

Food and Drug Administration 10903 New Hampshire Avenue Document Control Center – WO66-G609 Silver Spring, MD 20993-0002

## September 16, 2014

Zibo Yusheng Plastic Products Company Limited C/O Mr. Chu Xiaoan
Official Correspondent
Beijing Easy-Link Co.
Room 1606 Bldg 1 Yuan #209
Bei Si Huan Zhong Road, Haidian District
Beijing, PR China 100083

Re: K141108

Trade/Device Name: Powder-Free Vinyl Patient Examination Gloves, Clear (Non-Colored)

Regulation Number: 21 CFR 880.6250

Regulation Name: Patient Examination Glove

Regulatory Class: I Product Code: LYZ Dated: August 11, 2014 Received: August 14, 2014

#### Dear Mr. Xiaoan:

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. 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); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please contact the Division of Industry and Consumer Education at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address

http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to

http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm for the CDRH's Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

You may obtain other general information on your responsibilities under the Act from the Division of Industry and Consumer Education at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address

http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm.

Sincerely yours,

Tejashri Purohit-Sheth, M.D. Clinical Deputy Director

Tejashri Purohit-Sheth, M.D. Clinical Deputy Director DAGRID/ODE/CDRH FOR

Erin I. Keith, M.S.
Director
Division of Anesthesiology, General Hospital,
Respiratory, Infection Control and Dental Devices
Office of Device Evaluation
Center for Devices and
Radiological Health

Enclosure

## DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration

## **Indications for Use**

Form Approved: OMB No. 0910-0120 Expiration Date: January 31, 2017 See PRA Statement below.

510(k) Number (if known)			
K141108			
Device Name			
Powder Free Vinyl Patient Examination Gloves, Clear (non-colored)			
ndications for Use (Describe)			
Powder Free Vinyl Patient Examination Gloves, Clear (non-colo	ored) is a non-sterile disposable device intended for		
edical purposes that is worn on the examiner's hand or finger 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)		
— Flescription use (Fait 21 GFK 601 Subpait D)	Over-the-counter use (21 CFR 601 Subpart C)		
PLEASE DO NOT WRITE BELOW THIS LINE – CONTINUE ON A SEPARATE PAGE IF NEEDED.			
FOR FDA US	E ONLY		
Concurrence of Center for Devices and Radiological Health (CDRH) (S	ignature)		

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

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Department of Health and Human Services Food and Drug Administration Office of Chief Information Officer Paperwork Reduction Act (PRA) Staff PRAStaff@fda.hhs.gov

"An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB number."

## Section C

## 510(k) Summary

"This summary of 510(k) safety and effectiveness information is being submitted in accordance with the requirements of SMDA 1990 and 21 CFR 807.92."

"The as signed 510(k) number is: K141108 (applicant leave blank)

Premarket Notification [510(k)] Summary

[(a)(1)]. The summary contains on the first page, preferably on your letterhead paper, the submitter's name, address, phone and fax numbers, name of contact person, and date the summary was prepared:

**Submitter's name:** Zibo Yusheng Plastic Products Company Limited

**Submitter's address:** Li Wang Village, Zhoujia District, Hengtai County Zibo,

Shandong, 255414, China

**Phone number** : (86)533-3819127

**Fax number**: (86)533-3819127

Name of contact person: Cherry Xu

Date the summary was prepared: 2014-08-11

[(a)(2)]. The name of the device, including the trade or proprietary name if applicable, the common or usual name, and the classification name, if known

**Device Name**: Powder-Free Vinyl Patient Examination Gloves, Clear

(Non-colored)

**Proprietary/Trade name**: "Powder-Free Vinyl Patient Examination Gloves, Clear

(Non-colored)"

**Common Name:** Patient examination glove

Classification Name: Patient examination glove

**Device Classification:** I

**Regulation Number:** 21 CFR 880.6250

Panel: General Hospital (80)

**Product Code:** LYZ

[(a)(3)]. An identification of the legally marketed device to which your firm is claiming substantial equivalence.

Class I\* Powder-Free Vinyl Patient Examination Gloves, Clear (Non-colored) that meets all of the requirements of ASTM standard D 5250-06 (Reaffirmation 2011).

**Predicate device:** Powder free Vinyl Patient Examination Gloves, Clear(Non-colored), Tangshan Zhonghong Pulin Plastic Co.,Ltd.. K120968.

## [(a)(4)] A description of the device

**Device Description**: Powder-Free Vinyl Patient Examination Gloves, Clear (Non-colored) that meets all of the requirements of ASTM standard D 5250-06(Reaffirmation 2011).

## -- How the device functions:

PVC films form a barrier to body fluids and bloodborne Pathogens

### -- Scientific concepts that form the basis for the device

The PVC rubber is water tight under normal conditions of use. It's tensile properties cause it to conform to the hand, allowing movements necessary for a medical procedure.

-- Physical and performance characteristics such as design, materials and physical properties: PVC gloves are known to create a barrier to bloodborne pathogens and body fluids. ASTM conforming tensile properties create a glove that is strong and flexible. The glove is manufactured in accordance with the requirements of ASTM D5250 and ASTM D5151 requirements.

#### [(a)(5)] The summary describes the intended use of the device

**Device Intended Use:** Powder-Free Vinyl Patient Examination Gloves, Clear (Non-colored) is a disposable device intended for medical purposes that is worn on the examiner's hand or finger to prevent contamination between patient and examiner.

# [(a)(6)] A summary of the technological characteristics of new device compared to the predicate device.

The Powder-Free Vinyl Patient Examination Gloves, Clear (Non-colored) non sterile are summarized with the following technological characteristics compared to ASTM or equivalent standard.

Features & Description	Predicate Device	Subject Device	Result of Comparison
Company	Tangshan Zhonghong Pulin Plastic Co.,Ltd.	Zibo Yusheng Plastic Products Company Limited	
510(K) Number	K120968	K141108	
Product name	Powder Free Vinyl Patient	Powder-Free Vinyl Patient	Same
	Examination Gloves, Clear	Examination Gloves, Clear	
	(Non-colored)	(Non-colored)	
Product Code	LYZ	LYZ	Same
Size	Small/ Medium/	Small/ Medium/	Substantially
	Large/X large	Large/X large	equivalent
Intend for use	Powder free Vinyl Patient	Powder free Vinyl Patient	Substantially
	Examination Gloves,	Examination Gloves, Clear	equivalent
	Clear(Non-colored)is a	(Non-colored) is a disposable	
	disposable device intended for	device intended for medical	
	medical purposes that is worn on	purposes that is worn on the	
	the examiner's hand or finger to	examiner's hand or finger to	
	prevent contamination between	prevent contamination	
	patient and examiner.	between patient and examiner.	
Device	Meets ASTM D5250-06	Meets ASTM D5250 -06	Substantially
Description and	(Reapproved 2011)	(Reapproved 2011)	equivalent
Specifications			
Dimensions	Meets ASTM D5250-06	Meets ASTM D5250-06	Substantially
Length	(Reapproved 2011)	(Reapproved 2011)	equivalent
	≥230mm min.	230mm min for all sizes	
Dimensions	Meets ASTM D5250-06	Meets ASTM D5250-06	Substantially
Width	(Reapproved 2011)	(Reapproved 2011)	equivalent
		,	•
	Small 80-90 mm	Small 84-86 mm	
	Medium 90-100mm	Medium 94-96 mm	
	Large 100-110mm	Large 104-106mm	
	X large 110-120 mm	X large 113-115 mm	
Dimensions	Meets ASTM D5250-06	Meets ASTM D5250-06	
Thickness	(Reapproved 2011)	(Reapproved 2011)	

	Finger 0.05mm min.	Finger 0.05mm min.	
	Palm 0.08mm min.	Palm 0.08mm min.	
Physical Properties	Meets ASTM D5250-06	Meets ASTM D5250-06	Substantially
	(Reapproved 2011)	(Reapproved 2011)	equivalent
	Before aging/after aging	Before aging/after aging	
	Elongation ≥300%	Elongation≥300%	
	Tensile Strength≥11MPa	Tensile Strength≥11MPa	
Freedom from	Meets	Meets AST M	Substantially
Pinholes	• 21 CFR 800.20	D5151-06 (Reapproved 2011)	equivalent
	<ul> <li>ASTM D5250-06</li> </ul>		
	(Reapproved 2011)	Holes	
	• ASTM D 5151-06	Inspection Level I	
	(Reapproved 2011)	AQL2.5	
Residual Powder	Meets AST M	ASTM D6124-06	Substantially
residual I o waei	D6124-06 (Reaffirmation	(Reaffirmation 2011)	equivalent
	2011) (Real milation	Results generated values	oqui , arono
	2011)	below 2mg of residual powder	
Compare all	PVC	PVC	Substantially
	PVC	PVC	
materials used to fabricate the			equivalent
devices	D. I.	D. 1	0.1
Dusting or	PU	PU	Substantially
Donning Powder:			equivalent
Dusting or	PU	Surface Coating Agent	Substantially
Donning Powder:			equivalent
name			
Compare	Meets	Meets	Substantially
performance data	ASTM D5151-06	ASTM D5151-06	equivalent
supporting	(Reapproved 2011)	(Reapproved 2011)	
substantial	ASTM D5250-06	ASTM D5250-06	
equivalence	(Reapproved 2011)	(Reapproved 2011)	
	ASTM D6124-06	ASTM D6124-06	
	(Reaffirmation 2011)	(Reaffirmation 2011)	
Single Patient Use	Single Patient Use	Single Patient Use	Substantially
Single 1 attent USE	Single 1 attent OSC	Single 1 atient Osc	equivalent
Biocompatibility	SKIN IRRITATION DERMAL	Under the conditions of the	Substantially
Diocompanionity	and SENSITIZATION		equivalent
	STUDIES Meets ISO	study, the device is not a sensitizer or an irritant.	equivalent
	10993-10:2002/Amd.1:2006	sensitizer of an Iffitalit.	
	10993-10:2002/Amd.1:2006	CHANA ADDAM AMAGAA	
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		DERMAL and SENSITIZATION STUDIES Meets ISO 10993-10 Third	
Y 1 1 2 2		DERMAL and SENSITIZATION STUDIES Meets ISO 10993-10 Third Edition 2010-08-01	
Labeling for the	-Powder Free	DERMAL and SENSITIZATION STUDIES Meets ISO 10993-10 Third Edition 2010-08-01 -Powder Free	Substantially
legally marketed	-Powder Free -devices color:	DERMAL and SENSITIZATION STUDIES Meets ISO 10993-10 Third Edition 2010-08-01 -Powder Free -devices color:	Substantially equivalent
legally marketed device to which	-Powder Free -devices color: Clear(Non-colored)	DERMAL and SENSITIZATION STUDIES Meets ISO 10993-10 Third Edition 2010-08-01 -Powder Free -devices color: Clear(Non-colored)	
legally marketed device to which substantial	-Powder Free -devices color: Clear(Non-colored) -Patient Examination Glove	DERMAL and SENSITIZATION STUDIES Meets ISO 10993-10 Third Edition 2010-08-01  -Powder Free -devices color: Clear(Non-colored) -Patient Examination Glove	
legally marketed device to which substantial equivalence is	-Powder Free -devices color: Clear(Non-colored) -Patient Examination Glove -Non sterile	DERMAL and SENSITIZATION STUDIES Meets ISO 10993-10 Third Edition 2010-08-01  -Powder Free -devices color: Clear(Non-colored) -Patient Examination Glove -Non sterile	
legally marketed device to which substantial	-Powder Free -devices color: Clear(Non-colored) -Patient Examination Glove -Non sterile -Single Use Only	DERMAL and SENSITIZATION STUDIES Meets ISO 10993-10 Third Edition 2010-08-01  -Powder Free -devices color: Clear(Non-colored) -Patient Examination Glove -Non sterile -Single Use Only	
legally marketed device to which substantial equivalence is	-Powder Free -devices color: Clear(Non-colored) -Patient Examination Glove -Non sterile	DERMAL and SENSITIZATION STUDIES Meets ISO 10993-10 Third Edition 2010-08-01  -Powder Free -devices color: Clear(Non-colored) -Patient Examination Glove -Non sterile	

[(b)(1)] A brief discussion of the nonclinical submitted, reference, or relied on in the premarket notification submission for a determination of substantial equivalence.

Powder-Free Vinyl Patient Examination Gloves, Clear (Non-colored)meet requirements per ASTM D5250-06(Reaffirmation 2011), per ASTM D6124-06(Reaffirmation 2011), per 21 CFR 800.20 and ISO 10993-10 Third Edition 2010-08-01.

[(b)(2)] A brief discussion of the clinical submitted, reference, or relied on in the premarket notification submission for a determination of substantial equivalence.

Clinical data is not needed for gloves or for most devices cleared by the 510(k) process. [(b)(3)] The conclusions drawn from the nonclinical tests that demonstrate that the device is as safe, as effective, and performed as well or better than the legally marketed device identified in

(a)(3).

It can be concluded that the Powder Free Vinyl Patient Examination Gloves meet the ASTM standard or equivalent standard and FDA requirements for waterleak test on pinhole AQL, meet labeling claims and the Powder-Free Vinyl Patient Examination Gloves, Clear (Non-colored) is as safe, as effective, and performs as well as the predicate device, Powder free Vinyl Patient Examination Gloves, Clear(Non-colored), Tangshan Zhonghong Pulin Plastic Co.,Ltd. K120968.