



TEST REPORT NO 420119/24/GDY

Client NADARRA GmbH Alsterkamp 19 20149 Hamburg		Sample (according to declaration of Client) Sample description: Vitamin D 1000 IU Nadarra, 15 ml Batch: D1000N/25042024 Production date: 25.04.2024 Expiry date: 25.10.2025		
Sample reception date:	14.05.2024	Sample status: no objections		
Start of analysis	24.05.2024	Sample number: 284588/24/GDY/Z1		
End of analysis	31.05.2024	Sample received from the Client		
Test report date	10.07.2024			
Test Method	Unit	Result	Criteria	Statement of conformity
* Vitamin D3 ^{1) 2) 3)} PN-EN 12821:2009				
Vitamin D3 (cholecalciferol)	µg/dose	29,9 ± 4,5	-	-
	IU/dose	1190 ± 180	1000 +50%/-20%	Pass

1) Dose declared by the Client: 0,031 g.

2) Guidance Document for competent authorities for the control of compliance with EU legislation on: Regulation (EU) No 1169/2011 of the European Parliament and of the Council of 25 October 2011 on the provision of food information to consumers, amending Regulations (EC) No 1924/2006 and (EC) No 1925/2006 of the European Parliament and of the Council, and repealing Commission Directive 87/250/EEC, Council Directive 90/496/EEC, Commission Directive 1999/10/EC, Directive 2000/13/EC of the European Parliament and of the Council, Commission Directives 2002/67/EC and 2008/5/EC and Commission Regulation (EC) No 608/2004 and Council Directive 90/496/EEC of 24 September 1990 on nutrition labelling of foodstuffs and Directive 2002/46/EC of the European Parliament and of the Council of 10 June 2002 on the approximation of the laws of the Member States relating to food supplements with regard to the setting of tolerances for nutrient values declared on a label, December 2012. Table 2.

3) Client specification.

Identification of the change: sample description

Authorized by:

ID: 127, Analysis Expert, Vitamin Analysis Laboratory

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

Laboratory address:

Chwaszczyńska 180, 81-571 Gdynia

The results refer only to the samples received. When a measurement uncertainty is given, it is an expanded uncertainty estimated for a coverage factor $k=2$ at 95% confidence level and is not including sampling uncertainty, unless otherwise stated. When the conformity is stated J.S. Hamilton Poland Sp. z o.o. applies the simple acceptance decision rule in accordance with ILAC-G8:09/2019, unless otherwise reported. If the "result" column of the accredited method contains a record: "<" or ">", it means, that it is the test outcome directly related to the lower or upper limit of the measuring range of the accredited method, whereas the given expanded measurement uncertainty relates only to the lower or upper limit of the measuring range of the accredited method respectively. In such a case, the Laboratory presents the opinion and interpretation in the "statement of conformity" column, which is based on the obtained test outcome. This test report may not be copied in part without the prior written permission of J.S. Hamilton Poland Sp. z o.o. The responsibility of J.S. Hamilton Poland Sp. z o.o. is limited solely to the data issued in its original. J.S. Hamilton Poland Sp. z o.o. does not permit the use of the PCA accreditation symbol AB 079 by customers, subcontractors, external service providers and other third parties. For further information please refer to the PCA document DA-02. The service confirmed by this report is subject to the General Terms and Conditions of Services of J.S. Hamilton Poland Sp. z o.o. published on www.hamilton.com.pl.

* Test method accredited

Test performed by external provider

THE END OF THE REPORT