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**Gut microbiota and nutrient interactions with skin in psoriasis: A comprehensive review of animal and human studies**

Damiani G *et al*. Diet and psoriasis

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**Abstract**

The intestinal tract (*i.e.*, the gut), is where the body’s nutrients are absorbed, and is simultaneously inhabited by numerous microbes. An increasing body of literature suggests a crucial role for the gut microbiome in modulating systemic inflammatory disease. Psoriasis is a chronic systemic inflammatory disease and its pathogenesis is related to the interaction between genetic susceptibility, immune response and environmental triggers. The omics era has allowed physicians to assess different aspects of psoriasis pathogenesis such as the microbiome, infectome, and autoinfectome. Furthermore, diet appears to play an important role in modulating disease activity, perhaps by influencing gut microbes. Given these observations, we aimed to summarize the current knowledge regarding skin-microbiome-gut-nutrients and psoriasis.

**Key words**: Gut; Microbiota; Nutrients; Endotypes; Exposome; Psoriasis

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**Core tip:** Psoriasis is a chronic systemic inflammatory disease and its pathogenesis is related to the interaction between genetic susceptibility, immune response and environmental triggers. The omics era has allowed physicians to study psoriasis pathogenesis from different perspectives such as the microbiome, infectome, and autoinfectome. Furthermore, diet appears to play an important role in modulating disease activity. Given these observations, this review aimed to summarize the current knowledge on skin-microbiome-gut-nutrients and psoriasis.

**Introduction**

Psoriasis is alternatively regarded as an inflammatory[1], pruritic[2], autoimmune[3,4], or even autoinflammatory[5] systemic chronic disease affecting not only the skin but the whole body, providing a potential explanation for the numerous comorbidities discovered in relation to this disease[6-12]. The biomarker research performed in the past decades identified substantial inadequacies in describing the wide spectrum of psoriasis[13-15], and failed to explain the complexity of psoriasis endotypes[16]. The genetic background of psoriasis patients reveals that deficiency of the interleukin 1 (IL-1) receptor antagonist and deficiency of the IL-36 receptor antagonist are the only recognized mutations[17], conversely other psoriatic forms are considered complex diseases, due to the intricate interaction between inherited susceptibility alleles and environmental triggers[18]. In addition, epigenetic modifications seem to play a pivotal role in developing psoriasis, establishing environmental triggers as potential modulatory factors[19] as well as in response to anti-psoriatic therapies[20]. These new findings provide a rationale for observed treatment loss-of-response and biologically justify switching to alternative treatments in patient management[21-24]. Furthermore, an increased body of evidence suggests a crucial role for the gut microbiome in modulating psoriasis; linking the skin and gut microbiome[25]. The gut is the site where nutrients are absorbed and at the same time is inhabited by nutrient-modifying microbes. The Skin-Microbiome-Gut-Nutrient interaction is still only partially understood; therefore, this review aimed to summarize the current knowledge in this field.

**Psoriasis, diet and circadian rhythm**

Psoriasis is associated with metabolic and cardiovascular disease[6,26,27]. It has been widely demonstrated that genetic and environmental factors including nutrition, may strongly influence psoriatic pathogenesis and disease progression[28,29]. Increased body mass and a high fat diet may trigger as well as exacerbate psoriasis[29]. Moderate to severe psoriasis is frequently associated with numerous metabolic disorders including obesity, diabetes, dyslipidemia, metabolic syndrome and non-alcoholic fatty liver disease[30]. Fatty acids are increased in obese patients, leading to augmented inflammation and insulin resistance. Furthermore, obesity may also affect drug pharmacokinetics and pharmacodynamics[29]. The option to treat psoriasis patients with phototherapy is affected by body mass index; obese patients require an excessive amount of photosensitizing drug which may lead to toxicities. Obesity is also an important risk factor for psoriasis, given that the relationship between obesity and psoriasis is mutually interdependent[29]. Recent evidence has suggested an addition role of vitamin D in the pathogenesis of several inflammatory skin diseases including psoriasis[31]. An association between low levels of vitamin D and psoriasis has been described[31]. Vitamin D also participates in keratinocyte proliferation and maturation. Nevertheless, the potential value of vitamin D supplementation for psoriasis is still under debate[31]. Dietary antioxidants including omega 3 polyunsaturated fatty acids derived from fish oil, vitamin B12, vitamin D and selenium have the capacity to decrease oxidative stress and consequently lower reactive oxygen species production, and this could be relevant in the pathogenesis of psoriasis. Based on these findings, the Mediterranean diet has been proposed to slow disease progression[32,33]. All of these endogenous factors contribute to the composition of the so-called “exposome”, a measure of the whole complex of endogenous, ingested and not ingested, substances that interact with the body that are capable of perturbing, modifying and modulating vital functions[34]. Exposures capable of modulating psoriasis include tobacco, which increases severity, flare frequency, and even the incidence of the psoriasis[35], alcohol[36] and pollutants[37], even if the role of both alcohol and pollutants in psoriasis is still debatable. Interestingly, the use of marijuana was recently related to secukinumab resistance in a cohort of erythrodermic patients, focusing the attention on addiction screening during intake medical history of psoriatic patients[38]. Anti-psoriatic drugs are also capable of modulating taste, and consequently, the patient’s diet, as recently described for both methotrexate[39] and apremilast[40].

Since the melatoninergic system was discovered in the epidermis, circadian rhythm has been regarded as a possible modulator of inflammation[41], healing[42], aging[43], neuroendocrine[44], and neoplastic conditions[45]. The cyclic nature of sunlight which influences the suprachiasmatic nucleus (central clock) also influences other peripheral tissue, including skin (peripheral clock), in different ways that are slowly beginning to be understood by examining the mechanism(s) of their dysfunction[46,47]. The shift in circadian rhythm may be occasional and transitory, as in the case of jet-lag during intercontinental flights[48], or in the case of the Ramadan fasting months for Muslims[49,50], or more chronic, as in night shift workers[51,52]; however this effect is particularly evident in psoriatic patients. Interestingly, night-shift workers exhibited not only an increase in severity of psoriatic flares, but also an increased incidence of psoriasis, suggesting that shifting the circadian rhythm (*i.e.,* sleep and diet) may be a risk factor for psoriasis[52]. Sunlight, in the form of narrow band UVB (NB-UVB), may also be curative for psoriatic skin, allowing a reprograming of the circadian clock and inhibition of autoimmune phenomena[53], however skin may also marginally adapt to NB-UVB, an outcome termed photoadaptation[54]. Thus, sunlight is considered an integral part of the exposome.

**Psoriasis, nutrients and skin cancer**

The possible influence of nutrients on gene expression and clinical progression or remission of psoriasis is not fully explored and may represent the main challenge in approaching complementary therapy for psoriasis. It is reported that many dietetic factors may exert beneficial effects, while others can aggravate inflammatory and immune networks, thus leading to psoriasis comorbidities[55]. Previous studies have reported the positive effects of low-energy diets, vegetarian diets, formula diet weight loss programs, gluten-free or very low-calorie carbohydrate-free diet. It is believed that certain vitamins (*e.g.*, A, E and C), and oligo-elements (*e.g.*, iron, copper, manganese, zinc, and selenium) are anti-oxidants, leading to a reduction in oxidative stress and decreased production of reactive oxygen species[55]. In fact, the disruption of cell redox signaling and involvement of oxidative stress in the pathogenesis of psoriasis was previously suggested, indicating that the potential therapeutic use of dietary antioxidants in psoriasis may represent a novel complementary strategy[56]. Although limited data exist regarding the role of specific diet regimens in psoriasis, the main goal for clinicians is to reduce cardiac risk factors and obesity-related comorbidities.

Interestingly, psoriatic patients displayed a higher risk of cancer compared to the general population; furthermore, this increase is not fully explained by anti-psoriatic immunosuppressive therapies, so several real-life or ecological studies have suggested an intimate relation with diet[57]. This theory is supported by the evidence that different foods modify microRNA expression in psoriatic patients[28]. The influence of a restrictive caloric diet was documented to be beneficial in relapsing plaque psoriasis, and may also decrease the risk of cancer[58-60]. Some lipid components, such as omega-3 polyunsaturated fatty acids may protect cells against UV-induced DNA damage by increasing the expression of tumor-suppressor protein p53, thus promoting cell cycle arrest and preventing melanoma development[61-63]. Solid cancers in psoriasis have been reported, especially those linked to alcohol and smoking. A higher risk of non-melanoma skin cancers, especially squamous cell carcinoma has been shown, possibly as a result of previous exposure to 8-methoxypsoralen-ultraviolet-A (PUVA), cyclosporin, tumor necrosis factor-inhibitors and/or methotrexate[64-66]. Consideration of malignancy risk associated with individual treatments and personal nutritional phenotype may help clinicians to make optimal therapeutic decisions for individual patients.

**Mouse model of psoriasis and diet**

The "omics" era has provided researchers with new powerful techniques (*e.g.*, metagenomics) and has elucidated a myriad of dysregulated immune responses that are heavily associated with the gut microbiota. Previous research suggested that ingested nutrients heavily affect the body’s microbial composition and community. This has led to experimental approaches in mouse models of psoriasis. In recent years, nutrition and microbial influence has been heavily implicated in psoriasis onset and disease severity. The contribution of nutrients in mouse models has provided valuable insight into human disease regulation. For instance, 12-O-tetradecanoylphorbol-13-acetate (TPA), a known inflammatory signal transducer, can induce psoriasis-like skin lesions in mice, while lesions and proinflammatory cytokine expression were significantly reduced in TPA-induced psoriasis by tangerine-derived nutrient flavonoids: nobiletin (Nob) and 5-hydroxy-6,7,8,3′,4′-pentamethoxyflavone (5-HPMF)[67]. In addition, obesity, a result of poor nutrition, has been shown to exacerbate the severity of psoriasiform dermatitis in imiquimod-induced rodent models[68]. Collectively, these results support the need to elucidate nutritional impact on psoriasis severity. Research has suggested that monounsaturated fatty acid-rich diets, such as the Mediterranean Diet have anti-inflammatory effects and slow the progression of psoriasis in patients[33]. Poor nutrition has been associated with dysregulated metabolic functions and even more so, has been shown to be critical in skin disease metabolic homeostasis compared to healthy individuals. Previous research has shown evidence for biochemical skin barrier restoration through topical administration of solenopsin, a compound of fire ant venom chemically similar to ceramides, and its derivates by reducing inflammatory markers and improving acanthosis in KC-Tie2 mice, an established rodent-model of psoriasis[69].

**Psoriasis and THE microbiome**

Several lines of evidence confirmed the relationship between skin microorganisms and psoriatic lesions, such as *Group A β*-*hemolytic streptococcal infections* linked to guttate psoriasis[70]. Other microorganisms including *Staphylococcus aureus*, *Malassezia* and *Candida albicans* also appear to be involved in psoriasis pathogenesis[71]. The deep inter-relationship between the mycobiome and microbiome seem to act as a disease-modifier in psoriatic patients. Although it is well known that the gut and skin microbiome deeply interact, sparse information is available on the gut and skin mycobiome. Using high-throughput 16S rRNA gene sequencing, Alekseyenko *et al*[72], found thatpsoriatic plaques had an abundance of the following bacteria: *Corynebacterium*, *Propionibacterium*, *Staphylococcus*, and *Streptococcus*. Baker *et al*[73], found that peptidoglycan, a cell wall component of Gram-positive bacteria including *Streptococci* and *Staphylococci*, acts as a T cell activator in psoriasis. The authors observed that dermal papillae and cellular infiltrates of guttate and chronic plaque skin lesions had higher numbers of peptidoglycan-containing cells compared to non-lesional psoriatic skin. Psoriatic dermal *Streptococcal*- and *Staphylococcal*-specific CD4+ T cell lines proliferated and produced IFN-alfa in response to the *respective* peptidoglycan structures. Overall, these results suggest that peptidoglycans may be responsible for T cell activation in psoriasis. Moreover, some studies have linked gut microbiota and psoriasis.

Up to 10% of patients with inflammatory bowel disease (IBD) are diagnosed with psoriasis[74]. Patients with psoriasis have a 3-fold higher risk of developing Crohn’s disease as compared to the general population; and Crohn’s disease patients have a 7-fold higher risk of developing psoriasis[75]. Recently, Scher *et al*[76], using pyrosequencing, found that patients with psoriatic arthritis and patients with skin psoriasis had a decreased bacterial diversity and a reduced relative abundance of some bacterial taxa such as *Akkermansia*, *Ruminococcus*, and *Pseudobutyrivibrio,* as compared to healthy controls. Among the risk factors for psoriatic diseases summarized in Table 1, overall, the alteration of gut microbiota may translate into physiological consequences including poor regulation of intestinal immune responses that may then affect distant organ systems[77-85]. Given the gut microbiome’s influence on the Gut-Skin axis, probiotic supplementation may have a promising role in the management of psoriatic patients. On this point, Gueniche *et al*[77]*,* in a randomized double-blind placebo-controlled clinical study, showed that oral supplementation with the probiotic strain *Lactobacillus paracasei* decreased skin sensitivity and increased the rate of barrier function recovery.

**Infectomics and autoinfectomics in psoriasis**

The term “exposome” defines all environmental factors, including infectious and non-infectious agents, to which a human is exposed over a lifetime[80]. The “microbiota” is a term used to describe the 10-100 trillion symbiotic microbes harbored by each human; the “microbiome” consists of the genes that these microbes harbor[81]; the“infectome” is a part of the microbiome, referring to the collection of human exposure to infectious agents; the “autoinfectome” describes a part of the microbiome that includes the infectious agents linked to the presence of autoimmune diseases[82]. Figure 1 summarizes the main interactions between the exposome, microbiome, infectome and autoinfectome. Recently a systematic review, which included 933 psoriatic arthritis patients and 1611 controls, aimed to evaluate the link between infections (viral and bacterial infections) and the risk of psoriatic arthritis and reported a controversial result that exhibited a trend but failed to achieve significance[83]. However, differences exist between infection, colonization and dysbiosis, as suggested by several studies highlighting a different mycobiome and microbiome in psoriasis, psoriatic arthritis and control subjects[84]. In fact, a dysregulation in the ratio of *Firmicutes*/*Bacteroidetes* was highlighted in the gut microbiome of psoriatic patients; furthermore, *Actinobacteria* was reduced in the gut of psoriatic patients. Gut dysbiosis was also found to be related to skin dysbiosis as decreased beta-diversity in psoriatic skin microbiome is related to an increased risk of developing psoriatic arthritis, and skin flora are now regarded as possible sensitive and specific biomarkers to predict comorbidities in psoriatic patients[84]. The skin microbiota in psoriasis patients seems to be less diverse when compared to healthy persons with a decrease in *Coprococcus* species[76], and more recently *Akkermansia muciniphila*[85]. The characteristic proinflammatory mediators of psoriatic skin lesions have been reported to be the innate antimicrobial peptides and proteins (AMPs). AMPs are a diverse group of small molecules (12–100 amino acid residues) that constitute the primary effector system of innate immunity against microbes[86]. Although medical history certainly plays a crucial role in psoriasis management, it has some limitations such as recall bias. In particular, it has already been demonstrated that not all infections or even dysbiosis that are not clinically evident are still capable of triggering an immune/autoimmune response. Currently, there is limited, evolving information suggesting some benefit from fecal microbiota transplantation, although durability of response is a concern and currently under investigation[87,88].

**Treatment OF psoriasis: taking into account interactionS with diet and microbiome**

Drug therapies are an important consideration for alteration of the cutaneous and gut microbiome[78,89] as well as the mycobiome; however, few studies have explored this topic, and are summarized in Table 2.

To date IL-17/IL-17RA signaling has been demonstrated to be a key component in regulating *Candida* in the gut microbiome[79]. Furthermore, psoriatic arthritis and IBD have genetic and environmental similarities, highlighting that microbiome dysbiosis may affect autoimmune diseases[78]. T-cell activation is an important mechanism of psoriasis, and dysbiosis has been associated with the differentiation of T-cells into effector T cells with fewer regulatory T-cells resulting in changes in the levels of cytokines. In particular, Th17 inhibitors produced the best response compared to patients treated with tumor necrosis factor-α and IL-23 inhibitors[78]. It is interesting to note that this Th-17 mediated response may not translate to the skin, as the skin microbiome could prevent the development of psoriatic plaques in these individuals[3,79]. It is possible that transplanting fecal microbiota could improve or resolve the dysbiosis present in psoriatic arthritis[79]. Fecal microbiota transplants have been used with success in IBD. In fact, Kragsnaes *et al*[90] are currently exploring fecal microbiota transplantation (FMT) in patients with psoriatic arthritis currently on methotrexate to examine their treatment response. Evaluating evidence of the efficacy of FMT is likely due to be complicated by various factors including antibiotic use, prior psoriasis therapy, subtype of psoriasis and comorbidities which similarly affect the gut or skin microbiome[91]. Future studies are needed to identify potential modulators of the gut and skin microbiome, and identify which medications would be optimal for a patient’s individual microbiome signature.

Conclusion

The interaction of Microbiome-Gut-Nutrients in psoriasis is beginning to be understood with the advent of improved ‘omics technologies and their possible integration with each other in order to more precisely separate psoriasis patient endotypes. The transition from immune-targeted therapy to precision-based therapy will be based on the mix between biological signature, the endotype, and potential specific interaction within the exposome.

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**Footnotes**

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**Figure Legends**

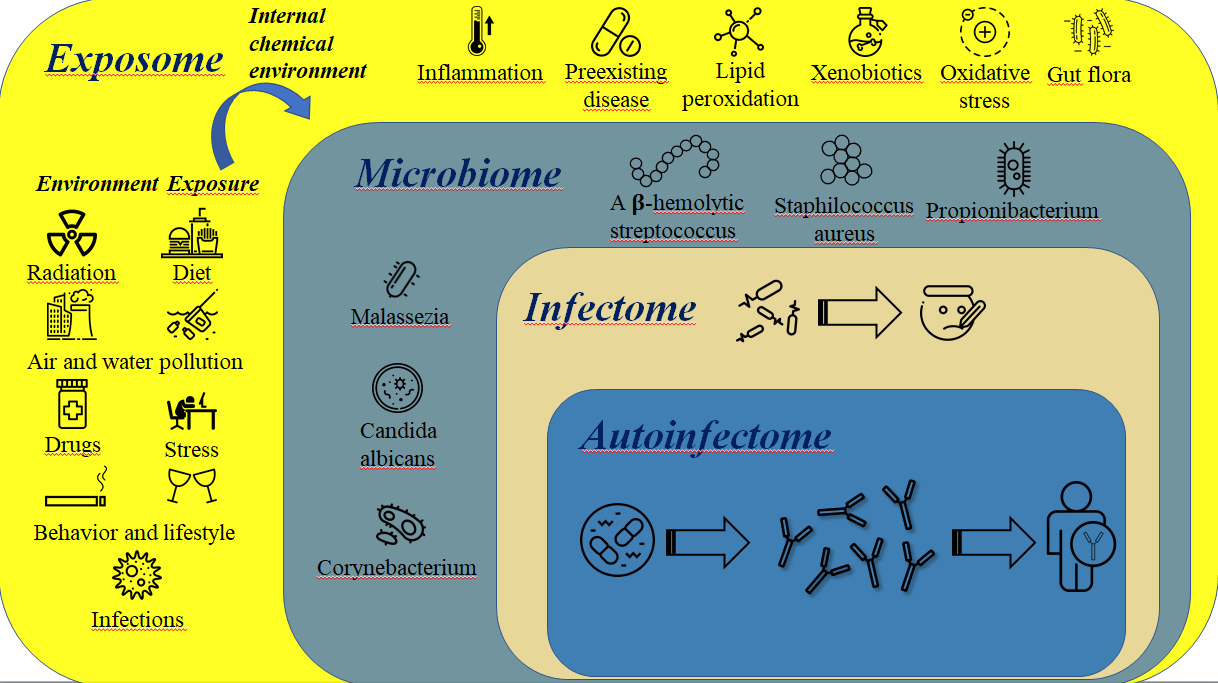
**Table 1 Synthesis of psoriasis risk factor~~s including~~ reporting reference number, first author surname, year of publication, population risk factors reported in each study and the type of psoriasis**

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Ref.** | **Year** | **Studied population** | **Psoriasis risk factors** | **Type of psoriasis** |
| Barrea *et al*[29] | 2016 | Human | Obesity | Not specified |
| Kanemaru *et al*[68] | 2015 | Animal - Mice | Obesity | Psoriasiform dermatitis |
| Dauden *et al*[30] | 2018 | Human | Metabolic disorders | Not specified |
| Barrea *et al*[31] | 2017 | Human | Reduction vitamin D | Not specified |
| Lin *et al*[56] | 2016 | Human | Oxidative stress | Not specified |
| Yang *et al*[67] | 2018 | Animal - Mice | 12-O-tetradecanoylphorbol-13-acetate administration | Psoriasis-like skin lesions |
| Ottman *et al*[70] | 2012 | Human | *Beta-hemolytic streptococcal* infections | Guttate psoriasis |
| Zeng *et al*[71] | 2017 | Human | *Staphylococcus aureus*, *Malassezia* and *Candida albicans* infections | Not specified |
| Alekseyenko *et al*[72] | 2013 | Human | *Corynebacterium, Propionibacterium, Staphylococcus,* and *Streptococcus* infections | Psoriatic plaques |
| Baker *et al*[73] | 2006 | Human | Higher numbers of peptidoglycan-containing cells | Guttate and chronic plaques |
| Oliveira Mde *et al*[75] | 2015 | Human | Inflammatory bowel disease (*i.e.*, Crohn’s disease) | Not specified |
| Scher *et al*[76] | 2015 | Human | Significant reduction in *Akkermansia*, *Ruminococcus*, and *Pseudobutyrivibrio* in gut microbiota | Psoriasic arthritis |
| Tan *et al*[85] | 2018 | Human | Reduction of *Coprococcus species* and *Akkermansia* *muciniphila* in gut microbiota | Psoriatic arthritis |

**Table 2** **Synthesis of psoriasis proposed treatments including reporting reference number, first author surname, year of publication, population risk factors reported in each study and the type of psoriasis**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Ref.** | **Year** | **Study type** | **Studied population** | **Proposed treatment** | **Type of psoriasis** |
| Barrea *et al*[32]  Phan *et al*[33] | 2015  2018 | Cohort | Human | Dietary antioxidants (omega 3 polyunsatured fatty acids derived from fish oil, vitamin B12 , vitamin D and selenium) | Not specified |
| Subbiah *et al*[55] | 2010 | Review | - | Low-energy diets, vegetarian diets, formula diet weight loss programs, gluten-free or very low-calorie carbohydrate-free diet | Not specified |
| Subbiah *et al*[55]  Lin *et al*[85] | 2010  2016 | Review | - | Dietary antioxidants: Vitamins (A, E and C), and oligo-elements (iron, copper, manganese, zinc, and selenium) | Not specified |
| Murzaku *et al*[61]  Upala *et al*[62]  Morken *et al*[63] | 2014  2017  2011 | Review  systematic review  pilot study | Human | Omega-3 polyunsaturated fatty acids | Not specified |
| Yang *et al*[67] | 2018 | RCT | Animal – mice | Nobiletin (Nob) and 5-hydroxy-6,7,8,3′,4′-pentamethoxyflavone (5-HPMF) | Not specified |
| Arbiser *et al*[69] | 2017 | RCT | Animal - mice | Topical administration of solenopsin analogs | Not specified |
| Gueniche *et al*[77] | 2014 | RCT | Human | Oral supplementation with the probiotic *Lactobacillus paracasei* | Not specified |
| Eppinga *et al*[78] | 2014 | Review | - | Th17, TNF-α and IL-22 inhibitors | Psoriatic arthritis |
| Castelino *et al*[79] | 2014 | Review | - | Transplant of fecal microbiome | Psoriatic arthritis |

RCT: Randomized controlled trial; TNF: tumor necrosis factor; IL: interleukin 1.



**Figure 1** **The figure describes the complex panorama of the exposome.** The exposome represent all environmental exposures, from infectious and noninfectious causes. Environmental exposure changes the internal chemical environment that can lead to alterations in the microbiome. The microbiome is the number of all genes of symbiotic microbes harbored by each human. The fraction of the microbiome concerning the collection of human exposure to infectious agents is represented by the infectome. The autoinfectome is the part of the infectome that contains the infectious agents causing autoimmune diseases.