







Test Report SL52025257428201TX-1 Date:September 04,2020 Page 1 of 5

MULTIBRANDS INTERNATIONAL LTD ROYDS HALL, ROYDS HALL LANE, BRADFORD, ENGLAND BD12 0EJ

THIS REPORT CANCELS AND SUPERSEDES THE TEST REPORT NO. SL52025257428201TX DATE: June 06,2020 ISSUED BY SGS (SHANGHAI) UPDATED SAMPLE INFORMATION&TEST INFORMATION

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

Sample Description : (A)Type IIR Medical mask

Style No. : FB2RM-U

Composition : (A)Non-woven fabrics

Sample Color : (A)Blue outside and white inside

Buyer : MULTIBRANDS INTERNATIONAL LTD
Manufacturer : MULTIBRANDS INTERNATIONAL LTD

Country of Destination : United Kingdom

Lot No./Batch No. : PO12168

Proposed Care Instruction:

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

Sample Receiving Date : May 13, 2020

Testing Period : May 13, 2020 - Jun 06, 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)

Dongjing Liu (Authorized Signatory)



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Test Result

### EN 14683:2019+AC:2019 Medical Face Masks-Requirements and Test Methods

### Clause 5.2 Performance Requirement

### Clause 5.2.2 Bacterial filtration efficiency (BFE)\*

(EN 14683 :2019 Annex B)

Conditioning Parameters : Minimum of 4 hours at 22.0°C and 85.0% R.H.

Dimensions of test specimen : 150 mm x 130 mm

Test Area : 133 cm²
Test Side : Inside
Flow Rate : 28.3 l/min
Positive Control Average : 1906 CFU
Negative Monitor Count : < 1 CFU

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

Remark: Performance Requirement: Type I≥95%, Type II≥98%, Type IIR≥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).



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## Clause 5.2.3 Breathability

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

Sample: A

Test Side : Randomly test in different location (1 around and 4 away from the centric

point) on each of the 5 masks

Pre-Conditioning : Minimum of 4 hours at 21±5°C and 85±5% R.H.

Test Area : 4.9 cm<sup>2</sup> Flow Rate : 8 l/min

Specimen No.	Test Area No.	Different Pressure for each tested area (Pa/cm²)	The average value for each test specimen (Pa/cm²)	
	1-1	51.1	, ,	
	1-2	45.0		
1	1-3	42.8	47	
	1-4	45.9		
	1-5	51.9		
	2-1	48.1	45	
	2-2	43.5		
2	2-3	44.6		
	2-4	43.8		
	2-5	46.7		
3	3-1	49.8	47	
	3-2	43.7		
	3-3	47.1		
	3-4	45.8		
	3-5	48.2		
4	4-1	46.2	47	
	4-2	44.8		
	4-3	45.7		
	4-4	47.6		
	4-5	48.2		
5	5-1	45.9	47	
	5-2	46.0		
	5-3	48.2		
	5-4	45.4		
	5-5	47.8		

# Remark:

- 1) Performance Requirement: Type I<40 Pa/cm², Type II<40 Pa/cm², Type IIR<60 Pa/cm²
- 2) The number of specimens that shall be tested is minimum 5, but can be greater and shall be increased if necessary to allow for an AQL(Acceptable Quality Level) of 4%.



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### Clause 5.2.4 Splash Resistance

(ISO 22609:2004)

Sample: A

Test Blood Pressure : 16.0kPa

Pre-Conditioning : Minimum of 4 hours at 21±5°C and 85±5% R.H.

Distance of the mask to the tip of cannula : 300±10mm

Test Specimen#	Penetration on inside surface	Conclusion	Test Specimen#	Penetration on inside surface	Conclusion
1	None Seen	Pass	17	None Seen	Pass
2	None Seen	Pass	18	None Seen	Pass
3	None Seen	Pass	19	None Seen	Pass
4	None Seen	Pass	20	None Seen	Pass
5	None Seen	Pass	21	None Seen	Pass
6	None Seen	Pass	22	None Seen	Pass
7	None Seen	Pass	23	None Seen	Pass
8	None Seen	Pass	24	None Seen	Pass
9	None Seen	Pass	25	None Seen	Pass
10	None Seen	Pass	26	None Seen	Pass
11	None Seen	Pass	27	None Seen	Pass
12	None Seen	Pass	28	None Seen	Pass
13	None Seen	Pass	29	None Seen	Pass
14	None Seen	Pass	30	None Seen	Pass
15	None Seen	Pass	31	None Seen	Pass
16	None Seen	Pass	32	None Seen	Pass
Number of Pass:		32			
Overall result:			Acceptable		

#### Remark:

- 1) Performance Requirement Type I: N/A, Type II: N/A, Type IIR: ≥16.0kPa
- 2) Test was conducted within 60s after removal from conditioning chamber.
- 3) 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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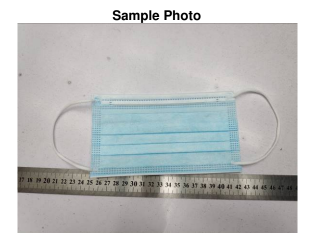
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### **Clause 5.2.5 Microbial Cleanliness**

(EN 14683:2019+AC:2019 Annex D and EN ISO 11737-1:2018)

	Mask Weight(g)	Total Bioburden, (cfu/mask)	Total Bioburden, (cfu/g)
Sample Number		,	( 0/
1#	3.31	39	11.78
2#	3.29	33	10.03
3#	3.32	24	7.23
4#	3.39	9	2.65
5#	3.33	18	5.41

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



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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