





中国认可 国际互认 检测 TESTING CNAS L2954

# **Final Report**

Report Number: SDWH-M202007193-4(E)

# **Skin Sensitization Test of**

# 5040TPU Low noise medical cable

According to ISO 10993-10:2010 Guinea Pig Maximization Test Sesame Oil Extract

Sponsor: Shenzhen YONGQIANGFU Industrial CO.,Ltd

Address: 2Bldg,No.2 Industrial park,Xinwei Village ,Dalang LongHua Town,Shenzhen City



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## Supplementary Explanation

(1) Please apply for rechecking within 15 days of receiving the report if there are any objections.

(2) Any erasure or without special inspection and testing seal renders the report null and void.

(3) The report is only valid when signed by the persons who edited, checked and approved it.

(4) The results relate only to the articles tested.

(5) The report shall not be reproduced except in full without the written approval of the institute.











## **Quality Assurance Statement**

The Quality Assurance Unit inspected/audited this study in compliance with the following GLP regulations:

Good Laboratory Practice (GLP) Regulation 21 CFR Part 58, U.S. Food and Drug Administration (FDA). The laboratory is exempt from the following provisions: 21 CFR Part 58.105 Test and Control Article Characterization, and Part 58.113 Mixtures of Articles with Carriers.

The Quality Assurance Unit conducted inspections on the following dates. The findings were reported to the Study Director and to the Testing Facility Management. The final report was reviewed by the Quality Assurance Unit. The final report accurately describes the test methods in accordance with standard operating procedures, and the results are consistent with raw data of non-clinical studies conducted according to the study protocol.

Inspections	Date of Inspection	Date Reported to Study Director	Date Reported to Testing Facility Management.
Study Protocol	2021-01-07	2021-01-07	2021-02-25
Study Procedure	2021-01-29 2021-02-02	2021-01-29 2021-02-02	2021-02-25
Raw Data	2021-02-25	2021-02-25	2021-02-25
Final Report	2021-02-25	2021-02-25	2021-02-25

Quality Assurance Unit:

Rian Xu

2021-02-25

Quality Assurance

Date

## **GLP** Compliance Statement

This study was fully in accordance with the technical requirements of the study protocol.

This study was conducted in compliance with Good Laboratory Practice (GLP) Regulation 21 CFR Part 58, U.S. Food and Drug Administration (FDA). The laboratory is exempt from the following provisions: 21 CFR Part 58.105 Test and Control Article Characterization, and Part 58.113 Mixtures of Articles with Carriers.

	C		C
Test A	rticle Receipt	2020-12-24	4
Protoco	Effective Date	2021-01-07	7
Technica	l Initiation Date	2021-01-07	7
Technical	<b>Completion Date</b>	2021-02-03	5
<b>Final Repor</b>	t Completion Date	2021-02-26	

## **Verification Dates**

Edited by:

Chennongrong

2021-02-24

Date

2021-02-26

2021-02-26

Date

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Date

Approved by:

Reviewed by:

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Study Director

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Authorized Signatory

Sanitation & Environment Technology Institute, Soochow

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## Summary

### 1 Test Article

Test Article Name	5040TPU Low noise medical cable	
Manufacturer	Shenzhen YONGQIANGFU Industrial CO.,Ltd	
Address	2Bldg,No.2 Industrial park,Xinwei Village ,Dalang LongHua Town,Shenzhen City	
Model	5040TPU	
Lot/Batch	N/A	

### 2 Main Reference

ISO 10993-10:2010 Biological evaluation of medical devices — Part 10: Tests for irritation and skin sensitization

### 3 Test Method

Potential skin sensitization of test article was evaluated using guinea pig maximization test in accordance with ISO 10993-10:2010 Biological evaluation of medical devices — Part 10: Tests for irritation and skin sensitization.

Study protocol number: SDWH-PROTOCOL-GLP-M202007193-4.

### 4 Conclusion

Under the conditions of this study, the test article extract showed no significant evidence of causing skin sensitization in the guinea pig. The positive rate of sensitization was 0%. No evidence of skin sensitization in guinea pigs was found.

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## **Test Report**

## 1 Purpose

The test was designed to evaluate the potential of a test article to cause skin sensitization. The test is used as a procedure for screening of contact allergens in guinea pigs and extrapolating the results to humans, but it does not establish the actual risk of sensitization.

## 2 Reference

ISO 10993-10:2010 Biological evaluation of medical devices — Part 10: Tests for irritation and skin sensitization

ISO 10993-12:2012 Biological evaluation of medical devices — Part 12: Sample preparation and reference materials

ISO 10993-2:2006 Biological evaluation of medical devices — Part 2: Animal welfare requirements

## 3 Compliance

Good Laboratory Practice Regulations, 21 CFR, Part 58.

ISO/IEC 17025:2017 General requirements for the competence of testing and calibration laboratories (CNAS—CL01 Accreditation criteria for the competence of testing and calibration laboratories) China National Accreditation Service for Conformity Assessment LABORATORY ACCREDITATION CERTIFICATE Registration No. CNAS L2954.

RB/T 214—2017 Competence assessment for inspection body and laboratory mandatory approval—General requirements for inspection body and laboratory Certification and Accreditation Administration of the People's Republic of China INSPECTION BODY AND LABORATORY MANDATORY APPROVAL Certificate No. CMA 180015144061.

## 4 Identification of Test and Control Articles

### 4.1 Test Article

Test Article Name	5040TPU Low nois	e medical cable					
Manufacturer Shenzhen YONGQIANGFU Industrial CO.,Ltd							
Address	2Bldg,No.2 Industr	ial park,Xinwei V	illage ,Dalang LongHua				
	Town,Shenzhen Cit	у					
Test Article Initial State	Non-sterile						
CAS Number	N/A						
Model	5040TPU						
Size	N/A						
Lot/Batch	N/A						
Raw Material	NA						
Packaging Material	N/A						
Physical State	Solid						
Color	Green						
Density	N/A						
Stability	NA						
Solubility	N/A						
Storage Condition	Room temperature						
Intended Use	N/A						
Additional Information	N/A						

The information about the test article was supplied by the sponsor wherever applicable. The Sponsor is responsible for all test article characterization data as specified in the GLP regulations.

#### 4.2 Control Article

#### 4.2.1 Negative Control

Name: Sesame oil (SO). Manufacturer: Ji'an Qingyuan District luyuanxiangliao. Co. Ltd Size: 5kg Lot/ Batch#: 20200928 Physical State: Oily liquid Color: Pale yellow Storage Condition: Room Temperature

#### 4.2.2 Positive Control

Name: 2, 4-Dinitrochlorobenzene (DNCB) Manufacturer: Chengdu Aikeda Chemical Reagent Co., Ltd. Size: 100g Lot/ Batch#: 201904101 Induction Concentration: 0.5% Challenge Concentration: 0.1% Solvent: Sesame oil Date prepared: Intradermal Induction Phase I :2020-11-30; Topical Induction Phase II: 2020-12-07; Challenge Phase: 2020-12-21 Physical State: Liquid Color: Light Yellow Storage Condition: Room Temperature

## 5 Equipment and Reagents

### 5.1 Equipment

Equipment Name	<b>Equipment Number</b>	<b>Calibration Expire</b>
Electronic scale	SDWH442	2021-04-25
Horizontal Large Capacity Constant Temperature Vibrator	SDWH2671	2021-12-23
Steel straight scale	SDWH463	2021-07-06
Vertical pressure steam sterilizer	SDWH2097	2021-03-25

### 5.2 Reagents

Reagent Name	Manufacturer	LOT
Freund's adjuvant, complete liquid	SIGMA	SLCC6223
Sodium dodecyl sulfate (SDS)	Ron reagent	RH178474

## 6 Identification of Test System

Species: Hartley guinea pig (Cavia Porcellus) Number: 15 (10 test +5 negative control) Sex: Male

Initial body weight:  $300 \sim 500$  g

Health status: healthy, not previously used in other experimental procedures Housing: animals were housed in groups in cages identified by a card indicating the lab number, test code and first treatment date, etc.

Animal identification: Stain with dyeing liquid

Cages: plastic cage

Acclimation period: 7 days under the same conditions as for the actual test

## 7 Animal Care and Maintenance

Animal source: Suzhou Experimental Animal Sci-Tech Co., Ltd. <Permit Code: SCXK (SU) 2020-0007>

Bedding: corncob, Suzhou Shuangshi Laboratory Animal Feed Science Co., Ltd.

Feed: guinea pig diet, Suzhou Experimental Animal Sci-Tech Co., Ltd.

Water: drinking water met the Standards for Drinking Water Quality GB 5749-2006

Animal room temperature:  $18 \sim 26$  °C

Animal room relative humidity:  $30\% \sim 70\%$ 

Lights: 12 h light/dark cycle, full-spectrum lighting

Personnel: associates involved were appropriately qualified and trained

Selection: only healthy, previously unused animals were selected

There were no known contaminants present in the feed, water, or bedding expected to interfere with the test data.

## 8 Justification of Test System and Route of Administration

The albino guinea pig has been used historically for sensitization studies (Magnusson and Kligman, 1970). The guinea pig is believed to be the most sensitive animal model for this type of study. The susceptibility of the guinea pig to a known sensitizing agent, 2,4-dinitrochlorobenzene (DNCB) has been substantiated at SDWH (listed in **Table 1** and **Table 2**).

The test article was extracted and administered in vivo through a medium compatible with the test system. Dermal application corresponds to the likely route of human exposure.

## 9 Experimental Design

### 9.1 Preparation of Extracts

#### 9.1.1 Pretreatment

No pretreatment required.

#### 9.1.2 Extraction

Under aseptic conditions, samples were taken according to the sampling method (Randomly take the cable jacket only for test). The extraction was performed with agitation in closed inert containers according to the extraction ratio listed in the following table (sample: extraction vehicle). The extraction vehicle was SO.

	^	<b>^</b>	Esta	act Procedu	Mo	Einal
	<b>Test Period</b>	Actual Sampling	EXU	lre	Final	
iest i criou		Actual Sampling	<b>Extract Ratio</b>	SO	Condition	Extract
	Intradermal	Surface area	$3 \text{ cm}^2$ : 1 mL	10.0 mL	50°C, 72 h	Clear
	Induction Phase I	$30 \text{ cm}^2$	5 cm . T mL	10.0 IIIL	50 C, 72 II	Cicai
	<b>Topical Induction</b>	Surface area	$3 \text{ cm}^2$ : 1 mL	10.0 mI	50°C, 72 h	Clear
	Phase II	$30 \text{ cm}^2$	5 cm . 1 mL	10.0 IIIL	50 C, 72 II	Cicai
	Challenge Phase	Surface area	$3 \text{ cm}^2$ : 1 mL	10.0 mL	50°C, 72 h	Clear
	Chancinge Fliase	$30 \text{ cm}^2$		10.0 IIIL	50  C, 72  H	Cical

The state of the extract did not change after extraction. The extract was stored at room temperature, and tested within 24 h, without the process of adjusting its pH value, filtering, centrifuging, diluting, etc.

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The vehicle (without the test article) was similarly prepared to serve as the control.

#### **9.2 Experimental Procedure**

#### 9.2.1 Animal Preparation and Grouping

On the first day of treatment, 15 guinea pigs were weighed and identified. The fur from the dorsoscapular area of the animals was removed with an electric clipper. Grouping as follow:

Group Name	<b>Group Size</b>	Gender
Test	10 animals	Male
Negative Control	5 animals	Male

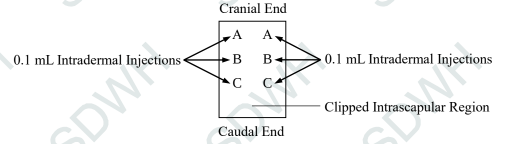
#### 9.2.2 Intradermal Induction Phase I

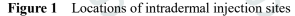
A pair of 0.1 mL intradermal injections was made for each of the following, into each animal, at the injection sites (A, B and C) as shown in Figure 1 in the clipped intrascapular region.

Site A: A 50:50 (V/V) stable emulsion of Freund's complete adjuvant mixed with the chosen solvent.

Site B: the test sample (undiluted extract); the control animals were injected with the solvent alone.

Site C: the test sample at the concentration used at site B, emulsified in a 50:50 (V/V) stable emulsion of Freund's complete adjuvant and the solvent (50%); the control animals were injected with an emulsion of the blank liquid with adjuvant.





#### 9.2.3 Topical Induction Phase II

The maximum concentration that can be achieved in Intradermal induction phase I did not produce irritation. Animals are pretreated with 10% sodium dodecyl sulfate (Solvent: Distilled water, Date prepared: 2020-09-17) ( $24 \pm 2$ ) h before the topical induction application.

At  $7 \pm 1$  d after completion of the intradermal induction phase, administer 0.5 mL test article extract by topical application to the intrascapular region of each animal, using a patch of area approximately 8 cm<sup>2</sup> (absorbent gauze), so as to cover the intradermal injection sites. Secure the patches with an occlusive dressing. Remove the dressings and patches after (48 ± 2) h. Treat the control animals similarly, using the blank liquid alone.

#### 9.2.4 Challenge Phase

At  $14\pm 1$  d after completion of the topical induction phase, challenge all test and control animals with the test sample. Administer 0.5ml test article extract and control article by topical application to sites that were not treated during the induction stage, using absorbent gauze (8 cm<sup>2</sup>) soaked in the test article extract and control article. Secure with an occlusive dressing. Remove the dressings and patches after (24±2) h.

#### 9.3 Observation of Animals

Observe the appearance of the challenge skin sites of the test and control animals  $(24 \pm 2)$  h and  $(48 \pm 2)$  h after removal of the dressings. Full-spectrum lighting was used to visualize the skin reactions. Describe and grade the skin reactions for erythema and oedema according to the Magnusson and Kligman grading given in the following table for each challenge site and at each

#### time interval.

 Magnusson and Kligman	scale	5
Patch Test Reaction	Grading Scale	
No visible change	0	
Discrete or patchy erythema	1	
Moderate and confluent erythema	2	
Intense erythema and/or swelling	3	

### 9.4 Evaluation of Results

Magnusson and Kligman grades of 1 or greater in the test group generally indicate sensitization, provided grades of less than 1 are seen in control animals.

If grades of 1 or greater are noted in control animals, then the reactions of test animals which exceed the most severe reaction in control animals are presumed to be due to sensitization.

If the response is equivocal, rechallenge is recommended to confirm the results from the first challenge.

The outcome of the test is presented as the frequency of positive challenge results in test and control animals.

## **10 Results**

The results of skin reaction after challenge were listed in **Table 3**. No skin sensitization reaction was found in the skin of guinea pigs using extracts of the test article, and the positive rate of sensitization was 0%.

The positive rate of sensitization in the positive control group was 100%, listed in **Table 1**. Clinical observations and weight changes of guinea pigs were listed in **Table 4**.

## **11 Conclusion**

Under the conditions of this study, the test article extract showed no significant evidence of causing skin sensitization in the guinea pig.

## 12 Record Storage

All raw data pertaining to this study and a copy of the final report are to be retained in designated SDWH archive.

## **13 Confidentiality Agreement**

Statements of confidentiality were as agreed upon prior to study initiation.

## **14 Deviation Statement**

There were no deviations from the approved study protocol which were judged to have any impact on the validity of the data.

## Annex 1 Test Data

	Table 1	Guine	ea pig ser	sitization d	ermal react	ions of po	sitive co	ntrol
Group	Animal Number	Bef I	4 ± 2) h ore Phas I Patch plication	e Fo Chall	l ± 2) h llowing enge Phase	Fo	8 ± 2) h llowing enge Pha	Positive Rate after Challenge
		Left	Righ	nt Test Sites	Control Sites	Test Sites	Contr Sites	-
	1	1	2	1	0	2	0	<
Positive	2	2	2	2	0	1	0	
Control	3	2	1	2	0	2	0	100%
Control	4	1	1	1	0	1	0	
	5	1	2	2	0	2	0	
	6	0	0	0	0	0	0	
	7	0	0	0	0	0	0	
Negative	8	0	0	0	0	0	0	-
Control	9	0	0	0	0	0	0	
	10	0	0	0	0	0	0	
Note: the	data of r	ositive	control	come fron	1 SDWH-	M20200	6469-2	(Completed Date:

Note: the data of positive control come from SDWH- M202006469-2 (Completed Date: 2020-12-25)

Table 2         Weigh change and clinical observation of positive control							
	Animal –	Wei	ight (g)	Clinical Observation Except			
Group	Number	Before Injection	After Experiment	Dermal Reactions			
	1	337	411	Normal			
D : 4:	2	333	415	Normal			
Positive Control	3	316	393	Normal			
Control	4	342	425	Normal			
	5	355	432	Normal			
	6	356	428	Normal			
	7	319	393	Normal			
Negative Control	8	347	425	Normal			
Control	9	330	411	Normal			
	10	341	412	Normal			

Note: the data of positive control come from SDWH- M202006469-2 (Completed Date: 2020-12-25)

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	Table 3         Guinea pig sensitization dermal reactions									
	Group	Animal Number	(24 ± 2) h Before Phase II Patch Application		(24 ± 2) h Following Challenge Phase		(48 ± 2) h Following Challenge Phase		Positive Rate after Challenge	
			Left	Right	Test Sites	Control Sites	Test Sites	Control Sites	Phase	
7		1	0	0	0	0	0	0	•	
	Test	2	0	0	0	0	0	0		
		3	0	0	-0	0	0	0		
		4	0	0	0	0	0	0		
		5	0	0	0	0	0	0	0%	
		6	0	0	0	0	0	0	070	
		7	0	0	0	0	0	0		
		8	0	0	0	0	0	0		
		9	0	0	0	0	0	0		
7		10	0	0	0	0	0	0		
	Negative Control	11	0	0	0	0	0	0		
		12	0	0	0	0	0	0		
		13	0	0	0	0	0	0	-	
		14	0	0	0	0	0	0		
		15	0	0	0	0	0	0		

		Table 4	Weigh chans	ge and clinical ob	servation		
		Animal —	· · · · ·	ht (g)			
	Group	Number	Before Injection	After Experiment	<ul> <li>Clinical Observation Except Dermal Reactions</li> </ul>		
		1	344	425	Normal		
		2	352	433	Normal		
		3	348	436	Normal		
		4	341	424	Normal		
	Test	5	325	400	Normal		
	Test	6	357	450	Normal		
		7	344	424	Normal		
		8	347	433	Normal		
		9	322	395	Normal		
		10	348	427	Normal		
	Negative Control	11	351	439	Normal		
		12	354	437	Normal		
		13	355	439	Normal		
		14	323	395	Normal		
		15	348	432	Normal		

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#### **Information Provided by Sponsor** Annex 3

#### 1 **Production Process**

Not supplied by sponsor.

# 2 Other Information Not supplied by sponsor.

End of Report













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