

ATTACHMENT TO THE TEST REPORT**Attachment No 19770/08/25**sample/samples No 19770/08/25
order No K/0/08/2025/236**DERMATOLOGICAL (PATCH TEST) – SENSITIVE SKIN**

Product name	Restorative Cream for Dry Flaky Skin
Date of test start	18-08-2025
Date of test end	22-08-2025
Report date	27-08-2025

Purpose of the study

Assessment of irritation / allergenic effect of the product

Microbial purity testingThe microbiological purity tests carried out in GBA POLSKA Laboratory / ~~delivered by the Customer.~~**Product characteristics**

Appearance:	Emulsion
Color:	Pink
Fragrance:	Characteristic of the raw materials used
Product purpose:	Skin Care
How to use:	2x daily. On clean skin .
Packaging:	Replacement

Qualitative product composition

INCI composition (*):

Aqua, Urea, Sodium Phytate, Sodium Levulinate, Sodium Anisate, Niacinamide, Aloe Barbadensis Leaf Juice Powder, Sucrose Stearate, Glycerin, Xanthan Gum, Glyceryl Stearate, Copernicia Cerifera (Carnauba) Wax, Sodium Stearoyl Lactylate, Cetearyl Alcohol, Oenothera Biennis (Evening Primrose) Oil, Helianthus Annuus (Sunflower) Seed Oil, Avena Sativa (Oat) Kernel Oil, Simmondsia Chinensis (Jojoba) Seed Oil, Cannabis

Sativa Seed Oil, Cannabidiol, Borago Officinalis Seed Oil, Cyanocobalamin, Methylsulfonylmethane, Ganoderma Lucidum Extract, Inonotus Obliquus (Mushroom) Extract, Cordyceps Sinensis Extract, Lactobacillus Ferment, Colloidal Oatmeal, Bisabolol, Tocopherol, Salicylic Acid, Salix Nigra (Willow) Bark Extract, Centella Asiatica Extract, Potassium Sorbate, Sodium Benzoate, Benzyl Alcohol.

(*) - The Customer bears full responsibility for compliance of samples delivered for testing with the declared qualitative composition. The Laboratory does not analyse the composition in regard to the current regulatory requirements.

The scope of tests in accordance with:

- » *The general principles of medical ethics in clinical research coming from Declaration of Helsinki (June 1964) and its successive amendments*
- » *Regulation of the European Parliament and of the Council (WE) No. 1223/2009 dated 30.11.2009 regarding cosmetic products*
- » *COLIPA Guidelines*
- » *„Product test Guidelines for the Assessment of Human Skin Compatibility 1997”*
- » *„Guidelines for the Evaluation of the Efficacy of Cosmetic Products 2008”*
- » *Memorandum on use of Human Data in risk assessment of skin sensitisation – SCCS/1567/15*
- » *The SCCS Notes of Guidance for the Testing of Cosmetic Ingredients and their Safety Evaluation 12th Revision – SCCS/1647/22*
- » *European Regulation (EU) 2016/679 and its successive amendments on the protection of individuals with regard to the processing of personal data and free movement of such data.*

Testing methodology

The study was conducted in accordance with the Research Procedure SOP-MET-029.

The tested substance is applied to the skin using special chambers fixed on a hypoallergenic adhesive (chambers IQ Ultimate®). The chambers are filled with the substances being tested and then affixed to the skin of the back in the shoulder blade area. During the tests, one cannot soak his/her back and should avoid sudden movements and sweating. Patches with test substances are left on the test skin for 48 hours.

After the removal of the adhesives with the chambers, the excess test substances are removed by pressing a paper towel onto the skin. The skin reaction is assessed after 15 minutes after removing the adhesives (after 48 hours) and after 72 hours according to a scale that is consistent with the scale generally accepted in dermatological tests. Additionally, in the case of positive skin reactions, the skin reaction is also assessed after 96 hours and on the following days until symptoms resolve.

Evaluation parameters

The patch test readings were made according to the International Contact Dermatitis Research Group (ICDRG) Graphical scale of the ICDRG patch test reading:



(-) negative reaction	no reaction
(?) doubtful reaction	subtle erythema, palpably undetectable erythematous spot
(+) weak reaction	palpable erythema, suggestive of mild edema / infiltration, clumping may occur, no blisters
(++) strong reaction	increased swelling, infiltration, clots, blisters
(+++) very strong reaction	blisters, erosions, ulceration
(IR) irritative reaction	shiny skin, dry skin, pimples, erythema

A product can be classified as non-irritant and compliant with the Skin Compatibility Test if the mean of the reactions does not exceed 0,10 in a group of test persons, according to the table below:

IRRITATION POTENTIAL		
$0,00 \leq \text{average} \leq 0,10$	POSITIVE REPORT	

RATING SCALE		
Negative result	0	0
Doubtful reaction	?	0,5
Weak reaction	+	1
Strong reaction	++	2
Very strong reaction	+++	3

Selection of Study Participants

The selection of Study Participants was carried out in accordance with internal procedure, SOP ORG 001, taking into account:

- »Helsinki Declaration of 1964 (with later additions)
- »Current Polish and European legal regulations
- »Cosmetics Europe guidelines using the inclusion and exclusion criteria

20 skin-healthy volunteers with sensitive skin were chosen to become subjects in the study. Volunteers filled out a detailed survey regarding their lifestyle, the current state of health, past illnesses, eating habits, medicines and the use of stimulants and the survey on coexisting skin problems (allergy issues

included). The volunteers were selected from this general panel on the basis of inclusion criteria and non-inclusion criteria specific to the study and on their ability to respect the constraints required by the study methodology. The application of the product was preceded by dermatology examination assessing, above the others, the type of skin and presence of any pathological changes on the skin.

GENERAL INCLUSION CRITERIA	Healthy subject
	Declaring to have a health coverage
	Signing an „informed consent form” for this study
	Skin without irritation and changes requiring pharmacological treatment
	Cooperative subject, aware of the necessity and duration of controls, free to ensure the visits to the investigating center

SPECIFIC INCLUSION CRITERIA	Gender	<i>Female and male</i>
	Age	<i>18+</i>
	Skin type	<i>Sensitive</i>
	Skin phototype (Fitzpatrick scale)	<i>I-IV</i>
	Other	Regular or occasional users of body cosmetics products.

GENERAL NON INCLUSION CRITERIA	Subjects who use any treatment on the studied zone
	Being in exclusion period
	Pregnant or breastfeeding woman or woman planning a pregnancy during the study
	Subject having a skin disease on the studied zone
	Subject exhibiting or having a known history of acute or chronic dermatological, medical and/or physical conditions that could influence the outcome of the test
	Subject considered by the investigator to be likely not compliant with the study methodology
	Subject undergoing treatment with antihistamines, nonsteroidal anti-inflammatory agents, corticosteroids and/or any other medications that could have interfered with the results of this study.
	Subject planning the hospitalization during the study period

All volunteers selected for the study met the requirements for inclusion in the study and signed consent for conscious participation in the study, and were informed about the purpose of the study, how it was conducted and about possible side effects. The skin, where the patch tests were applied, was healthy and free of skin lesions.

Results

<i>Volunteers characteristic</i>				<i>Results after 48h</i>	<i>Results after 72h</i>	
<i>No.</i>	<i>Sex</i>	<i>Age</i>	<i>Skin type</i>	<i>Skin reaction</i>	<i>Skin reaction</i>	
1	F	51	S	(-)	(-)	
2	F	61	S	(-)	(-)	
3	F	62	S	(-)	(-)	
4	M	34	S	(-)	(-)	
5	F	38	S	(-)	(-)	
6	F	44	S	(-)	(-)	
7	F	50	S	(-)	(-)	
8	F	42	S	(-)	(-)	
9	F	43	S	(-)	(-)	
10	F	45	S	(-)	(-)	
11	F	45	S	(-)	(-)	
12	M	71	S	(-)	(-)	
13	F	25	S	(-)	(-)	
14	F	64	S	(-)	(-)	
15	F	66	S	(-)	(-)	
16	F	51	S	(-)	(-)	
17	F	24	S	(-)	(-)	
18	F	53	S	(-)	(-)	
19	M	30	S	(-)	(-)	
20	F	49	S	(-)	(-)	
	<i>F</i>	17	<i>Age min.</i>	24	<i>Irritant potential</i>	0,00
	<i>M</i>	3	<i>Age max.</i>	71		

Legend: F – female, M – male, S – sensitive, (-) – no skin reaction

SUMMARY

In the group of 20 people who underwent the study, no positive contact allergy or irritant reaction was observed after using the product. The irritation potential score is $\leq 0,10$.

The lack of positive reactions indicates that the tested product does not show irritating and sensitizing effects in contact with the sensitive skin.

Based on the results of the tests performed, we conclude that the „Restorative Cream for Dry Flaky Skin” product meets the requirements of the Skin Compatibility Test and may be estimated as non-irritant.

The issued opinion does not apply to people who are allergic to any of the components of the product under evaluation.

Prepared on: 27-08-2025	Prepared: GBA POLSKA Employee no.: 2611	Authorized: GBA POLSKA Employee no.: 2599
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Authorized Results:

Dermatologist:

Dr Dominika Perron-Plusa - Specialist
in Dermatology and Venereology

**The appendix from the research is signed with a qualified electronic seal of GBA Polska
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