# Idaho State Police Forensic Services

Approval for Quality System Controlled Documents



Discipline/Name of Document: Toxicology 3.9.2 - Quantitation of Higher pKa Drug Compounds (FOR QUALITATIVE USE ONLY)

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Original Certificate did not document that the approval was only for reporting qualitative results.

# Idaho State Police Forensic Services

Approval for Quality System Controlled Documents



## Idaho State Police **Forensic Services** Toxicology Discipline

## **Section Three**

## **Blood Toxicology**

Liquid-Liquid Extraction Methods for Quantitative Gas Chromatography 3.9 3.9.2 Quantitation of Higher pKa Drug Compounds.

#### BACKGROUND 3.9.2.1

This method is a quantitative liquid-liquid extraction procedure for a variety of commonly encountered basic drugs that exhibit a pKa of ≥9 along with This method prepares an extract for confirmatory their metabolites. quantitative analysis with a gas chromatograph (GC) equipped with a mass selective (MSD) or nitrogen phosphorus detector (NPD).

#### PRINCIPLE 3.9.2.2

The method is based upon the principle of liquid/liquid extraction. The sample pH is adjusted with a pH 12 saturated borate buffer and extracted with n-butyl chloride. Following mixing and centrifugation the supernatant is transferred and 1%HCI in MeOH is added to prevent to loss of volatile analytes during solvent evaporation steps. This method may be performed with or without a back extraction for sample clean-up. If the backextraction is used, it must be applied to all calibrators, controls and samples. Quantitation is accomplished with a 5 to 6 point calibration curve using the appropriate internal standard to establish a response factor.

#### OUIPMENT AND SUPPLIES 3.9.2.3

2 3.9.2.3.1	pipetters, and appropriate tips, capable of accurate precise dispensing of volumes indicated.
3.9.2.3.2	Tube rocker
3.9.2.3.3	Evaporative concentrator equipped with nitrogen tank.
3.9.2.3.4	Vortex mixer
3.9.2.3.5	Laboratory centrifuge capable of ≥3200rpm
3,9,2,3.6	16 x 100mm round bottom glass screw-top tubes
3.9.2.3.7	Screw Cap for 16mm O.D. tubes

0,7,2,0,1	Boton Cup for Tommi of the contract of the con
3.9.2.3.8	GC/MS Automated Liquid Sampler (ALS) vials
3.9.2.3.9	GC/MS vial microinsert
3.9.2.3.10	GC equipped with Dual NPDs
3.9.2.3.11	GC equipped with a MSD
3.9.2.3.12	Non-polar Capillary Column (GC-NPD and GC-MSD)

and

100%-Dimethylsiloxane or a 5%-Diphenyl-95%-Dimethyl-

siloxane copolymer, 12.5 to 30M.

Mid-Polar Capillary Column (GC-NPD)

50% Phenyl, 50% methyl-polysiloxane copolymer, 12.5 to

		30M.	
3.9.2.4	3.9.2.4.1 3.9.2.4.2 3.9.2.4.3 3.9.2.4.4	mual section 5.1 Methanol (CepH 12 Satural n-Butyl chlor 1% Hydrochl	2 for solution preparation instructions. ertified ACS grade) ted Borate Buffer ide (Certified ACS grade) oric Acid in Methanol
3.9.2.5	<b>QUALITY</b> 3.9.2.5.1	ASSURANCE  Drug Stock S  1mg/mL (1µ  control prepa	MATERIAL solutions g/ $\mu$ L) drugs standards used for calibrator and ration must be obtained from different vendors.
	3.9.2.5.2		B Solutions 10ng/μL Add 100.0μL each 1mg/mL Stock Solution drug of interest to ≅9mL Methanol in a 10mL volumetric class A flask. QS to 10mL. Working drug solutions may be mixed or single compound depending on the compound's retention time.
8,	opert.	3.9.2.5.2.2	1ng/μL Add 1.0mL 10ng/μL working drug solution to ≅8mL Methanol in a 10mL volumetric class A flask. QS to 10mL.
		3.9.2.5.2.3	Working solutions are stable for 6 months when stored at 4°C.
		3.9.2.5.2.4	Store remaining stock solution in ALS vial in freezer.
	3.9.2.5.3	Select appro	dard Stock Solutions priate internal standard for drug of interest. For alysis, use deuterated standard when available.

3.9.2.3.13

1ng/µL Working Internal Standard Solution 3.9.2.5.4

Add 10µL each 1mg/mL or 100.0µL each 100µL/mL Stock Solution to to ≅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.

3.9.2.5.5 Negative Control Negative Whole Blood

#### **PROCEDURE** 3.9.2.6

Initial set-up 3.9.2.6.1

For each calibrator, control and case sample, label two sets of extraction tubes and an ALS vial with microinserts,

Calibration Standard Preparation 3.9.2.6.2

Add 2mL of negative whole blood to six 3.9.2.6.2.1 screw top extraction tubes.

Add the calibration below.

Level

3.9.262 Add the volume of working lng/µL working calibration standard as indicated in the chart

Level	Desired ng/mL	μL Working Standard
1	25	50
2	50	100

Add the volume of working 10ng/µL working calibration standard as indicated in the chart

Level	Desired ng/mL	μL Working Standard
3	100	20
4	250	50
5	500	100
6	1000	200

#### Positive Control Sample Preparation 3.9.2.6.3

Add indicated amount of working 10ng/µL 3,9,2,6,3,1 mixed control solution to 2mL negative whole blood.

Desired ng/ml	L μ <b>L Working</b>

	Control
75	15
750	150

3.9.2.6.3.2 Positive controls must be run in duplicate. For every 10 case samples prepare an additional control.

3.9.2.6.4	Negative Cont	rol Sample Preparation egative whole blood to screw top tube.
3.9.2.6.5	Case Sample F	
3.9.2.6.6	<u>Internal Stands</u> 3.9.2.6.6.1	ard Addition To calibrators, controls and case samples, add 200μL of internal standard mix. Vortex.
	3.9.2.6.6.2	Allow tubes to stand 15 minutes.
3.9.2.6.7	Extraction 3.9.2.6.7.1	Add 2mL pH 12 borate buffer. Vortex.
	3.9.2.6.7.2	Add 4mL n-butyl chloride into each tube, cap.
	3.9.2.6.7.3	Place tube on rocker for 10 minutes.
,10	3.9.2.6.7.4	Centrifuge for 10 minutes at 3200 - 3400 rpm.
etty o'	3.9.276, 7.5	Transfer the upper n-butyl chloride layer to second tube.
96	3.9.2.6.7.6	Add 50 $\mu$ L 1% HCl in methanol.
	3.9.2.6.7.7	Evaporate to dryness under a gentle stream of nitrogen at approximately 37°C.
3.9.2.6.8	Back-extraction 3.9.2.6.9.1	on Reconstitute with 50ul 100mM HCl.
	3.9.2.6.9.2	Add 1ml of n-Butyl Chloride. Vortex.

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3.9.2.6.9.3

3.9.2.6.9.4 Centrifuge for 5 minutes at 3200-3400 rpm.

Rock for 5 minutes.

	3.9.2.6.9.5	Discard upper n-Butyl Chloride layer.
	3.9.2.6.9.6	Add 2ml of pH 12 borate solution. Vortex
	3.9.2.6.9.7	Add 4 ml of n-Butyl Chloride.
	3.9.2.6.9.8	Rock for 5 minutes.
	3.9.2.6.9.9	Centrifuge for 5 minutes at 3200 - 3400 rpm.
	3.9.2.6.9.10	Transfer upper n-Butyl Chloride layer into screw-top tube.
	3.9.2.6.9.11	Evaporate under a gentle stream of nitrogen at approximately 37°C
3.9.2.6.10	Reconstitution 3.9.2.6.10.1	Add 50nL Methanol to the residue, vortex.
	3.9.2.6.10.2	Transfer extract to labeled ALS vial with microinsert.
3.9.2.6.11	Preparation for 3.9.2.6 11.1	or Analysis Run Unto Sequence log table, enter the sample case numbers, blanks and controls.
01/0	3,9,2 6,11.2	Load samples, standards, blank and controls into the quadrant rack as noted in the sequence table.
39.2.6.12	Analysis Para 3.9.2.6.12.1	meters  Refer to instrument METHOD printouts for analysis parameters.
	3.9.2.6.12.2	For GC-MSD appropriate SIM ions for quantitative analysis are selected from full scan analysis of standard. Selected ion monitoring at the corresponding retention time is configured accordingly.
	3.9.2.6.12.3	Current analysis method must be stored centrally as a hard or electronic copy.
3.9.2.6.13	GC-MS Calil 3.9.2.6.13.1	The calibration curve should be established with a minimum of four data points.

	3.9.2.6.13.2	All reported results must be bracketed by calibrators.
	3.9.2.6.13.3	Calibrators should be analyzed in order of increasing concentration.
	3.9.2.6.13.4	The least squares line resulting from the analysis of calibrators must have a coefficient of correlation of ≥0.98.
	3.9.2.6.13.5	If calibration standards are run in duplicate, it is not required that duplicate calibration points are included as long as the linearity requirement is met.
DEDODTING	G CRITERIA	X () ()
	J CKII EKIA	Control Control
3.9.2.7.1	Qualitative C	hromatographic Criteria
		sulls can be accepted when the following two
	criteria are me	
	1. The rete	ention time falls within the ±0.2 minute window
		ed by calibrators.
	For may	ss spectral data, SIM ion ratios for the analyte
	2, Put max	corresponding internal standard, established by
, 10	adil ust	for target and qualifier ions do not differ by
À,	Cambrac	ors for target and qualifier ions, do not differ by
, 0,	more th	an ±20%.
(23)	O	
39.2.7.2	Quantitative (	
<b>X</b>	3.9.2.7.2.1	Quantitative results can be accepted if the calculated concentration of all calibration standards and control samples are within ±20% of their respective concentrations.
	3.9.2.7.2.2	Quantitation is achieved through the plotting of the compound's response ratio versus the concentration for each calibrator.
	3.9.2.7.2.3	Quantitative values for case samples, calibrators and controls will be truncated for reporting purposes.

3.9.2.7

3,9.2,7.2,4

Limit of quantitation is the lowest calibrator.

3.9.2.7.2.5 If the concentration exceeds the calibration range, the sample can either be appropriately diluted with DI water for reanalysis or reported as greater than 1000ng/mL.

## 3.9.2.8 REPORTING OF RESULTS

3.9.2.8.1 Quantitative Value

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

3.9.2.8.2 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.

## 3.9.2.9 QUALITY ASSURANCE REQUIREMENTS

3.9.2.9.1 <u>General</u>

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

Refer to toxicology analytical method 5.1 for pipette calibration options.

Refer to toxicology analytical method 5.2 for balance calibration requirements.

Refer to toxicology analytical method 5.3.1 for GC-MSD maintenance guidelines.

3.9.2.1.5 Refer to toxicology analytical methods 5.8 and 5.10 for reference standard authentication and additional GC-MSD quality assurance requirements.

3.9.2.9.2 Monitoring of Control Values

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

### 3.9.2.10 ANALYSIS DOCUMENTATION

3.9.2.10.1 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.

3.9.2.10.2 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.

### 3.9.2.11 REFERENCES

- 3.9.2.11.1 Procedure for High pKa Drug Compounds, Courtesy of Jim Hutchison, Montana Department of Justice, Forensic Services Division, 2005.
- 3.9.2.11.2 Procedure for Back Extraction, Courtesy of Jim Hutchison, Montana Department of Justice, Forensic Services Division, 2006.
- 3.9.2.11.3 Strong Bases Extractions Screening SOP, Courtesy of Dr. Graham Jones, Office of the Chief Medical Examiner, Edmonton, Canada, 2003
- Jones, G., *Postmortem Toxicology*. pp. 98-102, in: Clarke's Analysis of Drugs and Poisons, 3rd Edition, Moffat, A.C., Osselton, M.D. and Widdop, B., eds., Pharmaceutical Press, 2004.
- Hearn, W.L. and Walls, H.C. Strategies for Postmortem Toxicology Investigation. pp. 937-939. *In*: "Drug Abuse Handbook" S.B. Karch, ed., CRC Press, Boca Raton, FL:1998.

Forensic Ser Toxicology I		
Section Thro Blood Toxic		
3.9.2	-Liquid Extract Liquid-Liquid Drug Compou	ion Methods for Quantitative Gas Chromatography Extraction Procedure for the Recovery of Higher pK nds.
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