

Cosmetic Product Safety Report

SQT Revitalizing Beauty Set—SQT Revitalizing Hydrolyzed Sponge Powder & SQT Fibronectin Repair Essence

This safety assessment relates to the formulation described below. If the information below is incorrect, please amend and resubmit for reassessment.

Hunan Sunshine Bio-Tech Co., Ltd
 Building E7, Lugu Yuyuan, No.27, Wenxuan Road,
 High-Tech Development Zone, Changsha, Hunan,
 China, 410000

Formulation Ref: N/A
Buyer/Final Retailer: N/A
Manufacturer: N/A

PRODUCT FORMULATION

The chemical names shown below refer to the raw materials used to formulate this product. The identity of the raw materials is not necessarily based on the International Nomenclature of Cosmetic Ingredients (INCI) and does not represent the INCI listing that must be shown on the product label and is for assessment purposes only. An outline INCI label can be prepared on request.

Chemical Name	Conc %	% Max % Max Active		CAS No	Einecs No
		Active	in Product		
AQUA	76.114	100	76.114	7732-18-5	231-791-2
HYDROLYZED SPONGE	14	100	14	-	-
GLYCERIN	8.572	100	8.572	56-81-5 / 8013-25-0	200-289-5
1,2-HEXANEDIOL	0.257	100	.257	6920-22-5	230-029-6
BETA-GLUCAN	0.257	100	.257	26874-89-5 / 53238-80-5 / 55965-23-6	258-443-2/ 310-127-6
HYDROXYACETOPHENONE	0.171	100	.171	99-93-4	202-802-8 (I)
SODIUM POLYGLUTAMATE	0.171	100	.171	28829-38-1	POLYMER
CALCIUM SILICATE	0.143	100	.143	1344-95-2	215-710-8
SODIUM SILICATE	0.143	100	.143	1344-09-8	215-687-4
CAPRYLHYDROXAMIC ACID	0.043	100	.043	7377-03-9	230-936-7
ETHYLHEXYLGLYCERIN	0.043	100	.043	70445-33-9	408-080-2
HYDROLYZED SODIUM HYALURONATE	0.043	100	.043	-	-
PROPYLENE GLYCOL	0.043	100	.043	57-55-6	200-338-0

Remark: Mix repair essence with sponge powder in 1:5.
 MSDS/SDS, CoA/TDS and toxicity profile (where applicable) were listed in the Appendix 1

LABELLED WARNINGS & INSTRUCTIONS OF USE

Keep away from eyes.



CONSUMER EXPOSURE

Product Class: Face serum

IFRA Product type: Women's Facial Creams / Lotions / Butter / Make-up of all types

IFRA Category: Category 5

Targeted Population: Children 14 years of age 50.4kg (Mean)

Amount per application/g:

Number of applications per day: Twice a day

Skin Surface Area of Application/cm²: 555

Physical form: Liquid

Total Amount applied per day/g: 1.54

Part of body exposed to undiluted product: Hands and face

Estimated Daily Exposure mg/kg/day: 24.14

Amount Per Unit Area of Skin per day mg/cm²/day: 2.70

Retention factor: 1.00

Exposure Time Neat: 720-960 Minutes

Exposure Time Dilute: Not Applicable

Exposure time Solvent Inhalation: Not Applicable

Exposure time Aerosol Inhalation: Not Applicable

MICROBIOLOGICAL QUALITY

To comply with "Guidelines on Microbiological Quality of the Finished Cosmetic Product" under SCCS's Notes of Guidance, the following limits apply to this product:

Category 2: Other cosmetic products. Total Aerobic Mesophilic Microorganisms (Bacteria plus yeast and mould) should not exceed 1000 cfu/g or ml in the product. Escherichia coli, Pseudomonas aeruginosa and Staphylococcus aureus must not be detectable in the cosmetic product

Microbiological specifications for the product have been supplied and meet the requirements specified therein for this type of product.

Microbiological specification test report or data was listed in the Appendix 2-1

This product is a single use cosmetic product which can be used only once and is then disposed of. Preservative challenge testing is therefore not applicable to the safety of this product and only microbiological quality testing of the finished cosmetic product is required.

STABILITY OF COSMETIC PRODUCT

It is assumed that the responsible person has selected all pertinent criteria required to evaluate the stability of this cosmetic product during reasonable foreseeable conditions of storage. The stability report provided by the supplier and based upon the conclusions made therein, this cosmetic product appears to be stable under reasonably foreseeable storage conditions.

Stability Test Report or Data of Cosmetic Product was listed in the Appendix 3

PACKAGING COMPATIBILITY

It is assumed that the responsible person has identified the most applicable testing required to determine the packaging stability and its interaction with the cosmetic product contained within it. Taking into consideration the information supplied to the assessor, there appears to be no immediate health concern due to the characteristics of packaging materials in direct contact with the final product.

Package Compatibility Test Report and/or data was listed in the Appendix 4

SERIOUS / UNDESIRABLE EFFECTS

The supplier has confirmed that no undesirable effects or serious undesirable effects of this cosmetic product have been reported or, where relevant, cosmetic products with a similar formulation.

Serious/ Undesirable Effects of Cosmetic Product Declaration was listed in Appendix 5

FRAGRANCE COMPOSITIONS

This formulation does not contain a synthetic fragrance and therefore a fragrance safety evaluation as per IFRA code of practice is not applicable to this product.

According to Cosmetic Regulation (EC) No. 1223/2009, a complete cosmetic product safety report shall, at a minimum, contain cosmetic product safety information and cosmetic product safety assessment. The former includes raw materials and packaging material specifications, toxicological profile of substance (taking into account of purity of substance) contained in the cosmetic product and any known undesirable effects of the cosmetic product.



TOXICOLOGICAL & REGULATORY REVIEW

The product is mix repair essence with sponge powder including solvent, skin conditioner, moisturizer for facial skin care by people aged 14 and above. None of the ingredients is on the prohibition list of the Cosmetic Regulation (EC) No. 1223/ 2009. Due to the absence of preservatives, production according to Good Manufacturing Practices is considered appropriate to minimize the risk of microbiological hazard.

Most of the ingredients are commonly used in cosmetic products and reviewed by CIR Panel. Based on the available NOAEL, the lowest MoS is more than 100 from hydroxyacetophenone. In addition, CIR confirmed that 1,2-hexanediol, sodium silicate, caprylhydroxamic acid, hydrolyzed sodium hyaluronate are safe for use at the current level. sodium polyglutamate is a synthetic polymer formed by the polymerization of glutamic acid, used as a skin conditioner, this polymer if a naturally occurring amino acid is not expected to present any risks to health when used in cosmetics. Calcium silicate may cause physical irritation to the eyes, nose and upper respiratory tract, as well dryness to the skin following prolonged contact, but the safe concern is not expected due to its low concentration.

According to above information, there is no safety concern for the ingredients used in this product. However, the possibility cannot be discounted that a small number of users may experience an allergic reaction or other idiosyncratic reaction to an ingredient in the formulation if they have been previously sensitized to the ingredient.

Limits of heavy metals were satisfied legal requirements and testing report was listed in the Appendix 7

Effects of the product as supplied on the skin

The formulation as supplied may cause slight skin irritation especially if exposure is prolonged and/or repeated.

There are low concentrations of substances present in this product which have allergenic activity. The concentrations present are sufficiently low for the level of use to ensure that people do not become sensitised. However, people who are already sensitised to a substance may react adversely to any product containing that substance even when present at extremely low concentrations.

Exposure to this product is unlikely to result in phototoxic effects.

Unlikely to cause damage to internal organs following absorption through the skin.

Effects of the product as supplied on the eye

Accidental exposure of the eye to the formulation as supplied may result in slight eye irritation.

Effects following ingestion of the product as supplied

The formulation as supplied if swallowed may cause slight, transient irritation to the mouth and upper digestive tract.

Effects of inhaling the product

Inhalation is an unlikely route of exposure

Overall Assessment Conclusion

The ingredients are legally permitted as per Cosmetic Regulation (EC) No 1223/2009 and its amendments and the safety assessment has been carried out in accordance to this regulation. They must comply with the relevant purity standards for cosmetic ingredients. It is assumed that these ingredients do not contain any undisclosed impurities/contaminants that would affect the conclusions reached. The product must be manufactured in accordance with EU Guidance on Good Manufacturing Practice.

Keep away from eyes.

Under normal or reasonably foreseeable conditions of use, product made to this formulation is unlikely to produce an abnormally high number of adverse reactions.

Leshuai Zhang, Professor, PhD, DABT, ERT, UKRT

Intertek GM Testing Services Zhuhai Co. Ltd.

6/F, R&D and Testing/B, Guangdong-Macau TCM Park commercial Service center, 2522 Huan Dao Bei Road, Hengqin New Area, Zhuhai, China

Date: 23 Nov 2022



SUMMARY OF TOXICOLOGICAL INFORMATION OF SUBSTANCES

Remark: Substance toxicological summary was listed in this section and used only for MoS calculation, please refer to Toxicological Profiles of Substances for full information.

Chemical Substance: AQUA

EU INCI NAME:AQUA

CAS: 7732-18-5
EINECS 231-791-2

Appearance: Liquid

Water Solubility: highly soluble

Function: Solvent

Melting Point: 0

Boiling Point: 100

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> Unclassified

EU CLP Harmonised Classification>

Unclassified

Systemic Exposure Dosage / Margin of Safety:SED Adult mg/kg bw/day: 19.53592 No NOAEL Available
SED Child mg/kg bw/day: 70.18895 No NOAEL Available
SED Baby mg/kg bw/day: 198.67044 No NOAEL Available**Toxicological Summary:**

Cosmetic function : Solvent. Simply water unlikely to cause irritation, allergy or harm. Used in many cosmetic products as a solvent and necessary to sustain biological life. The source of water should be known, monitored to GMP and either a deionised or high purity grade free from toxins, pollutants and bacteriological contamination should be used in cosmetic products.

Chemical Substance: HYDROLYZED SPONGE

EU INCI NAME:HYDROLYZED SPONGE

CAS: -
EINECS -

Function: SKIN CONDITIONING

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

EU DSD/DPD Classification>

EU CLP Harmonised Classification>

Systemic Exposure Dosage / Margin of Safety:SED Adult mg/kg bw/day: 3.59333 No NOAEL Available
SED Child mg/kg bw/day: 12.91017 No NOAEL Available
SED Baby mg/kg bw/day: 36.54237 No NOAEL Available**Toxicological Summary:**

Hydrolyzed Sponge is the hydrolysate of Sponge (q.v.) obtained by acid, enzyme or other method of hydrolysis.

Chemical Substance: GLYCERIN

EU INCI NAME:GLYCERIN

CAS: 56-81-5 / 8013-25-0
EINECS 200-289-5

Appearance: liquid

Log Kow: -1.76

Water Solubility: miscible with water

Function: Denaturant / Humectant / Perfuming / Solvent / Fragrance
Ingredient / Hair & Skin Conditioning Agent / Oral Care Agent
/ Skin Protectant / Viscosity Decreasing Agent

Melting Point: ~18°C

Boiling Point: 290°C

Vapour Pressure: <0.01 mm Hg @ 20°C

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> unclassified

EU CLP Harmonised Classification>

unclassified

Systemic Exposure Dosage / Margin of Safety:SED Adult mg/kg bw/day: 2.20014 MoS - Adult 60kg: 2081.6 NOAEL mg/kg bw day: 4580
SED Child mg/kg bw/day: 7.90471 MoS - Child 16.7kg: 579.4 NOAEL test method: 90-day oral
SED Baby mg/kg bw/day: 22.37437 MoS - Baby 5.9kg: 204.6**Toxicological Summary:**

The ingredient is not acutely toxic, a skin irritant, an eye irritant, a skin sensitizer, mutagenic, carcinogenic, a reproductive toxicant, nor is it bioaccumulative. No information was available on its phototoxicity. Based on this information and other scientific literature on this ingredient, safety concerns are not expected with this ingredient for use in cosmetics.

Chemical Substance: 1,2-HEXANEDIOL

EU INCI NAME:1,2-HEXANEDIOL

CAS: 6920-22-5
EINECS 230-029-6

Function: Solvent

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> Unclassified

EU CLP Harmonised Classification>

Systemic Exposure Dosage / Margin of Safety:SED Adult mg/kg bw/day: 0.06596 No NOAEL Available
SED Child mg/kg bw/day: 0.23699 No NOAEL Available
SED Baby mg/kg bw/day: 0.67081 No NOAEL Available**Toxicological Summary:**

A diol alcohol, Hexane diol has the formula $\text{CH}_3(\text{CH}_2)_3\text{CH}_2\text{CH}(\text{OH})\text{CH}_2\text{OH}$. This alcohol is widely used in cosmetic products and incorporation into skin formulations will be uneventful.

**Chemical Substance: BETA-GLUCAN**

EU INCI NAME: BETA-GLUCAN

CAS: 26874-89-5 / 53238-80-5 / 55965-23-6

EINECS 258-443-2 / 310-127-6

Function: Skin conditioning agent

Boiling Point: 865.2 °C at 760 mmHg

Appearance: powder

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> Unclassified

EU CLP Harmonised Classification> Unclassified

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.06596

MoS - Adult 60kg: 113699.5

NOAEL mg/kg bw day: 7500

SED Child mg/kg bw/day: 0.23699

MoS - Child 16.7kg: 31646.3

NOAEL test method: 99-114 wks in mice by oral

SED Baby mg/kg bw/day: 0.67081

MoS - Baby 5.9kg: 11180.4

Toxicological Summary:

Cosmetic function: skin conditioning. A polysaccharide consisting of b(1-3) linked glucose chains carrying b(1-6) linked glucose sidechains. Used to enhance the immune system and to lower blood cholesterol levels. When use in cosmetic products should be uneventful.

Chemical Substance: HYDROXYACETOPHENONE

EU INCI NAME: HYDROXYACETOPHENONE

CAS: 99-93-4

EINECS 202-802-8 (I)

Melting Point: 109 °C (REACH Dossiers, 2017)

Boiling Point: the normal boiling temperature could not be determined

Appearance: solid (REACH Dossiers, 2017)

Water Solubility: 10 g/L at 22 °C

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> Unclassified

EU CLP Harmonised Classification> Unclassified

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.04389

MoS - Adult 60kg: 1025.2

NOAEL mg/kg bw day: 45

SED Child mg/kg bw/day: 0.15768

MoS - Child 16.7kg: 285.3

NOAEL test method: 90 day to rats by oral

SED Baby mg/kg bw/day: 0.44633

MoS - Baby 5.9kg: 100.8

Toxicological Summary:

The ingredient is not acutely toxic, an eye irritant, a skin sensitizer, mutagenic, a reproductive toxicant, but it is an eye irritant. No safety concern at current levels of intake when used as a flavouring agent by JECFA (JECFA, 2017). Based on this information and other scientific literature on this ingredient, safety concerns are not expected with this ingredient for use in cosmetics when formulated to be non-irritating.

Chemical Substance: SODIUM POLYGLUTAMATE

EU INCI NAME: SODIUM POLYGLUTAMATE

CAS: 28829-38-1

EINECS polymer

Function: humectants

EU DSD/DPD Classification>

EU CLP Harmonised Classification>

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.04389

No NOAEL Available

SED Child mg/kg bw/day: 0.15768

No NOAEL Available

SED Baby mg/kg bw/day: 0.44633

No NOAEL Available

Toxicological Summary:

Synthetic polymer formed by the polymerization of glutamic acid. Used as a skin and hair conditioning agent this polymer if a naturally occurring amino acid is not expected to present any risks to health when used in cosmetics.

Chemical Substance: CALCIUM SILICATE

EU INCI NAME: CALCIUM SILICATE

CAS: 1344-95-2

EINECS 215-710-8

Function: absorbents / opacifiers / viscosity controlling agents

Cosmetic Regulatory Summary:

EU Cosmetics Status:

Regulatory Summary:

EU DSD/DPD Classification> unclassified

EU CLP Harmonised Classification>

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.03670

No NOAEL Available

SED Child mg/kg bw/day: 0.13186

No NOAEL Available

SED Baby mg/kg bw/day: 0.37325

No NOAEL Available

Toxicological Summary:

This material may cause physical irritation to the eyes, nose and upper respiratory tract, as well dryness to the skin following prolonged contact. However the material is of a size unlikely to be inhaled.

**Chemical Substance: SODIUM SILICATE**

EU INCI NAME: SODIUM SILICATE

CAS: 1344-09-8

EINECS 215-687-4

Cosmetic Regulatory Summary:**Regulatory Summary:**

EU DSD/DPD Classification> R36-37/38

EU CLP Harmonised Classification>

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.03670	No NOAEL Available
SED Child mg/kg bw/day: 0.13186	No NOAEL Available
SED Baby mg/kg bw/day: 0.37325	No NOAEL Available

Toxicological Summary:

A strongly alkaline material used to adjust the pH of products. As supplied irritating to skin and eyes.

A final assessment have been conducted on Potassium Silicate, Sodium Metasilicate, and Sodium Silicate.

These are used as corrosion inhibitors in cosmetics and also as chelators and buffer. It is reported that sodium metasilicate is currently used at concentrations ranging from 13-18% and sodium silicate 0.3-55%.

The silicate are also used in industrial cleaners such as detergents including laundry detergents. The corrosive properties of sodium silicate is determined by the molar ratio of SiO₂:Na₂O with the higher molar ratios being less alkaline. The toxicity is also related to the molar ratio.

Toxicological endpoints: LD50 (rat, oral) 847mg/kg-1349.3mg/kg (Sodium metasilicate).

Lesions reported in the oral cavity, pharynx, esophagus, stomach, larynx, lungs, and kidneys of dogs receiving 0.25 g/kg or more of a detergent containing sodium metasilicate.

Rats administered 464mg/kg of a 20% solution of varying ratios showed no signs of toxicity. 1000mg/kg and 2150mg/kg showed signs of gasping, dyspnea, and acute depression. Gross lesions observed in dogs (2.4 g/kg/day).

Dermal irritation ranged from negligible to severe depending on molar ratio and test species. Non sensitizing to the skin (LLNA) but delayed hyper sensitivity in mice.

Eye irritation (Potassium silicate) -: non irritating (rabbit). Sodium Metasilicate (42.4% H₂O) - corrosive (rabbit). Overall the silicate ranged from severely irritating to the eye to non irritating in some studies.

Mutagenicity: (Sodium Metasilicate): non mutagenic in bacterial cells. Some effects observed with a reduced number of offspring in rats when silica was administered in drinking water.

Three adult rats injected intratesticularly and subcutaneously with 0.8 mM/kg of Sodium Silicate showed no morphological changes in the testes and no effect on the residual spermatozoa in the ductus deferens.

Human studies: (Sodium Metasilicate 37%) Effects of skin irritation observed on intact and abraded skin. Sodium silicate (6-13%) - non irritating to human skin. Also negative in HRIPT (10% of a 40% solution in water) but showed irritation in a cumulative study under normal use conditions.

The CIR panel supported their use in cosmetic products when formulated to reduce the effects of irritation whilst considering they already have GRAS status and the limited dermal absorption.

References:

Int J Toxicol. 2005;24 Suppl 1:103-17.

CIR Compendium 2010

Chemical Substance: CAPRYLHYDROXAMIC ACID

EU INCI NAME: CAPRYLHYDROXAMIC ACID

CAS: 7377-03-9

EINECS 230-936-7

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

EU DSD/DPD Classification>

EU CLP Harmonised Classification>

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.01103	No NOAEL Available	NOAEL mg/kg bw day: -
SED Child mg/kg bw/day: 0.03965	No NOAEL Available	
SED Baby mg/kg bw/day: 0.11223	No NOAEL Available	

Toxicological Summary:

Function: CHELATING. At a low concentration used in cosmetic products, not expected to pose an adverse risk to health.

Chemical Substance: ETHYLHEXYLGLYCERIN

EU INCI NAME: OCTOXYGLYCERIN

CAS: 70445-33-9

EINECS 408-080-2

Function: Skin conditioning agent/ preservative

Appearance: Solid

Log Kow: 2.4 +/- 0.55

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> R41-52/53

EU CLP Harmonised Classification>

Eye Dam. 1

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.01103	MoS - Adult 60kg: 4530.3	NOAEL mg/kg bw day: 50
SED Child mg/kg bw/day: 0.03965	MoS - Child 16.7kg: 1260.9	NOAEL test method: subchronic oral toxicity study
SED Baby mg/kg bw/day: 0.11223	MoS - Baby 5.9kg: 445.4	

Toxicological Summary:

This ingredient is not acutely toxic. May cause mild skin irritation. Undiluted ethylhexylglycerin causes serious eye damage; 5% aqueous solution of ethylhexylglycerin was mildly irritating to eyes. It is not sensitizing, mutagenic or reproductive toxic.

**Chemical Substance: HYDROLYZED SODIUM HYALURONATE**CAS: -
EINECS -**Cosmetic Regulatory Summary:**

EU Cosmetics Status: Not controlled

EU DSD/DPD Classification>

EU CLP Harmonised Classification>

Systemic Exposure Dosage / Margin of Safety:SED Adult mg/kg bw/day: 0.01103 No NOAEL Available
SED Child mg/kg bw/day: 0.03965 No NOAEL Available
SED Baby mg/kg bw/day: 0.11223 No NOAEL Available**Toxicological Summary:****Chemical Substance: PROPYLENE GLYCOL**

EU INCI NAME: PROPYLENE GLYCOL

CAS: 57-55-6
EINECS 200-338-0Function: Humectant/Solvent
Skin Conditioning/Viscosity Controlling

Appearance: liquid

Log Kow: -0.78

Water Solubility: miscible

Melting Point: -60°C

Boiling Point: 187°C

Vapour Pressure: 0.07 mm/Hg

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> unclassified

EU CLP Harmonised Classification>

unclassified

Systemic Exposure Dosage / Margin of Safety:SED Adult mg/kg bw/day: 0.00630 MoS - Adult 60kg: 269758.3 NOAEL mg/kg bw day: 1700
SED Child mg/kg bw/day: 0.02264 MoS - Child 16.7kg: 75082.7 NOAEL test method: Chronic oral Toxicity to rat
SED Baby mg/kg bw/day: 0.06408 MoS - Baby 5.9kg: 26526.2**Toxicological Summary:**

The ingredient is not acutely toxic, mutagenic, a reproductive toxicant, and is not carcinogenic. It is not a dermal irritant based on in vivo animal tests and clinical trials with human subjects. It causes minimal eye irritation according to OECD 405 test. Based on this information and other scientific literature on this ingredient, safety concerns are not expected with this ingredient for use in cosmetics when formulated to be non-irritating.

Note: In the absence of NOAEL data, the Margin of Safety (MoS) has not been calculated. Unless otherwise determined and in the absence of literature or other experimental data, a Dermal Absorption (Dap) of 100% is taken as the worst case scenario. NOAEL: No Observed Adverse Effect Level; MoS: Margin of Safety; SED Systemic Exposure Dosage
Calculation of Margin of Safety: MoS = NOAEL / SED

Reference for skin surface area, exposures and application quantities are derived from:

1. RIVM Report 320104001/2006
2. References cited in Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients, 2006 revision
3. Exposure factors handbook 2009 Update
4. The SCCS's Notes of Guidance for the Testing of Cosmetic Ingredients and their Safety Evaluation 11th Revision SCCS/1628/21
5. Colipa Data SCCNFP/0321/02
6. McNamara et al, Food Chem. Tox: 2007, 45, 2086
7. Loretz et al, Food Chem. Tox: 2008, 46, 1516
- N.B. Exposure times have been taken from RIVM Report 320104001/2006
8. Body weights taken from Exposure factors handbook 2009 Update and mean values have been used unless specified otherwise
9. ConsExpo database
10. New default values for the spray model, RIVM, March 2010

This formulation has been safety assessed by Intertek with reference to Articles 3 and 10 of Cosmetic Regulation (EC) No. 1223/2009. The safety assessment is based on the chemical specification and toxicological profile of the ingredients as supplied at the time of assessment. The supplier to this safety assessment is advised to ask for a new safety evaluation if any change in formulation occurs, change in raw materials used, abnormally high number of adverse events are recorded, changes in recommended uses or other circumstances that may affect the safety of this product.

The REACH dossier, IUCLID Datasheet, OECD SIDS and EU RAR often include unpublished data, not otherwise available; however, it must be noted that the authorities have issued legal disclaimers for the databases. The disclaimer, among other cautions, stressed that the data in the dossier, datasheet, data set or report are not necessarily comprehensive, complete, accurate or up-to-date. Use of the data may also be subject to copyright laws. Therefore, data from the such resource are used as supporting information.

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Cosmetic Product Safety Report

SQT Revitalizing Beauty Set-SQT Repairing Mask

This safety assessment relates to the formulation described below. If the information below is incorrect, please amend and resubmit for reassessment.

Hunan Sunshine Bio-Tech Co., Ltd
 Building E7, Lugu Yuyuan, No.27, Wenxuan Road,
 High-Tech Development Zone, Changsha, Hunan,
 China, 410000

Formulation Ref: N/A

Buyer/Final Retailer: N/A

Manufacturer: N/A

PRODUCT FORMULATION

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AQUA	84.91	100	84.91	7732-18-5	231-791-2
GLYCERIN	6.6	100	6.6	56-81-5 / 8013-25-0	200-289-5
GLYCERETH-26	3.3	100	3.3	31694-55-0	-
LACTOBACILLUS/BEAN SEED EXTRACT/SODIUM GLUTAMATE FERMENT FILTRATE	2.1	100	2.1	-	-
NIACINAMIDE	0.6	100	.6	98-92-0	202-713-4
BETAINE	0.6	100	.6	107-43-7	203-490-6
1,2-HEXANEDIOL	0.16	100	.16	6920-22-5	230-029-6
ALLANTOIN	0.16	100	.16	97-59-6	202-592-8
BUTYLENE GLYCOL	0.15	100	.15	107-88-0	203-529-7
BETA-GLUCAN	0.3	100	.3	26874-89-5 / 53238-80-5 / 55965-23-6	258-443-2 / 310-127-6
XANTHAN GUM	0.2	100	.2	11138-66-2	234-394-2
METHYLPARABEN	0.2	100	.2	99-76-3	202-785-7
ETHYLHEXYLGLYCERIN	0.1	100	.1	70445-33-9	408-080-2
HYDROXYACETOPHENONE	0.1	100	.1	99-93-4	202-802-8 (I)
SODIUM POLYGLUTAMATE	0.08	100	.08	28829-38-1	POLYMER
HYDROXYPROPYL GUAR	0.05	100	.05	68442-94-4 / 39421-75-5	270-497-9
ARGININE	0.05	100	.05	74-79-3 / 7200-25-1	200-811-1 / 230-571-3
SCHIZOPHYLLAN	0.05	100	.05	9050-67-3	-
CAPRYLYL GLYCOL	0.05	100	.05	1117-86-8	214-254-7
GLUCOSE	0.05	100	.05	50-99-7	200-075-1
CAPRYLHYDROXAMIC ACID	0.05	100	.05	7377-03-9	230-936-7
PROPYLENE GLYCOL	0.05	100	.05	57-55-6	200-338-0
CARBOMER	0.05	100	.05	54182-57-9 / 9007-20-9 / 9003-01-4 / 76050-42-5 / 9062-04-8 / 9007-16-3 / 9007-17-4	POLYMER
SODIUM HYALURONATE	0.02	100	.02	9067-32-7	-
MENTHOL	0.02	100	.02	1490-04-6 / 2216-51-5 / 89-78-1 / 15356-70-4 / 98167-53-4	216-074-4 / 218-690-9 / 201-939-0 / 239-388-3

MSDS/SDS, CoA/TDS and toxicity profile (where applicable) were listed in the Appendix 1

LABELLED WARNINGS & INSTRUCTIONS OF USE



CONSUMER EXPOSURE

Product Class: Face Mask

IFRA Product type: Facial Masks

IFRA Category: Category 5

Targeted Population: Adult Female & Adult Males Mean value 60kg

Amount per application/g: 20.00

Number of applications per day: Twice per week

Skin Surface Area of Application/cm²: 565

Physical form: Liquid

Total Amount applied per day/g: 20.00

Part of body exposed to undiluted product: Hands and face

Estimated Daily Exposure mg/kg/day: -

Amount Per Unit Area of Skin per day mg/cm²/day: 35.50

Retention factor: 1.00

Exposure Time Neat: 20 Minutes

Exposure Time Dilute: Not Applicable

Exposure time Solvent Inhalation: Not Applicable

Exposure time Aerosol Inhalation: Not Applicable

MICROBIOLOGICAL QUALITY

To comply with "Guidelines on Microbiological Quality of the Finished Cosmetic Product" under SCCS's Notes of Guidance, the following limits apply to this product:

Category 2: Other cosmetic products. Total Aerobic Mesophilic Microorganisms (Bacteria plus yeast and mould) should not exceed 1000 cfu/g or ml in the product. Escherichia coli, Pseudomonas aeruginosa and Staphylococcus aureus must not be detectable in the cosmetic product

Microbiological specifications for the product have been supplied and meet the requirements specified therein for this type of product.

Microbiological specification test report or data was listed in the Appendix 2-1

This product is a single use cosmetic product which can be used only once and is then disposed of. Preservative challenge testing is therefore not applicable to the safety of this product and only microbiological quality testing of the finished cosmetic product is required.

STABILITY OF COSMETIC PRODUCT

It is assumed that the responsible person has selected all pertinent criteria required to evaluate the stability of this cosmetic product during reasonable foreseeable conditions of storage. The stability report provided by the supplier and based upon the conclusions made therein, this cosmetic product appears to be stable under reasonably foreseeable storage conditions.

Stability Test Report or Data of Cosmetic Product was listed in the Appendix 3

PACKAGING COMPATIBILITY

It is assumed that the responsible person has identified the most applicable testing required to determine the packaging stability and its interaction with the cosmetic product contained within it. Taking into consideration the information supplied to the assessor, there appears to be no immediate health concern due to the characteristics of packaging materials in direct contact with the final product.

Package Compatibility Test Report and/or data was listed in the Appendix 4

SERIOUS / UNDESIRABLE EFFECTS

The supplier has confirmed that no undesirable effects or serious undesirable effects of this cosmetic product have been reported or, where relevant, cosmetic products with a similar formulation.

Serious/ Undesirable Effects of Cosmetic Product Declaration was listed in Appendix 5

FRAGRANCE COMPOSITIONS

This formulation does not contain a synthetic fragrance and therefore a fragrance safety evaluation as per IFRA code of practice is not applicable to this product.

According to Cosmetic Regulation (EC) No. 1223/2009, a complete cosmetic product safety report shall, at a minimum, contain cosmetic product safety information and cosmetic product safety assessment. The former includes raw materials and packaging material specifications, toxicological profile of substance (taking into account of purity of substance) contained in the cosmetic product and any known undesirable effects of the cosmetic product.



TOXICOLOGICAL & REGULATORY REVIEW

The SQT Repairing Mask contains solvent, moisturizer, skin conditioner, adhesive agent and thickener, which is used on face. None of the ingredients is on the prohibition list of the Cosmetic Regulation (EC) No. 1223/ 2009. Most of the ingredients are commonly used in cosmetic products, reviewed by CIR panel and confirmed to be safe for use at the current level of use. Ingredients with available NOAEL have MoS no less than 100, indicating their high safety. Although the MoS of niacinamide is lower than 100, it is at a concentration of 0.6% with a level within the restrictions indicated in the CIR file (3%). Glycereth-26, 1,2-hexanediol, allantoin, arginine, glucose and caprylhydroxamic acid are lack of NOAEL, while they are safe in this product within suggested concentrations from CIR documents. Lactobacillus/bean seed extract/sodium glutamate ferment filtrate is a filtrate of the fermentation product of phaseolus radiatus seed extract and sodium glutamate by a food grade bacterium. As the bean seed is well-established food material and sodium glutamate is widely used as a food additive, the filtrate is unlikely to produce adverse effects. Sodium polyglutamate is a synthetic polymer formed by the polymerization of glutamic acid, this polymer is a naturally occurring amino acid and not expected to present any risks to health when used in cosmetics. Schizophyllan obtained from S. commune is known as a water-soluble homopolysaccharide widely used in skin care products, is unlikely to produce adverse effects when added in small amounts to this product.

Methylparaben, the preservative in this product, was approved for use and at 0.2% within the restrictions under EU Regulation, in which the maximal concentration of this ingredient is 0.4%.

Most of the ingredients used to formulate this product are well known ingredients. They are present at typical concentrations where they are unlikely to cause irritation or allergy. However, the possibility cannot be discounted that a small number of users may experience an allergic reaction or other idiosyncratic reaction to an ingredient in the formulation if they have been previously sensitized to the ingredient.

Manufacturer should ensure the grade of glycerin being used containing low level of diethylene glycol impurities (e.g. pharmaceutical grade).

Limits of heavy metals were satisfied legal requirements and testing report was listed in the Appendix 7

No existing studies from human volunteers were provided.

If used as directed, use of this product should be uneventful.

Effects of the product as supplied on the skin

The formulation as supplied may cause only minimal skin irritation even if exposure is prolonged and/or repeated.

Repeated exposure to the formulation as supplied is unlikely to produce allergy by skin contact.

Exposure to this product is unlikely to result in phototoxic effects.

Unlikely to cause damage to internal organs following absorption through the skin.

Effects of the product as supplied on the eye

Accidental exposure of the eye to the formulation as supplied may result in minimal eye irritation.

Effects following ingestion of the product as supplied

The formulation as supplied if swallowed is unlikely to cause harm.

Effects of inhaling the product

Inhalation is an unlikely route of exposure

Overall Assessment Conclusion

The ingredients are legally permitted as per Cosmetic Regulation (EC) No 1223/2009 and its amendments and the safety assessment has been carried out in accordance to this regulation. They must comply with the relevant purity standards for cosmetic ingredients. It is assumed that these ingredients do not contain any undisclosed impurities/contaminants that would affect the conclusions reached. The product must be manufactured in accordance with EU Guidance on Good Manufacturing Practice.

Under normal or reasonably foreseeable conditions of use, product made to this formulation is unlikely to produce an abnormally high number of adverse reactions.

Cosmetic Regulations Product Safety Assessor

Leshuai Zhang, Toxicologist, PhD, DABT, ERT, UKRT

Intertek Testing Services Shanghai Limited

2/F, Building No. 2 Shanghai Comalong Technology Service Park, 889 Yi Shan Road, Shanghai, China

Date: 29 Nov 2022



SUMMARY OF TOXICOLOGICAL INFORMATION OF SUBSTANCES

Remark: Substance toxicological summary was listed in this section and used only for MoS calculation, please refer to Toxicological Profiles of Substances for full information.

Chemical Substance: AQUA

EU INCI NAME:AQUA

CAS: 7732-18-5

EINECS 231-791-2

Appearance: Liquid

Water Solubility: highly soluble

Function: Solvent

Melting Point: 0

Boiling Point: 100

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> Unclassified

EU CLP Harmonised Classification>

Unclassified

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 283.03333 No NOAEL Available

SED Child mg/kg bw/day: 1016.88622 No NOAEL Available

SED Baby mg/kg bw/day: 2878.30508 No NOAEL Available

Toxicological Summary:

Cosmetic function : Solvent. Simply water unlikely to cause irritation, allergy or harm. Used in many cosmetic products as a solvent and necessary to sustain biological life. The source of water should be known, monitored to GMP and either a deionised or high purity grade free from toxins, pollutants and bacteriological contamination should be used in cosmetic products.

Chemical Substance: GLYCERIN

EU INCI NAME:GLYCERIN

CAS: 56-81-5 / 8013-25-0

EINECS 200-289-5

Appearance: liquid

Log Kow: -1.76

Water Solubility: miscible with water

Function: Denaturant / Humectant / Perfuming / Solvent / Fragrance
Ingredient / Hair & Skin Conditioning Agent / Oral Care Agent
/ Skin Protectant / Viscosity Decreasing Agent

Melting Point: ~18°C

Boiling Point: 290°C

Vapour Pressure: <0.01 mm Hg @ 20°C

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> unclassified

EU CLP Harmonised Classification>

unclassified

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 22.00000 MoS - Adult 60kg: 208.1

SED Child mg/kg bw/day: 79.04191 MoS - Child 16.7kg: 57.9

SED Baby mg/kg bw/day: 223.72881 MoS - Baby 5.9kg: 20.4

NOAEL mg/kg bw day: 4580

NOAEL test method: 90-day oral

Toxicological Summary:

The ingredient is not acutely toxic, a skin irritant, an eye irritant, a skin sensitizer, mutagenic, carcinogenic, a reproductive toxicant, nor is it bioaccumulative. No information was available on its phototoxicity. Based on this information and other scientific literature on this ingredient, safety concerns are not expected with this ingredient for use in cosmetics.

Chemical Substance: GLYCERETH-26

EU INCI NAME:GLYCERETH-26

CAS: 31694- 55-0

EINECS -

Appearance: Liquid

Water Solubility: 100

Function: humectants / solvents

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> Unclassified

EU CLP Harmonised Classification>

Not controlled

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 11.00000 No NOAEL Available

SED Child mg/kg bw/day: 39.52095 No NOAEL Available

SED Baby mg/kg bw/day: 111.86440 No NOAEL Available

Toxicological Summary:

Cosmetic Functions : Humectant / Solvent / Viscosity Controlling. An ethoxylated glycerol. From its structure it is unlikely to irritate the skin and eye or cause other adverse effects.

Chemical Substance: LACTOBACILLUS/BEAN SEED EXTRACT/SODIUM GLUTAMATE FERMENT FILTRATE

CAS: -

EINECS -

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

EU DSD/DPD Classification>

EU CLP Harmonised Classification>

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 7.00000 No NOAEL Available

SED Child mg/kg bw/day: 25.14970 No NOAEL Available

SED Baby mg/kg bw/day: 71.18644 No NOAEL Available

Toxicological Summary:

Cosmetic functions: hair conditioning, skin conditioning. Lactobacillus/Bean Seed Extract/Sodium Glutamate Ferment Filtrate is a filtrate of the fermentation product of Phaseolus Radiatus Seed Extract and Sodium Glutamate by the microorganism Lactobacillus. Unlikely to cause adverse effects when used in cosmetics.



Issued: 29 Nov 2022

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Chemical Substance: NIACINAMIDE

EU INCI NAME:NIACINAMIDE

CAS: 98-92-0

EINECS 202-713-4

Appearance: Powder (CIR, 2005)

Log Kow: -0.37 (CIR, 2005)

Water Solubility: Soluble (CIR, 2005)

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> R36

EU CLP Harmonised Classification> Unclassified

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 2.00000	MoS - Adult 60kg: 35.8	NOAEL mg/kg bw day: 71.6	
SED Child mg/kg bw/day: 7.18562	MoS - Child 16.7kg: 9.9	NOAEL test method:	28 -day oral in rat
SED Baby mg/kg bw/day: 20.33898	MoS - Baby 5.9kg: 3.5		

Toxicological Summary:

Skin irritation tests of up to 2.5% Niacinamide in rabbits produced only marginal irritation. Skin sensitization tests of Niacinamide at 5% during induction and 20% during challenge were negative in guinea pigs. Neither cosmetic ingredient was mutagenic in Ames tests, with or without metabolic activation. Niacinamide and Niacin are considered as GRAS by US FDA. The CIR Expert Panel considered that Niacinamide and Niacin are sufficiently similar from a toxicologic standpoint to combine the available data and reach a conclusion on the safety of both as cosmetic ingredients at present use (Niacinamide up to 3% and Niacin up to 0.1%).

Chemical Substance: BETAINE

EU INCI NAME:BETAINE

CAS: 107-43-7

EINECS 203-490-6

Appearance: crystalline powder

Log Kow: -3.1

Function: Antistatic/Hair & Skin Conditioning
Humectant/Viscosity Controlling

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> Unclassified

EU CLP Harmonised Classification> Unclassified

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 2.00000	MoS - Adult 60kg: 25000.0	NOAEL mg/kg bw day: 50	
SED Child mg/kg bw/day: 7.18562	MoS - Child 16.7kg: 6958.3	NOAEL test method:	maternal toxicity study
SED Baby mg/kg bw/day: 20.33898	MoS - Baby 5.9kg: 2458.3		

Toxicological Summary:

Cosmetic Functions : Antistatic / Hair & Skin Conditioning / Humectant / Viscosity Controlling. This is non-irritating to the skin and has no sensitisation potential. It has hair conditioning properties and reduces the irritancy of surfactants. The ingredient is not mutagenic or genotoxic, or carcinogenic.

Chemical Substance: 1,2-HEXANEDIOL

EU INCI NAME:1,2-HEXANEDIOL

CAS: 6920-22-5

EINECS 230-029-6

Function: Solvent

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> Unclassified

EU CLP Harmonised Classification>

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.53333	No NOAEL Available
SED Child mg/kg bw/day: 1.91616	No NOAEL Available
SED Baby mg/kg bw/day: 5.42372	No NOAEL Available

Toxicological Summary:

A diol alcohol, Hexane diol has the formula $\text{CH}_3(\text{CH}_2)_3\text{CH}_2\text{CH}(\text{OH})\text{CH}_2\text{OH}$. This alcohol is widely used in cosmetic products and incorporation into skin formulations will be uneventful.

Chemical Substance: ALLANTOIN

EU INCI NAME:ALLANTOIN

CAS: 97-59-6

EINECS 202-592-8

Appearance: white odorless powder

Water Solubility: 5260 mg/L

Function: Skin conditioning agent

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> unclassified

EU CLP Harmonised Classification> unclassified

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.53333	No NOAEL Available	NOAEL mg/kg bw day: -	
SED Child mg/kg bw/day: 1.91616	No NOAEL Available	NOAEL test method:	-
SED Baby mg/kg bw/day: 5.42372	No NOAEL Available		

Toxicological Summary:

Cosmetic functions : Skin Conditioning / Skin Protecting / Soothing. Low acute toxicity and minimal potential to irritate the skin and eyes, not a skin sensitiser. Also negative Ames test data. Suppliers data also states hazardous properties are relatively improbable and no description of any toxic symptoms. Unlikely to produce any adverse effects when used at typical concentrations in cosmetic products.



Issued: 29 Nov 2022

GZHH0047424502

Chemical Substance: BUTYLENE GLYCOL

EU INCI NAME: Butylene Glycol

CAS: 107-88-0

EINECS 203-529-7

Appearance: Viscous liquid

Log Kow: -0.29

Water Solubility: miscible

Function: humectants / solvents

Melting Point: -77°C

Boiling Point: 207.5 °C

Vapour Pressure: 0.08 at 20°C

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> Unclassified

EU CLP Harmonised Classification> Unclassified

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.50000	MoS - Adult 60kg: 14285.7	NOAEL mg/kg bw day: 6000	
SED Child mg/kg bw/day: 1.79640	MoS - Child 16.7kg: 3976.1	NOAEL test method:	90-days toxicity study to dogs
SED Baby mg/kg bw/day: 5.08474	MoS - Baby 5.9kg: 1404.7		

Toxicological Summary:

The ingredient is not acutely toxic via dermal and oral route; it is not a skin irritant, a skin sensitizer, mutagenic, carcinogenic, a reproductive toxicant, or photosensitizer. Low bioaccumulation potential based on study results. Undiluted butylenes glycol was not an eye irritant to rabbits, but was to humans. Based on this information and other scientific literature on this ingredient, safety concerns are not expected with this ingredient for use in cosmetics when formulated to be non-irritating.

Chemical Substance: BETA-GLUCAN

EU INCI NAME: BETA-GLUCAN

CAS: 26874-89-5 / 53238-80-5 / 55965-23-6

EINECS 258-443-2 / 310-127-6

Appearance: powder

Function: Skin conditioning agent

Boiling Point: 865.2 °C at 760 mmHg

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> Unclassified

EU CLP Harmonised Classification> Unclassified

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 1.00000	MoS - Adult 60kg: 7500.0	NOAEL mg/kg bw day: 7500	
SED Child mg/kg bw/day: 3.59281	MoS - Child 16.7kg: 2087.5	NOAEL test method:	99-114 wks in mice by oral
SED Baby mg/kg bw/day: 10.16949	MoS - Baby 5.9kg: 737.5		

Toxicological Summary:

Cosmetic function: skin conditioning. A polysaccharide consisting of b(1-3) linked glucose chains carrying b(1-6) linked glucose sidechains. Used to enhance the immune system and to lower blood cholesterol levels. When use in cosmetic products should be uneventful.

Chemical Substance: XANTHAN GUM

EU INCI NAME: XANTHAN GUM

CAS: 11138-66-2

EINECS 234-394-2

Appearance: Cream coloured powder (JECFA, 1999; CIR, 2012)

Water Solubility: Soluble (JECFA, 1999)

Function: Binders / Emulsion stabilisers / Viscosity controlling agents

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> unclassified

EU CLP Harmonised Classification> unclassified

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.66666	MoS - Adult 60kg: 1499.9	NOAEL mg/kg bw day: 1000	
SED Child mg/kg bw/day: 2.39520	MoS - Child 16.7kg: 417.4	NOAEL test method:	CD rats 104 weeks oral
SED Baby mg/kg bw/day: 6.77966	MoS - Baby 5.9kg: 147.5		

Toxicological Summary:

The ingredient is not acutely toxic through the oral and inhalation routes. It is not an ocular or skin irritant and is not sensitizing. It is not carcinogenic, reprotoxic and does not bioaccumulate in the body. No information is readily available on the mutagenicity, dermal absorption/ percutaneous potential as well as the acute dermal toxicity of the ingredient. It should be noted that this ingredient has been approved by the EU and FDA as a food additive. Based on this information and other scientific literature on this ingredient, safety concerns are not expected with this ingredient for use in cosmetics

Chemical Substance: METHYLPARABEN

EU INCI NAME: METHYLPARABEN

CAS: 99-76-3

EINECS 202-785-7

Appearance: Powder

Log Kow: 1.87 or 1.66

Water Solubility: Slightly soluble

Function: preservatives

Cosmetic Regulatory Summary:

EU Cosmetics Status: Approved preservative

Regulatory Summary:

EU DSD/DPD Classification> R36/37/38

EU CLP Harmonised Classification> Unclassified

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.66666	MoS - Adult 60kg: 40540.5	NOAEL mg/kg bw day: 1000	
SED Child mg/kg bw/day: 2.39520	MoS - Child 16.7kg: 11283.7	NOAEL test method:	96 weeks oral in rats
SED Baby mg/kg bw/day: 6.77966	MoS - Baby 5.9kg: 3986.4		

Toxicological Summary:

The ingredient is not acutely toxic by oral or dermal administration or by inhalation. It is not a skin irritant in humans at concentrations of 5% or less, but when treated with undiluted methylparaben, mild skin irritation occurs. 100% methylparaben causes slight, transient eye irritation; however, 0.2% is non-irritating to the eyes. Methylparaben is not a skin sensitizer, a mutagen, a carcinogen, a reproductive toxicant, bioaccumulative nor phototoxic. Based on this information and other scientific literature on this ingredient, safety concerns are not expected with this ingredient for use in cosmetics when formulated to be non-irritating.



Issued: 29 Nov 2022

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Chemical Substance: ETHYLHEXYLGLYCERIN

EU INCI NAME:OCTOXYGLYCERIN
CAS: 70445-33-9
EINECS 408-080-2

Function: Skin conditioning agent/ preservative

Appearance: Solid
Log Kow: 2.4 +/- 0.55

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> R41-52/53

EU CLP Harmonised Classification-> Eye Dam. 1

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.33333	MoS - Adult 60kg: 150.1	NOAEL mg/kg bw day: 50	
SED Child mg/kg bw/day: 1.19760	MoS - Child 16.7kg: 41.7	NOAEL test method:	subchronic oral toxicity study
SED Baby mg/kg bw/day: 3.38983	MoS - Baby 5.9kg: 14.7		

Toxicological Summary:

This ingredient is not acutely toxic. May cause mild skin irritation. Undiluted ethylhexylglycerin causes serious eye damage; 5% aqueous solution of ethylhexylglycerin was mildly irritating to eyes. It is not sensitizing, mutagenic or reproductive toxic.

Chemical Substance: HYDROXYACETOPHENONE

EU INCI NAME:HYDROXYACETOPHENONE
CAS: 99-93-4
EINECS 202-802-8 (I)

Melting Point: 109 °C (REACH Dossiers, 2017)

Appearance: solid (REACH Dossiers, 2017)

Boiling Point: the normal boiling temperature could not be determined (REACH Dossiers, 2017)

Water Solubility: 10 g/L at 22 °C

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> Unclassified

EU CLP Harmonised Classification-> Unclassified

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.33333	MoS - Adult 60kg: 135.1	NOAEL mg/kg bw day: 45	
SED Child mg/kg bw/day: 1.19760	MoS - Child 16.7kg: 37.5	NOAEL test method:	90 day to rats by oral
SED Baby mg/kg bw/day: 3.38983	MoS - Baby 5.9kg: 13.2		

Toxicological Summary:

The ingredient is not acutely toxic, an eye irritant, a skin sensitizer, mutagenic, a reproductive toxicant, but it is an eye irritant. No safety concern at current levels of intake when used as a flavouring agent by JECFA (JECFA, 2017). Based on this information and other scientific literature on this ingredient, safety concerns are not expected with this ingredient for use in cosmetics when formulated to be non-irritating.

Chemical Substance: SODIUM POLYGLUTAMATE

EU INCI NAME:SODIUM POLYGLUTAMATE
CAS: 28829-38-1
EINECS polymer

Function: humectants

EU DSD/DPD Classification->

EU CLP Harmonised Classification->

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.26666	No NOAEL Available
SED Child mg/kg bw/day: 0.95808	No NOAEL Available
SED Baby mg/kg bw/day: 2.71186	No NOAEL Available

Toxicological Summary:

Synthetic polymer formed by the polymerization of glutamic acid. Used as a skin and hair conditioning agent this polymer if a naturally occurring amino acid is not expected to present any risks to health when used in cosmetics.

Chemical Substance: HYDROXYPROPYL GUAR

EU INCI NAME:HYDROXYPROPYL GUAR
CAS: 68442-94-4/ 39421-75-5
EINECS 270-497-9

Function: Emulsion stabilising/ Viscosity controlling

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> Unclassified

EU CLP Harmonised Classification-> Unclassified

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.16666	MoS - Adult 60kg: 4799.9	NOAEL mg/kg bw day: 800	
SED Child mg/kg bw/day: 0.59880	MoS - Child 16.7kg: 1336.8	NOAEL test method:	guar gum maternal toxicity study in mice
SED Baby mg/kg bw/day: 1.69491	MoS - Baby 5.9kg: 472.9		

Toxicological Summary:

The hydroxypropyl derivative of guar gum. Has low potential to cause irritancy or allergy.



Issued: 29 Nov 2022

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Chemical Substance: ARGININE

EU INCI NAME: ARGININE

CAS: 74-79-3 / 7200-25-1
EINECS 200-811-1 / 230-571-3

Function: Antistatic/Hair & Skin Conditioning

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

EU DSD/DPD Classification>

EU CLP Harmonised Classification>

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.16666 No NOAEL Available
SED Child mg/kg bw/day: 0.59880 No NOAEL Available
SED Baby mg/kg bw/day: 1.69491 No NOAEL Available

Toxicological Summary:

Cosmetic Functions : Antistatic / Hair Conditioning / Masking / Skin Conditioning. An essential amino acid with low potential to cause irritancy or toxicity. It is unlikely to give rise to adverse effects when incorporated into a product.

Chemical Substance: SCHIZOPHYLLAN

EU INCI NAME: SCHIZOPHYLLAN

CAS: 9050-67-3
EINECS -

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

EU DSD/DPD Classification>

EU CLP Harmonised Classification>

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.16666 No NOAEL Available
SED Child mg/kg bw/day: 0.59880 No NOAEL Available
SED Baby mg/kg bw/day: 1.69491 No NOAEL Available

Toxicological Summary:

Cosmetic function: humectant. Schizophyllan is a polysaccharide produced by the fungus, Schizophyllum commune. It consists of three β -(1 \rightarrow 3)-linked D-glucopyranose residues, to one of which is attached a single β -(1 \rightarrow 6)-linked D-glucopyranosyl side chain. It is widely used in skin care cosmetics because it can quickly penetrate into the skin and be recognized by receptors on skin tissue cells and immune cells.

Chemical Substance: CAPRYLYL GLYCOL

EU INCI NAME: CAPRYLYL GLYCOL

CAS: 1117-86-8
EINECS 214-254-7

Function: emollients / humectants

Appearance: liquid

Log Kow: 1.316 \pm 0.215

Water Solubility: 4.4 g/l

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> Unclassified

EU CLP Harmonised Classification>

Unclassified

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.16666	MoS - Adult 60kg: 599.9	NOAEL mg/kg bw day: 100	
SED Child mg/kg bw/day: 0.59880	MoS - Child 16.7kg: 167.1	NOAEL test method:	28 day oral study
SED Baby mg/kg bw/day: 1.69491	MoS - Baby 5.9kg: 59.1		

Toxicological Summary:

The ingredient is not acutely toxic by oral, dermal or inhalation routes. It is non to severely dermal irritating, non to severely eye irritating and non skin sensitising. Caprylyl Glycol is non mutagenic/non genotoxic, non carcinogenic, non reproductive/non developmental toxic and non phototoxic or photosensitizing. It has no bioaccumulation potential. Based on this information and other scientific literature on this ingredient, safety concerns are not expected with this ingredient for use in cosmetics.

Chemical Substance: GLUCOSE

EU INCI NAME: GLUCOSE

CAS: 50-99-7
EINECS 200-075-1

Function: humectants

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> unclassified

EU CLP Harmonised Classification>

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.16666 No NOAEL Available
SED Child mg/kg bw/day: 0.59880 No NOAEL Available
SED Baby mg/kg bw/day: 1.69491 No NOAEL Available

Toxicological Summary:

Cosmetic Functions : Humectant / Flavoring Agent / Skin Conditioning Agent. A monosaccharide sugar with minimal irritant properties. Found in nature in fruits and other food sources and is an essential carbohydrate needed as an energy source. Glucose is rapidly absorbed and metabolized. Glucose is of low toxicity (LD50 \approx 8g/kg). Glucose is non irritating and non sensitizing and ingestion is not considered a threat to health.

**Chemical Substance: CAPRYLHYDROXAMIC ACID**

EU INCI NAME:CAPRYLHYDROXAMIC ACID

CAS: 7377-03-9

EINECS 230-936-7

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

EU DSD/DPD Classification>

EU CLP Harmonised Classification>

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.16666 No NOAEL Available NOAEL mg/kg bw day: -
SED Child mg/kg bw/day: 0.59880 No NOAEL Available
SED Baby mg/kg bw/day: 1.69491 No NOAEL Available

Toxicological Summary:

Function: CHELATING. At a low concentration used in cosmetic products, not expected to pose an adverse risk to health.

Chemical Substance: PROPYLENE GLYCOL

EU INCI NAME:PROPYLENE GLYCOL

CAS: 57-55-6

EINECS 200-338-0

Function: Humectant/Solvent
Skin Conditioning/Viscosity Controlling

Appearance: liquid

Melting Point: -60°C

Log Kow: -0.78

Boiling Point: 187°C

Water Solubility: miscible

Vapour Pressure: 0.07 mm/Hg

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> unclassified

EU CLP Harmonised Classification> unclassified

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.16666 MoS - Adult 60kg: 17863.3 NOAEL mg/kg bw day: 1700
SED Child mg/kg bw/day: 0.59880 MoS - Child 16.7kg: 4971.9 NOAEL test method: Chronic oral Toxicity to rat
SED Baby mg/kg bw/day: 1.69491 MoS - Baby 5.9kg: 1756.5

Toxicological Summary:

The ingredient is not acutely toxic, mutagenic, a reproductive toxicant, and is not carcinogenic. It is not a dermal irritant based on in vivo animal tests and clinical trials with human subjects. It causes minimal eye irritation according to OECD 405 test. Based on this information and other scientific literature on this ingredient, safety concerns are not expected with this ingredient for use in cosmetics when formulated to be non-irritating.

Chemical Substance: CARBOMER

EU INCI NAME:CARBOMER

CAS: 54182-57-9 / 9007-20-9 / 9003-01-4 / 76050-42-5 /

EINECS 9062-04-8 / 9007-16-3 / 9007-17-4

Function: Thickener

polymer

Appearance: gel/powder

Cosmetic Regulatory Summary:

EU Cosmetics Status: May contain benzene whose use is prohibited by Saudi legislation. Should be analyzed to ensure that no benzene is present.

Regulatory Summary:

EU DSD/DPD Classification> unclassified

EU CLP Harmonised Classification> unclassified

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.16666 MoS - Adult 60kg: 599.9 NOAEL mg/kg bw day: 100
SED Child mg/kg bw/day: 0.59880 MoS - Child 16.7kg: 167.1 NOAEL test method: Chronic oral study
SED Baby mg/kg bw/day: 1.69491 MoS - Baby 5.9kg: 59.1

Toxicological Summary:

The ingredient is not acutely toxic by oral or dermal routes. It is considered to be acutely harmful by inhalation route. It is non to minimally skin irritating, non to moderately eye irritating, non phototoxic/non photo-allergic and has no to low potential for skin sensitization. It has a low bioaccumulation potential. No information is readily available on the ingredient's mutagenicity, carcinogenicity, reproductive/developmental toxicity or dermal/percutaneous absorption. However, it has not been identified on any positive lists as having CMR potential (substitution of carcinogens, mutagens and reproductive toxins). In addition, being a large polymer, dermal absorption should not occur. Based on this information and other scientific literature on this ingredient, safety concerns are not expected with this ingredient for use in cosmetics.

Chemical Substance: SODIUM HYALURONATE

EU INCI NAME:SODIUM HYALURONATE

CAS: 9067-32-7

EINECS -

Function: Humectant / Skin Conditioning

Appearance: powder

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> unclassified

EU CLP Harmonised Classification> unclassified

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.06666 MoS - Adult 60kg: 899.9 NOAEL mg/kg bw day: 60
SED Child mg/kg bw/day: 0.23952 MoS - Child 16.7kg: 250.5 NOAEL test method: Reproductive / Developmental Toxicity study
SED Baby mg/kg bw/day: 0.67796 MoS - Baby 5.9kg: 88.4

Toxicological Summary:

The ingredient is not acutely toxic via oral, a skin irritant, an eye irritant, a skin sensitizer, mutagenic, a reproductive toxicant, No enough information about the carcinogenic, bioaccumulative and phototoxic. Hyaluronic acid does not penetrate the skin. Based on this information and other scientific literature on this ingredient, safety concerns are not expected with this ingredient for use in cosmetics.



Issued: 29 Nov 2022

GZHH0047424502

Chemical Substance: MENTHOL

EU INCI NAME: MENTHOL

CAS: 1490-04-6 / 2216-51-5 / 89-78-1 / 15356-70-4 / 98167
EINECS -53-4
216-074-4 / 218-690-9 / 201-939-0 / 239-388-3

Function: Masking/ Refreshing / Flavour

Appearance: Solid (OECD, 2003)

Log Kow: 3.4 (measured) (OECD, 2003)

Water Solubility: 431 mg/l at 20°C (OECD, 2003)

Cosmetic Regulatory Summary:

EU Cosmetics Status: Not controlled

Regulatory Summary:

EU DSD/DPD Classification> R38

EU CLP Harmonised Classification> Unclassified

Systemic Exposure Dosage / Margin of Safety:

SED Adult mg/kg bw/day: 0.06666	MoS - Adult 60kg: 8399.9	NOAEL mg/kg bw day: 560	
SED Child mg/kg bw/day: 0.23952	MoS - Child 16.7kg: 2338.3	NOAEL test method:	13 weeks oral in rat
SED Baby mg/kg bw/day: 0.67796	MoS - Baby 5.9kg: 825.9		

Toxicological Summary:

The ingredient is not acutely toxic through the oral, dermal and inhalation routes. It is moderately irritating to the skin and is an ocular irritant. It was not sensitizing in studies using guinea pigs and mice but showed low sensitizing potential in studies with human subjects. It penetrates the skin but does not bioaccumulate in the body. The ingredient is not mutagenic, carcinogenic or a reproductive toxicant. No information is readily available on the phototoxicity of the ingredient. Based on this information and other scientific literature on this ingredient, safety concerns are not expected with this ingredient for use in cosmetics.

Note: In the absence of NOAEL data, the Margin of Safety (MoS) has not been calculated. Unless otherwise determined and in the absence of literature or other experimental data, a Dermal Absorption (DAp) of 100% is taken as the worst case scenario.
NOAEL: No Observed Adverse Effect Level; MoS: Margin of Safety; SED Systemic Exposure Dosage
Calculation of Margin of Safety: $MoS = NOAEL / SED$

Reference for skin surface area, exposures and application quantities are derived from:

1. RIVM Report 320104001/2006
2. References cited in Dermal Sensitization Quantitative Risk Assessment (QRA) For Fragrance Ingredients, 2006 revision
3. Exposure factors handbook 2009 Update
4. The SCCS's Notes of Guidance for the Testing of Cosmetic Ingredients and their Safety Evaluation 10th Revision SCCS/1501/12
5. Colipa Data SCCNFP/0321/02
6. McNamara et al, Food Chem. Tox; 2007, 45, 2086
7. Loretz et al, Food Chem. Tox; 2008, 46, 1516
- N.B. Exposure times have been taken from RIVM Report 320104001/2006
8. Body weights taken from Exposure factors handbook 2009 Update and mean values have been used unless specified otherwise
9. ConsExpo database
10. New default values for the spray model, RIVM, March 2010

This formulation has been safety assessed by Intertek with reference to Articles 3 and 10 of Cosmetic Regulation (EC) No. 1223/2009. The safety assessment is based on the chemical specification and toxicological profile of the ingredients as supplied at the time of assessment.

The supplier to this safety assessment is advised to ask for a new safety evaluation if any change in formulation occurs, change in raw materials used, abnormally high number of adverse events are recorded, changes in recommended uses or other circumstances that may affect the safety of this product.

The REACH dossier, IUCLID Datasheet, OECD SIDS and EU RAR often include unpublished data, not otherwise available; however, it must be noted that the authorities have issued legal disclaimers for the databases. The disclaimer, among other cautions, stressed that the data in the dossier, datasheet, data set or report are not necessarily comprehensive, complete, accurate or up-to-date. Use of the data may also be subject to copyright laws. Therefore, data from the such resource are used as supporting information.

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Appendixes of Cosmetic Product Safety Report

For

SQT Revitalizing Beauty Set

(Include: SQT Revitalizing Hydrolyzed Sponge

Powder,SQT Fibronectin Repair Essence,SQT Repairing

Mask)

The testing report, declaration letter, SDS/MSDS, TDS, CoA, IFRA Certificate and other supportive document listed in this appendix were provided from client and delivered to risk assessor to conduct the CPSR, it is supplier' s responsibility to make sure the accuracy of the documents.

Appendix 1- Toxicological Profiles of Substances

1. *Toxicity summary*
2. *MSDS/SDS*
3. *TDS/CoA*

Appendix 2- Microbiological Quality Test Report of Cosmetic Product

1. *Microbiological specification test report or data*
2. *Preservative challenge test report or data*

Appendix 3- Stability Test Report or Data of Cosmetic Product

Appendix 4- Packaging Compatibility Test Report and/or data

1. *Container data*
2. *Outer Packaging material*

Appendix 5- Serious/ Undesirable Effects of Cosmetic Product Declaration or Report

Appendix 6- Fragrance

1. *IFRA Certificate*
2. *MSDS/SDS*
3. *Allergen declaration*

Appendix 7- Heavy Metal Test Report of Cosmetic Product

Appendix 8- Human Volunteers Studies

1. *Human volunteers study for the cosmetic product*
2. *Human volunteers study for raw material*

Appendix 9- Assessor's credentials

Appendix 1- Toxicological Profiles of Substances

1. Toxicity summary

Substance toxicological summary was listed in this report and detailed data are stored in Intertek owned in house database, could provide on specific request.

2. MSDS/SDS

See below report(s) if available

3. TDS/CoA

See below report(s) if available



MATERIAL SAFETY DATA SHEET (SQT Revitalizing Beauty Set)

1. CHEMICAL PRODUCT AND COMPANY IDENTIFICATION

Identification of the substance or preparation:

Product Name: SQT Revitalizing Beauty Set
Use of the substance/preparation: Cosmetic additives

Company identification:

Manufactured By: Hunan Sunshine Bio-Tech Co., Ltd
Unit 1, E7 building, No. 27
Wenxuan Road, High-Tech Development Zone
Changsha 410000, P.R.of China

Phone Number: 86-731-83991999
Email: info@sunshineextract.com

2. HAZARDOUS IDENTIFICATION

Potential Acute Health Effects:

Hazardous in case of eye contact (irritant), of ingestion, of inhalation. Slightly hazardous in case of skin contact (irritant, permeator).

Potential Chronic Health Effects:

CARCINOGENIC EFFECTS: Not available.
MUTAGENIC EFFECTS: Not available.
TERATOGENIC EFFECTS: Not available.
DEVELOPMENTAL TOXICITY: Not available.

3. COMPOSITION/INFORMATION ON INGREDIENT

Chemical Identity: Hydrolyzed Sponge
Purity: 99%
ELINCS #: N/A
CAS#: 14808-60-7

4. FIRST AID MEASURES

Inhalation: Move person to fresh air immediately.
Eye Contact: Irrigate surfaces thoroughly with water
Skin Contact: Rinse areas thoroughly with water
Ingestion: Rinse mouth thoroughly with water

5. FIRE FIGHTING MEASURES

Special Fire Fighting Procedures: Ordinary extinguishing process can be taken in case of fire.

Extinguishing Media: No prohibited media.

Protection for the person-related fire fighting: Wear or use normal protective equipment. No special clothing or equipment is required.

6. ACCIDENTAL RELEASE MEASURES

Personal precautions

Avoid dust formation.

Environmental precautions

Do not let product enter drains.

Methods for cleaning up

Sweep up and shovel. Keep in suitable, closed containers for disposal.

7. HANDLING AND STORAGE

Handling: Once the container is opened it should be used promptly, as coloration and decomposition may occur by moisture absorption.

Storage: Storage below room temperature preferred. Store tightly closed in cool, dry, dark and ventilated conditions to maintain the quality for long period.

8. EXPOSURE CONTROL PERSONAL PROTECTION

Desirable Concentration: Not established

Acceptable Concentration: Not established

Facility Care: No special care required

Protective Care: Not necessary during usual handling

9. PHYSICAL AND CHEMICAL PROPERTIES

Appearance: white powder

Odor: Characteristic

Taste: Characteristic

Color: white powder

Critical Temperature: Not available.

Specific Gravity: Not available.

Volatility: Not available.

Odor Threshold: Not available.

Water/Oil Dist. Coeff.: Not available.

Ionicity (in Water): Not available.

Dispersion Properties: Not available.

Solubility: Very slightly soluble in cold water.

10. STABILITY AND REACTIVITY

Stability: The product is stable.

Instability Temperature: Not available.

Conditions of Instability: Excess heat, incompatible materials

Incompatibility with various substances: Reactive with oxidizing agents.

Corrosivity: Non-corrosive in presence of glass.

Special Remarks on Reactivity: Not available.

Special Remarks on Corrosivity: Not available.

Polymerization: Will not occur.

11. TOXICOLOGICAL INFORMATION

Routes of Entry: Eye contact. Inhalation. Ingestion.

Toxicity to Animals:

LD50: Not available. LC50: Not available.

Chronic Effects on Humans: The substance is toxic to lungs, mucous membranes.

Other Toxic Effects on Humans:

Hazardous in case of ingestion, of inhalation. Slightly hazardous in case of skin contact (irritant, permeator).

Special Remarks on Toxicity to Animals: Not available.

Special Remarks on Chronic Effects on Humans: Not available.

Special Remarks on other Toxic Effects on Humans: Not available.

12. ECOLOGICAL INFORMATION

Ecotoxicity: Not available.

BOD5 and COD: Not available.

Products of Biodegradation:

Possibly hazardous short term degradation products are not likely. However, long term degradation products may arise.

Toxicity of the Products of Biodegradation: The products of degradation are less toxic than the product itself.

Special Remarks on the Products of Biodegradation: Not available.

13. DISPOSAL CONSIDERATION

Disposal Method:

Disposal should be made in accordance with federal, state and local regulation.

Contaminated packaging

Dispose of as unused product.

14. TRANSPORT INFORMATION

DOT (US)

Not dangerous goods

IMDG

Not dangerous goods

IATA

Not dangerous goods

15. REGULATORY INFORMATION

The Law Concerning the Examination and Regulation of Manufacture etc. of Chemical Substances

- The Pharmaceutical Affairs Law

16. OTHER INFORMATION

The above information is believed to be correct but does not purport to be all inclusive and shall be used only as a guide. The information in this document is based on the present state of our knowledge and is applicable to the product with regard to appropriate safety precautions. It does not represent any guarantee of the properties of the product. HUNAN SUNSHINE BIO-TECH CO., LTD shall not be held liable for any damage resulting from handling or from contact with the above product. See reverse side of invoice or packing slip for additional terms and conditions of sale.

Updated Jan.1, 2022

End of MSDS

Appendix 2- Microbiological Quality Test Report of Cosmetic Product

1. Microbiological specification test report or data

See below report(s) if available

2. Preservative challenge test report or data

This product is a single use cosmetic product which can be used only once and is then disposed of. Preservative challenge testing is therefore not applicable to the safety of this product and only microbiological quality testing of the finished cosmetic product is required.

Test Report

Number: GZHH00472059

Applicant: Hunan Sunshine Bio-Tech Co., Ltd
Building E7, Lugu Yuyuan, No.27, Wenxuan Road, High-Tech Development Zone, Changsha, Hunan, China, 410000

Date: Nov 01, 2022

Sample Description:

One (1) style of submitted sample said to be :
Item Name : **SQT Revitalizing Beauty Set.**
Country of Origin : China.
Date Sample Received : Oct 20, 2022
Testing Period : Oct 20, 2022 to Nov 01, 2022

Tests conducted:

As requested by the applicant, refer to attached page(s) for details.

Conclusion:

<u>Tested Sample</u>	<u>Standard</u>	<u>Result</u>
Tested component(s) of submitted sample(s)	The European Cosmetic Regulation (EC) No.1223/2009 Annex I Part A 3, Microbiological control criteria of the cosmetic products.	Pass
	With reference to the Notification of the German Federal Health Office Centre (BGA) up to 1996 on toxic elements analysis for cosmetics	Meet

Intertek GM Testing Service Zhuhai Co. Ltd.

Maggie Liu

Maggie Liu
Technical Supervisor
Healthcare and Beauty Products



Test Report

Number: GZHH00472059

Tests Conducted

- 1 Microbiological examination of non-sterile products : Microbial Enumeration Tests and tests for specified microorganisms of submitted sample in composite

With reference to British Pharmacopoeia (2020), Appendix XVI B2 & B1 and European Pharmacopoeia 10.0 Edition, Chapter 2.6.12 & 2.6.13.

Test Item		Result	Limit
		(1+2)	
(I)	Total Aerobic Microbial Count (per mL)	<10 CFU#	Category (B): (I) + (II) should be ≤1,000 CFU
(II)	Moulds and Yeasts Count (per mL)	<10 CFU#	
(III)	<i>Escherichia coli</i> (per mL)	Absence	Absence
(IV)	<i>Pseudomonas aeruginosa</i> (per mL)	Absence	Absence
(V)	<i>Staphylococcus aureus</i> (per mL)	Absence	Absence
(VI)	<i>Candida albicans</i> (per mL)	Absence	Absence
(VII)	Bile-Tolerant Gram-Negative Bacteria (per mL)	Absence	-
(VIII)	<i>Salmonella sp.</i> (per 10mL)	Absence	-
(IX)	<i>Clostridia sp.</i> (per mL)	Absence	-

Test Item		Result	Limit
		(3)	
(I)	Total Aerobic Microbial Count (per g)	<10 CFU#	Category (B): (I) + (II) should be ≤1,000 CFU
(II)	Moulds and Yeasts Count (per g)	<10 CFU#	
(III)	<i>Escherichia coli</i> (per g)	Absence	Absence
(IV)	<i>Pseudomonas aeruginosa</i> (per g)	Absence	Absence
(V)	<i>Staphylococcus aureus</i> (per g)	Absence	Absence
(VI)	<i>Candida albicans</i> (per g)	Absence	Absence
(VII)	Bile-Tolerant Gram-Negative Bacteria (per g)	Absence	-
(VIII)	<i>Salmonella sp.</i> (per 10g)	Absence	-
(IX)	<i>Clostridia sp.</i> (per g)	Absence	-



Test Report

Number: GZHH00472059

Tests Conducted

Test component(s):

- (1) White Powder
- (2) Transparent liquid
- (3) Yellow liquid

Remark :

- # = No colony was detected at the one-tenth dilution of the sample
- CFU = Colony Forming Unit
- < = Less than
- ≤ = Less than or equal to

The limit for test item (I) and (II) is with reference to the notes of guidelines on microbiological quality of the finished cosmetic products adopted by Scientific Committee on Consumer Safety (SCCS, 11th Rev., 2021) of the European Commission, Microbiological Quantitative Limits for Category B Cosmetic Products.

Category A: Cosmetic products specifically intended for children under 3 years, eye area and mucous membranes.

Category B: Other cosmetic products.

Sample received condition: sample (1) in closed container, : sample (2)、 (3) in closed bottle.

Note:

Because the above samples component(1) and (2) had been mixed and tested as one testing item as requested by the applicant, the testing result is for the mixed samples, instead of any individual one.



Test Report

Number: GZHH00472059

Tests Conducted

2 Toxic Element Analysis (Cosmetics)

Acid digestion for Antimony (Sb), Arsenic (As), Cadmium (Cd), Lead (Pb) and Mercury (Hg) and perspiration simulant extraction for Soluble Nickel (Ni), follow by Inductively Coupled Plasma Mass Spectrometry analysis.

Element	Result (ppm)		Reporting Limit (ppm)	Limit# (ppm)
	Test component(s)			
	(1+2)	(3)		
Total Antimony (Sb)	ND	ND	0.1	10
Total Arsenic (As)	ND	ND	0.1	5
Total Cadmium (Cd)	ND	ND	0.1	5
Total Lead (Pb)	ND	ND	0.1	20
Total Mercury (Hg)	ND	ND	0.1	1
Soluble Nickel (Ni)	ND	ND	0.1	10

Test component(s):

- (1) White Powder
- (2) Transparent liquid
- (3) Yellow liquid

Remark :

ppm = parts per million = mg/kg

= The limit is with reference to the Notification of the German Federal Health Office (BGA), Pg 28, No. 7, July 1985 and No. 7/1992 and No. 4/1996 for cosmetics

ND = Not detected (less than reporting limit)

End of report

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Appendix 3- Stability Test Report or Data of Cosmetic Product

See below report(s) if available

Heat resistance	At (40+1)°C , no significant differences in traits after return to room temperature.	Comply						
Cold resistance	At (5+1)°C, no significant differences in traits after return to room temperature.	Comply						
PH	4.0-8.5	6.0	6.1	6.2	6.0	6.0	6.1	6.0
Net content	Should comply with the regulations	Comply						
Total number of colonies	≤ 1000CFU/g	<10CFU/g						
Total Mold and Yeast	≤ 100CFU/g	<10CFU/g						
Conclusion	This product was tested according to QB/T 2872 and the results were in accordance with the regulations.							

Head of Quality : Phil

Reviewer: Peter

Inspector: Adam

Appendix 4- Packaging Compatibility Test Report and/or data

1. Container data

1.1 Basic information

No detail information was provided

2. Outer Packaging material

See below report(s) if available

Test Report

Number: GZHH00472093

Applicant: Hunan Sunshine Bio-Tech Co., Ltd
Building E7, Lugu Yuyuan, No.27, Wenxuan Road, High-Tech Development Zone, Changsha, Hunan, China, 410000

Date: Oct 28, 2022

Sample Description:

One (1) style of submitted sample said to be :
Item Name : **(1) 3ml clear glass vial**
(2) Butyl plug.
Country of Origin : China.
Date Sample Received : Oct 20, 2022
Testing Period : Oct 20, 2022 to Oct 28, 2022

Tests conducted:

As requested by the applicant, refer to attached page(s) for details.

Conclusion:

<u>Tested Sample</u>	<u>Standard</u>	<u>Result</u>
Tested component(s) of submitted sample(s)	Heavy Metals Content Requirement in Directive 94/62/EC and amendments on packaging and packaging waste	Pass

Intertek GM Testing Service Zhuhai Co. Ltd.

Maggie Liu

Maggie Liu
Technical Supervisor
Healthcare and Beauty Products



Test Report

Number: GZHH00472093

Tests Conducted

1 Toxic Elements Analysis

Alkaline digestion method was used and Hexavalent Chromium was determined by UV-Visible Spectrophotometry; Acid digestion method was used and Lead, Cadmium and Mercury were determined by Inductively Coupled Plasma Mass Spectrometry.

Element	Result (ppm)		Detection Limit (ppm)	Limit (ppm)
	Tested Component			
	(1)	(2)		
Lead (Pb)	62.6	12.9	5	--
Cadmium (Cd)	ND	ND	5	--
Mercury (Hg)	ND	ND	5	--
Chromium VI (Cr (VI))	ND	ND	1	--
Sum of Pb, Cd, Hg and Cr (VI)	62.6	12.9	--	100

Tested Component(s):

- (1) Transparent glass bottle
- (2) Gray plastic inner plug

ppm = part per million = mg/kg
ND = Not detected (less than detection limit)

Remark: Only detected element(s) is / are included in calculation of the sum.

End of report

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Test Report

Number: GZHH00472094

Applicant: Hunan Sunshine Bio-Tech Co., Ltd
Building E7, Lugu Yuyuan, No.27, Wenxuan Road, High-Tech Development Zone, Changsha, Hunan, China, 410000

Date: Oct 28, 2022

Sample Description:

One (1) style of submitted sample said to be :

Item Name : **(1) Aluminized facial mask bag**
(2) White facial mask cloth
(3) Pearlescent release film.

Country of Origin : China.

Date Sample Received : Oct 20, 2022

Testing Period : Oct 20, 2022 to Oct 28, 2022

Tests conducted:

As requested by the applicant, refer to attached page(s) for details.

Conclusion:

<u>Tested Sample</u>	<u>Standard</u>	<u>Result</u>
Tested component(s) of submitted sample(s)	Heavy Metals Content Requirement in Directive 94/62/EC and amendments on packaging and packaging waste	Pass

Intertek GM Testing Service Zhuhai Co. Ltd.

Maggie Liu

Maggie Liu
Technical Supervisor
Healthcare and Beauty Products



Test Report

Number: GZHH00472094

Tests Conducted

1 Toxic Elements Analysis

Alkaline digestion method was used and Hexavalent Chromium was determined by UV-Visible Spectrophotometry; Acid digestion method was used and Lead, Cadmium and Mercury were determined by Inductively Coupled Plasma Mass Spectrometry.

Element	Result (ppm)			Detection Limit (ppm)	Limit (ppm)
	Tested Component				
	(1)	(2)	(3)		
Lead (Pb)	ND	ND	ND	5	--
Cadmium (Cd)	ND	ND	ND	5	--
Mercury (Hg)	ND	ND	ND	5	--
Chromium VI (Cr (VI))	ND	ND	ND	1	--
Sum of Pb, Cd, Hg and Cr (VI)	ND	ND	ND	--	100

Tested Component(s):

- (1) Aluminized facial mask bag
- (2) White facial mask cloth
- (3) Pearlescent release film

ppm = part per million = mg/kg
ND = Not detected (less than detection limit)

Remark: Only detected element(s) is / are included in calculation of the sum.

End of report

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Appendix 5- Serious/ Undesirable Effects of Cosmetic Product Declaration or Report

See below report(s) if available

LETTER OF DECLARATION

To Whom It May Concern:

Product Name: SQT Revitalizing Beauty Set

Product: SQT Revitalizing Hydrolyzed Sponge Powder

Chemical Name	Trade Name	Concentration (%)
HYDROLYZED SPONGE	HYDROLYZED SPONGE	98
CALCIUM SILICATE	CALCIUM SILICATE	2
SODIUM SILICATE	SODIUM SILICATE	

Product: SQT Fibronectin Repair Essence

Chemical Name	Trade Name	Concentration (%)
AQUA	AQUA	80-90
GLYCERIN	GLYCERIN	8-10
BETA-GLUCAN	BETA-GLUCAN	1-2
AQUA	AQUA	
1,2-HEXANEDIOL	1,2-HEXANEDIOL	
HYDROXYACETOPHENO NE	HYDROXYACETOPHENO NE	
CAPRYLHYDROXAMIC ACID	CAPRYLHYDROXAMIC ACID	0.75-1.0
ETHYLHEXYLGLYCERIN	ETHYLHEXYLGLYCERIN	
1,2-HEXANEDIOL	1,2-HEXANEDIOL	
PROPYLENE GLYCOL	PROPYLENE GLYCOL	
AQUA	AQUA	
SODIUM POLYGLUTAMATE	SODIUM POLYGLUTAMATE	0.12-0.2
HYDROLYZED SODIUM HYALURONATE	HYDROLYZED SODIUM HYALURONATE	0.05-0.06

Product: SQT Repairing Mask

Chemical Name	Trade Name	Concentration (%)
WATER	WATER	77.103-85.67
GLYCERIN	GLYCERIN	6-6.6
GLYCERETH-26	GLYCERETH-26	3-3.3
LACTOBACILLUS/BEAN SEED EXTRACT/SODIUM GLUTAMATE FERMENT FILTRATE	LACTOBACILLUS/BEAN SEED EXTRACT/SODIUM GLUTAMATE FERMENT FILTRATE	2-2.2
BUTYLENE GLYCOL	BUTYLENE GLYCOL	

WATER	WATER	0.8-0.9
SCHIZOPHYLLAN	SCHIZOPHYLLAN	
1,2-HEXANEDIOL	1,2-HEXANEDIOL	
ETHYLHEXYLGLYCERIN	ETHYLHEXYLGLYCERIN	
CAPRYLYL GLYCOL	CAPRYLYL GLYCOL	
HYDROXYACETOPHENONE	HYDROXYACETOPHENONE	
BUTYLENE GLYCOL	BUTYLENE GLYCOL	
GLUCOSE	GLUCOSE	
NIACINAMIDE	NIACINAMIDE	0.5-0.6
BETAINE	BETAINE	0.5-0.6
CAPRYLHYDROXAMIC ACID	CAPRYLHYDROXAMIC ACID	0.5-0.6
ETHYLHEXYLGLYCERIN	ETHYLHEXYLGLYCERIN	
1,2-HEXANEDIOL	1,2-HEXANEDIOL	
PROPYLENE GLYCOL	PROPYLENE GLYCOL	
WATER	WATER	0.4-0.5
BETA-GLUCAN	BETA-GLUCAN	
WATER	WATER	
1,2-HEXANEDIOL	1,2-HEXANEDIOL	
HYDROXYACETOPHENONE	HYDROXYACETOPHENONE	0.15-0.165
ALLANTOIN	ALLANTOIN	
XANTHAN GUM	XANTHAN GUM	0.1-0.2
METHYLPARABEN	METHYLPARABEN	0.1-0.2
SODIUM POLYGLUTAMATE	SODIUM POLYGLUTAMATE	0.08-0.808
HYDROXYPROPYL GUAR	HYDROXYPROPYL GUAR	0.06-0.0606
CARBOMER	CARBOMER	0.05-0.0505
ARGININE	ARGININE	0.05-0.055
SODIUM HYALURONATE	SODIUM HYALURONATE	0.02-0.022
MENTHOLUM	MENTHOLUM	0.02-0.022

1. Animal testing and toxicity studies:

The raw material(s) used in the product and the finish product itself have not been subjected to any animals testing in order to meet the requirements of EU Cosmetic Regulation (EC) No 1223/2009.

2. Undesirable effects (UEs) and serious undesirable effects (SUEs)

The product or, where relevant, other cosmetic products have not been involved to any undesirable effects or serious undesirable effects as defined in the Article 21 of Regulation (EC) No 1223/2009.

Undesirable effects (UEs): "adverse reactions for human health attributable to the normal or reasonably foreseeable use of a cosmetic product."

Serious Undesirable effects (SUEs): "undesirable effects which result in temporary or permanent functional incapacity, disability, hospitalisation, congenital anomalies or an immediate vital risk or death."

I hereby confirmed that all the above information is complete and accurate and agree to immediately notify in writing of any changes to the above details.

Name: Qin Hao

Position: CEO

Date: Sept 29,2022

Company Address: Building E7, Lugu Yuyuan, No.27, Wenxuan Road, High-Tech Development Zone, Changsha, Hunan, China, 410000

Appendix 6- Fragrance

This formulation does not contain a synthetic fragrance and therefore a fragrance safety evaluation as per IFRA code of practice is not available to this product.

1. IFRA Certificate

See below report(s) if available

2. MSDS/SDS

See below report(s) if available

3. Allergen declaration

See below report(s) if available

Appendix 7- Heavy Metal Test Report of Cosmetic Product

See below report(s) if available

Test Report

Number: GZHH00472059

Applicant: Hunan Sunshine Bio-Tech Co., Ltd
Building E7, Lugu Yuyuan, No.27, Wenxuan Road, High-Tech Development Zone, Changsha, Hunan, China, 410000

Date: Nov 01, 2022

Sample Description:

One (1) style of submitted sample said to be :
Item Name : **SQT Revitalizing Beauty Set.**
Country of Origin : China.
Date Sample Received : Oct 20, 2022
Testing Period : Oct 20, 2022 to Nov 01, 2022

Tests conducted:

As requested by the applicant, refer to attached page(s) for details.

Conclusion:

<u>Tested Sample</u>	<u>Standard</u>	<u>Result</u>
Tested component(s) of submitted sample(s)	The European Cosmetic Regulation (EC) No.1223/2009 Annex I Part A 3, Microbiological control criteria of the cosmetic products.	Pass
	With reference to the Notification of the German Federal Health Office Centre (BGA) up to 1996 on toxic elements analysis for cosmetics	Meet

Intertek GM Testing Service Zhuhai Co. Ltd.

Maggie Liu

Maggie Liu
Technical Supervisor
Healthcare and Beauty Products



Test Report

Number: GZHH00472059

Tests Conducted

- 1 Microbiological examination of non-sterile products : Microbial Enumeration Tests and tests for specified microorganisms of submitted sample in composite

With reference to British Pharmacopoeia (2020), Appendix XVI B2 & B1 and European Pharmacopoeia 10.0 Edition, Chapter 2.6.12 & 2.6.13.

Test Item		Result	Limit
		(1+2)	
(I)	Total Aerobic Microbial Count (per mL)	<10 CFU#	Category (B): (I) + (II) should be ≤1,000 CFU
(II)	Moulds and Yeasts Count (per mL)	<10 CFU#	
(III)	<i>Escherichia coli</i> (per mL)	Absence	Absence
(IV)	<i>Pseudomonas aeruginosa</i> (per mL)	Absence	Absence
(V)	<i>Staphylococcus aureus</i> (per mL)	Absence	Absence
(VI)	<i>Candida albicans</i> (per mL)	Absence	Absence
(VII)	Bile-Tolerant Gram-Negative Bacteria (per mL)	Absence	-
(VIII)	<i>Salmonella sp.</i> (per 10mL)	Absence	-
(IX)	<i>Clostridia sp.</i> (per mL)	Absence	-

Test Item		Result	Limit
		(3)	
(I)	Total Aerobic Microbial Count (per g)	<10 CFU#	Category (B): (I) + (II) should be ≤1,000 CFU
(II)	Moulds and Yeasts Count (per g)	<10 CFU#	
(III)	<i>Escherichia coli</i> (per g)	Absence	Absence
(IV)	<i>Pseudomonas aeruginosa</i> (per g)	Absence	Absence
(V)	<i>Staphylococcus aureus</i> (per g)	Absence	Absence
(VI)	<i>Candida albicans</i> (per g)	Absence	Absence
(VII)	Bile-Tolerant Gram-Negative Bacteria (per g)	Absence	-
(VIII)	<i>Salmonella sp.</i> (per 10g)	Absence	-
(IX)	<i>Clostridia sp.</i> (per g)	Absence	-



Test Report

Number: GZHH00472059

Tests Conducted

Test component(s):

- (1) White Powder
- (2) Transparent liquid
- (3) Yellow liquid

Remark :

- # = No colony was detected at the one-tenth dilution of the sample
- CFU = Colony Forming Unit
- < = Less than
- ≤ = Less than or equal to

The limit for test item (I) and (II) is with reference to the notes of guidelines on microbiological quality of the finished cosmetic products adopted by Scientific Committee on Consumer Safety (SCCS, 11th Rev., 2021) of the European Commission, Microbiological Quantitative Limits for Category B Cosmetic Products.

Category A: Cosmetic products specifically intended for children under 3 years, eye area and mucous membranes.

Category B: Other cosmetic products.

Sample received condition: sample (1) in closed container, : sample (2)、 (3) in closed bottle.

Note:

Because the above samples component(1) and (2) had been mixed and tested as one testing item as requested by the applicant, the testing result is for the mixed samples, instead of any individual one.



Test Report

Number: GZHH00472059

Tests Conducted

2 Toxic Element Analysis (Cosmetics)

Acid digestion for Antimony (Sb), Arsenic (As), Cadmium (Cd), Lead (Pb) and Mercury (Hg) and perspiration simulant extraction for Soluble Nickel (Ni), follow by Inductively Coupled Plasma Mass Spectrometry analysis.

Element	Result (ppm)		Reporting Limit (ppm)	Limit# (ppm)
	Test component(s)			
	(1+2)	(3)		
Total Antimony (Sb)	ND	ND	0.1	10
Total Arsenic (As)	ND	ND	0.1	5
Total Cadmium (Cd)	ND	ND	0.1	5
Total Lead (Pb)	ND	ND	0.1	20
Total Mercury (Hg)	ND	ND	0.1	1
Soluble Nickel (Ni)	ND	ND	0.1	10

Test component(s):

- (1) White Powder
- (2) Transparent liquid
- (3) Yellow liquid

Remark :

ppm = parts per million = mg/kg

= The limit is with reference to the Notification of the German Federal Health Office (BGA), Pg 28, No. 7, July 1985 and No. 7/1992 and No. 4/1996 for cosmetics

ND = Not detected (less than reporting limit)

End of report

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Appendix 8- Human Volunteers Studies

1. Human volunteers study for the cosmetic product

No existing studies from human volunteers for finish product were provided

2. Human volunteers study for raw material

No existing studies from human volunteers for raw material(s) were provided

Appendix 9- Assessor's credentials

Leshuai Zhang, Toxicologist, Intertek China
Professor, PhD, DABT, ERT, UKRT, DCST

Education

Ph. D., Comparative Biomedical Sciences **Aug 2005 – May 2010**
Center for Chemical Toxicology Research and Pharmacokinetics, College of Veterinary Medicine,
North Carolina State University, Raleigh, North Carolina, USA

M. S., Molecular Biology **Sept 2002 – June 2005**
Department of applied Biology, East China University of Science and Technology &
Institute of Biochemistry and Cell Biology, Shanghai Institutes for Biological Sciences, Chinese
Academy of Science, Shanghai, China

B. S., Biochemistry **Sept 1998 – June 2002**
Department of applied Biology, East China University of Science and Technology

Certificate

DCST, Diplomat of Certified Toxicologist CST	Apr 2021
ERT, Europe Registered Toxicologist	Aug 2018
UKRT, UK Registered Toxicologist	Aug 2018
DABT, Diplomate of American Board of Toxicology	Oct 2015

Career Experience

Mar 2021 – Present, Toxicologist, Intertek China

February 2014 – Present, Professor in School of Radiation Medicine and Protection (SRMP),
Soochow University, Suzhou, Jiangsu Province, China

Research Interests: Polysaccharides from traditional medical herbs and tumor immunotherapy;
Bismuth compounds and nephrotoxicity; Hepatotoxicity and phospholipidosis by liver spheroids (3D
cell culture); Microcontact printing technology and cell backpack based drug delivery system

November 2012 – January 2014, Research Assistant Professor in the Nanotechnology Innovation
Center of Kansas State University.

Research Interests: Food safety (toxicity) on primary hepatocytes; Nanocorona and Nanotoxicology
studies

June 2010 – June 2012, Research Fellow in the Division for Drug Safety Research, Center for
Drug Evaluation and Research, Food and Drug Administration, supported by the Oak Ridge
Institute of Science and Education Fellowship Program. Under the supervision of Dr. Rodney
Rouse and Dr. Thomas Colatsky.

Research Description: Drug induced pancreatitis in vivo, biomarker evaluation and toxicity
mechanisms; Nanoparticle toxicity prediction in vitro; Calcium signaling in drug induced
cardiovascular injury

Aug 2005 – June 2010, Graduate Research Assistant, Center for Chemical Toxicology Research
and Pharmacokinetics, Department of Clinical Sciences, College of Veterinary Medicine, North
Carolina State University, Raleigh, North Carolina. Under the supervision of Nancy A. Monteiro-
Riviere.

Research Description: Quantum dot nanoparticle penetration and absorption in skin; Cytotoxicity of
nanoparticles via MTT/Cell Titer Blue/Cell Titer 96AQ/LDH assays, live/dead fluorescence markers
and apoptosis/necrosis markers, inflammatory factors release and reactive oxygen species (ROS);
Nanoparticle cellular uptake and mechanisms by human epidermal keratinocytes, dendritic cells
and mesenchymal stem cell derived adipose cells

Publications Citation > 1600

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Book and Chapters

Zhang L, Chen D. 2017. Chapter 7. Cellular uptake mechanisms of nanoparticles for biomedical imaging. In Shi D, Zhang B (eds.): *Nano Imaging: From Fundamental Principles to Translational Medical Applications*. The World Scientific Encyclopedia of Nanomedicine and Bioengineering I. World Scientific., pp. 241-272.

Zhang L, Z Xuan, Xing T : Experimental Techniques for Radiation Nanomedicine and Nanotoxicology, 2016. ISBN 978-7-5605-9318-0.

Monteiro-Riviere NA, **Zhang LW**. 2008. Assessment of quantum dot penetration into skin in different species under different mechanical actions. In Linkov I, Steevens J (eds.): Nanomaterials: Risks and Benefits. Springer, Dordrecht, Netherlands, pp. 41-52.

Journal Reviewers

Journal Name	IF	Review #
Biomaterials	10.3	2
ACS Applied Materials & Interfaces	8.5	9
Nanoscale	7	3
Particle and Fibre Toxicology	6.6	2
Wiley Interdisciplinary Reviews-Nanomedicine and Nanobiotechnology	6.1	8
Carbohydrate Polymer	6	4
Nanotoxicology	6	1
Biomacromolecules	5.7	1
Nanomedicine-Nanotechnology Biology and Medicine	5.6	8
Science of the Total Environment	5.6	1
International Journal of Biological Macromolecules	4.8	7
ACS Biomaterials Science & Engineering	4.5	1
International Journal of Nanomedicine	4.5	23
Scientific Reports	4	3
Toxicological Sciences	3.6	2
Metallomics	3.6	1
Toxicology	3.5	6
Toxicology letters	3.5	22
Cellular Immunology	3.3	3
Toxicology in vitro	3.1	31
Journal of Applied Toxicology	3.1	1
Archives of Pharmacal Research	2.5	1
Cancer Management and Research	2.2	1
Frontiers in Veterinary Science	2	1
IET Nanobiotechnology	1.9	1
Toxicology and Industrial Health	1.6	20
Toxicologic Pathology	1.4	3
Animal Biotechnology	1.3	1
International Journal of Toxicology	1.2	16
Journal of Nanoscience and Nanotechnology	1.1	1
Cutaneous and Ocular Toxicology	1.1	2
Nanoimpact		3
Nanotoday		2
Nanoscale Advances		1
Applied In Vitro Toxicology		1
Theranostics		1
Total		195

Funding Support

1. Design of cell backpacks by micro contact printing and their applications in tumor immunotherapy. National Natural Science Foundation of China #32171403, 2022/01-2025/12
2. Hepatotoxicity of copper sulfide nanoparticles. National Natural Science Foundation of China #31971319, 2020/01-2023/12
3. Bismuth nanomaterials and nephrotoxicity, National Natural Science Foundation of China #31771104, 2018/01-2021/12
4. Influence of Graphene oxide Derivatives on phospholipidosis, National Natural Science Foundation of China #81401511, 2015/01- 2017/12
5. Immunoregulatory function on herbal polysaccharide on dendritic cells, National Natural Science Foundation of China #81373950, 2014/01 - 2017/12

Awards and Scholarships

1. Outstanding young scholars awarded by Chinese Society of Toxicology (2020)
2. Battelle Memorial Research Award of the Dermal Toxicology Specialty Section at the 48th Annual Meeting of the National Society of Toxicology (SOT), Baltimore, MD, 2009. Research Proposal "Inhibition of multi-walled carbon nanotubes in human epidermal keratinocytes by lectin or niacinamide", \$2500.
3. First place award for the MB Research Award, at the 47th Annual Meeting of the National Society of Toxicology (SOT), Seattle, Washington, 2008.
4. Third place for best poster at the In Vitro and Alternative Methods Specialty Section at the 47th Annual Meeting of the National Society of Toxicology (SOT), Seattle, Washington, 2008.
5. Toxicology and Applied Pharmacology, Certificate of Recognition for one of Elsevier's Top 10 Cited Articles on Scopus 2007-2008.

Professional Associations and Activities

- | | |
|----------------|--|
| 2021 – Present | Associate Editor, Journal of Nanobiotechnology |
| 2021 – Present | Editor Board Member, Toxicology Research and Applications |
| 2016 – Present | Officer, Nanotoxicology Specialty Section, Chinese Society of Toxicology |
| 2012 – Present | Associate Editor, Toxicology and Industrial Health |
| 2012 – 2015 | Education Committee Officer, US Society of Toxicology |
| 2011 – 2012 | Officer, Nanotoxicology Specialty Section, US Society of Toxicology |
| 2009 – Present | Full membership, Sigma Xi Scientific Research Society |
| 2006 – Present | Membership in US Society of Toxicology |



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LESHUAI ZHANG

may use the title

ERT

**EUROPEAN
REGISTERED
TOXICOLOGIST**

whilst registered with the

UK

Register of Toxicology


Signature

June 26, 2018

Date

EUROTOX
Basle, SWITZERLAND

This is to certify that

Leshuai Zhang

has been registered with the

UK Register of Toxicologists

and is bound by the codes of conduct of the

**Royal Society of Biology
and
British Toxicology Society**

for the period

21st May 2018 to 20th May 2023

Lesley Stanley

**Dr Lesley Stanley, ERT
(Panel Chair)**



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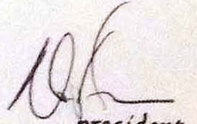
Leshuai Zhang

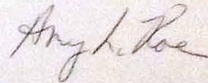
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Certified in General Toxicology



October 29, 2015


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corporate secretary



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EXECUTIVE DIRECTOR

Susie Masten

*Serving in a
personal capacity

August 2019

Dr. Leshuai Zhang
Guoliyuan Xincun 76-202
Nantong, 226001
China

Dear Dr. Zhang:

This letter is to inform you of the status of your recertification application.

Your application is in order and you passed the Literature Review assessment. Therefore, nothing further is required. In December of 2020 (**NOT 2019**) you will receive a letter and sticker affirming your recertification for five years.

Please note, Diplomates are strongly encouraged to record activities related to recertification on an ongoing basis via the ABT website.

If you have any questions, please contact the ABT office.

Sincerely,

Susie Masten
Executive Director