



### SUMMARY OF ANALYSIS (SAMPLE ID: SA39278)

Testing Location:	Customer ID: 2168	Order ID: OR11275	Sample Type: Primary
Arkansas	River Valley Relief MIPS	Lot Number:	Matrix: Edible
232 S. Broadview St.	5601 Old Greenwood Rd Suite C	M00065C13227865760	Mass: 1ea
Greenbrier, AR 72058	Fort Smith, AR 72903	Production Run:	Date Collected: 11/04/2024
License: ADH 113	License: 00065C	P20241031GHEE11	Date Received: 11/04/2024
Cultivar (Strain) or Sample De	Date Completed: 11/06/2024		

\*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. All contaminant action levels are referenced from the State of Arkansas MMJ testing guidelines.

Moisture Content (%)	Water Activity (aw)	PASS/FAIL
Not Tested	Not Tested	PASS

Moisture content/water activity action levels are referenced from the State of Arkansas MMJ testing guidelines.

Moisture content levels less than 15% are recommended but the sample does not fail. Water activity levels must be less than 0.65aw.

<u>Cannabinoi</u> Δ9-Τ Δ8-Τ CB( TOTAL TOTAL	HC HC G CBD THC	(%) 0.639 0.0368 0.0166 0.0122 0.639	mg/g 6.39 0.368 0.166 0.122 6.39	
TOTAL CANN	IABINOIDS	0.715	7.15	
<b><u>Terpenes</u></b> d-Limo α-Bisal Campl δ-3-Ca β-Caryopl	nene polol nene rene nyllene	(%) 0.000508	<b>µg/g</b> 5.08	
TOTAL TE		0.000508	5.08	
<u>Contaminants</u> Heavy Metals: Microbiology: Pesticides: Residual Solvents:	PASS/FAIL PASS PASS PASS PASS	Sample Picture Upon Receipt		



Scan the QR code to verify results.

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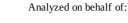


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Cultivar (Strain) or Sample	Date Completed: 11/06/2024		

### CANNABINOID (POTENCY) PROFILE (SOP: SOP-CANN-001)

Analysis Date/Time: 11/05/2024 1616 Analyst: PW Method: HPLC/DAD Instrument: Agilent 1100

<u>Cannabinoid</u>	<u>Result</u> (%)	<u>Result</u> (mg/g)	<u>LOD</u> (mg/g)	LOQ (mg/g)	<u>Result</u> (mg/ mL)	<u>Per</u> Serving (mg)	<u>Per</u> g <u>Unit</u> (mg)
CBC	ND	ND	0.00225	0.00526	-	-	-
CBCA	ND	ND	0.00700	0.0163	-	-	-
CBD	0.0122	0.122	0.0159	0.0371	-	0.183	11.0
CBDA	ND	ND	0.00585	0.0137	-	-	-
CBDV	ND	ND	0.00255	0.00595	-	-	-
CBDVA	ND	ND	0.00680	0.0159	-	-	-
CBG	0.0166	0.166	0.0103	0.0241	-	0.248	14.9
CBGA	ND	ND	0.0146	0.0173	-	-	-
CBL	ND	ND	0.0119	0.0278	-	-	-
CBN	0.0108	0.108	0.00548	0.0128	-	0.162	9.73
CBNA	ND	ND	0.00591	0.0138	-	-	-
Δ9-ΤΗC	0.639	6.39	0.00656	0.0153	-	9.58	575
$\Delta 8$ -THC	0.0368	0.368	0.0102	0.0239	-	0.552	33.1
THCA	ND	ND	0.00356	0.00832	-	-	-
THCV	ND	ND	0.00854	0.0199	-	-	-
THCVA	ND	ND	0.00273	0.00635	-	-	-
TOTAL	0.715	7.15			-	10.7	643
TOTAL CBC	-	-			-	-	-
TOTAL CBD	0.0122	0.122			-	0.183	11.0
TOTAL CBDV	-	-			-	-	-
TOTAL CBG	0.0166	0.166			-	0.248	14.9
TOTAL CBN	0.0108	0.108			-	0.162	9.73
TOTAL THC	0.639	6.39			-	9.58	575
TOTAL THCV	-	-			-	-	-

Abbreviations: DAD - Diode Array Detector, HPLC - High Pressure Liquid Chromatography, RL - Reporting Limit, RPD - Relative Percent Difference, RSD - Relative Standard Deviation, DET - Detected (less than LOQ), LOD - Limit of Detection, LOQ - Limit of Quantitation, UM - Measurement Uncertainty

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Moisture Content (%): -Water Activity (aw): -



SERVING MASS (g): 1.50 SERVINGS/UNIT: 60

"-" Not detected above LOD.

Deviations from standard 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 CBC = (CBCA x 0.877) + CBC Total CBD = (CBDA x 0.877) + CBD Total CBDV = (CBDVA x 0.867) + CBDV Total CBG = (CBGA x 0.878) + CBG Total CBN = (CBNA x 0.876) + CBN Total THC = (THCA x 0.877) +  $\Delta$ 9-THC Total THCV = (THCVA x 0.867) + THCV

Percentage results are reported by mass.

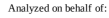
mg/g results are reported as mass component per mass material.

W. Felling, Ph.D. ory Dire











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License: ADH 113	License: 00065C	P20241031GHEE11	Date Received: 11/04/2024
Cultivar (Strain) or Sample D	Date Completed: 11/06/2024		

Analysis Date/Time:11/05/2024 1728 Analyst: KF

<b>TERPENOID PROFILE</b>
Method: GC/MS

Instrument: Agilent 7890/5975

			3
<u>Terpene</u>	<u>Result</u> (µg/g)	<u>Result</u> <u>(%)</u>	
α-Bisabolol	ND	-	
Camphene	ND	-	
δ-3-Carene	ND	-	
β-Caryophyllene	ND	-	
Caryophyllene oxide	ND	-	
p-Cymene	ND	-	
Eucalyptol	ND	-	
Geraniol	ND	-	
Guaiol	ND	-	
α-Humulene	ND	-	
Isopulegol	ND	-	
d-Limonene	5.08	0.000508	
Linalool	ND	-	
β-Myrcene	<loq< td=""><td>-</td><td></td></loq<>	-	
cis-Nerolidol	ND	-	
trans-Nerolidol	ND	-	
α-Ocimene	ND	-	
β-Ocimene	<loq< td=""><td>-</td><td></td></loq<>	-	
α-Pinene	ND	-	
β-Pinene	ND	-	
α-Terpinene	ND	-	
γ-Terpinene	ND	-	
Terpinolene	ND	-	
TOTAL	5.08	0.000508	



**Deviations from SOP:** 

None

Abbreviations: GC - Gas Chromatography, MS - Mass Spectrometry, RL - Reporting Limit Abbreviations: ND - Not Detected, , LOD - Limit of Detection, LOQ - Limit of Quantitation
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#### Reporting Limit (µg/g): 4.94

"-" Not detected above LOD.













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License: ADH 113	License: 00065C	P20241031GHEE11	<b>Date Received:</b> 11/04/2024
Cultivar (Strain) or Sample	<b>Date Completed:</b> 11/06/2024		

### **RESIDUAL SOLVENT PROFILE (SOP: SOP-RS-001)**

Analysis Date/Time:	11/04/20	24 2226		Ι	Method: HS/GC/MS			D	eviations	s from SOP:
Analyst: KF				I	nstrument: Agilent 78	90/5975	i i	Ν	one	
<u>Solvent</u>	<u>Result</u> (µg/g)	<u>LOD</u> (µg/g)	<u>LOQ</u> (µg/g)	<u>Action</u> Level (µg/g)	<u>Solvent</u>	<u>Result</u> (µg/g)	<u>LOD</u> (µg/g)	<u>LOQ</u> (µg/g)	<u>Action</u> Level (µg/g)	
Acetone (67-64-1)	-	35.3	70.6	5000	n-Heptane (142-82-5)	-	35.3	70.6	5000	
Acetonitrile (75-5-8)	-	35.3	70.6	410	n-Hexane (110-54-3)	-	12.4	24.7	290	
Benzene (71-43-2)	-	0.353	0.706	2	Isobutane (75-28-5)	-	35.3	70.6	5000	
n-Butane (106-97-2)	-	35.3	70.6	5000	Isopropanol (67-63-0)	-	35.3	70.6	5000	
1-Butanol (71-36-3)	-	35.3	70.6	5000	Isopropyl acetate	_	35.3	70.6	5000	
2-Butanol (78-92-2)	-	35.3	70.6	5000	(108-21-4)		00.0	, 010	0000	
2-Butanone (78-93-3)	-	35.3	70.6	5000	Isopropyl benzene (98-82-8)	-	3.53	7.06	70	
Cyclohexane (110-82-7)	-	35.3	70.6	3880	(98-62-6) Methanol (67-56-1)		35.3	70.6	3000	Color Key
1,2-Dimethoxyethane	_	3.53	7.06	100	2-Methylbutane (78-78-4)	_	35.3	70.6	5000	
(110-71-4) N,N-Dimethylacetamide (127-19-5)	-	35.3	70.6	1090	Methylene chloride (75-9-2)	-	35.3	70.6	600	RESULT < AL RESULT > AL
(127-15-5) 2,2-Dimethylbutane (75-83-2)	-	12.4	24.7	290	2-Methylpentane (107-83-5)	-	12.4	24.7	290	"-" not detected above 1/2 Action Level
2,3-Dimethylbutane		40.4		200	3-Methylpentane (96-10-0)	-	12.4	24.7	290	
(79-29-8)	-	12.4	24.7	290	n-Pentane (109-66-0)	-	35.3	70.6	5000	"*" - o,m,p-Xylene and
N,N-Dimethylformamide	_	35.3	70.6	880	1-Pentanol (71-41-0)	-	35.3	70.6	5000	Ethylbenzene
(68-12-2)					n-Propane (74-98-6)	-	35.3	70.6	5000	Action levels are
Dimethylsulfoxide (67-68-5)	-	35.3	70.6	5000	1-Propanol (71-23-8)	-	35.3	70.6	5000	referenced from the State of Arkansas
1,4-Dioxane (123-91-1)	-	35.3	70.6	380	Pyridine (110-86-1)	-	12.4	24.7	200	MMJ testing
Ethanol (64-17-5)	-	35.3	70.6	5000	Tetrahydrofuran (109-99-9)	-	35.3	70.6	720	guidelines.
2-Ethoxyethanol (110-80-5)	-	12.4	24.7	160	Tetramethylene sulfone (126-33-0)	-	12.4	24.7	160	A value of "-"
Ethyl ether (60-29-7)	-	35.3	70.6	5000	Toluene (108-88-3)	-	35.3	70.6	890	for the action level
Ethyl acetate (141-78-6)	-	35.3	70.6	5000	o-Xylene (95-47-6)	-	35.3	70.6	2170	means that analyte
Ethyl benzene (100-41-4)	-	35.3	70.6	2170	m,p-Xylene (108-38-3 or					is not currently
Ethylene glycol (107-21-1)	-	35.3	70.6	620	106-42-3)	-	35.3	70.6	2170	regulated by the
Ethylene oxide (75-21-8)	-	3.53	7.06	50	Xylenes* (1330-20-7)	-	43.3	86.7	2170	regulations referenced above.
<u>Solvent</u>		Synonym(s	<u>s)</u>		<u>Solvent</u>	<u>s</u>	Synonym(s)			
Acetonitrile		Methyl Cya	anide, ACN		Ethylene glycol	1	,2-Ethanediol			
1-Butanol		n-Butanol,	Butyl Alco	hol	Isobutane	2	-Methylpropa	ine		
2-Butanol		sec-Butyl a	lcohol		Isopropanol	2	-Propanol, IP/	Ą		
2-Butanone		Methyl eth	yl ketone, N	⁄IEK	Isopropyl Acetate	A	Acetic acid iso	propyl este	r	
1,2-Dimethoxyethane		Monoglym	ie		Methanol	Ν	Aethyl alcoho	l		
2,3-Dimethylbutane		Neohexane	<u>,</u>		2-Methylbutane	I	sopentane			
2,3-Dimethylbutane		Diisopropy	yl		Methylene chloride	Γ	Dichlorometha	ne		
N,N-Dimethylformamide		DMF			2-Methylpentane	I	sohexane			
Dimethysufoxide		DMSO			1-Pentanol	n	-Amyl alcoho	1		
2-Ethoxyethanol		Cellosolve	, Ethyl glyc	ol	1-Propanol	P	ropyl alcohol			
Ethyl ether		Diethyl eth	ner, Ether		Tetrahydrofuran	Г	THF			

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

Sulfolane

Dimethylbenzene

Tetramethylene sulfone

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Xylene



EtOAc

Phenylethane

Ethyl acetate

Ethyl benzene

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**Deviations from SOP:** 

None



# **CERTIFICATE OF ANALYSIS (SAMPLE ID: SA39278)**

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License: ADH 113	License: 00065C	P20241031GHEE11	<b>Date Received:</b> 11/04/2024
Cultivar (Strain) or Sample	<b>Date Completed:</b> 11/06/2024		

### Analysis Date/Time: 11/05/2024 1722 Analyst: KF

### **PESTICIDES PROFILE (SOP: SOP-PEST-001)**

Method: LC/MS/MS Instrument: Shimadzu LC-8050

<u>Pesticide</u>	<u>Result</u> (µg/g)	<u>LOD</u> (µg/g)	<u>LOQ</u> (µg/g)	<u>Action</u> Level (µg/g)	<u>Pesticide</u>	<u>Result</u> (µg/g)	<u>LOD</u> (µg/g)	<u>LOQ</u> (µg/g)	<u>Action</u> Level (µg/g)	
Abamectin (71751-41-2)	-	0.00988	0.0791	0.5	Kresoxim-methyl		0.00988	0.0791	0.4	
Acephate (30560-19-1)	-	0.00988	0.0791	0.4	(143390-89-0)		0.00900	0.0791	0.4	
Acequinocyl (57960-19-7)	-	0.00988	0.0791	2	Malathion (121-75-5)	-	0.00988	0.0791	0.2	
Acetamiprid	_	0.00988	0.0791	0.2	Metalaxyl (57837-19-1)	-	0.00988	0.0791	0.2	
(135410-20-7)					Methiocarb (2032-65-7)	-	0.00988	0.0791	0.2	<u>Color Key</u>
Aldicarb (116-06-3)	-	0.00988	0.0791	0.4	Methomyl (16752-77-5)	-	0.00988	0.0791	0.4	<b>RESULT &lt; AL</b>
Azoxystrobin (131860-33-8)	-	0.00988	0.0791	0.2	Methyl parathion (298-0-0)	-	0.00988	0.0791	0.2	<b>RESULT &gt; AL</b>
Bifenazate (149877-41-8)	-	0.00988	0.0791	0.2	MGK 264 (113-48-4)	-	0.00988	0.0791	0.2	"DET" detected less than LOQ
Bifenthrin (82657-04-3)	-	0.00988	0.0791	0.2	Myclobutanil	_	0.00988	0.0791	0.2	"-" not detected above
Boscalid (188425-85-6)	-	0.00988	0.0791	0.4	(88671-89-0)		0.00000	0.0701	0.5	LOD
Carbaryl (63-25-2)	-	0.00988	0.0791	0.2	Naled (300-76-5)	-	0.00988	0.0791	0.5	Permethrins measured as the
Carbofuran (1563-66-2)	-	0.00988	0.0791	0.2	Oxamyl (23135-22-0)	-	0.00988	0.0791	1	cumulative residue of the cis- and
Chlorantraniliprole (800008-45-7)	-	0.00988	0.0791	0.2	Paclobutrazol (76738-62-0)	-	0.00988	0.0791	0.4	<i>trans</i> - permethrin isomers.
Chlorfenapyr (122453-73-0)	-	0.00988	0.0791	1	Permethrins (52645-53-1)	-	0.00988	0.0791	0.2	Pyrethrins measured as the cumulative residue of the
Chlorpyrifos (2921-88-2)	-	0.00988	0.0791	0.2	Phosmet (732-11-6)	-	0.00988	0.0791	0.2	pyrethrin I, cinerin I, and jasmolin I isomers.
Clofentezine (74115-24-5)	-	0.00988	0.0791	0.2	Piperonyl butoxide	_	0.00988	0.0791	2	
Cyfluthrin (68359-37-5)	-	0.00988	0.0791	1	(51-03-6) Prallethrins (2331-36-9)		0.00988	0.0791	0.2	Action levels are referenced from the State of
Cypermethrin (52315-07-8)	-	0.00988	0.0791	1	Propiconazole	-	0.00988	0.0791	0.2	Arkansas MMJ testing
Daminozide (1596-84-5)	-	0.00988	0.0791	1	(60207-90-1))		0.00000	0.0701	0.2	guidelines.
DDVP (62-73-7)	-	0.00988	0.0791	0.1	Propoxur (114-26-1) Pyrethrins (8003-34-7)	-	0.00988 0.00988	0.0791 0.0791	0.2 1	A value of "-" for the action level
Diazinon (333-41-5)	-	0.00988	0.0791	0.2	Pyrethrins (8003-34-7) Pyridaben (96489-71-3)	-	0.00988	0.0791	0.2	means that analyte is not
Dimethoate (60-51-5)	-	0.00988	0.0791	0.2	Spinosad (168316-95-8)	-	0.00988	0.0791	0.2	currently regulated by the regulations referenced above.
Ethoprophos (13194-48-4)	-	0.00988	0.0791	0.2	Spiromesifen		0.00988	0.0791	0.2	Disclaimer: This information is
Etofenprox (80844-07-1)	-	0.00988	0.0791	0.4	(283594-90-1)					provided as a service and makes
Etoxazole (153233-91-1)	-	0.00988	0.0791	0.2	Spirotetramat (203313-25-1)	-	0.00988	0.0791	0.2	no claims of efficacy and/or safety of this product. Results are
Fenoxycarb (72490-01-8)	-	0.00988	0.0791	0.2	Spiroxamine					applicable only for the sample(s)
(E)-Fenpyroximate (134098-61-6)	-	0.00988	0.0791	0.4	(118134-30-8) Tebuconazole	-	0.00988	0.0791	0.4	analyzed and for the specific analysis conducted. This report is for informational purposes only
Fipronil (120068-37-3)	-	0.00988	0.0791	0.4	(80443-41-0)	-	0.00988	0.0791	0.4	and should not be used to
Flonicamid (158062-67-0)	-	0.00988	0.0791	1	Thiacloprid	_	0.00988	0.0791	0.2	diagnose, treat, or prevent any
Fludioxinil (131341-86-1)	-	0.00988	0.0791	0.4	(111988-49-9)		0.00000	5.0751	0.2	medical-related symptoms. The statements and results herein have
Hexythiazox (78587-05-0)	-	0.00988	0.0791	1	Thiamethoxam (153719-23-4)	-	0.00988	0.0791	0.2	not been approved and/or
Imazalil (35554-44-0)	-	0.00988	0.0791	0.2	Trifloxystrobin					endorsed by the FDA.
Imidacloprid (138261-41-3)	-	0.00988	0.0791	0.4	(141517-21-7)	-	0.00988	0.0791	0.2	

Abbreviations: LC - Liquid Chromatography, MS - Mass Spectrometry, RL - Reporting Limit, AL - Action Level, CAS - Chemical Abstract Services, LOD - Limit of Detection, LOQ - Limit of Quantification

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

Pesticide Propiconazole Propoxur <u>Synonym(s)</u> Tilt Baygon



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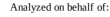


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Arkansas	River Valley Relief MIPS	Lot Number:	Matrix: Edible
232 S. Broadview St.	5601 Old Greenwood Rd Suite C	M00065C13227865760	Mass: 1ea
Greenbrier, AR 72058	Fort Smith, AR 72903	<b>Production Run:</b>	<b>Date Collected:</b> 11/04/2024
License: ADH 113	License: 00065C	P20241031GHEE11	Date Received: 11/04/2024
Cultivar (Strain) or Sample	<b>Date Completed:</b> 11/06/2024		

### HEAVY METAL PROFILE (SOP: SOP-ICP-200.6)

Analysis Date/T	ime: 11/05/2024	1722	Method: IC Instrumen	CP/MS <b>t:</b> Agilent 7500ce	<b>Deviations from SOP:</b> None	
Analyst: KF						
<u>Heavy Metal</u>	<u>Result</u> (µg/kg)	<u>LOD</u> (µg/kg)	<u>LOQ</u> (µg/kg)	<u>Action Level</u> (µg/kg)		
Arsenic (As)	-	58.1	92.1	200		
Cadmium (Cd)	-	58.1	92.1	200		
Lead (Pb)	-	58.1	92.1	500		
Mercury (Hg)	-	58.1	92.1	100		

*Abbreviations:* ICP - Inductively-Coupled Plasma, OES - Optical Emission Spectroscopy, DMA - Direct Mercury Analyzer, RL - Reporting Limit, AL - Action Level, LOD - Limit of Detection, LOQ - Limit of Quantitation

## <u>Color Key</u>



"DET" detected less than LOQ

"-" not detected above LOD

Action levels for heavy metals are referenced from the State of Arkansas MMJ testing guidelines.

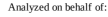
*Disclaimer:* This information is provided as a service and makes no claims of efficacy and/or safety of this product. Results are applicable only for the sample(s) analyzed and for the specific analysis conducted. This report is for informational purposes only and should not be used to diagnose, treat, or prevent any medical-related symptoms. The statements and results herein have not been approved and/or endorsed by the FDA.













Testing Location:	Customer ID: 2168	Sample ID: SA39278	Sample Type: Primary
Arkansas	<b>River Valley Relief MIPS</b>	Lot Number:	Matrix: Edible
232 S. Broadview St.	5601 Old Greenwood Rd Suite C	M00065C13227865760	Mass: 1ea
Greenbrier, AR 72058	Fort Smith, AR 72903	Production Run:	<b>Date Collected:</b> 11/04/2024
License: ADA 05_H273	License: 00065C	P20241031GHEE11	Date Received: 11/04/2024
Cultivar (Strain) or Sample	Date Completed: 11/06/2024		

### MICROBIOLOGICAL PROFILE (SOP: SOP-Micro-001)

Analysis Date/Time: 01/01/19 Analyst: PW		Hardy Diagnostics Compact ent: Thermo Incubator	Dry <b>Deviations from SOP:</b> None
Bacteria/Microbe	<u>Result</u> (CFU/g)	<u>Action Level</u> (CFU/g)	
Aerobic Plate Count	NT	-	
Coliforms, Total	Absent	1	
Escherichia Coli (E. Coli)	Absent	100	
Mold/Yeast	NT	-	
Pseudomonas aeruginosa	NT	-	
Salmonella spp.	NT	-	
Staphylococcus aureus	NT	-	

*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





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Action levels for microbiology are referenced from the State of Arkansas MMJ testing guidelines. A value of "-" for the action level means that analyte is not currently regulated by the regulations referenced above.

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