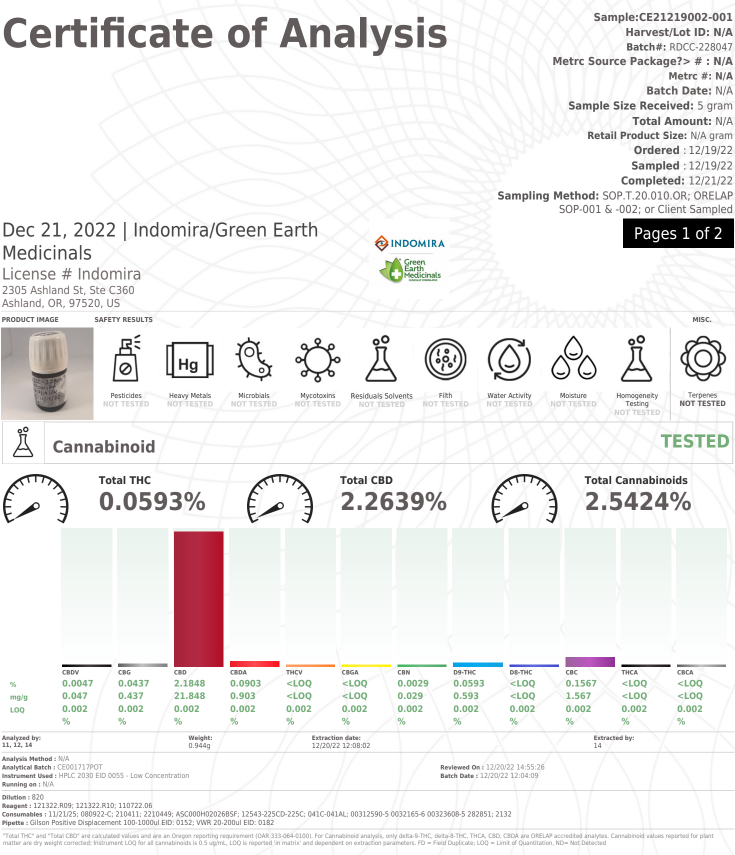


Central Point, OR, 97502, US

Kaycha Labs

CBD Oral Drops Cinnamon N/A Sample Type: Ingestible





This report shall not be reproduced, unless in its entirety, without written approval from Kaycha Labs. This report is an Kaycha Labs certification. Laboratory reports are for informational use only, unless indicated otherwise. The results relate only to the material or product analyzed. Test results are confidential unless explicitly waived otherwise. Void after 1 year from test end date. Cannabinoid content of batch material may vary depending on sampling error. IC=In-control QC parameter, NC=Non-controlled QC parameter, ND=Not Detected, NA=Not Analyzed, pom=Parts Per Million, pob=Parts Per Billion. Limit of Potection (LoD) and Limit Of Quantitation (LoQ) are terms used to describe the smallest concentration that can be reliably measured by an analytical procedure. RPD=Reproducibility of two measurements. Action Levels are State determined thresholds variable based on uncertainty of measurement (UM) for the analyte. The UM error is available from the lab upon request.The "Decision Rule" for the pass/fail does not include the UM. The limits are based on OAR 333-007, OAR 845-025.

Anthony Smith Lab Director State License # 010-10166277B9C ISO 17025 Accreditation # 99861

12/21/22

Signature

Signed On



Sample Type : Ingestible

N/A

Page 2 of 2

POTENCY BATCH QC REPORT



METHOD BLANK

Cannabinoid	LOQ	Result	Units
CBDV WET	0.002	0	%
CBDVA WET	0.002	NT	%
CBG_WET	0.002	0	%
CBD_WET	0.002	0	%
CBDA_WET	0.002	0	%
THCV WET	0.002	NT	%
CBGA_WET	0.002	0	%
CBN_WET	0.002	0	%
D9-THC_WET	0.002	0	%
D8-THC_WET	0.002	0	%
CBC_WET	0.002	0	%
THCA WET	0.002	0	%

Analytical Batch - CE001717POT

Instrument Used : HPLC 2030 EID 0055 - Low Concentration

ĥ
A

LCS

Cannabinoid	LOQ	Recovery	Units	Recovery Limits
CBG_WET	0.002	97.8	%	80-120
CBD_WET	0.002	106	%	90-110
CBDA_WET	0.002	100.9	%	90-110
CBGA_WET	0.002	100.5	%	80-120
CBN_WET	0.002	100.6	%	80-120
D9-THC_WET	0.002	101.2	%	90-110
D8-THC_WET	0.002	109.5	%	90-110
CBC_WET	0.002	99.4	%	80-120
THCA_WET	0.002	99.2	%	90-110

Analytical Batch - CE001717POT

Instrument Used : HPLC 2030 EID 0055 - Low Concentration

This report shall not be reproduced, unless in its entirety, without written approval from Kaycha Labs. This report is an Kaycha Labs certification. Laboratory reports are for informational use only, unless indicated otherwise. The results relate only to the material or product analyzed. Test results are confidential unless explicitly waived otherwise. Void after 1 year from test end date. Cannabinoid content of batch material may vary depending on sampling error. IC=In-control QC parameter, NC=Non-controlled QC parameter, ND=Not Detected, NA=Not Analyzed, pom=Parts Per Million, ppb=Parts Per Billion. Limit of Detection (LoD) and Limit Of Quantitation (LoQ) are terms used to describe the smallest concentration that can be reliably measured by an analytical procedure. RPD=Reproducibility of two measurements. Action Levels are State determined thresholds variable based on uncertainty of measurement (UM) for the analyte. The UM error is available from the lab upon request. The "Decision Rule" for the pass/fail does not include the UM. The limits are based on OAR 333-007, OAR 845-025.

Anthony Smith

Lab D State License # 010-10166277B9D ISO 17025 Accreditation # 99861



Signature

Signed On

KML Laboratories, Inc.



261 Great Northern Road **Bonners Ferry, Idaho 83805** Phone: 208-267-0818 Fax: 208-267-0878 Email: Info@kmlmicro.com

Certificate of Analysis

Green Earth Medicinals 2305 Ashland St. Suite C360 Ashland, OR 97520 Phone: 888-620-1110 Fax:

Invoice Number: 22.1496 PO Number: 12/19/2022 Received Date: 12/23/2022 Number of Samples: 01 Project Name: **Routine Testing**

Microbiology Report:

Lab #: 22-10801			Sample Lot: RDCC-228047		Sample Date: 12/19/2022	
Sample Name: CBD Oral Sublingual Drops Cinnamon			Additional ID:		Plated Date: 12/23/2022	
Qualifying Material Num	Qualifying Material Number: No QM			2		
Test Performed	Results	Units	Detection Limit	Method	Date Analyzed	
Aerobic Plate Count	<10	cfu/ml	10	USP 43-NF 38 <	<2021> 12/26/2022	
Coliforms	<10	cfu/ml	10	Bam C4 sec G	12/24/2022	
E. coli	absent	P/A	1	USP 43-NF 38 <	<2022> 12/26/2022	
Staph aureus	absent	P/A	1	USP 43-NF 38 <	<2022> 12/26/2022	
Yeast	<10	cfu/ml	10	USP 43-NF 38 <	<2021> 12/28/2022	
Mold	<10	cfu/ml	10	USP 43-NF 38 <	<2021> 12/28/2022	
Salmonella	absent	P/A	1	USP 43-NF 38 <	<2022> 12/27/2022	
Pseudo. aeruginosa	absent	P/A	1	USP 43-NF 38 <	(m62> 12/26/2022	
		1.00				

Poole ing Companies with Approved By: QA Director SMV 12/28/2022



Confidential

Malter

Page 1 of 2

This Certificate/Report shall not be reproduced, except in full, without the prior written consent of KML Laboratories, Inc. This report may include work not covered by KML's current ISO accreditation as indicated by ‡.

Note: On this date, this material met the specifications designated above, and is not known if statistically representative of the lot evaluated on a routine basis. This information is not intended to relieve the purchaser from its responsibility to determine the suitability of this material for purchaser's purposes, to comply with all laws and regulations regarding the safe use of this material. nd = none detected above the listed detection limit

KML Laboratories, Inc.



NS.

261 Great Northern Road **Bonners Ferry, Idaho 83805** Phone: 208-267-0818 Fax: 208-267-0878 Email: Info@kmlmicro.com

Certificate of Analysis

Green Earth Medicinals 2305 Ashland St. Suite C360 Ashland, OR 97520 Phone: 888-620-1110 Fax:

Invoice Number: 22.1496 PO Number: 12/19/2022 Received Date: 12/23/2022 Number of Samples: 01 Project Name: Routine Testing

Microbiology Report:

Lab #: Control 12232022	Additional ID: Negative Control Purposes	Plated Date: 12/23/2022
Sample Name: Control 12232022		
Qualifying Material Number: QM-06-001		

Test Performed	Results	Units	Detection Limit	Method Date Analy	zed
				Y	
Aerobic Plate Count	nd	cfu/ml	10	USP 43-NF 38 <2021> 12/26/2022	2
Coliforms	nd	cfu/ml	10	Bam C4 sec G 12/24/2022	2
E. coli	absent	P/A	1	USP 43-NF 38 <2022> 12/26/2022	2
Staph aureus	absent	P/A	1	USP 43-NF 38 <2022> 12/26/2022	2
Yeast	nd	cfu/ml	10	USP 43-NF 38 <2021> 12/28/2022	2
Mold	nd	cfu/ml	10	USP 43-NF 38 <2021> 12/28/2022	2
Salmonella	absent	P/A	1	USP 43-NF 38 <2022> 12/27/2022	2
Pseudo. aeruginosa	absent	P/A	1	USP 43-NF 38 <m62> 12/26/2022</m62>	2

on ering Companies with Approved By: QA Director SMV 12/28/202



Confidential

Malter

Page 2 of 2

This Certificate/Report shall not be reproduced, except in full, without the prior written consent of KML Laboratories, Inc. This report may include work not covered by KML's current ISO accreditation as indicated by ‡.

Note: On this date, this material met the specifications designated above, and is not known if statistically representative of the lot evaluated on a routine basis. This information is not intended to relieve the purchaser from its responsibility to determine the suitability of this material for purchaser's purposes, to comply with all laws and regulations regarding the safe use of this material. nd = none detected above the listed detection limit