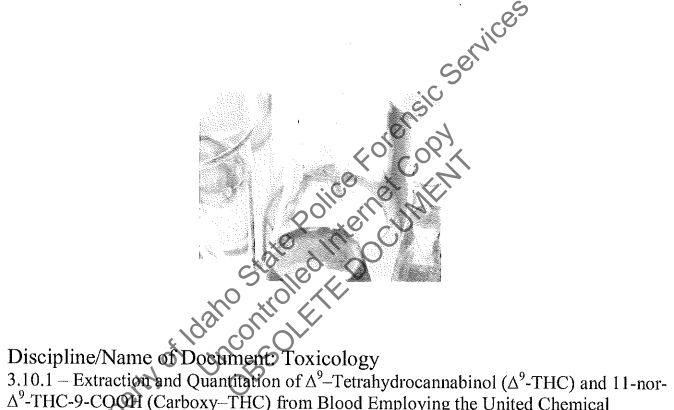
Idaho State Police Forensic Services

Approval for Quality System Controlled Documents



 Δ^9 -THC-9-COOH (Carboxy-THC) from Blood Employing the United Chemical Technologies (UCT) 200 mg CLEAN SCREEN® THC Extraction Column (FOR QUALITATIVE USE ONLY)

Revision Number: 1

Issue Date: 7/28/2008

APPROVED BY: (

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Section Three Blood Toxicology

3.10 SPE Methods for Quantitative GC/MSD Confirmation

3.10.1 Δ^9 -Tetrahydrocannabinol (Δ^9 -THC) and 11-nor- Δ^9 -THC-9-COOH (Carboxy-THC) from Blood Employing the United Chemical Technologies (UCT) 200 mg CLEAN SCREEN[®] THC Column

3.10.1.1 BACKGROUND

 Δ^9 -THC (Figure 1) is the chief psychoactive cannabinoid resulting from exposure to marijuana. Δ^9 -THC has a peak blood concentration within 5 to 15 minutes following smoking of a marijuana cigarette.^{3,4,5} This blood concentration drops rapidly after cessation of smoking.^{3,4} The level may fall to less than 5µL within 30 to 60 minutes although longer detection times have been reported.^{3,4} Detection of low dose (1.75%) post smoking Δ^9 -THC has been reported to vary from 3 to 12 hours. This detection window was based on a limit of quantitation of 0.5 ng/ml. The number, duration, and spacing of puffs, hold time, and inhalation volume all impact the degree of drug exposure and thus bioavailability. Longer detection times have been observed for frequent users. The Δ^{9} THC metabolite, 11-nor- Δ^{9} -THC-9-COOH (Carboxy-THC), concentration gradually increases and may plateau for several hours.⁴ There is poor correlation between blood Δ^9 -THC and psychoactive affects since the Δ^9 -THC concentrations begin to decline prior to the time of peak effects. Work continues on models using the relative amounts of Δ^9 -THC and Carboxy-THC to assist with establishing recent drug use.

Negative behavioral effects reported from exposure to marijuana include altered time perception, lack of concentration, impaired learning and memory which can lead to impairment of cognitive and performance tasks. Establishing impairment in an individual is based on evaluation of all available information in conjunction with the quantitative blood levels.

For additional background refer to analytical method 2.4.4 and provided references.

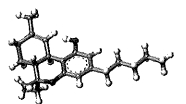


Figure 1.

3.10.1.2 PRINCIPLE

This procedure outlines the use of the 200mg United Chemical Technologies (UCT) CLEAN SCREEN® THC Column for the extraction from blood of the cannabinoids Δ^9 -THC and Carboxy-THC. The CLEAN SCREEN® THC column utilizes a copolymeric sorbent which combines a cationic exchanger and a hydrophobic functionality (reverse phase) to interact effectively, physically and chemically, with analytes of interest and minimally with interfering substances in the blood sample. The cation exchanger utilizes an anionic sorbent to bind to cations. Additional retention mechanisms include hydrophobic interactions and polar adsorption.

For the extraction and quantitation of Δ^9 -THC and Carboxy-THC, a deuterated internal standard of each is added to the sample. Blood proteins are precipitated with 10% Methanol-Acetonitrite solvent mix and are removed via centrifugation. The supernatant is transferred to a clean tube where the pH is adjusted with an Acetate buffer (pH 4.5), and loaded onto a pre-conditioned SPE column. The conditioning of the SPE column creates an environment, which allows for optimal interaction between the sorbent and the analytes of interest. The column is subsequently washed to selectively remove matrix components and interfering substances from the column. Next, the column is dried to remove traces of aqueous and organic solvents. After drying, the analytes of interest are eluted from the SPE column with an organic solvent mixture. Following elution and evaporation of the solvent, the extract is derivatized for compound confirmation using a gas chromatograph equipped with a mass selective detector (GC-MSD).

3.10.1.3 EQUIPMENT AND SUPPLIES

3.10.1.3.1	200mg CLEAN SCREEN® THC Extraction Column
3.10.1.32	Disposable inserts for SPE manifold ports.
3.10.1.3.3	Drybath or laboratory oven
3:10.1.3.4	Evaporative concentrator equipped with nitrogen tank.
3.10.1.3.5	Vortex mixer
3.10.1.3.6	Laboratory centrifuge capable of 3400rpm
3.10.1.3.7	Vacuum Manifold/ Vacuum pump
3.10.1.3.8	Fixed and adjustable volume single channel air displacement
	pipetters, and appropriate tips, capable of accurate and
	precise dispensing of volumes indicated.
3.10.1.3.9	16 x 100mm silanized glass tubes
3.10.1.3.10	Screw Cap for 16mm O.D. tubes
3.10.1.3.11	GC/MS Automated Liquid Sample (ALS) vials
3.10.1.3.12	Silanized GC/MS Vial Microinsert
3.10.1.3.13	Gas Chromatograph equipped with a quadrapole mass
	selective detector and a nonpolar capillary column with a
	phase composition comparable to 100%-

dimethylpolysiloxane or 95%-dimethyl-polysiloxane with 5%-diphenyl.

3.10.1.4 REAGENTS

Refer to manual sect	tion 5.12 for solution	n preparation instructions.
Teorer to mantitue soci	TOTE SIZE JOI DUCKTION	e proparator instructions

	in section of a joint solution propulation in all them
3.10.1.4.1	Deionized/distilled (DI) water
3.10.1.4.2	Methanol (Certified ACS Grade)
3.10.1.4.3	Hexane (Certified ACS Grade)
3.10.1.4.4	Ethyl Acetate (Certified ACS Grade)
3.10.1.4.5	Acetonitrile (Certified ACS Grade)
3.10,1,4,6	10% Methanol in Acetonitrile 100mM Acetate Buffer (pH 4.5)
3.10.1.4.7	100mM Acetate Buffer (pH 4.5)
3.10.1.4.8	100 317701
3.10.1.4.9	70:30 Hexane:Ethyl Acetate
3.10.1.4.10	70:30 100mM HCl: Acetonitrile C
3.10.1.4.11	BSTFA + 1% TMCS

QUALITY ASSURANCE MATERIAL 3.10.1.5

3.10.1.5.1 Calibrator and Control Solutions

3.10.1.5.1.1

Stock Solutions

Property of Janontroller Ca The source of a corresponding calibrator and control must be obtained from a different vendor.

Carboxy-THC and Δ^9 -THC

Concentration: 100µg/mL or 1000µg/mL

Working Solutions

Store remaining stock solution in ALS vial in freezer. Working solutions are stable for 6 months when stored at 4°C.

1ng/μL

As appropriate, add 100µL 100µg/mL or 10μL 1000μg/mL Stock Solution to approximately 9mL Methanol in a 10mL volumetric class A flask. OS to 10mL.

$0.1 \text{ng/}\mu\text{L}$

Add 1000µL lng/µL mixed working calibration solution to approximately 8mL Methanol in 10mL volumetric class A flask. QS to 10mL.

3.10.1.5.2

Internal Standard Solutions

3.10.1.5.2.1

Stock Solutions

 Δ^9 -THC-D₃ or Carboxy-THC-D₉

Concentration: 100µg/mL or 1000µg/mL

3.10.1.5.2.2

Working Internal Standard Solution [1ng/µL]

Add 100µL 100µg/mL or 10µL 1000µg/mL stock solutions to approximately 9mL Methanol in a 10mL volumetric class A flask. QS to 10mL. Solution is stable for six months when stored at 4°C.

3.10.1.5.3

Whole Blood Negative Control

Negative Whole Blood

3.10.1.6 PROCEDURE

3.10.1.6.1

Initial set-up

Label extraction tubes, 200mg CLEAN SCREEN® extraction columns, and GC/MSD vials with microinserts for calibrators, controls and case samples.

3.10.1.6.2

Calibrator Preparation

Use the same lot of negative blood used to prepare the negative control to prepare calibrators.

3.10.1.6.2.1

Add 1mL of negative whole blood to six extraction tubes.

3.10.1.6.2.

Add the volume of $0.1 \text{ng/}\mu\text{L}$ Δ^9 -THC and Carboxy-THC mixed calibrator working solution as indicated in the following table.

Level	Desired ng/mL	μL Working Reference Material
1	2.5	25
2	5	50
3	10	100

3.10.1.6.2.3

Add the volume of $lng/\mu L \Delta^9$ -THC and Carboxy-THC mixed calibrator working solution as indicated in the following table.

Level	Desired ng/mL	μL Working Reference Material
4	25	25
5	50	;C 50
6	1000	100

3.10.1.6.3 Positive Control Sample Preparation

Use the same lot of negative blood used to prepare the negative control for positive control preparation.

	_	1 60 COV	
	3.10.1.6.3.1		ve whole blood to two
	oc	extraction tubes.	
	3.10.1.6.3.2		amount of 0.1ng/μL
	CXOL 30	working mixed con	trol solution.
			1100
>	Shouth &	Desired ng/mL	μL Working Control
	, , , , , , , , , , , ,		
~ O.	UI BS	6.0	60.0
property of 18	3.10.1.6.3.3	Add indicated amo mixed control solut	unt of 1ng/μL working ion.
		Desired ng/mL	μL Working

Desired ng/mL	μL Working Control	
60.0	60.0	

3.10,1.6.4 Negative Control Sample Preparation

Add 1mL of negative whole blood into an extraction tube.

3.10.1.6.5 Case Sample Preparation

Add 1mL of blood into a labeled extraction tube.

3	3.10.1.6.6	Internal Standard 3.10.1.6.6.1	Addition To calibrators, controls and case samples, add 25μL of internal standard mix.
		3.10.1.6.6.2	Cap tube and vortex tube briefly.
		3.10.1.6.6.3	Allow tubes to stand 30 minutes for sample equilibration.
3	3.10.1.6.7	Protein Precipitati 3.10.1.6.7.1	While vortexing, with a glass pipette, add 2mL 10% methanol in acetonitrile dropwise to case, calibrator and control samples.
		3.10.1.6.7.2	Cap tube and continue vortexing tube for approximately 30 seconds.
		3,10,1,6,7.3	Centrifuge at 3400 rpm for 10 minutes.
opertyon		3.10.1.6.7.4	Decant organic supernatant into second labeled glass tube.
	. 83	3.10.1.67.5	Transfer tube to Evaporative Concentrator and evaporate under nitrogen at ≤40°C to approximately 1mL. Do not allow extract to go to dryness.
	ety of lo	\$.10.1.63.6	To the evaporated extract add 2mL 100mM acetate buffer (pH 4.5). Vortex briefly to mix.
0%		3.10.1.6.7.7	If necessary, centrifuge buffered solution for an additional 5 minutes at 3400 rpm to remove blood fragments or foam.
	3.10.1.6.8	<u>SPE Column Prep</u> 3.10.1.6.8.1	naration Insert labeled 200mg CLEAN SCREEN® THC extraction column into appropriate location on vacuum manifold.
		3.10.1.6.8.2	To each SPE column, add $3mL$ 70:30 Hexane:Ethyl acetate. Aspirate at ≤ 3 in. Hg to prevent sorbent drying.

Idaho State Police	Forensic Services	Toxicology Discipline Analytical Method
	3.10.1.6.8.3	To each SPE column, add 3mL methanol to the column. Aspirate at ≤ 3 in. Hg.
	3.10.1.6.8.4	To each SPE column, add 3mL deionized water to the column. Aspirate at \leq 3 in. Hg.
	3.10.1.6.8.5	To each SPE column, add 1mL 100mM HCl and aspirate at ≤ 3 in. Hg.
3.10.1.6.9		ading blood extract onto the SPE column and low or apply minimal vacuum.
3.10.1.6.10	Column Wash	50
5.10.1,0,10	3.10.1.6.10.1	To each SPE column, add $2mL$ of deionized water. Aspirate at ≤ 3 in. Hg.
	3.10,1.6,10.2	To each SPE column, add 2mL 70:30 100mM HCl. Acetonitrile. Aspirate at ≤ 3 in Hg.
3.10.1.6.11	Dry Disc	
		to \geq 10 in Hg (\geq 34 kPa) for \geq 5 minutes.
3.10.1.6.12	Compound Riutio	Open vacuum manifold, wipe collection
3/8	SI COURT	tips, and insert the collection rack containing the labeled glass tubes.
Property of I	3.10.1.C.12.2	To each SPE column, add 200uL hexane (important for elution solvent reception). Gravity flow only. Do not allow column to dry.
	3.10.1.6.12.3	To each SPE column, add 3mL 70:30 Hexane: Ethyl Acetate elution solvent to the column. Collect eluate with gravity

3.10.1.6.13 Eluate Evaporation

Transfer centrifuge tube to Evaporative Concentrator. Evaporate eluates to dryness, under a gentle stream of nitrogen at $\leq 40^{\circ}\text{C}$.

flow or apply minimal vacuum.

	3.10.1.6.14	<u>Derivatization</u> 3.10.1.6.14.1	In fume hood, add 40µL each ethyl acetate and BSTFA (1% TMCS) to each extract.
		3.10.1.6.14.2	Cap tubes and vortex briefly.
		3.10.1.6.14.3	Place tubes in dry bath or oven set at 70°C for 15 minutes.
		3.10,1.6.14,4	Remove tubes from oven and allow to cool to room temperature.
		3.10.1.6.14.5	Transfer derivative to labeled GC/MSD ALS vial with microinsert.
	3.10.1.6.15	Preparation for G	C-MS Run
		3.10.1.6.15.1	Perform an AUTOTUNE and TUNE EVALUATION.
		3.10.1.6.15.2	When tune values are acceptable, program SEQUENCE TABLE with sample, calibrator and control information.
		3.10.1.6.188	Load ALS vials into quadrant racks as indicated in the SEQUENCE TABLE.
	3.10.1.6.16	GC-MS Calibration	on Curve
	193	3.10.1.6.16.1	The calibration curve must be established with a minimum of five data points.
~(eky oi '	3.10 7 5.16.2	All reported results must be bracketed by calibrators.
X		3.10.1.6.16.3	Calibrators should be analyzed in order of increasing concentration.
		3.10.1.6.16.4	The least squares line resulting from the analysis of the calibrators must have a coefficient of correlation of ≥ 0.98 .
		3.10.1.6.16.5	If calibrators are run in duplicate, it is not required that duplicate calibration points are included as long as the linearity requirement is met.

3.10.1.7 GC and MSD ACQUISITION PARAMETERS

Critical parameters are specified below. Parameters not specified are at the discretion of the analyst and should be optimized for the particular GC-MSD instrument. Each laboratory should maintain a centrally stored printed or electronic copy of current and past GC-MSD methods. The data supporting the GC-MSD method should be stored centrally.

3.10.1.7.1 GC Temperature Parameter

Injection Port: 250°C or 260°C

3.10.1.7.2 MSD Instrument Parameters

Detector/Transfer Line: 280°C

3.10.1.7.3 ALS Parameters

Injection Volume: 1µL (1 stop)

Viscosity Delay: A minimum of 1 second

Solvent Washes (A & B): A minimum of 3 pre- and post-

wash rinses.

3.10.1.7.4 MS SIM Parameters

Target	Qualifier Ion	Qualifier Ion
Ion	1	2
386	371	315
374	306	389
371	473	488
380	482	497
	386 374 371	386 371 374 306 371 473

3.10.1.8 REPORTING CRITERIA

3.10.1.8.1 Qualitative Chromatographic Criteria

Acceptable retention time window established by calibrators is ± 0.2 minute.

3.10.1.8.2 Qualitative Mass Spectral SIM Criteria

Ion ratios for the analyte and its corresponding internal standard, established by calibrators for target and qualifier ions, must not differ by more than ±20%.

3.10.1.8.3 Quantitative Mass Spectral and Control Criteria

3.10.1.8.3.1 Quantitative results can be accepted if the calculated concentration of all calibrators

and control samples are within $\pm 20\%$ of their respective concentrations and the coefficient of variation (CV%) for replicates of control samples is $\leq 15\%$.

3.10.1.8.3.2

Quantitation is achieved through the plotting of the target ion response ratio versus the concentration for each calibrator.

3,10,1,8,3,3

Quantitative values for case samples, calibrators and controls will be truncated for reporting purposes.

3.10.1.8.3.4

Cut-off for Δ^9 -THC and Carboxy-THC is the lowest calibrator 2.5ng/mL, or the lowest calibrator that meets quality assurance requirements.

3.10.1.8.3.5

If the concentration exceeds the calibration range, the sample can either be appropriately diluted with negative whole blood for reanalysis or reported as greater than 190ng/mL.

3.10.1.9 REPORTING OF RESULTS

3.10.1.9.1

Ouantitative Value

Analysis results should be truncated and reported out without decimal places.

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Uncertainty Value

Based on the current uncertainty assessment, the +/- range should be included on the analysis report. Refer to method variation spreadsheet for current uncertainty figure.

3.10.1.10 QUALITY ASSURANCE REQUIREMENTS

3.10.1.10.1 General

3.10.1.10.1.1

Blood samples are to be stored under refrigeration after aliquots are removed for analysis.

3.10.1.10.1.2

Refer to toxicology manual section 5.1 for pipette calibration and intermediate check options.

	3.10.1.10.1.3	Refer to toxicology manual section 5.2 for balance calibration and intermediate check requirements.
	3.10.1.10.1.4	Refer to toxicology manual section 5.8 for additional GC-MSD quality assurance requirements.
	3.10.1.10.1.5	Refer to toxicology manual section 5.10 for reference material authentication requirements.
3.10.1.10.2	Per Analysis Rur	Control Requirements
	3.10.1.10.2.1	Solvent blank should follow the highest
		calibrator as well as each case sample.
	3.10.1.10.2.2	A minimum of two blood controls must be
		run per batch of samples. A control must
		be run for each additional 10 case samples.
2 10 1 10 2	Mantenton CO	CIVE OF SECTION
3.10.1.10.3	Monitoring of Co	1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1
		etion of analysis, input blood control values
	on spreadsneet u	sed to assess uncertainty for this method.
	~x\Q, \	

3.10.1.11 ANALYSIS DOCUMENTATION

3.10.1.11.1 A packet containing original data for controls and standards will be prepared for each analysis run and stored centrally in the laboratory where the analysis was performed until archiving.

A copy of controls and calibrators need not be included in individual case files. When necessary, a copy of the control and calibrator printouts can be prepared from the centrally stored document.

3.10.1.12 REFERENCES AND RECOMMENDED READING

- 3.10.1.12.1 UCT CLEAN SCREEN® Extraction Columns Application Manual.
- 3.10.1.12.2 Standard Operating Procedure for Blood SPE Cannabinoids, Montana Department of Justice Forensic Sciences Division.

- 3.10.1.12.3 Standard Operating Procedure for Blood SPE THC and Carboxy-THC GC/MSD Assay, Edmonton, Canada Office of the Chief Medical Examiners, 2003.
- 3.10.1.12.4 Huestis, M.A., Cannabis (Marijuana) Effects on Human Behavior and Performance, Forensic Science Rev. 14(1/2): 16-60, 2002.
- 3.10.1.12.5 Drummer, O.H., *Cannabis*, pp. 178-212. *in:* The Forensic Pharmacology of Drugs of Abuse, Arnold: London, 2001.
- Bock, Peter, Getting it right R&D method For science and Property of Idaho State Police Forensic Soll Internet Int 3.10.1.12.6 engineering, Academic Press, San Diego 2001.

Revision History

Section Three Blood Toxicology

- 3.10 Manual Solid Phase Extraction (SPE) Methods
 - 3.10.1 Extraction and Quantitation of THC and Carboxy-THC from Blood Employing the United Chemical Technologies (UCT) 200 mg CLEAN SCREEN® THC Extraction Column

		3 7
Revision No.	Issue Date	History/Comments
0	11-22-2006	Original Issue Method is approved for qualitative purposes only.
	:	Upon review of uncertainty determinations for
		quantitative analysis this method will be applied for intended use
1	07-28-2008	Clarified that negative blood used to prepare calibrators and positive controls is the same lot as
	5	used for negative control.
used for negative control.		
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