

May 28, 2022

Amazing Rubber Products Pvt Ltd % Manoj Zacharias Consultant Liberty Management Group Ltd. 75 Executive Dr. STE114 Aurora, Illinois 60504

Re: K212597

Trade/Device Name: Amazing+ Latex Examination Powder Free Gloves Regulation Number: 21 CFR 880.6250 Regulation Name: Non-Powdered Patient Examination Glove Regulatory Class: Class I, reserved Product Code: LYY Dated: May 2, 2022 Received: May 2, 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 <a href="https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpmn/pmn.cfm">https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpmn/pmn.cfm</a> 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 <a href="https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products">https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products</a>); 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 <u>https://www.fda.gov/medical-devices/medical-device-safety/medical-device-reporting-mdr-how-report-medical-device-problems</u>.

For comprehensive regulatory information about medical devices and radiation-emitting products, including information about labeling regulations, please see Device Advice (<u>https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance</u>) and CDRH Learn (<u>https://www.fda.gov/training-and-continuing-education/cdrh-learn</u>). 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 (<u>https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice</u>) for more information or contact DICE by email (<u>DICE@fda.hhs.gov</u>) 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

# **Indications for Use**

510(k) Number *(if known)* K212597

Device Name

Amazing+ Latex Examination Powder Free Gloves

#### Indications for Use (Describe)

Amazing+ Latex Examination Powder Free Gloves are disposable device intended for medical purpose that is worn on the examiner's hand 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.

#### \*DO NOT SEND YOUR COMPLETED FORM TO THE PRA STAFF EMAIL ADDRESS BELOW.\*

The burden time for this collection of information is estimated to average 79 hours per response, including the time to review instructions, search existing data sources, gather and maintain the data needed and complete and review the collection of information. Send comments regarding this burden estimate or any other aspect of this information collection, including suggestions for reducing this burden, to:

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."

# 510K SUMMARY K212597

#### As required by 21CFR§807.92(c)

#### A. APPLICANT INFORMATION

Applicant	Amazing Rubber Products Pvt. Ltd.
Address	Plot No 14c, Cochin Special Economic Zone , Kakkanad, Cochin-682 037, Kerala- India
Phone	9447053062
Fax	
E-mail	info@amazingglove.com
Contact Person	Mr. Jayasankar S
Designation	Executive Director
Contact Number	9447053062
Contact Email	info@amazingglove.com
Date Submitted	12 August 2021

#### **B. DEVICE IDENTIFICATION**

Name of the device	Amazing <sup>+</sup> Latex Examination Powder Free Gloves
Product proprietary or	Amazing <sup>+</sup>
trade name	
Common or usual name	Latex Examination Powder Free Gloves
Classification name	Patient Examination Gloves
Device Classification	Class I
Product Code	LYY
Regulation Number	21 CFR 880.6250
Review Panel	General Hospital
	-

#### C. PREDICATE DEVICE

6.1	JR MEDIC Blue Latex Examination Powder Free Gloves
Equivalency is claimed	
510(K) Number	K192329
Regulatory Class	Class I
Product code	LYY

#### **D. DESCRIPTION OF THE DEVICE:**

 $Amazing^+$  Latex Examination Powder Free Gloves are manufactured to meet all the current specifications listed under the ASTM Specification D3578-2019, Standard Specification for Rubber Examination Gloves. They are made from Natural Rubber Latex. These gloves are natural in color (No color is added) and are powder free.

#### E. INDICATIONS FOR USE/INTENDED USE OF THE DEVICE:

Amazing<sup>+</sup> Latex Examination Powder Free Gloves are disposable devices intended for medical purpose that are worn on the examiner's hand to prevent contamination between patient and examiner.

#### **Device Performance** Comparison Characteristics Standards Predicate Current K192329 K212597 510(K) Number ---Name of device JR MEDIC Amazing<sup>+</sup> ------Blue Latex Latex Examination Examination Powder Free Gloves Powder Free Gloves ASTM Dimensions-Length > 230 mmLength > 230 mm Length D3578-2019 Similar Size Average X-Small 305 Small 306 Medium 307 Large 308 X-Large 310 Dimensions- Width ASTM Width Min 95+/-Width Min 95+/-10 Similar D3578-2019 10 mm (for mm (for medium size) medium size) Size Average X-Small 76 Small 85 96 Medium Large 106 X-Large 116 Physical ASTM Before Ageing Before Ageing Similar Properties-D3578-2019 Tensile Strength Tensile Strength > 18 Mpa Tensile > 18 Mpa Size Actual Strength value X-Small 22.07 Small 22.15 Medium 22.22 Large 22.30 X-Large 22.32 After Ageing Tensile After Ageing Similar Strength Tensile Strength > 14 Mpa > 14 Mpa Size Actual value X-Small 18.49 Small 18.56 Medium 18.67 18.74 Large X-Large 18.76

### F. TECHNOLOGICAL CHARACTERISTICS

Characteristics	Standards	Device Performance				Comparison
		Predicate	Current			
510(K) Number		K192329	K212597			
Physical Properties-	ASTM	Before Ageing	Before Ageing			Similar
Ultimate Elongation	D3578-2019	Ultimate Elongation		Elongation		
		> 650%	Size	Actu	al value	_
			X-Small		858	
			Small		869	
			Medium		874	
			Large		880	
		After Ageing	X-Large		882	
		Ultimate Elongation				
		>500%	Ultimate	Elongation	> 500%	
			Size	Actu	ial value	
			X-Small		841	
			Small		848	
			Medium		854	
			Large		860	
			X-Large		862	-
Thickness	ASTM	Palm > 0.08 mm		m > 0.08 m	m	Similar
	D3578-2019	Finger > 0.08 mm	Fing	ger > 0.08 n	nm	
			Size	Palm	Finger	
				(Actual	(Actual	
				value)	value)	
			X-Small	0.16	0.22	
			Small	0.16	0.22	
			Medium	0.16	0.22	
			Large	0.16	0.22	
			X-Large	0.16	0.22	
Powder Free Residue	ASTM D6214	≤2 mg/glove	<	2 mg/glove		Similar
			Size	Resid		-
				powd	er content	
				(mg/g	glove)	
			X-Small		0.21	
			Small		0.21	
			Medium		0.22	-
			Large		0.22	-
			X-Large		0.22	-
	Primary Skin	Under the condition	Under the	e condition	of study	Same
	Irritation-ISO	of study, not an	n	ot an irritan	t	
	10993-10:2010( E)	irritant				
D:	Dermal	Under the conditions		ne condition		Same
Biocompatibility	Sensitization-ISO	of the study, not a	study	, not a sens	itizer	
	10993-10:2010( E)	sensitizer				
	In vitro	Under the conditions			Same	
	cytotoxicity ISO10993-5	of the study,	study	y, cytotoxic		
	:2009(E)	cytotoxic				
	Material mediated	Under the	Under th	e condition	s of the	Com-
	Pyrogenicity	conditions of the		/, non-pyrog		Same
	ISO 10993-	study non pyrogenic	Brudy	, non pyrog		
	11:2017(E) / USP	, r,8-me				
	41<151>					

Characteristics	Standards	Device Per	Comparison	
		Predicate	Current	<b>F</b>
510(K) Number		K192329	K212597	
Biocompatibility	Acute Systemic Toxicity Test ISO 10993-11:2017(E)	Under the conditions of study the device extracts do not pose a systemic toxicity concern	Under the conditions of study the device extracts do not pose a systemic toxicity concern	same
Water Tight (1000 ml)	ASTM D5151-2019	Passes AQL-1.5	Passes AQL-1.5	Same
Intended use		JR MEDIC Blue Latex Examination Powder Free Gloves are disposable devices intended for medical purpose that are won on the examiner's hand to prevent contamination between patient and examiner.	Amazing <sup>+</sup> Latex Examination Powder Free Gloves are disposable devices intended for medical purpose that are worn on the examiner's hand to prevent contamination between patient and examiner.	Same
Material	-	Natural Latex	Natural Latex	Identical
Color	-	Blue	Natural (No color is added)	different
Texture	-	Finger Texture	Finger texture	Identical
Size	ASTM D3578- 2019	Small, Medium, Large & X Large	X Small, Small, Medium, Large, X-Large	Similar
Single Use	Medical Glove Guidance Manual - Labeling	Single Use	Single Use	Same
Sterile/non sterile	-	Non sterile	Non sterile	Same
Powder/Powder free	-	Powder free	Powder free	Same
Label and Labeling	FDA Label requirements	Meets FDA's label and labeling requirements	Meets FDA's label and labeling requirements	Same
Manufacturer(s)	-	JR Engineering & Medical Technologies (M) SDN.BHD. Malaysia.	Amazing Rubber Products Pvt. Ltd.	

# G. SUMMARY OF NON-CLINICAL PERFORMANCE DATA

Test Method	Purpose	Acceptance Criteria	Result	
ASTM D3578-2019 Standard	To determine the	Min 230 mm for all sizes	X-Small 305 mm	
Specification for Rubber Examination	length of the gloves		Small 306 mm	
Gloves			Medium 307 mm	
			Large 308 mm	
			X-Large 310 mm	
ASTM D3578-2019 Standard	To determine the width	X-Small 70+/-10 mm	X-Small 76 mm	
Specification for Rubber Examination	of the gloves	Small 80+/-10mm	Small 85 mm	
Gloves	-	Medium 95+/-10 mm	Medium 96 mm	
		Large 111+/-10 mm	Large 106 mm	
		X-Large 115+/-10 mm	X-Large 116 mm	

Test Method	Purpose	Acceptance	Result			
		Criteria				
ASTM D3578-2019 Standard Specification for Rubber Examination Gloves	To determine the thickness of the gloves	Palm 0.08 mm min Finger 0.08 mm min for all sizes	Size X-Small Small Medium Large X-Large	Palm           0.16mm           0.16mm           0.16mm           0.16mm           0.16mm           0.16mm	<b>Finger</b> 0.22mm 0.22mm 0.22mm 0.22mm 0.22mm	
ASTM D3578-2019	To Determine the physical properties-Tensile strength	Before AgeingTensileStrength18MpaMinfor allsizesAfter AgeingTensileStrength14MpaMinfor allsizes	Size X-Small Small Medium Large X-Large	<b>Before</b> ageing 22.07 Mpa 22.15 Mpa 22.22 Mpa 22.30 Mpa 22.32 Mpa	After ageing 18.49 Mpa 18.56 Mpa 18.67 Mpa 18.74 Mpa 18.76 Mpa	
Standard Specification for Rubber Examination Gloves	To Determine the physical properties-Ultimate Elongation	Before AgeingUltimateElongation650%Minfor allsizesAfterAgeingUltimateElongation500%Minfor allsizes	Size X-Small Small Medium Large X-Large	Before ageing 858% 869% 874% 880% 880% 882%	After ageing 841% 848% 854% 860% 860%	
	To Determine the physical properties-stress at 500% Elongation	<b>Before Ageing</b> 5.5 Mpa Max for all sizes	Size X-Small Small Medium Large X-Large	<b>Before</b> ageing 5.1 Mpa 5.2 Mpa 5.2 Mpa 5.2 Mpa 5.2 Mpa	NA	
ASTM D5151-2019 Standard Test Method for Detection of Holes in Medical Gloves	To determine the holes in the gloves	AQL 2.5	Gloves Passes AQL 1.5		QL 1.5	
ASTM D6124-06 (Reapproved 2017) Standard Test Method for Residual Powder on Medical Gloves	To determine the residual powder in the gloves	2 Mg/Glove Max	Size X-Small Small Medium Large X-Large	-Small 0.21 mg/glov nall 0.21 mg/glov ledium 0.22 mg/glov 0.22 mg/glov		
ASTM D 5712-95 ( Re approved 2010) Standard Test Method for the Analysis of Protein in Natural Rubber	To determine the extractable protein in the gloves.	200 µg/ dm <sup>2</sup> Max for all sizes	Size X-Small Small Medium Large X-Large	Protein           ζ-Small         43.65 μ           βmall         43.65 μ           Δedium         43.65 μ           μarge         43.65 μ		

### The performance test data of the non-clinical tests meet following standards:

- > ASTMD 3578-2019 Standard Specification for Rubber Examination Gloves
- > ASTMD 5151-2019 Standard Test Method for Detection of Holes in Medical Gloves
- ASTMD 6124-06 (Reapproved 2017) Standard Test Method for Residual Powder on Medical Gloves
- ASTMD 5712-95 (Re approved 2010) Standard Test Method for the Analysis of Protein in Natural Rubber

# H. CLINICAL TESTING SUMMARY

This section is not applicable because clinical data not needed for gloves.

# I. CONCLUSION

The conclusion drawn from the nonclinical tests demonstrates that the subject device in 510(K) submission Amazing<sup>+</sup> Latex Examination Powder Free Gloves is as safe, as effective, and performs as well as or better than the legally marketed predicate device JR MEDIC Blue Latex Examination Powder Free Gloves (*K192329*).