

Goodpoint Chemicals
Urda Tee 2/1
Jälgimäe 76404
Estonia

EXPERT OPINION

Determination of the Mycobactericidal Activity of **NOVA 104 (GLOBACID AF MED)** according to EN 14348:2005

This expert opinion is based on the test report TR-23-0496 dated 22 August 2023.

The product **Nova 104 (Globacid AF Med)** was tested against the microorganism *Mycobacterium avium* ATCC 15769 and *Mycobacterium terrae* ATCC 15755 according to the test method EN 14348:2005, 'Chemical disinfectants and antiseptics – Quantitative suspension test for the evaluation of mycobactericidal activity of chemical disinfectants in the medical area including instrument disinfectants – Test method and requirements (Phase 2, step 1)'. This test method is accredited according to MS ISO/IEC 17025:2017.

When tested under the following conditions, **Nova 104 (Globacid AF Med)** demonstrated a mycobactericidal activity against tested organisms conforming to the EN 14348:2005:

Activity	Concentration	Contact Time	Test Temperature	Soiling
Tuberculocidal	100%	1 minute	20 °C	Clean condition
Mycobactericidal	100%	3 minutes	20 °C	Clean condition

Kuala Lumpur, 22 August 2023



Dr Marven Lee Cheng Shoou
Managing Director

Test Report No.: TR-23-0496

**Determination of the Mycobactericidal Activity of
NOVA 104 (GLOBACID AF MED) according to EN 14348:2005**

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Test Method

EN 14348:2005

Chemical disinfectants and antiseptics – Quantitative suspension test for the evaluation of mycobactericidal activity of chemical disinfectants in the medical area including instrument disinfectants – Test method and requirements (Phase 2, step 1)

Client

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Jälgimäe 76404
Estonia

Testing Laboratory

TECOLAB Sdn. Bhd.
J-2-6, Pusat Komersial Jalan Kuching
No. 115, Jalan Kepayang, Off Jalan Kuching
51200 Kuala Lumpur
Malaysia

Kuala Lumpur, 22 August 2023



Dr Marven Lee Cheng Shouu
Managing Director

IDENTIFICATION OF TESTING LABORATORY

TECOLAB Sdn. Bhd.
J-2-6, Pusat Komersial Jalan Kuching
No. 115, Jalan Kepayang, Off Jalan Kuching
51200 Kuala Lumpur
Malaysia

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IDENTIFICATION OF CLIENT

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IDENTIFICATION OF TEST ITEM

Test item name:	NOVA 104 (GLOBACID AF med)
Lab ID:	G007-23-007
Batch no.:	230457
Expiry date:	June 2026
Manufacturer:	Goodpoint Chemicals
Receipt date:	29 June 2023
Storage conditions:	Room temperature away from sunlight
Product diluent recommended by manufacturer:	Not specified
Active substances:	60% Isopropanol 15% Ethanol
Product appearance:	Clear, colourless liquid

TEST METHOD & VALIDATION

Test method:	EN 14348:2005 Chemical disinfectants and antiseptics – Quantitative suspension test for the evaluation of mycobactericidal activity of chemical disinfectants in the medical area including instrument disinfectants – Test method and requirements (Phase 2, step 1)
Inactivation method:	Dilution-neutralization method
Inactivator:	30 g/L Tween 80 30 g/L Saponin 3 g/L Lecithin

EXPERIMENTAL CONDITIONS

Date of test:	17 July 2023
Product diluent:	Distilled water
Concentration / contact time:	100%* / 1 minute \pm 5 seconds 100%* / 3 minutes \pm 10 seconds
Test temperature:	(20 \pm 1) °C
Interfering substance:	Clean condition (0.3 g/L bovine serum albumin)
Test organism:	<i>Mycobacterium terrae</i> ATCC 15755 <i>Mycobacterium avium</i> ATCC 15769
Incubation temperature:	(37 \pm 1) °C
Incubation period:	21 days
Appearance of the product dilutions:	Clear, colourless liquid
Stability and appearance of product dilutions during test:	Homogenous without any precipitate

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* The product can only be tested at a concentration of 80% or less as some dilution is always produced by adding the test organisms and interfering substance.

Test method accredited according to MS ISO/IEC 17025. This test report may not be reproduced, in whole or in part, without the prior permission of the laboratory. The test results relate only to the test item provided by the client.

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CONTROLS AND VALIDATION

Test Organism	Validation Suspension	Experimental Conditions Control	Neutralizer Control	Method Validation
<i>M. terrae</i> ATCC 15755	Nv/10: 100.0	A: 142.0	B: 140.0	C: 98.0
<i>M. avium</i> ATCC 15769	Nv/10: 41.0	A: 41.5	B: 24.5	C: 29.0

The control and validation tests A, B, and C were within the basic limits:

- The number of cells per mL in the validation suspension, Nv/10, must be between 30 and 160,
- A must be equal to or greater than $0.5 \times \text{Nv}/10$ to verify the absence of any lethal effect in the experimental conditions,
- B must be equal to or greater than $0.5 \times \text{Nv}/10$ to verify the absence of neutralizer toxicity, and
- C must be equal to or greater than $0.5 \times \text{Nv}/10$ to validate the dilution-neutralization method.

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TEST RESULTS

For each product concentration and contact time, the log reduction (lg R) is calculated using the formula $\lg R = \lg N_0 - \lg N_a$, in which:

- N_0 is the number of cells per mL in the test mixture at the beginning of the contact time, and
- N_a is the number of cells per mL in the test mixture at the end of the contact time and before neutralization.

Test organism: *Mycobacterium terrae* ATCC 15755

Test suspension, N	$N_0: 4.69 \times 10^9$ $\lg N_0: 8.67$
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Concentration / Contact Time	Test, N_a	Reduction, $\lg R = \lg N_0 - \lg N_a$
100%* / 1 minute	$N_a: <1.40 \times 10^2$ $\lg N_a: <2.15$	$\lg R: >6.52 \pm 0.11$ %R: >99.99997%
100%* / 3 minutes	$N_a: <1.40 \times 10^2$ $\lg N_a: <2.15$	$\lg R: >6.52 \pm 0.11$ %R: >99.99997%

Test organism: *Mycobacterium avium* ATCC 15769

Test suspension, N	$N_0: 3.01 \times 10^9$ $\lg N_0: 8.48$
-----------------------	--

Concentration / Contact Time	Test, N_a	Reduction, $\lg R = \lg N_0 - \lg N_a$
100%* / 3 minutes	$N_a: 1.55 \times 10^4$ $\lg N_a: 4.19$	$\lg R: 4.29 \pm 0.11$ %R: 99.995%

* The product can only be tested at a concentration of 80% or less as some dilution is always produced by adding the test organisms and interfering substance.

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CONCLUSION

The test item achieved a reduction of ≥ 4.00 log against the test organisms *Mycobacterium terrae* ATCC 15755 and *Mycobacterium avium* ATCC 15769 under the tested conditions.

Therefore, **NOVA 104 (GLOBACID AF med)** has demonstrated a mycobactericidal activity according to EN 14348:2005 under the following conditions:

Concentration	Contact Time	Test Temperature	Soiling	
100%*	1 minute	20 °C	Clean condition	(Excl. <i>M.avium</i>)
100%*	3 minutes	20 °C	Clean condition	

Kuala Lumpur, 22 August 2023



Neni Iffanida Ismail
Microbiologist

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INFORMATION ON MEASUREMENT UNCERTAINTY & DECISION RULE

The statement of conformity given by EN 14348:2005 states that the test item shall be considered to have passed EN 14348 if it demonstrates ≥ 4.00 log reduction under the defined conditions.

The laboratory employs the simple acceptance decision rule to account for the measurement uncertainty when stating the statement of conformity. The measurement uncertainty and conformance probability are shown in the raw data and are summarized as follows:

Test Organism	Concentration / Contact Time	Log Reduction	Conformance	Conformance Probability [†]
<i>M. terrae</i> ATCC 15755	100%* / 1 minute	$>6.52 \pm 0.11$	Yes	<0.001% chance of false acceptance
	100%* / 3 minutes	$>6.52 \pm 0.11$	Yes	<0.001% chance of false acceptance
<i>M. avium</i> ATCC 15769	100%* / 3 minutes	4.29 ± 0.11	Yes	<0.433% chance of false acceptance

* The product can only be tested at a concentration of 80% or less as some dilution is always produced by adding the test organisms and interfering substance.

[†] The conformance probability follows a normal distribution. Therefore, the percentage of conformance can never be zero or 100% due to the asymptotic tails.

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RAW DATA

Test Method:	EN 14348:2005		
Product Name:	NOVA 104 (GLOBACID AF med)	Batch No.:	230457
Product Diluent:	Distilled water	Lab ID:	G007-23-007
Appearance of Product Dilutions:	Clear, colourless solution		
Inactivation:	Dilution-neutralization	Dilution Method:	Standard (80%)
Neutralizer:	30 g/L Tween 80, 30 g/L Saponin, 3 g/L Lecithin		
Interfering Substance:	0.3 g/L bovine serum albumin		
Test Organism:	Mycobacterium terrae ATCC 15755	Plating Method:	Spread plate
Incubation Temperature (°C):	37	Passing Criteria (lg):	4.00
Testing Period:	17/07/2023	Tested By:	NII
		Measurement Uncertainty (±):	0.11
		Verified By:	CSE

Validation & Controls

Validation Suspension (N _v)	V _{C1}	V _{C2}	N _{v0} = 100.0 Limit: 30 ≤ N _{v0} ≤ 160
	107	93	
Validation Suspension (N _{VB})	V _{C1}	V _{C2}	N _{v0} = N _{VB} /1000 Limit: 30 ≤ N _{v0} ≤ 160
	-	-	
Experimental Conditions Control (A)	V _{C1}	V _{C2}	A = 142.0 Limit: A ≥ 0.5 × N _{v0}
	143	141	
Neutralizer Control (B)	V _{C1}	V _{C2}	B = 140.0 Limit: B ≥ 0.5 × N _{v0} or N _{VB} /1000
	139	141	
Method Validation (C)	V _{C1}	V _{C2}	C = 98.0 Limit: C ≥ 0.5 × N _{v0}
Conc.: 100%	87	109	

Test Suspension & Procedure

Test Suspension (N)	N	V _{C1}	V _{C2}	$\bar{x}_{wm} = N = 4.69E+09$ $N_0 = N/10$ Limit: 8.17 ≤ lg N ₀ ≤ 8.70
	10 ⁻⁷	480	474	
	10 ⁻⁸	42	35	

Product Concentration	Contact Time	Dilution	V _{C1}	V _{C2}	Na = \bar{x} or $\bar{x}_{wm} \times 10$	lg Na	lg R = lg N ₀ - lg Na	Conformance Probability
100%	1 min	10 ⁰	<14	<14	<1.40E+02	<2.15	>6.52 ± 0.11	>99.999%
		10 ⁻¹	<14	<14			>99.99997%	
		10 ⁻²	<14	<14				
		10 ⁻³	<14	<14				
100%	3 mins	10 ⁰	<14	<14	<1.40E+02	<2.15	>6.52 ± 0.11	>99.999%
		10 ⁻¹	<14	<14			>99.99997%	
		10 ⁻²	<14	<14				
		10 ⁻³	<14	<14				
		10 ⁰						
		10 ⁻¹						
		10 ⁻²						
		10 ⁻³						

Raw Data of Colony Count

	N _v		N _{vB}		A		B		C		N ⁻⁷		N ⁻⁸	
V _{C1}	56	51	-	-	68	75	63	76	47	40	238	242	20	22
V _{C2}	48	45	-	-	70	71	64	77	53	56	256	218	18	17

Product Concentration	Contact Time	Na ^x		Na ^{x-1}		Na ^{x-2}		Na ^{x-3}	
		V _{C1}	V _{C2}	V _{C1}	V _{C2}	V _{C1}	V _{C2}	V _{C1}	V _{C2}
		0	0	0	0	0	0	0	0
100%	1 min	0	0	0	0	0	0	0	0
		0	0	0	0	0	0	0	0
100%	3 mins	0	0	0	0	0	0	0	0
		0	0	0	0	0	0	0	0

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RAW DATA

Test Method:	EN 14348:2005			
Product Name:	NOVA 104 (GLOBACID AF med)		Batch No.:	230457
Product Diluent:	Distilled water		Lab ID:	G007-23-007
Appearance of Product Dilutions:	Clear, colourless solution			
Inactivation:	Dilution-neutralization	Dilution Method:	Standard (80%)	Test Temperature (°C): 20
Neutralizer:	30 g/L Tween 80, 30 g/L Saponin, 3 g/L Lecithin			
Interfering Substance:	0.3 g/L bovine serum albumin			
Test Organism:	Mycobacterium avium ATCC 15769		Plating Method:	Spread plate
Incubation Temperature (°C):	30	Passing Criteria (lg):	4.00	Measurement Uncertainty (±): 0.11
Testing Period:	17/07/2023	Tested By:	AZZ	Verified By: CSE

Validation & Controls

Validation Suspension (N _v)	V _{C1}	V _{C2}	N _{v0} = 41.0 Limit: 30 ≤ N _{v0} ≤ 160
Validation Suspension (N _{VB})	V _{C1}	V _{C2}	N _{v0} = N _{VB} /1000 Limit: 30 ≤ N _{v0} ≤ 160
Experimental Conditions Control (A)	V _{C1}	V _{C2}	A = 41.5 Limit: A ≥ 0.5 x N _v /10
Neutralizer Control (B)	V _{C1}	V _{C2}	B = 24.5 Limit: B ≥ 0.5 x N _v /10 or N _{VB} /1000
Method Validation (C)	V _{C1}	V _{C2}	C = 29.0 Limit: C ≥ 0.5 x N _v /10
Conc.: 100%	27	31	

Test Suspension & Procedure

Test Suspension (N)		N	V _{C1}	V _{C2}	$\bar{x}_{wm} = N = 3.01E+09$ $N_0 = N/10$ $lg N_0 = 8.48$ Limit: $8.17 \leq lg N_0 \leq 8.70$			
		10 ⁻⁷	302	299				
		10 ⁻⁸	28	34				
Product Concentration	Contact Time	Dilution	V _{C1}	V _{C2}	Na = \bar{x} or $\bar{x}_{wm} \times 10$	lg Na	lg R = lg N ₀ - lg Na	Conformance Probability
100%	3 mins	10 ⁰	>660	>660	1.55E+04	4.19	4.29 ± 0.11	99.567%
		10 ⁻¹	158	144				
		10 ⁻²	17	22			99.995%	
		10 ⁻³	<14	<14				
		10 ⁰						
		10 ⁻¹						
		10 ⁻²						
		10 ⁻³						
		10 ⁰						
		10 ⁻¹						
		10 ⁻²						
		10 ⁻³						

Raw Data of Colony Count

	N _v		N _{VB}		A		B		C		N ⁻⁷		N ⁻⁸	
V _{C1}	20	20	-	-	23	20	10	11	17	10	154	148	13	15
V _{C2}	23	19	-	-	19	21	13	15	12	19	161	138	20	14
Product Concentration	Contact Time		Na ^x		Na ^{x-1}		Na ^{x-2}		Na ^{x-3}					
100%	3 mins		V _{C1}	V _{C2}	V _{C1}	V _{C2}	V _{C1}	V _{C2}	V _{C1}	V _{C2}	V _{C1}	V _{C2}	V _{C1}	V _{C2}
			>330	>330	78	75	9	10	0	0				
			>330	>330	80	69	8	12	0	0				

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TEST PROCEDURE

1. Test Na: Determination of Mycobactericidal Concentrations

- 1.1 1.0 mL of the interfering substance was pipetted into a tube. 1.0 mL of the test suspension N ($1.5 - 5.0 \times 10^9$ cfu/mL) was added to the tube.
- 1.2 The stopwatch was started immediately and the tube was mixed and placed in a water bath controlled at test temperature θ for 2 minutes \pm 10 seconds.
- 1.3 At the end of the 2 minutes, 8.0 mL of the product test solution was added to the tube. The stopwatch was restarted at the beginning of the addition. The tube was mixed and placed in a water bath controlled at θ for the contact time t . Just before the end of t , the tube was mixed again.
- 1.4 At the end of t , 1.0 mL sample of the test mixture Na was transferred into a tube containing 8.0 mL of neutralizer and 1.0 mL of distilled water. The neutralizer tube was mixed and placed in a water bath controlled at $(20 \pm 1)^\circ\text{C}$.
- 1.5 After a neutralization time of 5 minutes \pm 10 seconds, the neutralizer tube was mixed and 1.0 mL of the neutralized test mixture Na (containing neutralizer, product test solution, interfering substance, and test suspension) was taken in duplicate and inoculated using the spread plate technique.
- 1.6 Additionally, 0.5 mL of the neutralized test mixture Na was transferred into a tube containing 4.5 mL of neutralizer to obtain a 10^{-1} dilution of Na . The mixture was diluted accordingly in neutralizer to produce 10^{-2} and 10^{-3} dilutions of Na . 1.0 mL of each dilution was taken in duplicate and inoculated using the spread plate technique.
- 1.7 The procedure was performed using other product test solutions at the same time.

2. Experimental Conditions Control A: Verification of the Absence of Any Lethal Effect in the Experimental Conditions

- 2.1 1.0 mL of the interfering substance used in the test Na was pipetted into a tube. 1.0 mL of the validation suspension N_v ($0.3 - 1.6 \times 10^3$ cfu/mL) was added to the tube.
- 2.2 The stopwatch was started immediately and the tube was mixed and placed in a water bath controlled at test temperature θ for 2 minutes \pm 10 seconds.
- 2.3 At the end of the 2 minutes, 8.0 mL of hard water (distilled water for ready-to-use product) was added to the tube. The stopwatch was restarted at the beginning of the addition. The tube was mixed and placed in a water bath controlled at θ for the contact time t . Just before the end of t , the tube was mixed again.
- 2.4 At the end of t , 1.0 mL sample of the test mixture A was taken in duplicate and inoculated using the spread plate technique.

3. Neutralizer Control B: Verification of the Absence of Toxicity of the Neutralizer

- 3.1 8.0 mL of the neutralizer used in the test Na and 1.0 mL of distilled water were pipetted into a tube. 1.0 mL of the validation suspension N_v was added to the tube.
- 3.2 The stopwatch was started at the beginning of the addition and the tube was mixed and placed in a water bath controlled at $(20 \pm 1)^\circ\text{C}$ for 5 minutes \pm 10 seconds. Just before the end of this time, the tube was mixed.

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- 3.3 At the end of the time, 1.0 mL sample of the test mixture *B* was taken in duplicate and inoculated using the spread plate technique.
4. Method Validation C: Validation of the Dilution-Neutralization Method
- 4.1 1.0 mL of the interfering substance used in the test *Na* was pipetted into a tube. 1.0 mL of diluent was added and then, starting a stopwatch, 8.0 mL of the product test solution of the highest concentration used in the test *Na* was added to the tube. The tube was mixed and placed in a water bath controlled at test temperature θ for contact time t . Just before the end of t , the tube was mixed again.
- 4.2 At the end of t , 1.0 mL of the mixture was transferred into a tube containing 8.0 mL of neutralizer used in the test *Na*. The stopwatch was restarted at the beginning of the addition. The tube was mixed and placed in a water bath controlled at $(20 \pm 1) ^\circ\text{C}$ for 5 minutes \pm 10 seconds.
- 4.3 1.0 mL of the validation suspension N_V was added. The stopwatch was restarted at the beginning of the addition. The tube was mixed and placed in a water bath controlled at $(20 \pm 1) ^\circ\text{C}$ for (30 ± 1) minutes. Just before the end of this time, the tube was mixed again.
- 4.4 At the end of this time, 1.0 mL sample of the test mixture *C* was taken in duplicate and inoculated using the spread plate technique.
5. Incubation and Counting
- 5.1 The plates were incubated for 21 days. The plates were counted to determine the number of cfu. Any plates which were not countable for any reason were discarded.
- 5.2 For each plate, the exact number of colonies were noted but any counts higher than 330 colonies were recorded as '>330'.
- 5.3 All experimental data were reported as V_C values, in which a V_C value is the number of cfu counted per 1.0 mL sample inoculated.
- 5.4 Only V_C values within the counting limits, i.e., 14 to 330 colonies, were taken into account for further calculation, except in the case of *Na*.