

ANALYTICAL LABORATORIES

microbiology - physicochemistry - sensory



GBA POLSKA Sp. z o.o. Member of GBA GROUP ul. Mochtyńska 65, 03-289 Warsaw, Poland

TEST REPORT No: B/0/08/2024/532/FM/3/EN

Customer:

7FIT s.a. 55-200 Stanowice, ul. Stanowice 28A

Order No:

B/0/08/2024/532 A - accredited methodology (accreditation no. AB 1095); reference - if the law so provides (the result can be used to assess compliance in the legally regulated area). AE - accredited methodology (accreditation no. AB 1095) of flexible scope – reference if the law so provides / equivalent to reference (the result can be used to assess compliance in the legally regulated area).

NA -	legally regulated area). non-accredited methodology, covered by	the PN-E	N ISO/IEC 1	7025:2018-02 system						
Materia	al/product tested: Dietary su	pplemei	nts							
Sample	collection address:	5	5-200 Stan	owice, Stanowice 82A						
Product name: 7NUTRITION OMEGA 3 100 softgels Date*: 23 August 2024										
Producer: 7NUTRITION										
Date of	production:	1	9/08/2024	9/08/2024						
Lot num	ber:	1	90824.1							
	g according to: - transported by: Shipping				Received by: GB.	A POLSKA emplo	oyee no.: 272	29		
Sample	no: 36409/08/24 Sample condition:	c	correct	Analysis start da	te: 23-08-2024 Analys	is end date:	31-08-2024	4		
Lab.	Analyzed parameter	Unit	Accred.	Test method	Requirement	Result	U	S		
Р	Total microbial count	cfu/g	AE	PN-EN ISO 4833-1:2013-12, PN- EN ISO 4833-1:2013-12/Ap1:2016- 11, PN-EN ISO 4833-1:2013- 12/A1:2022-06	no requirements	<1,0x101		-		
Р	Presence of Salmonella spp.	25g	AE	PN-EN ISO 6579-1:2017-04, PN- EN ISO 6579-1:2017-04/A1:2020- 09	no requirements	not detected in 25g		-		
Р	Presence of presumptive Escherichia coli	1g	AE	PN-ISO 7251:2006	no requirements	absent in 1g		-		
Р	Presence of coagulase-positive staphylococci (Staphylococcus aureus and other species)	lg	AE	PN-EN ISO 6888-3:2004, PN-EN ISO 6888-3:2004/AC:2005	no requirements	absent in 1g		-		
Р	Count of yeasts and moulds	cfu/g	AE	PN-ISO 7954:1999	no requirements	<1,0x10 ¹		-		
Ł	Lead	mg/kg	AE	PN-EN 15763:2010	≤3.0 ; mg/kg ; Start 2023/915	< 0,010	0,002	CONFORMING		
Ł	Mercury	mg/kg	AE	PN-EN 15763:2010	≤0.1 ; Start 2023/915	< 0,0010	0,0002	CONFORMING		
Ł	Cadmium	mg/kg	AE	PN-EN 15763:2010	no requirements	< 0,0020	0,0003	-		

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Lab.	Analyzed parameter	Unit	Accred.	Test method	Requirement	Result	U	S
Ł	An anisidine number	-	A	PN-EN ISO 6885:2016-04	no requirements	7,09	1,42	-
Ł	Peroxide value	meq O2/kg	А	PB-72/LF ed. 6 of 03.01.2022	no requirements	6,36	0,64	-
Ł	TOTOX indicator	-	NA	PN-EN ISO 6885:2016-04	no requirements	19,8		-

Date* - depending on the method of obtaining the sample by GBA POLSKA, it is the date of: collection (when the sample is collected only by a GBA POLSKA employee) or receipt (when the sample is collected from the

Date ' dependent of the method of obtaining the sample by OBA POLSKA, it is the due of: confected only by a CBA POLSKA employee) of receipt (when the sample is confected from the Customer by a GBA POLSKA employee) of receipt (when the sample is confected from the U - expanded measurement uncertainty at the level of confidence app. 95% and the coverage factor k=2, does not take into account the sampling uncertainty, except when indicated in the remarks. Measurement uncertainty is provided when it is important for the reliability of test results or compliance with requirements/specifications and at the request of the Customer. The 'test results' lower or higher than the measuring ranges of the methods are presented as "value of the lower limit of the measuring range or "> value of the upper limit of the measuring range", respectively. These values provide information about the research results. If expanded uncertainties are given with these test results, they apply to the lower or upper limit of the measuring range of the method. S – Statements of Conformity with the requirements or specifications relating to the results for the parameters indicated in a given row, where CONFORMING means conformity and NON CONFORMING means non-conformity with specification. The devicing must be customer and the risk associated with it as well as the identification of which specifications. The are methods are post are post-

conformity with specification. The decision rules agreed with the Customer and the risks associated with it, as well as the identification of which specifications, standards or parts thereof are met and which are not, are provided in the Remarks. In case of obtaining the "test results", the Statements of Conformity for those "test results" that are meet the requirements of PCA Communication No. 353 of August 24, 2021, it is carried out as part of the opinion and interpretation.

The results refer only to the tested samples (sampled or received - in accordance with the information presented in the Test Report). The information in italics included in the Test Report was provided by the Customer. The laboratory is not responsible for this information. The laboratory is not responsible for the method of sampling and the representativeness of the samples provided by the Customer for testing. The Test Report without the written approval of the Laboratory shall not be reproduced except in full.

The Laboratory does not store the samples after testing, unless otherwise agreed with the Customer. Place of performance of the tests ("Lab."): Ł - Łajski, ul. Kościelna 2a, 05-119 Legionowo, L - ul. Doświadczalna 50a, 20-280 Lublin, M - ul. Fabryczna 7, 41-404 Mysłowice, P - ul. Kazimierza Tymienieckiego 34, 60-681 Poznań, PS - in situ measurement.

NOTE: Original Test Report are issued in electronic form with the *.pdf extension, signed with a qualified electronic signature. Therefore, all prints, unless certified as true copies, are copies,

Remarks:

The tested sample meets the requirements indicated above as "conforming" in terms of the tested parameters.

In determining Statement of Conformity, the principle of simple acceptance described in the guidelines of document ILAC-G8-09/2019 has been applied. For results close to the tolerance/specification limit, the risk of false acceptance is up to 50%.

The second selective medium for detecting the presence of Salmonella spp. in accordance with PN-EN ISO 6579-1:2017-04, Mon-EN ISO 6579-1:2017-04/A1:2020-09 is RVS broth and Brilliance Salmonella/Agar. Coagulase is used to detect staphylococci-positive, Braid Parker RPF/agar was used.

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Created on:	Authorized result:	Authorized Test report:	
02-09-2024	GBA POLSKA employee no.: 2424 GBA POLSKA employee no.: 2642	Specialist in food and dietary supplements	Signed with a qualified electronic signature
	GBA POLSKA employee no.: 2813 GBA POLSKA employee no.: 2866	GBA POLSKA employee no: 2793	

Report prepared in a single copy

The end of the Test Report