



October 8, 2021

Guangdong Gynda Medical Technology Co., Ltd  
% Boyle Wang  
General Manager  
Shanghai Truthful Information Technology Co., Ltd.  
Room 608, No.738, Shangcheng Rd., Pudong  
Shanghai, Shanghai 200120  
China

Re: K212078

Trade/Device Name: Medical Examination Gloves  
Regulation Number: 21 CFR 880.6250  
Regulation Name: Non-Powdered Patient Examination Glove  
Regulatory Class: Class I, reserved  
Product Code: LZA  
Dated: September 6, 2021  
Received: September 10, 2021

Dear Boyle Wang:

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 Federal Register.

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-devices/medical-device-safety/medical-device-reporting-mdr-how-report-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>) for more information or contact DICE by email ([DICE@fda.hhs.gov](mailto:DICE@fda.hhs.gov)) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

For Clarence W. Murray, III, PhD  
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

## Indications for Use

510(k) Number (if known)  
K212078

Device Name  
Medical Examination Gloves

Indications for Use (Describe)

The Medical Examination Gloves are disposable devices intended for medical purposes that are worn on the examiner's hands to prevent contamination between patient and examiner.

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)

### CONTINUE ON A SEPARATE PAGE IF NEEDED.

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

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## 510(k) Summary

This summary of 510(k) safety and effectiveness information is being submitted in accordance with 21 CFR 807.92.

### **1.0 submitter's information**

Name: Guangdong Gynda Medical Technology Co.,Ltd  
Address: No.13, Quan'an Third Road, Phase 2 of High-tech Zone, Nanxiong City, Shaoguan City, Guangdong Province, 512400, China  
Phone Number: +86-20-37738661  
Contact: Olivia Chen  
Date of Preparation: 2021.05.27

### **Designated Submission Correspondent**

Mr. Boyle Wang  
Shanghai Truthful Information Technology Co., Ltd.  
Tel: +86-21-50313932  
Email: Info@truthful.com.cn

### **2.0 Device information**

Trade name: Medical Examination Gloves  
Common name: Patient Examination Gloves  
Classification name: Non-powdered patient examination glove  
Model(s): S, M, L, XL

### **3.0 Classification**

Production code: LZA  
Regulation number: 21CFR880.6250  
Classification: Class I  
Panel: General Hospital

### **4.0 Predicate device information**

Manufacturer: Ever Global (Vietnam) Enterprise Corp  
Device: Disposable Powder Free Nitrile Examination Glove, White/  
Blue/ Black/ Pink Color  
510(k) number: K171422

### 5.0 Intended use

The Medical Examination Gloves are disposable devices intended for medical purposes that are worn on the examiner’s hands to prevent contamination between patient and examiner.

### 6.0 Device description

The proposed device is Powder Free Medical Examination Gloves. The proposed device is blue. The design of proposed device is addressing the standards as ASTM D6124, ASTM D5151, and ASTM D6319. The proposed device is non-sterile.

### 7.0 Summary comparing technological characteristics with predicate device

**Table1-General Comparison**

Item	Proposed device	Predicated device	Remark
510(k) number	Pending	K171422	
Product Code	LZA	LZA	Same
Regulation No.	21CFR880.6250	21CFR880.6250	Same
Class	I	I	Same
Intended Use	The Medical Examination Gloves is a disposable device intended for medical purposes that is worn on the examiner’s hands to prevent contamination between patient and examiner.	The Disposable Powder Free Nitrile Examination Glove, White/ Blue/ Black/ Pink Color is a disposable device intended for medical purposes that is worn on the examiner’s hands to prevent contamination between patient and examiner.	Same
Powdered or Powered free	Powdered free	Powdered free	Same
Design Feature	ambidextrous	ambidextrous	Same
Labeling Information	Single-use indication, powder free, device color, device name, glove size and quantity, Medical Examination Gloves, Non-Sterile	Single-use indication, powder free, device color, device name, glove size and quantity, Disposable Powder Free Nitrile Examination Glove, Non-Sterile	Same

**Table2 Device Dimensions Comparison**

Predicate Device(K171422)	Designation	Size					Tolerance
		XS	S	M	L	XL	
	Length, mm	230	230	230	230	230	min
	Width, mm	75	85	95	105	115	±5
	Thickness, mm:						
	Finger	0.05					min
Palm	0.05					min	
Proposed Device	Designation	Size				Tolerance	
		S	M	L	XL		
	Length, mm	220	230	230	230	min	
	Width, mm	80	95	110	120	±10	
	Thickness, mm:						
	Finger	0.05					min
Palm	0.05					min	
Remark	Analysis1						

Analysis1: The sizes and tolerances of proposed device are different with those of the predicate, but they all meet the requirements of ASTM D6319-19, so the differences do not raise any new safety or performance questions.

**Table3 Performance Comparison**

Item			Proposed device	Predicated device	Remark
Colorant			blue	White/ Blue/ Black/ Pink	Analysis2
Physical Properties	Before Aging	Tensile Strength	14MPa, min	14MPa, min	SAME
		Ultimate Elongation	500%min	500%min	SAME
	After Aging	Tensile Strength	14MPa, min	14MPa, min	SAME
		Ultimate Elongation	400%min	400%min	SAME
	Comply with ASTM D6319			Comply with ASTM D6319	SAME
Freedom from Holes			Be free from holes when tested in accordance with ASTM D5151 AQL=2.5	Be free from holes when tested in accordance with ASTM D5151 AQL=2.5	SAME
Powder Content			0.15	Meet the requirements of ASTM D6124	SIMILAR

Analysis 2: The proposed device has different color to the predicate device, but all proposed devices are conducted the biocompatibility test, the test results shown that the color difference do not effect the safety of proposed device

**Table4 Safety Comparison**

Item		Proposed device	Predicated device	Remark
Material		Nitrile	Nitrile	SAME
Biocompatibility	Irritation	Under the conditions of the study, not an irritant	Comply with ISO10993-10	SAME
	Sensitization	Under conditions of the study, not a sensitizer.		
	Cytotoxicity	Under the conditions of the study, the device is potentially cytotoxic	Comply with ISO10993-5	Analysis3
	Systemic toxicity	Under the conditions of the study, the device does not elicit a systemic toxicity response in the model animal.	Complies with ISO 10993-11 Third edition 2017-09	
Label and Labeling		Meet FDA's Requirement	Meet FDA's Requirement	SAME

Analysis3: The proposed device is potentially cytotoxic, but all proposed devices are conducted the systemic toxicity test, the test results show that the proposed device is safe.

**8.0 Discussion of non-clinical testing**

The following tests were performed to evaluate the biocompatibility of the proposed device, the tests' results to show substantial equivalence to the predicate device:

ISO 10993-10:2010 Biological Evaluation Of Medical Devices - Part 10: Tests For Irritation And Skin Sensitization.

ISO 10993-5:2009 Biological Evaluation Of Medical Devices - Part 5: Tests For In Vitro Cytotoxicity

ISO 10993-11 Third edition 2017-09, Biological evaluation of medical devices - Part 11: Tests for systemic toxicity.

Clinical testing is not needed for this device.

**9.0 Discussion of performance testing (Bench)**

In summary, the performance testing of the subject device was conducted to adequately demonstrate the effectiveness of the device in accordance with the relevant test methods cited below:

ASTM D6124-06 (Reapproved 2017), Standard Test Method for Residual Powder on Medical Gloves

ASTM D5151-19, Standard Test Method for Detection of Holes in Medical Gloves.

ASTM D6319-19, Standard Specification For Medical Examination Gloves For Medical Application.

ASTM D7160-16, Standard Practice for Determination of Expiration Dating for Medical Gloves

## **10.0 Conclusion**

The conclusions drawn from the nonclinical tests demonstrate that the proposed device is as safe, as effective, and performs as well as the legally marketed predicated device.