

**Oklahoma City Location** 3680 E I-240 Service Rd Oklahoma City, OK 73135 (405) 595-0344 **Tulsa Location** 14720 E Admiral Pl, Ste H Tulsa, OK 74108 (918) 340-7139

#### SUMMARY OF ANALYSIS (SAMPLE ID: SA27622)

Testing Location:	Customer ID: 710	Order ID: OR8593	Sample Type: Primary	
OKC	Sunday Extracts	Lot Number:	Matrix: Concentrate	
3680 E. I-240 Service Rd.	1006 North Oak St	Not Entered	Mass: 2g	
Oklahoma City, OK 73135	Guthrie, OK 73044	Batch Number:	Date Collected: 05/26/2020	
License: LAAA-4Y4X-Z72Z	License: PAAA-NYIN-5SXK	RDS521MCGS415	Date Received: 05/26/2020	
Cultivar (Strain) or Sample Description: Mand Cookie x Grape SorbetDate Completed: 05/29/2020				

\*This page is simply a summary of the analysis performed. For analytical details, please consult the individual Certificate(s) of Analysis for each of the specific test(s) performed.

\*For Oklahoma, with the new OMMA rules to be effective April 1, 2020, limits on moisture are proposed at 15% and water activity at 0.65aw.

Moisture Content (%)	PASS/FAIL	Water Activity (aw)	PASS/FAIL
Not Tested	N/A	Not Tested	N/A
Cannabinoids (1	бор 3)	<u>(%)</u>	<u>mg</u> /g
THCa		82.5	825
CBC		0.302	3
<b>Δ9-</b> THC		0.255	3
TOTAL CBI	)		-
TOTAL THO		72.6	726
TOTAL CANNABI	NOIDS	83.3	833
<u>Terpenes (Top</u>	5)	<u>(%)</u>	<u>µg/g</u>
β-Caryophylle	ne	2.083	20827
α-Humulene		1.086	10862
d-Limonene		0.832	8317
β-Myrcene		0.440	4396
β-Pinene		0.219	2193
TOTAL TERPE	NES	5.59	55926
Contaminants	PASS/FAIL	Sample Picture Up	on Receipt
Heavy Metals:	PASS		

Contaminants	PASS/FAIL
Heavy Metals:	PASS
Microbiology:	PASS
Mycotoxins:	PASS
Pesticides:	PASS
<b>Residual Solvents:</b>	PASS
Visual Inspection:	PASS





Scan the QR code to verify results.

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REPORT OF LABORATORY ANALYSIS

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## **CANNABINOID (POTENCY) PROFILE**

Analysis Date/Time: 05/27/2020 1051 Analyst: WW

Method: HPLC/DAD Instrument: Agilent 1100 Moisture Content (%): -Water Activity (aw): -

<u>Cannabinoid</u>	<u>Result</u> (%)	<u>Result</u> (mg/g)	<u>Reporting Limit</u> (mg/g)	<u>Result</u> (mg/mL)	<u>Per Unit</u> (mg)	
CBD	-	-	0.498	-	-	
CBDa	-	-	0.498	-	-	
CBDv	-	-	0.498	-	-	
Δ9-ΤΗC	0.255	2.55	0.498	-	3	Constant State
Δ8-THC	0.203	2.03	0.498	-	2	
THCa	82.5	825	0.498	-	825	
THCv	-	-	0.498	-	-	
CBC	0.302	3.02	0.498	-	3	
CBG	-	-	0.498	-	-	120
CBGa	-	-	0.498	-	-	
CBN	-	-	0.498	-	-	
TOTAL	83.3	833		-	833	
TOTAL THC	72.6	726		-	726	
TOTAL CBD	-	-		-	-	

UNIT MASS (g): 1

#### "-" Not detected above RL.

ndard operating procedure: None

Recoveries for all analyte standards: 90-110% Replicate Uncertainties: <5% RSD, <20% RPD Sample/Reagent Blanks: <RL for all analytes

Values for plant matter are adjusted for moisture content.

Total THC = (THCa x 0.877) +  $\Delta$ 9-THC Total CBD = (CBDa x 0.877) + CBD

Percentage results are reported by mass. mg/g results are reported as mass component per mass material.

Abbreviations: UV - Ultraviolet, HPLC - High Pressure Liquid Chromatography, RL - Reporting Limit, RPD -Relative Percent Difference, RSD - Relative Standard Deviation



Kyle W. Felling, Ph.D. atory Directo

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www.FASTLaboratories.com

-	
	Deviations from stan

100 80 60 40 20 Δ9-THC Δ8-THC THCa THCv CBG CBGa CBC CBD CBDa CBDv CBN

**Cannabinoid Distribution** 

(% of Total Cannabinoids)

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**Cultivar (Strain) or Sample Description:** Mand Cookie x Grape Sorbet

#### **TERPENOID PROFILE**

Analysis Date/Time: Analyst: DJ	05/27/2020 1051		nod: HS/GC/FID (Internal Method-002) rument: Agilent 6890	<b>Deviations from SOP:</b> None
<u>Terpene</u>	<u>Result</u> (µg/g)	<u>Result</u> (%)		
α-Bisabolol	-	-		
Camphene	745	0.0745		
δ-3-Carene	317	0.0317		
β-Caryophyllene	20827	2.08		Rod Salaria Rod Salaria
Caryophyllene oxide	-	-		
p-Cymene	741	0.0741		
Eucalyptol	365	0.0365		
Geraniol	845	0.0845		
Guaiol	-	-	Abbreviatio	ns: HS - Headspace, GC -
α-Humulene	10862	1.09		atography, MS - Mass ry, RL - Reporting Limit
Isopulegol	-	-	•	ation is provided as a
d-Limonene Linalool	8317	0.832	service and	makes no claims of efficacy y of this product.
β-Myrcene	4396	0.44		applicable only for the
cis-Nerolidol		-	analysis co	nalyzed and for the specific nducted.
trans-Nerolidol	_	_		is for informational
$\alpha$ -Ocimene	671	0.0671		lly and should not be used to eat, or prevent any
β-Ocimene	1263	0.126	medical-rel	ated symptoms.
α-Pinene	1470	0.120		ents and results herein have proved and/or endorsed by
β-Pinene	2193	0.219	the FDA.	r
$\alpha$ -Terpinene	454	0.0454	ī	
γ-Terpinene	550	0.055		
Terpinolene	1910	0.191	"-" Not deta	ected above RL.
TOTAL	55926	5.59		<b>Reporting Limit (µg/g):</b> 19.9







Analysis Date/Time: 05/27/2020 1051

**Arkansas Location** 232 S Broadview St Greenbrier, AR 72058 (501) 679-2616

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Result <u>(µg/g)</u> Action Level

<u>(µg/g)</u>

1000

2

1000 60

60

430

1000

60

1000

# **RESIDUAL SOLVENT PROFILE**

Instrument: Agilent 6890

**Deviations from SOP:** 

None

<u>Solvent</u>	
Acetone (67-	(
D (= 1	

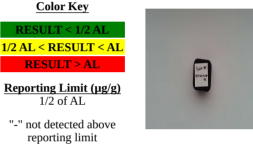
Analyst: CC

Acetone (67-64-1)
Benzene (71-43-2)
n-Butane (106-97-2)
2,2-Dimethylbutane (75-83-2)
2,3-Dimethylbutane (79-29-8)
Ethyl benzene (100-41-4)
n-Heptane (142-82-5)
n-Hexane (110-54-3)
Isobutane (75-28-5)
Isopropanol (67-63-0)
2-Methylbutane (78-78-4)
2-Methylpentane (107-83-5)
3-Methylpentane (96-10-0)
n-Pentane (109-66-0)
n-Propane (74-98-6)
Toluene (108-88-3)
o-Xylene (95-47-6)
m,p-Xylene (108-38-3 or 106-42-3)
Xylenes* (1330-20-7)

Method: USP <467>

"\*" - o,m,p-Xylene and

Ethylbenzene



Solvent	Synonym(s)	Solvent	Synonym(s)	Solvent	Synonym(s)
Acetonitrile	Methyl Cyanide, ACN	2-Ethoxyethanol	Cellosolve, Ethyl glycol	Methanol	Methyl alcohol
1-Butanol	n-Butanol, Butyl	Ethyl ether	Diethyl ether, Ether	2-Methylbutane	Isopentane
	Alcohol	Ethyl acetate	EtOAc	Methylene chloride	Dichloromethane
2-Butanol	sec-Butyl alcohol	Ethyl benzene	Phenylethane	2-Methylpentane	Isohexane
2-Butanone	Methyl ethyl ketone,	Ethylene glycol	1,2-Ethanediol	1-Pentanol	n-Amyl alcohol
	MEK	Ethylene oxide	Oxirane	1-Propanol	Propyl alcohol
1,2-Dimethoxyethane	Monoglyme	Isobutane	2-Methylpropane	Tetrahydrofuran	THF
2,3-Dimethylbutane	Neohexane		511	5	
2,3-Dimethylbutane	Diisopropyl	Isopropanol	2-Propanol, IPA	Tetramethylene sulfone	Sulfolane
N,N-Dimethylformamide	DMF	Isopropyl Acetate	Acetic acid isopropyl ester	Xylene	Dimethylbenzene
Dimethysufoxide	DMSO				

Abbreviations: HS-Headspace, GC-Gas Chromatography, MS-Mass Spectrometry, RL-Reporting Limit, AL-Action Level CAS-Chemical Abstract Services

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Analysis Date/Time: 05/27/2020 1051 Analyst: DJ

## **PESTICIDES PROFILE**

Method: LC/MS/MS Instrument: Waters Acquity/TQD

## **Deviations from SOP:**

None

Color Key

**RESULT < 1/2 AL** 

1/2 AL < RESULT < AL

**RESULT > AL** 

Reporting Limit (µg/g)

1/2 of AL

"-" not detected above

reporting limit

"\*" analyzed by GC/MS

(all others analyzed

by LC/MS/MS)

Permethrins measured as the cumulative residue of

the cis- and trans-

permethrin isomers.

Abamectin (71751-41-2)
Azoxystrobin (131860-33-8)
Bifenazate* (149877-41-8)
Etoxazole (153233-91-1)
Imazalil (35554-44-0)
Imidacloprid (138261-41-3)
Malathion (121-75-5)
Myclobutanil (88671-89-0)
Permethrins* (52645-53-1)
Spinosad A (168316-95-8)
Spinosad D (168316-95-8)
Spiromesifen (283594-90-1)
Spirotetramat (203313-25-1)
Tebuconazole (80443-41-0)

Pesticide

<u>Result</u> (µg/g)	<u>Action Level (µg/g)</u>
-	0.5
-	0.5
-	0.5
-	0.5
-	0.5
-	0.5
-	0.5
-	0.5
-	0.5
-	0.5
-	0.5
-	0.5
-	0.5
-	0.5

Abbreviations: LC - Liquid Chromatography, GC - Gas Chromatography, MS - Mass Spectrometry, RL - Reporting Limit, AL - Action Level, CAS - Chemical Abstract Services

Pesticide	<u>Synonym(s)</u>	Pesticide	<u>Synonym(s)</u>	Pesticide	Synonym(s)
Cyfluthrin	Baythroid	Myclobutanil	Systhane	Propiconazole	Tilt
DDVP	Dichlorvos	Naled	Dibrom	Propoxur	Baygon
Ethoprophos	Prophos	Phosmet	Imidan		

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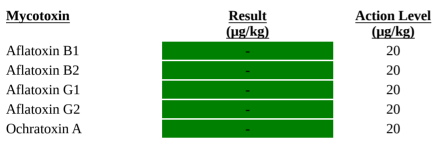
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#### **MYCOTOXIN PROFILE**

Analysis Date/Time: 05/27/2020 1051 Analyst: DJ

Method: LC/MS/MS Instrument: Waters Acquity/TQD **Deviations from SOP:** 

None





Abbreviations: EC - Escherichia Coli, CFU - Colony-Forming Unit, RL - Reporting Limit, AL - Action Level

#### **Color Key**



**Reporting Limit (CFU/g)** 1

"-" not detected above reporting limit

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Cultivar (Strain) or Sample	Date Completed: 05/29/2020		

Cultivar (Strain) or Sample Description: Mand Cookie x Grape Sorbet

**D** ... 1/

## **HEAVY METAL PROFILE**

A ....

Analysis Date/Time: 05/27/2020 1051 Analyst: CC

Method: ICP/MS Instrument: PerkinElmer Elan 9000

#### **Deviations from SOP:** None

**Heavy Metal** Arsenic (As) Cadmium (Cd)

Lead (Pb) Mercury (Hg)

<u>Result</u> (µg/kg)	<u>Action Level</u> <u>(μg/kg)</u>
-	400
-	440
-	1000
-	200



Abbreviations: ICP - Inductively-Coupled Plasma, OES - Optical Emission Spectroscopy, MS - Mass Spectroscopy, RL - Reporting Limit, AL - Action Level

#### Color Key

**RESULT** < 1/2 AI 1/2 AL < RESULT < AL **RESULT > AL** 

Reporting Limit (µg/kg) 50

"-" not detected above reporting limit

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Cultivar (Strain) or Sample Description: Mand Cookie x Grape Sorbet

## MICROBIOLOGICAL PROFILE

Analysis Date/Time: 05/27/2020 1051 Analyst: DJ

Method: Hardy Diagnostics CompactDry **Instrument:** Thermo Incubator

**Action Level** 

(CFU/g)

1 10000

1

1

**Deviations from SOP:** None

<b>Bacteria/Microbe</b>	Result	
	<u>(CFU/g)</u>	
Escherichia Coli (E. Coli)	Absent	
Mold/Yeast	Absent	
Salmonella spp.	Absent	
Staphylococcus Aureus	Absent	



Abbreviations: EC - Escherichia Coli, CFU - Colony-Forming Unit, RL - Reporting Limit, AL - Action Level, TNTC - Too Numerous To Count, NT - Not Tested Absent - Not Detected Above RL, Present - Detected Above RL

#### **Color Key**



**Reporting Limit (CFU/g)** 1

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