

Twenty One Cannabis



CERTIFICATE OF ANALYSIS

License #: 00000020LCVT89602592

Certificate: 1644

Sample ID: 2311SMAZ0285.0911 Batch #: 21

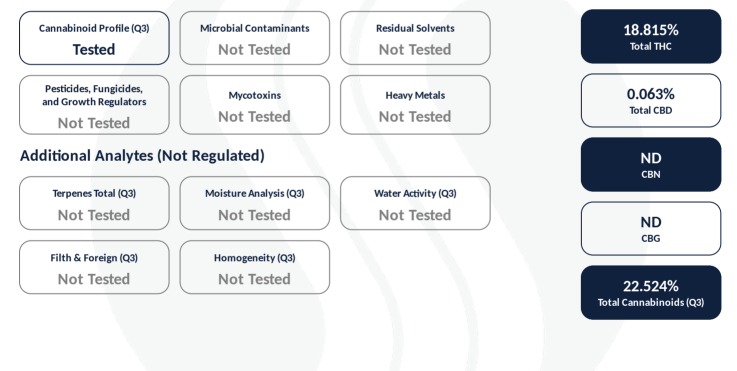
Hemp THCa Flower

Batch #: 21 Strain: 37 Mauie Wowie Parent Batch #: Sample Collected: 11/08/2023 08:31:00 Published: 11/13/2023 Sample ID: 2311SMAZ0285.0911 Amount Received: 2.8 g Sample Type: Flower - Cured Received: 11/09/2023



COMPLIANCE FOR RETAIL

Regulated Analytes



Ahmed Munshi

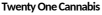
Technical Laboratory Director

AMMunshi

Smithers CTS Arizona LLC 734 W Highland Avenue, 2nd Floor Phoenix, AZ 85013 (602) 806-6930



The product associated with this COA has been tested by Smithers CTS Arizona LLC, using validated state certified testing methodologies as required by Arizona state law. This COA is governed by the terms and conditions listed on: https://www.smithers.com/arizona-terms-conditions





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Cannabinoid Profile			
HPLC	Tested		

Sample Prep

Batch Date: 11/08/2023 SOP: 418.AZ Batch Number: 325

Sample Analysis

Date: 11/09/2023 SOP: 417.AZ - HPLC Sample Weight: 0.101 g Volume: 40 mL

Analyte	LOD (mg/g)	LOQ (mg/g)	Dil.	Actual % (w/w)	mg/g	Qualifier
СВС	0.128	0.387	1	ND	ND	
CBD	0.128	0.387	1	ND	ND	
CBDA	0.128	0.387	1	0.071	0.715	
CBDV	0.128	0.387	1	ND	ND	
CBG	0.128	0.387	1	ND	ND	
CBGA	0.128	0.387	1	1.067	10.668	
CBN	0.128	0.387	1	ND	ND	
d8-THC	0.128	0.387	1	ND	ND	
d9-THC	0.128	0.387	1	0.285	2.853	
THCA	0.128	0.387	1	20.900	209.004	
THCV	0.128	0.387	1	ND	ND	

Cannabinoid Totals	Actual % (w/w)	mg/g	Qualifier
Total THC	18.815	188.150	
Total CBD	0.063	0.627	
Total Cannabinoids	22.524	225.240	Q3

Total THC = THC + (0.877 x THCA) and Total CBD = CBD + (0.877 x CBDA) ND = Not Detected, NT = Not Tested, <LOQ = Below Limit of Quantitation

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Qualifier Legend

- B1 The target analyte detected in the calibration is at or above the limit of quantitation, but the sample result for potency testing, is below the limit of quantitation.
- B2 The target analyte detected in the calibration blank, or the method blank is at or above the limit of quantitation, but the sample result when testing for pesticides, fungicides, herbicides, growth regulators, heavy metals, or residual solvents, is below the maximum allowable concentration for the analyte.
- D1 The limit of quantitation and the sample results were adjusted to reflect sample dilution.
- 1 The relative intensity of a characteristic ion in a sample analyte exceeded the acceptance with respect to the reference spectra, indicating interference.
- When testing for pesticides, fungicides, herbicides, growth regulators, heavy metals, or residual solvents, the percent recovery of a laboratory control sample is greater than the acceptance limits, but the sample's target analytes were not detected above the maximum allowable concentrations for the analytes in the sample.
- M1 The recovery from the matrix spike was high, but the recovery from the laboratory control sample was within acceptance criteria.
- M2 The recovery from the matrix spike was low, but the recovery from the laboratory control sample was within acceptance criteria.
- M3 The recovery from the matrix spike was unusable because the analyte concentration was disproportionate to the spike level, but the recovery from the laboratory control sample was within acceptance criteria.
- M4 The analysis of a spiked sample required a dilution such that the spike recovery calculation does not provide useful information, but the recovery from the associated laboratory control sample was within acceptance criteria.
- M5 The analyte concentration was determined by the method of standard addition, in which the standard is added directly to the aliquots of the analyzed sample.
- M6 A description of the variance is described in the final report of testing according to R9-17-404.06(B)(3)(d)(ii).
- Q1 Sample integrity was not maintained.
- Q2 The sample is heterogeneous, and sample homogeneity could not be readily achieved using routine laboratory practices.
- Q3 Testing result is for informational purposes only and cannot be used to satisfy dispensary testing requirements in R9-17-317.01(A) or labeling requirem
- R1 The relative percent difference for the laboratory control sample and duplicate exceeded the limit, but the recovery was within acceptance criteria.
- R2 The relative percent difference for a sample and duplicate exceeded the limit.
- V1 The recovery from continuing calibration verification standards exceeded the acceptance limits, but the sample's target analytes were not detected above the maximum allowable for the analytes in the sample.

Notes:

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