

Certificate of Analysis

Sample ID: 200603-004-003

Retention Sample ID: 5,847

Client ID: JCB Health Ltd

Client Address: 63 Craysfort Avenue, Blackrock, Dublin

 Batch:
 Milagro CBD 15%

 Date Receipt:
 2023/05/04 08:49:15

 Date Report Released:
 2023/05/08 12:23:54

 Sample Type:
 Liquid

Sample Type:

Package Condition:

Received By:

Sample Amount:

Tamper Proof:

Liquid
Intact
PMYBURGH
1 gram
No



Alexander Wrbka 2023/05/05 14:45:50

Test Name: Potency Cannabis(HPLC) (200603-004-003)

Specification: USP Specification (Oral)

Test Method: *TM006.3 Sampling Method: SH007

Analyte	Results		*Std Dev.	*Safety Limits	
CBC	ND	% Weight	% Weight	N/A	
CBD		% Weight	+- 0.441% Weight	N/A N/A	
CBD-A		% Weight	+- 0.120 % Weight	N/A	
CBD-V	ND	% Weight	% Weight	N/A	
CBG	ND	% Weight	% Weight	N/A	
CBG-A	ND	% Weight	% Weight	N/A	
CBN	ND	% Weight	% Weight	N/A	
Delta-9-THC	ND	% Weight	% Weight	N/A	
THC-A	ND	% Weight	% Weight	N/A	
THC-V	ND	% Weight	% Weight	N/A	
Total Potential CBD	18.209	% Weight	+- 0.547 % Weight	N/A	
Total Potential THC	ND	% Weight	% Weight	N/A	

Total Potential CBD

Total Potential CBD

18.21

CBC
0.00
CBG
0.00
CBGA
0.00
CBN
0.00
Total Potential CBD 18.21
Total Potential THC 0.00
Total: 18.21

Analyst:

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Analysed according to ISO 17025 Standards Email: info@NAFS.co.za Tel: 087 654 6273



*ND (Not Detected)

*Standard Deviation (Uncertainty of measurement of applicable duplicate sample

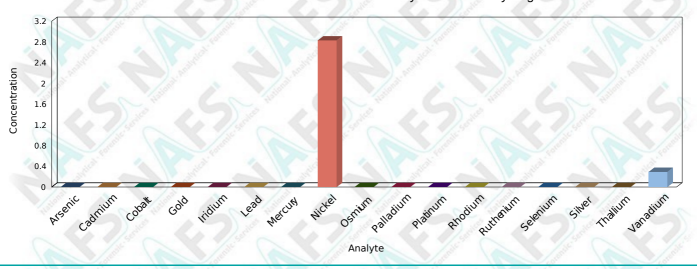
*Safety Limits (Regulatory limits specified are according to the United States pharmacopeia where applicab)e

Test Name: Heavy Metal Class 1,2 Analysis (200603-004-008)

Specification: USP Specification (Oral)

Test Method: *TM005.2 Sampling Method: SH007

Analyte	Re	sults	*Std	Dev.	*Safet	y Limits
Silver	ND	ppm		ppm	< 15	ppm PASS
Arsenic	ND	ppm		ppm	< 0.1	ppmPASS
Gold	ND	ppm		ppm	< 10	ppm PASS
Cadmium	ND	ppm		ppm	< 0.5	ppm PASS
Cobalt	ND	ppm		ppm	< 5	ppm PASS
Mercury	ND	ppm		ppm	< 3	ppm PASS
Iridium	ND	ppm		ppm	< 10	ppm PASS
Nickel	2.833	ppm	+- 0.03		< 20	ppm PASS
Osmium	ND	ppm		ppm	< 10	ppm PASS
Lead	ND	ppm		ppm	< 0.5	ppm PASS
Palladium	ND	ppm		ppm	< 10	ppm PASS
Platinum	ND	ppm		ppm	< 10	ppm PASS
Rhodium	ND	ppm		ppm	< 10	ppm PASS
Ruthenium	ND	ppm		ppm	< 10	ppm PASS
Selenium	ND	ppm		ppm	< 15	ppm PASS
Thallium	ND	ppm		ppm	< 0.8	ppm PASS
Vanadium	0.294	ppm	+- 0.004	4 ppm	< 10	ppm PASS
			Analyst:	Paul Myburgh	2023/05	/05 15:05:03



^{*}ND (Not Detected)

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^{*}The current test method version is Validated

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^{*}Standard Deviation (Uncertainty of measurement of applicable duplicate sample)

^{*}Safety Limits (Regulatory limits specified are according to the United States pharmacopeia where applicable)

^{*}OR (Outside Range) For quantitative analysis, calibration range is the concentration within which the analysis method is accurate. For limit tests the a single point calibration is employed with a range of +-15% across the specification limit. An OR analyte flag signifies that only semi-quantitative data has been reported.

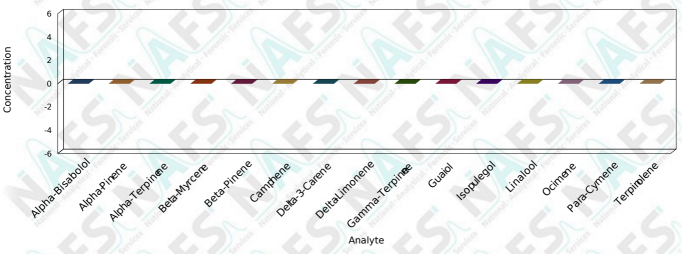


Test Name: Terpenes (200603-004-008)

Specification: USP Specification (Oral)

Test Method: *TM003.5 Sampling Method: SH007

Analyte	Results		*Std Dev.	*Safety Limits
Alpha-Bisabolol	ND	mg/g	mg/g	N/A
Alpha-Pinene	ND	mg/g	mg/g	N/A
Alpha-Terpinene	ND	mg/g	mg/g	N/A
Beta-Myrcene	ND	mg/g	mg/g	N/A
Beta-Pinene	ND	mg/g	mg/g	N/A
Camphene	ND	mg/g	mg/g	N/A
Delta-3-Carene	ND	mg/g	mg/g	N/A
Delta-Limonene	ND	mg/g	mg/g	N/A
Gamma-Terpinene	ND	mg/g	mg/g	N/A
Guaiol	ND	mg/g	mg/g	N/A
Isopulegol	ND	mg/g	mg/g	N/A
Linalool	ND	mg/g	mg/g	N/A
Ocimene	ND	mg/g	mg/g	N/A
Para-Cymene	ND	mg/g	mg/g	N/A
Terpinolene	ND	mg/g	mg/g	N/A



^{*}ND (Not Detected)

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^{*}The current test method version is Un-Validated

^{*}Standard Deviation (Uncertainty of measurement of applicable duplicate sample)

^{*}Safety Limits (Regulatory limits specified are according to the United States pharmacopeia where applicable)



Interpretations and Opinions:

None

Additions, Deviations & Exclusions:

None

It should be noted that NAFS will only analyze the sample/s received. This sample cannot be regarded as representative of entire batch or crop. It is the responsibility of the client to ensure a representative sample is taken in an appropriate tamper proof sample container. NAFS cannot be held liable for negligent handling, storage and transport of client samples, prior to receipt. Any complaints may be directed toward NAFS using info@nafs.co.za (QA005)

Quality Assurance Manager

Released By: Jeanette Leygonie
Released On: 2023/05/08 12:23:54

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