

### KİNGFA 金发科技

PARTICLE FILTERING
HALF MASK
EN149:2001+A1:2009
FFP2
KF-A F10(SC)





More protective

MORE COMFORTABLE

### KINGFA INTRODUCTION





Established in 1993

Research, production and sales of advanced polymer materials

**Listed** on Shanghai Stock Exchange in 2004



Over 6500 employees

Annual production capacity exceeds

2 million tons

# WITHIN 27 YEARS OF DEVELOPMENT KINGFA REALIZED:

(	4	Company Founders	)—(	Over <b>6500</b>	Employees	
(	20	Thousand CNY Capital	)—(	29.1	Billion Total Assets(CNY)	
(	1	Small Workshop	)—(	47	Subsidiary Companies	
(	0	CNY Sales Volume	)—(	29.2	Billion Sales Volume(CNY)	

## KF-A F10(SC) FFP2



# Color Box (30 pcs/box)

**Size:** 140\*120\*121mm **Gross Weight:** 290±10g



### Master Box (36 color boxes/ Master box)

Size: 585\*375\*385mm

Gross Weight: 14446 ± 500g



### Mask

**Size:**230\*120mm **Weight:**6.8±0.3g



Verify the validity with the QR code



**NB 2163** 

### **EU TYPE EXAMINATION CERTIFICATE**

Certificate No: 2163-PPE-884

Respiratory protective devices, filtering half masks to protect against particles manufactured by

Guangdong Kingfa Sci.&Tech. Co., Ltd.

28 Delong Avenue, Shijiao Town, Qingcheng District, Qingyuan City, Guangdong Province, China

are tested and evaluated according to

### EN 149:2001 + A1:2009 Respiratory Protective Devices -Filtering Half Masks to Protect Against Particles -Requirements, Testing, Marking

Based on the type examination conducted with the evaluation of test reports, technical file according to Personal Protective Equipment Regulation (EU) 2016/425 Annex 5, it is approved that the product meets the requirements of the regulation.

#### **Product Definition**

Brand Name: KINGFA Model: KF-A F10(SC) Filtering half mask

Classification: FFP2 NR

Here by the manufacturer is allowed to use notified body number (2163) and can fix CE mark, as shown below, on the Category III product models given above, with;

- Issuing an appropriate EU Declaration of Conformity according to Personal Protective Equipment Regulation (EU) 2016/425 Annex 9.
- Ongoing successful performance in fulfilment of the requirements set out in Personal Protective Equipment Regulation (EU) 2016/425 and harmonised standards, ensured by assessments based on Annex 7 (Module C2) or Annex 8 (Module D) of the regulation no later than 1 year from the beginning of serial production

This certificate is initially issued on 29/06/2020 and will be valid for 5 years, if there is no change in the relevant harmonised standard affecting the essential health and safety requirements.



Suat KAÇMAZ
UNIVERSAL CERTIFICATION
Director



Finnish Accreditation Service S003 (EN ISO/IEC 17065)

Certificate CN20/42082

The management system of

# Guangdong KINGFA SCI. &TECH. Co., Ltd.

No.28, Delong Avenue, Shijiao Town, Qingcheng District, Qingyuan City, Guangdong Province, 511545, P.R. China

has been assessed and certified as meeting the requirements of

### Regulation (EU) 2016/425

Module D

For the following activities

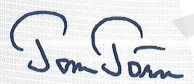
Manufacture of FFP1/FFP2 Protective Respirator (Note: all products marked CE0598 must have a valid EU Type Examination Certificates issued under Module B or a valid EC type examination certificate issued under Article 10 of the PPE Directive 89/686/EEC.)

This certificate is valid from 10 June 2020 until 9 June 2023 and remains valid subject to satisfactory surveillance audits.

Re certification audit due before 26 May 2023

Issue 1. Certified since 10 June 2020

Authorised by

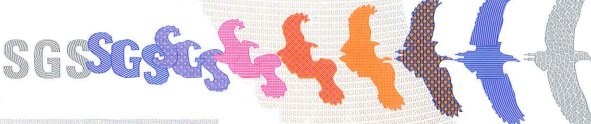


SGS FIMKO OY, Notified Body 0598

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### Certificate

The Certification Body of TÜV Rheinland LGA Products GmbH

hereby certifies that the organization

GuangDong Kingfa Science and Technology Co., Ltd. No.28, Delong Road, Qingcheng Dist. Qingyuan City 511545 Guangdong P.R. China

has established and applies a quality management system for medical devices for the following scope:

Design and Development, Manufacture and Distribution of Disposable Medical Face Masks (non-sterile)

Proof has been furnished that the requirements specified in

EN ISO 13485:2016

are fulfilled. The quality management system is subject to yearly surveillance.

Effective Date:

2020-07-13

Certificate Registration No.:

SX 60150441 0001

An audit was performed. Report No.: 17054679 002

This Certificate is valid until:

2023-07-12

Certification Body



Date 2020-07-13



TÜV Rheinland LGA Products GmbH - Tillystraße 2 - 90431 Nürnberg

Tel.: +49 221 806-1371 Fax: +49 221 806-3935 e-mail:cert-validity@de.tuv.com http://www.tuv.com/safety

# Certificate

Standard

ISO 9001:2015

Certificate Registr. No.

01 100 1430282

Certificate Holder:

GuangDong Kingfa Science and Technology Co., Ltd.

Unified Social Credit Code: 91441802077867032A

Registration Address: No. 28, Delong Road, Qingcheng Dist. Shijiao Town, Qingyuan City, 511545 Guangdong, P. R. China

Operation Address: same as above

Scope:

Design and Manufacturing of Modified Plastics;

Design and Manufacturing of Masks and Non-Powered Air-

Purifying Particle Respirator

Proof has been furnished by means of an audit that the

requirements of ISO 9001:2015 are met.

Validity:

The certificate is valid from 2020-07-19 until 2023-07-18. It remains valid subject to satisfactory surveillance audits.

First certification 2014

This certificate information can be searched on CNCA official

website http://www.cnca.gov.cn

2020-06-08

TÜV Rheinland Cert GmbH Am Grauen Stein · 51105 Köln







### FDA EUA



https://www.fda.gov/medical-devices/coronavirus-disease-2019-covid-19-emergency-use-authorizations-medical-devices/persona l-protective-equipment-euas#imported

### Appendix A: Authorized Imported, Non-NIOSH Approved Respirators Manufactured in China (Updated: July 23, 2020)

The table below includes a list of non-NIOSH respirators authorized by this Umbrella EUA for emergency use during the COVID-19 public health emergency.

As stated in the EUA, authorized respirators should be used in accordance with CDC's recommendations. For the most current CDC recommendations on optimizing respirator use, please visit CDC's webpage: Strategies for Optimizing the Supply of N95 Respirators.

	Search:	
<b>+</b>	Guangdong KINGFA SCI. & TECH. Co. Ltd.	KF-A F01, KF-A F10(SC)



June 16, 2020

GUANGDONG KINGFA SCI. & TECH. CO. LTD. 28 DELONG AVENUE, SHIJIAO TOWN QINGCHENG DISTRICT QINGYUAN CITY CN - CHINA

EUA201196

Re: FFRs Made in China

Dear David Wu:

This letter is in response to your request that the Food and Drug Administration (FDA) add your respirator model KF-A-F01 as an authorized respirator to the May 7, 2020 Emergency Use Authorization (EUA)<sup>1</sup>, which was issued under Section 564 of the Federal Food, Drug, and Cosmetic Act (the Act) (21 U.S.C. §360bbb-3). We have reviewed your email and determined that the models included meet the eligibility criteria in the May 7, 2020 EUA for non-NIOSH approved respirators made in China. As such, your respirator(s) is hereby added to Appendix A<sup>2</sup> as an authorized respirator.

Having concluded that the eligibility criteria are met, I am adding your respirators to Appendix A, as described in the Scope of Authorization (Section II). As such, the respirator is authorized for use by healthcare personnel in healthcare settings in accordance with CDC recommendations and subject to the Conditions of Authorization (Section IV) of the attached letter. We remind you that, among other things, you are required to meet the following labeling requirements:

#### Manufacturers

- A. Manufacturers of authorized respirators are required to publish the intended use and other instructions (such as fit testing, etc.) about all authorized models that are imported and authorized under this EUA on their website in English. Additionally, manufacturers must notify FDA by emailing FDA at <a href="mailto:CDRH-NonDiagnosticEUA-Templates@fda.hhs.gov">CDRH-NonDiagnosticEUA-Templates@fda.hhs.gov</a> of the website address (URL) that meets this condition. The subject line of this email should read "URL for FFR Made in China." FDA will make this information available to the public on its EUA website at <a href="https://www.fda.gov/medical-devices/emergency-situations-medical-devices/emergency-use-authorizations#covid19ppe">https://www.fda.gov/medical-devices/emergency-use-authorizations#covid19ppe</a>. Manufacturers must notify FDA of any changes to this page.
- B. In addition to the above electronic labeling condition, manufacturers of authorized respirators are additionally required to include a letter, in English, that can be distributed to each end user facility (e.g., each hospital, etc.) that receives the authorized respirator model. This letter must include the

<sup>&</sup>lt;sup>1</sup> The EUA Letter of Authorization is available at, https://www.fda.gov/media/136664/download.

<sup>&</sup>lt;sup>2</sup> Appendix A is available at, <a href="https://www.fda.gov/medical-devices/emergency-situations-medical-devices/emergency-use-authorizations">https://www.fda.gov/medical-devices/emergency-use-authorizations</a>.



authorized respirator's manufacturer, model, intended use, manufacturer's webpage (if applicable), etc.

Additionally, please be advised that if your firm does not have the appropriate fluid resistance testing, the respirator should not be labeled as "surgical."

Import information can be found on the <u>Information for Filing Personal Protective Equipment and Medical Devices</u> <u>During COVID-19 page</u>. If you need to resolve entry issues for shipments, please contact 301-796-0356 or <u>COVID19FDAIMPORTINQUIRIES@fda.hhs.gov</u>.

Sincerely,

Suzanne Schwartz, MD, MBA
Deputy Director (& Acting Office Director)
Office of Strategic Partnerships & Technology Innovation
Center for Devices and Radiological Health



#### **EU DECLARATION OF CONFORMITY**

This declaration of conformity is issued under the sole responsibility of the manufacturer:

Manufacturer and	GUANGDONG KINGFA SCI.&TECH. CO., LTD.
address	NO.28 Delong Avenue, Shijiao Town, Qingcheng District, Qingyuan City, Guangdong Province, China
Product name	Particle Filtering half mask
Model/ Serial No.	KF-A F10(SC) FFP2 NR
Applicable Regulation:	PPE Regulation 2016/425
Notified body for EU type-examination (Module B)	UNIVERSAL- NB 2163  Necip Fazıl Bulvarı Keyap Sitesi E2 Blok No:44/84 Yukarı Dudullu Ümraniye /  İSTANBUL / TÜRKİYE
Certificate number (Module B)	2163-PPE-884
Notified body for EU type-examination (Module D)	SGS FIMKO OY - NB 0598  Takomotie 8, FI-00380 Helsinki, Finland
Certificate number(Module D)	Certificate CN20/42082

We declared that given information on the above statement and attached documents/records are true and correct to the best of our knowledge.

Signed for and on behalf of:GUANGDONG KINGFA SCI.&TECH. CO., LTD.

(date of signature):2020-6-30

(title of signatory):General Manager

(signature):









(2020) WSZ FHL NO.W0708

Product Name _	Particle Filtering half mask
Client _	Guangdong KINGFA SCI.&TECH.Co.,Ltd.
Manufacturer _	
Test Type _	Entrusted inspection

Jiangsu Guojian Testing Technology Co., Ltd. 检验专用章



[2020] WSZ FHL NO.W0708

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D 1 4	Deuti-1- Eliterin - 1-161-	Specification	KF-A F10(SC)
Product name	Particle Filtering half mask	Brand	-
Client/Add/Tel	ue, Shijiao Town, Qingchen		
Manufacturer/ Add/Tel	_/_/_		
Sample grade	FFP2	Sample number	GWW0708-2020
Sample quantity	110 pcs	Receiving date of sample	21/05/2020
Test type	Entrusted inspection	Article number/Batch number/Style number	
Test date	23/05/2020~29/05/2020	Testing sites	Testing room
Sample state	Meeting the requirements of testing	Sample description	_
Test standard(s)	EN 149:2001+A1:2009 Respiratory particles- Requirements, testing, marking		g half masks to protect against
Test items	Visual inspection, practical performance carbon dioxide content of the inhalation filter material, breathing resistance, to	on air, material, head harnes	
Test conclusion	The sample upon testing, the test ite standard. The detail of test results see		多国健检测技术有强
Note	For the entrusted sample test, the tech undertaken for the test results of the		

Reviewer: 7/2

Chief Tester:

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S. No.	Test	item	Unit	Technical requirements		Test result	Single item decision	
	Visual	Packaging	<u></u>	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.	Packaging withstands mechanica damage and contamination.		0.10	
1	inspection	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.	Materials		Qualified	
		Head harness comfort		Head harness should be comfort.	Sample 1 has the feeling of comfortable wearing  Sample 2 has the feeling of comfortable wearing			
2	Practical performance	Security of fastenings	_	Fastenings are safe and reliable		: All fastenings are firm.	Qualified	
		Field of vision	_	Field of vision is acceptable	field	2: Having a wider visual		
3	Finish	of parts		Parts of the device likely to come into contact with the wearer shall have no sharp edges or burrs.	Parts of t	the device have no	Qualified	
4	Compatibil	ity with skin		Materials that may come into contact with the wearer's skin shall not be	A.R.	5 pcs all don't cause irritation	Qualified	
				known to be likely to cause irritation or any other adverse effect to health.	T.C.	5 pcs all don't cause irritation	Quanned	
5 Flammability		mask shall		When tested, the particle filtering half mask shall not burn or not to	A.R.	The Sample is burning. Burning time:0.1s The Sample is burning. Burning time:0.1s		
			continue to burn for more than 5s after removal from the flame.	T.C.	The Sample is burning. Burning time:0.1s The Sample is burning. Burning time:0.1s	Qualified		

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S.No.	Test	item	Unit	Technical requirements		Test result		Single item decision			
6	Carbon diox of the inha	and a second of	<u>}</u> _	≤1.0% (by volume)	Samp Samp Samp	le 2 le 3	0.596 0.604 0.602 0.60	0% 5%	Qualified		
7	Material		Material			After undergoing S.W., none of the particle filtering half masks shall have suffered mechanical failure of the facepiece or straps.	Sample 1: have mech Sample 2: have mech Sample 3: have mech	nanical fai neither fa nanical fai neither fa	lure acepiece n lure acepiece n	or straps	
				After undergoing S.W. and T.C., none of the particle filtering half masks shall not collapse.	Sa	ample 1: n	no collaps	e			
0				The head harness shall be designed so that the particle filtering half mask can be donned and removed easily.  The head harness shall be	A.R.		5 pieces half mas	-	Qualified		
8	Head n	Head harness		adjustab shall be hold the		adjustable or self-adjusting and shall be sufficiently robust to hold the particle filtering half mask firmly in position	T.C.		5 pieces half mas		Quanned
9	Field of	f vision		The field of vision is acceptable if determined so in practical performance tests.	The two samples both have a wider visual field		Qualified				
		Sodium		≤6%	A.R.	0.1%	0.1%	0.1%	Qualified		
10	Penetration of filter				M.S+T.C.	0.1%	0.2%	0.1%			
10	material	Paraffin		≤6%	A.R. S.W.	0.2%	0.2%	0.3%	Qualified		
		oil		×070	M.S+T.C.	0.276	0.5%	0.4%	Quanned		

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		Z THE TWO				7	Test	result		^							
S.No.	Test item		Unit	Technical requirements	Exercises	Facing directly ahead	Facing vertically upwards	Facing vertically downwards	Lying on the left side	Lying on the right side	Single item decision						
/ ,		5000				0.2	0.2	0.3	0.2	0.3							
					A.R.	0.2	0.3	0.2	0.3	0.2							
						0.3	0.2	0.3	0.2	0.3							
						0.3	0.2	0.3	0.2	0.3							
		Inhalation 30 L/min		≤0.7	S.W.	0.3	0.3	0.2	0.3	0.2	Qualified						
		30 L/IIIII				0.2	0.3	0.3	0.3	0.3							
			1			0.3	0.3	0.2	0.3	0.3							
					T.C.	0.2	0.3	0.3	0.2	0.3							
						0.3	0.2	0.3	0.3	0.2							
												1.2	1.3	1.3	1.3	1.2	
					A.R.	1.3	1.2	1.3	1.2	1.3							
						1.3	1.3	1.2	1.3	1.3							
						1.3	1.2	1.2	1.3	1.3							
11	Breathing	Inhalation 95 L/min	mbar	≤2.4	S.W.	1.2	1.3	1.3	1.2	1.3	Qualified						
	resistance	93 L/IIIII				1.3	1.3	1.3	1.3	1.2							
	/ /					1.3	1.2	1.3	1.2	1.3							
					T.C.	1.3	1.3	1.2	1.3	1.3							
	1					1.2	1.3	1.3	1.3	1.2							
						1.8	1.9	1.9	2.0	1.9							
		7			A.R.	1.8	1.8	2.0	1.9	1.8							
				1		1.9	1.9	1.9	1.9	1.9	7 /						
		/				1.9	1.9	1.9	1.9	1.9							
		Exhalation 160 L/min	120	≤3.0	S.W.	1.9	1.8	2.0	1.8	1.8	Qualified						
		100 L/IIIIn			200	1.8	1.9	1.9	1.9	1.9							
		1				1.9	1.8	2.0	1.9	1.9							
					T.C.	1.8	1.9	1.9	2.0	1.9							
					13,77	1.9	1.9	1.9	2.0	2.0							

Page 5 of 6 [2020] WSZ FHL NO.W0708 Single item **Technical** Test S.No. Unit Test result decision requirements item E2 E3 E5 TIL E1 E4 Exercises (%) (%) (%) (%) (%) (%) 1# 1.1 1.7 1.5 1.5 1.1 1.4 2# 1.4 2.2 2.0 2.3 1.6 1.9 At least 46 out of the 50 individual A.R. 3# 0.8 1.2 1.2 1.2 0.8 1.0 exercise results shall be not greater 4# 0.9 1.2 0.7 1.2 1.6 1.6 Total than 11%; inward And in addition, at Qualified 12 5# 1.0 1.7 1.7 2.0 1.3 1.5 least 8 out of the 10 leakage individual wearer 6# 1.0 1.9 1.7 1.7 1.3 1.5 arithmetic means for the total inward 7# 1.9 1.4 1.9 2.4 2.0 1.6 leakage shall be not greater than 8%. T.C. 0.7 0.6 1.3 1.4 1.3 1.1 0.6 1.4 1.5 0.7 1.1 1.2 10# 1.7 1.4 1.6 1.7 1.1 1.1

Thoond	and the same
 The end	

Note

### SUPPLEMENTARY TEST REPORT

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### Facial dimensions of ten test subjects:

Subject	Face Length	Face Width	Face Depth	Mouth Width
	(mm)	(mm)	(mm)	(mm)
1	120	130	109	59
2	122	140	115	65
3	119	160	139	55
4	112	122	119	63
5	110	130	118	60
6	115	119	110	59
7	112	123	113	55
8	103	130	100	50
9	118	139	130	63
10	115	129	120	50

The end —





### **Supplier Creditability & Capacity Audit Report**

Report:							
Supplier Name	Guangdong KINGFA SCI.&TEC	Guangdong KINGFA SCI.&TECH. Co., Ltd. 广东金发科技有限公司					
Supplier Address	No. 28, Delong Avenue, Shijiao Guangdong Province, China	No. 28, Delong Avenue, Shijiao Town, Qingcheng District, Qingyuan City, Guangdong Province, China					
Client Information	/						
Name of Assessor	James Lee	Reviewed by	Roger Wang				
Audited Date	04 May, 2020	Expiry Date	03 May, 2021				

#### **Assessment Scope:**

Section 1: Company Profile

Section 2: Personnel

Section 3: Main Market

Section 4: Manufacturing Ability

Section 5: Certificate

Section 6: Quality Control Management

Section 7: Development Plan

Section 8: Production Flow Chart

Section 9: Attachment

#### **Comments**

Guangdong KINGFA SCI.&TECH. Co., Ltd. is a trader and manufacturer combined company with 2097 employees; it was established in 2013, located in No. 28, Delong Avenue, Shijiao Town, Qingcheng District, Qingyuan City, Guangdong Province, China. They have passed ISO9001, ISO14001, OHSAS18001 certifications in 2017. Guangdong KINGFA SCI.&TECH. Co., Ltd. has successful foreign trading experience in Europe, North America and East Asia.

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### **Section 2: Personnel**

2.1 Company Org Chart					
		总经理			
		<b>管理者代</b>	表		
1					
生产部	品質部 技术部 部		销售 新 售 新		
2.2 Headcount and Key S					
According to	[ ] Attendance rec [ ] On-site observa		rs list		
	Department	Full time	Part time	Total	
	GM	1	0	1	
	Management Represents	1	0	1	
	Production Dept.	1666	0	1666	
	QC Dept.	80	0	80	
	Technology Dept.	20	0	20	
Headcount	Warehouse Dept.	220	0	220	
	Purchase Dept.	15	0	15	
	HR Dept.	15	0	15	
	Office	48	0	48	
	Marketing Dept.	24	0	24	
	Fin. Dept.	7	0	7	
	Total		2097		
	Full Name	Position	Working experience in this filed		
Key Staff	Mr. Hongtao Ning	General Manager	About 20 years working experience		
Ney Stall	Mr. Xiaojun Deng	Factory Director	About 15 years	working experience	
	Mr. Min Ding	Export Manager	8 years foreign	trading experience	
raining Procedure and Plan or Staff	[ X ] All staff [ ] No Training reco	[ ] Key station	on		
are there uniforms for all taff in company?	There are uniforms for				

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(86-21) 61156703 1 (86-21) 51152274 1 (86-21) 61156703 1 (86-21) 61152274

www.cn.ags.com e as ch@sgs.com



### **Section 3: Main Market**

#### 3.1 Foreign Trading Staff

There were 24 foreign trading members in the company.

Education Level	Headcount
Doctor	0
Master	19
University	5
Junior college	0
Technical secondary school	0

Working Experience	Headcount
Over 20 Years	0
Over 10 Years	12
Over 5 Years	12
2-5 Years	0
1Year	0

English Level	Headcount
TEM-8	0
CET-6	24
CET-4	0
CET-3	0
PETS-3	0

**Export means:** [X] Directly export through own export right

Export business operated by other foreign trading company

#### 3.2 Export Information

3.2 Export Information				
Item		Content		
	Area	% of Total Business Volume (last year)		
	North America	23.5		
	South America	0.12		
	West Europe	6.5		
	East Europe	0		
Maio Mantos	East Asia (Japanese/ Korea)	58		
Main Market	Africa	0		
	Australia	6.9		
	Southeast Asia	3		
	Mideast	0		
	Others	1.98		
	Domestic	0		
	Annual volume in last year	Confidential		
Sales Volume	Export volume in last year	Confidential		
	Estimated export in this year	Confidential		
Key Client	Confidential	Confidential		
Lead time	From PO Confirmation to Ex works	7-15 days		

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### **Section 4: Manufacturing Ability**

4.1 Main Facilities					
	Facility name	Brand/Model	Quantity	Year made	Condition
Please list the major machinery / utilities	Medical Mask Production Line 医用口罩生产线	Guoji	132	2020	Good
on site.	Protective Mask Production Line 防护口罩生产线	Kuaiyuda	80	2020	Good

4.2 Main Test Instruments						
	Facility name	Brand/Model	Quantity	Year made	Condition	
	Mask BFE Tester 口罩细菌过滤效率检 测仪	ZR-1000	1	2020	Good	
Please list the major	Mask Tensile Strength Tester 口罩拉力机	KT22	1	2020	Good	
test instruments on site.	Clean Bench 超净工作台	YJ-840	1	2020	Good	
	Mildew Incubator 霉菌培养箱	MJ-80	1	2020	Good	
	Constant Temperature Incubator 恒温培养箱	DHP-9082	1	2020	Good	

4.3 Output			
Output in last war	Product	Monthly output	Yearly output
Output in last year	N/A	N/A	N/A
Output in this year	Protective Mask/ Medical Mask (Non-sterile) 防护口罩/医用口罩 (非灭菌)	300,000,000 Pcs	N/A

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### **Section 5: Certificate**

5.1 Management System Certificate						
Certificate	Number	Expiry date	Certifying Body	Scope		
ISO 9001:2015	01 100 1430282	31 Oct., 2017	TUV Rheinland	Design and production of modified plastics		
ISO14001:2004	01 104 1430282	18 Jul., 2020	TUV Rheinland	Design and production of modified plastics		
OHSAS 18001:2007	01 113 1430282	18 Jul., 2020	TUV Rheinland	Design and production of modified plastics		

5.2 Product Certificate						
Certificate	Number	Issued date	Certifying Body	Product and model / type		
Test Report	20R000099 MT	23 Apr. 2020	GTT	Disposable medical mask(non-sterile) Standard EN14693:2019+ac:2019		
Test Report	(2020) WSZ FHL No. 2852	27 Mar., 2020	Jiangsu Guojian Testing Technology Co., Ltd.	Labor Protective Mask Standard: GB2626-2006		
FDA Registration	10065634	2020	FDA	Disposable Protective Mask Model: Adult; Protective Mask Model: KF-A(Adult)		

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**Section 6: Quality Control Management** 

lt a cor	Comtont	Grading			Observations (2	
Item	Content	Poor	Mid	Good	Observations /Comments	
6.1	Are the environmental conditions such as tidiness and cleanliness being controlled and suitable for the operation performed?			√	Refer to site observation; the environmental condition was suitable for the operation performed.	
6.2	Are the following items /documents provided at appropriate location and under control when necessary?  - Work Instructions /procedures  - Workmanship standard /acceptance  - Golden sample			V	Refer to site observation; there were documented work instructions, workmanship standard provided in the workshops.	
6.3	Does the company establish and implement an effective suppliers/ sub-contractors assessment procedure (which covers the acceptable criteria of supplier/ sub-contractor)?	X		<b>√</b>	The company had established this procedure for supplier assessment, latest record has been reviewed.	
6.4	Are written instructions available for incoming material inspections /testing? Is the relevant records maintained?			√	Refer to on-site observation; there were documented instructions for incoming material inspection. And inspection records were maintained well.	
6.5	Are written inspections /testing instructions available for finished products? Is the relevant records maintained?			1	The company had established the procedure for this inspection. And records were maintained well.	
6.6	Is there a procedure to conduct random product inspection after final packaging in place?			-1	All inspection procedures were implemented before packaging.	
6.7	Are non-conforming units clearly marked/ segregated to prevent accidental dispatch?			<b>V</b>	Refer to site observation; non-conforming units would be marked with label and placed in the non-conforming parts area	
6.8	Is there a clear procedure for handling customer complaint?				Refer to relevant documentation; the company had a clear procedure for handling customer complaint.	
6.9	Can the finished/packaged product be traced by lot identification to the appropriate raw materials test reports?			√	Auditor noted that the company had established this procedure for lot identification.	
6.10	Are corrective & preventive actions mechanism established and implemented effectively (including the suppliers/ sub-contractors' control, incoming inspection, process control, final inspection and customer complaint)?			<b>√</b>	The company had documented procedure for corrective & preventive actions mechanism and records were kept well.	

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### **Section 7: Development Plan**

7.1				
Item	Actions	Time Frame		
1	Enlarge the mask production capacity	2020		

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### **Section 8: Production Flow Chart**

#### 8.2 Product: Solar Module



1. Medical Mask (Non-sterile) Production 医用口罩(非灭菌)生产

2. Protective Mask Production 防护口罩生产



3. Lab. Testing 实验室检验



4.Packing 包装



5. Store 成品储存

N/A

N/A

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### **Section 9: Attachment**



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#### 9.2 Photos of Company and Product Sample

#### **Company Gate**

#### Office Building





Office

Lab.





**Testing Machine** 

**Testing Machine** 





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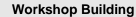
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#### **Workshop Building**







Workshop

Workshop





**Automatic Production line** 

**Automatic Production line** 





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