



CERTIFICATION OF FDA REGISTRATION

This certifies that:

has completed the FDA Establishment Registration and Device Listing with the US Food & Drug Administration, through

Shenzhen CCT Testing Technology Co., Ltd.

Owner/Operator Number: 10062984



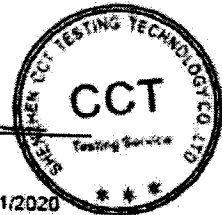
Device Listing#: See Next pager

CCT will confirm that such registration remains effective upon request and presentation of this certificate until the end of the calendar year stated above, unless said registration is terminated after issuance of this certificate. CCT makes no other representations or warranties, nor does this certificate make any representations or warranties to any person or entity other than the named certificate holder, for whose sole benefit it is issued. This certificate does not denote endorsement or approval of the certificate-holder's device or establishment by the U.S. Food and Drug Administration. CCT assumes no liability to any person or entity in connection with the foregoing.

Pursuant to 21 CFR 807.39, "Registration of a device establishment or assignment of a registration number does not in any way denote approval of the establishment or its products. Any representation that creates an impression of official approval because of registration or possession of a registration number is misleading and constitutes misbranding." The U.S. Food and Drug Administration does not issue a certificate of registration, nor does the U.S. Food and Drug Administration recognize a certificate of registration, CCT is not affiliated with the U.S. Food and Drug Administration.

Shenzhen CCT Testing Technology Co., Ltd.
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Stacy
Chief engineer
Issued: 03/15/2020
Expiration Date: 12/31/2020



Web: <http://www.fda.gov> Tel: 1-866-INFO-FDA (1-888-453-6332) e-mail: webmail@oc.fda.gov



CERTIFICATE OF CONFORMITY

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Certificate No.: CCT20030503QCS

Applicant.....:

Manufacturer.....:

Product.....: **Mask**
 Trade Name.....: **N/A**
 Model(s).....: **85mm×95mm, 175mm×95mm, 175mm×90mm, 170mm×90mm, 145mm×95mm, 145mm×90mm, 125mm×95mm, KN95, N95**

It is only valid in connection with the test report number CCT20030503QRS Identification of regulation/standards

EN 149: 2001+A1:2009

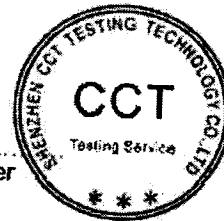
The EUT described above has been tested by us with the listed standards and found in compliance with the council PPE Directive 2016/425/EU. It is possible to use CE marking to demonstrate the compliance with this PPE Directive.

This certificate of conformity is based on a single evaluation of the submitted sample(s) of the above mentioned product. It does not imply an assessment of the whole production and other relevant directives have to be observed.

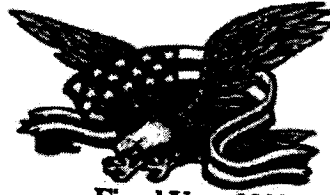


Toby

 Toby/Senior Manager
 Mar. 10, 2020



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Fiscal Year 2020

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Device Listing#:

Listing No	Code	Device Name	Proprietary Name
0375030	MDM	INSTRUMENT, MANUAL, SURGICAL, GENERAL USE	Face Mask
0375029	OEA	Non-surgical isolation gown	Protective clothing
0375028	HGY	Sunglasses (non-prescription including photosensitive)	Goggles
0375027	KHA	MASK, SCAVENGING	Mask

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Stany
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