

Prüfbericht-Nr.: Test Report No.:	60384703 001	Auftrags-Nr.: Order No.:	244246788	Seite 1 von 14 Page 1 of 14
Kunden-Referenz-Nr.: Client Reference No.:	2110099	Auftragsdatum Order date:	: 19.06.2020	
Auftraggeber: Client:	2nd Floor, 7th Bu	<b>Medical Products Co., L</b> ilding, Electronic Industry nic and Technological Dev 226010, China	Park, No. 9, Xinde	ong Road,
<b>Prüfgegenstand:</b> Test item:	Disposable Surg	gical Face Mask		
Bezeichnung / Typ-Nr.: Identification / Type No.:	EARLOOP			
Auftrags-Inhalt: Order content:	Type test			
Prüfgrundlage: Test specification:	EN 14683:2019+	AC:2019 (except for Cla	use 5.2.6 Biocon	npatibility)
Wareneingangsdatum: Date of receipt:	22.06.2020	8		
<b>Prüfmuster-Nr.:</b> Test sample No.:	A002851411-001	del		
<b>Prüfzeitraum:</b> Testing period:	23.06.2020 to 17.07.2020		<u>r</u>	
<b>Ort der Prüfung:</b> Place of testing:	See page 3			
<b>Prüflaboratorium:</b> Testing laboratory:	TÜV Rheinland (Shanghai) Co., L	.td.	001 001 001 001 001	
<b>Prüfergebnis*:</b> Test result*:	Pass			
<b>geprüft von</b> / tested by:		kontrolliert vo	n / reviewed by:	
2.07.2020 Rainbow Pan/PE	A	22.07.2020 Xiac	jun Ding/Reviewer	Kino jan Din
DatumName/StellungDateName/Position	Unterschrift Signature	<b>Datum</b> Date	Name/Stellung Name/Position	Unterschrift Signature
onstiges / Other: he test report consists of EN 1468 lause 5.2.6 Biocompatibility is not				
Zustand des Prüfgegenstar Condition of the test item at d	elivery:	Test item comple	ändig und unbeso ete and undamag	
Legende: 1 = sehr gut 2 = g P(ass) = entspricht o.g. Prüf		digend tspricht nicht o.g. Prüfgrundlage(n)	4 = ausreichend N/A = nicht anwendbar	5 = mangelhaft N/T = nicht getestet
Legend: 1 = very good 2 = g P(ass) = passed a.m. tests		5	4 = sufficient N/A = not applicable	5 = poor N/T = not tested
•	gtwerden.DieserB	ericht berechtigt nicht zur '	Verwendung eines	s Prüfzeichens.
his test report only relates to the a	a. m. test sample. Wi	thout permission of the test c s. This test report does not ei		



EN 14683:2019+AC:2019 Medical face masks —			
Req	uirements and test methods		
Report Reference No:	See cover page		
Date of issue:	See cover page		
Total number of pages::	See cover page		
Testing Laboratory:	TÜV Rheinland (Shanghai) Co., Ltd.		
Address:	No.177, 178, Lane 777 West Guangzhong Road, Jing'an District, Shanghai, China		
Applicant's name:	Nantong Jianan Medical Products Co., Ltd		
Address:	2nd Floor, 7th Building, Electronic Industry Park, No. 9, Xindong Road, Nantong Economic and Technological Development Zone, Jiangsu Province 226010, China		
Test specification:			
Standard:	EN 14683:2019+AC:2019		
Test procedure:	Type test		
Non-standard test method	N/A		
Test Report Form No:	EN 14683:2019+AC:2019_A		
Test Report Form Originator:	TÜV Rh (SZ)		
Master TRF:	2020-03		
Test item description:	Disposable Surgical Face Mask		
Trade Mark:	N/A		
Manufacturer:	Same as applicant		
Model/Type reference:	EARLOOP		
Classification:	Type IIR		

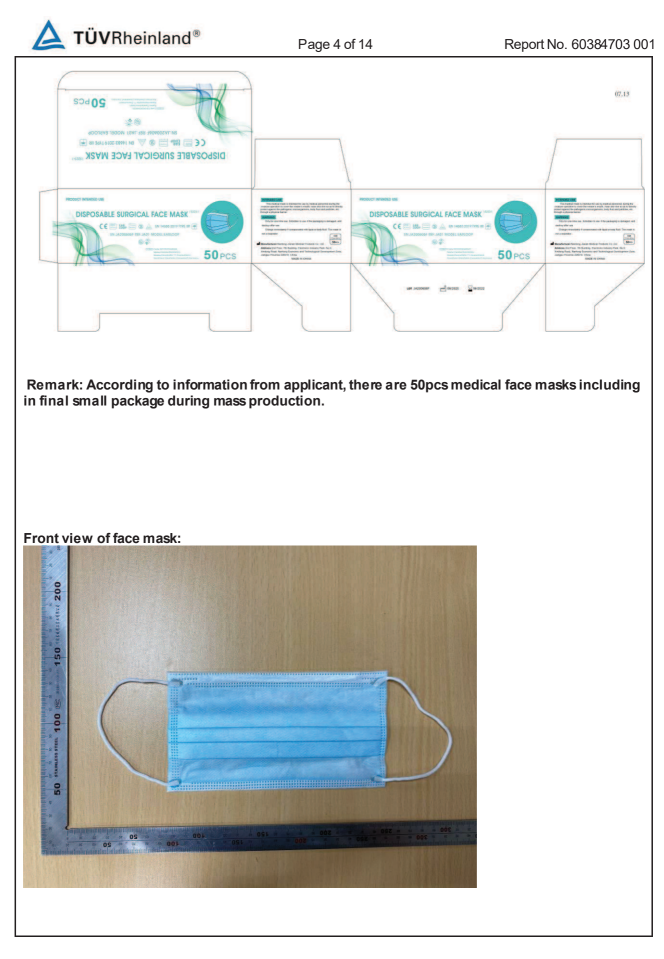


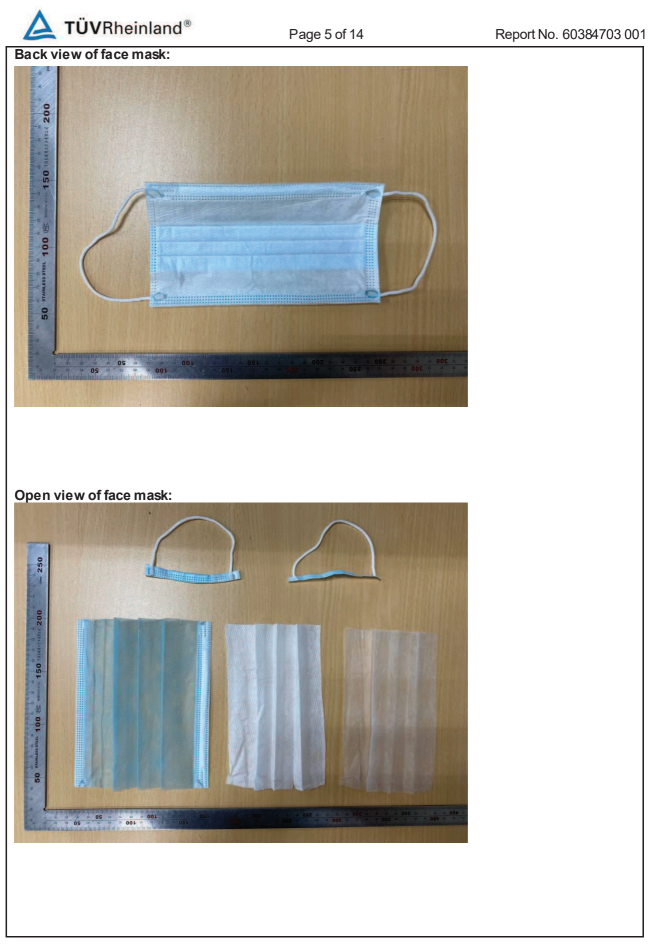
List of Attachments (including a total number of pages in	List of Attachments (including a total number of pages in each attachment):			
N/A				
Summary of testing:				
Tests performed (name of test and test clause):	Testing location:			
<b>Construction check was performed according to:</b> Clause 5.1.1 Materials and construction; Clause 5.1.2 Design	<b>TÜV Rheinland (Shanghai) Co., Ltd.</b> No.177, 178, Lane 777 West Guangzhong Road, Jing'an District, Shanghai, China			
<b>Other tests were performed:</b> Clause 5.2.2 Bacterial filtration efficiency; Clause 5.2.3 Breathability; Clause 5.2.4 Splash resistance; Clause 5.2.5 Microbial cleanliness	<b>Pony Testing Group Shanghai Co.,Ltd.</b> 2/3/4/6/F., Building 35, No.680, Guiping Road, Xuhui District, Shanghai, China			
Note: All tests listed as above have been conducted in the competent external lab under the supervision of a TUV engineer.				

# Copy of marking plate

The artwork below may be only a draft. The use of certification marks on a product must be authorized by the respective NCBs that own these marks.

Box:





QMF-RT-33008SHG

Revision number: 1.0





### Testing

Date of receipt of test item(s):	
Dates of tests performed:	See cover page
Possible test case verdicts:	
- test case does not apply to the test object: :	N/A
- test object does meet the requirement: :	P (Pass)
- test object was not evaluated for the requirement $\ldots$ :	N/E (collateral standards only)
- test object does not meet the requirement: :	F (Fail)
General remarks:	
"(See Attachment #)" refers to additional information a "(See appended table)" refers to a table appended to The tests results presented in this report relate only to	the report. o the object tested.

This report shall not be reproduced except in full without the written approval of the testing laboratory. List of test equipment must be kept on file and available for review.

Additional test data and/or information provided in the attachments to this report.

Throughout this report a  $\Box$  comma /  $\boxtimes$  point is used as the decimal separator.

Name and address of factory (ies) .....: Same as applicant

#### General product information:

The submitted samples are type IIR, non-sterile disposable surgical face mask which are intended for use by medical personnel during the invasive operation to cover the wearer's mouth, nose and chin so as to directly protect against the pathogenic microorganisms, body fluid and particles, etc. through a physical barrier.

Clause 5.2.6 Biocompatibility is not evaluated in this test report.

The test results are for reference only. Relevant certification may be needed if the mask is intended to be sold in Europe.



	EN 14683:2019+AC:20	19	
Clause	Requirement + Test	Result - Remark	Verdict
4	Classification		Р
	Medical face masks specified in this European Standard are classified into two types (Type I and Type II) according to bacterial filtration efficiency whereby Type II is further divided according to whether or not the mask is splash resistant. The 'R' signifies splash resistance.	Type IIR	Р
5	Requirements		Р
5.1	General		Р
5.1.1	Materials and construction		Р
	The medical face mask is a medical device, generally composed of a filter layer that is placed, bonded or moulded between layers of fabric.	Composed of a filter layer between layers of fabric	Р
	The medical face mask shall not disintegrate, split or tear during intended use.	Complied	Р
	In the selection of the filter and layer materials, attention shall be paid to cleanliness.	Considered	Р
5.1.2	Design		Р
	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.	Fitted closely over nose	Р
	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).	With a nose bridge	P
5.2	Performance requirements		Р
5.2.1	General		Р
	All tests shall be carried out on finished products or samples cut from finished products.	Complied	Р
5.2.2	Bacterial filtration efficiency (BFE)		Р
	When tested in accordance with Annex B, the BFE of the medical face mask shall conform to the minimum value given for the relevant type in Table 1.	See appended table 5.2.2	Р
	For thick and rigid masks such as rigid duckbill or cup masks the test method may not be suitable as a proper seal cannot be maintained in the cascade impactor. In these cases, another valid equivalent method shall be used to determine the BFE.	Not thick and rigid mask	N/A



	EN 14683:2019+AC:20	19	
Clause	Requirement + Test	Result - Remark	Verdict
	When a mask consists of two or more areas with different characteristics or different layer-composition, each panel or area shall be tested individually.	No such condition	N/A
	The lowest performing panel or area shall determine the BFE value of the complete mask		N/A
5.2.3	Breathability		Р
	When tested in accordance with Annex C, the differential pressure of the medical face mask shall conform to the value given for the relevant type in Table 1.	See appended table 5.2.3	Р
	If the use of a respiratory protective device as face mask is required in an operating theatre and/or other medical settings, it might not fulfil the performance requirements with regard to differential pressure as defined in this European Standard. In such case, the device should fulfil the requirement as specified in the relevant Personal Protective Equipment (PPE) standard(s).	No such respiratory protective device	N/A
5.2.4	Splash resistance		Р
	When tested in accordance with ISO 22609:2004 the resistance of the medical face mask to penetration of splashes of liquid shall conform to the minimum value given for Type IIR in Table 1.	See appended table 5.2.4	Р
5.2.5	Microbial cleanliness (Bioburden)		Р
	When tested according to EN ISO 11737-1:2018 the bioburden of the medical mask shall be $\leq$ 30 CFU/g tested (see Table 1).	See appended table 5.2.5	Р
5.2.6	Biocompatibility		N/E
	According to the definition and classification in EN ISO 10993-1:2009, a medical face mask is a surface device with limited contact.		N/E
	The manufacturer shall complete the evaluation of the medical face mask according to EN ISO 10993-1:2009 and determine the applicable toxicology testing regime.		N/E
	The results of testing should be documented according to the applicable parts of the EN ISO 10993 series.		N/E
	The test results shall be available upon request.		N/E
6	Marking, labelling and packaging		Р
	Annex I, §13, of the Medical Devices Directive (93/42/EEC) or Annex I, §23, of the Medical Device Regulation (EU) 2017/745 specifies the information that should be specified on the packaging in which the medical face mask is supplied.	Checked and complied	Р
	The following information shall be supplied:		Р
	a) number of this European Standard;	Marked on the label	Р

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	EN 14683:2019+AC:2019					
Clause	e Requirement + Test Result - Remark Verdic					
	b) type of mask (as indicated in Table 1).	Marked on the label	Р			
	EN ISO 15223-1:2016 and EN 1041:2008+A1:2013 should be considered.	Considered	Р			



	EN 14683:2019+AC:2019			
Clause	Requirement + Test	Result - Remark	Verdict	

5.2.2	5.2.2 TABLE: Bacterial filtration efficiency (BFE)					Р		
Batch/ lot no.:	Test Speci -men no.:	Dimension of the test specimen L x W (mm x mm)	test area <i>(</i> cm²)	Flow rate (I/min)	Mean of the total plate counts of the two positive controls	Mean plate count of the negative controls	BFE for each test specimen (%)	Remarks
A00285	1	150×135	95	28.3	1733	0	>99.9	Р
1411- 001	2	150×135	95	28.3	1733	0	99.8	Р
	3	150×135	95	28.3	1733	0	99.9	Р
	4	150×135	95	28.3	1733	0	99.9	Р
	5	150×135	95	28.3	1733	0	>99.9	Р

#### Supplementary information:

1, Each specimen was conditioned at <u>21.1</u> °C and <u>85.0</u> % relative humidity for <u>4</u> h to bring them into equilibrium with atmosphere prior to testing.

2, The side of the test specimen was facing towards the challenge aerosol: <u>face</u>

### Remark:

Limit value: Type I ≥95%; Type II≥98%; Type IIR ≥98%.

5.2.3 TABLE: Breathability (Differential pressure)						Р
Batch/ lot no.:	Test Specimer number- Test area number	Differential pressure for each test area (Pa/cm²)	The averaged differential pressure for each test specimen (Pa/cm <sup>2</sup> )	Flow rate (I/min)	Remarks	
A0028	1-1	49.7	48.6	8.0		Р
51411- 001	1-2	47.7		8.0		Р
	1-3	48.4		8.0		P
	1-4	47.7	-	8.0	Р	P
	1-5	49.6		8.0		P
	2-1	52.8	53.5	8.0		Р
	2-2	54.6		8.0		Р
	2-3	52.5		8.0		Р
	2-4	53.3		8.0		Р
	2-5	54.5		8.0		Р
	3-1	50.6	50.3	8.0		Р



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Clause	e Requirement + Test			Result - Remark Verdio		
	3-2	51.1		8.0	Р	
	3-3	49.1		8.0	Р	
	3-4	51.8		8.0	Р	
	3-5	49.0		8.0	Р	
	4-1	50.0	50.1	8.0	Р	
	4-2	49.1		8.0	Р	
	4-3	50.8		8.0	Р	
	4-4	51.1		8.0	Р	
	4-5	49.7		8.0	Р	
	5-1	52.7	52.5	8.0	Р	
	5-2	53.0		8.0	Р	
	5-3	51.8		8.0	Р	
	5-4	53.0		8.0	Р	
	5-5	51.8		8.0	Р	

## Supplementary information:

Each specimen was conditioned at 21.0 °C and 85.0 % relative humidity for 4 h to bring them into equilibrium with atmosphere prior to testing.

## Remark:

Limit value: Type I <40; Type II <40; Type IIR <60.

5.2.4	5.2.4 TABLE: Splash resistance				Р	
Batch/ lot	ot no.: Test mask The material of tested Test result I no.: mask (Pass/fail)		Re	emarks		
A002851411-001		1	Polypropylene fused jet filter layer	Pass		
		2	Polypropylene fused jet filter layer	Pass		
		3	Polypropylene fused jet filter layer	Pass		
		4	Polypropylene fused jet filter layer	Pass		
		5	Polypropylene fused jet filter layer	Pass		
		6	Polypropylene fused jet filter layer	Pass		
		7	Polypropylene fused jet	Pass		



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Clause	Requirement +	Test	F	Result - Remark	Verdict
			filter layer		
		8	Polypropylene fused je filter layer	et Pass	-
		9	Polypropylene fused je filter layer	et Pass	-
		10	Polypropylene fused je filter layer	et Pass	
		11	Polypropylene fused je filter layer	et Pass	
		12	Polypropylene fused je filter layer	et Pass	
		13	Polypropylene fused je filter layer	et Pass	
		14	Polypropylene fused je filter layer	et Pass	
		15	Polypropylene fused je filter layer	et Pass	
		16	Polypropylene fused je filter layer	et Pass	
		17	Polypropylene fused je filter layer	et Pass	
		18	Polypropylene fused je filter layer	et Pass	
		19	Polypropylene fused je filter layer	et Pass	
		20	Polypropylene fused je filter layer	et Pass	
		21	Polypropylene fused je filter layer	et Pass	
		22	Polypropylene fused je filter layer	et Pass	
		23	Polypropylene fused je filter layer	et Pass	
		24	Polypropylene fused je filter layer	et Pass	
		25	Polypropylene fused je filter layer	et Pass	
		26	Polypropylene fused je filter layer	et Pass	
		27	Polypropylene fused je filter layer	et Pass	



		EN 14683:2019+AC:201	19	
Clause	Requirement + Test		Result - Remark	Verdict
	28	Polypropylene fused j filter layer	iet Pass	
	29	Polypropylene fused j filter layer	iet Pass	
	30	Polypropylene fused j filter layer	iet Pass	-
	31	Polypropylene fused j filter layer	iet Pass	
	32	Polypropylene fused j filter layer	iet Pass	-

### Supplementary information:

1, Each specimen was conditioned at 21.0 °C and 85.0 % relative humidity for 4 h to bring them into equilibrium with atmosphere prior to testing.

2, The description of target area tested: the center of outside

- 3, Any technique used to enhance visual detection of synthetic blood: cotton swab
- 4, The temperature and relative humidity for testing: <u>24.3</u>°C and <u>81.2</u> %

5, Description of any pre-treatment techniques used: constant temperature and humidity machine was used

#### Remark:

Limit value: not required for Type I and Type II; Type IIR ≥16,0.

5.2.5	TABLE: Mi	TABLE: Microbial cleanliness (Bioburden)						
Batch/ lot no.:		Mask(under test) no.:	Weight of each mask (g)	Total bioburden per individual mask (CFU/g)	Remarks			
A002851411-001		1	2.87	24	Р			
		2	2.89	27	Р			
		3	2.84	19	Р			
		4	2.85	17	Р			
		5	2.83	11	Р			

Limit value: Type I ≤30; Type II ≤30; Type IIR ≤30.

# End of test report