

December 3, 2022

Safeskin Retailing (HK) Limited % Stephan Toupan President Dawa Medical LLC 7320 NW 12th Street Suite 103 Miami, Florida 33126

Re: K222715

Trade/Device Name: Black Nitrile Powder Free Patient Examination Glove, Non Sterile

Regulation Number: 21 CFR 880.6250

Regulation Name: Non-Powdered Patient Examination Glove

Regulatory Class: Class I, reserved

Product Code: LZA

Dated: September 5, 2022 Received: September 8, 2022

Dear Stephan Toupan:

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 -S

Bifeng Qian, M.D., Ph.D.
Assistant Director
DHT4B: Division of Infection Control
and Plastic Surgery Devices
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and Infection Control Devices
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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 See PRA Statement below.

K222715						
Device Name BLACK NITRILE POWDER-FREE PATIENT EXAMINATION GLO	VE, NON STERILE					
ndications for Use (Describe) A nitrile patient examination glove is a disposable device made of nitrile rubber intended for medical purposes that is worn on the examiner's hand or finger to prevent contamination between patient and examiner						
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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 SEPARAT	E PAGE IF NEEDED.					

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

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510(k) Summary K222715 As required by 21CFR§807.92(c)

1.0 Submitter:

Name : Alvin Ho

Address : Safeskin Retailing (HK) Limited

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Central, Hong Kong

Phone No. : +6012 826 5625

Date of Summary Prepared : 11 November 2022

2.0 Identification of the subject device:

Trade Name: : Black Nitrile Powder-Free Patient Examination Glove, Non-

Sterile

Common Name: : Patient Examination Gloves Classification Name : : Patient Examination Gloves

Device Classification :

Regulation Number: : 21 CFR 880.6250

Product Code : LZA

3.0 Predicate Device:

K190942

Trade Name: Disposable Powder Free Nitrile Examination Glove, Black Color

Company: Ever Growth (Vietnam) Co., Ltd

4.0 Description of The Subject Device:

Black Nitrile Powder-Free Patient Examination Glove, Non-Sterile is manufactured from nitrile rubber. Innersurface of gloves undergoes surface treatment process to produce a smooth surface that assists the user in donning the gloves with ease without using any lubricant such as powder or silicone on the glove surface. The glove is ambidextrous, i.e., can be worn on right or left hand.

5.0 Indication for use:

A Nitrile patient examination glove 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.

6.0 Comparison of the Technological Characteristics of the Device:

The Black Nitrile Powder-Free Patient Examination Gloves, Non-Sterile are summarized with the following technological characteristics compared to ASTM D6319 or equivalent standards as shown in Table 1

Table 1

		DEVICE			
CHARACTERISTICS	STANDARDS	PREDICATE	CURRENT	COMPARISON ANALYSIS	
		BLACK	BLACK		
510(k) Number	-	K190942	K222715		
Manufacturer(s)	-	Ever Growth (Vietnam) Co., Ltd	Safeskin Retailing (HK) Limited		
Material	ASTM D6319 Nitrile		Nitrile	Same	
Color	-	Black	Black	Same	
Sterility	-	Non-Sterile	Non-Sterile	Same	
Handedness - Ambidextrous		Ambidextrous	Ambidextrous	Same	
Physical Properties	ASTM D6319				
Before Aging Tensile Strength: Ultimate Elongation:		14Mpa, min 500% min	27.8 – 33.1 Mpa 530 -600%	Different but within the ASTM standard	
After Aging Tensile Strength: Ultimate Elongation:		14Mpa, min 400% min	31.5 – 35.3 Mpa 470- 550%	Different but within the ASTM standard	
Thickness: - Finger - Palm	ASTM D6319	0.05mm min 0.05mm min	Min 0.10mm for (XS, S, M, L, XL) Min 0.06mm for (XS, S, M, L, XL)	Different but within the ASTM standard	
Powder Free	ASTM D6124	< 2mg per glove	Below 2mg of residual powder	Similar	

		DEVICE	E PERFORMANCE		
CHARACTERISTICS	STANDARDS	PREDICATE	CURRENT	COMPARISON ANALYSIS	
		BLACK	BLACK		
	Primary Skin Irritation – ISO 10993-10:2010 (E) & Consumer Product Safety Commission Title 16. Chapter II, Part 1500	Passes	Under conditions of this study, the test material did not cause an irritant response. The Primary Irritant Response Category is deemed 'Negligible'	Same	
Biocompatibility	Dermal Sensitization- ISO 10993-10: 2010 (E) & Consumer Product Safety Commission, Title 16,Chapter II, Part 1500.3 (c) (4)	Passes	Under conditions of this study,the test material did not produce a skin sensitization effect in the guinea pigs.	Same	
	Cytotoxicity – MEM Elution, ISO 10993-5: 2009	Passes	The Systemic Toxicity study was conducted as the accepted alternative to the cytotoxic test	Different – but additional test of Acute Systemic Toxicity is conducted and passed	
	Acute Systemic Toxicity, ISO 10993- 11:2017 (E)	Not Tested	Under conditions of this study,the test item did not induce any systemic toxicity in Swiss albino mice.	Different. The subject glove was tested using systemic toxicity test and passed, but the Predicate did not have the test performed	

CHARACTERISTICS	STANDARDS	DEVICE PER	RFORMANCE		
		PREDICATE	CURRENT	COMPARISON ANALYSIS	
		BLACK	BLACK		
Watertight (1000ml)	ASTM D5151:2019	In accordance with ASTM D6319-10 and ASTM D5141-06 (Reapproved 2011), G-1, AQL 2.5	Meets • 21 CFR 800.20 • ASTM D6319-10 (Reapproved 2015) Tested in accordance with ASTM D5151 (Reapproved 2015) with acceptable results At an AQL 1.5	Different, but within ASTM standard; AQI 1.5 is more stringent than AQL 2.5	
Intended use	-	The Nitrile Powder Free Patient examination glove is a non-sterile disposable device intended for medical purpose that is worn on the examiners hands or finger to prevent contamination between patient and examiner.	A Nitrile patient examination glove is a disposable device made of nitrile rubber intended for medical purposes that is worn on the examiner's hand or finger to prevent contamination between patient and examiner.	Similar	
Size	Medical Glove Guidance Manual – Labeling	Extra Small Small Medium Large X Large	Extra Small Small Medium Large Extra Large	Same	
Single use	Medical Glove Guidance Manual – Labeling	Single Use	Single Use	Same	

7.0 Summary of Non-Clinical Testing

The performance test data of the non-clinical tests for this powder free nitrile examination glove is summarized as per below.

			Accep	Acceptance Criteria			Results	
Test Method	Standard	Purpose of Testing		Before aging	After aging	Before aging	After aging	Status
Physical Properties	ASTM D412 (Standard Test Method for Vulcanized Rubber and Thermoplastic Elastomers-Tension)	To evaluate the tensile (tension) properties of glove.	Tensile strength	Min 14.0 MPa	Min 14.0 MPa	XS - 28.9 S - 27.8 M - 28.7 L - 28.3 XL - 28.0	XS - 31.7 S - 32.1 M - 32.4 L - 31.9 XL - 31.5	Pass
			Ultimate elongation	Min 500%	Min 400%	XS - 540 S - 530 M - 560 L - 540 XL - 540	XS - 480 S - 470 M - 490 L - 480 XL - 490	Pass

Test Method	Standard	Purpose of Testing	Glove Size	Acceptance Criteria		Res	ults	Status
				Length	Min 240 mm	Length	244 mm	Pass
			X-Small	Width	70 ± 10 mm	Width	74.0 mm	Pass
			A-Siliali	Thickness	Finger – min 0.05mm	Thickness	0.10 mm	Pass
					Palm – min 0.05mm		0.06 mm	
				Length	Min 240 mm	Length	245 mm	Pass
		To measure the length, width and thickness ofglove	Small	Width	80 ± 10 mm	Width	85.0 mm	Pass
	ASTM D3767			Thickness	Finger – min 0.05mm	Thickness	0.10 mm	Pass
Dimension	mension Standard Practice for Rubber— Measurementof Dimensions				Palm – min 0.05mm		0.06 mm	
				Length	Min 240 mm	Length	244 mm	Pass
			Medium	Width	95 ± 10 mm	Width	95.0 mm	Pass
				Thickness	Finger – min 0.05mm	Thickness	0.10 mm	Pass
					Palm – min 0.05mm		0.06 mm	
				Length	Min 240 mm	Length	243 mm	Pass
			Large	Width	110 ± 10 mm	Width	106 mm	Pass

		Thickness	Finger – min 0.05mm	Thickness	0.10 mm	Pass
			Palm – min 0.05mm		0.06 mm	
		Length	Min 240 mm	Length	242 mm	Pass
	X-Large	Width	120 ± 10 mm	Width	117 mm	Pass
		Thickness	Finger – min 0.05mm	Thickness	0.10 mm	Pass
			Palm – min 0.05mm		0.06 mm	

Test Method	Standard	Purpose of Testing	Acceptance Criteria	Results	Status
Watertight	ASTM D5151 (Standard Test Method for Detection of Holesin Medical Gloves)	To detect holes that leak water and thereby compromise the usefulness of the glove.	Sample size: 315 pcs Inspection level: G1 AQL: 1.5, Acceptance No. 10	The batch size for this sampling is 150,001 to 500,000. Hence, according to the single sampling plan GI, the sample to be drawn is under code M equivalent to 315 pieces with accept 10 and reject 11 to be accepted under AQL 1.5. For Size XS, during the test, 1piece was found with leaks. Hence it falls within the acceptance criteria. For Size S, during the test, 3piece was found with leaks. Hence it falls within the acceptance criteria. For Size M, during the test, 2piece was found with leaks. Hence it falls within the acceptance criteria. For Size L, during the test, 3piece was found with leaks. Hence it falls within the acceptance criteria. For Size L, during the test, 3piece was found with leaks. Hence it falls within the acceptance criteria. For Size XL, during the test, 1piece was found with leaks. Hence it falls within the acceptance	Pass
				accepted under AQL 1.5. For Size XS, during the test, 1piece was found with leaks. Hence it falls within the acceptance criteria. For Size S, during the test, 3piece was found with leaks. Hence it falls within the acceptance criteria. For Size M, during the test, 2piece was found with leaks. Hence it falls within the acceptance criteria. For Size L, during the test, 3piece was found with leaks. Hence it falls within the acceptance criteria. For Size L, during the test, 3piece was found with leaks. Hence it falls within the acceptance criteria. For Size XL, during the test, 1piece was found with leaks. Hence it falls	

Test Method	Standard	Purpose of Testing	Acceptance Criteria	Results	Status
Residual Powder	ASTM D6124 (Standard Test Method for Residual Powder on Medical Gloves)	To determine the amount of residual and non-powder solids found on gloves	Less than 2 mg per glove Requirement : <2mg/glove	Result XS :0.26mg/glove Result S :0.32mg/glove Result M :0.28mg/glove Result L :0.34mg/glove Result XL :0.38mg/glove	Pass

8.0 Non-clinical performance testing methods full titles:

- ASTM D412 Test Methods for Vulcanized Rubber and Thermoplastic Elastomers— Tension
- ASTM D573 Test Method for Rubber—Deterioration in an Air Oven x ASTM D3578 Specification for Rubber Examination Gloves
- ASTM D6319 Standard Specification for Nitrile Examination Gloves for Medical Application
- ASTM D5151 Test Method for Detection of Holes in Medical Gloves
- ASTM D6124 Test Method for Residual Powder on Medical Gloves
- ISO 2859 Sampling Procedures and Tables for Inspection by Attributes Test results show that under the conditions of the testing, there is no difference inphysical attributes between the proposed device and the predicate device.
- ISO 10993 Part 10: Tests for Irritation and Sensitization. Both Skin Irritation and Dermal Magnuson/Kligman Sensitization performed.
- ISO 10993 Part 11: Tests for assessment of Systemic Toxicity

9.0 Summary of Clinical Testing

No clinical testing is included in this submission

10.0 Conclusion

The conclusion drawn from the non-clinical tests demonstrates that the subject "Black Nitrile Powder-Free Patient Examination Glove, Non-Sterile" is as safe, as effective, and performs as well as, or better than the legally marketed predicate device predicate K190942.