



Certificate of Analysis


502 Hemp 750mg/30mL FSD Vanilla Latte
Matrix: Derivative

Accession Number: 100721UD0008

Harvest/Lot ID: 5H10062116-01

Seed to Sale: *

Batch Date: 10/06/21

Batch #: 5H10062116-01

Sample Size Received: 10 ml

Retail Product Size: 10 ml

Ordered: 10/06/21

Completed: 10/13/21

Expires: 10/12/22

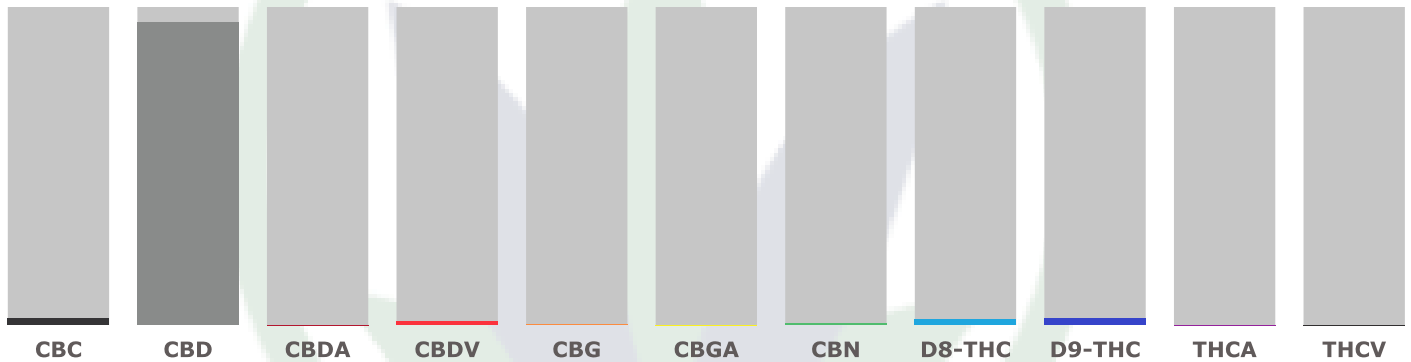
Sampling Method: SOP Client Method

Oct 13, 2021 | Commonwealth Extracts

 Louisville, Kentucky,
 (504) 419-7527

CANNABINOID RESULTS

Total THC 0.054%	Total CBD 2.664%	Total Cannabinoids 2.874%
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Conc.(wt%)	0.059	2.664	ND	0.030	0.006	ND	0.009	0.052	0.054	ND	ND
Conc.(mg/g)	0.590	26.640	ND	0.300	0.060	ND	0.090	0.520	0.540	ND	ND
LOQ	0.04	0.04	0.04	0.04	0.04	0.04	0.04	0.04	0.04	0.04	0.04

Analyzed by	Date	Instrument used	Analysis Method
TW	10/12/2021	Shimadzu HPLC w/ PDA	SOP.KY.02.012

Full spectrum cannabinoid analysis utilizing High Performance Liquid Chromatography with UV detection (HPLC-PDA). SOP.KY.02.005 for sample prep and SOP.KY.02.012 for analysis. % = %w/w = Percent (Weight of Analyte/Weight Product) Total Cannabinoids result reflects the absolute sum of all cannabinoids detected. **Total Potential THC/CBD is calculated using the following formulas to take into account the loss of a carboxyl group during decarboxylation Total THC = THC + (THCa*0.877) Total CBD = CBD + (CBDA*0.877)

This report shall not be reproduced, unless in its entirety, without written approval from BlueLeaf Laboratory. This report is an BlueLeaf Laboratory certification. 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, ppm=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 for human safety for consumption and/or inhalation. The result >99% are 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 F.S. Rule 64-4.310.

Daniel Burriss

 Lab Director
 State License # 19-05-02P
 ISO/IEC 17025:2017

10/13/21



Signature

Signed On