



UNIVERSITA' DEGLI STUDI DI PAVIA
DIPARTIMENTO DI MEDICINA INTERNA E TERAPIA MEDICA
(Direttore: Prof. Plinio Richelmi)



Open Patch Test

**Evaluation of the possible irritant power of a cosmetic product
according to the amended Draize classification**

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Peter Greven Physiaderm GmbH

**Peter Greven Physiaderm GmbH Physio UV 50 Spray
RO_27_01_20_CP**

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SCIENTIFIC TECHNICAL COMMITTEE

Bio Basic Europe S.r.l.

Plinio Richelmi, Ornella Pastoris, Fernando Marco Bianchi,
Antonella Colombo, Alessandra Di Benedetto, Evelyn Falconi
Klein, Laura Mainardi, Claudio Angelinetta, Daniela Gandini,
Antonella Praticò, Eliana Regola, Francesca Vallotto

MONITOR

SCIENTIFIC PERSON IN CHARGE

Prof. Plinio RICHELMI

Professore Ordinario di Farmacologia

Direttore del Dipartimento di Medicina Interna e Terapia Medica

Facoltà Medicina e Chirurgia

Università di Pavia

Via Ferrata 6- 27100 PAVIA

Experimenter

Dr. Fernando Marco BIANCHI

Dermatologist

consultant at CDC - Dermo-clinic Research Institute

Viale Misurata 59, Milano

Quality Control

Dr. Claudio ANGELINETTA

Master's degree in Chemistry at University of Milan,

M.Sc. Cosmetology at University of Milan

Technical Director BIO BASIC EUROPE S.r.l.

Via Antonio Panizzi, 10 20146 Milano

Person responsible for the report

Daniela GANDINI

Master's Degree in Biothecnology

University of Milan Bicocca

Clinical Test Reporting Officer – Safety

BIO BASIC EUROPE s.r.l.

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Based on our experience, we suggest to check every 3 years its compliance with the guidelines in force..



Title

Patch Test.

Evaluation of the possible irritant powers of a cosmetic product according to amended Draize classification.

Aim

To evaluate the tolerability of a cosmetic product through the individualization and the classification of its potential irritant power.

Legal information

In accordance with the current legislation and the declaration of Helsinki, all volunteers must be adequately informed of the aims, methods, clinical trial details, anticipated benefits and potential undesirable effects of the study. Each panellist must sign an informed consent form, which is managed and archived by applying the internal procedure of the Quality Management System of Bio Basic Europe S.r.l.

Contract information

Technical report performed by BIO BASIC EUROPE s.r.l. and Università degli Studi di Pavia.

Final technical report written by BIO BASIC EUROPE s.r.l. on behalf of Peter Greven Physiaderm GmbH

Experimentation performed at CDC - Dermo-clinic Research Institute

Materials and methods

The test will be carried out on 30 volunteers recruited according to the following inclusion and exclusion criteria: a) good state of health/absence of psychological and/or cognitive disorders; b) no dermatopathies and allergic pathologies (to cosmetics or other specific excipient), or other pathologies (as unknown irritant responses); c) no ongoing pharmacological treatments that could affect the result of the test; d) no participations in other clinical trial during the previous 30 days; e) signature of the informed consent form.

Applications of the samples

The products have been applied following their usual use : as they are (leave-on products) or with a 10% standard concentration (rinse-off products). Liquid products are applied on the skin in a dose of 20 µl. Solid and semisolid products are applied in a dose of 20 µg .

A plaster containing disks of blotting paper is an open patch test device.

For liquid products the plaster contains disks of blotting paper soaked with a certain quantity (20 µl) of the sample to be tested, for solid products the products to be tested are directly applied on skin(20 µg).

Execution of the test

The involved skin area (surface of the back) has been cleaned with a 70% alcoholic solution

- The cosmetic product has been applied in a quantity of 0,02 mg or ml/ cm² of skin.
- The involved skin area is occluded with the plaster
- The cosmetic product has been left in contact with the skin surface for 24 hours.
- The plaster is removed
- The cutaneous reactions have been analysed 15 minutes, 1 and 24 hours after plaster removal
- Negative control with physiological water

OPEN PATCH TEST

No of volunteers	30			
SEX	M (X)		F (X)	
AGE	18-70			
TIMES for TEST EXECUTION: - product application - evaluation of cutaneous reaction (ERYTHEMA and EDEMA) after the Plaster removal	1 st DAY	Product application with the Plaster and skin contact for 24 hours		
	2 nd DAY	Plaster removal and evaluation of the cutaneous reaction		
		15 min [X]	1 hour [X]	24 hours [X]
Product use	as it is (X)		with a 10% dilution (...)	

Liste der Inhaltsstoffe
 INGREDIENTS : Aqua, C12-15 Alkyl Benzoate, Glycerin, Ethylhexyl Salicylate, Isopropyl Palmitate, Ethylhexyl Triazone, Cetearyl Alcohol, Glyceryl Stearate, Butyl Methoxydibenzoylmethane, Bis-Ethylhexyloxyphenol Methoxyphenyl Triazine, Diethylhexyl Butamido Triazone, Phenylbenzimidazole Sulfonic Acid, Sodium Stearoyl Glutamate, 1,2-Hexanediol, Butyrospermum Parkii Butter, Tri-Biphenyl Triazine (Nano), Potassium Hydroxide, Xanthan Gum, Tocopheryl Acetate, Caprylyl Glycol, Ethylhexylglycerin, Decyl Glucoside, Decylene Glycol, Proctone Oamine, Butylene Glycol, Disodium Phosphate, Titanium Dioxide, Silver Chloride, Tocopherol

Alcohol Denat., C12-15 Alkyl Benzoate, Isopropyl Palmitate, Glycerin, Butyl Methoxydibenzoylmethane, Ethylhexyl Stearate, Ethylhexyl Triazone, Ethylhexyl Salicylate, Bis-Ethylhexyloxyphenol Methoxyphenyl Triazine, Diethylhexyl Butamido Triazone, Aqua, Phenylbenzimidazole Sulfonic Acid, Tocopheryl Acetate, Aminomethyl Propanol, Butane, Dimethyl Ether

Evaluation and calculation of the results

The reactions are evaluated according the arbitrary scale reported in table no. 1 .

Erythema and edema are evaluated after the Plaster removal.

The index of total skin irritation (IIM tot) comes out by calculating the average of the irritation indexes for erythemas (IIM Er) and edemas (IIM Ed) at 24 hours.

The product is classified according to table no. 2.

Table no. 1
Scale of evaluation

Erythema	
No erythema	0
Light erythema (hardly visible)	1
Clearly visible erythema	2
Moderate erythema	3
Serious erythema (dark red with possible formation of light eschars).	4
Edema	
No edema	0
Very light edema (hardly visible)	1
Light edema	2
Moderate edema (about 1mm raised skin)	3
Strong edema (extended swelling even beyond the application area)	4

Table no. 2
Classification of the index of average irritation (according to the amended Draize classification)

Index	Class
0,5 from 0,5 to 2,0 from 2,0 to 5,0 from 5,0 to 8,0	non irritant slightly irritant moderately irritant highly irritant

Results

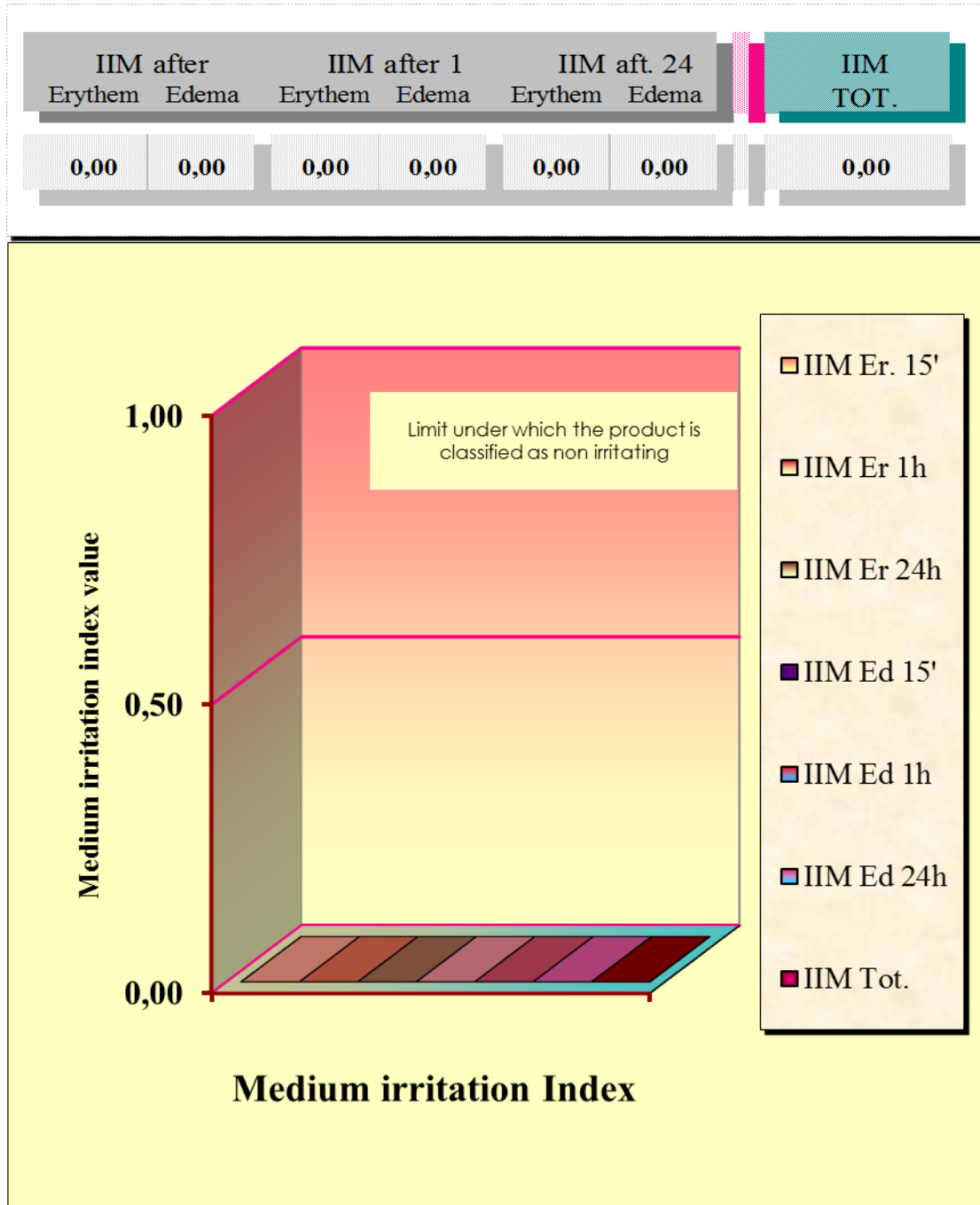
Values of edema and/or erythema

(scale of evaluation: see table no. 1)

Volunteers' references	SEX	Reaction after 15'		Reaction after 1 h		Reaction after 24 h	
		Erythema	Edema	Erythema	Edema	Erythema	Edema
1	F	0	0	0	0	0	0
2	F	0	0	0	0	0	0
3	F	0	0	0	0	0	0
4	F	0	0	0	0	0	0
5	F	0	0	0	0	0	0
6	F	0	0	0	0	0	0
7	F	0	0	0	0	0	0
8	F	0	0	0	0	0	0
9	F	0	0	0	0	0	0
10	M	0	0	0	0	0	0
11	F	0	0	0	0	0	0
12	M	0	0	0	0	0	0
13	F	0	0	0	0	0	0
14	M	0	0	0	0	0	0
15	F	0	0	0	0	0	0
16	F	0	0	0	0	0	0
17	F	0	0	0	0	0	0
18	M	0	0	0	0	0	0
19	F	0	0	0	0	0	0
20	F	0	0	0	0	0	0
21	F	0	0	0	0	0	0
22	F	0	0	0	0	0	0
23	F	0	0	0	0	0	0
24	F	0	0	0	0	0	0
25	F	0	0	0	0	0	0
26	F	0	0	0	0	0	0
27	F	0	0	0	0	0	0
28	F	0	0	0	0	0	0
29	M	0	0	0	0	0	0
30	M	0	0	0	0	0	0

Values and classification of the average irritation indexes

(See table no. 2)



Conclusions

The table and the graphs listed above summarize the values of the indexes of erythema and edema found out for each volunteer. We can state that product named:

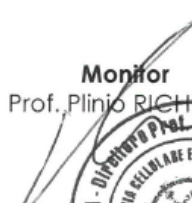
Peter Greven Physiaderm GmbH Physio UV 50 Spray RO_27_01_20_CP

has been dermatologically tested and it can be classified according to amended Draize classification :

NOT IRRITANT


Sperimentatore / Experimenter
Dott. Fernando Marco BIANCHI




Monitor
Prof. Plinio RICHELMI



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Via A. Panizzi, 10
MILANO ITALY