

March 1, 2021

Maquet GmbH % Mark Smith Regulatory Affairs Manager Getinge 45 Barbour Pond Road Wayne, New York 07470

Re: K201927

Trade/Device Name: Getinge GSS610N Series Steam Sterilizer Regulation Number: 21 CFR 880.6880 Regulation Name: Steam Sterilizer Regulatory Class: Class II Product Code: FLE Dated: February 1, 2021 Received: February 2, 2021

Dear Mark Smith:

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 <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,

Clarence W. Murray III, 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)* K201927

Device Name Getinge GSS610N Series Steam Sterilizer

Indications for Use (Describe)

The Getinge GSS610N Series Steam Sterilizer is intended for use by health care facilities to sterilize wrapped and unwrapped porous and nonporous heat and moisture stable items such as surgical instruments and linens by means of pressurized steam. The GSS610N Series Steam Sterilizer is available in 3 models differentiated by chamber length; GSS610N Model 610N10 (40.9 inch chamber), GSS610N Model 610N14 (51.1 inch chamber), and GSS610N Model 610N15 (60.6 inch chamber).

See "GSS610N Load chart table"

GSS610N Load chart table

Cycle Type	Factory Settings		gs	Load Configuration (Note 1)		ximum item hamber Len	
	Exp. Temp.	Exp. Time	Drying Time		610N10 1040 mm 40.9 in	610N14 1400 mm 51.1 in	610N15 1540 mm 60.6 in
P1 PREVAC 1	135.0°C (275.0°F)	3 min	16 min	Full Instrument packs	18	24	24
P2 PREVAC 2	135.0°C (275.0°F)	3 min	3 min	Full fabric packs	40	40	60
P3 PREVAC 4	132.2°C (270.0°F)	4 min	30 min	Full Instrument packs	18	24	24
P4 PREVAC 5	132.2°C (270.0°F)	4 min	5 min	Full fabric packs	40	40	60
P5 B & D TEST	134°C (273.0°F)	3 min, 30 sec	0 min	1 B&D Test Pack In an EMPTY chamber (other than loading accessories)	1 Test Pack	1 Test Pack	1 Test Pack
P6 GRAVITY 1	121.1°C (250.0°F)	30 min	45 min	Full Instrument packs Full fabric packs	18 40	24 40	24 60
P7 GRAVITY 2	135.0°C (275.0°F)	10 min	45 min	Full Instrument packs Full fabric packs	18 40	24 40	24 60
P8 GRAVITY 3	132.2°C (270.0°F)	15 min	45 min	Full Instrument packs Full fabric packs	18	24	24
P9 IUSS 1	135.0°C (275.0°F)	3 min	1 min	Full Instrument packs	1	1	1
Vac				Full fabric packs	1	1	1
P10 IUSS 2	135.0°C (270.0°F)	10 min	30 sec	Full Instrument packs	1	1	1
Grav				Full fabric packs	1	1	1
P11 IUSS 3	132.2°C (270.0°F)	4 min	1 min	Full Instrument packs	1	1	1
Grav				Full fabric packs	1	1	1
P12 Vented Bottles	121.1°C (250.0°F)	45 min	3kPa/min (0.44 psi/min) (Note 2)	Each container1000 mL (34 fl oz) or smaller (Note 3)	3	3	3

Getinge GSS610N Series Steam Sterilizer Cycles and Load Chart

NOTE: Liquid Cycles are not intended for the sterilization of liquids used for direct patient contact.

TABLE NOTES:

The load configurations listed in these tables are those used during testing validations of the sterilizer. These
configurations follow AAMI Standard ST8: Hospital steam sterilizers where applicable (fabric packs are process
challenge devices as described in ANSI/AAMI ST8 and were made to be consistent with the packs described in
ANSI/AAMI ST8). For guidance on processing loads in the sterilizer, refer to AAMI Standard ST79:
Comprehensive guide to steam sterilization and sterility assumance in health care facilities.

2. Cooldown rate

3. Use vented or open containers only.

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.

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510(k) Summary – K201927 Getinge GSS610N Series Steam Sterilizer

<u>Submitted by</u>	Maquet GmbH
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- <u>Contact person</u> Mark N. Smith Manager, Regulatory Affairs Phone: 585-272-5274 Email: <u>mark.n.smith@getinge.com</u>
- **510(k) number** K201927
- **Date Prepared** February 26, 2021

Proprietary Device Name

Trade Name: Getinge GSS610N Series Steam Sterilizer

Models: 610N10 40.9 inch (1000 mm) long chamber 610N14 51.1 inch (1400 mm) long chamber 610N15 60.6 inch (1540 mm) long chamber

Common Name: Steam Sterilizer

Classification: Steam Sterilizer (21CFR880.6880, Product Code 80 FLE)

Predicate Devices

[K172159] Getinge GSS67N Series Steam Sterilizer Getinge. Date 01/30/2018



Description of Device

The steam sterilizer is used by health care facilities to sterilize wrapped and unwrapped porous and nonporous heat and moisture stable items such as surgical instruments and linens by means of pressurized steam.

The Getinge GSS610N Series Steam Sterilizer employs both with gravity/downward displacement with positive pulse conditioning and pressure/vacuum pulsing for dynamic air removal. All cycle phases are sequenced and monitored by the control system, providing both audible and visual notification of deviation from certain operating parameters

Indications for Use

The Getinge GSS610N Series Steam Sterilizer is intended for use by health care facilities to sterilize wrapped and unwrapped porous and nonporous heat and moisture stable items such as surgical instruments and linens by means of pressurized steam. The GSS610N Series Steam Sterilizer is available in 3 models differentiated by chamber length; GSS610N Model 610N10 (40.9 inch chamber), GSS610N Model 610N14 (51.1 inch chamber), and GSS610N Model 610N15 (60.6 inch chamber).

Cycles and Load Chart

Getinge GS	S610N Series	Steam Ster	ilizer Cycle	es and Load	d Chart

Cycle Type	Factory Settings		gs	Load Configuration		ximum Items	•
			(Note 1)		Chamber Length		
	Exp.	Exp.	Drying		610N10	610N14	610N15
	Temp.	Time	Time		1040 mm	1400 mm	1540 mm
					40.9 in	51.1 in	60.6 in
P1	135.0°C	3 min	16 min	Full instrument packs	18	24	24
PREVAC 1	(275.0°F)						
P2	135.0°C	3 min	3 min	Full fabric packs	40	40	60
PREVAC 2	(275.0°F)						
P3	132.2°C	4 min	30 min	Full instrument packs	18	24	24
PREVAC 4	(270.0°F)						
P4	132.2°C	4 min	5 min	Full fabric packs	40	40	60
PREVAC 5	(270.0°F)						
P5	134°C	3 min,	0 min	1 B&D Test Pack in an	1 Test	1 Test	1 Test
B & D TEST	(273.0°F)	30 sec		EMPTY chamber (other	Pack	Pack	Pack
	· · · ·			than loading accessories)			
P6	121.1°C	30 min	45 min	Full instrument packs	18	24	24
GRAVITY 1	(250.0°F)			Full fabric packs	40	40	60
P7	135.0°C	10 min	45 min	Full instrument packs	18	24	24
GRAVITY 2	(275.0°F)			Full fabric packs	40	40	60
P8	132.2°C	15 min	45 min	Full instrument packs	18	24	24
GRAVITY 3	(270.0°F)			Full fabric packs	40	40	60
P9	135.0°C	3 min	1 min	Full instrument packs	1	1	1
IUSS 1	(275.0°F)						

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Vac				Full fabric packs	1	1	1
P10 IUSS 2	135.0°C (270.0°F)	10 min	30 sec	Full instrument packs	1	1	1
Grav				Full fabric packs	1	1	1
P11 IUSS 3	132.2°C (270.0°F)	4 min	1 min	Full instrument packs	1	1	1
Grav				Full fabric packs	1	1	1
P12 Vented Bottles	121.1°C (250.0°F)	45 min	3kPa/min (0.44 psi/min) (Note 2)	Each container 1000 mL (34 fl oz) or smaller (Note 3)	3	3	3

NOTE: Liquid Cycles are not intended for the sterilization of liquids used for direct patient contact.

TABLE NOTES:

1. The load configurations listed in these tables are those used during testing validations of the sterilizer. These configurations follow *AAMI Standard ST8: Hospital steam sterilizers* where applicable (fabric packs are process challenge devices as described in ANSI/AAMI ST8 and were made to be consistent with the packs described in ANSI/AAMI ST8). For guidance on processing loads in the sterilizer, refer to *AAMI Standard ST79: Comprehensive guide to steam sterilization and sterility assurance in health care facilities*.

2. Cooldown rate

3. Use vented or open containers only.

Technological Characteristics Table:

Shown below is a comparison of the subject device Getinge GSS610N Series Steam Sterilizer versus the predicate device (Getinge GSS67N Series Steam Sterilizer):

ltem	Getinge GSS610N [Subject device]	Getinge GSS67N [Predicate Device] K172159	Comparison
Pressure Vessel			_
Chamber Sizes	Model 610N10: 26.4" x 42.5" x 40.9" (672 x1080 x1040 mm)	Model 6710: 26.4" x 27.6" x 39.4" (672 x700 x1000 mm)	Different Chamber Sizes
	Model 610N14: 26.4" x 42.5" x 55.1" (672 x1080 x1400 mm)	Model 6713: 26.4" x 27.6" x 51.2" (672 x700 x1300 mm)	Different Chamber Sizes
	Model 610N15: 26.4" x 42.5" x 60.6" (672 x1080 x1540 mm)	Model 6717: 26.4" x 27.6" x 66.9" (672 x700 x1700 mm)	Different Chamber Sizes
Chamber Volumes	Model 610N10: Single Door 26.52 Cu Ft (751L), Double Door 26.59 Cu Ft (753L) Model 610N14: Single Door	Model 6710: Single Door 17 Cu Ft (481L), Double Door 16.5 Cu Ft (468L)	The chamber volumes correspond to the chamber sizes
	35.73 Cu Ft (1012L),		

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			-
	Double Door 35.77 Cu Ft (1013L)	Model 6713: Single Door 22 Cu Ft (621L), Double Door 21.5 Cu Ft (609L)	
	Model 610N15: Single Door	21.3 Cu Ft (009L)	
	39.3 Cu Ft (1113L), Double		
	Door 39.34 Cu Ft (1114L)	Model 6717: Single Door 28.6	
		Cu Ft (809L), Double Door	
		28.11 Cu Ft (796L)	
ASME Pressure Vessel	All pressure vessels are	All pressure vessels are built	Same
	built to ASME Sect. VIII,	to ASME Sect. VIII, Div. 1	
	Div. 1		
"U" Stamped Unfired Pressure Vessel)	Chamber: 45 psi Jacket: 45 psi	Chamber: 43.5 psi Jacket: 51 psi	The pressure rating differs slightly
Material of Jacket	Stainless Steel (SA240- 304)	Stainless Steel (SA240- 304)	Same
Material of Chamber	Stainless Steel (SA240- 316L)	Stainless Steel (SA240-316L)	Same
Vacuum method	Vacuum pump standard.	Vacuum pump standard.	Same
Chamber Closure			
Door Operation	Horizontal opening/closing	Vertical opening/closing door	Different door
	door design operated by	design with pneumatic	design.
	an electrical motor.	cylinder actuator	
Safety and Interlocks			
Door switch	Electro-mechanical logic	Electro-mechanical logic	Same
system/steam to			
chamber interlock			
Cycles			
Types of cycles offered	Prevac 132.2°C, 4	Prevac 132.2°C, 4 min;	Same
51 5	min; 135°C, 3 min,	135°C, 3 min,	
	Gravity 121°C, 30 min;	Gravity 121°C, 30 min;	
	135°C 10 min; 132.2°	135°C 10 min; 132.2° 15	
	15 min;	min;	
	IUSS 135°C, 3 min,;	IUSS 135°C, 3 min,;	
	135C, 10 min; 132.2°C	135C, 10 min; 132.2°C 4	
	4 min	min	
	Vented Bottles 121°C	Vented Bottles 121°C	
	45min.	45min.	
	BD Test	BD Test	
Maximum Load	Reference chart for	Reference chart for	Same
Capacities	maximum loads within	maximum loads within	
	GSS610N - up to 25lbs	GSS67N - up to 25lbs per	
	per tray	tray	
Utility Requirements			



Primary Electrical Connection	GSS610N are available for connection to: 208V 3ph 60Hz (Not GSS61015N with integrated electrical heated steam boiler) 460V 3ph 60Hz 480V 3ph 60Hz	GSS67N are available for connection to: 208V 3ph 60Hz 460V 3ph 60Hz 480V 3ph 60Hz 600V 3ph 60Hz 600V 3ph 60Hz (Not with integrated electrical heated steam boiler)	Similar electrical connections
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Summary of Non-Clinical Testing

Shown below is a summary of the non-clinical testing that was performed with this device:.

Test Performed	Device Description	Test Method	Acceptance Criteria	Results
Biological Perfo	ormance Tests			
Sterilization Efficacy Validation Biological Performance with a fabric PCD	GSS610N10 GSS610N14 GSS610N15	AAMI ST8:2013 (R2018) §5.5.2	Sterility Assurance Level (SAL) 10 ⁻⁶	Pass
Biological Performance with liquid loads	GSS610N10 GSS610N14 GSS610N15	AAMI ST8:2013 (R2018) §5.5.3	Sterility Assurance Level (SAL) 10 ⁻⁶	Pass
Biological Performance with a wrapped instrument PCD	GSS610N10 GSS610N14 GSS610N15	AAMI ST8:2013 (R2018) §5.5.4	Sterility Assurance Level (SAL) 10 ⁻⁶	Pass
Biological performance of immediate-use steam sterilization for single-wrapped or unwrapped nonporous items	GSS610N10 GSS610N14 GSS610N15	AAMI ST8:2013 (R2018) §5.5.5	Sterility Assurance Level (SAL) 10 ⁻⁶	Pass
Physical Perfor		1		
Chamber Temperature Profile	GSS610N10 GSS610N14 GSS610N15	AAMI ST8:2013 (R2018) §5.4.2.5	+3°C (or +6°F) and - 0°C (or -0°F) of the selected sterilization exposure temperature	Pass



Mechanical Air Removal Test	GSS610N10 GSS610N14 GSS610N15	AAMI ST8:2013 (R2018) §5.6.1	Load reaching exposuire temperature witin 10 secs	Pass
			Color change on BD chemical indicator sheet	Pass
Air Leak Rate Test	GSS610N10 GSS610N14 GSS610N15	AAMI ST8:2013 (R2018) §5.6.2	Average leak rate of 1 mmHg (0.13 kPa) (0.019 psia) per minute or less over the measured time interval.	Pass
Moisture Retention Rest	GSS610N10 GSS610N14 GSS610N15	AAMI ST8:2013 (R2018) §5.7	< 3% increase in presterilization test pack weight for fabric pack & <20% increase for wrapped instrument pack	Pass

Conclusion:

The conclusion drawn from the nonclinical test demonstrates that the subject device is as safe, as effective and performs as well as or better than the legally marketed predicate device, Class II (21 CFR 880.6880, Product code FLE)