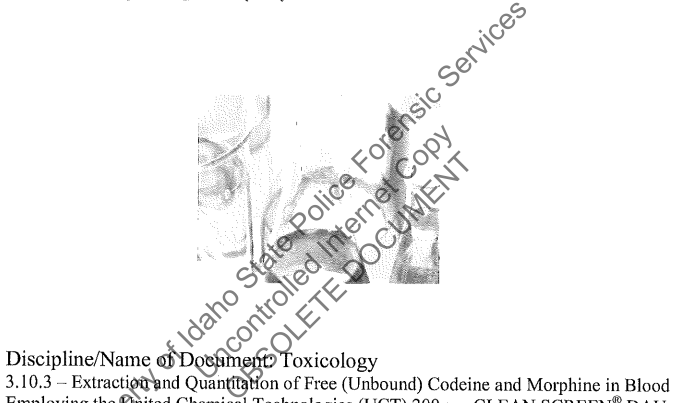
Idaho State Police Forensic Services

Approval for Quality System Controlled Documents



Employing the United Chemical Technologies (UCT) 200 mg CLEAN SCREEN® DAU Extraction Column (FOR QUALITATIVE USE ONLY)

Revision Number: 1

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Quality Manager

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Section Three Blood Toxicology

3.10 Manual Solid Phase Extraction (SPE) Methods

3.10.3 Extraction and Quantitation of Free (Unbound) Codeine and Morphine in Blood Employing the United Chemical Technologies (UCT) 200 mg CLEAN SCREEN® DAU Extraction Column

3.10.3.1 BACKGROUND

Refer to qualitative opiate analytical method 3.4.4 and provided references for information regarding the background and pharmacology of these compounds.²⁻⁵

3.10.3.2 PRINCIPLE

This procedure is based on a method developed by United Chemical Technology (UCT) which applies the UCT 200 mg CLEAN SCREEN® extraction column for the extraction of opiates from blood.¹ The CLEAN SCREEN® DAU 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 of opiate class drugs, the blood sample is diluted and adjusted to pH 6 with a phosphate buffer. After optional centrifugation, the sample is loaded onto a pre-conditioned SPE column. The blood pH is adjusted to maximize the ionic character of the analyte. The conditioning creates an environment, which allows for optimal interaction between the sorbent and the analytes of interest. The analyte is retained by ionic interaction of the amine functional groups present on the drug and the anionic sulfonic acid exchanger on the sorbent. The column is subsequently washed with water, a weak aqueous buffer and methanol to selectively remove matrix components and interfering substances from the column. The wash also disrupts the hydrophobic and adsorption interactions but not the ionically bound material. Next, the column is dried to remove traces of aqueous and organic solvents. When the column is dry the analytes of interest are eluted from the column with a basic organic solvent mixture. Following elution from the SPE column and evaporation, the extract is derivatized for confirmation on the GC/MSD. Quantitation is accomplished with a 5 to 6 point calibration curve using the corresponding deuterated standard to establish the response factor.

3.10.3.3	EQUIPMEN	IT AND SUPPLIES
	3.10.3.3.1	200mg CLEAN SCREEN® Extraction Column (ZSDAU020
		or ZCDAU020 or equivalent)
	3.10.3.3.2	Laboratory oven
	3.10.3.3.3	Evaporative concentrator equipped with nitrogen tank.
	3.10.3.3.4	Tube Rocker
	3.10.3.3.5	Vortex Mixer
	3.10.3.3.6	Laboratory centrifuge capable of 3400rpm
	3.10.3.3.7	Vacuum Manifold/pump
	3.10.3.3.8	Fixed and adjustable volume single channel air displacement
		pipetters, and appropriate tips, capable of accurate and
		precise dispensing of volumes indicated. pH indicator strips 16 x 100mm silanized glass tubes
	3.10.3.3.9	pH indicator strips
	3.10.3.3.10	16 x 100mm silanized glass tubes
	3.10.3.3.11	Screw Cap for 16mm O.D. tube
	3.10.3.3.12	GC/MS Automated Liquid Sample (ALS) vials
	3.10.3.3.13	Silanized GC/MS Vial Microinsert
	3.10.3.3.14	Gas Chromatograph (GC) equipped with a mass selective
		detector (MSD) and a nonpolar capillary column with a phase
		composition comparable to 100%-dimethylpolysiloxane or
		95%-dimethyl-polysiloxane with 5%-diphenyl.
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3.10.3.4 REAGENTS

J	Refer	· to	anal	vtical	met	hod	5.12	for solution	preparatio	n instructions.

	3.10.3.4.1	Deionized/distilled (DI) water
	3.10.3.4.2	Methanol (Certified ACS Grade)
	3.10.3.4.3	Methylene Chloride (Certified ACS Grade)
	3.10.3.4.4	Ethyl Acetate (Certified ACS Grade)
	3.10.3.4.5	Isopropanol (Certified ACS Grade)
	3.10.3,4.6	Ammonium Hydroxide (Certified ACS Grade)
	3.10.3.4.7	100mM Phosphate Buffer (pH 6.0)
7	3.10.3.4.8	100mM Acetate Buffer (pH 4.5)
	3.10.3.4.9	100mM Monobasic sodium phosphate
	3.10.3.4.10	100mM Dibasic sodium phosphate
	3.10.3.4.11	Elution Solvent
		Mix 20mL Isopropanol and 2mL Ammonium Hydroxide QS
		to 100mL with methylene chloride. pH should be 11-12.
		Make fresh.
	3.10.3.4.12	BSTFA + 1% TMCS

3.10.3.5 QUALITY ASSURANCE MATERIAL

3.10.3.5.1	Calibrator and Control Solutions		
	3 10 3 5 1 1	Stock Solutions	

The source of a corresponding calibrator and

control must be obtained from a different vendor.

Codeine and Morphine Concentration: 1mg/mL

3.10.3.5.1.2 Working Drug Solutions

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

<u>10ng/μL</u>

Add 100µL each Codeine and Morphine Stock Solution to ≥9mL Methanol in a 10mL volumetric class A flask. QS to 10mL.

Ing/µL

Add 1mL 10ng/µL working drug solution to ≅5mL Methanol in a 10mL volumetric class A flask: QS to 10mL.

3.10.3.5.2 <u>Internal Standard Solutions</u>

3.10.3.5.2.1

Stock Solutions

Cødeme-D₃ and Morphine-D₃
Concentration: 1mg/mL or 100μL/mL

3.10.35.2.2

Working Internal Standard Solution $\underline{1ng/\mu L}$

Add $10\mu L$ each 1mg/mL or $100\mu L$ each $100\mu L/mL$ Codeine-D3 and Morphine-D3 Stock Solution to $\cong 9mL$ Methanol in a 10mL volumetric class A flask. QS to 10mL. Store remaining stock solution in ALS vial in freezer. Working solution is stable for 6 months when stored at $4^{\circ}C$.

3.10.3.5.3 Commercial Whole Blood Controls

3.10.3.5.3.1 Negative Whole Blood

3.10.3.5.3.2 **Positive Whole Blood**

Control containing Codeine and Morphine each at a specified target concentration. Refer to package insert for verified value and expected range.

3.10.3.6 **PROCEDURE**

3.10,3,6,1 Initial set-up

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

3.10.3.6.2 Calibrator Preparation

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

Add 1mL of negative whole blood into six 3.10.3.6.2.1 screw-top extraction tubes.

Add the volume of 3.10.3.6.2.2 working Codeine-Morphine Ing/µL mixed calibrator solution to appropriate tube as indicated in the chart

	to appropulation to the total to the total total total total to the total tota	oriate tube	e as indicated in the chart
	Level	ng/ml	μL Working Reference Material
	1,0	25	25
	X2	3 0	50
	30	100	100
Property of Idahonting	Morphine to the ap	e 10ng/μL opropriate	of working Codeine and mixed calibrator solution tube as indicated in the
eky O.OB3	Level	ng/mL	μL Working Reference Material
1000	4	250	25
Ric	5	500	50
	6	1000	100

Level	ng/mL	μL Working Reference Material
4	250	25
5	500	50
6	1000	100

3.10.3.6.2.4 Additional or alternative concentrations may be used as necessary as long as the requirements in 3.10.3.6.15.1 are met.

3.10.3.6.3 Positive Control Sample Preparation

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

3.10.3.6.3.1 Add 1mL of negative whole blood to two screw-top tubes.

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Issued: 07-28-2008 BLOOD 3,10.3- Opiate SPE -Rev 1,doc Issuing Authority: Quality Manager 3.10.3.6.3.2 Add indicated amount of lng/µL working mixed control solution.

ng/mL	μL Working Control	
75	75	

3.10.3.6.3.3 Add indicated amount of 10ng/µL working mixed control solution.

ng/mL	L Working Control
750	75

3.10.3.6.3.4 Additional of alternative concentrations at the discretion of the analyst may be used as long as the requirements in 3.10.3.10.2 are

met

3.10.3.6.4 Negative Control Sample Preparation
Add 1mL of negative whole blood to a screw top tube.

3.10.3.6.5 <u>Case Sample Preparation</u>

B.10.3.6.5.1 Based on enzyme immunoassay screen results, samples may be diluted with negative whole blood prior to analysis.

3.10.3.6.5.2

The total volume of blood or diluted blood should be 1mL.

3.10.3.6.5.3 Add 1mL neat or diluted sample into a labeled 13x100mm screw top tube.

3.10.3.6.6 <u>Internal Standard Addition</u>

3.10.3.6.6.1 To prepare 100ng/mL internal standard add 10μL of 1mg/mL (1000ng/μL) or 100μL of (100ng/μL) 1ng/μL of internal standard mix to calibrators, controls and casework samples.

- 3.10.3.6.6.2 Cap tube and vortex tube briefly.
- 3.10.3.6.6.3 Allow tubes to stand 30 minutes for sample equilibration.

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3.10.3.6.7

Sample Preparation

3110/3/01/	3.10.3.6.7.1	Add 4mL DI water, vortex, let stand for 5 minutes.
	3.10.3.6.7.2	Centrifuge for 10 minutes @ 3400rpm.
	3.10.3.6.7.3	Transfer supernatant to second tube.
	3.10.3.6.7.4	Add 4mL 100mM phosphate buffer (pH 6.0), vortex.
	3.10.3.6.7.5	Sample pH should be 6.0 ±0.5. Adjust as necessary with 100mM Monobasic sodium phosphate or 100mM Dibasic sodium phosphate.
3.10.3.6.8		eparation necessary except for drying step, however, if e at ≤ 3 in. Hg to prevent sorbent drying.
	3.10.3.6.8.1	Insert labeled 200mg CLEAN SCREEN® extraction column onto the vacuum manifold.
	3.10.3.6.8.2 3.10.3.6.8.3	Add 3mL methanol to the column. Add 3mL DI water to the column.
10 to	3.10.36.8.4	Add 1mL 100mM Phosphate buffer (pH 6.00) to the column.
\$.10.3.6.9	Blood Extract I Load buffered or apply minim	blood onto column and allow to gravity flow
3.10.3.6.10	Column Clean- 3.10.3.6.10.1	up Add 2mL DI water to the column.
	3.10.3.6.10.2	Add 2mL 100mM Acetate buffer (pH 4.5) to the column. Aspirate.
	3.10.3.6.10.3	Add 3mL Methanol.
	3.10.3.6.10.4	Increase vacuum to ≥ 10 in. Hg (≥ 34 kPa) for ≥ 5 minutes (disc should be dry).

3.10.3,6.11	Compound Elur 3.10.3.6.11.1	tion Open vacuum manifold, wipe collection tips, and insert the collection rack containing the labeled tapered tip centrifuge tubes.
	3.10.3.6.11.2	Add 3mL elution solvent (3.10.3.4.12) to the column. Collect eluate with gravity flow or apply minimal vacuum.
3,10,3,6,12	Evaporate solv	rifuge tube to Evaporative Concentrator, vent to dryness under a sentle stream of roximately 37°C.
3.10.3.6.13	Derivatization	;C

3.10.3.6.13.1 In fume hood add 50µL ethyl acetate. Vortex for ≥15 seconds.

3.10.3.6.13.2 Add 50.0µLBSTFA + 1% TMCS.

3.10.3.6.13.3 Cap tubes and vortex briefly.

3.10.3.6.134 Place tubes in 70°C oven for 20 minutes.

3.10.3.6.13.5 Remove from heat and allow to cool to room temperature.

3.10.36.13.6 Transfer derivative to labeled GC/MSD ALS vial with microinsert.

3,10,3,6,14 Preparation for GC-MS Run

3.10.3.6.14.1 Perform an AUTOTUNE and TUNE EVALUATION. Evaluate applying acceptance criteria outlines in analytical method 5.3.1.

3.10.3.6.14.2 When tune values are acceptable, program SEQUENCE TABLE with sample, calibrator and control information.

3.10.3.6.14.3 Load ALS vials into quadrant racks as indicated in the SEQUENCE TABLE.

3.10.3.6.15 GC-MS Calibration Curve

3.10.3.6.15.1 The calibration curve should be established with a minimum of four data points.

- 3.10.3.6.15.2 All reported results must be bracketed by calibrators.
- 3.10.3.6.15.3 Calibrators should be analyzed in order of increasing concentration.
- 3.10.3.6.15.4 The least squares line resulting from the analysis of calibrators must have a coefficient of correlation of ≥0.98.
- 3.10.3.6.15.5 If calibrators are run in duplicate, it is not required that duplicate calibration points are included as long as the linearity requirement is met.

3.10.3.7 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.

- 3.10.3.7.1 GC Temperature Parameter
 Injection Port: 250° or 260°C
- 3.10.3.7.2 <u>MSD Instrument Parameters</u> Detector/Transfer Line: 280°C
- 3.10.3.7.3 ALS Parameters

Injection Volume: 1µL (1 stop)

Viscosity Delay: A minimum of 3 seconds

Solvent Washes (A & B): A minimum of 4 pre- and post-wash rinses.

3.10.3.7.4 MS SIM Parameters

Analyte	Target	Qualifier	Qualifier
	Ion	Ion 1	Ion 2
Morphine	429	287	324
Morphine-D3	432	290	327
Codeine	371	234	343
Codeine-D3	374	237	346

3.10.3.8 REPORTING CRITERIA

3.10.3.8.1 Qualitative Chromatographic and SIM Criteria

> 3.10.3.8.1.1 Qualitative results can be accepted when the following two criteria are met.

- 1. The retention time falls within the ± 0.2 minute window established by calibrators.
- 2. Ion ratios for the analyte and its corresponding internal standard, established by calibrators for target and qualifier ions, do not differ by more than ±20%.

3.10.3.8.2 Quantitative Mass Spectral Criteria

3.10.3.8,2,1 Quantitative results can be accepted if the calculated concentration of all calibrator and control samples are within ±20% of their

3.10.3.8.2.4 Administrative limit of detection of the target on response ratio versus the concentration for each calibrator.

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

Administrative limit of detection of the concentration for each calibrator.

3.10.3.8.2.5 If the concentration exceeds the calibration range, the sample must be appropriately diluted with negative whole blood for reanalysis.

3.10.3.9 REPORTING OF RESULTS

3.10.3.9.1 **Quantitative Value**

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

3.10.3.9.2 Uncertainty Value

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

3.10.3.10 QUALITY ASSURANCE REQUIREMENTS

3.10.3.10.1 General

- 3.10.3.10.1.1 Blood samples are to be stored under refrigeration after aliquots are removed for analysis.
- Refer to toxicology manual section 5.1 for 3.10.3.10.1.2 pipette calibration and intermediate check options.
- Refer to toxicology manual section 5.2 for 3.10.3.10.1.3 balance calibration and intermediate check requirements.
- 3.10.3.10.1.4 Refer to toxicology manual section 5.8 for additional GC-MSD quality assurance requirements.
- Refer to toxicology manual section 5.10 for reference material authentication requirements.

Run Quality Requirements

- Solvent blank should follow the highest calibrator as well as each case sample.
- 3.10.3.10.20 P 3.10.3.10.2.2 A minimum of two commercially obtained blood controls and the spiked controls described in section 3.10.3.6.3 must be run per batch of samples.
 - 3.10.3.10.2.3 In addition to the four blood controls indicated above, for each additional 10 case samples, one control must be run. preparation of controls is outlined in section 3.10.3.6.3. Additional concentrations may be used.

3.10.3.10.3 Monitoring of Control Values

Upon the completion of analysis, input blood control values on spreadsheet used to assess uncertainty for this method.

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3.10.3.11 ANALYSIS DOCUMENTATION

- 3.10.3.11.1 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.
- 3.10.3.11.2 A copy of controls and standards need not be included in individual case files. When necessary, a copy of the control and standard printouts can be prepared from the centrally stored document.

3.10.3.12 REFERENCES AND RECOMMENDED READING

- 3.10.3.12.1 Telepchak, M.J., August, T.F. and Chaney, G., Drug Methods for the Toxicology Lab, pp. 227 230. *in:* Forensic and Clinical Applications of Solid Phase Extraction, Humana Press: New Jersey, 2004.
- 3.10.3.12.2 Stout, P.R. and Farrell, L.J. Opioids Effects on Human Performance and Beliavior, Forensic Science Rev. 15(1): 29 60, 2003.
- 3.10.3.12.3 Drummer, O.H., *Opioids* pp. 219 265. *in:* The Forensic Pharmacology of Drugs of Abuse, Arnold: London, 2001.
- 3.10.3.12.4 Kerrigan, S and Goldberger, B.A., *Opioids.* pp. 187 206. in: Principles of Forensic Toxicology. Levine, B. ed., AACC. 2003.
- 3.10.3.12.5 Baselt, R.C., Codeine, pp. 262 265. and Morphine, pp. 759 763. in: Disposition of Toxic Drugs and Chemicals in Man, Biomedical Publications: Foster City, CA. Seventh ed., 2004.

Revision History

Section Three Blood Toxicology

3.10 Manual Solid Phase Extraction (SPE) Methods

3.10.3 Extraction and Quantitation of Free (Unbound) Codeine and Morphine in Blood Employing the United Chemical Technologies (UCT) 200 mg CLEAN SCREEN® DAU Extraction Column

Revision No.	Issue Date	Revision/Comments Original Issue
0	11-21-2006	Original Issue
	11 21 2000	Original Issue
1	07-28-2008	Clarified that negative blood used to prepare
		calibrators and positive controls is the same lot as used for negative control.
Proper	y of ldahoon	Clarified that negative blood used to prepare calibrators and positive controls is the same lot as used for negative control.