

VitaMoment GmbH

Barnerstraße 14 d
22765 Hamburg



Our Sign : RRu
Date : 03.06.2026

Certificate of analysis 25035873 - 001

Sample name : Prosta-Komplex

Marking of sample : Charge 77461, 31.05.2028

Customer No. : none

Packaging : Commercial package/plastic vessel

Sample amount : 6 x 33,28 g

Shipping of sample : Courier Service

Sample entry : 24.07.2025

Entrance temperature : Room temperature

Sample taken : by sender

Begin/end of analysis : 24.07.2025 / 07.08.2025

The test results apply only to the test items described in the report. No responsibility is accepted for the validity of the results if any data or information provided by the customer may affect them. Data provided by the customer are clearly identified. The laboratory assumes no responsibility for the sampling including minimum quantities unless it was carried out by samplers from a company within the GBA Group or on its behalf. In this case, the results apply to the sample as received. The test report may not be published or reproduced, in whole or in part, without the written consent of the issuing company. The general terms and conditions are available at <https://www.gba-group.com/en/general-terms-and-conditions/>.

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Dok.-Nr.: ML 510-01 # 2 V1 E, 511, 19.02.2026

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seit 1989

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Test Results

Microbiological Test	Result	Unit
Total Plate Count	<10	cfu/g
Yeasts / moulds		
Yeasts	<10	cfu/g
Moulds	<10	cfu/g
Enterobacteriaceae	<10	cfu/g
E. coli	<10	cfu/g
Salmonella	negative	/ 25 g

Chemical/Physical Test	Result	Unit	Declaration	± MU	MU[%]	MU Source	ML
Lead	0,032	mg/kg		0,0064	20	I	3
Cadmium	0,011	mg/kg		0,0022	20	I	1
Mercury	<0,010	mg/kg			25	I	0,1
Arsenic	<0,040	mg/kg			20	I	
PAH							
Benzo(a)anthracene	<1,0	µg/kg			30	I	
Chrysen	<1,0	µg/kg			30	I	
Benzo(b)fluoranthene	<1,0	µg/kg			30	I	
Benzo(a)pyrene	<1,0	µg/kg			30	I	10
Sum of PAH	not detectable	µg/kg			30	I	50
Biotin	63,1	µg/ daily serving	50	16	25	I	
Selenium	109	µg/ daily serving	80	22	20	I	
Zinc	13,2	mg/ daily serving	10	2,6	20	I	
Weight per dosage form	0,56	g		0,0056	1	VII	
Daily serving	2,0	capsule(s)					
Iodine	0,062	mg/100 g		0,015	24		
Amino acid spectrum, free							
Histidine	15,7	mg/ daily serving	10	4,7	30	I	

Maximum levels for food supplements according to VO (EU) 2023/915

Assessment:

Results of microbiological analysis are unobjectionable regarding the tests carried out.

The product meets the requirements of Regulation (EU) 2023/915 regarding maximum levels of heavy metals and polycyclic aromatic hydrocarbons (PAH) in food supplements.

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Results of vitamins and minerals analyses meet the details of the nutrition labelling regarding the tests carried out (cf. Guidance document of the EU Commission with regard to the setting of tolerances for nutrient values declared on a label in food supplements from December 2012).

The measured content of Histidine corresponds to the declared value with sufficient accuracy.

Hamburg, 03.06.2026

This test report is done automatically and is valid without signature.

Methods

Parameter	Method	DR
Total Plate Count	DIN EN ISO 4833-2: 2022-05 ^a ₀	m
Yeasts / moulds	BIOKAR Diagnostics, Symphony-Agar BM20208/BM19108: 2022-11 ^a ; validated according to EN ISO 16140-2 against EN ISO 21527-1/-2 2008-11 ₀	m
Enterobacteriaceae	Biomerieux, Rebecca-Agar AEB520020/AEB150022: 2020-09 ^a ; validated according to EN ISO 16140-2 against ISO 21528-2 2017-07 ₀	m
E. coli	Biomerieux, Rebecca-Agar AEB520020/AEB150022: 2020-09 ^a ; validated according to EN ISO 16140-2 against ISO 16649-2 2001-07 ₀	m
Salmonella	DIN EN ISO 6579-1: 2020-08 ^a ₀	m
Lead	DIN EN 15763, ICP-MS: 2010-04 ^a ₅	y
Cadmium	DIN EN 15763, ICP-MS: 2010-04 ^a ₅	y
Mercury	DIN EN 15763, ICP-MS: 2010-04 ^a ₅	y
Arsenic	DIN EN 15763, ICP-MS: 2010-04 ^a ₅	y
Pressure pulping	§ 64 LFGB L 00.00-19/1: 2015-06 ^a ₀	q
PAH	HH-MA-M 02-105 # U, HPLC-FLD: 2023-06 ^a ₀	y
Sum of PAH	berechnet α	
Biotin	HH-MA-M 02-160, LC-MS/MS: 2024-03 ^a ₀	z
Selenium	DIN EN 15763, mod., ICP-MS: 2010-04 ^a ₅	y
Zinc	DIN EN 15763, mod., ICP-MS: 2010-04 ^a ₅	y
Weight per dosage form	HH-MA-M 10-030, gravimetric: 2021-11 ^a ₀	z
Iodine	PNTe/LQM/FYQ/316: 2025-05 ^a ₅₈	z
Amino acid spectrum, free	HH-MA-M 02-183, LC-MS/MS: 2025-07 ^a ₀	y

The methods marked with ^a are accredited methods of the performing laboratory.

Testing laboratory: ₀GBA Hamburg ₅GBA Pinneberg α automatically calculated from the system ₅₈LQM

MU-Source:

I: According to DIN ISO 11352 as expanded, combined measurement uncertainty with $k = 2$ (95 %), sampling not included

VII: According to expert estimation

Decision rules:

m: The conformity assessment of microbiological measured values is performed without considering additional analytical measurands.

y: In conformity assessment, measurement uncertainty is disregarded for measured values below the tolerance limit. For measured

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values exceeding the tolerance limit, measurement uncertainty is subtracted from the measured value. If no conformity assessment is performed, measurement uncertainty serves as informational data only.

q: The conformity assessment of qualitative measurement values (positive/negative, conforms/ does not conform) is performed without considering additional analytical measurands.

z: In conformity assessment, measurement uncertainty is disregarded and serves as informational data only.