

Work Order	3212.1
Setup-Code	181029-10290-2801-01



Test Report

JIS Z 2801:2012 (Mod)

Antimicrobial products – Test for antimicrobial activity and efficacy

Test Object:

Coated Leneta-Foil vs. Pseudomonas aeruginosa EDCC 5272



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Report on Findings

Client: Address:	Nano-Care UK (Signo-Nanoc PO Box 225, Oswestry, Shrop	are UK Ltd) oshire, SY10 1DL, UK +44 1691 654 282
Work order no.:	3212.1	
Test object:	Coated Leneta-Foil vs. Pseud	domonas aeruginosa EDCC 5272
Sample description:	coated foil	
Date of receipt of sample:	2018-Oct-22	
Type of test:	JIS Z 2801:2012 Antimicrobia and efficacy	al products – Test for antimicrobial activity
Test Germ:	Pseudomonas aeruginosa ED	OCC 5272
Test laboratory:	QualityLabs BT GmbH	
Address:	Neumeyerstrasse 46a 90411 Nuremberg, Germany	
Setup-Code:	181029-10290-2801-01	
Sample material:	n.b.	
No. of pages in report:	7	
Report on findings Place at to the client: Recipi		Nuremberg, 2018-Nov-07 Nano-Care Deutschland GmbH
Laboratory Director:	Harald Gerauer, Laboratory Direc QualityLabs BT GmbH	etor
Released:	Markus Zehe, Managing Director QualityLabs BT GmbH	



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Declaration on Quality Assurance

This investigation was performed and supervised according to the standard operating procedure "SOP zu JIS Z 2801:2012 (Mod)" by QualityLabs BT GmbH. The laboratory and process are continually monitored by independent, external authorities, as well as by internal audits.

Archiving

A copy of the test report, a protocol of the measurement as well as the accompanying correspondence and business records are archived by QualityLabs BT GmbH. The retention period is at least 10 years.

Test description

Anti-bacterial activity is determined in accordance with a modified version of JIS Z 2801:2012.

During the test, a thin liquid-film containing the bacteria $(1.25 \times 10^4 \, \text{CFU} \, / \, \text{cm}^2)$ is applied directly to the test sample (Standard: 5 cm x 5 cm). To avoid desiccation a foil (Standard: 4cm x 4cm, Stomacher Bags) is applied. Immediately after inoculation, the bacteria from the reference sample are separated from the sample and the enveloping foil surfaces using ultrasound and vortex devices and the number of viable germs (CFU – colony-forming units) is determined (t_0 value). A further set of reference samples and samples given anti-microbial treatment is incubated with bacteria in a liquid-film and the enveloping foil in a damp environment at 37°C. After 24 hours, the bacteria are separated from the sample surfaces using ultrasound and vortex devices and the number of viable germs is determined (t_{24} value).



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Assessment of antimicrobial activity

A logarithmic germ reduction of ≥ 3 log scales of the antimicrobial sample in comparison to the respective reference is used as assessment criterion to pass the antimicrobial test.

Germ reduktion [log scales]	Antibacterial activity
< 3	Not sufficient antimicrobial activity
≥3	Sufficient antimicrobial activity



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References to Testconditions

Testconditions				
Sample size	25	cm ²		
Foil size	16	cm ²		
Volume Inoculum	300	μl		
Sample cleaning		-		

References to deviations, preincubations, special test conditions

NONE



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Test Results

	Sample Name	Sample Code		t ₀ (cells/cm²)		t	₂₄ (cells/cm²)	Reduction [%]	Log Reduction
1	Reference	102902310180011	5.3 x 10 ⁴	4.9 x 10 ⁴	4.2 x 10 ⁴	4.1 x 10⁵	3.5 x 10⁵	4.7 x 10 ⁵		Reference
2	Liquid Guard 2,9%	102902310180012				< 1.0 x 10 ¹	< 1.0 x 10 ¹	< 1.0 x 10 ¹	> 99.99	> 4

^{*}see "Interpretation of Results", page 6

Test strain	Pseudomonas aeruginosa EDCC 5272
Initial cell count inoculum / cm²	1.25 x 10⁴
Initials of the editor	MZ
Measurement ended on	Nov-11-2018



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NONE

Interpretation of the results based on the measurements

NONE

Editor	Mr Zehe	Crosschecked: Mr Mannala	
Faitor.	IVIT Zene	Crosschecked, Mr. Manhaia	

References

JIS Z 2801:2012 Antimicrobial products – Test for antimicrobial activity and efficacy