						DATE	2.02.2021
						DOC.NO	FR-FD-BYM-0
baymed <sup>®</sup> "we work for your health"	PRODUC	T SPECIE	FICATIO	<b>IN SHEE</b>	T 🗖	PAGE NO	1
						REV.NO	
			DRODUCT NA	NAC		REV.DATE	
			PRODUCT NA				
	STANE	OARD SURGICAL G	GOWN - SMS 43	gsm Sterile - Fu	ll Ultrasonic		
	c				A		
			E		V/VI	VVVI	
	<b>S</b>	M	L	XL 135.0	<b>XXL</b>	XXXL	
	<b>A</b> 125,0	125,0	<b>L</b> 130,0	135,0	140,0	145,0	
	A       125,0         B       33,0	125,0 34,0	<b>L</b> 130,0 36,0	135,0 36,0	140,0 37,0	145,0 39,0	
	<b>A</b> 125,0	125,0	<b>L</b> 130,0	135,0	140,0	145,0	
	A       125,0         B       33,0         C       57,0	125,0 34,0	L 130,0 36,0 59,0	135,0 36,0	140,0 37,0 63,0	145,0 39,0	
	A       125,0         B       33,0         C       57,0	125,0 34,0 58,0	<b>L</b> 130,0 36,0	135,0 36,0 60,0	140,0 37,0	145,0 39,0 65,0	

UDI	PRODUCT NAME	SIZE	REF. CODE
8681744101325	8681744101325 Standard Surgical Gown		SG-01201-01
8681744101318	Standard Surgical Gown	М	SG-01201-02
8681744101301	Standard Surgical Gown	L	SG-01201-03
8681744101332	Standard Surgical Gown	XL	SG-01201-04
8681744101288	Standard Surgical Gown	XXL	SG-01201-05
8681744101295	Standard Surgical Gown	XXXL	SG-01201-06

		BAY	TEKS TEKNIK TEKSTIL SAN. VE TIC	. A.Ş.	TITLE: STAN	NDARD SURGICAL	. GOWN - SMS 43 gsr	n Sterile - Full Ultrasoni	
	UNIT: cm				SIZE: A4				
Tolerances vary according to customer demand. If the custome			does not hav	e a special reque	est, the tolerance val	ue in the specification			
TOLERAN	CE % ± 2	is accept	ed.						
DRAWING: DATE:		١L	departmantion NAM		NAME	SING	DATE	DWG NO:	
			PRODUCTION	A.AKAI	2			1	
		РРК	QUALITY CONTROL	K.KARAG	UN			Technical File	
		Ā						1	

LCH LCH	OTEKS LABORATUVAR ve GÖZETİM HİZMETLERİ A.Ş. enyurt Firuzköy Bulvarı No:29 34325 Avcılar İstanbul/ TÜRKİYE TEST REPORT DENEY RAPORU	Тея         Тоя         Состания         Состани
	BAYTEKS TEKSTİL SAN. VE TİC. A.Ş.	
Customer name:		
Address:	ORGANİZE SAN.BÖLG. 19 NOLU CAD. NO:	9 MERKEZ/KILIS
Buyer name:	TSE GAZİANTEP BELGELENDİRME MÜDÜ	RLÜĞÜ/İBRAHİM AÇAR
Contact Person: Order No:	KADİR KARAGÜL	
Article No:		
Name and identity of test item:	- Blue non-woven surgical gown	
The date of receipt of test item: Re-submitted/re-confirmation	01.03.2021	
date: Date of test: Remarks:	01.03.2021-11.03.2021	
Sampling:	The results given in this report belong to the reco	eived sample by vendor.
End-Use:		
Care Label:	Not specified.	
Number of pages of the report:	6	
a transferration and provide a transferration of the second		

The Turkish Accreditation Agency (TURKAK) is signatory to the multilateral agreements of the European co-operation for the Accreditation (EA) and of the International Laboratory Accreditation (ILAC) for the Mutual recognition of test reports.

EKOTEKS LABORATUVAR ve GÖZETİM HİZMETLERİ A.Ş. accredited by TÜRKAK under registration number [AB-0583-T] for ISO 17025:2017 as test laboratory.

The test and/or measurement results, the uncertainties (if applicable) with confidence probability and test methods are given on the following pages which are part of this report.

Seal Seal	Date 11.03.2021	Customer Representative Zahide TAPAN	Head of Testing Laboratory Sevim A. RAZAK 11.03/2021
1 mil			

This report shall not be reproduced other than in full except with the permission of the laboratory. Testing reports without signature and seal are not valid.

AB-0583-T	
21007884- ING	
03-21	

REQUIRED TESTS	RESULT	COMMENTS	
PHYSICAL PROPERTIES	p		
Water Permeability	P P		
Lint and Other Particles Generation From	r		
Nonwoven			
MICROBIOLOGICAL TESTS	Р		
Wet-Bacterial Penetration	p		
Dry-Bacterial Penetration	P		
Microbial Cleanliness (Bioburden)	r		
P: Pass			
E. Fail			
R: Refer to retailer technologist. Test results were evaluated according to EN 13	3795-1:2019 Standard Pe	rformance Properties limit values	

REMARK: Original samples are kept for 3 months and all technical records are kept for 5 years unless otherwise specified. If requested, measurement uncertainty will be reported. But unless otherwise specified, measurement uncertainty is not considered while stating compliance with specification or limit values The reported uncertainty is based on a standard uncertainty multiplied by a coverage factor k=2, providing a level of confidence of approximately 95 %. The declaration of conformity was given in accordance with the Simple Acceptance Decision Rule. Tests marked (\*) in this report are not included in the accreditation schedule.



This report shall not be reproduced other than in full except with the permission of the laboratory. *Testing reports without signature and seal are not valid.* 

AB-0583-7	
21007884 ING	
03-21	

REQUIREMENT

 $\geq$  20 cm H<sub>2</sub>O

## TEST RESULTS

WATER PERMEABILITY; ISO 811:2018

Hydrostatic Head Tester, Textest marka Fx 3000 model Temperature of water 20°C. Pressure increase ratio 10 mbar/min. Performed in the conditioned room  $(20\pm2^{\circ}C-65\%\pm4)$ 

	RESULT
Sample 1	54,1 cm H <sub>2</sub> O
Sample 2	56,2 cm H <sub>2</sub> O 53,7 cm H <sub>2</sub> O
Sample 3	63,7 cm H <sub>2</sub> O
Sample 4 Sample 5	60,1 cm H <sub>2</sub> O
	57,5 cm H <sub>2</sub> O

Average

## MICROBIAL CLEANLINESS (Bioburden)

## Test Metod: Ref: EN ISO 11737-1:2018 /TS EN ISO 11737-1:2018

The sample is put in extraciton liquid after shaking well, inoculated on the agar. After incubation at  $30 \pm 1$  ° C for 72 hours, growth microorganisms are counted on the agar.

	RESULTS	<u>REQUIREMENT</u> ≤300 cfu/100 cm <sup>2</sup>
obial cleanliness (cfu/ 100	14 cfu/100 cm <sup>2</sup>	\$300 clu/ 100 clil

\*cfu= Colony forming unit.

_	
	AB-0583-T
	21007884- ING
	03-21

#### **TEST RESULT**

Test Method: BS EN 22610: 2006 (Surgical drapes, garments and fresh air clothes used as medical devices for patients, hospital staff and equipment - Test method for determination of resistance to wet

A test sample is placed on the agar plate on a rotating disc. Bacteria carrier material and coating film are placed on the test sample and all parts are fixed on the disk. A finger is placed on the test sample to apply a certain force (3N  $\pm$  0.02). The finger moves on the test sample over the entire surface of the agar within 15 minutes. 5 studies are carried out for 15 minutes. 6. The study is repeated by inverting the sample.

Sample amount:	5 pieces 25x25cm2 30 μm thin, 25x25cm2 Polyurethane Film
Carrier Material:	30 μm thin, 25x25cm2 r orygerenance r mm   25x25cm2 HDPE Film
Coating Material:	Staphylococcus aureus ATCC 29213
Microorganism: Bacterial Concentration (kob / ml):	5x10 <sup>3</sup> kob / ml
Incubation Conditions:	$(36 \pm 1)$ ° C 48 hours

	RESUL	Penetrati	on Rate
Number of Populating	g Bacteria (ciu)	R <sub>CUM1</sub>	0,04
X1	45		0,09
Ya	59	R <sub>CUM2</sub>	0,17
<u>X2</u>	93	R <sub>CUM3</sub>	0,28
X3	124	R <sub>CUM4</sub>	
X4	135	R <sub>CUM5</sub>	0,40
X5			
7	659	1115	

X1 ......... X5: Number of colonies growing in 5 parallel petri in the same sample Z: number of colonies growing in the sixth petri dish

 $T: X_1 + X_2 + X_3 + X_4 + X_5 + Z$ 

#### RCUM1 = X1/T $R_{CUM2} = (X2 + X1)/T$ Rcum3 = (X3 + X2 + X1)/T Rcum4 = (X4 + X3 + X2 + X1)/T Rcum5 = (X5 + X4 + X3 + X2 + X1)/T

	BARRIER INDEX (IB)	Expected value
	Result	>28
and the second second second second second second second second second second second second second second second	4,99	≥2,0

IB = 6 - (CUM1 + CUM2 + CUM3 + CUM4 + CUM5)

AB-05	83 <b>-</b> T
21007 IN	
03	-21

# Test Method: ISO 22612: 2005 (Clothing for protection against infectious agents - Test method for

**resistance to dry microbial penetration**) Samples and containers are sterilized. Agar plates are placed in each container. Samples are placed aseptically in the apparatus. The covers are closed. After making a pot in the sample with the piston, the pistons are removed and  $0.5 \text{ g} \pm 0.1 \text{ g}$  are added to five samples from the powder contaminated with bacteria and the six to the non-contaminated powder. Then all openings are closed with a plastic bag. The device is operated to give 20,800 vibrations per minute. The test time is 30 minutes. After the test is over, all agar plates are incubated at 35 ° C for 24 hours.

	6 pieces 20x20 cm <sup>2</sup>		
Sample amount:	Bacillus subtilis ATCC 9372		
Mikroorganism:			
Bacterial concentration (cfu/ml):	ntration (cfu/ml): 1×10 <sup>8</sup>		
ncubation conditions:	35°C / 24 hours		
	RESULTS		
Numb	er of Populationg Bacteria (cfu)		
1			
		)	
2		)	
3		0	
4		0	
5		0	
6 (Control)		0	
Total		-	
Logarithm		a sublicated according to	
Logano Auraiaal gowps and	drapes - Requirements and test methods a	re evaluated according to	
* EN 13795-1:2019 Surgical gowns and	u		
Table-1.	RESULT	. 1.1/2/100	
		Expected Value	
	It (cfu/g)	≤300 cfu/g	
0	cfu/g		

AB-0583-T
21007884- ING
03-21

#### TEST RESULTS LINT AND OTHER PARTICLES GENERATION FROM NONWOWEN;

Test Metod: ISO 9073-10: 2003 (\*)

5 test samples that in cross direction are maintained to twisting and compression action with Gelbo Flex for inner and outer surface in a clean room condition (according to ISO 14644-1 Class 5).

Lint and particles detached from the sample are counted with counter device and classified to size range.

Min. measuring size of SOLAIR 3100 particles measuring device: 0,3 µm,

Max. measuring size of SOLAIR 3100 particles measuring device: 25 μm,

Air flow:  $28,3 \pm 1,4$  L/min

Working mode: 30 s x 10 consecutive periods

SAMPLE, INNER SURFACE (3 μm - 25 μm)Total linting:8Standard deviation: 5Coefficient of variation: 62%Coefficient of linting (CL):1	SAMPLE, OUTER SURFACE (3 μm - 25 μm)Total linting:44Standard deviation:35Coefficient of variation: 81%Coefficient of linting (CL): 2	
CAMPLE	MATERIAL (TOTAL)	

#### SAMPLE, MATERL

	and the second sec	
Total linting	51	
Total linting	.7	
Coefficient of linting (CL)*	.2	$C_{1,2} \in C_{2,2}$ (log 10) should be $\leq 4$ for analysis of critical

\*According to EN ISO EN ISO 13795-1:2019, Coefficient of linting (CL) (log 1 product area and less critical product area of both standard performance and high performance testing. both standard performance and high performance testing.