

SUBJECT Physical & Microbiological Test

TEST LOCATION TÜV SÜD China

TÜV SÜD Products Testing (Shanghai) Co., Ltd. B-3/4, No.1999 Du Hui Road, Minhang District

Shanghai 201108, P.R. China

CLIENT NAME Anhui Tiankang Medical Technology Co.,Ltd

CLIENT ADDRESS No.228 Weiyi Road Development Zone Tianchang City 239300 Anhui China

TEST PERIOD 10-Apr -2020~18-Apr-2020

Prepared By

Bella Xu

(Bella Xu)
Report Drafter

Authorized By

Leo Bu)
Authorized Signatory

Note: (1) General Terms & Conditions as mentioned overleaf. (2) The results relate only to the items tested. (3) The test report shall not be reproduced except in full without the written approval of the laboratory. (4) Without the agreement of the laboratory, the client is not authorized to use the test results for unapproved propaganda.

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TEST REPORT

Sample Description Single-use Medical Face Mask

Sample Quantity 45 pieces Lot Number/Batch Code 200308 Specification With earloop

Size

Type of Mask Type IIR

Brand Name

Remark: The above information was provided by applicant.

Summary of Test Results

No.	Test Item	Test Standard	Judgement
1	Bacterial Filtration Efficiency (BFE) Test	EN 14683:2019+AC:2019(E) Annex B	Pass
2	Differential Pressure Test	EN 14683:2019+AC:2019(E) Annex C	Pass
3	Synthetic Blood Penetration Test	ISO 22609:2004	Pass
4	Microbial Cleanliness Test	EN 14683:2019+AC:2019(E) Annex D	Pass

Note: Pass = Meet customer requirements;

Fail = Fail customer requirements;

= No comment;

N.D. = Not detected.





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Results

No.	Test Item	Test Result
		Specimen 1#: 99.4%
		Specimen 2#: 99.0%
1	Bacterial Filtration Efficiency (BFE) Test	Specimen 3#: 99.5%
		Specimen 4#: 99.5%
		Specimen 5#: 99.1%
2	Differential Pressure Test	49.8 Pa/cm ²
3	Synthetic Blood Penetration Test	Specimen 1#~13#: None seen
	.0	Specimen 1#: <1 CFU/g
		Specimen 2#: <1 CFU/g
4	Microbial Cleanliness Test	Specimen 3#: <1 CFU/g
		Specimen 4#: <1 CFU/g
	(E)	Specimen 5#: <1 CFU/g

Bacterial Filtration Efficiency (BFE) Test

1. Purpose

For evaluating the bacterial filtration efficiency (BFE) of mask.

2. Sample description was given by client

Sample description : Single-use Medical Face Mask

Specification : With earloop Lot Number : 200308 Sample Receiving Date : 2020-04-10

3. Test Method

EN 14683:2019+AC:2019(E) Annex B

4. Apparatus and materials

- 4.1 Staphylococcus aureus ATCC 6538.
- 4.2 Peptone water.
- 4.3 Tryptic Soy Broth(TSB).
- 4.4 Tryptic Soy Agar(TSA).
- 4.5 Bacterial filtration efficiency test apparatus.
- 4.6 Six-stage viable particle Anderson sampler.
- 4.7 Flow meters.

5. Test specimen

- 5.1 As requested by client, take a total of 5 test specimens.
- 5.2 Prior to testing, condition all test specimens for a minimum of 4 h at (21±5)°C and (85±5)% relative humidity.

Chemical/Microbiology Laboratory: TÜV SÜD Products Testing (Shanghai) Co., Ltd.

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6. Procedure

- 6.1 Preparation of the bacterial challenge: Dilute the cultutre in peptone water to achieve a concentration of approximately 5×10⁵ CFU/mL.
- 6.2 Adjust the flow rate through the Anderson sampler to 28.3 L/min.
- 6.3 Deliver the challenge to the nebulizer using a syringe pump. Purge tubing and nebulizer of air bubbles.
- 6.4 Perform a positive control run without a test specime to determine the number of viable aerosol particles being generated. The mean particle size (MPS) of the aerosol will also be calculated from the results of these positive control plates.
 - 6.4.1 Initiate the aerosol challenge by turning on the air pressure and pump connected to the nebulizer. Immediaterly begin sampling the aerosol using the Anderson sampler.
 - 6.4.2 Time the challenge suspension to be delivered to the nebulizer for 1 min.
 - 6.4.3 Time the air pressure and Anderson sampler to run for 2 min.
 - 6.4.4 At the conclusion of the positive control ran, remove plates from the Anderson sampler.
- 6.5 Place new agar plates into Anderson sampler and clamp the test specimen into the top of the Anderson sampler, with the inside of the specimen facing towards the bacterial challenge (test area: 77cm²).
- 6.6 Repeat the challenge procedure for each test specimen.
- 6.7 Repeat a positive control after completion of the sample set.
- 6.8 Perform a negative control run by collecting a 2 min sample of air from the aerosol chamber. No bacterial challenge should be pumped into the nebulizer during the collection of the negative control.
- 6.9 Incubate agar plates at (37±2)°C for (20 to 52) h.
- 6.10 Count each of the six-stage plates of the Anderson sampler.

7. Calculation

Total the count from each of the six plates for the test specimens and positive controls, as specified by the manufacture of Anderson sampler. The filtration efficiency percentages are calculated as follows:

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

T is the total plate count for the test specimen.

C is the mean of the total plate counts for the two positive controls.

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8. Test results*

P Value	Positive	Positive	Negative	Specimen	Specimen	Specimen	Specimen	Specimen
Stage	Control (A)	Control (B)	Control	1#	2#	3#	4#	5#
Number								
1	50	234	0	0	0	0	0	0
2	82	116	0	0 %	0	0	0	0
3	151	406	0	0	0	0	0	0
4	277	644	0	0	0	0	0	0
5	1241	547	0	11	8	5	8	11
6	513	721	0	5	17	7	5	12
Total (T), CFU	2314	2668	<1	16	25	12	13	23
Average (C), CFU	2.5x10 ³ = ((PA+PB) / 2	/			11000		
BFE ,%	- GIII.		(ED)	99.4	99.0	99.5	99.5	99.1
Requirements	OLIO	// \		≥	98		.000	
Remarks	P is the value of corresponding corrected particle counts as specified by the manufacturer of the cascade impactor. T is the total of P value for the test specimen. C is the mean of the total of P value of the two positive controls.				er of the			



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Differential pressure Test

1.Purpose

The purpose of the test was to measure the differential pressure of masks.

2. Sample description was given by client

Sample description : Single-use Medical Face Mask

Specification : With earloop Lot Number : 200308 Sample Receiving Date : 2020-04-10

3.Test Method

EN 14683:2019+AC:2019(E) Annex C

4. Apparatus and materials

Differential pressure testing instrument

5.Test specimen

- 5.1 Test specimen are complete masks or shall be cut from masks. Each specimen shall be able to provide 5 different circular test areas of 2.5 cm in diameter.
- 5.2 Prior to testing, condition all test specimens for a minimum of 4 h at (21 ± 5) °C and (85 ± 5) % relative humidity.

6. Procedure

- 6.1 Without a specimen in place, the holder is closed and the differential manometer is zeroed. The pump is started and the flow of air adjusted to 8 L/min.
- 6.2 The pretreated specimen is placed across the orifice (total area 4.9cm², test area diameter 25mm) and clamped into place so as to minimize air leaks.
- 6.3 Due to the presence of an alignment system the tested area of the specimen should be perfectly in line and across the flow of air.
- 6.4 The differential pressure is read directly.
- 6.5 The procedure described in steps 6.1-6.4 is carried out on 5 different areas of the mask and readings averaged.

Results:

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Specimen	Test Results* (Pa/cm²)	Average (Pa/cm²)	Requirements	Judgement
1#	48.3			
2#	52.4	lui.		
3#	46.3	49.8	< 60	Pass
4#	54.2	(E),		
5#	47.7			

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Synthetic Blood Penetration Test

1.Purpose

For evaluation of resistance of masks to penetration by a fixed volume of synthetic blood at a high velocity.

2. Sample description was given by client

Sample description : Single-use Medical Face Mask

Specification : With earloop

Lot Number : 200308 Sample Receiving Date : 2020-04-10

3.Test Method

ISO 22609:2004

4. Apparatus and materials

- 4.1 Synthetic blood.
- 4.2 Tensiometer.
- 4.3 Synthetic blood penetration test apparatus;
- 4.4 Targeting plate.
- 4.5 Air pressure source.
- 4.6 Ruler.
- 4.7 Balance.
- 4.8 Controlled temperature and humidity chamber.

5.Test specimen

- 5.1 As requested by client, take a total of 13 test specimens.
- 5.2 Prior to testing, condition all test specimens for a minimum of 4h at (21±5)°C and (85±5) % relative humidity.

6.Procedure

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- 6.1 Prepare the synthetic blood (40~44 mN/m) for the test.
- 6.2 Determine the density of the synthetic blood.
- 6.3 Fill the reservoir with new synthetic blood.
- 6.4 Position the test specimen 30.5 cm (12 in.) from the exit of the canula.
- 6.5 Set the reservoir pressure to the approximate pressure.
- 6.6 Place the targeting plate approximately 1 cm away from the mask.
- 6.7 Set the valve timer to 0.5 s. Collect and weigh the amount of fluid delivered (before the targeting hole).

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- 6.8 Set the valve timer to 1.5 s. Collect and weigh the amount of fluid delivered (before the targeting hole).
- 6.9 Calculate the difference in weight of the two spurts. For a test fluid with a density of 1.003, Table 1 gives the target difference in weight plus lower and upper limits for a velocity range within 2% of the target.

Table 1 Target weight difference

Fluid Pressure (mmHg)	Weight difference for 1 s difference in spurt duration (g)			
	Min.	Target	Max.	
120	3.002	3.063	3.124	

- 6.10 Adjust the reservoir pressure and repeat steps 6.7 to 6.9 until the weight difference is within the target range.
- 6.11 Record the weight difference for the spurts exiting the nozzle.
- 6.12 Record the pressure in the reservoir.
- 6.13 Set the valve time to 0.5 s. Collect and weigh the amount of fluid passing through the targeting hole.
- 6.14 Set the valve time to 1.5 s. Collect and weigh the amount of fluid passing through the targeting hole.
- 6.15 The difference in weight between the 0.5 s and 1.5 s spurts through the targeting plate shall be within $+2\% \sim -5\%$ of the difference in weight from the nozzle.
- 6.16 If the differential weight is less than 95 % of the weight difference exiting the nozzle, check the aim of the stream to make sure it is passing cleanly through the targeting hole.
- 6.17 If the differential weight is more than 102 % of the weight difference exiting the nozzle, repeat the weight measurements exiting the nozzle (steps 6.7 to 6.11).
- 6.18 For standard synthetic blood, the timer duration can be estimated using the formula: (p is the density of the test fluid.) $t = 0.5 + (2 \times p - g)$ at 0.5 s) / (g at 1.5 s - g at 0.5 s).
- 6.19 Record the timer setting to use as the starting point for subsequent testing.
- 6.20 Mount a test specimen on the specimen holding fixture. If the mask contains pleats, spread the pleats out when mounting the mask onto the fixture to present a single layer of material as the target
- 6.21 Squirt the synthetic blood onto the test specimen for the calculated time. Ensure that the synthetic blood hits the target area of mask.
- 6.22 Inspect the inside surface for synthetic blood penetration within 10 s of squirting the synthetic blood against the target area.
- 6.23 Report the results (none / penetration) for each test specimen at the test pressure.

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Results:

Specimen	Test Results*	Requirements	Judgement
1#	None Seen		Pass
2#	None Seen		Pass
3#	None Seen	Cilip)	Pass
4#	None Seen	20	Pass
5#	None Seen	1,0,	Pass
6#	None Seen	. Cilip	Pass
7#	None Seen	Pass Pressure at 16.0 kPa (120mmHg)	Pass
8#	None Seen	(1201111111g)	Pass
9#	None Seen	181	Pass
10#	None Seen	111111111111111111111111111111111111111	Pass
11#	None Seen		Pass
12#	None Seen	l'is.	Pass
13#	None Seen	1 14	Pass



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Microbial Cleanliness Test

1. Purpose

The purpose of the test was to measure microbial cleanliness of mask.

2. Sample description was given by client

Sample description : Single-use Medical Face Mask

Specification : With earloop

Lot Number : 200308 Sample Receiving Date : 2020-04-10

3. Test Method

According to EN ISO 11737-1:2018 to determine the microbial cleanliness of mask material, and refer to the procedure as described in EN 14683:2019+AC:2019(E) Annex D

4. Apparatus and materials

- 4.1 Orbital shaker.
- 4.2 0.45 um filter.
- 4.3 Tryptic Soy Agar (TSA).
- 4.4 Sabouraud Dextrose Ager (SDA) with chloramphenicol.
- 4.5 Formula of Extraction Liquid: 1g/L peptone, 5g/L NaCl and 2g/L Tween 20.
- 4.6 Extraction apparatus.

5. Test specimen

- 5.1 As requested by client, take a total of 5 mask samples.
- 5.2 Mask samples for testing are provided in the original primary packaging.
- 5.3 Condition at (18 to 26) °C and (45 to 65)% relative humidity during testing.

6. Procedure

- 6.1 Five test specimens are selected from the top, bottom and 3 randomly chosen marks.
- 6.2 The mask is aseptically removed from the packaging and placed in a sterile 500 mL bottle containing 300 mL of extraction liquid.
- 6.3 The bottle is laid down on an orbital shaker and shaken for 5 min at 250 rpm.
- 6.4 After extracting, 100mL of the extraction liquid is filtered through a 0.45 um 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 SDA for fungi enumeration.
- 6.5 The plates are incubated for 3 days at 30°C and 7 days at (20 to 25)°C for TSA and SDA plates respectively.
- 6.6 Calculate the colonies of each agar plate.

7. Calculation

For each test specimen calculate the microbial cleanliness as follows by counting the total colonies of the TSA and SDA plates.

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Results*:

Specimen	Colonies of the TSA Plate	Colonies of the SDA Plate	Microbial Cleanliness, (CFU/g)	Requirements	Judgement
1#	0	0	<1		
2#	0	0	<1	According to EN ISO 11737-1:2018 the	
3#	0	0	<1	microbial cleanliness of	Pass
4#	0	0	<1	the mask shall be ≤30 CFU/g tested.	
5#	0	0	<1		

Note:

1.*denotes this test was carried out by external laboratory assessed as competent.

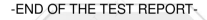
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2. This report is for internal use only such as internal scientific research ,education, quality control, product R&D.







Certificate

The Certification Body of TÜV Rheinland LGA Products GmbH

hereby certifies that the organization

Anhui Tiankang Medical Technology Co., Ltd. No. 228 Weiyi Road Economic Development Zone Tianchang City 239300 Anhui China

has established and applies a quality management system for medical devices for the following scope:

Manufacture and Distribution of Medical Devices
(see attachment for products and additional site included)

Proof has been furnished that the requirements specified in

EN ISO 13485:2016

are fulfilled. The quality management system is subject to yearly surveillance.

Effective Date: 2018-09-17

Certificate Registration No.: SX 60131062 0001

An audit was performed. Report No.: 15096003 004

This Certificate is valid until: 2021-09-16

Certification Body



Date 2018-09-17



TÜV Rheinland LGA Products GmbH - Tillystraße 2 - 90431 Nürnberg

Tel.: +49 221 806-1371 Fax: +49 221 806-3935 e-mail:cert-validity@de.tuv.com http://www.tuv.com/safety

1.0 EC Declaration of Conformity

Name: Anhui Tiankang Medical Technology Co., Ltd.

Manufacturer:

Add: No. 228 Weiyi Road, Economic Development Zone, Tianchang

City, 239300 Anhui, China

European

Name: MedPath GmbH

Representative:

Add: Mies-van-der-Rohe-Strasse 8, 80807 Munich, Germany

Product Name:

Single-use Medical Face Mask

(Type I,Type II,Type IIR)

Object of the declaration:

Types/Sizes

9cm,17cm x8cm,17cm x 9cm,17cm x 9.5cm,17cm x 10cm,17.5cm x 8cm,17.5cm x 9cm,17.5cm x 9.5cm,17.5cm x 10cm,18cm x

8cm,18cm x 9cm,18cm x9 .5cm,18cm x 10cm,19.5cm x 8cm,19.5cm x 9cm,19.5cm x 9.5cm,19.5cm x 10cm

UMDNS Code: 12-447

Classification (MDD, Annex IX): I, rule 1
Conformity Assessment Route: Annex VII

We herewith declare in sole responsibility that the above mentioned products meet the transposition into national law, the provisions of the following EC Council Directives and Standards such as EN 14683:2019+AC:2019 etc. All supporting documentations are retained under the premises of the manufacturer. We are exclusively responsible for the declaration of conformity.

DIRECTIVES

General applicable directives:

Medical Device Directive: COUNCIL DIRECTIVE 93/42/EEC

Start of CE Marking: 2020-03-25

Place of Issue: Tianchang, CHINA

Date of Issue: 2020-04-23

Signature:

Mr. Baodong Bai

Position: General Manager



EC-Registration Certificate

Directive 93/42/EEC on Medical Devices (MDD), Article 14 No. R A001 29 Rev. 01

MedPath

Manufacturer: Anhui Tiankang Medical Technology Co., Ltd.

No. 228 Weiyi Road, Economic Development Zone, Tianchang City, 239300 Anhui, China

Inyuy

Product

See Appendix A

CE

Category(ies):

This is to certify that, in accordance of the Medical Device Directive 93/42/EEC (amended by 2007/47/EC), MedPath GmbH agrees to perform all duties and responsibilities as the Authorized Representative for the aforementioned manufacturer as stipulated and demanded by the aforementioned Directive. The German Competent Authority is notified of the manufacturer's medical device(s) and has allocated registration numbers shown in Appendix A. The manufacturer has provided MedPath GmbH with the appropriate Declaration(s) of Conformity confirming that the medical device(s) fulfills/fulfill the applicable requirements of the aforementioned Directive.

MedPath GmbH Mies-van-der-Rohe-Strasse 8-D-80807 München

Tel.089-189174474 · Fax 089-54858884

Date, 2020-03-28

MedPath GmbH





Appendix A: Product Category(ies)

MedPath

No.	Name	Class	UMDNS Code	Form No.	Registration No.
1	Single-use Medical Face Mask	I	12-447	00297923	to be issued
2	Non-woven Coveralls	I	15-223	00297925	to be issued
3	Non-woven Isolation Gowns	1	15-037	00297927	to be issued



MedPath GmbH Mies-van-der-Rohe-Strasse 8 · D-80807 München Tel.089-189174474 · Fax 089-54858884



BfArM - Bundesinstitut für Arzneimittel und Medizinprodukte

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4 von 9 BfArM: MP Anzeigen (MPA) © DIMDI

Dokumentnummer	00159181	

Anzeige

Registrierdatum	2020-05-13
Registriernummer	DE/CA61/1M50/121
Typ der Anzeige	Erstanzeige
Anzeigender nach § 25 MPG	Bevollmächtigter
Formularnummer	00297923

Angaben zum Anzeigenden

Code	DE/0000047823
Bezeichnung	MedPath GmbH
Staat	Deutschland
Ort	München
Postleitzahl	80807
Straße, Haus-Nr.	Mies-van-der-Rohe-Strasse 8
Land	Bayern
Telefon	089 189174474
E-Mail	info@medpath.pro

Zuständige Behörde

Code	DE/CA61
Bezeichnung	Regierung von Oberbayern
Staat	Deutschland
Land	Bayern
Ort	München
Postleitzahl	80534
Straße, Haus-Nr.	Maximilianstraße 39
Telefon	+49-89-21760
Telefax	+49-89-21762914
E-Mail	medizinprodukteanzeigeverfahren@reg-ob.bayern.de

Hersteller

Bezeichnung	Anhui Tiankang Medical Technology Co., Ltd.
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Ort	Tianchang City
Postleitzahl	239300
Straße, Haus-Nr.	No. 228 Weiyi Road, Economic Development Zone
Telefon	+86 13705505106
E-Mail	zy@tkmedical.com

Medizinprodukt

Produkttyp	nichtaktives Medizinprodukt
Klasse	I
App (Software auf mobilen Endgeräten)	Nein
Handelsname	Single-use Medical Face Mask
Nomenklaturcode	12-447
Nomenklaturbezeichnung	Maske
Kategorie	10 Produkte zum Einmalgebrauch

zurück in der Dokumentausgabe blättern weiter



Anhui Tiankang Medical Technology Co., Ltd

No.228 Weiyi Road Economic Development Zone Tianchang City Anhui China

bestätigt

IEDAU INTERNATIONAL GmbH

Pallaswiesenstrasse 63, 64293 Darmstadt Deutschland Hessenring 32, 64546 Mörfelden-Walldorf Deutschland

die Zusammenarbeit in Europa



Markenautorisierungszertifizierung

代理经销授权书

Hiermit wird bescheinigt, dass die **IEDAU International GmbH** mit Sitz in Pallaswiesenstr. 63 64293 Darmstadt, Hessenring 32 64546 Mörfelden-Walldorf Deutschland offiziell zur Nutzung der Marke "TKMD" berechtigt und als **AUTORISIERTER VERTRIEBSPARTNER** in **EUROPA** ist.

Die IEDAU International GmbH hat die Verantwortung die Marke und die Produkte zu vermarkten, zu vertreiben, zu verkaufen und technische Unterstützung anzubieten.

Diese Vereinbarung gilt vom 01.01.2020 bis zum 31.12.2020.

兹有位于安徽省天长市经济开发区纬一路228号的安徽天康医疗科技股份有限公司,特授权位于德国黑森州达姆施塔特的IEDAU International GmbH为我公司在欧洲市场的代理经销商,IEDAU International GmbH有权使用我司的"TKMD"商标并经营销售我公司制造的口罩、医用人员用面罩等产品。

特此授权。

授权期限: 自2020年4月1日起至2021年3月31日止。

Für Fragen wenden Sie sich bitte an/ 如有疑问请联系:

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Pallaswiesenstr. 63 64293 Darmstadt Deutschland

Hessenring 32 64546 Mörfelden-Walldorf Deutschland

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Anhui Tiankang Medical Technology Co., and

制造商:安徽天康医疗科技服份有限公司

Unterschrift Geschäftsführung

Datum/日期

2024-4,13

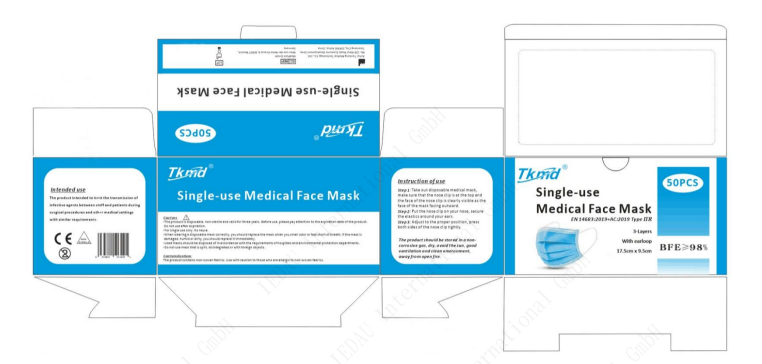
为有限公司



EDAN INTERNATIONAL CHOI

CACI

Sechs Grundansichten



Vorderansicht



Rückansicht



Single-use Medical Face Mask



- Cautions /!\
 -The product is disposable, non-sterile and valid for three years. Before use, please pay attention to the expiration date of the product.
- Do not use after expiration.
- -For single use only. No reuse.
- -When wearing a disposable mask correctly, you should replace the mask when you smell odor or feel short of breath; if the mask is damaged, humid or dirty, you should replace it immediately.
- Used masks should be disposed of in accordance with the requirements of hospitals and environmental protection departments.
- Do not use mask that is split, disintegrated or with foreign objects.

The product contains non-woven fabrics. Use with caution to those who are allergic to non-woven fabrics.

Draufsicht





Single-use Medical Face Mask



Anhui Tiankang Medical Technology Co., Ltd. No. 228 Weiyi Road, Economic Development Zone, Tianchang City, 239300 Anhui, China

Mies-van-der-Rohe-Strasse 8, 80807 Munich, Germany



200403 20200403 20230402

Ansicht von unten



Seitenansicht von links

Instruction of use

Step 1: Take out disposable medical mask, make sure that the nose clip is at the top and the face of the nose clip is clearly visible as the face of the mask facing outward.

Step 2: Put the nose clip on your nose, secure the elastics around your ears.

Step 3: Adjust to the proper position, press both sides of the nose clip tightly.





(() EN 14683:2019

NON-STERILE

Seitenansicht von rechts

