

**PARTIAL TEST REPORT NO 623324/26/GDA/2**
Translation of the report NO 623324/26/GDA/1 of 13.07.2026

Client DNL Equity Partners Spółka z o.o ul. Świętojańska 87/3 81-389 Gdynia		Sample (according to declaration of Client) Sample description: Alchemy Élixir
Sample reception date	17.10.2024	Sample status: no objections Sample number: 648571/24/GDA Sample received from the Client
Start of analysis	18.10.2024	
End of analysis	24.10.2024	
Test report date	13.07.2026	

Test Method	Unit	Result
* Presence of Staphylococcus aureus in 1 g ²⁾ PN-EN ISO 22718:2016-01; PN-EN ISO 22718:2016-01/A1:2023-01	in 1 g	Absent
* Presence of Escherichia coli in 1 g ¹⁾ PN-EN ISO 21150:2016-01; PN-EN ISO 21150:2016-01/A1:2023-03	in 1 g	Absent
* Presence of Pseudomonas aeruginosa in 1 g ⁵⁾ PN-EN ISO 22717:2016-01; PN-EN ISO 22717:2016-01/A1:2023-03	in 1 g	Absent
* Presence of Candida albicans in 1 g ⁶⁾ PN-EN ISO 18416:2016-01; PN-EN ISO 18416:2016-01/A1:2023-03	in 1 g	Absent
* Aerobic mesophilic bacteria at 32,5°C ⁴⁾ PN-EN ISO 21149:2017-07; PN-EN ISO 21149:2017-07/A1:2023-01	cfu/g	4,0x10 ¹
* Number of yeasts and moulds at 25°C ³⁾ PN-EN ISO 16212:2017-08; PN-EN ISO 16212:2017-08/A1:2023-01	cfu/g	<1,0x10 ¹

- 1) Neutralizer: Eugon LT (1 g of the sample in 9 ml of the neutralizer)
Culture media: MacConkey Agar
Test strains: Escherichia coli ATCC 8739
Volume of calibrated suspension: 0.1 ml of the calibrated inoculum Nv - 1.0x10² - 5.0x10² cfu/g
Result of the neutralization: presence of microbial growth in the test mixture with the neutralizer and formulation with 0.1 ml inoculum Nv and absence of microbial growth in the test mixture with the neutralizer and formulation
The effectiveness of the neutralization has been confirmed.
- 2) Neutralizer: Eugon LT (1 g of the sample in 9 ml of the neutralizer)
Culture media: Baird Parker Agar
Test strains: Staphylococcus aureus ATCC 6538
Volume of calibrated suspension: 0.1 ml of the calibrated inoculum Nv - 1.0x10² - 5.0x10² cfu/g
Result of the neutralization: presence of microbial growth in the test mixture with the neutralizer and formulation with 0.1 ml inoculum Nv and absence of microbial growth in the test mixture with the neutralizer and formulation
The effectiveness of the neutralization has been confirmed.



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- 3) Neutralizer: Eugon LT (1 g of the sample in 9 ml of the neutralizer)
Culture media: SDCA (Sabouraud dextrose Agar with chloramphenicol)
Test strains: *Candida albicans* ATCC 10231
Volume of calibrated suspension: 1 ml of the calibrated inoculum $N_v - 1.0 \times 10^3 - 3.0 \times 10^3$ cfu/g
Result of the neutralization: $N_{vw} \geq 0,5 N_{vk}$, where: N_{vw} - number of microorganisms present in the test mixture with the neutralizer and formulation and N_{vk} - number of microorganisms present in the test mixture with the neutralizer in the absence of the formulation
The effectiveness of the neutralization has been confirmed.
- 4) Neutralizer: Eugon LT (1 g of the sample in 9 ml of the neutralizer)
Culture media: TSA (Tryptic Soy Agar)
Test strains: *Staphylococcus aureus* ATCC 6538, *Pseudomonas aeruginosa* ATCC 9027
Volume of calibrated suspension: 1 ml of the calibrated inoculum $N_v - 1.0 \times 10^3 - 3.0 \times 10^3$ cfu/g
Result of the neutralization: $N_{vw} \geq 0,5 N_{vk}$, where: N_{vw} - number of microorganisms present in the test mixture with the neutralizer and formulation and N_{vk} - number of microorganisms present in the test mixture with the neutralizer in the absence of the formulation
The effectiveness of the neutralization has been confirmed.
- 5) Neutralizer: Eugon LT (1 g of the sample in 9 ml of the neutralizer)
Culture media: Cetrimide Agar
Test strains: *Pseudomonas aeruginosa* ATCC 9027
Volume of calibrated suspension: 0.1 ml of the calibrated inoculum $N_v - 1.0 \times 10^2 - 5.0 \times 10^2$ cfu/g
Result of the neutralization: presence of microbial growth in the test mixture with the neutralizer and formulation with 0.1 ml inoculum N_v and absence of microbial growth in the test mixture with the neutralizer and formulation
The effectiveness of the neutralization has been confirmed.
- 6) Neutralizer: Eugon LT (1 g of the sample in 9 ml of the neutralizer)
Culture media: SDCA (Sabouraud dextrose Agar with chloramphenicol)
Test strains: *Candida albicans* ATCC 10231
Volume of calibrated suspension: 0.1 ml of the calibrated inoculum $N_v - 1.0 \times 10^2 - 5.0 \times 10^2$ cfu/g
Result of the neutralization: presence of microbial growth in the test mixture with the neutralizer and formulation with 0.1 ml inoculum N_v and absence of microbial growth in the test mixture with the neutralizer and formulation
The effectiveness of the neutralization has been confirmed.

Identification of the change: sample description

Test report authorized by:

ID: 175, Senior Analysis Specialist, Cosmetics Microbiology Laboratory

The test report bears the certified electronic seal of J.S. Hamilton Poland Sp. z o.o.

Laboratory address:

Goździków 1, 43-100 Tychy

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* Test method accredited

Test performed by external provider

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