



12423 NE Whitaker Way
Portland, OR 97230
503-254-1794



Report Number: 23-013558/D002.R000
Report Date: 11/27/2023
ORELAP#: OR100028
Purchase Order:
Received: 11/16/23 12:12

Customer: NW Natural Goods
Product identity: HEMP - PR 0066
Client/Metric ID: .
Laboratory ID: 23-013558-0001

Summary

Potency:

Analyte per 4g	Result	Limits	Units	Status	
CBC per 4g	0.196		mg/4g		CBD-Total per Serving Size 21.0 mg/4g
CBD per 4g	21.0		mg/4g		
CBG per 4g	10.4		mg/4g		THC-Total per Serving Size <LOQ
(Reported in milligrams per serving)					

Residual Solvents:

All analytes passing and less than LOQ.

Pesticides:

Analyte	Result (mg/kg)	Limits (mg/kg)	Status
Multi-Residue Pesticide Profile	< LOQ for all analytes		

Metals:

Less than LOQ for all analytes.

Microbiology:

Less than LOQ for all analytes.



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Received: 11/16/23 12:12

Customer: NW Natural Goods

Product identity: HEMP - PR 0066

Client/Metric ID: .

Sample Date:

Laboratory ID: 23-013558-0001

Evidence of Cooling: No

Temp: 18.6

Relinquished by: ramos

Serving Size #1: 4 g

Sample Results

Potency per 4g					
Method: J AOAC 2015 V98-6 (mod) ^b		Units mg/se		Batch: 2312958	
				Analyze: 11/17/23 8:47:00 PM	
Analyte	Result	Limits	Units	LOQ	Notes
CBC per 4g	0.196		mg/4g	0.122	
CBC-A per 4g	< LOQ		mg/4g	0.122	
CBC-Total per 4g	< LOQ		mg/4g	0.228	
CBD per 4g	21.0		mg/4g	0.122	
CBD-A per 4g	< LOQ		mg/4g	0.122	
CBD-Total per 4g	21.0		mg/4g	0.228	
CBDV per 4g	< LOQ		mg/4g	0.122	
CBDV-A per 4g	< LOQ		mg/4g	0.122	
CBDV-Total per 4g	< LOQ		mg/4g	0.227	
CBE per 4g	< LOQ		mg/4g	0.122	
CBG per 4g	10.4		mg/4g	0.122	
CBG-A per 4g	< LOQ		mg/4g	0.122	
CBG-Total per 4g	10.4		mg/4g	0.227	
CBL per 4g	< LOQ		mg/4g	0.122	
CBL-A per 4g	< LOQ		mg/4g	0.122	
CBL-Total per 4g	< LOQ		mg/4g	0.228	
CBN per 4g	< LOQ		mg/4g	0.122	
CBT per 4g	< LOQ		mg/4g	0.122	
Δ8-THCV per 4g	< LOQ		mg/4g	0.122	
Δ10-THC-9R per 4g	< LOQ		mg/4g	0.122	
Δ10-THC-9S per 4g	< LOQ		mg/4g	0.122	
Δ10-THC-Total per 4g	< LOQ		mg/4g	0.243	
Δ8-THC per 4g	< LOQ		mg/4g	0.122	
Δ9-THC per 4g	< LOQ		mg/4g	0.122	
delta-9-THCP per 4g	< LOQ		mg/4g	0.122	
exo-THC per 4g	< LOQ		mg/4g	0.122	
THC-A per 4g	< LOQ		mg/4g	0.122	
THC-Total per 4g	< LOQ		mg/4g	0.228	
THCV per 4g	< LOQ		mg/4g	0.122	
THCV-A per 4g	< LOQ		mg/4g	0.122	
THCV-Total per 4g	< LOQ		mg/4g	0.229	
Total Cannabinoids per 4g	31.6		mg/4g		



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Microbiology

Analyte	Result	Limits	Units	LOQ	Batch	Analyzed Method	Status	Notes
E.coli	< LOQ		cfu/g	10	2312884	11/19/23 AOAC 991.14 (Petrifilm) ^P		
Total Coliforms	< LOQ		cfu/g	10	2312884	11/19/23 AOAC 991.14 (Petrifilm) ^P		
Mold (RAPID Petrifilm)	< LOQ		cfu/g	10	2312885	11/20/23 AOAC 2014.05 (RAPID) ^P		
Yeast (RAPID Petrifilm)	< LOQ		cfu/g	10	2312885	11/20/23 AOAC 2014.05 (RAPID) ^P		

Solvents Method: Residual Solvents by GC/MS^P Units µg/g Batch 2313025 Analyze 11/22/23 10:08 AM

Analyte	Result	Limits	LOQ	Status	Notes	Analyte	Result	Limits	LOQ	Status	Notes
1,4-Dioxane	< LOQ	380	100	pass		2-Butanol	< LOQ	5000	200	pass	
2-Ethoxyethanol	< LOQ	160	30.0	pass		2-Methylbutane (Isopentane)	< LOQ		200		
2-Methylpentane	< LOQ		30.0			2-Propanol (IPA)	< LOQ	5000	200	pass	
2,2-Dimethyl butane	< LOQ		30.0			2,2-Dimethylpropane (neo-pentane)	< LOQ		200		
2,3-Dimethyl butane	< LOQ		30.0			3-Methylpentane	< LOQ		30.0		
Acetone	< LOQ	5000	200	pass		Acetonitrile	< LOQ	410	100	pass	
Benzene	< LOQ	2.00	1.00	pass		Butanes (sum)	< LOQ	5000	400	pass	
Cyclohexane	< LOQ	3880	200	pass		Ethyl acetate	< LOQ	5000	200	pass	
Ethyl benzene	< LOQ		200			Ethyl ether	< LOQ	5000	200	pass	
Ethylene glycol	< LOQ	620	200	pass		Ethylene oxide	< LOQ	50.0	20.0	pass	
Hexanes (sum)	< LOQ	290	150	pass		Isopropyl acetate	< LOQ	5000	200	pass	
Isopropyl benzene (Cumene)	< LOQ	70.0	30.0	pass		m,p-Xylene	< LOQ		200		
Methanol	< LOQ	3000	200	pass		Methylene chloride	< LOQ	600	60.0	pass	
Methylpropane (Isobutane)	< LOQ		200			n-Butane	< LOQ		200		
n-Heptane	< LOQ	5000	200	pass		n-Hexane	< LOQ		30.0		
n-Pentane	< LOQ		200			o-Xylene	< LOQ		200		
Pentanes (sum)	< LOQ	5000	600	pass		Propane	< LOQ	5000	200	pass	
Tetrahydrofuran	< LOQ	720	100	pass		Toluene	< LOQ	890	100	pass	
Total Xylenes	< LOQ		400			Total Xylenes and Ethyl benzene	< LOQ	2170	600	pass	

Pesticides Method: AOAC 2007.01 & EN 15662 (mod)^P Units mg/kg Batch 2313035 Analyze 11/22/23 12:14 PM

Analyte	Result	Limits	Status	Notes
Multi-Residue Pesticide Profile	< LOQ for all analytes			



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Metals

Analyte	Result	Limits	Units	LOQ	Batch	Analyzed Method	Status	Notes
Arsenic*	< LOQ	0.200	mg/kg	0.0156	2313045	11/22/23 AOAC 2013.06 (mod.) ^p	pass	
Cadmium*	< LOQ	0.200	mg/kg	0.0156	2313045	11/22/23 AOAC 2013.06 (mod.) ^p	pass	
Lead*	< LOQ	0.500	mg/kg	0.0156	2313045	11/22/23 AOAC 2013.06 (mod.) ^p	pass	
Mercury*	< LOQ	0.100	mg/kg	0.00778	2313045	11/22/23 AOAC 2013.06 (mod.) ^p	pass	

Nutrition

Analyte	Result	Limits	Units	LOQ	Batch	Analyzed Method	Status	Notes
Moisture (Loss on Drying)	17.9		g/100g	0.10	2312977	11/20/23 AOAC 925.10 (mod.) ^p		
Water Activity	0.670		Aw	0.030	2312903	11/17/23 AOAC 978.18 ^p		



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Abbreviations

Limits: Action Levels per OAR-333-007-0400, OAR-333-007-0210, OAR-333-007-0220, CCR title 16-division 42. BCC-section 5723

Limit(s) of Quantitation (LOQ): The minimum levels, concentrations, or quantities of a target variable (e.g., target analyte) that can be reported with a specified degree of confidence.

Ⓟ = ISO/IEC 17025:2017 accredited method.

Ⓢ = TNI accredited analyte.

Units of Measure

cfu/g = Colony forming units per gram

g = g

g/100g = Grams per 100 Grams

µg/g = Microgram per gram

mg/kg = Milligram per kilogram = parts per million (ppm)

mg/4g = Milligram per 4g

% = Percentage of sample

Aw = Water Activity

% wt = µg/g divided by 10,000

Approved Signatory

Derrick Tanner
General Manager



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Compound	LOQ (mg/kg)	Compound	LOQ (mg/kg)	Compound	LOQ (mg/kg)
Abamec in	0.100	Cle hodim	0.050	ndrin	0.100
Acepha e	0.100	Cle hodim Sul one	0.050	PN	0.050
Acequinocyl	0.100	Cle hodim Sul oxide	0.050	PTC	0.100
Ace amiprid	0.020	Clo en ezine	0.020	s envalera e/ envalera e	0.200
Ace ochlor	0.100	Clomazone	0.020	aconazole	0.100
Acrina hrin	0.100	Clo hianidin	0.200	hal luralin	0.100
Alachlor	0.100	Coumaphos	0.050	hio encarb	0.050
Aldicarb	0.100	Cro oxyphos	0.020	hion	0.200
Aldicarb sul oxide	0.100	Cyanazine	0.020	hirimol	0.100
Aldoxycarb (Aldicarb-sul one)	0.100	Cyano enphos	0.020	ho umesa e	0.050
Aldrin	0.100	Cyan raniliprole	0.050	hoprophos	0.020
Ame ocr radin	0.020	Cyazo amid	0.020	o enprox	0.020
Ame ryn	0.500	Cycloa e	0.100	oxazole	0.020
Aspon	0.100	Cy lu hrin	0.200	ridiazole	0.100
Asulam	0.100	Cyhalo hrin, lambda	0.200	rim os	0.020
A razine	0.100	Cymoxanil	0.050	amoxadone	0.200
A razine-dese hyl	0.100	Cyperme hrin	0.200	amphur	0.100
Azinphos-e hyl	0.020	Cyprodinil	0.100	enamidone	0.020
Azinphos-me hyl	0.020	Dac hal	0.100	enamiphos	0.020
Azoxys robin	0.020	Daminozide	0.100	enamiphos sul one	0.020
Benalaxyl	0.020	DCPMU	0.050	enamiphos sul oxide	0.020
Bendiocarb	0.020	DDD, o,p'-	0.100	enazaquin	0.100
Ben luralin	0.100	DDD, p,p'-	0.100	enbuconazole	0.100
Benoxacor	0.050	DD , o,p'-	0.100	enchlorphos	0.100
Bensulide	0.050	DD , p,p'-	0.100	enchlorphos-oxon	0.100
B C alpha isomer	0.100	DDT, o,p'-	0.100	enhexamid	0.100
B C be a isomer	0.100	DDT, p,p'-	0.100	eni ro hion	0.100
B C del a isomer	0.500	D (Tribu os)	0.100	enobucarb	0.050
Bi enaza e	0.020	Del ame hrin	0.100	enoxy carb	0.020
Bi en hrin	0.020	Desmedipham	0.100	enpropa hrin	0.050
Boscalid	0.020	Dialla e	0.100	enpyroxima e	0.020
Bromophos-e hyl	0.100	Diazinon	0.020	enson	0.100
Bromophos-me hyl	0.200	Diazoxon	0.100	ensul o hion	0.020
Bromopropyla e	0.100	Dichlobenil	0.100	ensul o hion oxon	0.020
Bromuconazole	0.100	Dichlo luanid	0.100	ensul o hion sul one	0.100
Bupirima e	0.020	Dichlorvos	0.100	Fensulfothion-oxon-sulfone	0.020
Bupro ezin	0.050	Diclobu razol	0.050	en hion	0.050
Bu achlor	0.500	Dico ol	0.100	en hion oxon	0.020
Bu ralin	0.200	Dicro ophos	0.050	en hion oxon sul one	0.100
Bu yla e	0.100	Dieldrin	0.100	en hion sul one	0.050
Cadusa os	0.020	Die ho encarb	0.020	enuron	0.020
Cap an	1.000	Die hyl oluamide (D T)	0.050	ipronil	0.100
Carbaryl	0.050	Di enoconazole	0.100	lonicamid	0.100
Carbendazim	0.100	Dime henamid	0.050	luchloralin	0.100
Carbo uran	0.020	Dime hoa e	0.050	lucy hrina e	0.100
Carbopheno hion	0.200	Dime homorph	0.050	ludioxonil	0.200
Carboxin	0.020	Diniconazole	0.200	lu enace	0.020
Car en razone-e hyl	0.100	Dino e uran	0.200	lumioxazin	0.100
Chloran raniliprole	0.020	Dioxa hion	0.100	luome uron	0.020
Chlordane, cis-	0.200	Diphenamid	0.020	luopicolide	0.050
Chlordane, rans-	0.200	Diphenylamine	0.100	luopyram	0.020
Chlor enapyr	0.500	Disul o on	0.100	luoxas robin	0.050
Chlor enson	0.200	Disul o on sul one	0.100	lupyradi urone	0.020
Chlor envinphos	0.050	Disul o on sul oxide	0.100	luridone	0.100
Chlorobenzila e	0.100	Diuron	0.050	lusilazole	0.020
Chloroneb	0.200	di enphos	0.050	lu olanil	0.020
Chlorpyri os	0.050	ndosul an alpha	0.200	lu ria ol	0.020
Chlorpyri os-me hyl	0.200	ndosul an be a	0.200	luvalina e, au-	0.100
C PC	1.000	ndosul an sul a e	0.100	luxapyroxad	0.020



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Cannab s Mu t -Res due Prof e, L m ts of Quant tat on

Compound	LOQ (mg/kg)	Compound	LOQ (mg/kg)	Compound	LOQ (mg/kg)
omesa en	0.100	Mexacarba e	0.020	Propamocarb	0.050
ono os	0.100	MGK 264	0.020	Propanil	0.050
orchlor enuron	0.050	Mirex	0.100	Propargi e	0.050
orme ana e	0.050	Molina e	0.050	Propazine	0.020
ura hiocarb	0.020	Monocro ophos	0.100	Prope amphos	0.050
ep achlor	0.100	Monolinuron	0.020	Propham	0.050
ep achlor epoxide	0.100	Myclobu anil	0.050	Propiconazole	0.050
ep enphos	0.100	Naled	0.100	Propoxur	0.050
exachlorobenzene	0.100	Napropamide	0.050	Propoxycarbazono Na	0.050
exaconazole	0.100	Neburon	0.020	Propyzamide	0.050
exazinone	0.100	Ni rapyrin	0.100	Pro hio os	0.100
exy hiazox	0.020	Nor lurazon	0.050	Pyraclos robin	0.020
mazalil	0.100	Ome hoa e	0.100	Pyrazophos	0.050
midacloprid	0.100	O-Phenylphenol	0.100	Pyre hrins	0.050
ndazi lam	0.020	Oxadixyl	0.100	Pyridaben	0.020
ndoxacarb	0.020	Oxamyl	0.100	Pyrida ol	0.100
proben os	0.100	Oxamyl-oxime	0.100	Pyrida e	0.020
prodione	0.100	Oxychlorthane	0.100	Pyrimo hanil	0.050
sobenzan	0.100	Oxydeme on-Me hyl	0.100	Pyriproxi en	0.020
socarbophos	0.500	Oxy hioquinox	0.200	Pyroxasul one	0.020
sodrin	0.100	Paclobu razol	0.050	Pyroxulam	0.020
so enphos	0.050	Paraaxon-e hyl	0.020	Quinalphos	0.050
so enphos-me hyl	0.020	Paraaxon me hyl	0.100	Quinoxy en	0.050
so enphos oxon	0.050	Para hion e hyl	0.100	Quin ozene (PCNB)	0.200
soproc carb	0.020	Para hion me hyl	0.200	Resme hrin	0.050
sopropalin	0.200	Penconazole	0.050	Ro enone	0.050
sopro hiolane	0.050	Pendime halin	0.050	S421	0.100
sopro uron	0.050	Pen lu en	0.020	Simazine	0.100
soxaben	0.050	Pen achloroaniline	0.100	Sime ryn	0.200
soxa lu ole	0.050	Pen achloroanisole	0.100	Spine oram	0.020
Kresoxim-me hyl	0.050	Pen achlorobenzene (PCB)	0.100	Spinosa d	0.050
ac o en	0.500	Pentachlorothioanisole (PCTA)	0.100	Spirodiclo en	0.100
enacil	0.100	Pen hiopyrad	0.020	Spiromesi en	0.050
indane (gamma B C)	0.100	Perme hrin	0.050	Spiro e rama	0.050
inuron	0.020	Per hane	0.100	Spiroxamine	0.020
Malaaxon	0.050	Phenmedipham	0.050	Sul o ep	0.050
Mala hion	0.050	Phen hoa e	0.050	Sul oxa lor	0.050
Mandipropamid	0.020	Phora e	0.050	Sulpro os	0.020
Mecarbam	0.020	Phora e Sul one	0.050	Tebuconazole	0.100
Mepanipyrim	0.050	Phora e Sul oxide	0.050	Tebu enozide	0.020
Merphos	0.500	Phosalone	0.050	Tebu hiuron	0.020
Me alaxyl	0.050	Phosme	0.100	Tecnazene	0.100
Me aldehyde	0.050	Phosphamidon	0.050	Te lu hrin	0.100
Me conazole	0.100	Phoxim	0.050	Terbu os	0.020
Me hacri os	0.100	Pinoxaden	0.020	Terbu os sul one	0.050
Me hamidophos	0.050	Piperonyl bu oxide	0.050	Terbu os sul oxide	0.050
Me hida hion	0.050	Pirimicarb	0.020	Terbu hylazine	0.020
Me hiocarb	0.050	Pirimiphos-me hyl	0.050	Terbu ryn	0.020
Me hiocarb sul one	0.100	Pirimiphos-e hyl	0.020	Te rachlorvinphos	0.050
Me hiocarb sul oxide	0.100	Pralle hrin	0.100	Te raconazole	0.050
Me homyl	0.100	Prochloraz	0.020	Te radi on	0.200
Me hoxychlor	0.100	Procyimidone	0.100	Te rame hrin	0.050
Me hoxy enozide	0.020	Pro eno os	0.100	Te rasul	0.100
Me obromuron	0.050	Pro luralin	0.100	Thiabendazole	0.100
Me olachlor	0.100	Promecarb	0.050	Thiabendazole, 5-hydroxy	0.100
Me olcarb	0.050	Prome on	0.100	Thiacloprid	0.050
Me ra enone	0.050	Prome ryn	0.020	Thiame hoxam	0.100
Me ribuzin	0.100	Propachlor	0.020	Thiobencarb	0.050
Mevinphos	0.100			Thiodicarb	0.050
				Thiophana e-me hyl	0.050



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Compound	LOQ (mg/kg)	Compound	LOQ (mg/kg)	Compound	LOQ (mg/kg)
Tolclo os-me hyl	0.100	Triazophos	0.020	Tri loxys robin	0.020
Tri orin	0.100	Tolyl luanid	0.050	Tri iconazole	0.050
Tralkoxydim	0.100	Tridiphane	0.500	Vinclozolin	0.100
Triadime on	0.050	Tri lumizole	0.020	Zoxamide	0.020
Trialla e	0.100	Tri luralin	0.100		

LOQ= Limit of Quantitation, mg/kg

Factors affecting the LOQ include instrument sensitivity or a particular analyte, sample size, moisture content (percent solids) of the sample, effectiveness of the cleanup on the sample extract, and especially the type of sample matrix.



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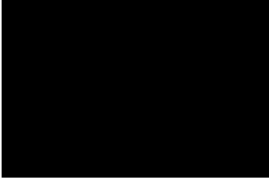
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**Hemp & Cannabis
Chain of Custody**

**Northwest-Natural
Goods-1700084244**

ORELAP ID: **OR1000028** ANAB ISO 17025 ID: **ATI508**

Contact Information Company: <u>Northwest Natural Goods</u> 		Project Details Turnaround Time: <u>5 Business Days / Rep. For Micro Testing / Standard</u> Sample Relinquishment Options: <u>Click-Up Request</u> Compliance: <u>Compliance</u> Project Name / ID: <u>HEMP-PR0066</u> Cannabis Type (select if applicable): <u>Industrial</u>			Testing H0010 - Potency Cannabinoid Basic + Extended Profile F0020 - Fastfold - Multi-Residue Profile H0008 - Residual Solvents - CF H0013 - Heavy Metals Profile (As, Cd, Pb & Hg) M075 - Total Coliforms - EColi P003 - Yeast and Mold N150 - Moisture Loss on Drying N300 - Water Activity								
#	Sample Name / Test	Material	Amount Provided	Reporting Unit	Serving Size	H0010	F0020	H0008	H0013	M075	P003	N150	N300
1	HEMP-PR0066	Edible	20 units for sale	mg/g & mg/-serving	g	✓	✓	✓	✓	✓	✓	✓	✓

Relinquished By	Date	Time	Temp., °C	Received By	Date	Time	Received Temp., °C	Evidence of Cooling?
<i>Amber Johnson</i>	<i>11/15/2023</i>	<i>13:37</i>		<i>BR</i>	<i>11/16/2023</i>	<i>10:37</i>		<i>No</i>
<i>BR</i>	<i>11/16/2023</i>	<i>11:17</i>	<i>18.6</i>	<i>BR</i>	<i>11/16/2023</i>	<i>12:12</i>		<i>No</i>

Samples submitted to Columbia Laboratories with testing requirements constitute an agreement for services in accordance with the [current terms of service](#) associated with NSCOC. By signing Relinquished by you are agreeing to these terms.

Columbia Laboratories
2423 NE Whitaker Way
Portland, OR 97233

P: (503) 254-1794 / Fax: (503) 254-1462
<http://www.columbialaboratories.com>

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Revision: 4 Document ID: 7148
 Legacy ID: Worksheet Validated 04/20/2021

Laboratory Quality Control Results

JAOAC2015 V986 Batch ID: 2312958

Laboratory Control Sample									
Analyte	LCS	Result	Spike	Units	% Rec	Limits	Evaluation	Notes	
CBDA	2	0.0329	0.0329	%	100	80.0 - 120	Acceptable		
CBDA	2	0.0324	0.0324	%	100	80.0 - 120	Acceptable		
CBE	2	0.0348	0.0349	%	99.6	80.0 - 120	Acceptable		
CBDA	1	0.0311	0.0311	%	100	90.0 - 110	Acceptable		
CBSA	1	0.0317	0.0313	%	101	80.0 - 120	Acceptable		
CBS	1	0.0334	0.0332	%	101	80.0 - 120	Acceptable		
CB	1	0.0317	0.0319	%	99.6	90.0 - 110	Acceptable		
THCV	2	0.0330	0.0337	%	97.9	80.0 - 120	Acceptable		
deltaTHCV	2	0.0276	0.0271	%	102	80.0 - 120	Acceptable		
THCVA	2	0.0317	0.0317	%	100.0	80.0 - 120	Acceptable		
CBN	1	0.0335	0.0332	%	101	80.0 - 120	Acceptable		
exo-THC	2	0.0304	0.0308	%	98.8	80.0 - 120	Acceptable		
deltaTHC	1	0.0329	0.0320	%	103	90.0 - 110	Acceptable		
deltaTHC	1	0.0271	0.0277	%	97.9	90.0 - 110	Acceptable		
9SdeltaTHC	1	0.0328	0.0328	%	100	80.0 - 120	Acceptable		
CB	2	0.0335	0.0351	%	95.4	80.0 - 120	Acceptable		
9SHHC	3	0.0294	0.0292	%	100	80.0 - 120	Acceptable		
9RdeltaTHC	1	0.0311	0.0314	%	99.2	80.0 - 120	Acceptable		
CB	2	0.0340	0.0341	%	99.7	80.0 - 120	Acceptable		
9RHHC	3	0.0868	0.0883	%	98.4	80.0 - 120	Acceptable		
THCA	1	0.0311	0.0303	%	103	90.0 - 110	Acceptable		
CBSA	2	0.0329	0.0336	%	98.0	80.0 - 120	Acceptable		
CBSA	2	0.0329	0.0336	%	97.9	80.0 - 120	Acceptable		
deltaTHCP	2	0.0325	0.0331	%	98.2	80.0 - 120	Acceptable		
deltaTHCO	3	0.0340	0.0334	%	102	80.0 - 120	Acceptable		
CB	2	0.0333	0.0337	%	98.7	80.0 - 120	Acceptable		
deltaTHCO	3	0.0326	0.0329	%	99.2	80.0 - 120	Acceptable		

Method Blank

Analyte	Result	LOQ	Units	Limits	Evaluation	Notes
CBDA	<LOQ	0.00316	%	< 0.00316	Acceptable	
CBDA	<LOQ	0.00316	%	< 0.00316	Acceptable	
CBE	<LOQ	0.00316	%	< 0.00316	Acceptable	
CBDA	<LOQ	0.00316	%	< 0.00316	Acceptable	
CBSA	<LOQ	0.00316	%	< 0.00316	Acceptable	
CBS	<LOQ	0.00316	%	< 0.00316	Acceptable	
CB	<LOQ	0.00316	%	< 0.00316	Acceptable	
THCV	<LOQ	0.00316	%	< 0.00316	Acceptable	
deltaTHCV	<LOQ	0.00316	%	< 0.00316	Acceptable	
THCVA	<LOQ	0.00316	%	< 0.00316	Acceptable	
CBN	<LOQ	0.00316	%	< 0.00316	Acceptable	
exo-THC	<LOQ	0.00316	%	< 0.00316	Acceptable	
deltaTHC	<LOQ	0.00316	%	< 0.00316	Acceptable	
deltaTHC	<LOQ	0.00316	%	< 0.00316	Acceptable	
9SdeltaTHC	<LOQ	0.00316	%	< 0.00316	Acceptable	
CB	<LOQ	0.00316	%	< 0.00316	Acceptable	
9SHHC	<LOQ	0.00316	%	< 0.00316	Acceptable	
9RdeltaTHC	<LOQ	0.00316	%	< 0.00316	Acceptable	
CB	<LOQ	0.00316	%	< 0.00316	Acceptable	
9RHHC	<LOQ	0.00316	%	< 0.00316	Acceptable	
THCA	<LOQ	0.00316	%	< 0.00316	Acceptable	
CBSA	<LOQ	0.00316	%	< 0.00316	Acceptable	
CBSA	<LOQ	0.00316	%	< 0.00316	Acceptable	
deltaTHCP	<LOQ	0.00316	%	< 0.00316	Acceptable	
deltaTHCO	<LOQ	0.00316	%	< 0.00316	Acceptable	
CB	<LOQ	0.00316	%	< 0.00316	Acceptable	
deltaTHCO	<LOQ	0.00316	%	< 0.00316	Acceptable	

Abbreviations

ND - None Detected at or above MRL
 RPD - Relative Percent Difference
 LOQ - Limit of Quantitation

Units of Measure:

% - Percent



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Laboratory Quality Control Results

JAOAC2015 V986		Batch ID: 2312958						
Sample Duplicate		Sample ID: 23-0132800002						
Analyte	Result	Org. Result	LOQ	Units	RPD	Limits	Evaluation	Notes
CBDA	0.0111	0.0113	0.00318	%	1.27	< 20	Acceptable	
CBDV	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	
CBE	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	
CBDA	0.0215	0.0211	0.00318	%	1.78	< 20	Acceptable	
CBGA	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	
CBG	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	
CB	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	
THCV	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	
d8THCV	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	
THCVA	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	
CBN	0.00674	0.00622	0.00318	%	8.09	< 20	Acceptable	
exo-THC	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	
d9THC	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	
d8THC	0.650	0.621	0.00318	%	4.70	< 20	Acceptable	
9Sd10THC	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	
CB	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	
9SHHC	0.0800	0.0791	0.00318	%	1.11	< 20	Acceptable	
9Rd10THC	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	
CB	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	
9RHHC	0.199	0.194	0.00318	%	2.44	< 20	Acceptable	
THCA	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	
CBGA	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	
CBGA	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	
d9THCP	0.0168	0.0172	0.00318	%	2.75	< 20	Acceptable	
d8THCO	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	
CB	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	
d9THCO	<LOQ	<LOQ	0.00318	%	NA	< 20	Acceptable	

Abbreviations

ND - None Detected at or above MRL
 RPD - Relative Percent Difference
 LOQ - Limit of Quantitation

Units of Measure:

% - Percent



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Laboratory Quality Control Results

Residual Solvents				Batch D: 2313025					
Method Blank				Laboratory Control Sample					
Analyte	Result	LOQ	Notes	Result	Spike	Units	% Rec	Limits	Notes
Propane	ND	< 200		476	584	µg/g	81.5	60 - 120	
Isobutane	ND	< 200		533	767	µg/g	69.5	60 - 120	
Butane	ND	< 200		499	782	µg/g	63.8	60 - 120	
2,2-Dimethylpropane	ND	< 200		726	939	µg/g	77.3	60 - 120	
Methanol	ND	< 200		1820	1600	µg/g	113.8	60 - 120	
Ethylene Oxide	ND	< 30		39.3	57.1	µg/g	68.8	60 - 120	
2-Methylbutane	ND	< 200		1750	1600	µg/g	109.4	60 - 120	
Pentane	ND	< 200		1760	1600	µg/g	110.0	60 - 120	
Ethanol	ND	< 200		1720	1600	µg/g	107.5	70 - 130	
Ethyl Ether	ND	< 200		1700	1600	µg/g	106.3	60 - 120	
2,2-Dimethylbutane	ND	< 30		170	161	µg/g	105.6	60 - 120	
Acetone	ND	< 200		1740	1600	µg/g	108.8	60 - 120	
2-Propanol	ND	< 200		1720	1600	µg/g	107.5	60 - 120	
Ethyl Formate	ND	< 500		1150	1600	µg/g	71.9	70 - 130	
Acetonitrile	ND	< 100		530	488	µg/g	108.6	60 - 120	
Methyl Acetate	ND	< 500		1500	1610	µg/g	93.2	70 - 130	
2,3-Dimethylbutane	ND	< 30		172	163	µg/g	105.5	60 - 120	
Dichloromethane	ND	< 60		477	488	µg/g	97.7	60 - 120	
2-Methylpentane	ND	< 30		150	161	µg/g	93.2	60 - 120	
MTBE	ND	< 500		1540	1650	µg/g	93.3	70 - 130	
3-Methylpentane	ND	< 30		167	162	µg/g	103.1	60 - 120	
Hexane	ND	< 30		168	161	µg/g	104.3	60 - 120	
1-Propanol	ND	< 500		1700	1620	µg/g	104.9	70 - 130	
Methyl ethyl ketone	ND	< 500		1530	1610	µg/g	95.0	70 - 130	
Ethyl acetate	ND	< 200		1740	1610	µg/g	108.1	60 - 120	
2-Butanol	ND	< 200		1690	1610	µg/g	105.0	60 - 120	
Tetrahydrofuran	ND	< 100		499	483	µg/g	103.3	60 - 120	
Cyclohexane	ND	< 200		1660	1600	µg/g	103.8	60 - 120	
2-methyl-1-propanol	ND	< 500		1350	1600	µg/g	84.4	70 - 130	
Benzene	ND	< 1		5.11	4.99	µg/g	102.4	60 - 120	
Isopropyl Acetate	ND	< 200		1770	1600	µg/g	110.6	60 - 120	
Heptane	ND	< 200		1750	1600	µg/g	109.4	60 - 120	
1-Butanol	ND	< 500		1480	1610	µg/g	90.7	70 - 130	
Propyl Acetate	ND	< 500		1590	1610	µg/g	98.8	70 - 130	
1,4-Dioxane	ND	< 100		470	480	µg/g	97.9	60 - 120	
2-Ethoxyethanol	ND	< 30		134	161	µg/g	83.2	60 - 120	
Methylisobutylketone	ND	< 500		1620	1610	µg/g	100.6	70 - 130	
3-Methyl-1-butanol	ND	< 500		1520	1610	µg/g	94.4	70 - 130	
Ethylene Glycol	ND	< 200		197	481	µg/g	41.0	60 - 120	Q6
Toluene	ND	< 100		487	483	µg/g	100.8	60 - 120	
Isobutyl Acetate	ND	< 500		1600	1610	µg/g	99.4	70 - 130	
1-Pentanol	ND	< 500		1570	1610	µg/g	97.5	70 - 130	
Butyl Acetate	ND	< 500		1580	1600	µg/g	97.5	70 - 130	
Ethylbenzene	ND	< 200		988	982	µg/g	100.4	60 - 120	
m,p-Xylene	ND	< 200		981	972	µg/g	100.9	60 - 120	
o-Xylene	ND	< 200		942	965	µg/g	97.6	60 - 120	
Cumene	ND	< 30		165	169	µg/g	97.6	60 - 120	
Anisole	ND	< 500		1320	1600	µg/g	82.5	70 - 130	
DMF	ND	< 500		1070	1600	µg/g	66.9	70 - 130	Q6
1,2-dimethoxyethane	ND	< 50		151	163	µg/g	92.6	70 - 130	
Triethylamine	ND	< 500		1190	1600	µg/g	74.4	70 - 130	
N,N-dimethylformamide	ND	< 150		400	482	µg/g	83.0	70 - 130	
N,N-dimethylacetamide	ND	< 150		478	483	µg/g	99.0	70 - 130	
Pyridine	ND	< 50		129	161	µg/g	80.1	70 - 130	
Silofane	ND	< 50		96.6	163	µg/g	59.3	70 - 130	Q6
1,2-Dichloroethane	ND	< 1		1.13	1	µg/g	113.0	70 - 130	
Chloroform	ND	< 1		0.994	1	µg/g	99.4	70 - 130	
Trichloroethylene	ND	< 1		0.867	1	µg/g	86.7	70 - 130	
1,1,1-Trichloroethane	ND	< 1		1.08	1	µg/g	108.0	70 - 130	



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QC- Sample Duplicate		Sample ID: 23-013428-0001						
Analyte	Result	Org. Result	LOQ Units	RPD	Limits	Accept/ Fail	Notes	
Propane	ND	ND	200 µg/g	0.0	< 20	Acceptable		
Isobutane	ND	ND	200 µg/g	0.0	< 20	Acceptable		
Butane	ND	ND	200 µg/g	0.0	< 20	Acceptable		
2,2-Dimethylpropane	ND	ND	200 µg/g	0.0	< 20	Acceptable		
Methanol	ND	ND	200 µg/g	0.0	< 20	Acceptable		
Ethylene Oxide	ND	ND	30 µg/g	0.0	< 20	Acceptable		
2-Methylbutane	ND	ND	200 µg/g	0.0	< 20	Acceptable		
Pentane	ND	ND	200 µg/g	0.0	< 20	Acceptable		
Ethanol	ND	ND	200 µg/g	0.0	< 20	Acceptable		
Ethyl Ether	ND	ND	200 µg/g	0.0	< 20	Acceptable		
2,2-Dimethylbutane	ND	ND	30 µg/g	0.0	< 20	Acceptable		
Acetone	ND	ND	200 µg/g	0.0	< 20	Acceptable		
2-Propanol	ND	ND	200 µg/g	0.0	< 20	Acceptable		
Ethyl Formate	ND	ND	500 µg/g	0.0	< 20	Acceptable		
Acetonitrile	ND	ND	100 µg/g	0.0	< 20	Acceptable		
Methyl Acetate	ND	ND	500 µg/g	0.0	< 20	Acceptable		
2,3-Dimethylbutane	ND	ND	30 µg/g	0.0	< 20	Acceptable		
Dichloromethane	ND	ND	60 µg/g	0.0	< 20	Acceptable		
2-Methylpentane	ND	ND	30 µg/g	0.0	< 20	Acceptable		
MTBE	ND	ND	500 µg/g	0.0	< 20	Acceptable		
3-Methylpentane	ND	ND	30 µg/g	0.0	< 20	Acceptable		
Hexane	ND	ND	30 µg/g	0.0	< 20	Acceptable		
1-Propanol	ND	ND	500 µg/g	0.0	< 20	Acceptable		
Methylethylketone	ND	ND	500 µg/g	0.0	< 20	Acceptable		
Ethyl acetate	ND	ND	200 µg/g	0.0	< 20	Acceptable		
2-Butanol	ND	ND	200 µg/g	0.0	< 20	Acceptable		
Tetrahydrofuran	ND	ND	100 µg/g	0.0	< 20	Acceptable		
Cyclohexane	ND	ND	200 µg/g	0.0	< 20	Acceptable		
2-methyl-1-propanol	ND	ND	500 µg/g	0.0	< 20	Acceptable		
Benzene	ND	ND	1 µg/g	0.0	< 20	Acceptable		
Isopropyl Acetate	ND	ND	200 µg/g	0.0	< 20	Acceptable		
Heptane	ND	ND	200 µg/g	0.0	< 20	Acceptable		
1-Butanol	ND	ND	500 µg/g	0.0	< 20	Acceptable		
Propyl Acetate	ND	ND	500 µg/g	0.0	< 20	Acceptable		
1,4-Dioxane	ND	ND	100 µg/g	0.0	< 20	Acceptable		
2-Ethoxyethanol	ND	ND	30 µg/g	0.0	< 20	Acceptable		
Methylisobutylketone	ND	ND	500 µg/g	0.0	< 20	Acceptable		
3-Methyl-1-butanol	ND	ND	500 µg/g	0.0	< 20	Acceptable		
Ethylene Glycol	ND	ND	200 µg/g	0.0	< 20	Acceptable		
Toluene	ND	ND	100 µg/g	0.0	< 20	Acceptable		
Isobutyl Acetate	ND	ND	500 µg/g	0.0	< 20	Acceptable		
1-Pentanol	ND	ND	500 µg/g	0.0	< 20	Acceptable		
Butyl Acetate	ND	ND	500 µg/g	0.0	< 20	Acceptable		
Ethylbenzene	ND	ND	200 µg/g	0.0	< 20	Acceptable		
m,p-Xylene	ND	ND	200 µg/g	0.0	< 20	Acceptable		
o-Xylene	ND	ND	200 µg/g	0.0	< 20	Acceptable		
Cumene	ND	ND	30 µg/g	0.0	< 20	Acceptable		
Anisole	ND	ND	500 µg/g	0.0	< 20	Acceptable		
DMSO	ND	ND	500 µg/g	0.0	< 20	Acceptable		
1,2-dimethoxyethane	ND	ND	50 µg/g	0.0	< 20	Acceptable		
Triethylamine	ND	ND	500 µg/g	0.0	< 20	Acceptable		
N,N-dimethylformamide	ND	ND	150 µg/g	0.0	< 20	Acceptable		
N,N-dimethylacetamide	ND	ND	150 µg/g	0.0	< 20	Acceptable		
Pyridine	ND	ND	50 µg/g	0.0	< 20	Acceptable		
Sulfolane	ND	ND	50 µg/g	0.0	< 20	Acceptable		
1,2-Dichloroethane	ND	ND	1 µg/g	0.0	< 20	Acceptable		
Chloroform	ND	ND	1 µg/g	0.0	< 20	Acceptable		
Trichloroethylene	ND	ND	1 µg/g	0.0	< 20	Acceptable		
1,1-Dichloroethane	ND	ND	1 µg/g	0.0	< 20	Acceptable		

Abbreviations

ND - None Detected at or above MRL
 RPD- Relative Percent Difference
 LOQ - Limit of Quantitation
 Q6- Quality control outside QClimits. Data acceptable based on remaining QC.

Units of Measure:

µg/g- Microgram per gram or ppm



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Explanation of QC Flag Comments:

Code	Explanation
Q	Matrix interferences affecting spike or surrogate recoveries.
Q1	Quality control result biased high. Only non-detect samples reported.
Q2	Quality control outside QC limits. Data considered estimate.
Q3	Sample concentration greater than four times the amount spiked.
Q4	Non-homogenous sample matrix, affecting RPD result and/or % recoveries.
Q5	Spike results above calibration curve.
Q6	Quality control outside QC limits. Data acceptable based on remaining QC.
R	Relative percent difference (RPD) outside control limit.
R1	RPD non-calculable, as sample or duplicate results are less than five times the LOQ.
R2	Sample replicates RPD non-calculable, as only one replicate is within the analytical range.
LOQ1	Quantitation level raised due to low sample volume and/or dilution.
LOQ2	Quantitation level raised due to matrix interference.
B	Analyte detected in method blank, but not in associated samples.
B1	The sample concentration is greater than 5 times the blank concentration.
B2	The sample concentration is less than 5 times the blank concentration.