



FFP3 CE1463 EN 149:2001+A1:2009









FFP3
(€1463
EN 149:2001+A1:2009



FEATURES

Ergonomic design, comfortable and good protection; Adjustable ear loop, no hurt to ears; With valve, easy breath High filtration efficiency Material:non-woven fabrics/Meltblown fabric/ mask belt



anti-virus, anti dust, anti droplets etc.

型 号	描述	包 装	认 证 号
CARE0750	FFP3 FFP3 NR EN 149:2001 +A1:2009	1PC/BAG 20PCS/BOX 50BOXES/CARTON	CE 1463





FITTING INSTRUCTIONS: See figure:

- 1. Cup respirator in one hand with nose piece at fingertips, allow headbands to hang freely below hand.
- 2. Hold respirator under chin, with nose piece up.
- 3. Locate the upper strap across the crown of the head and the lower strap below the ears .
- 4. Straps must not be twisted.
- 5. Using both hands, mould nose clip to the shape of the lower part of the nose to unsure a close fit and good seal. Pinching the nose clip using only one hand may result in less effective respirator performance.
- 6. The seal of the respirator on the face should be fitchecked before intering the workplace.
- 7. If the user feels the bottom strap is too loose and it is not possible to achieve a satisfactory fit of the mask a knot can be tied in the strap.



DISCARD AND REPLACE THE MASK IF:

- 1. The mask is removed whilst in the contaminated area.
- 2. Cloggong of the respirator causes breathing difficulties.
- 3. The mask becomes damaged.
- 4. For respirators protecing against vapours, the smell of vapours present becomes detectable.

FAILURE TO FOLLOW THE INSTRUCTIONS AND WARNINGS ON THE USE OF THIS MASK DURING ALL TIMES OF EXPOSURE CAN REDUCE THE EFFECTIVENESS OF THE MASK AND COULD RESULT IN ILLESS OR DISABILITY.

MANUFACTURER INDENTIFICATIONS:

Technical information Following the standard EN 149:2001 +A 1:2009

非医用Non Medical Mask

Technical Specification: EN149:2001 +A 1:2009

生产批号/Lot No.

主要成分:无纺布70%,熔喷布30%

Main components: 70%Non-woven fabric,

30%melt-blown fabric 生产日期/Manufacturing day 有效期/shelf life:3 years

T: 86-750-3082005 F: 86-750-3082039

Made in China













CERTYFIKAT BADANIA TYPU UE (MODUŁ B) EU TYPE-EXAMINATION CERTIFICATE (MODULE B)

Nr. CW/PPER/16/11/2020

ZAŚWIADCZA SIĘ,

že Polski Rejestr Statków S.A. (PRS) przeprowadził procedurę badania typu wymienionego niżej wyrobu i stwierdził jego zgodność z wymaganiami określonymi w załączniku V do Rozporządzenia Parlamentu Europejskiego i Rady (UE) 2016/425 (PPE) w sprawie środków ochrony indywidualnej oraz uchylenia dyrektywy Rady 89/686/EWG, ze zmianami.

THIS IS TO CERTIFY

that Polski Rejestr Statków S.A. (PRS) did undertake the EU type-examination procedure for the product identified below which was found to be in compliance with the requirements of Annex V to the Regulation (EU) 2016/425 of the European Parliament and of the Council of 9 March 2016 on personal protective equipment and repealing Council Directive 89/686/EEC, as amended.

Wnioskodawca Applicant

Careable Biotechnology Co., Ltd. Building O, no. 3, Hongxing road,

Jiangmen City, China.

Producent Manufacturer Careable Biotechnology Co., Ltd. Building O, no. 3, Hongxing road,

Jiangmen City, China.

Typ wyrobu Product type

Sprzęt ochrony układu oddechowego. Półmaski filtrujące do ochrony przed cząstkami.

Respiratory protective devices. Filtering half masks to protect against particles.

Opis wyrobu Product description

Półmaska filtrująca, model: CARE 0750 (klasa FFP3 NR). Filtering half mask, Model: CARE 0750 (class FFP3 NR).

Specified standards

PN-EN 149+A1:2010 EN 149:2001+A1:2009

Niniejszy certyfikat pozostaje ważny do czasu unieważnienia przy zachowaniu warunków uznania (patrz str. 2).
This certificate remains valid uniess cancelled or revoked, provided the approval conditions (see page 2) are complied with This certificate remains valid unless cancelled or revoked, prov

Data ważności Expiry date

2025-11-04

Dyrektor Pionu Certyfikacji Certification Division Director

Michał Chudziński

Gdańsk, 2020-11-05

Nr jednostki notyfikowanej No. of notified bady

1463

Polski Rejestr Statków S.A. al. Gen. Józefa Hallera 126 80-416 Gdańsk, Poland

tel. (+48) (58) 346 17 00 fax (+48) (58) 341 77 69 e-mail: dc@prs.pl www: http://www.prs.pl/

Form. 8/PCW-01/PPER 2020-03-26

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CW/PPER/16/11/2020

Wykaz dokumentacji List of documents

- 1. Instrukcja użytkowania zatwierdzona przez PRS S.A. dnia 2020-10-27.
- Ocena ryzyka zatwierdzona przez PRS S.A. dnia 2020-10-27.
- 3. Dokumentacja techniczna "Półmaski filtrującej, model: CARE 0750" zatwierdzony przez PRS S.A. dnia 2020-10-27.
- 4. Raport z badań nr JKF20025818 wydany przez Zhejiang Academy of Science and Technology for Inspection & Quarantine (Technology Center of Hangzhou Customs District/Zhejiang Lead Product Technical Co., Ltd. z akredytacją CNAS L0354 z dnia 2020-11-04.
- 5. Sprawozdanie z przeglądu PRS S.A. nr CW/MoK/PPER/229/2020 z dnia 2020-11-04.
- 1. Instuction of use approved by PRS S.A. on 2020-10-27.
- 2. Risk analysis approved by PRS S.A. on 2020-10-27.
- 3. Technical documentation "Filtering half mask, Model: CARE 0750" approved by PRS S.A. on 2020-10-27
- 4. Test report No. JKF20025818 issued by Zhejiang Academy of Science and Technology for Inspection & Quarantine (Technology Center of Hangzhou Customs District/Zhejiang Lead Product Technical Co., Ltd. with CNAS accreditation no. L0354 dated on 2020-11-04.
- 5. PRS S.A. Survey Report No. CW/MoK/PPER/226/2020 dated on 2020-11-04.

(inne niż podane na stronie 1) Places of production (different than given on page 1)

Approval limitations

- półmaska filtrująca z regulowanym klipsem na nos,
- półmaska filtrująca wykonana z 4 warstwowej włókniny z filtrem z tkaniny,
- półmaska filtrująca wyposażona w zauszniki,
- wymiary: 160 mm ± 5 mm x 120 mm ± 5 mm,
- docelowa grupa użytkowa: dorośli dla obu płci.
- kolor: maska biała, zauszniki białe.
- Półmaska filtrująca przeznaczona do jednorazowego użytku.
- 3. Dokumentacja techniczna zatwierdzona w języku angielskim.
- Produkt ten nie może być stosowany jako maska przeciwgazowa w środowisku toksycznym.
- Półmaska filtrująca nie jest przeznaczona do użytkowania medycznego i chirurgicznego.

1. Specifications:

- protective mask with adjustable nose clip,
- protective mask made with 4 layers non-woven fabric with melt-blown fabric filter,
- protective mask with ear loops,
- size: 160 mm ± 5 mm x 120 mm ± 5 mm.
- target group: unisex,
- color: mask white, ear loops white, nose clip silver.
- 2. Filtering half mask shall not be used for more than one shift.
- 3. Technical documentation approved in English.
- 4. This product can not be used as a gas mask in a toxic environment.
- 5. Filtering half mask can not be used for medical and surgical purposes.

Warunki uznania

- Warunki uznania
 Approval conditions

 1 Niniejszy certyfikat straci ważność po wprowadzeniu zmlan lub modyfikacji w wyrobie bez uprzedniego uzgodnienia z PRS.
 This certificate becomes invalid after changes or modifications to the product without prior agreement with PRS.

 2 Znak zgodności może być umieszczony na uznanym wyrobie oraz może być wystawiona deklaracja zgodności tylko pod warunkiem, że łącznie z badaniem typu UE zostanie przeprowadzona ocena zgodności produkcji pod nadzorem jednostki notyfikowanej, według załącznika VII lub VIII

badanies type a cooline processes against production for industrial periodic interpretation of conformity issued provided the production is assessed under surveillance of a natified body according to Annex VII or VIII of the a/m Regulation.

Form. 8/PCW-01/PPER 2020-03-26







TEST REPORT



Report No.: JKF20025818

Applicant: Careable Biotechnology Co., Ltd

Zhejiang Academy of Science and Technology for Inspection and Quarantine
Add: No. 398, Jianshe 3 Road, Sacostan Discret Hangzhou, Zhejiang, China
Tel: +86 0571 8352 7 27585/197 (China) Cosite: www.zaiq.org.cn







Paget Bpages

Report No.: JKF 20025818 Report date: 2020-11-04

500	Sample Names	Filtering half mask				
Sample Information	Size:	Care0750				
шкинаки	Brand:	Fuxi Care				
	Applicants	Careable Biot	echnology Co., Ltd			
Customer	Address	Building O, no.3, hongxing road, jiangmen city (002)				
Information	Manufacturer:	Careable Biotechnology Co., Ltd				
	Manufacturer addresss	Building O, no.3, hongxing road, jiangmen city (002)				
The informa	tion are confirmed by testi	ng organizatio	6	00-		
	Date of sample received:	2020-10-29	Testing period:	2020-10-29 to 2020-11-04		
	Quantity:	110 Pieces				
Test	Sample description:	White mask				
Information	Basis of judgment:	EN 149:2001+A1:2009 FFP3 NR Respiratory protective devices—Filtering half masks to protect against particles —Requirements, testing, marking				
Test Conclusion	The items tested meet the	requirements of	EN 149:2001+A1:2	2009 FFP3 NR		
Test Result	Please refer to next pages.					
Remark	ý					

Edit:

12+++2

Ye yiwen

Sign:

Zhao dong

*** End of this page***







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Test Results:

Clause 7.5 Material

(EN 149:2001+A1:2009 Clause 8.2 & 8.3.1 & 8.3.2)

Requirement	Results	Rating
Materials used shall be suitable to withstand handling and wear over the period for which the particle filtering half mask is designed to be used. After undergoing the conditioning described in 8.3.1 none of the particle filtering half masks shall have suffered mechanical failure of the facepiece or straps.		Pass
When conditioned in accordance with 8.3.1 and 8.3.2 the particle filtering half mask shall not collapse. Any material from the filter media released by the air flow through the filter shall not constitute a hazard or unisance for the wearer.	Comply	Pass

Clause 7.6 Cleaning and disinfecting

(EN 149:2001+A1:2009 Clause 8.4 & 8.5 & 8.11)

Requirement	Results	Rating
If the particle filtering half mask is designed to be re-usable, the materials used shall withstand the cleaning and disinfecting agents and procedures to be specified by the manufacturer. With reference to 7.9.2, after cleaning and disinfecting the re-usable particle filtering half mask shall satisfy the penetration requirement of the relevant class.	Not applicable (Not designed to be re-usable)	N/A

Clause 7.7 Practical performance

(EN 149:2001+A1:2009 Clause 8.4)

Requirement	Results	Rating
The particle filtering half mask shall undergo practical performance tests under realistic conditions. These general tests serve the purpose of checking the equipment for imperfections that cannot be determined by the tests described elsewhere in this standard.	No imperfections	Pass

Clause 7.8 Finish of parts

CEN 149:2001+A1:2009 Clause 8.2

Requirement	Results	Rating
Parts of the device likely to come into contact with the wearer shall have no sharp	No sharp edges or	Pass
edges or burrs.	burrs	rass







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Clause 7.9.1 Total inward leakage

(EN 149:2001+A1:2009 Clause 8.5)

Requirement	Results	Rating
For particle filtering half masks fitted in accordance with the manufacturer's information, at least 46 out of the 50 individual exercise results (i.e. 10 subjects x 5 exercises) for total inward leakage shall be not greater than: 25% for FFP1, 11% for FFP2, 5% for FFP3 and, in addition, at least 8 out of the 10 individual wearer arithmetic means for the total inward leakage shall be not greater than: 22% for FFP1, 8% for FFP2, 2% for FFP3	49 out of the 50 individual exercise ≤5% 8 out of the 10 individual wearer arithmetic means ≤2%	Pass

Table 7.9.1-A Inward leakage test data

Subject	Sample No.	Condition	Walk (%)	Head side/side (%)	Head up/down (%)	Talk (%)	Walk. (%)	Mean (%)
CQQ	1		1.707	1.795	1.658	3.014	1.783	1.991
WIJ	2	1. 1	1.699	1.708	1.831	2.816	1.866	1.984
WG	3	As received	1.692	1,831	1.744	2.819	1.721	1.961
ZJH	4		1.803	1.842	1.792	2,657	1.800	1.979
TLB	5		1.691	1,688	1.735	2,480	1.772	1.873
ZMY	6		2.483	2.483	2,533	5.487	3.076	3.212
LJF	7	+	1.948	1.966	1.976	3.530	2.071	2.298
HML	8	Temperature	1.609	1.718	1.831	2.609	1.931	1.940
RK	9	conditioned	1,700	1.789	1.831	2.554	1.816	1.938
ZD	10		1,810	1.726	1.677	2.681	1.839	1.947

Table 7.9. I-B Facial dimensions

Subject	Face Length (mm)	Face Width (mm)	Face Depth (mm)	Mouth Width (mm)
CQQ	136	167	125	65
WLJ	132	159	110	60
WG	120	152	109	57
ZJH	122	150	104	50
TLB	125	152	111	57
ZMY	137	150	120	60
LJF	125	135	90	55
HML	124	130	115	55
RK.	112	161	146	50
ZD	116	160	115	- 55











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Clause 7.9.2 Penetration of filter material

(EN 149:2001+A1:2009 Clause 8.11 & EN 13274-7:2019)

	Requirement		Results	Rating
enetration of the for				
Classification	Sodium chloride test 95 L/min	Paraffin oil test 95 L/min	Detail refer to	Pass
FFP1	≤20%	≤20%	Table 7.9.2	
FFP2	≤6%	≤6%		
FFP3	€1%	≤1%		

Table 7.9.2 Penetration of filter material

Aerosol	Condition	Sample No.	Penetration (%)
		n n	0.027
	As received	12	0.050
		13	0.063
Sodium chloride test	Simulated wearing	14	0:077
	treatment	15	0.081
	псавнен	16	0.063 0.077 0.081 0.056 0.089 0.147 0.173 0.040 0.195 0.083 0.110
	Mechanical strength+ Temperature conditioned	17	0.089
		18	0.147
	Temperature conditioned	19	0.173
	As received	20	0.040
		21	0.195
		22	0.083
	Cincileted marries	23	0.110
Paraffin oil test	Simulated wearing	24	0.088
1-24-5	treatment	25	0.012
	Mr. Anna Salamanaka	26	0.741
	Mechanical strength+	27	0.837
	Temperature conditioned	28	0.469

Clause 7.10 Compatibility with skin

(EN 149:2001+A1:2009 Clause 8.4 & 8.5)

Requirement	Results	Rating
Materials that may come into contact with the wearer's skin shall not be known to be likely to cause irritation or any other adverse effect to health.	No irritation or any other adverse effect to health	Pass







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Clause 7.11 Flammability

(EN 149:2001+A1:2009 Clause 8.6)

Requirement	Results	Rating
When tested, the particle filtering half mask shall not burn or not to continue to burn for more than 5s after removal from the flame.	Detail refer to Table 7.11	Pass

Table 7.11 Flammability

Condition	Sample No.	Result
According .	29	Not barn
As received.	30	Not burn
The second second second	31	Not burn
Temperature conditioned	32	Not burn

Clause 7,12 Carbon dioxide content of the inhalation air

(EN 149:2001+A1:2009 Clause 8.7)

Requirement	Results	Rating
The carbon dioxide content of the inhalation air (dead space) shall not exceed an average of 1.0 % (by volume).	Detail refer to Table 7.12	Pass

Table 7.12 Carbon dioxide content of the inhalation air

Condition	Sample No.		Result (%)
	33	0.54	i i i i i i i i i i i i i i i i i i i
As received	34	0.46	Mean value: 0.50
	35	0.51	

Clause 7.13 Head harness

(EN 149:2001+A1:2009 Clause 8.4 & 8.5)

Requirement	Results	Rating
The head harness shall be designed so that the particle filtering half mask can be donned and removed easily.		
The head hamess shall be adjustable or self-adjusting and shall be sufficiently robust	Comply	Pass
to hold the particle filtering half mask firmly in position and be capable of		
maintaining total inward leakage requirements for the device.		

Clause 7.14 Field of vision

(EN 149:2001+A1:2009 Clause 8.4)

Requirement	Results	Rating
The field of vision is acceptable if determined so in practical performance tests.	Comply	Pass







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Clause 7.15 Exhalation valve

(EN 149:2001+A1:2009 Clause 8.2 & 8.9.1 & 8.3.4 & 8.8)

Requirement	Results	Rating
A particle filtering half mask may have one or more exhalation valve(s), which shall function correctly in all orientations. If an exhalation valve is provided it shall be protected against or be resistant to dirt and mechanical damage and may be shrouded or may include any other device that may be necessary for the particle filtering half mask to comply with 7.9. Exhalation valve(s), if fitted, shall continue to operate correctly after a continuous exhalation flow of 300 L/min over a period of 30 s. When the exhalation valve housing is attached to the faceblank, it shall withstand axially a tensile force of 10 N applied for 10 s.	Not applicable (No exhalation valve)	N/A

Clause 7.16 Breathing resistance

(EN 149:2001+A1:2009 Clause 8.9

	Requir	rement		Results	Rating
ments of the filte		filtering half ma	sk shall meet the		
	Maximum	permitted resista	ance (mbar)	The second	Pass
Classification	Inhal	ation	Exhalation	Detail refer to	
	301/min	95L/min	160L/min	Table 7.16	
FFP1	0,6	2.1	3.0		
FFP2	0.7	2.4	3.0		
FFP3	1.0	3.0	3.0		

Table 7-16 Broathing resistance (mbar

Test item	Condition	Sample No.	A	В	C	D	E
		36	0.53	0.53	0.54	0.54	0.54
	As received	37	0.54	0.54	0.55	0.54	0.55
Inhalation (30 L/min) Simulated wearing treatment Temperature conditioned		38	0.58	0.58	0.58	0.58	0.59
		39	0.52	0.52	0.52	0.52	0.52
		40	0.52	0.52	0.52	0.52	0.52
	treatment	41	0.52	0.52	0.52	0.52	0,52
	Menal Pateria	42	0.52	0.51	0.51	0.52	0.51
	1737 (800 00000)	43	0.51	0.51	0.52	0.51	0.51
	conditioned	44	0.51	0.52	0.52	0.51	0.52







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Test item	Condition	Sample No.	A	B	C	D	E
		36	2.19	2.18	2.18	2.19	2.20
	As received	37	2,17	2.19	2.18	2.18	2.17
		38	2,24	2.24	2.26	2.24	2.25
Auto Contract	When the State of the second	39	2,23	2.24	2.23	2.23	2.24
Inhalation	Simulated wearing	40	2.02	2.03	2.03	2.02	2.02
(95 L/min)	treatment	41	2.13	2.13	2:13	2.11	2.12
	Thomas and the	42	2,03	2.05	2.05	2.04	2.04
	Temperature	43	2,00	2.02	2.02	2.01	2.02
	conditioned	44	2.05	2.03	2.03	2.02	2.04
	As received	36	2.92	2.91	2.89	2,89	2.90
		37	2.97	2.96	2.95	2,96	2,97
		38	2.97	2.94	2.98	2.97	2.99
Exhalation	CANADA CONTRACTOR OF THE PARTY	39	2.95	2.95	2.92	2,93	2.94
		40	2.88	2.86	2,86	2,88	2.86
(160 L/min)	treatment	41	2.97	2.98	2.98	2.96	2.97
	Tomachistone	42	2.84	2.83	2.84	2.81	2.82
	Temperature	43	2.67	2.69	2,66	2.68	2.67
	conditioned	44	2.82	2.84	2.83	2.82	2,84

A: facing directly ahead; B: facing vertically upwards; C: facing vertically downwards; D: lying on the left side; E: lying on the right side

Clause 7.17 Clogging

(EN 149:2001+A1:2009 Clause 8.9 & 8.10)

Requirement	Results	Rating
7.17.21 Valved particle filtering half masks After clogging the inhalation resistances shall not exceed FFP1:4mbar, FFP2:5mbar, FFP3:7mbar at 95 L/min continuous flow; The exhalation resistance shall not exceed 3mbar at 160 L/min continuous flow. 7.17.2.2 Valveless particle filtering half masks After clogging the inhalation and exhalation resistances shall not exceed FFP1;3mbar, FFP2;4mbar, FFP3:5mbar at 95 L/min continuous flow. 7.17.3 Penetration of filter material: All types (valved and valveless) of particle filtering half masks claimed to meet the clogging requirement shall also meet the requirements given in 7.9.2, for the Penetration test according to EN 13274-7, after the clogging treatment.	Not applicable (Single shift use only)	NA

Clause 7.18 Demountable parts

(EN 149;2001+A12009 Clause 8.2)

Requirement	Results	Rating
All demountable parts (if fitted) shall be readily connected and secured, where possible by hand.	Comply	Pass







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Sample photo





*** End of Report***





STATEMENT

- Our organization guarantees impartiality, independence and honesty of inspection, and is responsible for the content of report, except for the information provided by the client. The client shall not use the test results for improper publicity without authorization.
- Our organization shall not be responsible for the authenticity of the information provided by the client, nor shall bear the risks arising in the process of sample delivery. Test result is only responsible for the sample.
- This report is invalid without the dedicated seal for inspection and testing report and the paging seal.
- 4. This report is invalid without the signature of the approver (authorized signatory).
- 5. Test report is invalid if altered.
- The duplicate report without the "dedicated seal for inspection and testing" of the institution is invalid.
- 7. Each page of the report is an integral part of the report. Our organization shall not be responsible for any misunderstanding or consequences arising from the improper use of the test report by the user.
- 8. Without the CMA seal, the report is invalid for social certification.

Test institute: Zhejiang Academy of Science and Technology for Inspection and Quarantine

Add: No. 398, Jianshe 3 Road, Xiaoshan District, Hangzhou, Zhejiang, China

Tel: +86 0571 8352 7187/185/193

Website: www.zaiq.org.cn











CERTYFIKAT BADANIA TYPU UE (MODUŁ B) EU TYPE-EXAMINATION CERTIFICATE (MODULE B)

No. CW/PPER/16/11/2020

ZAŚWIADCZA SIĘ,

Ze Polski Rejestr Statków S.A. (PRS) przeprowadził przcedunę badania typu wymienionego niżej wyrobu i stwierdził jego zgodność z wymaganiami określonymi w załączniku V do Rozporządzenia Parlamentu Europejskiego i Rady (UE) 2016/425 (PPE) w sprawie środków ochrony indywidualnej oraz uchylenia dyrektywy Rady 89/686/EWG, ze zmianami.

that Polski Rejestr Statków S.A. (PRS) did undertake the EU type-examination pracedure for the product identified below which was found to be in compliance with the requirements of Annex V to the Regulation (EU) 2016/425 of the European Parliament and of the Council of 9 March 2016 an personal protective equipment and repealing Council Directive 89/686/EEC, as amended.

Wnioskodawca Applicant

Careable Biotechnology Co., Ltd. Building O, no. 3, Hongxing road,

Jiangmen City, China.

Producent Manufactures

Careable Biotechnology Co., Ltd. Building O, no. 3, Hongxing road,

Jiangmen City, China.

Product type

Sprzęt ochrony układu oddechowego. Półmaski filtrujące do ochrony przed cząstkami.

Respiratory protective devices. Filtering half masks to protect against particles.

Opis wyrobu Product description Pólmaska filtrująca, model: CARE 0750 (klasa FFP3 NR).

Filtering half mask, Model: CARE 0750 (class FFP3 NR).

Zastosowane normy Specified standards

PN-EN 149+A1:2010 EN 149:2001+A1:2009

Niniejszy certyfikat pozostaje ważny do czasu unieważnienia przy zachowaniu warunków uznania (patrz str. 2). This certificade cemains volid uniess conceiled or revoked, provided the appraval conditions (see page 2) are com

Data ważności

Expiry date

2025-11-04

Dyrektor Pionu Certyfikacji Certification Division Director

Michał Chudziński

Gdańsk, 2020-11-05



Nr jednostki natyfikowanej No. of notified body

1463

Polski Rejest/ Statków S. al. Gen. Józefa Hallera 126 80-416 Gdańsk, Poland

tel. (+48) (58) 346 17 00 fax (+48) (58) 341 77 69 e-mail: dc@prs.pl www.http://www.prs.pl/

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CW/PPER/16/11/2020

Wykaz dokumentacji List of documents

- 1. Instrukcja użytkowania zatwierdzona przez PRS S.A. dnia 2020-10-27.
- 2. Ocena ryzyka zatwierdzona przez PRS S.A. dnia 2020-10-27.
- 3. Dokumentacja techniczna "Półmaski filtrującej, model: CARE 0750" zatwierdzony przez PRS S.A. dnla 2020-10-27.
- Raport z badań nr JKF20025818 wydany przez Zhejiang Academy of Science and Technology for Inspection & Quarantine (Technology Center of Hangzhou Customs District/Zhejiang Lead Product Technical Co., Ltd. z akredytacją CNAS L0354 z dnia 2020-11-04.
- 5. Sprawozdanie z przeglądu PRS S.A. nr CW/MoK/PPER/229/2020 z dnia 2020-11-04
- Instuction of use approved by PRS S.A. on 2020-10-27.
- 2. Risk analysis approved by PRS S.A. on 2020-10-27.
- 3. Technical documentation "Filtering half mask, Model: CARE 0750" approved by PRS S.A. on 2020-10-27.
- 4. Test report No. IKF20025818 issued by Zhejiang Academy of Science and Technology for Inspection & Quarantine (Technology Center of Hangzhou Customs District/Zhejiang Lead Product Technical Co., Ltd. with CNAS accreditation no. L0354 dated on 2020-11-04.
- 5. PRS S.A. Survey Report No. CW/MoK/PPER/226/2020 dated on 2020-11-04.

Miejsca produkcji (inne niž podane na stronie 1) Places of production (different than given an page 1)

Ograniczenia uznania

1. Dane techniczne:

- półmaska filtrująca z regulowanym klipsem na nos,
- półmaska filtrująca wykonana z 4 warstwowej włókniny z filtrem z tkaniny,
- półmaska filtrująca wyposażona w zauszniki.
- wymiary: 160 mm ± 5 mm x 120 mm ± 5 mm,
- docelowa grupa użytkowa: dorośli dla obu płci,
- kolor: maska biała, zauszniki białe.
- 2. Półmaska filtrująca przeznaczona do jednorazowego użytku.
- 3. Dokumentacja techniczna zatwierdzona w języku angielskim.
- 4. Produkt ten nie może być stosowany jako maska przeciwgazowa w środowisku toksycznym.
- 5. Półmaska filtrująca nie jest przeznaczona do użytkowania medycznego i chirurgicznego.

1. Specifications:

- protective mask with adjustable nose clip,
- protective mask made with 4 layers non-woven fabric with melt-blown fabric filter,
- protective mask with ear loops,
- size: 160 mm ± 5 mm x 120 mm ± 5 mm,
- target group: unisex.
- color: mask white, ear loops white, nose clip silver.
- 2. Filtering half mask shall not be used for more than one shift.
- 3. Technical documentation approved in English.
- 4. This product can not be used as a gas mask in a toxic environment.
- 5. Filtering half mask can not be used for medical and surgical purposes.

Warunki uznania

- Approval conditions

 1 Miniejszy certyfikat straci ważność po wprowadzeniu zmian lub modyfikacji w wyrobie boz uprzedniego uzgodnienia z PRS.

 This certificate becomes invalid ofter changes or modifications to the product without prior agreement with PRS.

 2 Znak zgodności może być umieszczony na uznanym wyrobie oraz może być wystawiana deklaracja zgodności tylko pod warunidem, że łącznie z badaniem typu UE zostanie przeprowadzona ocena zgodności produkcji pod nadzorem jednostki notyfikowanej, według załącznika VII lub VIII wymienionego wyżej rozporządzenia.

wyministration by wyter responsections.

The Mark of Conformity may only be affixed to the above type approved product and a manufacturer's Deplaration of Conformity issued provided the production is assessed under surveillance of a notified bady according to Annex VII or VIII of the a/m Regulation.

Form 8/PCW-01/PPER 2020-03-26