

August 3, 2022

Makrite Industries, Inc.
% Ivy Wang
Technical Manager
Shanghai Sungo Management Consulting Company Limited
14th Floor, 1500# Century Avenue
Shanghai, 200122
China

Re: K220550

Trade/Device Name: Disposable Surgical Face Mask (M663BE, M663BT)

Regulation Number: 21 CFR 878.4040 Regulation Name: Surgical apparel

Regulatory Class: Class II

Product Code: FXX

Dear Ivy Wang:

The Food and Drug Administration (FDA) is sending this letter to notify you of an administrative change related to your previous substantial equivalence (SE) determination letter dated June 14, 2022. Specifically, FDA is updating this SE Letter for a typographical error in the company name as an administrative correction.

Please note that the 510(k) submission was not re-reviewed. For questions regarding this letter please contact Brent Showalter, Ph.D., OHT6: Office of Orthopedic Devices, 240-402-1840, <u>brent.showalter@fda.hhs.gov</u>.

Sincerely,

Brent Showalter -S

Brent Showalter, Ph.D.
Assistant Director
DHT6B: Division of Spinal Devices
OHT6: Office of Orthopedic Devices
Office of Product Evaluation and Quality
Center for Devices and Radiological Health



June 14, 2022

Makrite Industries Co, Inc. % Ivy Wang Technical Manager Shanghai Sungo Management Consulting Company Limited 14th Floor, 1500# Century Avenue Shanghai, 200122 China

Re: K220550

Trade/Device Name: Disposable Surgical Face Mask (M663BE, M663BT)

Regulation Number: 21 CFR 878.4040 Regulation Name: Surgical Apparel

Regulatory Class: Class II Product Code: FXX Dated: May 10, 2022 Received: May 16, 2022

Dear Ivy 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 <u>Federal Register</u>.

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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/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) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

Katherine D. Kavlock -S

for
Brent Showalter, Ph.D.
Assistant Director
DHT6B: Division of Spinal Devices
OHT6: Office of Orthopedic 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.

rotect both the patient and healthcare personnel from						
ransferring of microorganisms, body fluids and particulate material. These face masks are intended for use in infection ontrol practices to reduce the potential exposure to blood and body fluids. This is a single use, disposable device(s), rovided non-sterile.						
_						
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."

510(K) Summary

Prepared date: 2022-02-16

A. Applicant:

MAKRITE INDUSTRIES, INC.

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Contact Person: Bob Wen Tel: +886-2-26982419 Fax: +886-2-26982423

Email: bobwen@makrite.com.tw

Submission Correspondent: Primary contact: Ms. Ivy Wang

Shanghai SUNGO Management Consulting Co., Ltd.

Room 1401, Dongfang Building, 1500# Century Ave., Shanghai 200122, China

Tel: +86-21-58817802

Email: haiyu.wang@sungoglobal.com Secondary contact: Mr. Raymond Luo

Room 1401, Dongfang Building, 1500# Century Ave., Shanghai 200122, China

Tel: +86-21-68828050

Email: fda.sungo@gmail.com

B. Device:

Trade Name: Disposable Surgical Face Mask Common Name: Disposable Surgical Mask

Model: M663BE, M663BT

Regulatory Information

Classification Name: Surgical Face Mask

Classification: Class II Product code: FXX

Regulation Number: 878.4040 Review Panel: Surgical Apparel

C. Predicate device:

K210433

Surgical Face Mask

Wuhan Dymex Healthcare Co., Ltd.

D. Indications for use:

The Disposable Surgical Face Masks are intended to be worn to protect both the patient and healthcare personnel from transferring of microorganisms, body fluids and particulate material. These face masks are intended for use in infection control practices to reduce the potential exposure to blood and body fluids. This is a single use, disposable device(s), provided non-sterile.

E. Device Description:

The Disposable Surgical Face Mask is blue color, single use, four-layer, flat-folded masks with nose piece and ear loops or ties. The blue colorant is polypropylene (PP) master batch.

The body of the mask is composed of four layers: the inner(4th layer) and outer(1st layer) layers are made of spun-bond polypropylene, the 2nd layer is made of polyethylene film and the 3rd layer is made of melt blown polypropylene. The nose piece is made of Iron core coated with polypropylene, ear loop is made of Nylon and Spandex, and the ties are made of spun-bond polypropylene.

Each mask contains ear loops or ties to secure the mask over the user's face and mouth with a bendable nose piece to firmly fit over the nose. This device is not made with natural rubber latex.

The disposable surgical face masks are sold non-sterile and are intended to be single use, disposable devices.

F. Comparison with predicate device

Table 1 General Comparison

Device		Proposed Device	Predicate Device	Result	
Manufacturer		MAKRITE INDUSTRIES, INC.	Wuhan Dymex Healthcare Co., Ltd.	-	
510K number		-	K210433	-	
Model na	me	Disposable Surgical Face Mask	Surgical Face Mask	Same	
Classification		Class II Device, FXX (21 CFR878.4040)	Class II Device, FXX (21 CFR878.4040)	Same	
Intended use		The Disposable Surgical Face Masks are intended to be worn to protect both the patient and healthcare personnel from transferring of microorganisms, body fluids and particulate material. These face masks are intended for use in infection control practices to reduce the potential exposure to blood and body fluids. This is a single use, disposable device(s), provided non-sterile.	The Surgical Face Masks are intended to be worn to protect both the patient and healthcare personnel from transfer of microorganisms, body fluids and particulate material. These face masks are intended for use in infection control practices to reduce the potential exposure to blood and body fluids. This is a single use, disposable device(s), provided non-sterile.	Same	
Model		Ear loop, Tie-on, Flat pleated, 4 layers	Ear loop, Tie-on, Flat pleated, 3 layers	Different	
Material	Outer layer	Spun-bond polypropylene	Spun-bond polypropylene	Same	
	Middle layer	2 nd layer: polyethylene film	/	Different	

		3 rd layer: Melt blown polypropylene	Melt-blown Polypropylene	Same
	Inner Spun-bond polypropylene layer nonwoven fabric		Spun-bond polypropylene	Same
Nose clip Ear loops		Iron core coated with polypropylene	Malleable polyethylene wire	Different
		Nylon, Spandex	Spandex	Different
	Ties	Spun-bond polypropylene	Spunbond Polypropylene	Same
Color		Blue	Blue	Same
Dimension (Length)		17.5cm+/-0.45cm	17.5cm+/-0.2cm	Similar
Dimension (Width)		9.5cm+/-0.45cm	9.5cm+/-0.2cm	Similar
OTC use		Yes	Yes	Same
Sterility		Non-Sterile	Non-Sterile	Same
Use		Single Use, Disposable	Single Use, Disposable	Same
ASTM F2100 level		Level 3	Level 3	Same
Fluid Resistance Performance ASTM F1862		32 out of 32 pass at 160 mmHg	32 out of 32 pass at 160 mmHg	Same
	Particulate Filtration Efficiency ASTM ≥ 98% F2299		≥ 98%	Same
Bacterial Efficiency F2101	Efficiency ASTM $\geq 98\%$		≥ 98%	Same
Differential Pressure (Delta P) EN 14683 < 6.0mmH ₂ O/cm ² Annex C		< 6.0mmH ₂ O/cm ²	Same	
Flammab 1610 16	Flammability 16 CFR Class 1 1610 16		Class 1	Same
Biocompa	Biocompatibility ISO10993		ISO10993	Same

G. Non-Clinical Test Conclusion

Non-clinical tests were conducted to verify that the proposed device met all design specification for the standards and test methods. The test results demonstrated that the proposed device complies with the following standards and the requirements stated in the Guidance for Industry and FDA Staff: Surgical Masks – Premarket Notification [510(k)] Submission issued on March 5, 2004:

- ➤ ISO 10993-5: 2009 Biological Evaluation of Medical Devices -- Part 5: Tests For In Vitro Cytotoxicity
- ➤ ISO 10993-10: 2010 Biological Evaluation of Medical Devices Part 10: Tests for Irritation and Skin Sensitization

- ➤ ASTM F2100, Standard Specification for Performance of Materials Used in Medical Face Masks
- ASTM F1862, Standard Test Method for Resistance of Medical Face Masks to Penetration by Synthetic Blood (Horizontal Projection of Fixed Volume at A Known Velocity);
- ➤ EN 14683, Medical Face Masks—Requirements and Test Methods;
- ASTM F2101, Standard Test Method for Evaluating the Bacterial Filtration Efficiency (BFE) Of Medical Face Mask Materials, Using A Biological Aerosol of Staphylococcus Aureus;
- ASTM F2299, stand test method for determining the initial efficiency of materials used in medical face masks to penetration by particulates using latex spheres;
- ➤ 16 CFR 1610, Standard for the Flammability of clothing textiles;

Table 2 - Performance Testing & Biocompatibility

	Table 2 - Performance Testing & Biocompatibility						
Test Methodology	Purpose	Acceptance Criteria:	Result				
	1	ASTM F2100 Level 3					
Fluid Resistance		29 out of 32 pass at	Pass				
		160 mmHg for level 3	32 out of 32 pass at 160				
			mmHg, 3 lots				
Particulate	The manage of		Pass				
Filtration	The purpose of the performance	≥ 98%	Average 99.929%, 99.947%,				
Efficiency			99.978%				
Bacterial Filtration	testing is to demonstrate the		Pass				
Efficiency		. ≥ 98%	Average 99.8%, 99.9%,				
	functionality of		99.8%				
Differential	the subject		Pass				
Pressure	device. < 6.0mmH ₂ O/o	< (0 II 0 / 2	Average 3.7mmH ₂ O/cm ² ,				
		< 0.0mmH2O/cm²	3.8mmH ₂ O/cm ² ,				
			3.8mmH ₂ O/cm ²				
Flammability		Class 1	Pass, Class 1				
Cytotoxicity	The purpose of	Non-cytotoxic	Under the conditions of the				
	the testing is to		study, the device is				
	demonstrate the		non-cytotoxic.				
Irritation	safety of the	Non-irritating	Under the conditions of the				
	subject device.		study, the device is				
			non-irritating.				
Sensitization		Non-sensitizing	Under the conditions of the				
			study, the device is				
			non-sensitizing				

H. Clinical Test Conclusion

No clinical study is included in this submission.

I. Conclusion

The conclusions drawn from the nonclinical tests demonstrate that the subject device is as safe, as effective, and performs as well as or better than the legally marketed predicate device K210433.