

REF HS0501A

# FFP2 Maske / FFP2 Mask

Persönliche Schutzmaske

**Personal Protective Mask** 



CE2797

EN 149: 2001+A1: 2009

CARTON DIMENSION (FFP2 - White - 20er)

HYGISUN®

1 PC/OPP

20 PCS/box

1000 PCS/carton

20000 PCS/pallet

EAN: 4260676530003

**(E**<sub>2797</sub>



# **FFP2-MASKE**

PERSÖNLICHE SCHUTZMASKE EN149:2001+A1:2009

HYCISUN

REF HS0501A

Faltbare Partikel -Atemschutzmaske

Hohe Filtrationseffizienz Geringer Atemwiderstand Bequem zu tragen

20Stück @ Einmalgebrauch

C€ 2797

C€ 2797

20stiick & Einmalgebrauch

**C**€ 2797

Baquem zu frogei

### **Bedienungsanleitung Fitting instructions**



Nehmen Sie die Moske an den Ohrschlaufen in die Hand und drücken Sie diese mit dem Bägel auf den Nasenrücken gegen ihr Gesicht, während Sie die Ohrschlaufen finter ihren Ohnen positionieren.

Take the mask by the ear loops in your hand and press it against your face with the strap on the bridge of your nase while you position the ear



Formen Sie den Bügel mit beiden Händen in die Form übrer Nase. hope the nose dip into the shope of your nose with both hands



Atemschutzmaske und atmen Sie kräftig aus. Wenn Luft um ihre Nase

Test the correct fit. Put both hunds over the respirator and exhale errefully. When air flows out around your nose, gress the nose dip

### HINWEIS ZUR VERWENDUNG: / NOTICE FOR USE:

Bitte verwenden Sie dieses Produkt nicht in der Nithe einer Feuerquelle. Please do not use this product near fire sources.

Da es sich bei diesem Produkt um eine Einwegmaske handelt, kann es nicht durch Waschen wiederverwendet werden.

As this product is a disposable mask, it cannot be reused through washing.

Von hohen Temperaturen und Lufffeuchtigkeit fernhalten und an einem sauberen Ort

Keep it away from high temperature and humidity and keep it in dean place. Persönliche Schutzmaske, Nicht medizinisch.

Verwenden Sie einzeln verpackte Produkte, sobald diese ausgepackt sind. Use individually packaged products as soon as they are unpacked.

Large & growy trans heigh transcribure und humidly and havin it in them piece. For soutche Schaltmerier, Nicht meditanisch.

# FFP2-MASK

PERSONAL PROTECTIVE MASK EN149:2001+A1:2009



REF HS0501A

# **Foldable Particulate** Respirator

High filtration efficiency Low respitationy resistance More comfortable to wear

20PCS ® Single Use

C € 2797

C€ 2797

② Single Use

**C**€ 2797

### FFP2 Maske EN149:2001+A1:2009



MPORTANT: The respiratory protection mask FFP2 is designed to

### **ANWENDUNG: / APPLICATION:**

Die Macke wird in der Schutzendustrie bei Steubentwicklung, wahrend des Bioss zur Steubernklung, beim Metaliguss, Steinobbox, in der Bektrenik, Pformazie, der physikolischen Verurbeitung unb beim Schleiden verwendet und belet einen guten Schutz geges Stendisfatine, Durst und PMZ-S. Kenn waksom vor Pollensliergien, Verusdertragging zw. schützen.

It is used in the industry for class generation during construction, dust prevention, mutal custing, stone nating, electronics, phonocaratical, physical processing and grinding. It also offers good protection against southstarms, have and PAZ-5. Can affectively protect polion oflergy, views transmission, etc.

### VERFALLSDATUM: / EXPIRATION DATE:

 $\label{logical_logical} Logerteuthigkeit \le 80\%, Haltburkeit. 2. Notre in trackenen Innenträumen. The sturces temperature is: <math>20-23\%$  C the sturage is moderate  $\le 80\%$ . The

the storage temperature is  $-20^{\circ}-38^{\circ}$  (, the storage is moderate  $\leq 80\%$ , This validity period is 2 year in the dry indoor environment.



Hunan Dreaming Cloud E-Commerce CO., Ltd Block 1, Smart Tech Park, 57# Hunapsing Avenue, Changsha Economic and Technological Development Zone, Changsha, Hornan, China



Sunbeam International GmbH Schumanstr. 12, 52146 Würselen, Germany



Sunbeam International GmbH Stannastr, 12, 521 fé Wassler, Germany





# HYGININV® FFP2 Maske Michi medizinisch Entag:2001+A1:2009

# FFP2-MASKE Hichr moulzintsch EN149:2001+A1:2009 Faltbare Partikel -Nemschubmaske

HYCINIM®

SEA CTURES







# Persönliche Schutzmaske Personal Protective Mask

1Stück/ Piece

1Stück/ Piece



QUALIFIED CERTIFICATE

产品名称	FFP2个人防护口罩(不作为医疗防护使用)
Product Name	FFP2 personal protective mask (For personal protective only)
执行标准	EN149: 2001 + A1: 2009
Standard	EN149: 2001 + A1: 2009
产品规格	16×10.5厘米
Product Size	16×10.5CM
材质	30%熔喷布+70%无纺布
Material	30% MELT-BY CONTRABRICE AUTOMA NOVEN FABRIC
质检员	QUALITY PASSED B A A B B
Checker	The same of the sa
生产日期	and connection.
Date	QC:01
生产批号	PD: SEE ON PACKAGE
Lot code	LOT: SEE ON PACKAGE
使用周期	一次性
Usege count	Single use
有效期	2年
Period of validity	2 years
生产企业	湖南云想生活电子商务有限公司
Manufacturer	Hunan Dreaming Cloud E-Commerce CO., Ltd
生产地址	长沙经济技术开发区黄兴大道南段
	57号星为创芯园1栋501号
Address	Block 1, Smart Tech Park, S7# Huangking Avenue Changsha
	Economic and Technological Development Zone, Changoliu
	Hunan, China

#### 储存条件:本品应储存温度-20~38°C,储存湿度≤80% 避光干燥的室内环境下,通风良好,无腐蚀性气体的清洁 环境内,如贮存不当导致发霉变质禁用。

STONAGE CONDITIONS AND METHODS: The storage temperature is "20-39"C, and the relative humidity is not incredition 80% velicities that and dean environment without companyas, please of on no use if the product gets mildreved or deteriorished due to improper storage.

### 本产品为个人防护用品,不作为医疗防护使用

THIS PRODUCT IS LINDER PERSONAL PROTECTIVE EQUIPMENT DIRECTIVE (PRE) FOR PERSONAL PROTECTIVE ONLY

MADE IN CHINA

MADE IN CH

PROPRIET FAMILY FROM THE PARKET WAS THE PROPRIET FOR THE PROPRIET OF THE PROPRIET FROM THE



#### ANLEITUNG

Norm:

Dieses Produkt entspricht der Norm EN149:2001 + A1:2009 für Atemschutzgeräte – Halbmaske zur Filterung zum Schutz vor Partiskeln, Diese Filtermasken sind gemäß der Verordnung der Europäischen Kommission (EU) 2016/425 über PSA als Persönliche Schutzausrüstung in der Kategorie III eingestuft und entsprechend gekennzeichnet

#### Bestimmungsgemäße Verwendung:

Die Staubmaske ist als Kategorie FFP 2 eingestuft. Sie schützt vor Partikeln, Nebel, Rauch und Aerosolen auf Ölbasis. Die Verpackung schützt die Maske vor der Verwendung, Schützt wirksam vor Pollen. Die Maske kann nur zum persönlichen Schutz verwendet werden, nicht für medizinische Zwecke. Maske nicht bei der Brandbekämpfung und in explosionsgesfährdeten Bereichen nutzen.

#### Dichtsitztest

- 1. Bedecken Sie die Maske vorsichtig mit beiden Händen ohne den Dichtsitz zu verändern.
- 2.stark Ausatmen:
- 3.Bei einer Leckage im Nasenbereich, den Nasenbügel neu anpassen. Dichtsitzprüfung wiederholen.
- 4.Bei einer Leckage am Maskenrand, den Sitz der Bänder überprüfen und anpassen. Dichtsitzprüfung

Wenn Sie KEINEN richtigen Dichtsitz erreichen können, betreten Sie NICHT den Gefahrenbereich. Informieren Sie ihren Vorgesetzten.

#### Warnungen und Einschränkungen:

- Vergewissern Sie sich immer, dass das Produkt:
- Geeignet ist für die Anwendung;
- Korrekt angelegt ist:
- Während des gesamten Aufenthalts im Gefahrenbereich getragen wird;
- Ersetzt wird, wenn notwendig.
- Richtige Auswahl, Schulung, Gebrauch und gegebenenfalls Reinigung sind die Voraussetzungen dafür,
- dass das Produkt den Anwender vor bestimmten luftgetragenen Gefahrstoffen schützt.
  Die Nichtbefolgung aller Anweisungen zur Anwendung der Maske und/oder die Fehlbenutzung
- während des Aufenthaltes im Gefahrenbereich kann die Gesundheit des Anwenders beeinträchtigen und zu schweren Erkrankungen oder Dauerschäden führen.
- Beachten Sie bei der Auswahl und richtigen Anwendung nationale Bestimmungen und alle mitgeliefer ten Informationen.
   Vor Gebrauch muss der Anwender, in Übereinstimmung mit den nationalen Regeln, in der funktions.
- Vor Gebrauch muss der Anwender, in Übereinstimmung mit den nationalen Regeln, in der funktions gerechten Handhabung geschult sein.
- · Dieses Produkt schützt nicht vor Gasen und Dämpfen.
- Verwenden Sie die Maske nicht in Umgebungen mit weniger als 19.5% Sauerstoff.
- Verwenden Sie die Masken nicht in Umgebungen mit unbekannten Gefahrstoffen oder Konzentrationen, die die zulässigen Höchstwerte übersteigen.
- Verwenden Sie die Maske nicht, wenn Gesichtshaare im Bereich des Dichtrandes einen korrekten
- Dichtsitz der Maske verhindern.
  Verlassen Sie sofort den belasteten Bereich, wenn:
- a) Das Atmen schwer fällt.
- b) Schwindel oder andere Beschwerden auftreten.
- c) Die Maske beschädigt wird.
- d) Geruch oder Geschmack des Gefahrstoffs oder eine Reizung auftritt.
- Entsorgen und ersetzen Sie die Maske, wenn sie beschädigt ist, der Atemwiederstand stark erhöht ist oder am Ende einer Schicht.
- Die Maske darf niemals verändert oder repariert werden.
- Die Maske ist zum einmaligen Gebrauch vorgesehen und ist danach entsprechend der nationalen Vorgaben zu entsorgen.

#### Transport und Lagerung:

Die Partikelmasken haben eine Lagerdauer von 2 lahren. Das Ende der Lagerdauer ist auf der Verpackung angegeben. Vergewissens Fis sich vor Gebrauch immer, dass das Produkt noch innerhalb der Lagerdauer liegt. Das Produkt sollte saubet, trocken und im Temperaturbereich von -20°C bis +38°C bei einer maximalen et Luffteuchtügket von 80% gelagert werden. Für Lagerung

und Transport die Originalverpackung verwenden. Nicht direkter Sonnenstrahlung aussetzen.







# 取得国外标准认证或注册的非医用口罩生产企业清单 Name List of Non-Medical Use Face Masks Companies with

序 号 No.	生产企业 Company	统一社会信用代码 Uniform Social Credit Code	国外注册认证情况 Status of Certification / Authorization in Other Countires
977	は初れ付金料金料金料 Lunyang Kelijian Technology Co.,Ltd.	91400500MA400BX51L	German Made EUA
278	公共市庁務款主収券を限心司 Xianton Qianteng Life Sering Equipment Co.,Ltd.	RTHOSOBHMANNABOLISE	CE
379	湖南云想生活电子商务有限公司 Hunan Dreaming Cloud E-Commerce CO., Ltd	91430105MA4LAAUW8C	СЕ
380	维分得表用医疗用品有限企用 Linnytungung Meiden Modierd Supplier Co., Ltd.	91320724596800367H	Œ
501.	原来海尼斯丹基祖布联合司 Nonjing Homey Medical Apparatus and Instruments Ox.Ltd.	91530136MA30W250948	CE CE
582	II 声音速度等有限心用 Sangus Daoying Clothing Co., Ltd	911009Q1MATNIGUE28	cr



Block 1, Smart Tech Park, 57# Huangxing Avenue, Changsha Economic and Technological Development Zone,Changsha, Hunan, China

### **EU-KONFORMITÄTSERKLÄRUNG**

Diese Konformitätserklärung wurde unter der alleinigen Verantwortung des Herstellers Hunan Dreaming Cloud E-Commerce CO., Ltd.

Block 1, Smart Tech Park, 57 # Huangxing Avenue, Changsha Economic and Technological Development Zone, Changsha, Hunan, China

ausgestellt.

EG-Vertreter: Sunbeam International GmbH, Schumanstr.12, Würselen 52146 Deutschland

Hiermit wird erklärt, dass die folgende persönliche Schutzausrüstung (PSA)

Produktbeschreibung: HYGISUN Partikelfilter-Halbmaske

Produktmodell (e): HS0501A FFP2 NR ohne Ventil

den Bestimmungen der folgenden europäischen Verordnung entspricht:

### **PSA-Verordnung (Persönliche Schutzausrüstung)**

Das Modell entspricht den Bestimmungen der Verordnung (EU) 2016/425, PSA zur Verwendung durch Angehörige der Gesundheitsberufe gemäß der Empfehlung der Kommission 2020/403 und der Nationalen Norm zur Umsetzung der harmonisierten europäischen Normnummer (n):

EN 149: 2001 + A1: 2009

und ist identisch mit der PSA, die Gegenstand einer EU-Typprüfung ist (Modul B der Verordnung (EU) 2016/425), auf die auf der Zertifikatsnummer verwiesen wird:

Zertifikat Nr.: CE 750475 (Ausstellungsdatum: 09/06/2021)

herausgegeben von BSI Group Niederlande BV

John M. Keynesplein 9, 1066 EP, Amsterdam, Niederlande (Notified Body No. 2797)

und entspricht den Verfahren in Modul C2 der Verordnung (EU) 2016/425 unter der Überwachung der BSI Group The Netherlands BV (Notified Body Nr. 2797), auf die auf dem vom BSI ausgestelltem Zertifikat CE 750476 (Ausstellungsdatum: 09/06/2021) verwiesen wird.

Changsha, China, 19.06

OuYang Zhouya

(Nachname Name)

Qualitätsmanager

Hunan Dreaming Cloud E-Commerce CO., Ltd.







# **EU Type Examination Certificate**

This is to certify that: Sunbeam International GmbH

> Schumanstr. 12 Würselen 52146 Germany

Holds Certificate Number: CE 750475

In respect of:

Respiratory protective devices - Filtering half masks to protect against particles -To EN 149:2001+A1:2009 **Model: HYGISUN HS0501A.** 

on the basis that BSI carried out the relevant Type Examination procedures under the requirements with the Regulation (EU) 2016/425 of the European Parliament and Council relating to Personal Protective Equipment Regulation (PPE) Annex V (Module B) and meets the relevant health and safety requirements specified in Annex II

For and on behalf of BSI, a Notified Body for the above Regulation (Notified Body Number 2797):

Drs. Dave Hagenaars, Managing Director

First Issued: 2021-06-09 Latest Issue: 2021-06-09 Effective Date: 2021-06-09 Expiry Date: 2026-06-09

Page: 1 of 3



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# **EU Type Examination Certificate**

No. CE 750475

### **Product Specification**

**Product Type:** Filtering half masks to protect against particles.

Model: HYGISUN HS0501A.

**Product description:** The particulate respirator is designed to protect against solid and non-volatile liquid

particles.

The masks are a single size, non-sterile, non-valved product held on the face by a

pair of elasticated ear loops.

The masks are intended for single shift use as denoted by the classification symbol

NR.

**Technical specification:** EN 149:2001+A1:2009 - Respiratory Protective Devices -

Filtering half masks to protect against particles.

FFP2 NR. EN 149 classification:

First Issued: 2021-06-09 Effective Date: 2021-06-09 Latest Issue: 2021-06-09 Expiry Date: 2026-06-09

Page: 2 of 3

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# **EU Type Examination Certificate**

No. CE 750475

### **Certificate Administration Details**

Technical File reference: TCF.02.

### **Certificate Amendment Record:**

Issue date	Comments	BSI Review Number
June 2021	First issue under PPE Regulation (EU) 2016/425. Product initially Certified as a "Covid-19" mask by BSI, Certificate CE 730303 refers.	2797:2021:3339407

### **Certificate validity**

The Certificate holder is responsible for ensuring that the Notified Body is advised of changes to any aspect of the overall processes utilised in the manufacture of the product, failure to do so could invalidate the Certificate in respect of product manufactured following the introduction of such changes.

The validity of the Certificate for the products is also dependent on the maintenance of the EU Conformity to Type based on Internal Production Control plus supervised product checks at random intervals, Annex VII (Module C2) as referenced on BSI issued Certificate CE 750476.

First Issued: 2021-06-09 Effective Date: 2021-06-09
Latest Issue: 2021-06-09 Expiry Date: 2026-06-09

Page: 3 of 3

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To check its validity telephone +31203460780. An electronic certificate can be authenticated online.







# Conformity to Type based on Internal Production Control plus supervised product checks at random intervals

This is to certify that: Sunbeam International GmbH

> Schumanstr. 12 Würselen 52146 Germany

Holds Certificate Number: CE 750476

In respect of:

For the manufacture of respiratory protective devices -Filtering half masks to protect against particles - To EN 149:2001+A1:2009.

on the basis that BSI carried out the supervised production checks at random intervals under the requirements with the Regulation (EU) 2016/425 of the European Parliament and Council relating to Personal Protective Equipment Regulation (PPE) Annex VII (Module C2)

For and on behalf of BSI, a Notified Body for the above Regulation (Notified Body Number 2797):

Drs. Dave Hagenaars, Managing Director

First Issued: 2021-06-09 Latest Issue: 2021-06-09 Effective Date: 2021-06-09 Expiry Date: 2026-06-09

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# Conformity to Type based on Internal Production Control plus supervised product checks at random intervals

No. CE 750476

### Model produced by:

Hunan Dreaming Cloud E-Commerce CO., Ltd Block 1, Smart Tech Park, 57# Huangxing Avenue, Changsha Economic and Technological Development Zone, Changsha, Hunan, China

### **Product details**

The respiratory protective device covered by the scope of this Module C2 Certificate and the Technical Specification to which the product is manufactured are as follows:

**Product type:** Respiratory protective device – Filtering half masks to protect against particles.

Model: HYGISUN HS0501A.

**Technical Specification:** EN 149:2001+A1:2009 – Respiratory Protective Devices -

Filtering half masks to protect against particles.

**EN 149 classifications:** FFP2 NR.

### **Certificate Administration Details:**

### **Certificate Amendment Record:**

Issue date	Comments	<b>BSI Review No.</b>
June 2021	First issue.  Referenced product initially Certified as a "Covid-19" mask by BSI, with the associated BSI issued Module C2 Certificate CE 730304.	2797:21:3339408

### **Certificate validity**

The Certificate holder is responsible for ensuring that the Notified Body is advised of changes to any aspects of the overall quality system utilized in the manufacture of the products, failure to do so could invalidate the Certificate in respect of product manufactured after the introduction of such changes.

First Issued: 2021-06-09 Effective Date: 2021-06-09
Latest Issue: 2021-06-09 Expiry Date: 2026-06-09

Page: 2 of 2

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# Test Report 3339405.

# Sunbeam International GmbH.



### Introduction.

This report has been prepared by D. Key and relates to the activity detailed below:

Job/Registration Details		Client Details		
Job number:	3339405	Sunbeam International GmbH		
Job type:	lesting samples submitted	Schumanstr. 12 Würselen		
Start Date:	17/01/2021	52146		
Test type:	Туре	Germany		
Sample ID: 10195243				
Registration:	CE 730303			
Scheme:	Negative Pressure RPE			
Protocol:	PP123			
Scheme Manager:	Nathan Shipley			

The report has been approved for issue by T Wicksey – Senior Test Engineer

Approved For Issue	
22/5<	
	Issue Date: 22 March 2021

# Objectives.

This is an independent Type Test evaluation to BS EN 149:2001+A1:2009. This report covers the gap testing from the BSI COVID-19 filtering face piece technical specification, for COVID-19 masks for use by healthcare workers. See BSI Test Report 3220780 for the BSI COVID-19 filtering face piece technical specification test results.

# Product Scope.

Respiratory protective device- Filtering half masks to protect against particles.

# Report Summary.

The samples were received on 18 December 2020 and the testing was started on 17 January 2021.

The samples submitted complied with the requirements of the test work conducted.



# Test Samples.

Sample ID	ER Number	Description
1 to 37	10195243	Model: HYGISUN HS0501A FFP2 NR

# Description of Test Samples.

### **Sample Description**

Model: HYGISUN HS0501A FFP2 NR. Valveless vertical fold flat particle filtering half mask with elastic earloops and removable plastic earloop clip



# Test Requirements.

### BS EN 149:2001 + A1:2009

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

CLAUSE	REQUIREMENTS	ASSESSMENT			
7	Requirements	-			
7.1	General	-			
7.2	Nominal values and tolerances	-			
7.3	Visual Inspection	Pass (1)			
7.4	Packaging	N/T (1)			
7.5	Material	Pass			
7.6	Cleaning and disinfecting	N/A (2)			
7.7	Practical performance	N/T (3)			
7.8	Finish of parts	Pass			
7.9	Leakage	-			
7.9.1	Total inward leakage	Pass (3)			
7.9.2	Penetration of filter material	Pass (3)			
7.10	Compatibility with skin	Pass			
7.11	Flammability	Pass			
7.12	Carbon dioxide content of inhalation air	N/T (3)			
7.13	Head harness	Pass			
7.14	Field of vision	Pass			
7.15	Exhalation valves	N/A (4)			
7.16	Breathing resistance	Pass (3)			
7.17	Clogging	N/A (4)			
7.18	Demountable parts	N/A (4)			
9	Marking	N/T (1)			
10	Information to be supplied by the manufacturer	N/T (1)			
Appendix A	A - Test Panel Data				
Product Ph	Product Photographs				

- (1) Packaging, Marking and Information not assessed as requested by BSI Product Certification
- (2) Single use mask
- (3) See also results from BSI COVID-19 filtering face piece technical specification testing, BSI Test Report number 3220780.
- (4) Not a design feature of this product





# Glossary of Terms.

Pass: Complies. Tested by BSI engineers at BSI laboratories

Pass 1: Complies. Witnessed by BSI engineers in manufacturers laboratory.

Pass 2: Complies. Tests carried out by third party lab; results accepted by BSI.

Pass\*: Report resulted in uncertainty and states that Compliance is more probable than non-compliance.

Fail: Non-compliance. Product does not meet the requirements of this clause.

Fail\*: Report resulted in uncertainty and states that Non-compliance is more probable than compliance.

N/T: Not Tested N/A: Not Applicable AR: As Received

TC: Temperature Conditioned

SW: Simulated Wear FT: Flow Tested

MS: Mechanical strength

MMDF: Manufacturer's Minimum Design Flow

### Conditions of Issue.

This Test Report is issued subject to the conditions stated in current issue of 'BSI Terms of Service'. The results contained herein apply only to the particular sample(s) tested and to the specific tests carried out, as detailed in this Test Report. The issuing of this Test Report does not indicate any measure of Approval, Certification, Supervision, Control or Surveillance by BSI of any product. No extract, abridgement or abstraction from a Test Report may be published or used to advertise a product without the written consent of BSI, who reserve the absolute right to agree or reject all or any of the details of any items or publicity for which consent may be sought.

Should you wish to speak with BSI in relation to this report, please contact Customer Services on +44 (0)8450 80 9000.

BSI Kitemark House Maylands Avenue Hemel Hempstead Hertfordshire HP2 4SQ



Opinions and Interpretations expressed herein are outside the scope of our UKAS accreditation.

Unless otherwise stated, any results not obtained from testing in a BSI laboratory are outside the scope of our UKAS accreditation.



# Test Results.

### BS EN 149:2001 + A1:2009

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

CLAUSE	REQUIREMENTS	ASSESSMENT
7.1	General	
	In all tests all samples shall meet the requirements.	-
7.2	Nominal values and tolerances	
	Unless otherwise specified, the values stated in this European Standard are expressed as nominal values. Except for temperature limits, values, which are not stated as maxima or minima, shall be subject to a tolerance of $\pm 5\%$ . Unless otherwise specified, the ambient temperature for testing shall be (16 – 32) °C, and the temperature limits shall be subject to an accuracy of $\pm$ 1°C.	-
7.3	Visual Inspection	
	The visual inspection shall also include the marking and the information supplied by the manufacturer.	Pass (1)
7.5	Material	
	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.	Pass
	After undergoing the conditioning described in clause 8.3.1 of the standard none of the particle filtering half masks shall have suffered mechanical failure of the facepiece or straps.	
	Three particle filtering half masks shall be tested.	Pass
	When conditioned in accordance with 8.3.1 and 8.3.2 the particle filtering half mask shall not collapse.	Pass
	Any material from the filter media released by the air flow through the filter shall not constitute a hazard or nuisance for the wearer.	Pass
	Testing shall be done in accordance with 8.2.	
7.8	Finish of parts	
	Parts of the device likely to come into contact with the wearer shall have no sharp edges or burrs.	Pass
	Testing shall be done in accordance with 8.2.	

(1) Marking and user information were not assessed as requested by BSI Product Certification



CLAUSE	REQUIREMENTS	ASSESSMENT
7.9	Leakage	
7.9.1	Total inward leakage	
	The laboratory tests shall indicate that the particle filtering half mask can be used by the wearer to protect with high probability against the potential hazard to be expected.	Pass (1) See Table A
	The total inward leakage consists of three components: face seal leakage, exhalation valve leakage (if exhalation valve fitted) and filter penetration.	See Table A
	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 EED1	

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

Testing shall be done in accordance with 8.5.

**Table A:** Clause 7.9.1 - Total inward leakage.

			Inward leakage (%).					
Test	Sample		Α	В	С	D	E	Average
candidate		condition	Walking	Walking with head side to side	Walking with head up & down	Walking and talking	Walking	
LM2	8	TC	7.2973	3.9167	5.7087	3.2365	5.1736	5.0666
SI1	9	TC	0.2104	0.2047	0.2353	0.2276	0.1237	0.2003
KH1	10	TC	0.2112	0.2296	0.2503	0.5231	0.6973	0.3823
CB1	11	TC	3.0178	1.8510	8.0745	1.8897	5.6798	4.1025
JW1 (2)	12	TC	0.5893	0.9512	0.7366	0.4300	0.6535	0.6721

<sup>(1)</sup> Results for the remaining 'as received' samples are covered in BSI Test Report number 3220780 for the BSI COVID-19 filtering face piece technical specification testing.

(2) Earloop clip used.



CLAUSE	REQUIREMENTS	ASSESSMENT
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### 7.9.2 Penetration of filter material

The penetration of the filter of the particle filtering half mask shall meet the requirements of Table  ${\bf 1}$ 

A total of 9 samples of particle filtering half masks shall be tested for each aerosol. Testing in accordance with 8.11 using the Penetration test according to EN 13274-7, shall be performed on:

Pass (1) See Tables B and C

3 samples as received,

3 samples after the simulated wearing treatment described in 8.3.1.

Testing in accordance with 8.11 using the Exposure test with a specified mass of test aerosol of 120 mg, and for particle filtering devices claimed to be re-usable additionally the Storage test, according to EN 13274-7, shall be performed:

Pass (1) See Table D and E

for non-re-usable devices on:

3 samples after the test for mechanical strength in accordance with 8.3.3 followed by temperature conditioning in accordance with 8.3.2.

for re-usable devices on:

3 samples after the test for mechanical strength in accordance with 8.3.3 followed by temperature conditioning in accordance with 8.3.2 and followed by one cleaning and disinfecting cycle according to the manufacturer's instruction.

N/A (2)

**Table B:** Clause 8.11 - Sodium Chloride penetration test.

Cample	Pre-test	Continuous flow Penetration		tion (%)
Sample	condition	(l/min)	Limit	Measured
16	SW	95	6.0	0.1635
17	SW	95	6.0	0.1645
18	SW	95	6.0	0.1542

**Table C:** Clause 8.11 - Paraffin oil penetration test.

Cample	Pre-test	Continuous flow	Penetra	tion (%)
Sample	condition	(l/min)	Limit	Measured
22	SW	95	6.0	1.1895
23	SW	95	6.0	1.9665
24	SW	95	6.0	1.6080

Results for the remaining 'as received' samples are covered in BSI Test Report number 3220780 for the BSI COVID-19 filtering face piece technical specification testing.

<sup>(2)</sup> Not a design feature of this product.



CLAUSE	REQUIREMENTS	ASSESSMENT
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7.9.2 Penetration of filter material (continued)

**Table D:** Clause 8.11. Exposure test Sodium Chloride.

	Sample 28 MS TC	Sample 29 MS TC	Sample 30 MS TC	
Flow through filter		95 l/min		
Elapsed time (minutes)	Measured penetration % (Maximum permitted penetration 6.0 %)			
5	0.239798 (1)	0.100650 (1)	0.149081 (1)	
10	0.192890	0.079177	0.121274	
15	0.141387 0.065270 0.09		0.096979	
20	0.090230	0.052679	0.076271	
25	0.056086	0.041193	0.054319	
30	0.033608 0.032297 0.03879		0.038793	
Result	Pass	Pass	Pass	

<sup>(1)</sup> The reading at which 5 subsequent sampling intervals showed a declining filter penetration. The testing was terminated without the 120mg exposure limit being reached, as permitted by BS EN 13274-7.



CLAUSE	REQUIREMENTS	ASSESSMENT	l
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7.9.2 Penetration of filter material (continued)

**Table E:** Clause 8.11Paraffin oil exposure test.

	Sample 25 MS TC	Sample 26 MS TC	Sample 27 MS TC
Flow through filter		95 l/min	
Elapsed time (minutes)	(Maxim	Measured penetration % num permitted penetratio	
3	2.1110	1.5950	1.9965
5	2.2125	1.6255	2.0845
10	2.4525	1.9880	2.3595
15	2.5900	1.9280	2.4440
20	3.0625     2.0000     2.66       3.2990     2.1675     2.76       3.2725     2.2115     2.91		2.5800
25			2.6650
30			2.7670
35			2.9130
40			3.0045
45	3.6010	2.3765	3.0950
50	3.6315	2.3965	3.1885
55	3.7715 2.4610 3.220		3.2285
60	3.8520	2.5240	3.3385
(1)	4.0125	2.5060	3.3755
Result	Pass	Pass	Pass

<sup>(1)</sup> A loading of 120 mg was achieved after a period of 63 minutes, 10 seconds had elapsed.



CLAUSE	REQUIREMENTS	1		ASSESSMENT
7.10	Compatibility w			
	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.			Pass
	Testing shall be do	one in accordance with 8.4 and 8.5.		
7.11	Flammability			
	The material used shall not present a danger for the wearer and shall not be of a highly flammable nature.			
	•	hen tested, the particle filtering half mask shall not burn or not continue to burn for more than seconds after removal from the flame.		
	The particle filtering half mask does not have to be usable after the test.			
	Testing shall be done in accordance with 8.6.			
	Table F: Clause 8.6 – Flammability.			_
	Sample	Area exposed	Comments	
	34 AR	Filter material, welding.	Did not ignite.	

### 7.13 Head harness

35 AR

36 TC

37 TC

The head harness shall be designed so that the particle filtering half mask can be donned and removed easily.

Earloop, vertical welding.

Filter material, welding.

Earloop, vertical welding.

The head harness shall be adjustable or self-adjusting and shall be sufficiently robust to hold the particle filtering half mask firmly in position and be capable of maintaining total inward leakage requirements for the device.

Pass

Did not ignite.

Did not ignite.

Did not ignite.

Testing shall be done in accordance with 8.4 and 8.5.

### 7.14 Field of vision

The field of vision is acceptable if determined so in practical performance tests.

Pass

Testing shall be done in accordance with 8.4.



CLAUSE	REQUIREMENTS	ASSESSMENT
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### 7.16 Breathing resistance

The breathing resistances apply to valved and valveless particle filtering half masks and shall meet the requirements of Table 2.

Testing shall be done in accordance with 8.9.

A total of 9 valveless particle filtering half masks shall be tested:

3 as received, 3 after temperature conditioning in accordance with 8.3.2 and 3 after the test for simulated wearing in accordance with 8.3.1.

Pass\* (1) (2) See Tables G, H and I

Testing shall be done in accordance with 8.9.

A total of 12 valved particle filtering half masks shall be tested: 3 as received, 3 after temperature conditioning in accordance with 8.3.2, 3 after the test for simulated wearing in accordance with 8.3.1, and 3 after the flow conditioning in accordance with 8.3.4.

N/A(3)

Testing shall be done in accordance with 8.9.

**Table G:** Clause 8.9 – Breathing resistance. Inhalation resistance at a continuous flow.

Sample	Pre-test condition	Flow (I/min)	Limit (mbar)	Measured (mbar)
16	SW	30	0.7	0.54
17	SW	30	0.7	0.50
18	SW	30	0.7	0.53
31	TC	30	0.7	0.48
32	TC	30	0.7	0.47
33	TC	30	0.7	0.46

**Table H:** Clause 8.9 – Breathing resistance. Inhalation resistance at a continuous flow.

Sample	Pre-test condition	Flow (I/min)	Limit (mbar)	Measured (mbar)
16	SW	95	2.4	1.89
17	SW	95	2.4	1.87
18	SW	95	2.4	1.89
31	TC	95	2.4	1.79
32	TC	95	2.4	1.79
33	TC	95	2.4	1.72

<sup>(1)</sup> Results for the remaining 'as received' samples are covered in BSI Test Report number 3220780 for the BSI COVID-19 filtering face piece technical specification testing.

(3) Not a design feature of this product.

<sup>(2)</sup> Results for exhalation resistance are within the uncertainty of measurement, but compliance is more probable than non-compliance.



CLAUSE	REQUIREMENTS	ASSESSMENT	I
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### 7.16 Breathing resistance (continued)

**Table I:** Clause 8.9 – Breathing resistance. Exhalation resistance at a continuous flow, measured in five orientations with the highest value recorded.

		I .		
Sample	Pre-test condition	Flow (I/min)	Limit (mbar)	Measured (mbar)
16	SW	160	3.0	2.87
17	SW	160	3.0	2.96
18	SW	160	3.0	2.95
31	TC	160	3.0	2.89
32	TC	160	3.0	2.84
33	TC	160	3.0	2.79

# Appendix A. – Test Panel Data

Test Candidate	Facial Dimensions (mm)					
	Length of face	Width of face	Face depth	Width of mouth	Head Circumference	Gender
JW1	116	126	122	48	570	Male
SI1	121	135	142	48	575	Male
LM2	110	148	125	47	567	Male
KH1	112	142	115	60	585	Male
CB1	117	147	130	57	566	Male

Note: All candidates were clean shaven

# bsi.

# Product photographs.





Front view Side view



Inside view

\*\*\* End of Report \*\*\*



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Test Report No.: 244315789a 001

Client: SUNBEAM INTERNATIONAL GMBH

Contact Information: Schumanstr. 12, 52146 Würselen, Germany

Contact Person: Edward Zhao

# Sample Description As Declared:

No. Of Sample : 80 pcs

Product Description : Personal Protective Respitator Mask

Product Type : Single shift use only

Material : -

Colour : White Lot No./Batch Code : -

Lot No./Batch Code : Buyer Name : -

Trademark : HYGISUN
Type-identifying : HS0501A
Claimed Classification : FFP2 NR

Manufacturer : Hunan Dreaming Cloud E-Commerce Co., Ltd.

Country of Origin : - Sales Destination (Country) : -

Test Type : Full Test

Test Specification : EN 149:2001 + A1:2009 Respiratory Protective Devices - Filtering Half

Masks to Protect Against Particles - Requirements, Testing and Marking

Other Information : -

Sample Obtaining Method: Sending by customer

**Delivery Condition:** Apparent good, samples tested as received

**Sample Receiving date:** 2021-03-04 & 2021-04-21

**Testing Period:** 2021-03-04 to 2021-04-01 & 2021-04-21 to 2021-04-27

Place of Testing: Textiles laboratory Shanghai

For and on behalf of TÜV Rheinland (Shanghai) Co., Ltd.

2021-04-30 Carmen Yan / Department Manager

Date Name/Position

Sample information is provided by customer. Test result is drawn according to the kind and extent of tests performed.

This test report relates to the above mentioned test sample. Without permission of the test center this test report is not permitted to be duplicated in extracts. This test report does not entitle to carry any safety mark on this or similar products.

'Decision Rule" document announced in our website (https://www.tuv.com/landingpage/en/qm-gcn/) describes the statement of conformity and its rule of enforcement for test results are applicable throughout this test report.



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# **Summary of Test Results:**

7.3 Visual Inspection P 7.4 Package P 7.5 Material P 7.6 Cleaning And Disinfection N/A 7.7 Practical Performance P 7.8 Finish Of Parts P 7.9.1 Leakage P
7.5 Material P  7.6 Cleaning And Disinfection N/A  7.7 Practical Performance P  7.8 Finish Of Parts P  7.9.1 Leakage P
7.6 Cleaning And Disinfection N/A 7.7 Practical Performance P 7.8 Finish Of Parts P 7.9.1 Leakage P
7.7 Practical Performance P 7.8 Finish Of Parts P 7.9.1 Leakage P
7.8 Finish Of Parts P 7.9.1 Leakage P
7.9.1 Leakage P
7.9.2 Penetration Of Filter Material P
7.10 Compatibility With Skin P
7.11 Flammability P
7.12 Carbon Dioxide Content Of The Inhalation Air P
7.13 Head Harness P
7.14 Field Of Vision P
7.15 Exhalation Valve(s) N/A
7.16 Breathing Resistance P
7.17 Clogging N/A
7.18 Demountable Parts N/A
10 Information To Be Supplied By The Manufacturer P
9 Marking P

Note: P = Pass F = Fail

# = No Comment - = Did Not Perform N/R = Not Request N/A = Not Applicable

# **Material List:**

Material No.	Material	Color	Location	Remark
M001	Whole Product	White	Personal Protective Respitator Mask	Received on 2021.03.04
M001'	Whole Product	White	Personal Protective Respitator Mask	Received on 2021.04.21



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# **Visual Inspection**

Test Method: EN 149:2001+A1:2009 Clause 8.2

Clause	Item	M001
7.3	The visual inspection shall also include the marking and the information supplied by the manufacturer.	Pass
7.4	Particle filtering half masks shall be offered for sale packaged in such a way that they are protected against mechanical damage and contamination before use.	Pass
7.5	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.	Pass
	After undergoing the conditioning described in 8.3.1 none of the particle filtering half masks shall have suffered mechanical failure of the face piece or straps.	Pass
	When conditioned in accordance with 8.3.1 and 8.3.2 the particle filtering half mask shall not collapse.	Pass
	Any material from the filter media released by the air flow through the filter shall not constitute a hazard or nuisance for the wearer.	Pass
7.8	Parts of the device likely to come into contact with the wearer shall have no sharp edges or burrs	Pass
7.18	All demountable parts (if fitted) shall be readily connected and secured, where possible by hand.	N/A

# Remark:

N/A: Due to no relevent information/material

N/R: Due to not request



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## **Practical Performance**

Test Method: EN 149:2001+A1:2009 Clause 8.4 & 8.5

Clause	Item	M001
7.7	Wearing	Pass
7.7	Walking test	Pass
7.7	Work simulation test	Pass
7.10	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	Pass
7.13	The head harness shall be designed so that the particle filtering half mask can be donned and removed easily. The head harness shall be adjustable or self-adjusting and shall be sufficiently robust to hold the particle filtering half mask firmly in position and be capable of maintaining total inward leakage requirements for the device	Pass
7.14	The field of vision is acceptable if determined so in practical performance tests	Pass

# Remark:

N/A: Due to no relevent information/material

N/R: Due to not request



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Clause 7.9.1: Leakage

Test Method : EN 149:2001+A1:2009 Clause 8.5

Requirement : FFP2 :

At least 46 out of the 50 individual exercise results for total inward leakage ≤ 11% At least 8 out of the 10 individual wearer arithmetic means for the total inward

leakage ≤ 8%

	M001										
			Leakage (%)								
Condition	Specimen No.	Subject	Walk	Head Side/Side	Head Up/Down	Talk	Walk	Mean			
	1	ВМ	4.927	7.304	9.711	5.581	2.803	6.065			
	2	ACH	3.824	6.874	8.145	8.664	5.217	6.545			
As received	3	ML	4.128	6.229	8.225	7.422	3.877	5.976			
	4	LLC	3.397	6.785	8.199	6.357	4.012	5.734			
	5	DG	3.981	6.932	8.902	7.559	4.331	6.341			
	6	SG	4.104	5.181	10.648	7.685	3.493	6.222			
	7	YL	6.247	5.487	8.375	8.247	6.027	6.877			
After conditioning	8	KQ	5.525	6.028	9.084	8.122	5.021	6.756			
	9	KXH	6.001	6.439	9.119	8.074	5.387	7.004			
	10	YY	5.743	6.009	8.911	7.936	5.111	6.742			
Conclusion	on				Pass						

	Facial Dimension Of Subject (mm)										
Subject	Subject BM ACH ML LLC DG SG YL KQ KXH YY LL								LL		
Face length	135	127	120	120	130	135	115	120	130	130	121
Face width	160	159	133	140	145	155	135	135	155	165	163
Face Depth	130	122	115	115	132	132	118	115	120	143	142
Mouth Width	52	55	52	50	50	55	48	50	52	50	45



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# Clause 7.9.2: Penetration Of Filter Material

Test method : EN 149:2001+A1:2009 Clause 8.11

Requirement : FFP2: ≤6%

	M	001	
Aerosol	Condition	Specimen No.	Penetration (%)
	As received	1	0.048
Sodium chloride	As received	2	0.223
	As received	3	0.226
	Simulated wearing treatment	4	0.568
	Simulated wearing treatment	5	0.483
	Simulated wearing treatment	6	0.439
Penetration	Mechanical strength + Temperature conditioned @ Exposure test of 120mg	7	0.322
	Mechanical strength + Temperature conditioned @ Exposure test of 120mg	8	0.282
	Mechanical strength + Temperature conditioned @ Exposure test of 120mg	9	0.289
	As received	10	0.566
	As received	11	0.536
	As received	12	0.521
	Simulated wearing treatment	13	0.586
	Simulated wearing treatment	14	0.623
Paraffin oil	Simulated wearing treatment	15	0.637
Penetration	Mechanical strength + Temperature conditioned @ Exposure test of 120mg	16	0.984
	Mechanical strength + Temperature conditioned @ Exposure test of 120mg	17	2.392
	Mechanical strength + Temperature conditioned @ Exposure test of 120mg	18	1.664
Conclusion		Pass	



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Clause 7.11: Flammability

: EN 149:2001+A1:2009 Clause 8.6 Test method

Requirement : ≤5s

M001										
Item	Condition	Specimen No.	Test results							
	As received	1	DNI							
	As received	2	DNI							
Afterflame time (s)	After conditioning	3	DNI							
	After conditioning	4	DNI							
Cor	nclusion	Pass								

Remark:

DNI-Do not ignite



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# Clause 7.12: Carbon Dioxide Content Of The Inhalation Air

: EN 149:2001+A1:2009 Clause 8.7 Test Method

Requirement : ≤1%

M001								
Item	Condition	Test results						
	As received	Specimen 1	0.58					
• • • • • • • • • • • • • • • • • • • •	As received	Specimen 2	0.59					
Content (%)	As received	Specimen 3	0.61					
	As received	Mean	0.60					
Cor	nclusion	Pass						



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Clause 7.16: Breathing Resistance

Test Method : EN 149:2001+A1:2009 Clause 8.9

Requirement : FFP2:

Inhalation: 30l/min: ≤0.7mbar Inhalation: 95l/min: ≤2.4mbar Exhalation: 160l/min: ≤3.0mbar

	M001'															
Flow rate (I/	min)		Resistance (mbar)													
A			Sp	ecime	n 1			Specimen 2				Specimen 3				
As receive	eu -	А	В	С	D	Е	А	В	С	D	Е	Α	В	С	D	Е
Inhalation	30	0.4	0.4	0.4	0.4	0.4	0.5	0.5	0.5	0.5	0.5	0.4	0.4	0.4	0.4	0.4
IIIIIaialiOII	95	1.3	1.3	1.3	1.3	1.3	1.4	1.4	1.4	1.4	1.4	1.4	1.4	1.4	1.4	1.4
Exhalation	160	2.0	2.0	2.0	2.0	2.0	2.2	2.2	2.2	2.2	2.2	2.2	2.2	2.2	2.2	2.2
Simulated we	aring	Specimen 4				Specimen 5				Specimen 6						
treatmen	it	А	В	С	D	Е	А	В	С	D	Ш	А	В	С	D	Е
Inhalation	30	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5
IIIIIaiaiioii	95	1.5	1.5	1.5	1.5	1.5	1.4	1.4	1.4	1.4	1.4	1.5	1.5	1.5	1.5	1.5
Exhalation	160	2.4	2.4	2.4	2.4	2.4	2.3	2.3	2.3	2.3	2.3	2.3	2.3	2.3	2.3	2.3
Temperatu	ıre		Specimen 7				Specimen 8				Specimen 9					
conditione	ed	А	В	O	D	Е	А	В	С	D	Ш	Α	В	С	D	Е
Inhalation	30	0.4	0.4	0.4	0.4	0.4	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5
innaiation	95	1.4	1.4	1.4	1.4	1.4	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5
Exhalation	160	2.3	2.3	2.3	2.3	2.3	2.4	2.4	2.4	2.4	2.4	2.2	2.2	2.2	2.2	2.2
Conclusion Pass																

Remark: A: facing directly ahead;

B: facing vertically upwards;

C: facing vertically downwards;

D: lying on the left side;

E: lying on the right side



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# Marking

Test Method: EN 149:2001+A1:2009 Clause 9

M001	
9.1 Packaging	
The following information shall be clearly and durably marked on the smallest comme packaging or legible through it if the packaging is transparent.	rcially available
9.1.1 The name, trademark or other means of identification of the manufacturer or supplier.	Present
9.1.2 Type-identifying marking.	Present
9.1.3 Classification The appropriate class (FFP1, FFP2 or FFP3) followed by a single space and then: "NR" if the particle filtering half mask is limited to single shift use only. Example: FFP3 NR, or "R" if the particle filtering half mask is re-usable. Example: FFP2 R D.	Present
9.1.4 The number and year of publication of this European Standard.	Present
9.1.5 At least the year of end of shelf life. The end of shelf life may be informed by a pictogram as shown in Figure 12a, where yyyy/mm indicates the year and month.	Present
9.1.6 The sentence 'see information supplied by the manufacturer', at least in the official language(s) of the country of destination, or by using the pictogram as shown in Figure 12b.	Present
9.1.7 The manufacturer's recommended conditions of storage (at least the temperature and humidity) or equivalent pictogram, as shown in Figures 12c and 12d.	Present
9.1.8 The packaging of those particle filtering half masks passing the dolomite clogging test shall be additionally marked with the letter "D". ID This letter shall follow the classification marking preceded by a single space.	N/A
9.2 Particle filtering half mask	
Particle filtering half masks complying with this European Standard shall be clearly an with the following:	d durably marked
9.2.1 The name, trademark or other means of identification of the manufacturer or supplier.	Present
9.2.2 Type-identifying marking.	Present
9.2.3 The number and year of publication of this European Standard.	Present
9.2.4 Classification The appropriate class (FFP1, FFP2 or FFP3) followed by a single space and then: "NR" if the particle filtering half mask is limited to single shift use only. Example: FFP3 NR, or "R" if the particle filtering half mask is re-usable. Example: FFP2 R D.	Present
9.2.5 If appropriate the letter D (dolomite) in accordance with clogging performance. This letter shall follow the classification marking preceded by a single space.	N/A
9.2.6 Sub-assemblies and components with considerable bearing on safety shall be marked so that they can be identified.	N/A



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# Remark:

- The evaluation is based on artwork. 1.
- N/A: Not applicable



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# Information To Be Supplied By The Manufacturer

Test Method: EN 149:2001+A1:2009 Clause 10

M001	
10.1 Information supplied by the manufacturer shall accompany every smallest commercial available package	Present
10.2 Information supplied by the manufacturer shall be at least in the official language(s) of the country of destination	Present
10.3 The information supplied by the manufacturer shall contain all information ned qualified persons on	cessary for trained and
- application/limitations	Present
- the meaning of any colour coding	N/A
- checks prior to use	Present
- donning, fitting	Present
- use	Present
- maintenance (e.g. cleaning, disinfecting), if applicable	N/A
- storage	Present
- the meaning of any symbols/pictograms used	Present
of the equipment	
10.4 The information shall be clear and comprehensible. If helpful, illustrations, part numbers, marking shall be added	Present
10.5 Warning shall be given against problems likely to be encountered, for example	e:
- fit of particle filtering half mask (check prior to use)	Present
- it is unlikely that the requirements for leakage will be achieved if facial hair passes under the face seal	Present
- air quality (contaminants, oxygen deficiency)	Present
- use of equipment in explosive atmosphere	Present
10.6 The information shall provide recommendations as to when the particle filtering half mask shall be discarded	Present
10.7 For devices marked "NR", a warning shall be given that the particle filtering half mask shall not be used for more than one shift	Present

## Remark:

- 1. The evaluation is based on artwork.
- 2. N/A: Not applicable



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# Photo(s):











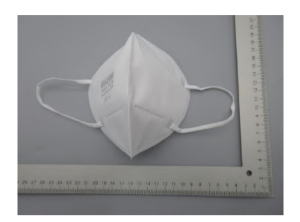




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# Photo(s):







## General Terms and Conditions of Business of TÜV Rheinland in Greater China

- These General Terms and Conditions of Business of TÜV Rheinland in Greater China ("GTCB") is made between the client and one or more member entities of TÜV Rheinland in Greater China as applicable as the case may be c'TÜV Rheinland'. The Greater China hereof refers to Mainland China, Hong Kong and Taiwan.The client hereof includes:
- a natural person capable to form legally binding contracts under the applicable laws who concludes the contract not for the purpose of a daily use;
- (ii) the incorporated or unincorporated entity duly organized, validly existing and capable to form legally binding contracts under the applicable law.
- 1.2 The following terms and conditions apply to agreed services including consultancy services, information, deliveries and similar services as well as ancillary services and other secondary obligations provided within the scope of contract performance.
- Any standard terms and conditions of the client of any nature shall not apply and sha hereby be expressly excluded. No standard contractual terms and conditions of the clien shall form part of the contract even if TÜV Rheinland does not explicitly object to them.
- In the context of an ongoing business relationship with the client, this GTCB shall also apply to future contracts with the client without TÜV Rheinland having to refer to them separately

Unless otherwise agreed, all quotations submitted by  $T\ddot{U}V$  Rheinland can be changed by  $T\ddot{U}V$  Rheinland without notice prior to its acceptance and confirmation by the other party.

## Coming into effect and duration of contracts

- The contract shall come into effect for the agreed terms upon the quotation letter of TÜV Rheinland or a separate contractual document being signed by both contracting parties, or upon the works requested by the client being carried out by TÜV Rheinland. If the client instructs TÜV Rheinland without receiving a quotation from TÜV Rheinland (quotation). TÜV Rheinland, in its sice discretion, entitled to accept the order by giving written cof such acceptance (including notice sent via electronic means) or by performing the requested service.
- 3.2 The contract term starts upon the coming into effect of the contract in accordance with article 3.1 and shall continue for the term agreed in the contract.
- 3.3 If the contract provides for an extension of the contract term, the contract term will be extended by the term provided for in the contract unless terminated in writing by either party with a six-week notice prior to the end of the contractual term.

### Scope of services

- The scope and type of the services to be provided by TÜV Rheinland shall be specified in the contractually agreed service scope of TÜV Rheinland by both parties. If no such separate service scope of TÜV Rheinland exists, then the written confirmation of order by TÜV Rheinland shall be decisive for the service to be provided.
- 4.2 The agreed services shall be performed in compliance with the regulations in force at the time the contract is entered into.
- TÜV Rheinland is entitled to determine, in its sole discretion, the method and nature of the assessment unless otherwise agreed in writing or if mandatory provisions require a specific procedure to be followed.
- On execution of the work there shall be no simultaneous assumption of any guarar the correctness (proper quality) and working order of either tested or examined parts the installation as a whole and its upstream and/or downstream processes, organiss use and application in accordance with regulations, nor of the systems on which installation is based. In particular, 70th heinland shall assume no responsibility for construction, selection of materials and assembly of installations examined, nor to use and application in accordance with regulations, unless these questions are exprovered by the contract.
- 4.5 In the case of inspection work, TÜV Rheinland shall not be responsible for the accuracy or checking of the safety programmes or safety regulations on which the inspections are based, unless otherwise expressly agreed in writing.
- 4.6 If mandatory legal regulations and standards or official requirements for the agreed service scope change after conclusion of the contract, with a written notice to the client, TUV Rheinland shall be entitled to additional remuneration for resulting additional expenses.
- 4.7The services to be provided by TÜV Rheinland under the contract are agreed exclusively with the client. A contract of third parties with the services of TÜV Rheinland, as well as making available of and justifying confidence in the work results (test reports, test results, expert reports, etc.) is not part of the agreed services. This also applies if the client passes or work results in full or in extracts to third parties in accordance with clause 11.4.

## Performance periods/dates

- The contractually agreed periods/dates of performance are based on estimates of involved which are prepared in line with the details provided by the client. They be binding if being confirmed as binding by TÜV Rheinland in writing.
- If binding periods of performance have been agreed, these periods shall not commence until the client has submitted all required documents to TÜV Rheinland.
- 5.3 Articles 5.1 and 5.2 also apply, even without express approval by the client, to all extensions of agreed periods/dates of performance not caused by TÜV Rheinland.
- 5.4TÜV Rheinland is not responsible for a delay in performance, in particular if the client has not fulfillided his duties to cooperate in accordance with clause 6.1 or has not done so in time and, in particular, has not provided TÜV Rheinland with all documents and information required for the performance of the service as specified in the contract.
- 5.5If the performance of TÜV Rheinland is delayed due to unforeseeable circumstances such as force majeure, strikes, business disruptions, governmental regulations, transport obstacles, etc., TÜV Rheinland is entitled to postpone performance for a reasonable period of time which corresponds at least to the duration of the hindrance plus any time period which may be required to resume performance.

## The client's obligation to cooperate

- The client shall guarantee that all cooperation required on its part, its agents or third parties will be provided in good time and at no cost to  $T\ddot{U}V$  Rheinland.
- 6.2 Design documents, supplies, auxiliary staff, etc. necessary for performance of the services shall be made available free of charge by the client. Moreover, collaborative action of the client must be undertaken in accordance with legal provisions, stardards, safety regulations and accident prevention instructions. And the client represents and warrants that:
  - a) it has required statutory qualifications:
  - b) the product, service or management system to be certified complies with applicable laws and regulations; and
  - c) it doesn't have any illegal and dishonest behaviours or is not included in the list of Enterprises with Serious Illegal and Dishonest Acts of People's Republic of China.
  - If the client breaches the aforesaid representations and warranties, TÜV Rheinland is entitled to i) immediately terminate the contract/order without prior notice; and ii) withdraw the issued testing report/certificates if any.
- The client shall bear any additional cost incurred on account of work having to be redone or being delayed as a result of late, incorrect or incomplete information provided by or lack of proper cooperation from the client. Even where a fixed or maximum price is agreed, TÜV Rheinland shall be entitled to charge extra fees for such additional expense.

- If the scope of performance is not laid down in writing when the order is placed, invoicing shall be based on costs actually incurred. If no price is agreed in writing, invoicing shall be made in accordance with the price list of TÜV Rheinland valid at the time of performance.
- 7.2 Unless otherwise agreed, work shall be invoiced according to the progress of the work.
- 7.3 If the execution of an order extends over more than one month and the value of the contract or the agreed fixed price exceeds £2,500.00 or equivalent value in local currency, TÜV Rheinland may demand payments on account or in instalments.

- All invoice amounts shall be due for payment without deduction on receipt of the invoice. No discounts and rebates shall be granted.
- Payments shall be made to the bank account of TÜV Rheinland as indicated on the invoice, stating the invoice and client numbers.
- 8.3 In cases of default of payment, TÜV Rheinland shall be entitled to claim default interest at the applicable short term loan interest rate publicly announced by a reputable commercial bank in the country where TÜV Rheinland is located. At the same time, TÜV Rheinland reserves the right to claim further damages.
- Should the client default in payment of the invoice despite being granted a reasonable grace period, TÜV Rheinland shall be entitled to cancel the contract, withdraw the certificate, claim damages for non-performance and refuse to continue performance of the
- 8.5 The provisions set forth in article 8.4 shall also apply in cases involving returned cheques, cessation of payment, commencement of insolvency proceedings against the client's assets or cases in which the commencement of insolvency proceedings has been dismissed due to lack of assets.

- 8.6 Objections to the invoices of TÜV Rheinland shall be submitted in writing within two w of receipt of the invoice
- 8.7 TÜV Rheinland shall be entitled to demand appropriate advance payments
- 8.7 IUV kneinland shall be entitled to desire fieles at the beginning of a month if overheads and/or purchase costs have increased. In this case, TÜV Rheinland shall notify the client in writing of the rise in fees. This notification shall be issued one month prior to the date on which the rise in fees. This notification shall be issued one month prior to the date on which the rise in fees shall come into effect (period of notice of changes in fees). If the rise in fees remains under 5% per contractual year, the client shall not have the right to terminate the contract. If the rise in fees exceed 5% per contractual year, the client shall be described to the right to terminate the contract in the rise in fees acceed 5% per contractual year, the client shall be described to the right to terminate the contract is not terminated, the changed fees shall be deemed to have been agreed upon by the time of the expiry of the notice period.
- 8.9 Only legally established and undisputed claims may be offset against claims by TÜV Rheinland.

- 9.1 Any part of the work result ordered which is complete in itself may be presented by TÜV Rheinland for acceptance as an instalment. The client shall be obliged to accept it interesting the complete of the complete or the complete or
- 9.2 If acceptance is required or contractually agreed in an individual case, this shall be deemed to have taken place two (2) weeks after completion and handover of the work, unless the client refuses acceptance within this period stating at least one fundmental breach of contract by TÜV Rheinland.
- 9.3 The client is not entitled to refuse acceptance due to insignificant breach of contract by TÜV Rheinland.
- 9.4 If acceptance is excluded according to the nature of the work performance of TÜV Rheinland, the completion of the work shall take its place.
- rnemiano, the completion of the work shall take its place.

  9. If the claim was unable to make use of the time windows provided for within the scope of contribution procedure for auditing/performance by TUV. Rheinland and the certificate severe to be provided to the provided provided to t
- 9.6 Insofar as the client has undertaken in the contract to accept services, TÜV Rheinland shall also be entitled to charge lump-sun damages in the amount of 10% of the order amount as compensation for expenses if the service is not called within one year after the order has been placed. The client reserves the right to prove that the TÜV Rheinland has incurred no damage whatsoever or only a considerably lower damage than the above mentioned tump and the contraction of the place of the reserves the reser

- 10. Confidentiality
  10.1-for the purpose of these terms and conditions, "confidential information" means all information, documents, images, drawings, know-how, data, samples and project documentation which one party (the "disclosing party") hands over, transfers or otherw discloses to the other party (the "foceiving party"), and the confidential information reducing performance of work by TUV Rheinfand, including product testing data, defects, conformity to the technical standard and related reports. Confidential information is exp not the data and know-how collected, compled or otherwise obtained by TUV Rheinfand (non-personal) within the scope of the provision of services by TUV Rheinfand. TUV Rheinfand is entitled to store, use, further develop and pass on the data obtained in connection with the provision of services for the purposes of developing new services, improving services and analysing the provision of services.
- 10.2 The disclosing party shall mark all confidential information disclosed in written form as confidential before passing it onto the receiving party. The same applies to confidential information is disclosed orally, the receiving party shall be appropriately information is disclosed orally, the receiving party shall be appropriately informed in advance and the disclosing party shall confirm in writing the confidentiality nature of the information within five working days of oral disclosure. Where the disclosing party fails to do so within the stipulated period, the receiving party shall not take any confidentiality holigations her enurient towards such information.
- 10.3 All confidential information which the disclosing party transmits or otherwise discloses to the receiving party and which is created during performance of work by TÜV Rheinland:

a)may only be used by the receiving party for the purposes of performing the contract, unless expressly otherwise agreed in writing by the disclosing party;

b)may not be copied, distributed, published or otherwise disclosed by the receiving party, unless this is necessary for fulfilling the purpose of the contract or TÜV Rheinland is requir to pass on confidential information, inspection reports or documentation to the governmen authorities, judicial court, accreditation bodies or third parties that are involved in the

communities treated by the receiving party with the same level of confidentiality as the party uses to protect its own confidential information, but never with a lesser level of confidentiality than that which is reasonably required.

- 10.4 The receiving party may disclose any confidential information received from the disclosing party only to those of its employees who need this information to perform the services required for the contract. The receiving party undertakes to obligh these employees to observe the same level of secrecy as set forth in this confidentiality clause.
- 10.5 Information for which the receiving party can furnish proof that:
  - a)it was generally known at the time of disclosure or has become general knowledge without violation of this confidentiality clause by the receiving party; or
  - b)it was disclosed to the receiving party by a third party entitled to disclose this information; or c)the receiving party already possessed this information prior to disclosure by the disclosing party; or
  - d)the receiving party developed it itself, irrespective of disclosure by the disclosing party, sha not be deemed to constitute "confidential information" as defined in this confidentiality clause
- 10.6 All confidential information shall remain the property of the disclosing party. The receiving party hereby agrees to immediately (i) return all confidential information, including all copie party hereby agrees to immediately (i) return all confidential information, including all copies, to the disclosing party, and/or (ii) on request by the disclosing party, to destroy all confidential information, including all copies, and confirm the destruction of this confidential information the disclosing party in writing, at any time if so requested by the disclosing party but at the latest and without special request after termination or expiry of the contract. This does not extend to include reports and certificates prepared for the client solety for the purpose of fulfilling the obligations under the contract, which shall remain with the client. However, TUV Rheinland is entitled to make file copies of such reports, certificates and confidential information that forms the basis for preparing these reports and certificates in order to evidence the correctness of its results and for general documentation purposes required by laws, regulations and the requirements of working procedures of TÜV Rheinland.
- 10.7 From the start of the contract and for a period of three years after termination or expiry of the contract, the receiving party shall maintain strict secrecy of all confidential information and shall not disclose this information to any third parties or use it for itself.

# 11. Copyrights and rights of use, publications

- 11.1 TÜV Rheinland shall retain all exclusive copyrights in the reports, expert reports/opinions, reports/results, results, calculations, presentations etc. prepared by TÜV Rheinland, unit otherwise agreed by the parties in a separeta agreement. As the owner of the copyright TÜV Rheinland is free to grant others the right to use the work results for individual or types of use tright of use?
- 11.2 The client receives a simple, unlimited, non-transferable, non-sublicensable right of use to the contents of the work results produced within the scope of the contract, unless otherwise agreed by the parties in a separate agreement. The client may only use such reports, expert reports/opinions, test reports/results, results calculations, presentations etc. prepared within the scope of the contract for the contractually agreed purpose.
- 11.3 The transfer of right of use of the generated work results regulated in clause 11.2. of the GTCB is subject to full payment of the remuneration agreed in favour of TÛV Rheinland.
- 11.4 The client may use work results only complete and unshortened. The client may only pass on the work results in full unless TÜV Rheinland has given its prior written consent to the partial passing on of work results
- 11.5 Any publication or duplication of the work results for advertising purposes or any further u the work results beyond the scope regulaed in clause 11.2 needs the prior written appror T/U Rheinland in each individual case.
- 11.6 TÜV Reinland may revoke a once given approval according to clause 11.5 at any time without stating reasons. In this case, the client is obliged to stop the transfer of the work results immediately at his own expense and, as far as possible, to withdraw publications.
- The consent of  $T\ddot{U}V$  Rheinland to publication or duplication of the work results does not entitle the client to use the corporate logo, corporate design or test/centification mark of  $T\ddot{U}V$

## 12 Liability of TÜV Rheinland

12.1 Irrespective of the legal basis, to the fullest extent permitted by applicable law, in the event of a breach of contractual obligations or tort, the liability of TÜV Rheinland for all damages, losses and reimbursement of expenses caused by TÜV Rheinland, its legal representatives and/or employees shall be limited to: (i) in the case of a contract with a fixed overall fee, three times the overall fee for the entire contract; (ii) in the case of a contract or annually recurring services, the agreed annual fee; (iii) in the case of a contract or annually recurring services, the agreed annual fee; (iii) in the case of a contract or entire the contract of the co

orders, three times of the fee for the individual order under which the damages or losses have occurred. Notwithstanding the above, in the event that the total and accumulated liability accumulated lia calculated according to the foregoing provisions exceeds 2.5 Million Euro or equiva amount in local currency, the total and accumulated liability of TÜV Rheinland shall be limited to and shall not exceed the said 2.5 Million Euro or equivalent amount in

- 12.2 The limitation of liability according to article 12.1 above shall not apply to damages losses caused by malice, intent or gross negligence on the part of TÜV Rheinland vicarious agents. Such limitation shall not apply to damages for a person's death, pirjury or illness.
- 12.3 In cases involving a fundamental breach of contract, TÜV Rheinland will be liable even w minor negligence is involved. For this purpose, a "fundamental breach" is breach of a man contractual obligation, the performance of which permits the due performance of the cont Any claim for damages for a fundamental breach of contract shall be limited to the amou damages reasonably foreseen as a possible consequence of such breach of contract a time of the breach (reasonably foreseeable damages), unless any of the circumstal described in article 12.2 applies.
- 12.4 TÜV Rheinland shall not be liable for the acts of the personnel made available by the client to support TÜV Rheinland in the performance of its services under the contract, unless such personnel made available is regarded as vicanious agent of TÜV Rheinland. IT TÜV Rheinland is not liable for the acts of the personnel made available by the client under the foregoing provision, the client shall indemnify TÜV Rheinland against any claims made by third parties arising from or in connection with such personnel's acts.
- 12.5 Unless otherwise contractually agreed in writing, TÜV Rheinland shall only be liable under the contract to the client.
- 12.6 The limitation periods for claims for damages shall be based on statutory provisions
- 12.7 None of the provisions of this article 12 changes the burden of proof to the disadvantage of the client

- 13.1When passing on the services provided by TÜV Rheinland or parts thereof to third parties in Greater China or other regions, the client must comply with the respectively applicable regulations of national and international export control tab.
- 13.2The performance of a contract with the client is subject to the proviso that there are no obstacles to performance due to national or international foreign trade legislations or embargos and/or sanctions, in the event of a violation, TDV Pheniand shall be entitled to terminate the contract with immediate effect and the client shall compensate for the fosses incured thereof by TDV Rehelland.

### 14. Data protection notice

Data protection notice

TÜV Rheinland processes personal data of the client for the purpose of fulfilling this contract. In addition, TÜV Rheinland also processes the data for other legal purposes in accordance with the relevant legal basis. The personal data of the client will only be disclosed to other natural or legal persons if the legal requirements are met. This also applies to transfers to third countries. The personal data will be deleted immediately as soon as a corresponding reason for deletion arises. Data subjects may exercise the following rights: right of objection, right of oretification, right of recessing limitation, right of objection, right of objection, right of the processing limitation, right to objection, right of data transferability. In addition, persons concerned by the data processing have the right to reveloc heir consent at any time with effect for the future, as well as the right to file a complaint with the competent data protection supervisor guildrow; For further death of the processor, please refer to the respective data protection formation. You can contact the Group Data Protection Officer of TÜV Rheinland by e-mail at datenschutz@de.tuv.com or by post at the following address: TÜV Rheinland AG, c/o Group Data Protection Officer, Am Grauen Stein, 51105 Cologne, Germany.

### 15. Test material: transport risk and storage

- 15.1The risk and costs for freight and transport of documents or test material to and from TÜV Rheinland as well as the costs of necessary disposal measures shall be borne by the client.
- 15.2Any destroyed and otherwise worthless test material will be disposed of by TÜV Rheinland for the client at the expense of the client, unless otherwise agreed.
- 15.3Undamaged test material shall be stored by TÜV Rheinland for four (4) weeks after completion of the test. If a longer storage period is desired, TÜV Rheinland charges an appropriate storage fee.
- 15.4After the expiry of the 4 weeks or any longer period agreed upon, the test material will be disposed of by TÜV Rheinland for the client for a fee in accordance with clause 15.2.

- 16.1 Notwithstanding clause 3.3 of the GTCB, TÜV Rheinland and the client are entitled to te the contract in its entirety or, in the case of services combined in one contract, eac combined parts of the contract individually and independently of the continuation remaining services with six (6) months notice to the end of the contractually agreed te
- 16.2For good causes, TÜV Rheinland may consider giving a written notice to the client to terminate the contract which includes but not limited to the following:
  - a) the client does not immediately notify TÜV Rheinland of changes in the conditions within the company which are relevant for certification or signs of such changes;
  - b) the client misuses the certificate or certification mark or uses it in violation of the contract;
  - c) in the event of several consecutive delays in payment (at least three times);
  - d) a substantial deterioration of the financial circumstances of the client occurs and as a result the payment claims of TÜV Rheinland under the contract are considerably endangered and TÜV Rheinland cannot reasonably be expected to continue the contractual relationship.
- 16.3.In the event of termination with written notice by TÜV Rheinland for good cause. TÜV Rheinland shall be entitled to a lump-sum claim for damages against the client if the conditions of a claim for damages sex sit. In this case, the client shall owe 15% of the remuneration to be paid until the end of the fixed contract term as lump-sum compensation. The client reserves the right to prove that there is no damage or a considerably lower damage, TÜV Rheinland reserves the right to prove a considerably higher damage in individual cases.
- 16.4TÜV Rheinland is also entitled to terminate the contract with written notice if the client has not been able to make use of the time windows for auditing /service provision provided by TÜV Rheinland within the scope of a certification procedure and the certificate therefore has to be withdrawn (for example during the performance of monitoring audits). Clause 16.3 applies

## 17. Partial invalidity, written form, place of jurisdiction and dispute resolution

- 17.1 All amendments and supplements must be in writing in order to be effective. This also applies to amendments and supplements to this clause 17.1.
- 17.2 Should one or several of the provisions under the contract and/or these terms and condition be or become ineffective, the contracting parties shall replace the invalid provision with legally valid provision that comes closest to the content of the invalid provision in legal a commercial terms.
- 17.3 Unless otherwise stipulated in the contract, the governing law of the contract and these terms and conditions shall be chosen following the rules as below:
  - a)if TÜV Rheinland in question is legally registered and existing in the People's Republic of China, the contracting parties hereby agree that the contract and these terms and conditions shall be governed by the laws of the People's Republic of China.
  - b)if TÜV Rheinland in question is legally registered and existing in Taiwan, the contracting parties hereby agree that the contract and these terms and conditions shall be governed by the laws of Taiwan.
- c)if TÜV Rheinland in question is legally registered and existing in Hong Kong, the contracting parties hereby agree that the contract and these terms and conditions shall be governed by the laws of Hong Kong.
- 17.4 Any dispute in connection with the contract and these terms and conditions or the execution thereof shall be settled friendly through negotiations. Unless otherwise stipulated in the contract, if no settlement or no agreement in respect of the extension of the negotiation period can be reached within two months of the arising of the dispute, that Despute shall be submitted:
  - ajin the case of TÜV Rheinland in question being legally registered and existing in the People's Republic of China, to China International Economic and Trade Arbitration Commission (CIETAC) to be settled by arbitration under the Arbitration Rules of CIETAC in force when the arbitration is submitted. The arbitration shall take place in Beijing, Shanghai, Shenzhen or Chongqing as appropriately chosen by the claiming party.
  - b)in the case of TÜV Rheinland in question being legally registered and existing in Taiwan, to Chinese Arbitration Association Taipel Branch to be arbitrated in accordance with its then current Rules of Arbitration. The arbitration shall take place in Taipei.
  - c)in the case of TÜV Rheinland being legally registered and existing in Hong Kong, to Hong Kong International Abitration Centre (HKIAC) to be settled by arbitration under the HKIAC Administered Abitration Rules in force when the Notice of Abitration is submitted in accordance with these rules. The arbitration shall take place in Hong Kong.
  - The decision of the relevant arbitration tribunal shall be final and binding on both parties. The arbitration fee shall be borne by the losing party.