

## **REPORT NO 273/10/2022**

Cus	stomer <sup>1</sup> :			BLUE	TARGET LTD SPÓŁKA K Górska 15, 84-230		AWG				
	stomer order no. <sup>1</sup> : applicable)				no data						
Order no.: 357/09/			/2022	/2022 Order date:			15.09.2022				
San	Sample type <sup>1</sup> : Diet supplement				Sample code:		41/15/09/22				
	poratory name and dress:	Ekolabos sp. z o.o. ul. Duńska 9, 54-427 Wrocław									
Tes	sting purpose <sup>1</sup> :		Customer's own needs								
Tes	st initiation date:	15.09	2022	022 Test end date: 06.10.2022							
Rep	porting date:		06.10.2022								
				San	npling						
Sampling person <sup>1</sup> : Object taken and o Custor				by the	Sampling method <sup>1</sup> :	No data					
San	npling location <sup>1</sup> :	No data			Sampling point <sup>1</sup> :		No data				
Sample register date <sup>1</sup> : 15.0			2022 Sample condition: Unreservedly								
Sampling protocol:			No data								
				Sample	description						
San	nple name:				APOLLOS HEGEMONY Alpha GPC 90 caps						
LOT	Т:		08/2025								
Pro	oduction date:	No data									
Expiration date:			31.08.2025								
Pac	ckaging type:					original					
Ma	terial:					No data					
Dat	Date of receipt of the sample:			15.09.2022							
			Tes	ts done in	the laboratory						
No.	Tested parameter	Test method			Result [±uncertainty]	Unit	Authorizing person	The highest allowable value or range	Statement of assesment		
1	Detection od Salmonella spp.	PN-EN ISO 6579-1:2017-04+ A1:2020-09		NA	not detected in 25g	-	BD	-	-		
2	Total number of microorganisms	s PN-EN ISO 4833-1:2013- 12/Ap1:2016-11E		NA	40	cfu/1 g	BD	-	-		
3	Enumeration of Enterobacteriaceae	PN-ISO 21528-2:2017-08		NA	<10	cfu/1 g	BD	-	-		
4	Enumeration of yeasts and mould	PN-ISO 21527-1:2009		NA	<10	cfu/1 g	BD	-	-		
5	Detection of Staphylococcus aureus	PN-EN ISO 6888- 3:2004/AC:2005		NA	not detected in 1g	-	BD	-	-		
6	Detection of Listeria monocytogenes	PN-EN ISO 11290-1:2017-07		NA	not detected in 25g	-	BD	not detected in 25g	-		
7	Enumeration of Escherichia coli B-glucuronidase positive	li PN-ISO 16649-2:2004		NA	<10	cfu/1 g	BD	-	-		
8	Mercury	PN-EN 15763:2010		A P1	<0,001	mg/kg	P1	-	-		
9	Lead	PN-EN 15763	:2010	A P1	0,024 [+/-0,004]	mg/kg	P1	-	-		

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Signed: Qualified Electronic Signature (QES) Mateusz Latosiński Ekolabos



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## PO-11/Z-03 issue 3th dated 16.05.2022

	Tests done in the laboratory													
No.	Tested parameter	Test method		Result [±uncertainty]	Unit	Authorizing person	The highest allowable value or range	Statement of assesment						
10	Cadmium	PN-EN 15763:2010	A P1	<0,002	mg/kg	P1	-	-						
11	Arsenic	PN-EN 15763:2010	A P1	<0,010	mg/kg	P1	-	-						

## Legend/Explanations:

Employees authorizing test results:

BD - Bartłomiej Dudek

P1 - testing performed by an external service provider AB 1095

Testing methods marked with a symbol: A - accredited tests, NA - non-accredited tests, S- tests approved by PPIS in Wrocław, decision 5169/21 of 18.10.2021 r. and decision 1824/22 of 22.03.2022, R- the reference method may, if the law so provides, be used to statement of conformity in a regulated area, EP - tests performed by an external service provider.

(W) standard removed from the Polish Standards catalog without replacement.

\* Maximum acceptable value or range in order to COMMISSION REGULATION (EC) No 2073/2005 of 15 November 2005 on microbiological criteria for foodstuffs 'Data declared by the Customer, information obtained from the Customer may affect the validity of the result

In case of sampling and delivery of the sample by the Client, the Laboratory is not responsible for the process of sampling and transportation of the samples for testing.

Results marked with a below (<) sign and an above (>) sign mean that the results lie below and above the range.

The test results apply only to the sample received and tested.

NZ result a statement of nonconformity, ZG result a statement of conformity.

The laboratory provides uncertainty: at the customer's request, when conformity to requirements is evaluated, when it is relevant to the validity and applicability of the test results. The laboratory reports the uncertainty as expanded uncertainty with a confidence level of 95% and a coverage factor of k=2. Where the sample is provided by the customer, the uncertainty does not include the sampling.

Uncertainty of testing and statement of conformity was agreed with the customer at the ordering.

The Laboratory is responsible for all information in the Test Report, except for information provided by the Client.

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The original Test Report is issued in electronic form with the extension\*pdf, signed with a qualified electronic signature. Therefore, all hard copies, unless certified as true copies of the original, are copies.

Complaints must be received no more than 14 days from the date of mailing of the Test Report. Complaints may be received at: biuro@ekolabos.pl.

Reported by:

Elżbieta Świerczek - Płuska Environmental Project Specialist / Cosmetic Product Research Specialist **Report approved by:** Mateusz Latosiński Customer Services Manager / Key Account Specialist

---END OF THE REPORT---

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