

AB-1804-T

BT-2024-003835

06.24

## ANALYSIS REPORT

<b>Purpose of Analysis</b> : Private Request	<b>Report Number</b> : BT-2024-003835
<b>Sample requested by:</b>	<b>Date and Time of Report</b> : 28.06.2024 14:42
<b>Sample Detail:</b>	
<b>Name</b> : EVLY PHARMA COSMETICS-The Purest Solutions	<b>Name</b> : Peptide Complex
<b>Address</b> : Şerifali Mah. Söyleşi Sok. No:41/A Ümraniye /İSTANBUL	<b>Qty/Pcs - Temp. (C)</b> : 10 g
<b>Authorized Person</b> :	<b>Packing</b> : Company Packaging
<b>Phone/Fax</b> :	<b>Date of Prod./Exp.</b> : 09/2023 - 09/2026
<b>Sender</b> : EVLY PHARMA COSMETICS-The Purest Solutions	<b>Lot Number</b> : - 008
<b>Manufacturer</b> : EVLY PHARMA COSMETICS-The Purest	<b>Brand</b> :
<b>Offerr No</b> :	<b>Date Received</b> : 25.06.2024
	<b>Date Started</b> : 25.06.2024
	<b>Date Finished</b> : 28.06.2024

## RESULT

Name of Analysis	Result	Unit	U	Rec.	LOQ	LVS	D.R.	Reference Ranges	Method/s	Conformity
Arsenic	<0,01	mg/kg			<0,01	2		< 5	In House Metot	Passed
Lead	<0,01	mg/kg			<0,01	2		< 20	In House Metot	Passed
Cadmium	<0,01	mg/kg			<0,01	2		< 5	In House Metot	Passed
Mercury	<0,01	mg/kg			<0,01	2		< 1	In House Metot	Passed
Antimony	<0,01	mg/kg			<0,01	2		< 10	In House Metot	Passed
Color and Physical Condition Detection *	Suitable.	-				1		It should have its own characteristics	TS EN ISO 5492	Passed

### DESCRIPTION

#### DECISION RULE (D.R.)

:

#### Limit Value Source (LVS)

1 - Evaluated according to product specification.

2 - Compliance has been evaluated according to the Pharmaceuticals and Medical Devices Agency Guideline on Heavy Metal Impurities in Cosmetic Products.

#### U. Uncertainty of Measurement

#### Rec. Recovery

#### LOQ. Limit of Quantification

- When the conformity assessment regarding the test results is given, the regulations, standards, specifications, contracts, etc., if any. The decision rule specified in the documents is used. If there is no decision rule specified in the legislation, the Simple Decision Rule is applied without considering the measurement uncertainty.

-The uncertainties specified in the report are k=2, expanded uncertainty at the 95% confidence interval.

-The results are valid as the sample is received and we are not responsible for the sampling phase. The laboratory cannot be held responsible for the information given by the customer.

#### REVISION INFORMATION

:

\*\*\* Analysis marked with \*\*\* are within the scope of accreditation.

1. BiYOTEST Laboratory and Consulting Services Ltd., which operates as an Analysis laboratory. Şti. is accredited by TURKAK according to AB-1804-T and TS EN ISO/IEC 17025 standard. The Turkish Accreditation Agency (TURKAK) has signed a Multilateral Agreement with the European Accreditation Association (EA) and a mutual recognition agreement with the International Laboratory Accreditation Association (ILAC) on the recognition of test reports.

2. The results of the Analysis are valid for the above-mentioned sample sent to the laboratory by the company/institution/individual.

3. Descriptive information in the test report that affects the validity of the results has been declared by the customer. Our laboratory is not responsible for any losses/legal obligations that may occur due to the accuracy and use of this information.

4. No part of this test report can be used alone or separately, can not be copied, reproduced or published in whole or in part without the written permission of the laboratory.

5. This report cannot be used for advertising purposes, unsigned and unsealed reports are invalid.

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P03-T08-F01/

Rev.02/15.12.2022

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**Tuğba ÖZKAN**

**Head of Sample Submission and  
Reporting Department**

BIYOTEST LABORATUVARLARI VE  
DANIŞMANLIK HİZMETLERİ LTD. ŞTİ.  
Gümüşpala Mah. Kaynata Sok. No:2 Kat:6  
Avcılar / İSTANBUL  
Avciilar V.D. 781549767

**Approved**  
**28.06.2024 14:42**

**Sema YUMAK**  
**Biologist**  
**Manager Of Laboratory**

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