Evaluation of microbial content of some sunscreen creams in Iran’s market

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Received: Feb 5, 2015, Revised: May 27, 2015, Accepted: Jun 28, 2015

Abstract
The risk of microbial contamination in the cosmetic products especially in smuggled preparations and transmission of it to consumers is very high. In this study, the microbial content and the pollution of some sunscreen creams in the market and one sample in official market as witness were evaluated. The microbial content (bacterial total count, fungal count, and presence of Pseudomonas aeruginosa, Staphylococcus aureus and Entrobacter) of 5 samples of sunscreen cream in the market and two samples in official market were evaluated by two methods (pour plate and Multiple tube technique). All samples showed high microbial and fungal contamination. Entrobacters was observed in all samples. Staphylococcus aureus was recognized in one of the non-standard sunscreens. High level of contamination in sunscreen creams, can affect consumers, health. It seems that low grade raw materials, and insufficient manufacturing surveillance in production process are the main factors in the contamination.

Keywords: Cosmetic, sunscreen, microbial content, Pseudomonas aeruginosa, Staphylococcus aureus

Pharm Biomed Res 2015; 1(2): 30-34 DOI: 10.18869/acadpub.pbr.1.2.30
Pseudomonas aeroginosa should not be observed in these products (9). The cosmetic products may be spoiled in two ways: in manufacturing process or by during consumer use (8). Raw materials and manufacturing apparatus are the main sources for microbial contamination. This contamination might be causes changes in the composition, odor, or color of the products (8, 10). Contamination of cosmetic products directly may affects on human health as a result of the formation of metabolites microbial harmful and spoilage of products (3). Legislation and introduction of GMP (Good Manufacturing Practice) has improved the microbiological standards, but a contaminated cosmetic product has serious consequences for the consumers (8). By use of water and raw material with suitable quality, GMP leads to preparation of products with lower microbial contamination. The suitable storage and the use of noninvasive packages (such as tubes, pumps or narrow orifice containers) causes of reducing contamination levels during storage and using products remains (11). The inclusion of essential minerals, growth factors, moisture content and acidity provides favorable environment for microbial growth (12). Only a few studies have been done on the microbial quality of sunscreens and there are very limited data available in Iran. The present study was performed to determine the microbial quality of sunscreens products (illegal and non-illegal products) available in Iran market.

Materials and methods
In this study, the microbial content of seven sunscreen creams (five foreign creams from informal and non-standard market and two creams as control from Iranian products which are presented drugstore) was evaluated. All of the samples were analyzed to detect the presence of total bacterial and fungal count (yeast and mold). The presence of Staphylococcus aureus, Pseudomonas aeroginosa, and Enterobacters were investigated based on United States Pharmacopeia (13). The surface of three sample containers from each cream were swabbed and disinfected by 70% (v/v) ethanol before opening. For tubed products a large sample was extruded into a sterile plate and mixed thoroughly with a sterile spatula. Samples were opened and weighed under the laminar air flow with aseptic conditions. To determine the microbial content of the samples and ensure the absence of antimicrobial effect of potential products and the possibility of microorganisms growth, preliminary experiment with inoculation of Staphylococcus aureus (PTCC 1112) and Pseudomonas aeroginosa (PTCC 1074) were performed on all samples. Lack of growth of inoculated microorganisms on the plates, showed the action of preservative and was proved necessityof the use of 3% Tween 80 and 0.5% lecithin for neutralization of preservative effect. Pour plate and multiple-tube methods were used for total microbial counts. In the Plate Method, one gram was aseptically taken from each product and placed into a 9-ml sterile normal saline solution as diluents with 3% Tween 80 and 0.5% lecithin thus giving a 10⁻¹ dilution. Bacterial challenge levels and yeast and mold were 10⁴ CFU/g of products. One milliliter of above suspension was transferred into two sterile plates, under aseptic conditions. For aerobic bacterial colony counts, 20 milliliter of sterile Soybean Casein Digest Agar (SCDA) (Merck Co, Germany) was added at 45 °C to each plate aseptically. The plates were rotated to completely disperse and then incubated at 35 °C for 24-48 h. Sabouraud
Dextrose Agar (SDA) (Merck Co, Germany) was used for yeast and mold cultivation, and incubated at 25 °C for 5-7 days. After incubation, the number of colonies was recorded for each plate. In the multiple-tubes method, one gram was aseptically taken from each product and placed into a 9-mL sterile normal saline solution as diluents with 3% Tween 80 and 0.5% lecithin. One milliliter of this suspension was transferred into tubes containing Soybean Casein Digest Broth (SCDB) (Merck Co, Germany) according to the method, aseptically and then were incubated at 35 °C for 48 h than tubes with no growth were reported negative for the presence of microorganisms.

For detecting prohibited microorganisms according to the USP, one gram was taken from each product under condition aseptic and placed into a 10-mL sterile normal saline solution as diluent with 3% Tween 80 and 0.5% lecithin and was mixed well and tubes were incubated at 35 °C for 48 h. After incubation, tubes with no growth were reported negative for the presence of Staphylococcus aureus, Pseudomonas aeruginosa and Enterobacters, then a loop of medium with positive growth was transferred to Mannitol-salt Agar (Merck Co, Germany) medium for Staphylococcus aureus detection. This process was performed to Cetrimide Agar and Mac. conkey agar (Merck Co, Germany) for Pseudomonas aeruginosa and Entrobacter detection respectively. After incubation at 35 °C for 48 h, the plates were checked. In the case of growth on these selective mediums, the sample was reported positive for the presence of related microorganism.

**Results**

The results of the plate method showed that microbial count in 60% (3 creams) of the non-standard samples and all of control samples contamination was more than $10^4$CFU/g (Table 1). The results of total count of all samples based on multiple tube method showed that all of samples were outside the standard range (contamination >$10^3$CFU.g$^{-1}$ or mL$^{-1}$).

Fungal (yeast and mold) contamination was observed in all of sunscreens and total count was more than $10^3$ CFU.g$^{-1}$ or mL$^{-1}$. Enterobacter contamination was observed in all samples and in 14.29% (1 of 7 creams) of cases but Staphylococcus aureus contamination was seen only in one of the non-standard sunscreen cream. *Pseudomonas aeruginosa* was not observed in these preparations.

**Discussion**

Hygienic characteristics of creams and other cosmetic products to provide the consumer with regard to the standards defined in each country should be at an acceptable level. Condition of manufacturing, GMP observation and antimicrobial preservative are the main factors in microbial contamination and level of hygienic state of cosmetics and toiletries, according to criteria of National Institute of Standards (14). Other researches proved that one of the main recall reasons is microbial contamination of cosmetic products (3-5).

Non-compliance with hygiene issues in the manufacturing processes of product and hygienic products with packaging non-standard and inadequate can provide easily background of microbial contamination of product. *Staphylococcus aureus* was one of pathogenic bacteria isolated in this study that according to the USP should not be seen in topical preparations. It is from the most factors important skin pathogens that can gain antibiotic resistance genes through chromosomal and non-chromosomal multiple ways and causing is irreparable effects on consumers (14).
Staphylococcus aureus has a special ability to colonize on the skin of patients with eczema and atopic dermatitis; and bacteria colonization is from major factors in aggravate skin lesions (15). Haft-baradaran, et al. showed that 40% of the Iranian sunscreen creams, 73.3% of the imported products, and 43.3% of the formulated products at the time of purchase contained at least one of the objectionable microorganisms of the samples showed contamination with Staphylococcus aureus, Pseudomonas aeruginosa. This study showed yeasts and molds contamination in evaluated creams too (3).

Keshavarz et al. study showed microbial contamination in 46% of the 135 moisturizing cream. The most contamination related was to Pseudomonas aerogenosa. Percentage of contamination to Pseudomonas aerogenosa, Staphylococcus aureus, coagulase-negative staphylococci, yeast, and mold were 35.5, 6.4, 26, 24.1, 6.4 and 1.6% respectively (14).

Lundov et al. reported 68% contamination in some moisturizing creams; 30% of these creams were contaminated to Staphylococcus aureus (16).

Behravan et al. showed that the percentage of contamination to Gram positive Bacilli, Staphylococcus aureus and non-Escherichia coli Gram-negative microorganisms in the used cosmetic creams, was 54, 38 and 8% respectively. This contamination in unused cosmetic creams was 38, 25 and 0% respectively. This study proved that 17% of unused cream and 10% of used products were in the acceptable range of microbial content. According to Cosmetic, the requirements for cosmetic products set by the FDA, The cosmetic Toiletry and Perfumery Association Ltd. (CTPA) guidelines, and Cosmetic Toiletry and Fragrance Association Inc. (CTFA) guidelines and Cosmetic, Products with a high number of microorganisms or products containing pathogenic microorganisms would be considered as spoiled (6).

In the present study all samples (100%) were contaminated with pathogenic microorganisms Enterobacter and counts of molds and yeasts (Table 1) were not in the acceptable range. Samples contained Staphylococcus aureus of pathogenic bacteria (14.29%) was among samples of the informal market in Iran.

The important note in this study was high levels of contamination in the sunscreen creams before use and contamination in the creams was, in during manufacture and

<table>
<thead>
<tr>
<th>Sunscreen cream code</th>
<th>Total bacterial count (CFU/g)</th>
<th>Yeast and mold counts (CFU/g)</th>
<th>Isolated microorganisms</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Plate method</td>
<td>Multiple-tube method</td>
<td></td>
</tr>
<tr>
<td>Sample 1</td>
<td>1×10^2</td>
<td>&gt;1100</td>
<td>3×10^3 Enterobacter</td>
</tr>
<tr>
<td>Sample 2</td>
<td>6×10^2</td>
<td>&gt;1100</td>
<td>3×10^3 Enterobacter</td>
</tr>
<tr>
<td>Sample 3</td>
<td>1×10^5</td>
<td>&gt;1100</td>
<td>6×10^3 Enterobacter, Staphylococcus aureus</td>
</tr>
<tr>
<td>Sample 4</td>
<td>2×10^4</td>
<td>&gt;1100</td>
<td>1×10^4 Enterobacter</td>
</tr>
<tr>
<td>Sample 5</td>
<td>4×10^4</td>
<td>&gt;1100</td>
<td>1×10^4 Enterobacter</td>
</tr>
<tr>
<td>Control 1</td>
<td>6×10^5</td>
<td>&gt;1100</td>
<td>6.4×10^5 Enterobacter</td>
</tr>
<tr>
<td>Control 2</td>
<td>3.1×10^5</td>
<td>&gt;1100</td>
<td>1.5×10^5 Enterobacter</td>
</tr>
</tbody>
</table>
Microbial contamination of sunscreen creams

before reaching to the consumer which emphasis has on necessity to improve functional the quality assurance system in the manufacture process, packaging and adequate preservation and other factors involved in contamination (2).

Conclusion

The high microbial contamination was observed in Iranian and imported sunscreen creams. Microbial contamination of sunscreen creams is a potential health danger to consumers. It appears that it is necessary to inspect and supervise the products during manufacture and packaging and adequate preservation. It is strongly recommended to control and regulate cosmetics by health organization to ensure quality and safety of this type of products.

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Acknowledgment

This article was written based on MSc. thesis of MA Narges Sedghi Sharif Abad, and was supported by a grant from the research council of the Mazandaran University of Medical Sciences.

Conflict of interest

The authors declared no potential conflict of interest with respect to the authorship, and/or publication of this study.