and FOS (n = 27) or FOS only (n = 27) | Symbiotic combination was superior to the prebiotic alone |
| 8            | Yesilova et al.—2012 [84]   | Children suffering from a moderate-to-severe AD, 1-13 years of age (n = 40) | Double-blind, randomized, placebo-controlled study | The probiotic group (n = 20) was administered with a probiotic complex containing *B. bifidum*, *L. acidophilus*, *L. casei*, and *L. salivarius* for 8 weeks. The placebo group (n = 20) was administered skim milk powder and dextrose | Probiotics to be effective in reducing SCORAD index, serum IL-5, IL-6, IFN-γ, and total serum IgE levels but not effective in reducing serum IL-2, IL-4, IL-10, ECP, or TNF-α levels |
| 9            | Yang et al.—2014 [85]       | Children aged 2-9 years with AD (n = 100)     | Randomized, double-blind, placebo-controlled, parallel trial | Randomly allocated to the probiotics (L. casei, L. rhamnosus, L. plantarum, B. lactis) (n = 50) or placebo (n = 50) groups for 6 weeks | Probiotics successfully colonized in the intestine; but additional effects were not found |
| 10           | Wang et al.—2015 [86]       | Children aged 1-18 years with moderate-to-severe AD (n = 210) | Double-blind, prospective, randomized placebo-controlled study | The groups received *L. paracasei* (n = 55) or *L. fermentum* (n = 55) or *L. paracasei* and *L. fermentum* mixture (n = 55) or placebo (n = 55) for 3 months | Supplementation of a probiotic mixture was associated with clinical improvement |

Abbreviations: AD, atopic dermatitis; *B.*, *Bifidobacterium*; ECP, Eosinophil cationic protein; FOS, fructo-oligosaccharides; IFNγ, Interferon gamma; IgE, immunoglobulin E; IL-2, interleukin 2; IL-4, interleukin 4; IL-5, interleukin 5; IL-6, interleukin 6; IL-10, interleukin 10; *L.*, *Lactobacillus*; SCORAD, Scoring Atopic Dermatitis; TNF-α, tumor necrosis factor alpha.
| No. of Study | Author | Patient Population (Number) | Type of Study | Intervention | Results |
|--------------|--------|-----------------------------|---------------|--------------|---------|
| 1            | Roessler et al.—2008 [87] | Adults with AD (n = 15) and healthy adults (n = 15) | Double-blind, placebo-controlled, randomized cross-over study | Probiotic containing a combination of probiotics *L. paracasei* Lpc-37, *L. acidophilus* 74-2 and *B. lactis* DGCC 420 in healthy volunteers (n = 15) and in patients with AD (n = 15) given over 8 weeks | Probiotic bacteria transiently colonized the intestines |
| 2            | Yoshida et al.—2010 [88]  | Adults with AD (n = 24) | Randomized, placebo-controlled study | Intervention group (n = 16) were given either *B. breve* strain YY or patients received placebo (n = 8) for 8 weeks | Probiotic was beneficial for the treatment of AD |
| 3            | Drago et al.—2012 [89]    | Adult patients between 18 and 46 years with moderate-to-severe AD (n = 38) | Parallel-group double-blind placebo-controlled randomised trial | Clinical efficacy of the intake of *L. salivarius* LS01 (n = 19) in the treatment of adult patients with AD | Positively modified clinical and immunologic status and life quality |
| 4            | Iemoli et al.—2012 [90]   | Adult AD patients (n = 48) | Randomized double-blinded active treatment versus placebo study | Intake of a combination of two probiotics: *L. salivarius* LS01 and *B. breve* BR03 for 12 weeks in the probiotic group (n = 16) | Beneficial effects for clinical and immunologic alterations |
| 5            | Matsumoto et al. [91]     | Adult patients with AD (n = 44) | Randomized controlled trial | Patients were randomly assigned to receive LKMS12 (n = 22) or a placebo (n = 22) | LKMS12 exerted antipruritic effects by increasing kynurenic acid production |
| 6            | Drago et al.—2014 [92]    | Adult patients with AD (n = 25) | Prospective, controlled pilot trial | *L. salivarius*, *Str. thermophilus* ST10 and tara gum intake for half of participants (n = 13). The other half of the participants (n = 12) received placebo for 1 month | The combination of tara gum and probiotics increases the efficacy of other probiotic strains |
| 7            | Nakatsuji et al.—2021 [93]| Adult patients with AD (n = 54) | Double-blinded, randomized trial | 1-week trial of topical *Staphylococcus hominis* A9 (Sha9) or vehicle on the forearm skin of 54 adults with *S. aureus*-positive AD | Participants receiving Sha9 had fewer adverse events associated with AD; eczema severity was not significantly different when evaluated in all participants treated with Sha9 but a significant decrease in S. aureus and increased Sha9 DNA were seen |

Abbreviations: AD, atopic dermatitis; *B.*, *Bifidobacterium*; *L.*, *Lactobacillus*; LKMS12, *Bifidobacterium animalis* subsp. *lactis* LKM512; *S.*, *Staphylococcus*; *Str.*, *Streptococcus*.
Table 6. Probiotic application in psoriasis.

| No. of 
Study | Author                           | Patient Population (Number) | Type of Study                                    | Intervention                                                                 | Results                                                                 |
|---------|----------------------------------|-----------------------------|--------------------------------------------------|-----------------------------------------------------------------------------|--------------------------------------------------------------------------|
| 1       | Vijayashankar, Raghunath.—2012 [94] | A patient with generalised pustular psoriasis (n = 1) | Case report                                      | L. sporogene, one sachet thrice daily                                     | In 15 days, the fever subsided, lesions started involuting and no new lesions appeared |
| 2       | Groeger et al.—2013 [95]         | Patients with psoriasis (n = 26), patients with ulcerative colitis and chronic fatigue syndrome (n = 70), healthy volunteers (n = 35) | Randomized, double-blind, placebo-controlled    | Sachets containing B. infantis 35264 (n = 63) or placebo containing maltodextran (n = 55) daily for 8 weeks | Significant decrease in CRP and TNF-α levels |
| 3       | Navarro-Lopez et al.—2019 [96]   | 18–70 year old adults with plaque psoriasis (n = 90) | Randomized, double-blind, placebo-controlled    | Participant were randomized into probiotic (n = 45) and placebo (n = 45) groups. In the probiotic group capsule containing a mixture of 3 probiotic strains in 1:1:1 ratio (B. longum CECT 7347, B. lactis CECT 8145 and L. rhamnosus CECT 8361) was given for 12 weeks | Lower risk of relapse following the administration of probiotic bacteria, which reduced PASI75 in 66.7% of the patients. In the placebo group, 41.9% of patients showed reduction. In PGA index 48.9% of the probiotic group reached a score of 0 or 1 compared to 30.2% in the placebo group |
| 4       | Chen et al.—2017 [97]            | Male BALB/c; imiquimod-induced epidermal hyperplasia and psoriasis-like skin inflammation (n = 24) | Animal study                                    | In the intervention group mice were fed orally with different doses of L. pentosus GMNL-77 or with the vehicle control (distilled water) for 7 consecutive days | Improvement of skin symptoms, decreased TNF-α, IL-6, IL-23, IL-17A/F, and IL-22 levels in the skin, and reduced number of IL-17- and IL-22-producing CD4+ T cells |
| 5       | Rather et al.—2018 [98]          | Mice with imiquimod-induced psoriasis-like skin inflammation (n = 30) | Animal study                                    | Mice divided into five different groups, 6 mice each: control group, imiquimod group, imiquimod+vaseline group, imiquimod+clobetasol group, and imiquimod+ethanolic extract of L. sakei Probio65 | Significant inhibition of imiquimod-induced skin inflammation |
Table 6. Cont.

| No. of Study | Author | Patient Population (Number) | Type of Study | Intervention | Results |
|--------------|--------|-----------------------------|---------------|--------------|---------|
| 6            | Lu et al.—2021 [99] | Female BALB/c mice (n = 60) | Animal study | Mice were separated into 10 groups (6 included in each group): control group, imiquimod group, methotrexate positive control group and probiotic groups (seven groups); CCFM667 *B. adolescentis*, CCFM1078 *B. breve*, CCFM1148 *B. animalis*, CCFM1147 and CCFM1074 *L. paracasei*, CCFM1032 and CCFM1040 *L. reuteri* | Four probiotic bacteria groups ameliorated psoriasis-like pathological characteristics and suppressed the release of IL-23/T helper cell 17 axis-related inflammatory cytokines |
| 7            | Ogawa C. et al.—2021 [100] | Mice with imiquimod-induced psoriasis | Animal study | Mice were administered *L. mesenteroides* for 21 days alongside the topical application of imiquimod on the dorsal skin for 6 consecutive days | Suppressed erythema, scaling, upregulated IL-17 production, increased levels of plasma deoxycholic acid, altered the faecal microbiota composition |

Abbreviations: *B.*, *Bifidobacterium*; CRP, C-reactive protein; IL-6, interleukin 6; IL-17, interleukin 17; *L.*, *Lactobacillus*; *L. mesenteroides*, *Leuconostoc mesenteroides*; *L. reuteri*, *Limosilactobacillus reuteri*; PASI75, Psoriasis Area Severity Index 75%; PGA, Physician Global Assessment.

Table 7. Probiotic application in chronic ulcers.

| No. of Study | Author | Patients (Number) | Type of Study | Intervention | Results |
|--------------|--------|-------------------|---------------|--------------|---------|
| HUMAN MODEL—PROBIOTIC SUPPLEMENTATION | | | | | |
| 1            | Peral et al.—2010 [101] | Patients aged 40–70 years of age; patients suffered from type 2 diabetes mellitus (n = 14); non-diabetic (n = 20); inclusion criteria: venous ulcer; infection and no signs of healing in the past 3 months, despite conventional medical treatment | Intervenional study | Wounds were treated with topical applications of a whole culture of *L. plantarum* ATCC; the culture was applied once-daily over a period of 10 days | After 30 days of treatment, a reduction of more than 90% of the wound area was observed in 43% and 50% of the diabetic and non-diabetic patients, respectively |
| 2            | Mohseni et al.—2018 [102] | Patients aged 40-85 years old with grade 3 diabetic foot ulcer (n = 60) | Randomized, double-blind, placebo-controlled trial | Participants were randomly divided into two groups (n = 30/group) to receive either probiotic or placebo daily for 12 weeks. The probiotic mix consisted of *L. acidophilus*, *L. casei*, *L. fermentum*, *B. bifidum* | Beneficial effects on ulcer size, glycaemic control, total cholesterol, CRP, plasma nitric oxide, total antioxidant capacity and malondialdehyde levels |
| No. of Study | Study  | Author            | Patients (Number)                                                                 | Type of Study          | Intervention                                                                                           | Results                                                                                       |
|-------------|--------|-------------------|----------------------------------------------------------------------------------|------------------------|--------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|
| 3           | 3      | Venosi et al.—2019 [103] | 83-year-old woman with a critical limb ischemia and an infected difficult-to-treat ulcerated cutaneous lesion of the right leg | Case report            | Mixture of probiotic bacteria (lyophilized powder sachets, containing *Lactobacillus plantarum*, *Lactobacillus acidophilus* and *Str. thermophilus*) against different bacteria species *K. pneumonia*, *P. mirabilis* and *E. faecalis* | Treatment was effective against the three bacteria species                                      |
| 4           | 4      | Jones et al.—2012 [104]  | New Zealand white rabbit (n = 4)                                                 | Animal study           | The wounds were treated with control or gNO-producing patches designed to produce gNO levels. Wounds are not infected (1. and 2. rabbit) or infected (3. and 4. rabbit). Wounds are treated with placebo (1. and 3. rabbit) or with gNO producing patches (2. and 4. rabbit) | Histological analysis showed improved wound healing in gNO-producing patch-treated rabbits |
| 5           | 5      | Stefia et al.—2020 [105] | C57BL/6 wild type wounded mice (n = 30)                                           | Randomized controlled trial in mice | Mice were wounded and divided into 3 groups (n = 10/group); receiving topical applications Pluronic gel containing either vehicle alone or the supernatant fractions prepared from *F. prausnitzii* strains A2-165 or AHMP21 | Probiotic can regulate wound inflammation and accelerate wound closure                         |
| 6           | 6      | Kusumaningsih et al.—2021 [106] | Male Wistar rats (n = 36)                                                        | Animal study           | Rats were wounded and divided into 6 groups (n = 6/group); (1) a control group over 3 days, (2) a group that used distilled water over 7 days, (3) a group that underwent topical treatment over 3 days, (4) a group that used probiotic (*L. casei*) administered topically over 7 days, (5) a group that underwent systemic treatment over 3 days (6) a group that took oral probiotics for the traumatic ulcers over 7 days | Significant differences were observed in the number of fibroblasts and blood vessels          |

### ANIMAL MODEL—PROBIOTICS SUPPLEMENTATION AND TOPICAL APPLICATION

| No. of Study | Study  | Author            | Patients (Number)                                                                 | Type of Study          | Intervention                                                                                           | Results                                                                                       |
|-------------|--------|-------------------|----------------------------------------------------------------------------------|------------------------|--------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|
| 3           | 3      | Venosi et al.—2019 [103] | 83-year-old woman with a critical limb ischemia and an infected difficult-to-treat ulcerated cutaneous lesion of the right leg | Case report            | Mixture of probiotic bacteria (lyophilized powder sachets, containing *Lactobacillus plantarum*, *Lactobacillus acidophilus* and *Str. thermophilus*) against different bacteria species *K. pneumonia*, *P. mirabilis* and *E. faecalis* | Treatment was effective against the three bacteria species                                      |
| 4           | 4      | Jones et al.—2012 [104]  | New Zealand white rabbit (n = 4)                                                 | Animal study           | The wounds were treated with control or gNO-producing patches designed to produce gNO levels. Wounds are not infected (1. and 2. rabbit) or infected (3. and 4. rabbit). Wounds are treated with placebo (1. and 3. rabbit) or with gNO producing patches (2. and 4. rabbit) | Histological analysis showed improved wound healing in gNO-producing patch-treated rabbits |
| 5           | 5      | Stefia et al.—2020 [105] | C57BL/6 wild type wounded mice (n = 30)                                           | Randomized controlled trial in mice | Mice were wounded and divided into 3 groups (n = 10/group); receiving topical applications Pluronic gel containing either vehicle alone or the supernatant fractions prepared from *F. prausnitzii* strains A2-165 or AHMP21 | Probiotic can regulate wound inflammation and accelerate wound closure                         |
| 6           | 6      | Kusumaningsih et al.—2021 [106] | Male Wistar rats (n = 36)                                                        | Animal study           | Rats were wounded and divided into 6 groups (n = 6/group); (1) a control group over 3 days, (2) a group that used distilled water over 7 days, (3) a group that underwent topical treatment over 3 days, (4) a group that used probiotic (*L. casei*) administered topically over 7 days, (5) a group that underwent systemic treatment over 3 days (6) a group that took oral probiotics for the traumatic ulcers over 7 days | Significant differences were observed in the number of fibroblasts and blood vessels          |

### IN VITRO STUDIES—PROBIOTICS APPLICATION

| No. of Study | Study  | Author            | Patients (Number)                                                                 | Type of Study          | Intervention                                                                                           | Results                                                                                       |
|-------------|--------|-------------------|----------------------------------------------------------------------------------|------------------------|--------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|
| 7           | 7      | Vågesjö et al.—2018 [107] | Human skin wound model/mice                                                      | In vitro model of wound reepithelialization | Wounds were treated daily with saline solution, control *Lactobacillus reuteri* or CXCL12-expressing *L. reuteri* or *L. lacti* | Promising therapeutic approach for non-healing wounds                                          |
| 8           | 8      | Coman et al.—2020 [108] | Pathogenic bacteria were isolated from chronic ulcerative lesions               | In vitro study         | To evaluate probiotic efficacy of SYNBIIO (1:1 combination of *L. rhamnosus IMC 501* and *L paracasei IMC 502*) in wound infections | Good antimicrobial capacity and adhesion percentage to human keratinocyte cells and fibroblasts |

Abbreviations: *B*. *Bifidobacterium*; CRP, C-reactive protein; *E.* *Enterococcus*; *F.* *Faecalibacterium*; gaseous nitric oxide, gNO; *K.* *Klebsiella*; *L.* *Lactobacillus*; *P.* *Proteus*; *Str.* *Streptococcus*. 

Table 7. Cont.
Table 8. Probiotic treatment in SD.

| No. of Study | Author | Patient Population (Number) | Type of Study | Intervention | Results |
|--------------|--------|-----------------------------|---------------|--------------|---------|
| 1            | Guéniche et al.—2008 [109] | Volunteers aged 6 to 70 years suffering from SD (n = 60) | Prospective, double-blind, placebo-controlled | A cream containing a 5% lysate of the nonpathogenic bacteria *V. filiformis* (n = 30) or a vehicle cream applied once daily for 4 weeks (n = 30) | Significant improvement of SD |
| 2            | Reygagne et al.—2017 [110] | Male volunteers aged 18 to 60 years with moderate-to-severe dandruff (n = 60) | Randomized, placebo-controlled study | A sachet containing ST11 (n = 30) or a placebo (n = 30) administered orally for 56 days | Significantly reduced severity of dandruff |

Abbreviations: SD, seborrheic dermatitis; ST11, *Lactobacillus paracasei* ST11; *V. filiformis*, *Vitreoscilla filiformis*.

Table 9. Probiotic application of burns in humans and animal models.

| No. of Study | Author | Patient Population (Number) | Type of Study | Intervention | Results |
|--------------|--------|-----------------------------|---------------|--------------|---------|
| **HUMAN MODEL—PROBIOTICS ADMINISTERED ORALLY OR TOPICALLY** |
| 1            | Peral et al.—2009 [111] | Patients with second and third-degree burns (n = 80) | Case-control study | Patients were separated into 2 groups: in the topical probiotic group patients (n = 38) received *L. plantarum* ATCC 10241. In the control group patients (n = 42) received 1% SD-Ag cream for 10 days | Topical probiotic treatment of 2nd degree burn patients was as effective as SD-Ag decreasing pathogen load |
| 2            | Stefanatou et al.—[112] | 34-year-old woman suffering from extensive deep-partial and full thickness thermal burns | Case report | *S. boulardii* administered for nearly 2 months | Probiotic sepsis due to fungaemia in a critically ill burn patient which resulted in death |
| 3            | Thomson et al.—2012 [113] | 47-year-old lady with 54% deep-dermal and full-thickness flame burns to her neck, chest, upper abdomen and upper limbs | Case report | Oral administration of *L. casei shirota* for 2 weeks after infection which occurred 5 months after burn | Pathogen of the wound changed from multidrug resistant to multidrug sensitive strain |
| 4            | Mayes T et al.—2015 [114] | Less than 22 years old acutely burned patients, and were admitted/consented within 10 days of burn injury (n = 20) | Randomized, double-blind, placebo-controlled | The treatment group received LGG (n = 10). The control group received placebo (n = 10). Investigational products were administered via nasoduodenal feeding tube twice daily | Improved gastrointestinal outcomes and reduced time to wound healing |
Table 9, Cont.

| No. of Study | Author                          | Patient Population (Number)                                                                 | Type of Study                          | Intervention                                                                                           | Results                                                                 |
|--------------|---------------------------------|-------------------------------------------------------------------------------------------|----------------------------------------|--------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------|
| 5            | El-Ghazely et al.—2016 [115]    | Thermally-injured pediatric patients with total body surface burns between 20-50% and depth between 5-10% (n = 40) | Randomized, double-blinded, controlled trial | Participants were separated into 2 groups; probiotic group (n = 20) received probiotic preparations of *L. fermentum* and *L. delbrueckii* and placebo control group (n = 20) | Decreased infection incidence in the probiotic group                   |
| 6            | Perdanakusuma et al.—2019 [116] | Burn patients (n = 16)                                                                    | Randomized, placebo-controlled trial   | Patients were randomized into three treatment groups. Oral administration of either a placebo, a *L. reuteri* probiotic, or a *B. infantis* 35624 probiotic for 14 consecutive days | *B. infantis* 35624 single-strain probiotic was not significantly superior to *L. reuteri protectis* in altering intestinal immunity after burns |
| 7            | Fleming et al.—2019 [117]       | Burn patients aged 18 to 89, who were hospitalized for at least 2 weeks, no formal protocol of antibiotics use was established (n = 108) | Retrospective study                    | Oral administration of >1 million colony-forming units per day of *L. acidophilus* and *L. rhamnosus* | No improvements in patient outcomes but increased incidence of diarrhea |

**ANIMAL MODEL—PROBIOTICS ADMINISTERED ORALLY**

| No. of Study | Author                          | Patient Population (Number)                                                                 | Type of Study            | Intervention                                                                                           | Results                                                                 |
|--------------|---------------------------------|-------------------------------------------------------------------------------------------|--------------------------|--------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------|
| 8            | Herek et al.—2004 [118]         | Male albino rats (n = 23)                                                                  | Animal study             | The rats were divided into sham burn group (n = 7), burn + Ampicillin-sulbactam group (n = 8), burn + Ampicillin-sulbactam + probiotic (*S. boulardii*) group (n = 8) administered twice daily for 5 days | Decreased incidence of antibiotic-induced bacterial translocation      |
| 9            | Gong et al.—2017 [119]          | Healthy male Wistar rats (n = 60)                                                         | Animal study             | The rats were divided into groups: burn model group (n = 15)—normal saline; glutamine treatment group (n = 15)—glutamine + normal saline; probiotics treatment group (n = 15)—probiotic + normal saline; glutamine and probiotics combined treatment group (n = 15)—lutamine + normal saline. All were administered once daily for 7 days | Glutamine and probiotics together significantly inhibited nitric oxide (NO) content and reduced levels of the inflammatory factors |

**ANIMAL MODEL—PROBIOTICS ADMINISTERED TOPICALLY OR LOCALLY**

| No. of Study | Author                          | Patient Population (Number)                                                                 | Type of Study         | Intervention                                                                                           | Results                                                                 |
|--------------|---------------------------------|-------------------------------------------------------------------------------------------|-----------------------|--------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------|
| 10           | Valdez et al.—2005 [120]        | Adult inbred BALB/c mice                                                                  | Animal study          | *L. plantarum* ATCC 10241 injection into burned area on 3, 4, 5, 7 and 9 days                         | Samples from skin, liver and spleen taken after 5, 10 and 15 days demonstrated inhibition of *P. aeruginosa* colonisation |
Table 9. Cont.

| No. of Study | Author                        | Patient Population (Number) | Type of Study | Intervention                                                                                                                                                                                                 | Results                                           |
|--------------|-------------------------------|-----------------------------|---------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------|
| 11           | Brachkova et al.—2011 [121]   | Male Wistar rats (n = 25)   | Animal study  | Rats were randomly allocated into groups: non-burned control rats (n = 2); burned control rats (n = 6); burned skin covered with films containing *L. plantarum* (n = 3); burned skin onto with a suspension of *P. aeruginosa* (n = 7); burned skin, contaminated with *P. aeruginosa*, and covered with films containing *L. plantarum* ATCC 8014(7) | Reduction of pathogen load                       |
| 12           | Argenta et al.—2016 [122]     | Female C57 BL/b mice (n = 38)| Animal study  | The mice were divided into groups. Injured sites were treated with vehicle (burn wound control), probiotics (*L. plantarum* ATCC 1024 only, pathogenic bacteria (*P. aeruginosa*) only, or probiotics + pathogen (*Lactobacillus* and *P. aeruginosa*) for five days | Lower mortality rate and inhibition of pathogenic bacteria |
| 13           | Satish et al.—2017 [123]      | Male Dutch Belted rabbits  | Animal study  | Each rabbit had four burn wounds created on its dorsum—the four burn wound conditions therefore were: (1) Burn wound only; (2) *L. plantarum* ATCC 10241 only; (3) *P. aeruginosa* only; (4) *L. plantarum* + *P. aeruginosa* | Curtailed severity and length of infection, reduced scarring |
| 14           | Sürmeli et al.—2019 [124]     | Rats (n = 35)                | Animal study  | Rats were divided into groups (n = 7/group): control group; *L. plantarum* applied immediately after the burn and then MRSA inoculated; MRSA applied immediately after the burn and then *L. plantarum* inoculated; control of *L. plantarum*; control of MRSA | Probiotic showed protective role in non-infected burn wounds |
| 15           | Khan et al.—2019 [125]        | Male BALB/c mice (n = 30)   | Animal study  | The mice were randomized into negative (untreated), positive (silver sulfadiazine cream), vehicle (biodispersion and nanoscaffold), and experimental bioscaffold groups (n = 6/group). Treatments were applied locally on 2, 6, 10, and 14 days postburn—application of probiotic (*E. mundtii* QAUEM2808) | Accelerated epithelialization, collagen deposition, and hair follicle formation and inhibit pathogens |
### Table 9. Cont.

| No. of Study | Author                | Patient Population (Number) | Type of Study                          | Intervention                                                                 | Results                                           |
|--------------|-----------------------|-----------------------------|----------------------------------------|------------------------------------------------------------------------------|--------------------------------------------------|
| 16           | Olguin et al.—2005    | Burn patients (n = 21)      | Randomized, double-blind, placebo-controlled | 6 g of oligofructose (study group) or sucrose as placebo (control group) during 15 days | No effect on gastrointestinal permeability        |

**Abbreviations:** B., Bifidobacterium; E., Enterococcus; L., Lactobacillus; LGG, Lactobacillus rhamnosus GG; MRSA, Methicillin-resistant Staphylococcus aureus; P., Pseudomonas; S., Saccharomyces; SD-Ag, Silversulphadiazine.

### Table 10. Pre- and probiotic treatment of acne in humans, animal models and in vitro studies.

| No. of Study | Author                | Patient Population (Number) | Type of Study                          | Intervention                                                                 | Results                                           |
|--------------|-----------------------|-----------------------------|----------------------------------------|------------------------------------------------------------------------------|--------------------------------------------------|
| 1            | Kim et al.—2010 [35]  | Patients with acne (n = 36) | Randomized, double-blind, placebo-controlled | Fermented milk with lactoferrin daily (n = 18) or fermented milk only (n = 18) for 12 weeks supplemented orally | Improvement of acne with a selective decrease of triacylglycerols in skin surface lipids |
| 2            | Fabroccini et al.—2016 [127] | Patients with acne (n = 20) | Placebo–controlled trial              | Over a 12-week period, the probiotic group (n = 10) consumed a liquid supplement containing LSP1, placebo group (n = 10) | Normalised skin expression of genes involved in insulin signalling and improvement of acne |
| 3            | Dall’Oglio et al.—2018 [128] | Female patients with mild to moderate acne (n = 12) | Proof of concept pilot trial          | Prebiotic oral supplementation with food supplement containing FOS and GOS for 3 months | Positive effects on glycemic and lipid metabolic parameters |
| 4            | Rahmayani et al.—2019 [129] | Patients with acne aged between 17 and 25 years old (n = 33) | Pre-experimental clinical study with a pretest-posttest design | Oral mix of probiotics was given to individuals for 30 days—B. lactis W51, B. lactis W52, L. acidophilus W55, L. casei W56, L. W57, L. lactis W58 | Elevated serum IL-10 levels |
| No. of Study | Author | Patient Population (Number) | Type of Study | Intervention | Results |
|--------------|--------|-----------------------------|---------------|--------------|---------|
| 5            | Kang et al.—2009 [130] | 12 years of age or older patients with acne (n = 70) | Double-blind, randomized, placebo-controlled trial | E. faecalis SL-5 lotion (n = 35) or placebo lotion (n = 35) to apply twice a day for 8 weeks | Reduced number of inflammatory lesions |
| 6            | AOBiome LLC.—2019 [131] | Adult patients with mild to moderate acne (n = 388) | Double-blind, randomized, placebo-controlled trial | Probiotic (N. eutropha) or placebo spray to saturate the entire face in the morning and at night for 12 weeks | 2-point reduction in IGA of acne severity compared to vehicle control |
| 7            | De Los Angeles Mosquera Tayupanta et al.—2019 [132] | Patients from 15 to 20 years old, with previous diagnosis of type II acne (n = 20) | Interventional study | First the evaluation of the in vitro antagonistic effect of L. acidophilus against C. acnes was performed, then topical application | Decrease in the population of C. acnes |

**IN VITRO STUDIES**

| No. of Study | Author | Patient Population (Number) | Type of Study | Intervention | Results |
|--------------|--------|-----------------------------|---------------|--------------|---------|
| 8            | Al-Ghazzewi et al.—2010 [133] | - | In vitro study | The symbiotic ability of probiotic bacteria and konjac glucomannan hydrolysates to inhibit acne-inducing bacterium, C. acnes growth was studied in vitro | Inhibition of the growth of C. acnes, which was significantly enhanced by the presence of prebiotic |
| 9            | Kang et al.—2012 [134] | - | In vitro study | Study examined the effects of L. reuteri strains (KCTC 3594, KCTC 3678, KCTC 3679) on the proliferation of C. acnes and S. epidermidis | Control of the growth of bacteria involved in acne inflammation and prevent acne |
| 10           | Lee et al.—2012 [135] | - | In vitro study | Activity of Bifidobacterium spp. against C. acnes KCTC3320 using the co-culture method was investigated | Bifidobacterium spp. could be used as an effective treatment and reduced the risk of acne development |
| 11           | Khalfallah et al.—2021 [136] | - | In vitro study | Two type of Str. salivarius strains and one L. plantarum were tested for production of antimicrobials-target organisms used were C. acnes, S. aureus, and P. aeruginosa | Probiotic containing could be topically applied without the need for a regular antibiotic treatment or as an adjunctive therapy |

Abbreviations: B., Bifidobacterium; BMI, body mass index; C., Cutibacterium; E., Enterococcus; FOS, fructo-oligosaccharides; GOS, galacto-oligosaccharides; IL-10, interleukin 10; LSP1, Lactobacillus rhamnosus SP1; L., Lactobacillus; N., Nitrosomonas; S., Staphylococcus; Str., Streptococcus; P., Pseudomonas; spp., species.
4. Discussion

4.1. Atopic Dermatitis

AD is a common, chronic inflammatory skin disease, affecting almost 3% of adults and up to 10–20% of the child population, with an increasing prevalence. The onset usually occurs during the first year of life. AD is characterized by dry skin, pruritus and recurrent eczematous lesions. The severity of AD may be assessed by SCORAD (scoring atopic dermatitis) severity score [137]. AD is often associated with other atopic diseases: allergic rhinitis and asthma [138]. The skin and gut microbiome in adult AD patients is affected among others by maternal diet during pregnancy, the mode of delivery, antibiotics taken during pregnancy and in infancy, chronic exposure to allergens [139]. It is estimated that in over 90% of cases both lesional and non-lesional skin of the patients is colonised with S. aureus in AD, compared with less than 5% of healthy individuals. Moreover, in the affected areas, the abundance of S. aureus was associated with disease severity [140]. Increase in fungal diversity and the presence of unique anaerobic bacterial species such as Clostridium and Serratia spp. was also found on the skin of AD patients [13].

Prevention of the development of atopic dermatitis using probiotics. Eight out of 21 studies reported a decreased occurrence of AD in the probiotics group. Most studies investigated L. rhamnosus GG (LGG). The positive impact of probiotics has been proven by Kalliomaki et al. According to the results, AD is diagnosed in 46 of 132 (35%) children aged two years, with the frequency of AD in the probiotic group (LGG) half that of the placebo group (15/64 [23%] vs. 31/68 [46%]) [25]. Wickens et al. in their study examined the L. rhamnosus HN001 (HN001) and HN019 influence on AD, founding the probiotic group with significantly lower cumulative prevalence of eczema and skin prick test sensitization [29]. Another study by Wickens et al. proved that mother and child intervention with HN001 probiotic supplementation was associated with a reduction in eczema and SCORAD. Note that maternal-only HN001 supplementation did not significantly reduce the prevalence of eczema in the infant by 12 months [43]. In 2006, Rautava et al. examined the use of LGG and Bifidobacterium lactis Bb-12 (Bb-12) in a 12-month follow-up trial. AD developed in 4/32 (13%) of the infants receiving probiotics and 8/40 (20%) of those receiving placebo [26]. Six years later, Rautava et al. confirmed the impact of daily probiotics intake (either the combination of L. rhamnosus LPR and B. longum BL999 (BL999) or the combination of L. Paracasei ST11 (ST11) and BL999), showing the risk of developing AD during the first 24 months of life significantly reduced in infants of mothers receiving probiotics [38]. Schmidt et al. carried out another study involving the supplementation of LGG and Bb-12, resulting in lower incidence of AD (4.2%) in the probiotic versus placebo group (11.5%) [45] Kim et al. proved that in the probiotic group the occurrence of AD was significantly reduced compared to the placebo group at 12 months of age (36.4% vs. 62.9%) [141]. Lau et al. showed that Escherichia coli and Enterococcus faecalis significantly reduced the incidence of AD development in the subgroup of high risk infants. Ten percent (15/154) of infants in the active group developed AD compared to 19% in the placebo group. This was more pronounced in the group of infants with paternal heredity for atopy (11% vs. 32%) [40]. The study of West et al. investigated the use of L. paracasei F-19 and found that the cumulative incidence of eczema at 13 months of age was 9/84 in the probiotic and 19/87 in the placebo groups [32].

However, 13 out of 21 trials showed that the administration of probiotics had no impact on prevalence of AD. Studies by Allen et al., Dotterud et al., Huurre et al. and Plummer et al. showed that a mix of bacterial strains was given and revealed a similar frequency of diagnosed AD both in the study and control groups. They also showed no effect of the use of probiotics in pregnant mother and infants to avoid the development of AD [30,36,41,44]. The study by Niers et al. provided interesting results in which parental-reported eczema during the first three months of life was significantly lower in the intervention group compared with placebo, 6/50 vs. 15/52. After three months, the incidence of AD was similar in both groups [33]. The use of probiotic (L. reuteri) in the group of pregnant women
and infants was evaluated by Abrahamsson et al. Despite the cumulative incidence of AD was similar in the \textit{L. reuteri} and the placebo groups (36\% vs. 34\%), IgE-associated eczema was less common in the \textit{L. reuteri} group, although the difference was only statistically significant during the second year of life (8\% vs. 20\%) [27]. Conclusions from the studies by Boyle et al., Cabana et al., Kopp et al. and Ou et al. evaluating the effect of LGG on pregnant mothers showed that there was no difference between the probiotic group and placebo in the appearance of AD among the infants [31,37,39,42]. Soh et al. examined the incidence of AD in infants receiving probiotics (\textit{B. longi} and \textit{L. rhamnosus}). The incidence of eczema in the probiotic group was similar to that in the placebo group (22\% vs. 25\%). The median SCORAD at 12 months was 17.10 in the probiotic group and 11.60 in the placebo group [34].

**Prebiotics in the prevention of atopic dermatitis.** The number of studies on prebiotics in the prevention of AD is limited and they present inconsistent results. Studies investigated only a few prebiotic compounds: combination of galacto-oligosaccharide (GOS) and fructo-oligosaccharide (FOS), acidic oligosaccharides, polydextrose (PDX), different content of lactose, oligofructose plus inulin. Among the nine studies included in this review, five have shown the positive effect of prebiotics in the prevention of the development of eczema. The rest of the studies showed no significant differences in group of infants fed with or without prebiotics.

Positive effects of the administration of prebiotics has been shown by Ziegler et al., who investigated the administration of a GOS and PDX mix and found a statistical difference in the occurrence of eczema (prebiotics vs. control: 18 vs. 7\%) [47]. The same combination was used by Pontes et al. in their study, who reported a lower number of allergic diseases including AD in the analyzed group receiving prebiotics [53]. Three trials investigating the relationship between GOS supplementation and preventing eczema showed a decreased risk of developing AD [46,48,49]. However, Gruber et al. found that a formula containing a mixture of neutral oligosaccharides can be also effective in prevention of AD [50]. Wopereis et al. also presented a beneficial impact in the prevention of AD and modulation of gut microbiota by using a partially hydrolyzed formula containing short-chain GOS and long-chain FOS and pectin-derived acidic oligosaccharides [56]. No differences between prebiotic groups and control groups have been found in four studies. Two of them investigated a mix of FOS and GOS [51,52]. Ranucci et al. used a mixture of GOS and PDX in their trial. There were no significant differences in the cumulative incidence, intensity and duration of AD among the investigated groups of patients [55]. A study by Boyle et al. on prebiotic containing FOS has shown that prebiotics did not prevent AD in high-risk infants during the first 12 months of life [54].

**Role of probiotics and prebiotics in AD treatment.** 20 of 27 studies on probiotics revealed improving SCORAD in AD patients compared to placebo. One of the first studies on probiotic treatment in AD, that found using probiotics may have positive impact on the course of AD, was the study by Isolauri et al. The aim of their study was to evaluate the effects of probiotics use with Bb-12 or LGG on infants with AD. The results showed that by using probiotics, the skin condition improves. SCORAD decreased in the Bb-12 group to 0, and in the LGG group to 1 versus the SCORAD of 13.4 in the placebo group [58]. Drago et al. evaluated the influence of \textit{S. thermophilus} ST10 and tara gum on the SCORAD score. The score decreased significantly in the probiotic group after one month and the index was significantly lower in the probiotic group than in the placebo group [92]. Ivakhnenko et al. evaluated both the use of Bb-12 and \textit{Streptococcus thermophilus} for 4 weeks. The results showed significant improvement of SCORAD in the probiotic group compared to placebo [73]. Wu et al. proofed that the SCORAD index declined from baseline after two months in the LGG group [76]. Brouwer, Folster-Holst and Kirjavainen showed similar effects of this LGG bacteria in AD patients [59,63,64]. Studies which resulted in a significant decrease in the SCORAD index in AD patients by using probiotics containing single strains or combined bacterial strains, including \textit{L. salivarius} LS01, were also those conducted
by Drago et al. [89] and Iemoli et al., [90] *L. acidophilus* DDS-1—by Gerasimov et al. [80],
*L. plantarum*—by Han et al. [82], *L. fermentum*—by Weston et al. [61] and *L. sakei*—by
Woo et al. [81] Yang et al. randomly assigned their patients to the probiotic-receiving groups
(*L. casei, L. rhamnosus, L. plantarum, and B. lactis*) or placebo groups for six weeks. The
result of their trial was a significant clinical improvement in the skin condition among the
probiotic groups [85]. Two of the studies also proved a positive impact on the SCORAD
score by using probiotics with *B. breve*. Taniuchi et al. and Yoshida et al. showed a
significant improvement in skin conditions during the study in the probiotic group [62,88].
Three studies assessing the impact of using bacteria mix in probiotic groups revealed a
significant improvement in SCORAD scores [84,86,87].

Seven out of 27 studies in children showed no significant differences in SCORAD
scores between the probiotic and placebo groups after treatment. Lin et al. proved that the
SCORAD index was not significantly reduced in the *B. bifidum* group versus controls [74].
Grüber et al. suggested that AD improved after four weeks of supplementation (LGG vs.
placebo); however, the difference was not significant [65]. Similar results were obtained by
Viljanen et al.: the SCORAD score decreased by 65%, but with no statistically significant
differences between treatment groups [60]. Sistek et al. evaluated the role of *L. rhamnosus*
and *B. lactic*. Their findings stated that there is no significant difference between probi-
otic and placebo groups [78]. Rosenfeldt et al. examined probiotic *Lactobacillus* strains
(lyophilized *L. rhamnosus* 19070-2 and *L. reuteri* DSM 122460) in combination for six weeks
in 1- to 13-year-old children with AD. The total SCORAD index in this trial did not change
significantly [77]. The results of the study by Göbel et al. study on *L. acidophilus* and *B. lactis*
Bi-07 (Bi-07) were that there was no benefit for the probiotics on the severity of AD. How-
ever, a post hoc analysis showed a significant reduction in severity of AD in the Bi-07 group
and possible positive effects of this probiotic strain could be of further interest [68]. Gore
et al. in their trail compared the effects of using *B. lactis* and *L. paracasei*. No significant
differences were observed between the groups after 12-week treatment-period [71].

Five out of 36 studies did not estimate the SCORAD score, but evaluated other factors,
such as puritus. Matsumo et al. in their study found that *Bifidobacterium animalis* subsp.*lactic* LKM512 may reduce pruritus by increasing expression of metabolite kynurenic
acid [91]. The results obtained by Majama et al. suggest that probiotic bacteria may
improve endogenous barrier mechanisms in patients with AD and those with food allergies
due to breast feeding and may be useful in AD treatment [57]. Studies by Flinterman et al. [66], as well as by Guo et al., suggested that in vitro IgE production is decreased in the probiotic group compared to placebo [75]. Nermes et al. found that
the levels of IgA and IgM-sectreting cells decreased significantly in the probiotic group
compared to placebo. The baseline-adjusted ratios for treated to untreated patients after
one month were 0.59 for IgA- and 0.53 for IgM-secreting cells [69].

**Synbiotics.** Five publications on the use of synbiotics were found; however, only one
of them by Farid et al. reported a significant reduction of the SCORAD score [70]. Passeron
et al. compared the effects of probiotics (*L. rhamnosus* Lcr35) and synbiotics in children
over two years old. The study showed no statistical differences regarding SCORAD scores
between the two groups [79]. Shafiei et al. showed that there is no significant difference in
the mean decrease of total SCORAD between placebo (22.3) and synbiotic groups (24.2) [72].
A significantly greater SCORAD score improvement was found in the symbiotic group
of infants with IgE-associated AD by van der Aa et al. [67] Wu et al. also found that a
combination of *L. salivarius* and FOS resulted in lower SCORAD in a comparison with the
control [83].

The role of microbiome composition in allergic diseases is well-known, with lower bio-
diversity found as a factor inducing their development. Modulating the microbiome with
probiotics balances the gut microflora, protects the function of intestinal barrier and lowers
the level of pro-inflammatory cytokines produced. Probiotics also influence Toll-like recep-
tors, which play an important role in T-cell differentiation and the development of allergic
reactions. As skin colonization with *S. aureus* plays an important role in AD, a promising new perspective of displacing it with more desirable species is also considered [93].

4.2. Psoriasis

Psoriasis is a common inflammatory disease that affects around 2–3% of the population [142]. It manifests with papulosquamous skin lesions with variable distribution and severity [143]. The pathogenesis of the disease is not yet fully elucidated. However, it is known that genetic, immunological and environmental factors may act as triggering factors, making the keratinocytes start secreting pro-inflammatory cytokines [14]. In the skin lesions, increased abundance of *Streptococcus* spp., *Corynebacterium* spp., *Cutibacterium* spp., *Staphylococcus* spp., *Finegoldia* spp. and *Neisseria* spp. can be found. The biodiversity of microbiota is generally decreased in moderate-to-severe psoriatic patients in contrary to mild psoriatic patients [5].

Based on the reported alterations in gut microbiome, attempts were made to use probiotics and prebiotics in the treatment of psoriasis. Two original studies, one case report and four mice studies were published. In the case report described by Vijayashankar and Raghunath, a supplementation with *L. sporogenes* for 15 days alleviated the symptoms accompanying the sudden onset of generalised pustular psoriasis in a 47-year-old female [94]. Groeger et al. demonstrated a significant decrease in serum CRP, TNF-α levels in psoriatic patients administered with *B. infantis* 35264 for 8 weeks [95]. In the study by Navarro-Lopez et al., supplementation with *B. longum* CECT 7347, *B. lactis* CECT 8145 and *L. rhamnosus* CECT 8361 for 12 weeks resulted in a significant reduction in PASI scores [96]. In three out of four mice studies, the probiotics were administered orally, while in one study it was administered topically. In all studies, the psoriasis-like skin inflammation was induced by topical application of imiquimod. Chen et al. found that in administering *L. pentosus* GMNL-77, both for five or seven days, causes a reduction in erythroematous scaling lesions, decreases TNF-α, IL-6, IL-23, IL-17A/F and IL-22 levels in the skin, decreases spleen weight and reduces the number of IL-17- and IL-22-producing CD4+ T cells in the spleen [97]. In the study by Lu et al., seven different groups of six mice each were given different strains of probiotics. *B. adolescentis* CCFM667, *B. breve* CCFM1078, *Lactocaseibacillus paracasei* CCFM1074, and *Limosilactobacillus reuteri* CCFM1132 ameliorated psoriasis-like pathological characteristics and suppressed the release of IL-23/T helper cell 17 (Th17) axis-related inflammatory cytokines. On the contrary, *B. animalis* CCFM1148, *L. paracasei* CCFM1147 and *L. reuteri* CCFM1040 neither alleviated the pathological characteristics nor reduced the levels of inflammatory cytokines [99]. Ogawa et al. showed that administering *Leuconostoc mesenteroides* NTM048 to imiquimod-induced mice suppressed erythema, scaling, upregulated IL-17 production, increased the levels of plasma deoxycholic acid and altered the faecal microbiota composition. Changes in the gut microbiome were indicated by the increased abundance of *Akkermansia* and a decreased abundance of *Staphylococcus* and *Streptococcus* [100]. The only study concerning a topical application of probiotics was conducted by Rather et al. Application of ethanol extract (SEL001) isolated from *L. sakei* proBio-65 resulted in an inhibition of the imiquimod-induced changes in the skin, as well as decreased IL-19, IL-17A and IL-23 levels [98]. It was shown that the gut microbiome plays an important role in the pathogenesis of psoriasis—patients suffering from this disease present with an increased amount of *Bacteroidetes* and decreased levels of *Firmicutes*, *Proteobacteria* and *Actinobacteria*, probably altering the intestinal barrier integrity, T-cell response and population-type balance, chemotaxis along with carbohydrate, cobalamin, and iron metabolism [144].

4.3. Chronic Ulcers

The use of probiotics as a novel treatment for diabetic foot ulcers (DFU) was first published in 2014. It was suggested that the application of probiotic agents would enable the healing of diabetic ulcers and would prevent diabetic foot infections by activating Toll-like receptors and producing β-defensins, which stimulate skin immune functions [145].
Mohseni et al. investigated the advantages of probiotics in patients with DFU. After the 12-week intervention of probiotic supplementation (L. acidophilus, L. casei, L. fermentum, B. bifidum), it had beneficial effects on the DFU size. It also decreased the serum total cholesterol and CRP and increased plasma nitric oxide (NO) and total plasma antioxidant capacity [102].

Most research was carried out using in vitro models, e.g., the effectiveness of a probiotic based on L. rhamnosus and L. paracasei strains in a 1:1 ratio against microorganisms previously isolated from chronic ulcerative lesions. Following the administration of probiotics, the growth of bacteria, compared to the control, was lower in the case of such bacteria as P. aeruginosa, C. striatum, A. baumanii, S. aureus, P. mirabilis in 75%, in the case of Candida parapsilosis in 93.75%, while in the case of E. faecalis 18.75%, and 50% for the mixed flora of the mentioned pathogens. The ability to co-aggregate all pathogens that could prevent adhesion and invasion was also shown [108]. Kusumaningsih et al. investigated the differences in the number of fibroblast cells and blood vessels after the administration of the probiotic L. casei shirota topically and systemically during the onset of the healing of traumatic ulcers in Wistar rats. The number of fibroblasts and new blood vessels were significantly higher in the two intervention groups in a comparison with the control group [106]. A further study investigated wounded New Zealand rabbits infected with S. aureus and treated with L. fermentum, which secretes gaseous NO. The day after the procedure, treatment with the patch with a probiotic agent started and lasted for 21 or 20 days. Morphometric analysis of the ulcer healing revealed that it was significantly accelerated with this treatment method in both infected and uninfected ischemic wounds [104]. Stefia et al. compared the effects of two different strains of Faecalibacterium prausnitzii (SPA and SPAH) for immune cell activity and wound healing in mice. They found that the presence of these strains in the gut exhibited significantly higher patterns of reepithelialization compared to controls by inhibiting NF-κB activation. It resulted in decreased wound proinflammatory cytokine expression and induced myofibroblast and collagen transitions [105]. In another model, L. reuteri was transformed with a plasmid containing the genetic material of the C-X-C Motif Chemokine Ligand 12 chemokine involved in accelerated wound healing. Additionally, the lactic acid produced by the probiotic bacterium lowered the pH and increased the bioavailability of the chemokine. This strain was applied to wounds in mice, accelerating ulcer healing, epithelialization, and wound closure [107].

The most relevant original work was published in 2010. L. plantarum was used in the treatment of chronic leg ulcers. The probiotic was applied to ulcers in 14 patients with diabetes and 20 non-diabetic patients. After 30 days of follow-up, 90% of the extent of ulceration had resolved in 43% of diabetic patients and 50% of non-diabetic patients. A decrease in CFU of S. aureus, S. epidermidis and P. aeruginosa was also noted. It was found that probiotics disrupt biofilm, regulate IL-8 levels and modulate the immune system [101]. In addition, Venosi et al. reported a case of an old woman who was successfully treated with a topical administration of probiotics for an ischemic and infected (K. pneumoniae, E. faecalis and P. mirabilis) chronic wound. The patient received a mixture of probiotics (L. plantarum, L. acidophilus and S. thermophiles) three times/week [103].

4.4. Seborrheic Dermatitis

Seborrheic dermatitis (SD) characterized by erythematous, scaling plaques on the face, chest and scalp [146,147]. It is assumed that the underlying cause of the disease is the excessive activity of sebaceous glands and concomitant infection with Malassezia spp. [146]. Research indicates an increased number of Malassezia strains in the seborrheic area and a satisfactory therapeutic effect of antifungal formulas [147]. Currently, it seems that SD is the result of the skin’s response to free fatty acids produced by M. furfur, which elicit an inflammatory response from keratinocytes [148,149]. M. furfur also possesses the ability to produce metabolites, which stimulate the aryl hydrocarbon receptor and thus may modulate the function of antigen-presenting cells [150].
There are limited data on the effects of probiotics and the modulation of the cutaneous microbiome on the course of SD. The use of superficial *Vitreoscilla filiformis* preparation in a double-blind study involving 60 patients with SD resulted in a reduction of itching, erythema and scaling. At the cellular and subcellular level, the lysate of these bacteria resulted in an increase in the activity of IL-10 produced by dendritic cells of the skin and an increase in the activity of regulatory T lymphocytes [109]. Another study involving the oral administration of ST11 demonstrated a significant reduction in symptoms, which at the subcellular level was also accompanied by a shift in immune activity consisting, as before, in an increase in IL-10 production [110]. These examples confirm the possible benefits of using both forms of probiotics in the group of patients with SD.

4.5. Burns

The analysis of studies conducted both in animal models and in clinical trials, mostly showed at least partial positive effect of the use of probiotics on the healing of infected wounds by inhibiting microbiome growth, microfilm formation and interbacterial communication [151].

Among the various used bacterial strains, the most evidence exists for *L. plantarum*. Peral et al. established the effectiveness of *L. plantarum* probiotic treatment with a topical application in human patients. *L. plantarum* would compete with bacterial pathogens and would be able to promote tissue repair [111]. El-Ghazzey et al. studied the effect of *L. fermentum* and *L. delbruekii* treatment in pediatric post-burn patients. They conclude that probiotic administration is safe to use and improves wound healing [115]. In a case report, oral application of *L. casei* resulted in the appearance of multi-drug sensitive *P. aeruginosa* instead of an extremely drug-resistant strain [113]. Perdanakusuma et al. demonstrated that *B. infantis* 35624 single-strain probiotics were more effective compared to *Lactobacillus reuteri protectis* in altering intestinal immunity [116].

Valdez et al. has been shown in adult inbred BALB/c mice that *L. plantarum* and/or its by-products could be a potential therapeutic agent for *P. aeruginosa* burn infections [120]. Brachkova et al. found that the application of calcium alginate films containing *L. plantarum* reduced *P. aeruginosa* in a rat model of burns [121]. Argenta et al. proved in mice that probiotic therapy (*L. plantarum*) suppressed the induction of TNF-α, IL-6 and IL-10 in liver and inhibited the accumulation of the pathogen in remote organs [122]. Satish et al. *L. plantarum* as a therapeutic agent alleviates burn wound infection and scarring after burn injury in rabbits [123]. Sürmeli et al. demonstrated that *L. plantarum* has a protective role in non-infected burn wounds against meticillin-resistant *Staphylococcus aureus* (MRSA). Additionally, the therapeutic effect of *L. plantarum* was not shown in MRSA infection [124]. Herek et al. investigated that the *Saccharomyces boulardii* could effectively decrease the incidence of antibiotic-induced bacterial translocation in burned rats [118].

Khan et al. demonstrated the importance of the method of probiotic application in a thermal burn mouse model. The use of the bioskeleton compared to traditional forms of probiotic application resulted in acceleration of epithelialization, collagen production and formation of hair follicles, as well as an inhibitor on the growth of pathogenic bacteria, reducing infection and accelerating wound healing [125]. As a result of burns due to systemic stress, the intestinal barrier is significantly impaired, resulting in inflammation and oxidative stress, leading to the destruction of the intestinal barrier and abnormal intestinal function. The studies on the animal model of burns show that the application of glutamine and probiotics reduced the apoptosis of the intestinal epithelial cells [119].

Fleming et al. performed a retrospective study in connection with preventing potential antibiotic-associated *C. difficile* colitis by giving probiotics to burned patients in a critical condition. Otherwise, they found no significant difference in *C. difficile* infection between the control group and the intervention group [117]. Oglin et al. proposed that the regular intake of prebiotics might help to increase the gastrointestinal permeability in burn patients. Following the application of oligofructose (OF), they found no difference between the control and OF groups [126].
Due to the damaged intestinal barrier and the impaired immune system function caused by burns, there is a potential risk that probiotic bacteria may translocate and ultimately result in infection. Mayes et al. demonstrated the efficacy and safety of probiotics in the pediatric population hospitalized due to skin burns [114]. However, there are known cases of severe infections and probiotic-induced sepsis in critically ill people [112].

4.6. Acne

Acne is a chronic skin disease, affecting the pilosebaceous units, with multifactorial pathogenesis including hormonal influence, the immunological state of the host, diet, deregulation of insulin-like growth factor, excessive sebum production and FoxO1 deficiency [152,153]. Considering the pathogenesis of acne, Cutibacterium acnes has been implicated as an important pathogenic factor. Fitz-Gibbon et al. compared the Cutibacterium strains in patients suffering from acne and healthy individuals, finding remarkable differences [154]. More and more evidence suggests that dysbiosis on the phylotype/strain level leading to a diversity loss is also a major factor in the pathogenesis of acne [155]. The role of the gut microbiome in acne is also raised, as a study conducted in 2018 showed that patients with acne present with lower gut microbiota diversity (abundance of Firmicutes, Clostridium, Clostridiales, Lachnospiraceae, Ruminococcaceae increased Bacteroides levels) [156].

A limited number of studies concerning probiotics and prebiotics use in acne is available. Yet, it is known that the beneficial components of the microflora may ameliorate skin lesions via the suppression of the Treg cell population. In addition, the suppression of B and Th cells due to the modulation of inflammatory cytokine production along with increasing IgA and butyrate secretion may also have an important effect [144]. A clinical trial investigating oral supplementation of L. rhamnosus SP1 (LSP1) had been reported to bring health benefits to the patients such as LSP1 normalized skin expression of genes involved in insulin signalling and an improvement in the appearance of adult acne [127]. However, a mix of B. lactis W51, B. lactis W52, L. acidophilus W55, L. casei W56, L. salivarius W57, and L. lactis W58 was reported to be a trigger for elevated IL-10 serum levels [129]. The results concerning oral prebiotics supplementation remain more consistent, as both lactoferrin as well as GOS and FOS were associated with positive effects [35,128]. Topical application of probiotic-enriched formulas also seem to have a promise: all of the analyzed studies involving the use of E. faecalis SL-5 [130], Nitrosomonas europa [131] or L. acidophilus showed improvement in the skin condition. L. acidophilus was also reported to decrease the population of C. acnes [132]. These findings were also confirmed in in vitro studies: Al-Ghazzewi et al. showed that probiotic bacteria and konjac glucomannan hydrolysates inhibit C. acnes growth [133]. Similar effects were reported by Kang et al., who investigated the properties of L. reuteri on the proliferation of C. acnes and S. epidermidis [134]. Bifidobacterium spp. [135], as well as two S. salivarius strains and one L. plantarum strain, were also reported to show antimicrobial activity in in vitro studies against C. acnes and other pathogens [136].

4.7. Limitations

The present review focuses on a subject that is relatively new and is still not investigated in full detail. One of the major limitations is the small number of publications reporting clinical studies, especially multi-center, double-blinded, placebo-controlled clinical trials. The number of patients in the presented studies were usually low and many studies involved animal models, which cannot be extrapolated to humans. Since the exact pattern composition of a “healthy microbiome” is impossible to establish, there are no objective measures to investigate a universal model. The bacteria used in different studies presented various genera and properties; moreover, they were derived from different sources, often with no exact information on the method of production, storage and other properties. Moreover, the skin diseases presented in the paper were chosen based on their duration and the number of studies available; however, single studies show that the
microbiome modulation, e.g., via a fecal microbiota transplant, may be also effective in the treatment of other dermatological conditions, for example in alopecia areata [157].

5. Conclusions

It can be stated that the microbiome plays an important role in dermatological diseases, at the same time as being an attractive target of therapeutic interaction. This may contribute to the promotion of beneficial (from the point of view of inflammation) activation of the immune system, a reduction of the inflammatory state and, above all, could constitute a physical barrier to the colonization of the skin by pathogenic bacteria. The perspective of treating skin diseases with microbiome modulation via oral and topical probiotics, prebiotics or symbiotics are becoming a part of reality.

There is a growing number of studies into the beneficial effects of probiotics in patients with atopic diseases. It is estimated that the oral application of probiotics or prebiotics during delivery or in the first months of life could delay or alleviate the appearance of AD in infants. From another point of view, probiotics could have the potential to reduce the SCORAD index as a treatment method. On the basis of the available evidence, a recommendation on probiotic intake in order to avoid AD cannot be currently made. Administering probiotics may influence the composition of the gut microbiome, which is more and more often considered to be a factor in the development of psoriasis. The suspected efficacy of probiotics in alleviating the course of psoriasis may be connected to lowering the levels of plasma pro-inflammatory cytokines. Since the data and the amount of research on this topic are limited, it still requires new, randomized, placebo-controlled trials, which would gain an insight into the pathogenesis and novel strategies of psoriasis treatment. There are very limited data available at the moment in the context of chronic ulcers. The positive effects of probiotics were shown mainly in studies focusing on ulcers resulting from diabetes complications. Probiotics may prevent or reduce the infection of burned wounds. Most research has focused on the *L. plantarum* and has showed at least a partial positive effect of the use of probiotics on the healing of infected wounds by inhibiting pathogen growth, microfilm formation and interbacterial communication. Concerning SD and acne, the very limited available data on probiotic administration have showed inconsistent results.

The studies have shown that probiotics and prebiotics both administered orally or applied topically may have a positive influence on the course of skin diseases. Despite the continuous increase in promising data on the effectiveness of the use of probiotics and prebiotics, further clinical trials are needed to assess the efficacy and long-term safety profile of probiotics and prebiotics in the treatment of patients with dermatological diseases.

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