Evaluation of a Product Containing Xyloglucan and Pea Protein on Skin Barrier Permeability

Michela Campolo    Marika Lanza    Alessia Filippone    Irene Paterniti
Giovanna Casili    Sarah A. Scuderi    Alessio Ardizzone    Salvatore Cuzzocrea
Emanuela Esposito
Department of Chemical, Biological, Pharmaceutical and Environmental Sciences, University of Messina, Messina, Italy

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Abstract
Objective: The skin acts as a mechanical and protective barrier against viral, fungal, and bacterial infections. Skin conditions such as atopic dermatitis and psoriasis are characterized by alterations of the skin barrier, often caused by injury and by bacterial infections. In the last years, non-pharmacological interventions have gained great importance in epidermis-related diseases. Xyloglucan (XG) is a polysaccharide that possesses a “mucin-like” molecular structure that confers mucoadhesive properties, allowing XG-containing formulations to act as a protective barrier for the management of different diseases. Moreover, there is also increasing interest in the use of proteins due to their film-forming features. This study aimed to evaluate the barrier-protective properties of a product containing XG and pea protein (PP) in an in vitro model, assessing its effects on the membrane permeability of keratinocytes infected by Staphylococcus aureus.

Methods: HaCaT keratinocytes were pretreated with XG and PP for 3 h and then infected with S. aureus cells (10⁶ bacteria/well) at a multiplicity of infection of 10 for 1 h. The number of bacterial colonies and membrane integrity were measured, respectively. Results: We observed that pretreatment with XG and PP in human HaCaT keratinocytes infected with S. aureus significantly increased trans-epithelial electrical resistance (a marker of skin barrier function) measurement, reduced lucifer yellow (a marker of membrane integrity) permeation across the monolayer, and released lactate dehydrogenase (a marker of tissue damage). Moreover, XG and PP pretreatment was able to reduce bacterial adherence, avoiding S. aureus infection. Conclusion: In summary, we demonstrated that the product containing XG and PP was able to maintain barrier permeability preserving its integrity, and therefore, it can be considered as an interesting approach for the management of epidermis-related diseases.

Introduction
The skin is constantly exposed to a variety of external noxious agents. One of the most important functions of the epidermis is to generate a permeable barrier against viral, fungal, and bacterial infections and prevent excess trans-epidermal water loss [1, 2]. The loss of these functions leads to the colonization of different bacteria, in-
including *Staphylococcus aureus* [3, 4]. *S. aureus* uses adhesion molecules to adhere to the skin producing extracellular vesicles containing immunogenic compounds and serine proteases that can further damage the skin barrier, all of which alter the epidermal permeability [5]. Therefore, appropriately formulated skin products are needed to maintain skin barrier integrity, preventing acute or chronic inflammatory and infectious skin diseases. Xyloglucan (XG) belongs to a new class of products defined as “mucosal protectors,” which form a protective film and exercise a mechanical effect [6]. XG is a hemicellulose of plant origin, extracted from seeds of the tamarind tree (*Tamarindus indica*). It has been shown that XG is able to restore physiological functions of the intestinal wall [7]; XG forms a protective film which facilitates the mucosal resistance to pathological invaders, restoring normal function. This barrier effect decreases the permeability of the intestinal mucosa and avoids the adherence and proliferation of pathogens [7]. Furthermore, in vitro studies have shown that XG increases trans-epithelial electrical resistance (TEER) following *Escherichia coli* infection [7]. The TEER is an index that assesses good functionality of the mucosal tight junctions and therefore confirms the ability of XG to neutralize mucosal permeability after *E. coli* infection [8]. Moreover, considerable importance was given to the proteins extracted from the *Pisum sativum* plant. Pea protein (PP) is a fiber-rich legume that is low in fat and boasts an extraordinary aminoacidic profile conferring to PP favorable health properties due to its film-forming features [6, 9]. Given the beneficial protective properties of XG and PP towards mucosal barriers, the aim of this study was to evaluate the barrier-protective properties of a product containing XG and PP in an in vitro model, testing the permeability of keratinocyte monolayers upon *S. aureus* infection.

**Materials and Methods**

**Materials**

All chemicals were obtained from the highest grade of commercial sources. The product containing XG and PP was kindly provided by DEVINTEC SAGL (Lugano, Switzerland). *S. aureus* (ATCC 29213) was purchased by ATCC materials resource. Human immortalized keratinocytes (HaCaT cell line) (CLS Cell Lines Service, 300493) were used for the study. For Franz diffusion cell studies, we used XG+PP at the dose of 200 mg, dissolved in a physiological buffer.

**S. aureus Culture**

For the experimental infection, *S. aureus* strain was grown to the exponential phase (about $1 \times 10^9$ CFU/mL) in brain heart infusion broth at 37° C overnight with shaking, and harvested by centrifugation (5,000 g for 5 min), washed (3× in PBS), and suspended to the required number in fresh, serum- and antibiotic-free RPMI 1640 medium. The viable count for infection was made through a spread plate technique.

**S. aureus Infection**

HaCaT keratinocytes were grown in RPMI 1640 medium supplied with 10% FBS at 37° C in 5% CO₂. The keratinocytes monolayers were grown in 12-well tissue culture plates and infected with SA cells ($10^9$ bacteria/well) at a multiplicity of infection of 10 for 1 h at 37° C [10]. Extracellular bacteria were removed by washing the keratinocytes 3 times with PBS, the medium was replaced with 1% FBS-RPMI 1640 medium containing 50 µg/mL of gentamicin to kill the extracellular SA bacteria for 1 h. Subsequently, the cells were washed 3 times with PBS to remove the gentamicin. XG+PP at chosen concentrations was added to cells 3 h before SA infection (1 h at 37° C). After that, infected cells were washed 3 times with PBS and then incubated for 15 min at 37° C with 0.02% Triton X-100 to lyse the keratinocytes and release the intracellular bacteria into the lysate. Various dilutions of the lysates were plated on brain heart infusion agar plates and incubated overnight at 37° C. The colonies were counted to determine the rate of infection. To analyze the number of surviving bacteria compared to the initial number of intracellular bacteria (time zero), the relative number of CFU was calculated as follows: number of CFU at 4 h/number of CFU at time zero.

**Experimental Groups**

Group 1: control + vehicle: PBS was added to HaCaT (n = 12)

Group 2: control + XG+PP: XG+PP was added to HaCaT (n = 12)

Group 3: *S. aureus*: keratinocyte monolayers were infected with *S. aureus* cells ($10^9$ bacteria/well) at a multiplicity of infection of 10 for 1 h at 37° C (n = 12)

Group 4: *S. aureus* + XG+PP: XG+PP was added to cells 3 h before SA infection (n = 12)

**LDH Assay**

The membrane integrity of the keratinocytes was determined by measuring the lactate dehydrogenase (LDH) activity present in the culture supernatant following *S. aureus* infection. The assay was performed using a CytoTox 96 nonradioactive cytotoxicity assay kit (Promega, Madison, WI, USA), according to the manufacturer’s instructions. LDH was measured at 492 nm absorbance using a standard 96-well plate reader. LDH release was expressed as units per liter of supernatant. The percentage of cytotoxicity was calculated as follows: ((experimental release – spontaneous release) / (total release – spontaneous release)) × 100, where a spontaneous release is the amount of LDH activity in supernatants of cells incubated in medium alone, and total release is the LDH activity measured in cell lysates [11].

**Lucifer Yellow Permeation**

Lucifer yellow (LY) was used as a membrane integrity marker. Cells were plated onto 96-transwell plates with polycarbonate membrane inserts at a seeding density of 50,000 cells/cm². The cells were grown to confluence and treated on the 5th day with dilutions of XG+PP for 4 h, then were infected with SA for 1 h. The monolayer integrity was evaluated by measuring the LY permeation across the barrier, as previously described [12]. The LY con-
centrations in the apical and basolateral divisions were determined by measuring their fluorescence intensities compared to the standard curves of LY, at excitation and emission wavelengths of 485 and 529 nm, respectively, using a microplate reader.

**TEER Measurement**

A quantitative measure describing the barrier integrity and the electrical, ohmic resistance of the cell layer was performed. The traditional method for measuring TEER values is to use STX2 electrodes using a 4-point measurement system on Transwell permeable inserts [13]. One set of electrodes was placed in the basolateral compartment and the other was placed in the apical compartment [14]. For measurement of TEER, the cells were seeded at a high plating density (1.25 × 10^5 cells/cm^2) on clear polyester Transwell permeable supports (Corning Glass) in growth medium. Medium (apical and basal) was changed every 24 h before and during treatments. The association XG+PP was added to the culture medium when TEER reached a plateau (2–3 weeks after seeding, ∼3,000 ohms × cm^2). An EVOM epithelial volt/ohmmeter with STX (chopstick) electrodes (World Precision Instruments, Sarasota, FL, USA) was used to measure TEER.

**Franz Diffusion Cell Studies on Hairless Mouse Skin Infected by S. aureus**

The association XG+PP barrier effect on hairless mouse skin infected by S. aureus was observed using the Franz diffusion cell method [15]. Mouse skin was set up in Franz cells apparatus. Briefly, frozen (−20°C) skin was defrosted, the underlying fat was removed, then the epidermis was cleaned with distilled water and dried. The chamber receptor cell was filled with 1 mL of PBS and allowed to equilibrate at 32 ± 2°C in the heated magnetic block for 30 min. The skin was cut into 3.5-cm diameter pieces sufficient to cover the area of the receptor and was fixed between the donor and the receptor chamber [16]. The association XG+PP was placed on the skin surface in the donor compartment and was brushed to cover the whole surface uniformly. S. aureus was harvested and suspended in PBS to determine the bacterial density (1.5 × 10^7 CFU/100 µL) and then was added to the upper well of the skin. After 30 min, the skin surface was washed twice with 6 mL of PBS to recover S. aureus and then serial dilutions of the washings were added to BD Mannitol Salt Agar plates followed by CFU counting to evaluate S. aureus proliferation [17]. After 24 h, the formation of colonies was observed. A control experiment was performed by adding on the skin surface only S. aureus; CFUs were counted after washing as explained above.

**Statistical Analysis**

All values are expressed as mean ± SEM. The results were analyzed by one-way ANOVA followed by a Bonferroni post hoc test for multiple comparisons. A value of p ≤ 0.05 was predetermined as the criterion of significance.
Results

Evaluation of S. aureus CFU on Infected Keratinocytes
The relative number of CFU was estimated by plating out the lysate of infected keratinocytes and counting the number of CFU at each time point (T0 and T4h). The relative number of CFU is the difference between the initial number of intracellular bacteria and the number at 4 h. As shown by Figure 1, XG+PP was able to avoid S. aureus infection by reducing bacteria adherence (Fig. 1).

Evaluation of Keratinocyte Membrane Integrity with LDH Assay and LY Permeation Test
The release of LDH was performed to investigate keratinocyte membrane integrity. LDH release significantly increased in the cell culture supernatant after incubation with S. aureus compared with the control group. The addition of XG+PP was able to counteract the damaging effects of S. aureus infection reducing the percentage of LDH released in HaCaT keratinocytes (Fig. 2a). A LY permeation study was performed to test the effects of XG+PP on HaCaT cell monolayer intactness. S. aureus infection, after 24 h, significantly increased the paracellular flux of LY, while application of XG+PP reduced LY permeation across the monolayer (Fig. 2b).

Evaluation of Skin Barrier Function with TEER Test
HaCaT infection with S. aureus significantly reduced TEER measurement, while the administration of XG+PP prevented TEER decrease, enabling the maintenance of skin barrier function and integrity (Fig. 3).

Evaluation of Barrier Effect on Mouse Skin Infected by S. aureus Using Franz Diffusion Cell
The XG+PP barrier effect on hairless mouse skin infected by S. aureus was observed using the Franz diffusion cell method. The bacterial burden was evaluated by plating serial dilutions of S. aureus onto the BD Mannitol Salt Agar dish counting the CFU to evaluate bacterial proliferation. It was observed that XG+PP decreased the attachment and penetration of S. aureus onto the surface of the skin, as demonstrated by the increased number of CFU counted, as shown in Figure 4.

Discussion and Conclusions
The disruption of the skin barrier function was recently associated with a variety of diseases, such as atopic dermatitis, contributing to a greater risk of contracting bacterial infections, in particular by S. aureus [18, 19]. Therefore, compounds able to preserve barrier integrity could represent a promising approach to counteract skin pathologies. The aim of this study was to demonstrate the barrier-protective properties of a product containing XG and PP through an in vitro permeability model of keratinocyte membrane infected by S. aureus. Our data demonstrated that the number of internalized bacterial colo-
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Skin from male CD1 mice (Envigo, Milan, Italy) weighing 20–25 g were used for Franz cell study. This study was approved by the University of Messina Review Board for the care of animals (Protocol number 8/U-apr16). Animal care was in conformity with current legislation for the protection of animals used for scientific purposes (Directive 2010/63/EU, April 9, 2016) and the ARRIVE guidelines.

Conflict of Interest Statement

The study was financed by Devintec SAGL, Switzerland. The authors have no other relevant affiliations or financial involvement with any organization or entity with a financial interest, or financial conflict with the subject matter or materials discussed in the manuscript apart from those disclosed. No writing assistance was utilized in the production of the manuscript.

Author Contributions

E.E. and S.C. planned the experiments. M.L., A.F., G.C., S.A.S., and A.A. performed experiments. M.C. and I.P. analyzed the results and prepared the manuscript. All authors read and approved the final manuscript.

Data Availability Statement

The authors declare that all data and materials supporting the findings of this study are available within the article. The data that support the findings of this study are available from the corresponding author upon reasonable request.
References


