

EU Declaration of Conformity

Manufacturer: Xiantao Yi-Ya Protective Products Co.,Ltd.
Mian dong Road, Hefeng Village, Xiliuhe Town,
Xiantao City, Hubei Province, China
Tel +86 18696471067
Daisy@xtyiya.com

European Representative: Luxus Lebenswelt GmbH
Kochstr. 1, 47877, Willich, Germany
DIMDI Code DE/0000047791
Tax Number DE305829099
Contact Person Lin Sun
Tel/Fax 0049-1715605732
E-mail Info.m@luxuslw.de

Product Name Disposable face mask/Disposable protective mask
Disposable Medical face mask
Test Article YY-1, YY-2, YY-3, YY-4, YY-5

Classification(MDD, Annex VIII) : Class 1, Rule 1

We herein declare that the above mentioned products meet the transposition into national law, the provisions of the following EC Council Directives and Standards. All supporting documentations are retained under the premises of the manufacturer. it bears the mark



General applicable regulations, directives:

Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices, amending Directive 2001/83/EC, Regulation (EC) No 178/2002 and Regulation (EC) No 1223/2009 and repealing Council Directives 90/385/EEC and 93/42/EEC.

Applied standards, common specification, guidance:

EN 14683:2019+AC:2019, EN ISO 15223-1:2016, EN 1041:2008, EN ISO 14971:2012, EN62366-1:2015+AC:2015, ISO 10993- 1:2018, ISO 10993-5:2009, ISO 10993-10:2010, ASTM D4169-2016, MDCG 2019-15.

13th Dec, 2019

Zhao Junpeng /General Manager
Xiantao Yi-Ya Protective Products Co.,Ltd.



Zhao Junpeng

Synthetic Blood Penetration Resistance Final Report

Test Article: Disposable medical mask - 1
Disposable medical mask - 2
Disposable medical mask - 3
Disposable medical mask - 4
Disposable medical mask - 5
Study Number: 1289150-S01.1 Amended
Study Received Date: 16 Apr 2020
Study Completion Date: 27 Apr 2020
Testing Facility: Nelson Laboratories, LLC
6280 S. Redwood Rd.
Salt Lake City, UT 84123 U.S.A.
Test Procedure(s): Standard Test Protocol (STP) Number: STP0012 Rev 09
Deviation(s): None

Summary: This procedure was performed to evaluate surgical facemasks and other types of protective clothing materials designed to protect against fluid penetration. The purpose of this procedure is to simulate an arterial spray and evaluate the effectiveness of the test article in protecting the user from possible exposure to blood and other body fluids. The distance from the target area surface to the tip of the cannula is 30.5 cm. A test volume of 2 mL of synthetic blood was employed using the targeting plate method.

This test method was designed to comply with ASTM F1862 and ISO 22609 (as referenced in EN 14683:2019 and AS4381:2015) with the following exception: ISO 22609 requires testing to be performed in an environment with a temperature of $21 \pm 5^\circ\text{C}$ and a relative humidity of $85 \pm 10\%$. Instead, testing was performed at ambient conditions within one minute of removal from the environmental chamber held at those parameters.

All test method acceptance criteria were met. Testing was performed in compliance with US FDA good manufacturing practice (GMP) regulations 21 CFR Parts 210, 211 and 820.

Number of Test Articles Tested: 32 per test pressure
Number of Test Articles Passed: 32
Test Side: Outside
Pre-Conditioning: Minimum of 4 hours at $21 \pm 5^\circ\text{C}$ and $85 \pm 5\%$ relative humidity (RH)
Test Conditions: 22.2°C and 21% RH



Shang Shuang for
Study Director _____ James W. Luskin

07 May 2020
Amended Report Date _____



1289150-S01

Results: Per ASTM F1862 and ISO 22609, an acceptable quality limit of 4.0% is met for a normal single sampling plan when ≥ 29 of 32 test articles show passing results.

Test Pressure: 120 mmHg (16.0 kPa)

Test Article Number	Synthetic Blood Penetration
1-32	None Seen

Amendment Justification: At the request of the sponsor, the company name and address were updated. Additionally, the initial report was separated into individual reports by test pressure.

Microbial Cleanliness (Bioburden) of Medical Masks Final Report

Test Article: Disposable medical mask -1
Disposable medical mask - 2
Disposable medical mask - 3
Disposable medical mask - 4
Disposable medical mask - 5

Study Number: 1289157-S01

Study Received Date: 16 Apr 2020

Testing Facility: Nelson Laboratories, LLC
6280 S. Redwood Rd.
Salt Lake City, UT 84123 U.S.A.

Test Procedure(s): Standard Test Protocol (STP) Number: STP0036 Rev 15
Customer Specification Sheet (CSS) Number: 202002111 Rev 01

Deviation(s): None

Summary: The testing was conducted in accordance with EN 14683:2019, with the exception of approximate volumes of eluent used when performing the extraction procedure and a temperature range of 30-35°C used for aerobic incubation.

When bioburden results are calculated using a software program, manual calculations may differ slightly due to rounding. The counts determined on products are colony forming units and may not always reflect individual microorganisms. The sponsor performs any statistical analysis and determines the acceptable limits. Testing was performed in compliance with US FDA good manufacturing practice (GMP) regulations 21 CFR Parts 210, 211 and 820.



Carl Danielson electronically approved for
Study Director

Robert Putnam

29 Apr 2020 17:53 (+00:00)
Study Completion Date and Time

Results:

Unit Number	Weight (g)	Aerobic	Fungal	Total Bioburden (CFU/mask)	Total Bioburden (CFU/g)
1	3.2	<3	<3	<6.1	<1.9
2	3.1	<3	<3	<5.8	<1.9
3	3.1	<3	<3	<6.1	<2.0
4	3.1	<3	<3	<5.8	<1.9
5	3.1	<3	<3	<6.0	<1.9
Recovery Efficiency	UTD ^a				

< = No Organisms Detected

UTD = Unable to Determine

Note: The results are reported as colony forming units per test article.

^a UTD due to zero count on the first rinse. An alternative method or inoculated product recovery efficiency is recommended.

Method Suitability:

Organism	Percentage
<i>Bacillus atrophaeus</i>	88%

Test Method Acceptance Criteria: If applicable, anaerobic controls are acceptable for the bioburden test results. The number of masks to be tested shall be a minimum of 5 or more to meet an acceptable quality level of 4%. The bioburden of the medical mask shall be < 30 CFU/g tested.

Procedure:

- Positive Controls/Monitors: *Bacillus atrophaeus*
- Extract Fluid: Peptone Tween[®]
- Extract Fluid Volume: ~300 mL
- Extract Method: Orbital Shaking for 15 minutes at 250 rpm
- Plating Method: Membrane Filtration
- Agar Medium: Potato Dextrose Agar
Tryptic Soy Agar
- Recovery Efficiency: Exhaustive Rinse Method
- Aerobic Bacteria: Plates were incubated 3 - 7 days at 30-35°C, then enumerated.
- Fungal: Plates were incubated 5 - 7 days at 20-25°C, then enumerated.

Bacterial Filtration Efficiency (BFE) and Differential Pressure (Delta P) Final Report

Test Article: Disposable medical mask -1
Disposable medical mask - 2
Disposable medical mask - 3
Disposable medical mask - 4
Disposable medical mask - 5
Study Number: 1289151-S02
Study Received Date: 16 Apr 2020
Testing Facility: Nelson Laboratories, LLC
6280 S. Redwood Rd.
Salt Lake City, UT 84123 U.S.A.
Test Procedure(s): Standard Test Protocol (STP) Number: STP0004 Rev 18
Deviation(s): None

Summary: The BFE test is performed to determine the filtration efficiency of test articles by comparing the bacterial control counts upstream of the test article to the bacterial counts downstream. A suspension of *Staphylococcus aureus* was aerosolized using a nebulizer and delivered to the test article at a constant flow rate and fixed air pressure. The challenge delivery was maintained at $1.7 - 3.0 \times 10^3$ colony forming units (CFU) with a mean particle size (MPS) of $3.0 \pm 0.3 \mu\text{m}$. The aerosols were drawn through a six-stage, viable particle, Andersen sampler for collection. This test method complies with ASTM F2101-19 and EN 14683:2019, Annex B.

The Delta P test is performed to determine the breathability of test articles by measuring the differential air pressure on either side of the test article using a manometer, at a constant flow rate. The Delta P test complies with EN 14683:2019, Annex C and ASTM F2100-19.

All test method acceptance criteria were met. Testing was performed in compliance with US FDA good manufacturing practice (GMP) regulations 21 CFR Parts 210, 211 and 820.

Test Side: Inside
BFE Test Area: $\sim 40 \text{ cm}^2$
BFE Flow Rate: 28.3 Liters per minute (L/min)
Delta P Flow Rate: 8 L/min
Conditioning Parameters: $85 \pm 5\%$ relative humidity (RH) and $21 \pm 5^\circ\text{C}$ for a minimum of 4 hours
Test Article Dimensions: $\sim 177 \text{ mm} \times \sim 94 \text{ mm}$
Positive Control Average: 1.7×10^3 CFU
Negative Monitor Count: < 1 CFU
MPS: $2.9 \mu\text{m}$



Trang Truong electronically approved for
Study Director

James Luskin

19 May 2020 00:42 (+00:00)
Study Completion Date and Time

Results:

Test Article Number	Percent BFE (%)
1	99.8
2	99.6
3	99.9
4	>99.9
5	99.8

Test Article Number	Delta P (mm H ₂ O/cm ²)	Delta P (Pa/cm ²)
1	4.4	43.6
2	4.2	41.5
3	4.3	42.1
4	4.7	46.4
5	4.2	41.4

The filtration efficiency percentages were calculated using the following equation:

$$\% BFE = \frac{C - T}{C} \times 100$$

C = Positive control average

T = Plate count total recovered downstream of the test article

Note: The plate count total is available upon request



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CNAS L0599

Test Report **SL52025245933401TX** Date: May 15, 2020 Page 1 of 3
 XIANTAO JIAJING PROTECTIVE PRODUCTS CO.,LTD
 7-17-301 XIANDAI ZHONGJIA SCIENCE AND TECHNOLOGY CITY, SHAZUI STREET, XIANTAO CITY

The following sample(s) was/were submitted and identified on behalf of the client as:

Sample Description : (A) Disposable face mask

SGS Internal Ref No. : O-CQAFL202000392405

Sample Color : (A) Blue

Test Performed : Selected test(s) as requested by applicant

Sample Receiving Date : Apr 26, 2020

Testing Period : Apr 26, 2020 - May 15, 2020

Test Result(s) : Unless otherwise stated the results shown in this test report refer only to the sample(s) tested, for further details, please refer to the following page(s).

Signed for and on behalf of
 SGS-CSTC Standards Technical Services (Shanghai) Co., Ltd Testing Center

Sara Guo (Account Executive)



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Test Report

SL52025245933401TX

Date: May 15, 2020

Page 2 of 3

Test Result

Medical Face Masks-Requirements and Test Methods
(EN 14683:2019)

Clause 5.2.2 Bacterial filtration efficiency (BFE)*
(EN 14683 :2019 Annex B)

(BFE), %	1#	2#	3#	4#	5#
	99.6	99.6	99.6	99.8	99.7

Remark: Performance Requirement: Type I $\geq 95\%$, Type II $\geq 98\%$, Type IIR $\geq 98\%$

* This test standard is not within the accredited scope in SGS Shanghai testing centre, it is carried out by external laboratory accredited by CMA (China Metrology Accreditation).

Clause 5.2.3 Breathability (Differential Pressure)
(EN 14683 :2019 Annex C, Flow rate 8 l/min)

Differential pressure ΔP (Pa/cm ²)	1#	2#	3#	4#	5#
	34	34	37	36	36

Remark: Performance Requirement: Type I < 40 Pa/cm², Type II < 40 Pa/cm², Type IIR < 60 Pa/cm²

Clause 5.2.4 Splash Resistance
(ISO 22609 :2004, Pressure 16.0 kPa)

Penetration on inside surface							
1#	2#	3#	4#	5#	6#	7#	8#
Pass	Pass	Pass	Pass	Pass	Pass	Pass	Pass
9#	10#	11#	12#	13#	14#	15#	16#
Pass	Pass	Pass	Pass	Pass	Pass	Pass	Pass
17#	18#	19#	20#	21#	22#	23#	24#
Pass	Pass	Pass	Pass	Pass	Pass	Pass	Pass
25#	26#	27#	28#	29#	30#	31#	32#
Pass	Pass	Pass	Pass	Pass	Pass	Pass	Pass
Number of Pass:	32						
Overall result:	Acceptable						

Remark:

- 1) Performance Requirement Type I: N/A, Type II: N/A, Type IIR: $\geq 16.0kPa$
- 2) Distance of the medical face mask target area surface to the tip of cannula is $300 \pm 10mm$.
- 3) Condition and Test temperature $(21 \pm 5)^{\circ}C$, relative humidity $(85 \pm 10)\%$
- 4) An acceptable quality limit of 4.0% is met for a single sampling plan when 29 or more of the 32 tested specimens show pass results



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Clause 5.2.5 Microbial Cleanliness
(EN 14683: 2019 Annex D)

Sample A	1#	2#	3#	4#	5#
CFU/g	1	<1	<1	<1	<1

Remark: Performance Requirement: Type I≤30 CFU/g, Type II≤30 CFU/g, Type IIR≤30 CFU/g

Sample Photo



The statement of conformity in this test report is only based on measured values by the laboratory and does not take their uncertainties into consideration.

End of Report



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TEST REPORT

(Electronic version)



No:200076624

VERIFICATION WEBSITE: www.gtgc.net.cn

VERIFICATION CODE: MFSD-8322-54



ISSUE DATE:2020-04-20

APPLICANT: XIANTAO YI-YA PROTECTIVE PRODUCTS CO.,LTD
ADDRESS: MIANDONG ROAD HEFENG VILLAGE XILIUHE TOWN XIANTAO CITY, HUBEI PROVINCE, CHINA

INFORMATION CONFIRMED BY APPLICANT:

ORDINARY PROTECTIVE MASK
QUANTITY: THIRTY-FIVE PIECES
COLOUR: BLUE

DATE RECEIVED/DATE TEST STARTED: 2020-04-14

CONCLUSION:

BACTERIAL FILTRATION EFFICIENCY	M
PARTICLE FILTRATION EFFICIENCY	M
AIRFLOW RESISTANCE	M

NOTE: "M" -MEET THE STANDARD'S REQUIREMENT "F" -FAIL TO MEET THE STANDARD'S REQUIREMENT
"---" -NO COMMENT

REMARK:

THE DECISION INDICATORS ARE DERIVED FROM THE GROUP STANDARD REQUIRED BY CLIENT (T/CTCA 7-2019). OUR INSPECTION CAPACITY AUTHORIZED BY CMA COVERS THE INSPECTION ITEMS T/CTCA 7-2019.
THIS REPORT IS THE ENGLISH TRANSLATION VERSION OF THE REPORT 200076623.
ALL THE TESTED ITEMS ARE TESTED UNDER THE STANDARD CONDITION (EXCEPT FOR INDICATION).
COPIES OF THE REPORT ARE VALID ONLY RE-STAMPED.
THE EXPERIMENT WAS CARRIED OUT AT No.1, ZHUJIANG ROAD, PANYU DISTRICT, GUANGZHOU, GUANGDONG, P. R. CHINA.

APPROVED BY:
Yuan Liu ENGINEER

刘圆

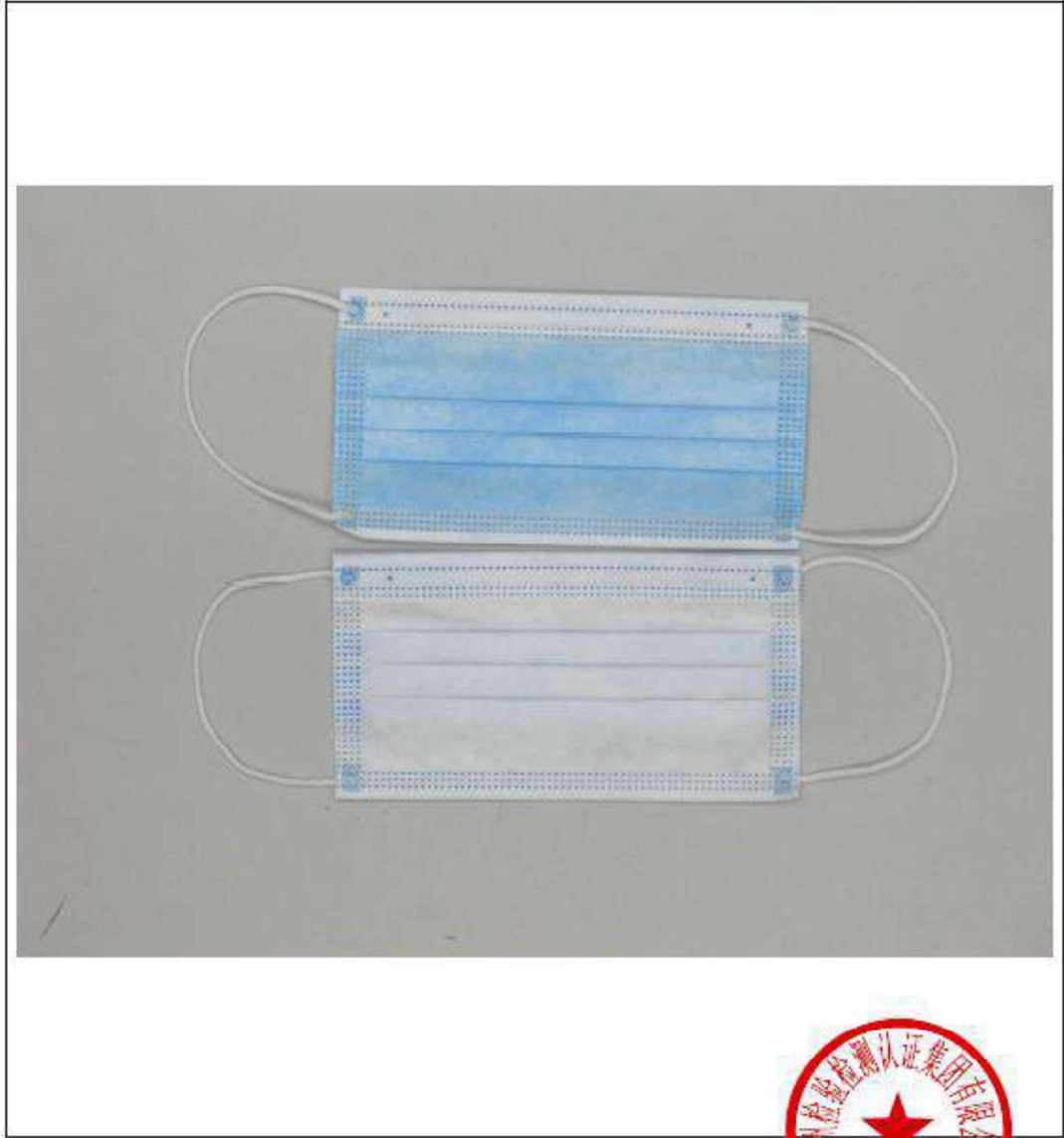


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TEST REPORT

(Electronic version)

No:200076624



TEST REPORT

(Electronic version)

No:200076624

BACTERIAL FILTRATION EFFICIENCY(%)

(YY 0469-2011 ANNEX B, TEST BACTERIA: STAPHYLOCOCCUS AUREUS ATCC 6538, TEST AREA:
40cm², FLOW RATE: 28.3L/min, MEAN PARTICLE SIZE: 3.0 μm, RESULT OF THE POSITIVE
CONTROL: 1.9×10⁸ CFU, RESULT OF THE NEGATIVE CONTROL: <1CFU)

		REQUIREMENT
BFE ₁	99.6	≥95
BFE ₂	99.4	(T/CTCA 7-2019)
BFE ₃	99.2	

PARTICLE FILTRATION EFFICIENCY(%)

(YY 0469-2011 5.6.2, AIR FLOW: 30L/min, AEROSOL: NaCl, AEROSOL CONCENTRATION:
15mg/m³, TEMP: 23.2℃, RH: 36.3%)

		REQUIREMENT
MINIMUM	96.02	≥80
		(T/CTCA 7-2019)

AIRFLOW RESISTANCE(Pa)

(YY 0469-2011 5.6.2, AIR FLOW: 30L/min, AEROSOL: NaCl, AEROSOL CONCENTRATION:
15mg/m³, TEMP: 23.2℃, RH: 36.3%)

		REQUIREMENT
MAXIMUM	40.0	≤80
		(T/CTCA 7-2019)



—End of Report—

PAGE 3 OF 3

Test Report

(Electronic version)

Verification Website: www.gttc.net.cn

Verification Code: FVMR-0184-54

No: 20R000519

Issue Date: 2020-04-24

Applicant: XIANTAO YI-YA PROTECTIVE PRODUCTS CO.,LTD
 Address: MIANDONG ROAD HEFENG VILLAGE XILIUHE TOWN XIANTAO CITY,HUBEI PROVINCE,CHINA

Information confirmed by applicant:

Disposable medical mask(non-sterile)

Quantity: sixty pieces

Lot number: 042020

Model: YY-101 YY-102 YY-103(submission no.: YY-101)

Size: 175mm×95mm

Classification: Type II R

Standard Adopted:

EN 14683:2019+AC:2019 <Medical face masks-Requirements and test method>

Date Received/Date Test Started: 2020-04-09

Conclusion:

Bacterial filtration efficiency (BFE)	M
Microbial cleanliness	M
Differential pressure	M
Splash resistance pressure	M
Materials and construction	M
Design	M
General	M

Note: "M"-Meet the standard's requirement "F"-Fail to meet the standard's requirement "--"-No comment

Remark:

All the tested items are tested under the standard condition (except for indication).

Copies of the report are valid only re-stamped.

The experiment was carried out at No 1, Zhujiang Road, Panyu District, Guangzhou, Guangdong, P.R.China.

Approved By:

ZiShan Guo

ZiShan Guo Senior Engineer



Test Report

(Electronic version)

No: 20R000519



Test Report

(Electronic version)

No: 20R000519

Bacterial filtration efficiency (BFE)

Test method: EN 14683: 2019+AC: 2019 Annex B

Test principle:

A specimen of the mask material is clamped between a six-stage cascade impactor and an aerosol chamber. An aerosol of *Staphylococcus aureus* is introduced into the aerosol chamber and drawn through the mask material and the impactor under vacuum. The bacterial filtration efficiency (BFE) of the mask is given by the number of colony forming units passing through the medical face mask material expressed as a percentage of the number of colony forming units present in the challenge aerosol.

Test equipment:

- Incubator
- Electronic balance
- Autoclave
- Experimental system for bacterial filtration efficiency (BFE) of mask

The environmental conditions of the laboratory and test condition:

- Total bacteria: 0 CFU/plate
- Total fungi: 0 CFU/plate
- Blank experiment: Aseptic growth
- Test environment temperature: 24.5°C, Relative humidity: 56.0%
- Culture medium: TSA agar medium
- Culture temperature: 37°C, Culture time: 48h
- Test bacteria : *staphylococcus aureus* ATCC 6538
- Concentration of bacterium: 5.0×10^5 CFU /ml
- Positive control average (C): 1.9×10^5 CFU
- Negative monitor count: <1 CFU
- Test area: 40 cm²
- Dimensions of the test specimens: 15cm×15cm
- Flow rate: 28.3 l/min
- Pretreatment: Condition each specimen for 4 h by exposure to a temperature of (21±5)°C and a relative humidity of (85±5)%
- Mean particle size: 3.0 μm
- The medical face mask in contact with the bacterial challenge: inside



Test Report

(Electronic version)

No: 20R000519

Results:

Sample	T	BFE (%)	Requirement (%)	Classification	Conclusion
1	9	99.53	≥98 EN 14683:2019+AC:2019	Type II R	Pass
2	8	99.58			
3	6	99.68			
4	11	99.42			
5	14	99.26			

Remarks:

For each test specimen calculate the bacterial filtration efficiency B, as a percentage, using the following formula:

$$B = (C - T) / C \times 100$$

where

B is bacterial filtration efficiency (BFE), %;

C is positive control average;

T is the total plate count for the test specimen.



Test Report

(Electronic version)

No: 20R000519

Microbial cleanliness

Test method: EN ISO 11737-1:2018, Membrane filtration

Test principle:

Take the required samples from the original packaging. Weigh a certain amount of sample and placed in a sterile 500 ml bottle containing 300 ml of extraction liquid (1 g/l Peptone, 5 g/l NaCl and 2 g/l Tween 20). The bottle is laid down on an orbital shaker and shaken for 5 min at 250 rpm. After this extraction step, 100 ml of the extraction liquid is filtered through a 0.45 µm filter and laid down on a TSA plate for the total viable aerobic microbial count. Another 100 ml aliquot of the same extraction liquid is filtered in the same way and the filter plated on Sabouraud Dextrose agar (SDA) for fungi enumeration. The plates are incubated for 3 days at 30°C and 7 days at (20 to 25)°C for TSA and SDA plates respectively. The total bioburden is expressed by addition of the TSA and SDA counts.

Test equipment:

Constant temperature incubator

Electronic balance

Pressure steam sterilizer

Biosafety cabinet

The environmental conditions of the laboratory and test condition:

Test environment temperature: 24.5°C, Relative humidity: 56.0%

Test environment monitoring: total bacteria: 0 CFU/plate, total fungi: 0 CFU/plate, blank experiment: aseptic growth



Test Report

(Electronic version)

No: 20R000519

Results:

Microbial	Measured value (CFU/g)	Microbial cleanliness (CFU/g)	Requirement (CFU/g)	Classification	Conclusion
Bacteria	0	0	≤30 EN 14683:2019+AC:2019	Type II R	Pass
Fungi	0				



Test Report

(Electronic version)

No: 20R000519

Differential pressure

Test method: EN 14683:2019+AC:2019 Annex C

Test principle:

This procedure was performed to evaluate the differential pressure of the medical face mask material by measuring the air exchange pressure through a measured surface area at a constant air flow rate.

Test equipment:

GTTC-YLC-1 Apparatus for measuring differential pressure

The environmental conditions of the laboratory and test condition:

Air flow: 8 l/min

Test area: 4.9cm²

Pretreatment: Condition each specimen for a minimum of 4 h by exposure to a temperature of (21±5) °C and a relative humidity of (85±5)%

General location of the areas of the mask the differential measurements: specimen center



Test Report

(Electronic version)

No: 20R000519

Results:

Sample	Measured value (Pa)	Differential pressure (Pa/cm ²)	Requirement (Pa/cm ²)	Classification	Conclusion
1	167	33.7	< 60 EN 14683:2019+AC:2019	Type II R	Pass
2	163				
3	172				
4	150				
5	174				
Average	165				



Test Report

(Electronic version)

No: 20R000519

Splash resistance pressure

Test method: ISO 22609:2004

Test principle:

A specimen medical face mask is supported on an apparatus. A volume of synthetic blood is sprayed horizontally at the specimen mask to simulate the scenario of a mask being splashed by a punctured blood vessel. The volume of fluid, distance to impact, orifice size and fluid velocity are defined in this method and intended to be consistent with this health care scenario. Any evidence of synthetic blood penetration on the side of the medical face mask contacting the wearer's face constitutes failure. Results are reported as "pass/fail". Specimen medical face masks are evaluated at a total of three different velocities corresponding to human blood pressures of 10.6 kPa, 16.0 kPa, and 21.3 kPa. Test results are reported at each velocity and the medical face mask is rated at the highest corresponding blood pressure for which medical face mask specimens demonstrate an acceptable quality limit of 4.0.

Test equipment:

Test apparatus for synthetic blood penetration LFY-227

Air compressor

Graduated cylinder

Electronic balance

Targeting plate

The environmental conditions of the laboratory and test condition:

Condition each specimen for a minimum of 4 h by exposure to a temperature of $(21\pm 5)^\circ\text{C}$ and a relative humidity of $(85\pm 5)\%$

Surface tension of synthetic blood: 0.042 N/m

Pressure: 16.0 kPa

Velocity: 550 cm/s



Test Report

(Electronic version)

No: 20R000519

Results:

Sample	Measured value	Requirement (kPa)	Classification	Conclusion
	Pressure			
	16.0 kPa			
1	pass	≥16.0 EN 14683:2019+AC:2019	Type II R	Pass
2	pass			
3	pass			
4	pass			
5	pass			
6	pass			
7	pass			
8	pass			
9	pass			
10	pass			
11	pass			
12	pass			
13	pass			
14	pass			
15	pass			
16	pass			
17	pass			
18	pass			
19	pass			
20	pass			
21	pass			
22	pass			
23	pass			
24	pass			
25	pass			
26	pass			
27	pass			
28	pass			
29	pass			
30	pass			
31	pass			
32	pass			
Final result	pass			

Remarks:

An acceptable quality limit of 4.0 % is met for a single sampling plan when 29 or more of the show "pass" results.



Test Report

(Electronic version)

No: 20R000519

Materials and construction

Test Method: EN 14683:2019+AC:2019 5.1.1

Results:

Requirement	Conclusion
The medical face mask is a medical device, generally composed of a filter layer that is placed, bonded or moulded between layers of fabric.	Pass
The medical face mask shall not disintegrate, split or tear during intended use.	Pass
In the selection of the filter and layer materials, attention shall be paid to cleanliness.	Pass



Test Report

(Electronic version)

No: 20R000519

Design

Test Method: EN 14683:2019+AC:2019 5.1.2

Results:

Requirement	Conclusion
The medical face mask shall have a means by which it can be fitted closely over the nose, mouth and chin of the wearer and which ensures that the mask fits closely at the sides.	Pass
Medical face masks may have different shapes and constructions as well as additional features such as a face shield (to protect the wearer against splashes and droplets) with or without anti-fog function, or a nose bridge (to enhance fit by conforming to the nose contours).	Pass



Test Report

(Electronic version)

No: 20R000519

General

Test Method: EN 14683:2019+AC:2019 5.2.1

Results:

Requirement	Conclusion
All tests shall be carried out on finished products or samples cut from finished products.	Pass



——End of Report——

Suche **Suchergebnis** Dokumentausgabe Merkliste (0)

Anzeigen Medizinprodukte (MPA)  

[zurück](#) [weiter](#)

1 von 1 BfArM: MP Anzeigen (MPA) © DIMDI

Dokumentnummer	00162689
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Anzeige

Registrierdatum	2020-06-18
Registriernummer	DE/CA20/01-Luxuslebenswelt-468/20
Formblatttyp	Erstanzeige Medizinprodukt
Anzeigender nach § 25 MPG	Bevollmächtigter
Formularnummer	00304496
Frühere Formularnummer	00304496

Angaben zum Anzeigenden

Code	DE/0000047791
Bezeichnung	Luxus Lebenswelt GmbH
Staat	Deutschland
Ort	Willich
Postleitzahl	47877
Straße, Haus-Nr.	Kochstr. 1
Land	Nordrhein-Westfalen
Telefon	0049-1715605732
E-Mail	info.m@luxuslw.de

Zuständige Behörde

Code	DE/CA20
Bezeichnung	Bezirksregierung Düsseldorf
Zusatz	Dezernat 24
Staat	Deutschland
Land	Nordrhein-Westfalen
Ort	Düsseldorf
Postleitzahl	40474
Straße, Haus-Nr.	Cecilienallee 2
Telefon	+49-211-4750
Telefax	+49-211-4752671
E-Mail	dez24.mpg@brd.nrw.de
Bearbeiter	Frau Nadine Schlingmeier
Bearbeiter Telefon	0211-475-3853

Hersteller

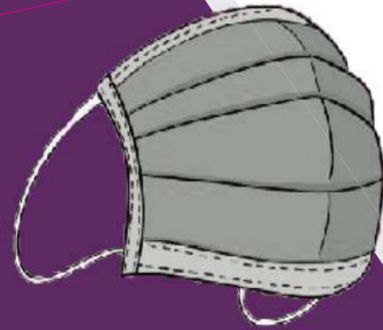
Bezeichnung	XianTao JiaJing Protective Products Co., Ltd
Staat	China
Ort	XianTao
Postleitzahl	433000
Straße, Haus-Nr.	7-17-301,Modern Sino-Canada Science & Technology Industry Park,XianTao, Hubei Province
Telefon	0086-18696471067
Telefax	0086-728-3326822
E-Mail	daisy@xtyiya.com

Medizinprodukt

Produkttyp	nichtaktives Medizinprodukt
Klasse	I
App (Software auf mobilen Endgeräten)	Nein
Handelsname	Disposable Face mask
Allgemeine Produktbezeichnung	Disposable Face mask
Nomenklaturcode	15-230
Nomenklaturbezeichnung	Maske, sonstige
Kategorie	10 Produkte zum Einmalgebrauch
Kurzbeschreibung	Es kann von Arbeitern in der Umwelt mit hoher Partikelkonzentration getragen werden und kann grundlegende Mikroorganismen und Keime isolieren. Es besteht aus schmelzgeblasenem Stoff und Polypropylen-Vliesstoff.
Kurzbeschreibung in Englisch	It can be worn by workers in the environment with large particle concentration, and can isolate basic microorganisms and germs. It is composed of melt blown cloth and polypropylene non-woven cloth.

Typ II R

Medizinische Masken Typ II R – EN 14683



50x
Medizinische Masken
Masques Médicaux
Mascherine Mediche
Medical Face Masks

Product information

- High quality and filtration capacity
- High wearing comfort
- Low breathing resistance
- 100% latex-free and fiberglass-free
- Hypoallergenic

Warnings

This mask is not a respirator. It does not protect the wearer from contagious diseases such as the flu. For single use only. Do not use masks that are damaged or soiled with body fluids.

Anwendungsinformationen

Vor dem Anziehen der Maske die Hände mit Seife oder Desinfektionsmittel waschen.

Maske über Nase und Mund legen. Blaue Seite aussen. Der mit Draht verstärkte Teil ist oben über dem Nasenrücken. Ohrschlaufen um jedes Ohr platzieren.

Unteren Teil der Maske bis unter das Kinn ziehen und den Nasenbügel in die richtige Passform bringen, sodass die Maske überall eng an der Haut anliegt.

Es wird empfohlen, die Maske zu wechseln, wenn sie feucht wird. Die Maske kann im Haushaltsabfall entsorgt werden.

Informations sur la demande

Se laver les mains avec du savon ou du désinfectant avant de mettre le masque.

Mettez le masque de manière à ce qu'il recouvre le nez et la bouche. Côté bleu à l'extérieur. La partie renforcée par du fil de fer doit être placée au-dessus de l'arête du nez. Placez une fixation auriculaire élastique autour de chaque oreille.

Tirez la partie inférieure du masque jusqu'en dessous du menton et réglez le pince-nez afin que le masque s'adapte parfaitement à la peau.

Il est recommandé de changer le masque dès qu'il est mouillé. Le masque peut être jeté dans les ordures ménagères.

Informazioni sull'applicazione

Lavare le mani con sapone o disinfettante prima di indossare la maschera.

Mettere la maschera sul naso e sulla bocca. Lato blu fuori. La parte rinforzata con filo metallico si trova sopra il ponte del naso. Mettere un occhietta per orecchie ad ogni orecchie.

Tirare la parte inferiore della maschera fino a sotto il mento e regolare la barra nasale in modo che la maschera aderisca perfettamente alla pelle ovunque.

Si consiglia di cambiare la maschera quando si bagna. La maschera può essere smaltita nei rifiuti domestici.

Application Information

Wash hands with soap or disinfectant before putting on the mask.

Put the mask over nose and mouth. Blue side outside. The wire rein-forced part is above the bridge of the nose. Place earloop around each ear.

Pull the lower part of the mask up to under the chin and adjust the nose bar to the correct fit so that the mask fits tightly against the skin everywhere.

It is recommended to change the mask as soon as it becomes wet. The mask can be disposed of in household waste.

Leistungsanforderungen für Medizinische Masken Exigences de performance pour les Masques Médicaux Requisiti di prestazione per le Mascherine Mediche Performance requirements for Medical Masks

Klassifizierung / Classification / Classificazione / Classificazione		Type I	Type II	Type II R
Bakterielle Filterleistung / Efficacité de filtration bactérienne / Prestazioni del filtro batterico / Bacterial Filtration Efficiency	BFE	≥ 95 %	≥ 98 %	≥ 98 %
Druckdifferenz / Pression différentielle / Differenza di pressione / Differential pressure	pa/cm ²	< 40	< 40	< 60
Druck des Spritzwiderstands / Pression de la résistance aux pulvérisations / Pressione della resistenza agli spruzzi / Splash Resistance Pressure	kPa	–*	–*	≥ 120*
Mikrobiologische Reinheit / Propreté microbienne / Purezza microbiologica / Microbial cleanliness	cfu/g	≤ 30	≤ 30	≤ 30

* Typ II R ist ein flüssigkeitsresistenter Typ. / Le type II R est un type résistant aux liquides
 Tipo II R è un tipo resistente ai liquidi. / Type II R is a liquid resistant type.

Die in dieser Verpackung enthaltenen medizinischen Masken dienen vor allem dem Fremdschutz und schützen das Gegenüber des Maskenträgers vor der Exposition möglicher infektiöser Tröpfchen. Primäre Zweckbestimmung: Anwendung im medizinischen Umfeld um die Patienten zu schützen. Diese medizinischen Masken schützen bei festem Sitz begrenzt auch den Träger der Maske, dies ist jedoch nicht die primäre Zweckbestimmung.



Medizinische Masken
Masques Médicaux
Mascherine Mediche
Medical Face Masks
 50x

Medizinische Masken Typ II R – EN 14683

REF AN-00-2-R
 LOT 201218
 12-2020
 12-2023

Produkt Medizinische Masken Typ II R, EN14683:2019, 3-Schichten, Ohrschlaufe
 Masques Médicaux Type II R, EN14683:2019, 3-plis, fixation auriculaire élastique
 Maschere Mediche Tipo II R, EN14683:2019, 3-strati, occhiella per orecchie
 Medical Face Mask Type II R, EN14683:2019, 3-layers, Ear Loop
 Medische maskers, 3 lagen, oortje / Medicinska masker, 3 lager, öronslinga
 Maski medyczne, 3 warstwy, pętla na ucho / Medicinske maske, 3 sloja, ušna petlja
 Medicinska masker, 3 lager, öronslinga / Orvosi maszkok, 3 réteg, fülhurok

Importeur Stöckli Medical AG, Länggasse 4, CH-6208 Oberkirch
 info@stoecklimedical.ch, +41 41 925 66 55

Xiantao Yi-Ya Protective Products Co., Ltd.
 Miandong Road Hefeng Village Xiliuhe Town,
 Xiantao City, Hubei Province, China

EC REP Luxus Lebenswelt GmbH, Kochstrasse 1, 47877, Willich, Germany

