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AB 1334

Hero Pro LTD
 Office 312B
 High Street North
 182-184 London E6 2JA
 WIELKA BRYTANIA

Issue date 31.12.2024



Analytical report AR-24-RE-136777-01

Sample code 122-2024-00243348

* Type of sample	Apatil Max - 60vcaps
* Prescriber	Hero Pro LTD
* Purchase order date	17.12.2024
* Client Purchase order nr.	APCiM
Reception date	20.12.2024
Transport by	Courier
Sample condition	acceptable
Transport condition	at ambient temp.
* Date of sampling	17.12.2024
* Sampling Person	Principal
* Type of sampling	no data
* Purpose of the testing	not specified
* Best before date	12.2026
* Batch number	APM02122024
Number of tested samples	1
Start analysis	23.12.2024
End Analysis	31.12.2024

Results / Outcomes

UM276	Aerobic plate count, Pour plate technique (A)		
Method	PN-EN ISO 4833-1:2013-12+A1:2022-06, Total aerobic bacteria 30°C <10>30000000 /g (1-5) PC-Guss-ISO 4833, E-Cultural technique (non-chromogenic media)		
Aerobic Plate Count 30°C	< 10	cfu/g	
UM2PF	The presence of Salmonella spp., Breeding method with biochemical and serological confirmation (A)		
Method	PN-EN ISO 6579-1:2017-04+A1:2020-09, D-Cultural technique (non-chromogenic media)		
Salmonella	Not Detected	/25 g	
UMIMW	Coagulase positive Staphylococcus D Abs Pres /1 g ISO 6888-3 (A)		
Method	PN-EN ISO 6888-3:2004+AC:2005, D-Cultural technique (MPN tubes)		
Coagulase positive Staphylococcus	Not Detected	/1 g	
UMLS5	The presence of presumptive Escherichia coli, Breeding method with biochemical confirmation (A)		

Method PN-ISO 7251:2006, D-Cultural technique (non-chromogenic media)
Escherichia Coli Not Detected /1 g

ZM02A Yeasts & Moulds E [PB/MBK/40] <10 >1 500 000 /g (1-4) PYM-PF 3M™ Petrifilm™ Rapid Yeast and Mold Cou
(A)

Method PB/MBK/40 issue 01 of 13.12.2017
Yeasts & Moulds < 10 cfu/g

A = Method accredited

x = Data provided by the customer



Authorized by:
Agnieszka Siemińska - Senior Assistant

Approved by Magdalena Matysiewicz
Analytical Service Manager

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AB 1334

Hero Pro LTD
Office 312B
High Street North
182-184 London E6 2JA
WIELKA BRYTANIA

Issue date 27.12.2024



Analytical report AR-24-RE-136399-01

Sample code 122-2024-00243350

x Type of sample	Apatil Max - 60vcaps
x Prescriber	Hero Pro LTD
x Purchase order date	17.12.2024
x Client Purchase order nr.	APCiM
Reception date	20.12.2024
Transport by	Courier
Sample condition	acceptable
Transport condition	at ambient temp.
x Date of sampling	17.12.2024
x Sampling Person	Principal
x Type of sampling	no data
x Purpose of the testing	not specified
x Batch number	APM02122024
Number of tested samples	1
Start analysis	20.12.2024
End Analysis	27.12.2024

Results / Outcomes

E8006 Cadmium content (A)

Method PN-EN 15763:2010, Inductively coupled plasma ionization mass spectrometry method (ICP-MS)
Eurofins Polska SP. z o.o. (Łódź), accreditation nr. AB 1334

Cadmium content	0,006	mg/kg	± 0,002
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E8007 Arsenic content (A)

Method PN-EN 15763:2010, Inductively coupled plasma ionization mass spectrometry method (ICP-MS)
Eurofins Polska SP. z o.o. (Łódź), accreditation nr. AB 1334

Arsenic content	<0,005	* mg/kg	0.005 ± 0.001
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E8008 Lead content (A)

Method PN-EN 15763:2010, Inductively coupled plasma ionization mass spectrometry method (ICP-MS)

Eurofins Polska SP. z o.o. (Łódź), accreditation nr. AB 1334

Lead content 0,026 mg/kg $\pm 0,007$

E8009 Mercury content (A)

Method PN-EN 15763:2010, Inductively coupled plasma ionization mass spectrometry method (ICP-MS)

Eurofins Polska SP. z o.o. (Łódź), accreditation nr. AB 1334

Mercury content <0,005 * mg/kg 0.005 ± 0.001

* = less than the limit of quantification

A = Method accredited

+/- Uncertainty of measurement presented as expanded uncertainty of measurement (95%; k=2). For test results given in the form „<or>”, the given uncertainty of measurement applies only to the lower or upper limit of the accredited measuring range, respectively.

x = Data provided by the customer



Authorized by:
Adrianna Syrowa - Laboratory Technician - Chemical Department

Approved by Aneta Zawila
Junior Analytical Service Manager

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TEST REPORT NO 798495/24/GDY

Client HERO PRO LTD Office 312b, 182 -184 High Street North E6 2JA London		Sample (according to declaration of Client) Sample description: Apatil Max - 60vcaps Batch: APM02122024 Production date: 01.12.2024 Expiry date: 31.12.2026
Sample reception date:	20.12.2024	Sample status: no objections Sample received from the Client
Start of analysis	21.12.2024	
End of analysis	30.12.2024	
Test report date	30.12.2024	

Test Method	Unit	Result
* Content of elements ¹⁾ PN-EN 15763:2010		
Lead (Pb)	mg/kg	0,026 ± 0,007
Arsenic (As)	mg/kg	< 0,010 (0,010 ± 0,002)
Cadmium (Cd)	mg/kg	0,0068 ± 0,0016
Mercury (Hg)	mg/kg	< 0,0010 (0,0010 ± 0,0002)
* Aerobic colony count at 30°C PN-EN ISO 4833-1:2013-12; PN-EN ISO 4833-1:2013-12/A1:2022-06	cfu/g	<1,0x10 ¹
* Number of yeasts and moulds at 25°C PN-ISO 7954:1999 (withdrawn)	cfu/g	<1,0x10 ¹
* Presence of coagulase-positive staphylococci (Staphylococcus aureus and other species) in 1 g PN-EN ISO 6888-3:2004; PN-EN ISO 6888-3:2004/AC:2005	in 1 g	Not detected
* Presence of Escherichia coli in 1 g PN-ISO 7251:2006	in 1 g	Not detected
* Presence of Salmonella spp. in 25 g PN-EN ISO 6579-1:2017-04; PN-EN ISO 6579-1:2017-04/A1:2020-09	in 25 g	Not detected
* Presence of Listeria monocytogenes in 25 g PN-EN ISO 11290-1:2017-07	in 25 g	Not detected

¹⁾ The lower limit of the measuring range of the accredited method, which is also the limit of quantification set by the Laboratory.

Authorized by:

ID: 106, Analysis Expert, Microbiology Laboratory

ID: 295, Analysis Expert, Spectrometry Laboratory

ID: 368, Analysis Expert, Microbiology 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



HAMILTON



AB 079

TEST REPORT NO 798495/24/GDY

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* Test method accredited

Test performed by external provider

THE END OF THE REPORT