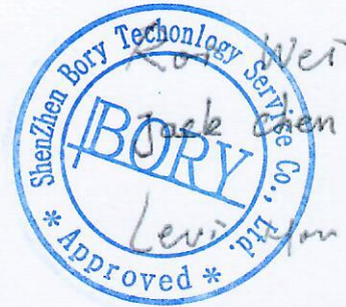


TEST REPORT EN 14683:2019 Medical face masks - Requirements and test methods	
Report Number	B2003TR9132S02
Tested by (+ signature)	Roi Wei
Compiled by (+ signature)	Jack Chen
Approved by (+ signature)	Levi Hou
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Total number of pages	17 pages
Testing Laboratory	Shenzhen Bory Technology service Co., Ltd.
Address	3F, Baijiasheng Business building, BaoTian Industrial, Qianjin 2 Rd., Bao'an District, Shenzhen, Guangdong, China
Address	As above
Applicant's name	Fujian Opooger Technology Co.Ltd
Address	No:1-2 A Business Building Hanjiang Aquatic Wholesale Market Hanxi Office Hanjiang Region Putian Fujian China
Test specification:	
Standard	EN 14683:2019
Test procedure	CE
Non-standard test method	N/A
Test Report Form No	EN 14683:2019
Test Report Form(s) Originator	BORY
Master TRF	Dated 2020-02
The test results presented in this report relate only to the object tested. This report shall not be reproduced except in full, without the written approval of the Laboratory. The authenticity of this Test Report and its contents can be verified by contacting the Laboratory, responsible for this Test Report.	
Test item description	KN 95 mask/face mask/mouth mask/Stereoscopic respirator
Trade Mark	OPOGER
Manufacturer	Putian City Hongtian Industrial Trading Co.LTD
Address	No:1-2 10th Building Canglin Area Hanjiang Region Putian City
Model/Type reference	OPOGER KN95



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List of Attachments (including a total number of pages in each attachment):

-- Attachment 1: One pages for Photo documentation.

Summary of testing:**Tests performed (name of test and test clause):**

-- EN 14683:2019

Medical face masks - Requirements and test methods

Testing location:

3F, Baijiasheng Business building, BaoTian Industrial, Qianjin 2 Rd., Bao'an District, Shenzhen, Guangdong, China

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.

(Additional requirements for markings. See 1.7 NOTE)

OPOGER

KN 95 mask/face mask/mouth mask/

Stereoscopic respirator

OPOGER KN95

EN 14683: 2019



Type I



<p>Possible test case verdicts:</p> <ul style="list-style-type: none"> - test case does not apply to the test object : N/A(or N) - test object does meet the requirement : P (Pass) - test object does not meet the requirement : F (Fail)
<p>Testing..... :</p> <p>Date of receipt of test item..... : Feb. 24, 2020</p> <p>Date (s) of performance of tests : Feb. 24, 2020 – Mar.26, 2020</p>
<p>General remarks:</p>
<p>Throughout this report a <input checked="" type="checkbox"/> comma / <input type="checkbox"/> point is used as the decimal separator.</p>
<p>General product information:</p> <p>The product is intended to be used as medical face masks intended to lirrít the transmission of infective agents from staff to patients during: surgical procedures and other medical settings with similar requirements.</p>

EN 14638:2019			
Clause	Requirement Test	Result-Remark	Verdict
4	Classification		P
	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	<input checked="" type="checkbox"/> Type I <input type="checkbox"/> Type II	P
5	Requirements		P
5.1	General		P
5.1.1	Materials and construction		P
	The medical face mask is a medical device, generally composed of a filter layer that is placed, bonded or moulded between layers of fabric.	Filter layer composed	P
	The medical face mask shall not disintegrate, split or tear during intended use.	Not disintegrate, split or tear	P
	In the selection of the filter and layer materials, attention shall be paid to cleanliness (absence of particulate matter).	Cleanliness has been considered.	P
5.1.2	Design		P
	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.		P
	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).		P
5.2	Performance requirements		P
5.2.1	All tests shall be carried out on finished products or samples cut from finished products, if applicable in their sterile state.	Carried out on finished product.	P
5.2.2	Bacterial filtration efficiency (BFE)	Type I; $\geq 95\%$	P
	When tested in accordance with Annex B, the bacterial filtration efficiency (BFE) of the medical face mask shall conform to the minimum value given for the relevant type in Table 1.	BFE was tested in accordance with Annex B.	P
5.2.3	Breathability	$< 29,4 \text{ Pa/cm}^2$	P
	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.	Breathability was tested in accordance with Annex C.	P

EN 14638:2019			
Clause	Requirement Test	Result-Remark	Verdict
5.2.4	Splash resistance	Not required.	N/A
	When tested in accordance with ISO 22609 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		N/A
5.2.5	Microbial cleanliness (Bioburden)	≤ 30 cfu/g	P
	When tested according to EN ISO 11737-1 the bioburden of the medical mask shall be ≤ 30 cfu/g tested.	Bioburden was tested according to EN ISO 11737- 1.	P
	EN ISO 11737-1 specifies requirements and provides guidance for the enumeration and microbial characterisation of the population of viable microorganisms on or in a medical device, component, raw material or package.		P
	To determine the mask's bioburden according to EN ISO 11737-1, follow the procedure below:		P
	<input type="checkbox"/>The number of masks that shall be tested is minimum 5 (five), but can be greater if necessary to allow for an AQL of 4 %.	5 specimens	P
	Weigh each mask prior testing	Weights were recorded	P
	The full mask is aseptically removed from the packaging and placed in a sterile 500 ml bottle containing 300 ml of extraction liquid (1 g/l Peptone, 5 g/l NaCl & 2 g/l polysorbate surfactant 20 [e.g. Tween 20, Alkest TW 20]).	Extraction liquid: 300 ml.	P
	The bottle is laid down on an orbital shaker and shaken for 5 min at 250 rpm	Shaken: 250 rpm	P
	After this extraction step, 100 ml of the extraction liquid is filtered through a 0,45 µ filter and laid down on a TSA plate for the total viable aerobic microbial count.	Filter: 0,45 µ	P
	Another 100 ml aliquot of the same extraction liquid is filtered in the same way and the filter plated on Sabouraud Dextrose agar (SDA) with chloramphenicol for fungi enumeration.		P
	The plates are incubated for 3 days at 30 °C and 7 days at (20 – 25) °C for TSA and SDA plates respectively.	30 °C: 3 days; 20 °C– 25 °C: 7 days.	P
	The total bioburden is expressed by addition of the TSA and SDA counts.		P
	In the report, indicate the total bioburden per mask and based on the mask weigh, the total bioburden per gram tested.		P
5.2.6	Biocompatibility	In accordance with EN ISO 10993-1.	P

EN 14638:2019			
Clause	Requirement Test	Result-Remark	Verdict
	According to the definition and classification in EN ISO 10993-1, a medical face mask is a surface device with limited contact.		P
	The manufacturer shall complete the evaluation of the medical face mask according to EN ISO 10993-1 and determine the applicable toxicology testing regime.	In accordance with EN ISO 10993-1.	P
	The results of testing should be documented according to the applicable parts of the EN ISO 10993 series.		P
	The test results shall be available upon request.		P
	As a minimum, EN ISO 10993-5 and EN ISO 10993-10 shall be considered. 5.2.7 Summary of performance requirements		P
5.2.7	Summary of performance requirements	a Bacterial filtration efficiency (BFE): $\geq 95\%$; b Differential pressure: $< 29,4 \text{ Pa/cm}^2$; c Splash resistance pressure: Not required; d Microbial cleanliness: $\leq 30 \text{ cfu/g}$.	P

Table 1 - Performance requirements for medical face masks			
Test	Type Ia	Type II	Type IIR
Bacterial filtration efficiency (BFE), (%)	≥ 95	≥ 98	≥ 98
Differential pressure (Pa/cm ²)	$< 29,4$	$< 29,4$	$< 49,0$
Splash resistance pressure (kPa)	Not required	Not required	$\geq 16,0$
Microbial cleanliness (cfu/g)	≤ 30	≤ 30	≤ 30
<p>a Type I medical face masks should only be used for patients and other persons to reduce the risk of spread of infections particularly in epidemic or pandemic situations.</p> <p>Type I masks are not intended for use by healthcare professionals in an operating room or in other medical settings with similar requirements.</p>			

6	Labelling and information to be supplied		P
	Annex I, §13, of the Medical Devices Directive (93/42/EEC) specifies the information that has to be specified on the packaging in which the medical face mask is supplied.		P
	The following information shall be supplied in addition:		P
	a).... umber of this European Standard	EN 14683:2019	P

EN 14638:2019			
Clause	Requirement Test	Result-Remark	Verdict
	b).... type of mask (as indicated in Table 1).	Type I	P
	EN ISO 15223-1 and EN 1041 should be considered.		P
A	Information for users		P
	When breathing, speaking, coughing, sneezing etc., one releases smaller or larger amounts of droplets of secretions from the mucous membranes in the mouth and nose.		P
	The majority of the nuclei are between 0,5 µm and 12 µm in diameter and especially the larger droplets can contain micro-organisms from the source site.	Nuclei: 0,5 µm - 12 µm.	P
	Nuclei can subsequently spread through the air to a susceptible site such as an open operating wound or sterile equipment.		P
	The medical face masks intended to be used in operating rooms and health care settings with similar requirements are designed to protect the entire working environment.	It was designed to protect the entire working environment	P
	This standard describes two types of medical face masks with associated protection levels.	Type I medical face mask.	P
	As a minimum, Type I medical face masks are used for patients in order to reduce the risk of the spread of infections, particularly in epidemic or pandemic situations.		P
	Type II masks are principally intended for use by healthcare professionals in an operating room or other medical settings with similar requirements.	Type I medical face mask	N/A
	A special case, also covered by the European Medical Devices legislation, is that in which the wearer wishes to protect him/herself against splashes of potentially contaminated fluids and particles that are created in the surgical environment, e.g. by the use of electro-cautery devices.		P
	If the intended use of the mask is to protect the wearer against infective agents (bacteria, viruses or fungi), the use of a respirator device should be considered		N/A
	Performance requirements for respirators are the scope of EN 149	In accordance with EN 149.	P
	The level of efficiency offered by a mask depends on a number of factors such as the filtration efficiency, quality of the material and the fit of the mask on the wearer's face.		P
	Different designs are suited for different applications and the careful choice of mask is therefore important in order to achieve the desired result.		P

EN 14638:2019			
Clause	Requirement Test	Result-Remark	Verdict
	The filtration capacity of mask materials can vary depending on the filter media.		P
	The fit of masks varies considerably from those which are held in place by ear loops fastened behind the wearer's ears to those with tie bands around the head and a nose clamp that can be shaped to the wearer's nose.	Can be shaped to the wearer's nose.	P
	The effect of a very good or less good fit can be tested in vivo whereas the filtration efficiency may be reproducibly tested in vitro.	Effect can be tested in vivo	P
	The considerable variations in results when masks are tested in vivo results in the need for large groups of test subjects and observations.	No any variations.	N/A
	It is thus usual to characterise mask performance using in vitro tests of the material from which the mask is made		P
	It is, however, important to consider the fit of the mask carefully when a mask for a certain application is chosen. Users should request such information from their suppliers.		P
	A further factor to be considered is the capacity of the mask to absorb moisture from the exhaled air and thereby to maintain its performance over a longer period of time.		P
	The more advanced designs easily maintain their performance throughout even very long operations whereas the less advanced ones are intended only for short procedures.		P
	The contamination risk resulting from hand contact with a used mask means that it is essential that the mask is taken off and disposed of when no longer worn over nose and mouth.	No longer worn over nose and mouth.	P
	When there is a further need for protection then a new mask should be put on.	No further need for protection.	N/A
	Touching a used face mask or putting on a new one should always be followed by a full hand disinfection procedure and a used mask should always be disposed of when no longer needed or between two procedures.		P
	In summary, to use an appropriate mask is an effective means to protect the working environment from droplet contamination from nose and throat during health care procedures.		P
	Masks with very different performance are, however, available.	Without very different performance.	N/A
	Therefore such factors as infection risk and mask fit		P

EN 14638:2019			
Clause	Requirement Test	Result-Remark	Verdict
	should be carefully considered when choosing a mask.		
B	Method for in-vitro determination of bacterial filtration efficiency (BFE)		P
	WARNING:		P
	<input type="checkbox"/>Staphylococcus aureus is a pathogen.		P
	<input type="checkbox"/>The relevant national provisions by law and hygienic instructions when dealing with pathogens shall be complied with.		P
B.1	Principle		P
	A specimen of the mask material is clamped between a six-stage cascade impactor and an aerosol chamber		P
	An aerosol of Staphylococcus aureus is introduced into the aerosol chamber and drawn through the mask material and the impactor under vacuum.		P
	The bacterial filtration efficiency 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.		P
B.2	Reagents and materials		P
B.2.1	General		P
	B.2.2 and B.2.3 describe commercially available solutions of tryptic soy agar and tryptic soy broth.		P
	Other variants may be suitable.		P
B.2.2	Tryptic soy agar		P
	Formula/liter:		P
	<input type="checkbox"/>Enzymatic digest of casein.....	15g	P
	<input type="checkbox"/>Enzymatic digest of soybean meal....	5g	P
	<input type="checkbox"/>Sodium chloride.....	5g	P
	<input type="checkbox"/>Agar.....	15g	P
	<input type="checkbox"/>Final pH.....	7,3 ± 0,2 at 25 °C	P
B.2.3	Tryptic soy broth		P
	Formula/liter		P
	<input type="checkbox"/>Enzymatic digest of casein	17g	P
	<input type="checkbox"/>Enzymatic digest of soybean meal.....	3g	P
	<input type="checkbox"/>Sodium chloride	5g	P
	<input type="checkbox"/> Dipotassium phosphate	2.5g	P

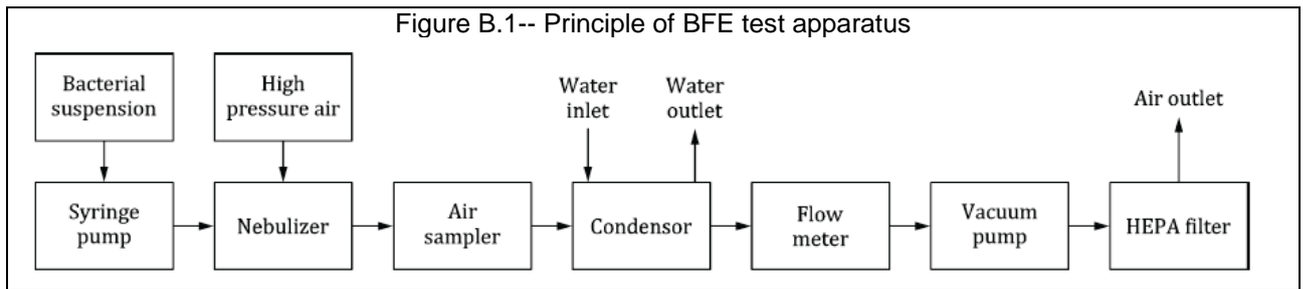
EN 14638:2019			
Clause	Requirement Test	Result-Remark	Verdict
	<input type="checkbox"/> Dextrose..	2.5g	P
	<input type="checkbox"/>Final pH....	7,3 ± 0,2 at 25 °C	P
B.2.4	Peptone water		P
	Formula/liter		P
	<input type="checkbox"/>Peptone..	10g	P
	<input type="checkbox"/>Sodium chloride..	5g	P
	<input type="checkbox"/>Final pH....	7,2 ± 0,2 at 25 °C	P
B.2.5	Culture of Staphylococcus aureus ATCC 6538, growing on tryptic soy agar slants		P
B.3	Apparatus		P
B.3.1	Six stage cascade impactor		P
B.3.2	Nebulizer, capable of delivering particles with a mean size of (3,0 ± 0,3) µm when in contact with the impactor	3,1 µm	P
B.3.3	Aerosol chamber, glass, 600 mm long and 80 mm in external diameter	Long: 600 mm; Diameter: 80 mm.	P
B.3.4	Flow meters, capable of measuring a flow rate of 28,3 l/min	Flow rate: 28,3 l/min.	P
B.3.5	Pressure gauge, capable of measuring a pressure of 35 kPa to an accuracy of ± 1 kPa	35 kPa	P
B.3.6	Erlenmeyer flasks, 250 ml and 500 ml capacity		P
B.3.7	Peristaltic or syringe pump, capable of delivering 0,01 ml/min	0,01 ml/min	P
B.3.8	Vacuum pump, capable of maintaining a flow rate of 57 l/min	Flow rate: 57 l/min.	P
B.4	Test apparatus		P
B.4.1	Six stage cascade impactor , the arrangement is specified in Table B.1.		P
B.4.2	Nebulizer , capable of delivering particles with a mean size of (3,0 ± 0,3) µm when in contact with the cascade impactor.		P
B.4.3	Aerosol chamber , glass, 600 mm long and 80 mm in external diameter.		P
B.4.4	Flow meters , capable of measuring a flow rate of 28,3 l/min.		P
B.4.5	Pressure gauge , capable of measuring a pressure of 35 kPa to an accuracy of ± 1 kPa.		P
B.4.6	Erlenmeyer flasks , 250 ml and 500 ml capacity.		P
B.4.7	Peristaltic or syringe pump , capable of delivering 0,01 ml/min.		P

EN 14638:2019			
Clause	Requirement Test	Result-Remark	Verdict
B.4.7	Vacuum pump , capable of maintaining a flow rate of 57 l/min.		P
B.5	Test specimens		P
	Test specimens shall be cut from complete masks.		P
	Each specimen shall be minimum 100 mm by 100 mm and shall include all layers of the mask in the order in which they are placed in the complete mask.		P
	The number of specimens that shall be tested is minimum 5 (five), but can be greater and shall be increased if necessary to allow for an AQL of 4 %.	5 specimens	P
	All specimens tested shall be taken from representative areas to incorporate all/any variation in construction.		P
	Unless otherwise specified, the testing shall be performed with the inside of the medical face mask in contact with the bacterial challenge.		P
	Each test specimen shall be conditioned at $(21 \pm 5) ^\circ\text{C}$ and $(85 \pm 5) \%$ relative humidity for the time required to bring them into equilibrium with atmosphere prior to testing.		P
B.6	Preparation of bacterial challenge	37 °C for 24 h	P
	Staphylococcus aureus (see B.2.4) shall be inoculated into 30 ml tryptic soy broth in an Erlenmeyer flask and incubated with mild shaking at a temperature of $(37 \pm 2) ^\circ\text{C}$ for (24 ± 2) h.	5×10^5 cfu/ml	P
	The culture shall then be diluted in peptone water to give a concentration of approximately 5×10^5 cfu/ml.	2300 cfu	P
	The bacterial challenge shall be maintained at $1,7 \times 10^3$ to $3,0 \times 10^3$ CFU per test.		P
	The bacterial challenge shall be determined on the basis of experience and previous positive control plates (see B.7.3) and the dilution of the challenge suspension adjusted accordingly.		P
	The mean particle size in the bacterial challenge shall be maintained at $(3,0 \pm 0,3) \mu\text{m}$ (see B.7.9).	3,1 μm	P
B.7	Procedure		P
B.7.1			
	Assemble the apparatus in accordance with the flow chart shown in Figure B.1.		P
B.7.2	Deliver the bacterial challenge to the nebulizer using the peristaltic or syringe pump.		P
B.7.3	Perform a positive control run without a test specimen. Initiate the bacterial challenge by turning on the vacuum pump and adjust the flow rate through the	Flow rate: 28,3 l/min.	P

EN 14638:2019			
Clause	Requirement Test	Result-Remark	Verdict
	cascade impactor to 28,3 l/min.		
	Deliver the bacterial challenge for 1 min		P
	Maintain the airflow through the impactor for 2 min.	2 min	P
	Then remove the plates from the impactor.		P
	Ensure that each plate is numbered to indicate its position in the impactor.		P
B.7.4	Place fresh plates in the impactor, fix a test specimen in place and repeat the above procedure.		P
B.7.5	Repeat this procedure for each test specimen.		P
B.7.6	After the last test specimen has been tested, perform a further positive control run		P
B.7.7	Perform a negative control run by passing air, without addition of the bacterial challenge, through the cascade impactor for 2 min.	2 min	P
B.7.8	Incubate all the plates at (37 ± 2) °C for (20 to 52) h. 3	37 °C for 48 h	P
B.7.9	For each specimen and control run, count the number of colonies on each plate and add up the counts to give the total number of cfu collected by the impactor using the “positive hole” conversion table1) in accordance with the instructions of the cascade impactor manufacturer (stages 3 to 6).		P
	For the two positive control runs, take the mean of the two totals		P
	From the positive control plates calculate the mean particle size of the bacterial challenge aerosol using the “positive hole” conversion table in accordance with the instructions of the cascade impactor manufacturer.		P
B.8	Calculation of bacterial filtration efficiency		P
	For each test specimen calculate the bacterial filtration efficiency B, as a percentage, using the formula.	$B = (C - T) / C \times 100$ <p>C is the mean of the total plate counts for the two positive control runs;</p> <p>T is the total plate count for the test specimen</p>	P
B.9	Test report	<p>a number and date of this European Standard;</p> <p>b lot number or batch code of the masks tested;</p> <p>c dimensions of the test specimens and the</p>	P

EN 14638:2019			
Clause	Requirement Test	Result-Remark	Verdict
		size of the area tested; d which side of the test specimen was facing towards the challenge aerosol; e flow rate during testing; f mean of the total plate counts of the two positive controls; g mean plate count of the negative controls; h bacterial filtration efficiency for each test specimen.	

Figure B.1-- Principle of BFE test apparatus



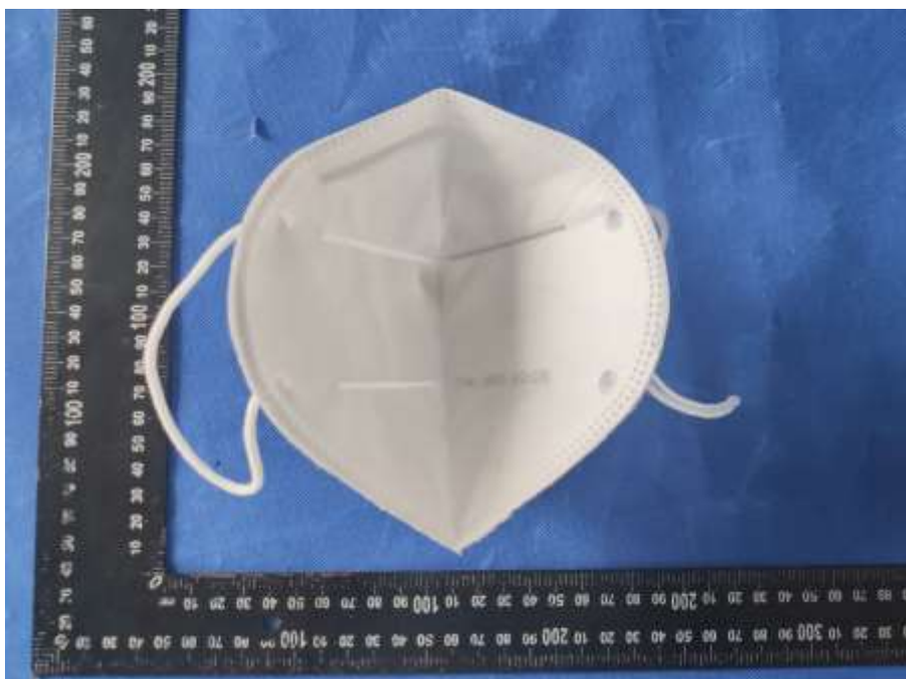
C	Method for determination of breathability (differential pressure)	P
C.1	Principle	P
	A device which measures the differential pressure required to draw air through a measured surface area at a constant air flow rate is used to measure the air exchange pressure of the medical face mask material, as shown in Figure C.1.	P
	Water-filled manometers (M1 and M2) are used to measure the differential pressure.	P

EN 14638:2019			
Clause	Requirement Test	Result-Remark	Verdict
	A flow meter is used for measurement of the airflow.		P
	An electric vacuum pump draws air through the apparatus and a needle valve is used to adjust the airflow rate		P
C.2	Apparatus		P
C2.1	Flow meter, capable of measuring an airflow of 8 l/min	Airflow: 8 l/min	P
C2.2	Manometers, M1 and M2 or differential manomete		P
C2.3	Electric vacuum pump including a pressure buffer tank		P
C2.4	Valve permitting the adjustment of the flow rate		P
C2.5	Sample holder		P
C2.5.1	The sample holder shall consist of a mechanical clamping system and alignment of the top and bottom holder.		P
C2.5.2	The sample holder shall consist of a mechanism to adjust the clamping pressure. A system with thread of screw can be used either at the bottom or top part of the sample holder.		P
C2.5.3	The internal diameter of the top holder and the bottom holder in the contact area with the filter material shall be (25 ± 1) mm.		P
C2.5.4	The seal of the top and bottom holder onto the filter material shall consist of a metal-metal contact.		P
C2.5.5	Validation of the test apparatus shall consist of a leak test. A second flow meter (12) placed immediately before the valve (10) will allow for evaluation of an air leak within the test apparatus. With the sample holder closed, start the pump and adjust the flow meter to read 8 l/min on the first flow meter (2). If no leaks are present both flow meters should read 8 l/min.		P
C.3	Test specimens		P
	Test specimens are complete masks or shall be cut from masks.	Complete mask.	P
	Each specimen shall be able to provide 5 different circular test areas of 2,5 cm in diameter.	Diameter: 2,5 cm	P
	If one specimen cannot provide 5 test areas of 2,5 cm in diameter, the number of test areas retrieved should be representative for the entire mask.	Diameter: 2,5 cm	P
	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 of 4 %.	5 specimens	P
	All specimens tested shall be taken from areas representative from the mask to incorporate all/any variation in construction.		P

EN 14638:2019			
Clause	Requirement Test	Result-Remark	Verdict
	Each test specimen shall be conditioned at $(21 \pm 5) ^\circ\text{C}$ and $(85 \pm 5) \%$ relative humidity for the time required to bring them into equilibrium with atmosphere prior to testing.	$(21 \pm 2) ^\circ\text{C}$; $(85 \pm 2) \%$	P
C.4	Procedure		P
C.4.1	The test specimen is placed across the 2,5 cm diameter orifice (total area 4,9 cm ²) and clamped into place so as to minimise air leaks and that the tested area of the specimen will be in line and across the flow of air.	Area: 4,9 cm ²	P
C.4.2	The pump is started and the flow of air adjusted to 8 l/min.		P
C.4.3	The manometers M1 and M2 are read and recorded.		P
C.4.4	The procedure described in steps C.4.1 through C.4.3 is carried out on 5 (or appropriate number of) different areas of the mask and the readings averaged.		P
C.5	Calculation of differential pressure		P
	For each test specimen calculate the differential pressure ΔP as.	$\Delta P = (X_{m1} - X_{m2})/4,9$	P
C.6	Test report	a number and date of this European Standard; b lot number or batch code of the masks tested; c flow rate during testing; d differential pressure for each test specimen.	P
ZA	Relationship between this European Standard and the Essential Requirements of EU Directive 93/42/EEC on medical devices		P
	This European Standard has been prepared under a mandate given to CEN by the European Commission Union to provide a means of conforming to the essential requirements of New Approach EU Directive 93/42/EEC concerning medical devices.		P
	Once this standard is cited in the Official Journal of the European Union under that Directive and has been implemented as a national standard in at least one Member State, compliance with the clauses of this standard given in Table ZA.1 confers, within the limits of the scope of this standard, a presumption of conformity with the corresponding Essential Requirements of that Directive and associated EFTA		P

EN 14638:2019			
Clause	Requirement Test	Result-Remark	Verdict
	regulations.		
ZA.1	Correspondence between this European Standard and EU Directive 93/42/EEC concerning medical devices		P
	Clause/subclause of this European Standard; 5.1.1, 5.1.2, 5.2.1, 5.2.2, 5.2.3, 6.....	Corresponding Essential Requirement of Directive 93/42/EEC: 8.1	P
	Clause/subclause of this European Standard; 5.2.2....	Corresponding Essential Requirement of Directive 93/42/EEC: 9.2	P
	Clause/subclause of this European Standard; 6.	Corresponding Essential Requirement of Directive 93/42/EEC: 13	P
	WARNING 1 — Presumption of conformity stays valid only as long as a reference to this European Standard is maintained in the list published in the Official Journal of the European Union. Users of this standard should consult frequently the latest list published in the Official Journal of the European Union.		P
	WARNING 2 —Other requirements and other EU Directives may be applicable to the product(s) falling within the scope of this standard.		P

Photo-documentation



===== End of test report=====