

June 16, 2022

Thanh Cong Pharmaceutical and Trading Company Limited % Manoj Zacharias
US Agent
Liberty Management Group Limited
75 Executive Drive, Suite 114
Aurora, Illinois 60504

Re: K221186

Trade/Device Name: TC Gloves - Powder Free Nitrile Examination Gloves

Regulation Number: 21 CFR 880.6250

Regulation Name: Non-Powdered Patient Examination Glove

Regulatory Class: Class I, reserved

Product Code: LZA Dated: March 29, 2022 Received: April 25, 2022

Dear Manoj Zacharias:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. Although this letter refers to your product as a device, please be aware that some cleared products may instead be combination products. The 510(k) Premarket Notification Database located at https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpmn/pmn.cfm identifies combination product submissions. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the <u>Federal Register</u>.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal

statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803) for devices or postmarketing safety reporting (21 CFR 4, Subpart B) for combination products (see https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820) for devices or current good manufacturing practices (21 CFR 4, Subpart A) for combination products; and, if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to https://www.fda.gov/medical-device-problems.

For comprehensive regulatory information about medical devices and radiation-emitting products, including information about labeling regulations, please see Device Advice (https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance) and CDRH Learn (https://www.fda.gov/training-and-continuing-education/cdrh-learn). Additionally, you may contact the Division of Industry and Consumer Education (DICE) to ask a question about a specific regulatory topic. See the DICE website (https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice">https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice) for more information or contact DICE by email (DICE@fda.hhs.gov) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

Bifeng Qian, M.D., Ph.D.
Acting Assistant Director
DHT4B: Division of Infection Control
and Plastic Surgery Devices
OHT4: Office of Surgical
and Infection Control Devices
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure

DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration

Indications for Use

510(k) Number (if known)

Form Approved: OMB No. 0910-0120 Expiration Date: 06/30/2023

Expiration Date: 06/30/2023 See PRA Statement below.

K221186	
Device Name TC Gloves- Powder Free Nitrile Examination Gloves	
Indications for Use (Describe)	
TC Gloves- Powder Free Nitrile Examination Gloves is a disposable the examiner's hand to prevent contamination between patient and	
Type of Use (Select one or both, as applicable)	
Prescription Use (Part 21 CFR 801 Subpart D)	☑ Over-The-Counter Use (21 CFR 801 Subpart C)

This section applies only to requirements of the Paperwork Reduction Act of 1995.

CONTINUE ON A SEPARATE PAGE IF NEEDED.

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510(k) SUMMARY (K221186)

[AS REQUIRED BY 21CFR807.92]

I. SUBMITTER DETAILS

510(k) Owner's Name	THANH CONG PHARMACEUTICAL AND TRADING COMPANY LIMITED	
Address	KM 6+ 200 National Road No.38, Hap Linh, Bac Ninh City, Bac Ninh province, Vietnam	
Contact person	Ms. Nguyễn Thị Tú Anh	
Contact Designation	Director of Production	
Contact Phone Number	098 171 2992	
Contact Email	tuanhnguyentcpharma@gmail.com	
Date of Summary Prepared	15-06-2022	

II. DEVICE DETAILS

Brand Name	TC Gloves
Device Common Name Powder Free Nitrile Examination Gloves	
Device Classification name	Non-powdered patient examination glove
Regulation Number	21 CFR 880.6250
Class	I
Product Code	LZA

III. PREDICATE DEVICE DETAILS

Predicate Device Name	Jr Medic Blue Nitrile Examination Gloves Powder Free
510(k) Number	K192333
Regulation Number	21 CFR 880.6250
Class	I
Product Code	LZA



IV. DEVICE DESCRIPTION

TC Gloves- Powder Free Nitrile Examination Gloves is a Class I device bearing the product code LZA (21CFR 880.6250). They meet all the current specifications listed under the ASTM D6319 -19, Standard Specification for Nitrile Examination Gloves for Medical Application. They are made from acrylonitrile-butadiene copolymer dispersion. These gloves are blue in color having Finger Texture, Ambidextrous, Single use and are powder free. The product is non-sterile. Sizes available - S, M, L and XL.

V. INDICATION FOR USE

TC Gloves- Powder Free Nitrile Examination Gloves is a disposable device intended for medical purpose that is worn on the examiner's hand to prevent contamination between patient and examiner.

VI. COMPARISON OF TECHNOLOGICAL CHARACTERISTICS WITH THE PREDICATE DEVICE

Table 1: General Comparison

SI. No	Features compared	Proposed Device	Predicate Device	Result				
	General Information							
1.	510(k) Number	K221186	K192333	-				
2.	Manufacturer	Thanh Cong Pharmaceutical And Trading Company Limited	JR Engineering & Medical Technologies	-				
3.	Classification	I	I	Same				
4.	Regulation number	21 CFR 880.6250	21 CFR 880.6250	Same				
5.	Product Code	LZA	LZA	Same				
6.	Indication For Use	TC Gloves-Powder Free Nitrile Examination Gloves is a disposable device intended for medical purpose that is worn on the examiner's hand to prevent contamination between patient and examiner.	Jr Medic Blue Nitrile Examination Gloves Powder Free are intended for medical purposes that is worn on the examiner's hands to prevent contamination between patient and examiner.	Same				
7.	Material	Nitrile	Nitrile	Same				
8.	Color	Blue	Blue	Same				
9.	Texture	Finger Texture	Finger texture	Same				
10.	Ambidextrous	Yes	Yes	Same				
11.	Size	S, M, L, XL	XS, S, M, L, XL	Similar-1				



SI. No	Features compared		Proposed Device	Predicate Device	Result		
12.	OTC Use		OTC Use		Yes	Yes	Same
13.	Reusability		Single use	Single use	Same		
14.	Steril	lity	Non- sterile	Non- sterile	Same		
15.	Shelf	Life	3 years	Data Not available	-		
16.	Dime	nsions	Length Min 230 mm Width Min 95+/-10 mm (for medium size)	Length Min 230 mm Width Min 95±10 mm (for medium size)	Same		
17.	Thick	ness	Palm min 0.05 mm Finger min 0.05 mm	Palm min 0.05 mm Finger min 0.05 mm	Same		
18.	. Physical Properties		Before Aging Tensile Strength min 14 Mpa Ultimate Elongation Min 500% After Aging Tensile Strength min 14 Mpa Ultimate Elongation Min 400%	Before Aging Tensile Strength min 14 Mpa Ultimate Elongation Min 500% After Aging Tensile Strength min 14 Mpa Ultimate Elongation Min 400%	Same		
19.	Detection of Holes		Passes AQL 2.5	Passes AQL 1.5	Different-1		
20.	Powder Free Residue		≤2 mg/glove	≤2 mg/glove	Same		
	In Vitro Cytotoxicity		Under the conditions of the study, cytotoxic to L-929 cells. Additional Testing was performed to determine if this was a systemic toxicity concern.	Under the conditions of the study, cytotoxic to L-929 cells. Additional Testing was performed to determine if this was a systemic toxicity concern.	Same		
	Study	Study	Study	Skin Sensitization	Under the conditions of the study not a sensitizer	Under the conditions of the study not a sensitizer	Same
21.	Biocompatibility Study	Skin Irritation	Under the condition of study not an irritant	Under the condition of study not an irritant	Same		
	Biocom	Material mediated pyrogenicity	Under the conditions of the study, the device did not demonstrate a material mediated Pyrogenicity response.	Under the conditions of the study, the device did not demonstrate a material mediated Pyrogenicity response.	Same		
		Acute systemic toxicity	Under the condition of study, the device extracts do not pose a systemic toxicity.	Under the condition of study, the device extracts do not pose a systemic toxicity.	Same		



There are no significant differences between the two products and are identical in terms of intended use, materials, design, manufacturing methods. Both devices meet the ASTM standard D6319-19.

VII. PERFORMANCE DATA

A. Non- Clinical Data

Performance Tests

TC Gloves- Powder Free Nitrile Examination Gloves is subjected to the following performance tests according to the requirements of Guidance for Industry and FDA Staff - Medical Glove Guidance Manual and found to be safe and efficient with respect to its intended use:

- Dimension
- Physical property
- Barrier property tests- Detection of Holes in Medical Gloves
- Powder Free Residue

Table 2: Performance Testing Summary

SI No.	Tests	Prop	osed Device	actual Data	Acceptance Criteria		
		Size	Length	Width	Size	Length	Width
		S	241.4 mm	86.6 mm	S		80 mm±10
	<u>Dimension</u>	М	240.6 mm	96.5 mm	М	220	95 mm ±10
	Length, Width and	L	243.2 mm	106.4 mm	L	230mm min	110 mm ±10
	Thickness	XL	242.7 mm	113.8 mm	XL		120 mm ±10
1.	ASTM D6319-19 Standard		Thicknes	SS		Thickne	ss
	Specification for	Size	Palm	Finger	Size	Palm	Finger
	Nitrile Examination Gloves for Medical	S	0.14 mm	0.16 mm	S		0.05 mm min
	Application	М	0.14 mm	0.15 mm	М	0.05 mm min	
		L	0.14 mm	0.15 mm	L	0.03 11111 111111	0.03 11111 111111
		XL	0.13 mm	0.16 mm	XL		1
			Tensile stre	ngth	Tensile strength		
		Size	Before ageing	After ageing	Size	Before ageing	After ageing
	Physical property	S	15.7 MPa	15.0 MPa	S		14 MPa Min for all sizes
	Tensile strength	М	15.5 MPa	15.3 MPa	М	14 MPa Min for all	
	and Ultimate	L	15.4 MPa	14.9 MPa	L	sizes	
2.	Elongation	XL	15.2 MPa	15.3 MPa	XL		
۷.	ASTM D6319-19 Standard		Ultimate elon	gation		Ultimate eloi	ngation
	Specification for	Size	Before ageing	After ageing	Size	Before ageing	After ageing
	Nitrile Examination Gloves for Medical	S	515.8%	469.2%	S		
	Application	М	514%	510%	М	500% Min for all	400%Min for all
		L	512%	510%	L	sizes sizes	sizes
		XL	517%	511%	XL		



SI No.	Tests Pro		Proposed Device actual Data		Acceptance Criteria
	<u>Detection of Holes</u> <u>in Medical Gloves</u>	Size		Size	
	ASTM D6319-19	S		S	
3.	/ASTM D5151-19 Standard Test	М	AQL 2.5	М	AQL 2.5
	Method for Detection of Holes in Medical Gloves	L		L	
		XL		XL	
	<u>Powder Free</u> <u>Residue</u>	Size	Residual powder content	Size	Residual powder content
	ASTM D6124-06 (Reapproved 2017) Standard Test	S	0.41 mg	S	
4.		М	0.23 mg	М	≤ 2 mg/Glove Max
	Method for Residual Powder on Medical	L	0.57 mg	L	S 2 mg/ Glove Max
	Gloves	XL	0.37 mg	XL	

B. Biocompatibility

The materials used in the TC Gloves- Powder Free Nitrile Examination Gloves are biocompatible based on the biocompatibility tests mentioned in the Guidance for Industry and FDA Staff - Medical Glove Guidance Manual:

- Invitro Cytotoxicity
- Skin Sensitization
- Skin Irritation
- Acute Systemic Toxicity
- Material-Mediated Pyrogenicity

These tests are performed according to ISO 10993-1:2018, Biological Evaluation of Medical Devices - Part 1, Evaluation and Testing within a Risk Management Process.

Table 3: Biocompatibility Test Summary

SI. No	Test Performed	Proposed Device	Acceptance Criteria	Result
1.	In Vitro Cytotoxicity	Under the conditions of the study, cytotoxic to L-929 cells. Additional Testing was performed to determine if this was a systemic toxicity concern	Under the conditions of the study, not cytotoxic.	-
2.	Skin Sensitization	Under the conditions of the study not a sensitizer	Under the conditions of the study not a sensitizer	Pass
3.	Skin Irritation	Under the condition of study not an irritant	Under the condition of study not an irritant	Pass



4.	Acute Systemic Toxicity	Under the conditions of the study, the device extracts do not pose a systemic toxicity concern	Under the conditions of the study, the device extracts do not pose a systemic toxicity concern	Pass
5.	Material Mediated Pyrogenicity	Under the conditions of the study, the device did not demonstrate a material mediated pyrogenicity response.	Under the conditions of the study, the device did not demonstrate a material mediated pyrogenicity response.	Pass

C. Shelf Life Study

Expiration Date: Three years from the date of Manufacture

A provisional expiration date of three years from the date of manufacturing is provided based on the shelf life study conducted as per ASTM D7160-16 Standard Practice for Determination of Expiration Dating for Medical Gloves.

D. Clinical Test Data

The clinical study was not conducted as clinical data is not needed for TC Gloves- Powder Free Nitrile Examination Gloves.

VIII. CONCLUSION

The conclusion drawn from the non-clinical tests demonstrates that the subject device, TC Gloves-Powder Free Nitrile Examination Gloves are as safe, as effective and perform as well as or better than legally marketed predicate device in K192333.