



**50 SUBJECT HUMAN REPEAT INSULT PATCH TEST FOR SENSITIVE SKIN
FOR SKIN IRRITATION AND SKIN SENSITIZATION EVALUATION**

Date: October 11, 2023

Revision #1 Date: October 13, 2023

Study No.: Panel 13

Sponsor: Center for Beauty Salon Supply
9985 82nd Way
Seminole, FL 33777

1.0 Objective: To determine the irritation and sensitization (contact allergy) potential of a test material after repeated application to the skin of human subjects with sensitive skin.

2.0 Test Material:

2.1 Test Material Description:

Date Received: July 21, 2023

Received From: Center for Beauty Salon Supply

Number Of Test Samples Received: 14

Label On Test Samples: Sa'sHa

Accession No.: 1220486

2.2 Handling:

Upon arrival at ALS Pharmaceutical, Beauty, and Personal Care the test material was assigned a unique laboratory code number and entered into a daily log identifying the lot number, sample description, sponsor, date received and tests requested.

Samples will be retained for a period of thirty (30) days beyond submission of final report unless otherwise specified by the sponsor or, if sample is known to be in support of governmental applications, in which case representative retained samples are kept two (2) years beyond final report submission.

Sample disposition will be conducted in compliance with appropriate federal, state and local ordinances.

3.0 Panel Selection:

3.1 Standards for Inclusion in a Study:

- Individuals who were not currently under a doctor's care.
- Individuals who were free of any dermatological or systemic disorder that would interfere with the results, at the discretion of the Investigator.
- Individuals who were free of any acute or chronic disease that would interfere with or increase the risk of study participation.
- Individuals who completed a preliminary medical history form mandated by ALS and were in general good health.
- Individuals who read, understood and signed an informed consent document relating to the specific type of study.
- Individuals who were able to cooperate with the Investigator and research staff, and were willing to have test materials applied according to the protocol, and complete the full course of the study.
- Individuals with self-perceived sensitive skin.

3.2 Standards for Exclusion from a Study:

- Individuals who were under 18 years of age.
- Individuals who were currently under a doctor's care.
- Individuals who were Fitzpatrick V or VI.
- Individuals who were currently taking any medication (topical or systemic) that might mask or interfere with the test results.
- Individuals who had a history of any acute or chronic disease that might interfere with or increase the risk associated with study participation.
- Individuals who were diagnosed with chronic skin allergies.
- Female volunteers who indicated that they were pregnant or nursing.

3.3 Recruitment:

Panel selection was accomplished by advertisements in local periodicals, community bulletin boards, phone solicitation, electronic media or any combination thereof.

3.4 Informed Consent and Medical History Forms:

An informed consent was obtained from each volunteer prior to initiating the study describing reasons for the study, possible adverse effects, associated risks and potential benefits of the treatment and their limits of liability. Panelists signed and dated the informed consent document to indicate their authorization to proceed and acknowledge their understanding of the contents. Each subject was assigned a permanent identification number and completed an extensive medical history form. These forms along with the signed consent forms are available for inspection on the premises of ALS only. [Reference 21 CFR Ch. 1 Part 50, Subpart B]

The parties agree to comply with applicable state and federal privacy laws for the use and disclosure of a subject's personal health information by taking reasonable steps to protect the confidentiality of this information. This obligation shall survive the termination or expiration of this Agreement.

4.0 Population Demographics:

Number of subjects enrolled	90
Number of subjects completing study	57
Age Range	18 - 64
Sex	
Male	08
Female	49
Sensitive Skin	
Yes	57
No	0
Fitzpatrick Skin Type*	
1 – always burn, does not tan	0
2 – burn easily, tan slightly	06
3 – burn moderately, tan progressively	26
4 – burn a little, always tan	25
5 – rarely burn, tan intensely	0
6 – never burn, tan very intensely	0

*Agache P., Humbert P.. Measuring the skin. (p 473, table 48.1) Springer-Verlag Berlin Heidelberg, 2004, (p 473, table 48.1)

5.0 Equipment:

Test materials to be tested under occlusive conditions were placed on an adhesive tape with paper filter discs with 1.0 cm² (Adhesive tapes from 3M Company – Durapore (Code 1538) and Blenderm (Code 1525) or placed on an 8-millimeter aluminum Finn Chamber® (Epitest Ltd. Oy, Tuusula, Finland) supported on Scanpor® Tape (Norgesplaster A/S, Kristiansand, Norway) or an 8-millimeter filter paper coated aluminum Finn Chamber® AQUA supported on a thin flexible transparent polyurethane rectangular film coated on one side with a medical grade acrylic adhesive, consistent with adhesive used in state-of-the-art hypoallergenic surgical tapes or a 7mm IQ-ULTRA® closed cell system which is made of additive-free polyethylene plastic foam with a filter paper incorporated (It is supplied in units of 10 chambers on a hypoallergenic non-

woven adhesive tape; the width of the tape is 52mm and the length is 118mm) or other equivalents.

Test materials to be tested under semi-occlusive conditions were placed on an adhesive tape with paper filter discs with 1.0 cm² (Adhesive tapes from 3M Company – Durapore (Code 1538) or placed on a test strip with a Rayon/Polypropylene pad or on a 7.5mm filter paper disc affixed to a strip of hypoallergenic tape (Johnson & Johnson 1 inch First Aid Cloth Tape).

Test materials to be tested in an open patch were applied and rubbed directly onto the back of the subject.

Approximately 0.02-0.05 mL (in case of liquids) and/or 0.02-0.05 gm (in case of solids) of the test material was used for the study. Liquid test material was dispensed on a paper disk, which fit in the patch chamber.

6.0 Procedure:

- Subjects were requested to bathe or wash as usual before arrival at the facility.
- Patches containing the test material were then affixed directly to the skin of the intrascapular regions of the back, to the right or left of the midline and subjects were dismissed with instructions not to wet or expose the test area to direct sunlight.
- Patches remained in place for 48 hours during the week and for 72 hours during weekends.
- This procedure was repeated until a series of nine (9) consecutive, 48-hour/72-hour exposures had been made three (3) times a week for three (3) consecutive weeks.
- Prior to each reapplication, the test sites evaluated by trained laboratory personnel.
- Following a 10 day rest period a retest/challenge dose was applied once to a previously unexposed test site and left for approximately 48 hours. Test sites were evaluated by trained laboratory personnel 48 and 96 hours after application.
- In the event of an adverse reaction, the area of erythema and edema were measured. Edema is estimated by the evaluation of the skin with respect to the contour of the unaffected normal skin.
- Subjects were instructed to report any delayed reactions that might occur after the final reading.

- Clients will be notified immediately in the case of an adverse reaction and a determination is made as to treatment program if necessary.

7.0 Scoring:

Scoring scale and definition of symbols shown below are based on the scoring scheme according to the International Contact Dermatitis Research Group scoring scale [Rietschel, R.L., Fowler, J.F., Ed., Fisher's Contact Dermatitis (fourth ed.). Baltimore, Williams & Wilkins, 1995] listed below:

- 0** no reaction (negative)
- 1** mild erythema
- 2** clear erythema
- 3** erythema + edema + papules
- 4** erythema + edema + papules + vesicles

- D** Site discontinued
- Dc** Subject discontinued voluntarily
- Dcl** Subject discontinued per Investigator

NOTE: Clinical evaluations are performed by an ALS investigator or designee trained in the clinical evaluation of the skin. Whenever feasible, the same individual will do the scoring of all the subjects throughout the study and will be blinded to the treatment assignments and any previous scores.

8.0 Results:

Accession No.: 1220486

Test Material Description: Sa'sHa

Patch Description: Semi-Occlusive (Diluted to 3% in DI Water)

Subject Information					Induction									Challenge	
No.	Sex	Age	Skin Type	Sensitive Skin	1	2	3	4	5	6	7	8	9	1	2
001	F	45	4	Yes	0	0	0	0	0	0	0	0	0	0	0
002	F	32	3	Yes	0	0	0	0	0	0	0	0	0	0	0
003	F	45	3	Yes	0	0	0	0	0	0	0	0	0	0	0
004	F	42	4	Yes	0	0	0	0	0	0	0	0	0	0	0
005	F	55	2	Yes	0	0	0	0	0	0	0	0	0	0	0
006	F	28	3	Yes	0	0	0	0	0	0	0	0	0	0	0
007	F	62	4	Yes	0	0	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc
008	F	55	3	Yes	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc
009	F	54	4	Yes	0	0	0	0	0	0	0	0	0	0	0
010	F	26	3	Yes	0	0	0	0	0	0	0	0	0	0	0
011	F	43	3	Yes	0	0	0	0	0	0	0	0	0	0	0
012	F	40	4	Yes	0	0	0	0	0	0	0	0	0	0	0
013*	F	32	3	Yes	0	0	0	0	0	0	Dcl	Dcl	Dcl	Dcl	Dcl
014	F	56	4	Yes	0	0	0	0	0	0	0	0	0	0	0
015*	M	47	3	Yes	0	0	0	Dcl	Dcl	Dcl	Dcl	Dcl	Dcl	Dcl	Dcl
016	F	53	2	Yes	0	0	0	0	0	0	0	0	0	0	0
017	M	28	4	Yes	0	0	0	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc

* Dropped due to study criteria

Subject Information					Induction									Challenge	
No.	Sex	Age	Skin Type	Sensitive Skin	1	2	3	4	5	6	7	8	9	1	2
018	F	28	4	Yes	0	0	0	0	Dc	Dc	Dc	Dc	Dc	Dc	Dc
019	F	24	2	Yes	0	0	0	0	0	0	0	0	0	0	0
020	F	43	4	Yes	0	0	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc
021	F	41	3	Yes	0	0	0	0	0	0	0	0	0	0	0
022	F	26	4	Yes	0	0	0	0	0	0	0	0	0	0	0
023	F	63	2	Yes	0	0	0	0	0	0	0	0	0	0	0
024	M	25	3	Yes	0	0	0	0	0	0	0	0	0	0	0
025	F	44	4	Yes	0	0	0	0	0	0	0	0	0	0	0
026	F	31	4	Yes	0	0	0	0	Dc	Dc	Dc	Dc	Dc	Dc	Dc
027	F	31	4	Yes	0	0	0	0	0	0	Dc	Dc	Dc	Dc	Dc
028*	F	44	3	Yes	0	0	0	0	Dcl	Dcl	Dcl	Dcl	Dcl	Dcl	Dcl
029	F	32	4	Yes	0	0	0	0	0	0	0	0	0	0	0
030	F	50	4	Yes	0	0	0	0	0	0	0	0	0	0	0
031	F	27	3	Yes	0	0	0	0	0	0	0	0	0	0	0
032	F	44	4	Yes	0	0	0	0	0	0	0	0	0	0	0
033	F	62	4	Yes	0	0	0	0	Dc	Dc	Dc	Dc	Dc	Dc	Dc
034	F	59	3	Yes	0	0	0	0	0	0	0	0	0	0	0
035	F	52	3	Yes	0	0	0	0	0	0	0	0	0	0	0
036	M	58	3	Yes	0	0	0	0	0	0	0	0	0	0	0
037*	F	56	2	Yes	0	0	0	Dcl	Dcl	Dcl	Dcl	Dcl	Dcl	Dcl	Dcl
038	F	54	3	Yes	0	0	0	0	0	0	0	0	0	0	0

* Dropped due to study criteria

Subject Information					Induction									Challenge	
No.	Sex	Age	Skin Type	Sensitive Skin	1	2	3	4	5	6	7	8	9	1	2
039	F	49	3	Yes	0	0	0	0	0	Dc	Dc	Dc	Dc	Dc	Dc
040	M	39	4	Yes	0	0	0	0	0	0	0	0	0	0	0
041	M	29	4	Yes	0	0	0	0	0	0	0	0	0	0	0
042	F	53	4	Yes	0	0	0	0	0	0	0	0	0	0	0
043	F	25	4	Yes	0	0	0	0	0	0	0	0	0	0	0
044	F	39	3	Yes	0	0	0	0	0	0	0	0	0	0	0
045	F	37	3	Yes	0	0	0	0	0	0	0	0	0	0	0
046	F	24	4	Yes	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc
047	M	23	3	Yes	0	0	0	0	0	0	0	0	0	0	0
048	F	45	3	Yes	0	0	0	0	0	0	0	0	0	0	0
049	F	20	4	Yes	0	0	0	0	0	0	0	0	0	0	0
050	F	42	3	Yes	0	0	0	0	0	0	0	0	0	0	0
051	F	59	2	Yes	0	0	0	0	0	0	0	0	0	0	0
052	F	60	2	Yes	0	0	0	0	0	0	0	0	0	0	0
053	F	22	4	Yes	0	0	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc
054	F	25	4	Yes	0	0	0	0	0	0	0	0	0	0	0
055	F	19	4	Yes	0	0	0	0	0	0	0	0	0	0	0
056	F	33	4	Yes	0	0	0	0	0	0	0	0	0	0	0
057	F	33	4	Yes	0	0	0	0	0	0	0	0	0	0	0
058	F	55	3	Yes	0	0	0	0	0	0	0	0	0	0	0
059	F	48	4	Yes	0	0	0	0	0	0	0	0	0	0	0

* Dropped due to study criteria

Subject Information					Induction									Challenge	
No.	Sex	Age	Skin Type	Sensitive Skin	1	2	3	4	5	6	7	8	9	1	2
060	F	47	4	Yes	0	0	0	0	0	0	0	0	0	0	0
061	F	32	4	Yes	0	0	0	0	0	0	0	0	0	0	0
062	F	44	3	Yes	0	0	0	0	Dc	Dc	Dc	Dc	Dc	Dc	Dc
063	F	22	4	Yes	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc
064	M	60	3	Yes	0	0	0	0	0	0	0	0	0	0	0
065	M	25	3	Yes	0	0	0	0	0	0	0	0	0	0	0
066	F	18	4	Yes	0	0	0	0	0	0	0	0	0	0	0
067	F	29	3	Yes	0	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc
068	F	46	3	Yes	0	0	0	0	0	0	0	0	0	0	0
069	F	48	3	Yes	0	0	0	0	0	0	0	0	0	0	0
070	F	51	3	Yes	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc
071	F	21	4	Yes	0	0	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc
072	M	46	3	Yes	0	0	0	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc
073	F	42	3	Yes	0	0	0	0	Dc	Dc	Dc	Dc	Dc	Dc	Dc
074	F	58	3	Yes	0	0	0	0	0	Dc	Dc	Dc	Dc	Dc	Dc
075	M	18	3	Yes	0	0	0	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc
076	F	19	4	Yes	0	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc
077	F	35	3	Yes	0	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc
078	F	63	3	Yes	0	0	0	0	0	0	0	0	0	0	0
079	M	23	3	Yes	0	0	0	0	0	0	0	0	0	0	0
080	F	27	3	Yes	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc
081	F	41	4	Yes	0	0	0	0	0	0	0	0	0	0	0
082*	F	20	2	Yes	Dcl	Dcl	Dcl	Dcl	Dcl	Dcl	Dcl	Dcl	Dcl	Dcl	Dcl

* Dropped due to study criteria

Subject Information					Induction									Challenge	
No.	Sex	Age	Skin Type	Sensitive Skin	1	2	3	4	5	6	7	8	9	1	2
083	M	30	3	Yes	0	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc	Dc
084	F	32	4	Yes	0	0	0	0	0	0	Dc	Dc	Dc	Dc	Dc
085	F	40	3	Yes	0	0	0	0	0	0	0	0	0	Dc	Dc
086	F	30	4	Yes	0	0	0	0	0	0	0	0	0	Dc	Dc
087	F	64	3	Yes	0	0	0	0	0	0	0	0	0	0	0
088	F	31	3	Yes	0	0	0	0	0	0	0	0	0	0	0
089	F	26	4	Yes	0	0	0	0	0	0	0	0	Dc	Dc	Dc
090	F	37	4	Yes	0	0	0	0	0	0	0	0	0	0	0

* Dropped due to study criteria

9.0 Evaluation Period:

The study was conducted from August 28, 2023 to October 06, 2023.

10.0 Observations:

There were four non-product related adverse reactions reported during the course of this study.

Adverse Event #1: Subject 01ZHG (013), a 32 years old female, came into the facility on 13september2023 for visit 8 presenting erythema on the adhesive application area, non related to the product. Sub-investigator determined that she presented an individual predisposition related to irritation to adhesive tape (patch test) and decided to discontinue the subject from the study. Non medical attention was needed.

Adverse Event #2: Subject 028QY (015), a 47 years old male, came into the facility on 06september2023 for visit 5 presenting redness and burning on the adhesive application area, non related to the product. Sub-investigator determined that he presented an individual predisposition related to irritation to adhesive tape (patch test) and decided to discontinue the subject from the study. Non medical attention was needed.

Adverse Event #3: Subject 06JY2 (028), a 44 years old female, came into the facility on 08september2023 for visit 6 presenting erythema and abrasions on the adhesive application area, non related to the product. Sub-investigator determined that she presented an individual predisposition related to irritation to adhesive tape (patch test) and decided to discontinue the subject from the study. Non medical attention was needed.

Adverse Event #4: Subject 34R3K (082), a 20 years old female, came into the facility on 30august2023 for visit 2 presenting erythema on the adhesive application area, non related to the product. Sub-investigator determined that she presented an individual predisposition related to irritation to adhesive tape (patch test) and decided to discontinue the subject from the study. Non medical attention was needed.

11.0 Study Archives:

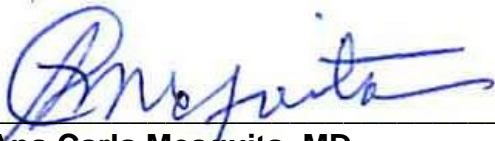
All original samples, raw data sheets, technician's notebooks, correspondence files and copies of final reports and remaining specimens will be maintained on premises of ALS in limited access storage files marked "Archive".

12.0 Conclusions:


The test product was dermatologist tested and under the conditions of the study, there was no indication of a potential to elicit dermal irritation or sensitization (contact allergy) on individuals with self-perceived sensitive skin, noted for Sa'sHa, Accession No. 1220486.

REPORT REVISION HISTORY

Revision Number	Date of Revision	Revision Description
1	13 Oct 2023	Dermatologist signature was added.



Dr. Ana Carla Mesquita, MD
Consulting Dermatologist



Vivian Pessoto Rosa
Principal Investigator