Idaho State Police Forensic Services Toxicology Section



### Section Four

**Analysis of Alcohol and Common Volatile Solvents** 

4.1 Quantitative Analysis for Ethanol and Qualitative Analysis for Other Volatiles by Dual Column Headspace Gas Chromatography

### 4.1.1 BACKGROUND

Fermented beverages such as beer and wine have been known and used by humans since prehistoric times. Ethanol abuse is often manifest in driving under the influence (DUI) problems, which is a worldwide concern. The National Highway Traffic Safety Administration (NHTSA) estimates that alcohol was involved in 41% of fatal automobile crashes and 7% of all crashes in 1995. chronic alcoholism also contributes to ethanol related deaths. Ethanol consumed on a regular basis can lead to the development of alcoholic hepatitis which can progress into cirrhosis, liver failure, and death. Chronic excessive ingestion of ethanol is directly associated with serious neurologic and mental disorders such as brain damage, memory loss, sleep disturbances and psychoses. Alcohol is also involved in a high percentage of domestic disputes many of which result in injury and/or death.

Notwithstanding the public perception that ethanol is stimulatory, ethanol is classified as a *Central Nervous System Depressant*. Ethanol is a psychoactive drug that is similar in most respects to sedative-hypnotic compounds.<sup>4</sup> The first mental processes to be affected are those that depend on training and previous experience.<sup>7</sup> The individual's memory, concentration, and insight are dulled and subsequently lost. The person may become overly confident and exhibit uncontrolled mood swings and/or emotional outbursts.<sup>7</sup> The effects of ethanol and other central nervous system depressants are additive, resulting in more sedation and greater impairment of driving ability.<sup>4</sup>

Ethanol is rapidly and completely absorbed from the stomach, small intestine and colon. The mechanism of absorption is a simple diffusion process, that is, alcohol moving from a region of higher to a region of lower concentration. Alcohol is soluble in both water and fat, a property that facilitates its diffusion through biological membranes. The major amount of absorption takes place in the small intestine due to its large surface area, good blood supply and thin walled membrane. The time from the last drink to peak concentrations can range between 30 and 90 minutes, depending upon the individual's stomach contents. Alcohol absorption is slowed by the presence of food in the stomach. The time period required for gastric emptying is a prime factor that contributes to

the wide variety of absorption rates of ingested ethanol observed in different individuals and under different conditions.<sup>2,7</sup> Hence, the extent of absorption in the stomach and small intestine is a function of the amount of ethanol at that site, the vascularity of the site and the surface area in contract with the blood supply.<sup>2</sup> Other factors that affect the absorption of ethanol include the type of beverage, the alcohol content and any disease state that affects normal gastric function.<sup>2</sup>

Upon absorption, ethanol is distributed to all the water containing regions of the body. Within the blood there can be significant differences between arterial and venous blood depending upon the absorption status of the individual.<sup>2</sup> In the absorptive phase, the arterial blood ethanol concentration exceeds the venous blood ethanol concentration. Analysis of venous blood therefore, underestimates the brain alcohol concentration of the individual at this point. When absorption is complete there is little difference in ethanol concentration between and arterial and venous blood.<sup>2</sup>

90 to 98 percent of ethanol is completely oxidized in the liver by reacting with the cofactor nicotinamide adenine dinucleotide (NAD) facilitated by alcohol dehydrogenase to produce acetaldehyde. Acetaldehyde is then acted upon by aldehyde dehydrogenase to form acetic acid which goes onto form carbon dioxide and water (figure 1). The amount of ethanol oxidized per unit time is roughly proportional to body weight and probably to liver weight. The remaining (unoxidized) alcohol is excreted unchanged in urine, expired air, saliva and sweat. The average elimination rate of ethanol is 0.015 g/dL/hour from men and 0.018 g/dL/hour for women.<sup>2</sup> In addition to gender, chronic abuse, ethanol use combined with prescription drugs and certain genetic factors can also influence the elimination rate.<sup>2,6,7</sup>

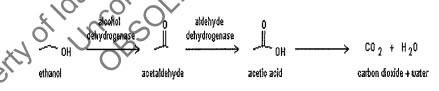


Figure 1. Metabolism of Ethanol.

Methanol (wood alcohol) causes relatively little intoxication compared to ethanol.<sup>2,6</sup> Its harmful affects are due to the direct result of its metabolism to formaldehyde (embalming fluid) and subsequently to formic acid. These metabolites lead to the destruction of neural cells, particularly the optic nerve, which can result in blindness.<sup>2,6</sup>

### 4.1.2 PRINCIPLE

This method describes the analysis of aqueous samples for the presence of volatile compounds including methanol, ethanol,

acetaldehyde, acetone, isopropanol and related compounds, via a headspace sampling gas chromatographic method. Samples, controls and standards are sealed into vials that contain an aqueous 1-propanol internal standard solution and heated by the headspace analyzer. As described in Henry's Law, in a closed container at a given temperature, a direct (proportional) relationship exists between the amount of a volatile substance dissolved in a liquid and the amount of the volatile substance in the headspace vapor above the solution. An aliquot of the vapor is injected into a gas chromatograph (GC) in a dual column configuration. The GC serves to separate out the components of the solution as a function of their chemical properties. The separated components are identified on the basis of the retention time determined for each of the columns. Quantitation is accomplished through area percent data obtained from a flame ionization detector (FID). The quantitative result is based on a minimum of a threepoint calibration curve, which uses the peak area ratio between the analyte and the internal standard.

### 4.1.3 EQUIPMENT

- Perkin Elmer Auto System XL Gas Chromatograph (GC) 4.1.3.1
- 4,1,3,2 **Columns** 
  - Restek Rtx<sup>®</sup>-BAC1 (#18003: 30 meter X 0.32mm 4.1.3.2.1 inner diameter (ID), 1.8µm film thickness (FT)) or equivalent column
  - Restek Rtx® BAC2 (#18002: 30 meter X 0.32mm 4.1.3.2.2 2 µm film thickness (FT)) or equivalent rolumn
- Perkin Elmer HS-40 or HS-110 Headspace Autosampler (figures 2



Figure 2. HS-40

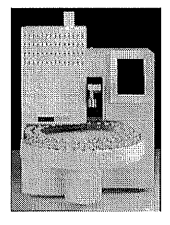


Figure 2. HS-110

- 4.1.3.4 PE Workstation Software, TotalChrom Version 6.2.0 or more recent version/upgrade.
- 4.1.3.5 Hand Crimper (P-E B003-8134 or equivalent)

4.1.3.6	Hamilton MICROLAB 503A or equivalent semi-automatic Dilutor/Pipetter equipped with sample and reagent syringes capable of dispensing 250µL and 2000µL, respectively.
4.1.3.7	Glassware 4.1.3.7.1 GC-Headspace vials (P-E B010-4236 or equivalent) Safety Closures {PTBE septa, crimp caps and star springs} (P-E B010-4240 or equivalent)
CONTROLS	AND CALIBRATORS
4.1.4.1	Whole Blood Ethanol Control (LiquiSP <sub>x</sub> <sup>™</sup> or equivalent).
4.1.4.2	Aqueous Ethanol Standards (g/100mL) 0.025, 0.05, 0.08, 0.10, 0.20, 0.30, and 0.40 (Cerilliant or equivalent)
4.1.4.3	Multicomponent alcohol Calibration Kit (Cerilliant #A-054 or equivalent)
REAGENTS	4° 07
4.1.5.1	1-Propanol (Acros/Fisher Scientific #23207-0010, #A996-1 or
4,1,5,1	equivalent)
4.1.5.2	Acetone (Fisher #A929-1 or equivalent)
4.1.5.3	Acetaldehyde (Fisher #01004-250 or equivalent)
4.1.5.4	Isopropanol (2-Propanol) (Fisher #A416-500 or equivalent)
4.1.5.5	Methanol (Fisher #A454-1 or equivalent)
4.1.5.6	Ammonium Sulfate (Fisher #A702-500 or equivalent)
4.1.5.7	Sodium Fluoride (Fisher #S299-500 or equivalent)
	daho nitie E
SAFETY CO	NCERNS
4.1.6.1	Blood samples should be processed according to safety guidelines
Obely,	in the Chemical Hygiene and Safety Manual.
REAGENT P	PREPARATION
	eparation of all reagents on reagent log.
4.1.7.1	Internal Standard Solution
	{0.03g/dL 1-propanol in 1.0M (NH <sub>4</sub> ) <sub>2</sub> SO <sub>4</sub> }
	4.1.7.1.1 1.0M (NH <sub>4</sub> ) <sub>2</sub> SO <sub>4</sub> Dissolve 132.14g (NH <sub>4</sub> ) <sub>2</sub> SO <sub>4</sub> in distilled water.
	Dilute to 1L.  4.1.7.1.2  Dilute to 1L.  0.03g/dL 1-propanol in 1.0M (NH <sub>4</sub> ) <sub>2</sub> SO <sub>4</sub> Add approximately 800mL of 1.0M (NH <sub>4</sub> ) <sub>2</sub> SO <sub>4</sub> to a 1000mL volumetric flask.  Add 1g sodium fluoride {optional}.

4.1.4

4.1.5

4.1.6

4.1.7

- Add 375μL 1-propanol. QS to 1000mL with 1.0M (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>.
- 4.1.7.1.3 Solution is stable for 3 months.

# Volatile Standard Mix Solution 4.1.7.2.1 Add approximately 200 mL of DI water to a 250-mL volumetric flask. 4.1.7.2.2 Add the following volatiles, as indicated: 100 μL acetaldehyde 100 μL acetone 500 μL methanol 500 μL isopropanol 500 μL ethanol 4.1.7.2.3 QS to 250-mL.

Solution is stable for 1 year.

# 4.1.8 ANALYSIS PROCEDURE

4.1.8.1

4.1.7.2.4

General

4.1.8.2.6

	4.1.8.1.1	Bring calibrators, controls, internal standard and
		samples to room temperature.
	4.1.8.1.2	Gather necessary vials, closures and ancillary
		supplies in or near laminar flow hood.
	4.1.8.1.3	Sample preparation should take place in a laminar
	5	flow hood,
	0	(O), (V)
4.1.8.2	Quality Conti	<u>01</u>
	4.1.8.2.1	Ethanol calibration standards must be run prior to
		The analysis of each batch of samples. A minimum
	0,0,0	of three points of calibration should be established.
	4.1.8.2.2	An internal standard blank should follow the last
·00.		ethanol calibrator.
40%	4.1.8.2.3	A blood or aqueous control sample must be run
Q\		after every 10 case samples. A minimum of two
*		blood controls must be run per batch of samples.
	4.1.8.2.4	Refer to package insert for manufacturer blood
	7,1,0,2,7	control ranges.
	41005	
	4.1.8.2.5	Values obtained from aqueous control and whole
		blood control samples must agree ± 10% of their

target values.

or

the

the retention of other volatiles of interest.

Solution

Periodically run either the Volatile Standard Mix

Calibration Kit solution to determine and monitor

Multicomponent

Alcohol

	4.1.8.2.7	Record values  Analysis QC l	s for blood control samples in Batch	
	4.1.8.2.8	· -	y basis calculate the mean, standard	
	7,1.0,2.0	-	ative standard deviation (CV%) and	
		•	acy of the control samples. The data	
			to generate a mean quality control	
		chart.	to generate a mean quanty control	
	4,1,8,2,9		control lots should be analyzed a	
	4.1.0.2.3		nine times prior to official use.	
			mean, standard deviation, relative	
		standard dard	ation (CV%) and percent accuracy of	
		the control sar	nnlag	
		the control sai	inples,	
4,1,8,3	Dinattar/Dilute	or Cet un	nples.  ver.  inquire as to the sizes of installed	
4,1,0,3	Pipetter/Dilute		7709	
	4.1.8.3.1	Switch on pov	inquire age to the gizeg of installed	
	4.1.8.3.2		miduito no to mie primo or minimon	
			ect the correct size for sample syringe	
	4 1 0 2 2		gent syringe [left].	
	4.1.8.3.3		to volume option. Select 250µL for	
			ge [right] and 2000μL for reagent	
	41004	syringe [left].	XX 10 dl d	
	4.1.8.3.4	Scroll down	o speed option. Verify that syringe	
	41005	speed is on de		
	4.1.8.3.5	Prime the flu		
	CX.	bubbles are of	bserved.	
4.1.8.4	Control and Mixed Standard			
1111011		Water Blank		
	4.1.8.4.1	4.1.8.4.1.1	Label test vial with water blank.	
1	* 10, 200 ×	4.1.8.4.1.2	Add 2000µL DI water to labeled test	
, (	0, 11, 00	5.1.0.7.1.2	tube.	
Ex	000	4.1.8.4.1.3	Seal <b>immediately</b> with crimp cap as	
00/		1,110,112,10	illustrated in figure 4.	
blobetr,				
010	4.1.8.4.2	Internal Stand	ard Blank	
		4.1.8.4.2.1	Label test vial with ISTD blank.	
		4.1.8.4.2.2	Use Pipetter/Dilutor to dispense	
			2000µL of internal standard (ISTD)	
			into labeled headspace vial.	
		4.1.8.4.2.3	Seal immediately with crimp cap as	
			illustrated in figure 4.	
			<b>5</b>	
	4.1.8.4.3	Blood Control	•	
		4.1.8.4.3.1	Label two headspace vials for blood	
			control 1 and 2.	

- 4.1.8,4.3.2 Use Pipetter/Dilutor to dispense 250μL of blood control and 2000μL of internal standard (ISTD) into each labeled headspace vial. 4.1.8.4.3.3 Seal immediately with crimp cap as illustrated in figure 4. Aqueous Controls 4.1.8.4.4.1 appropriate number headspace vials for aqueous controls (1, 2,...).Use Pipetter/Dilutor to 4.1.8.4.3.2 dispense 250µL of aqueous control and
  - into each labeled headspace vial.
    4.1.8.4.3.3 Seal **immediately** with crimp cap as illustrated in figure 4.

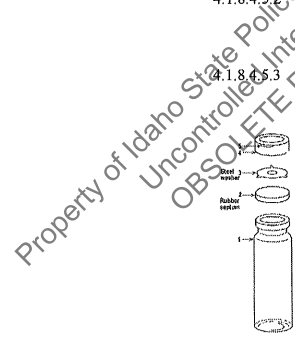
2000µL of internal standard (ISTD)

# 4.1.8.4.5 <u>Mixed Other Volatiles Solution</u>

4.1.8.4.5.1 Label test vial with mixed volatiles.

4.1.8.4.5.2 Use Pipetter/Dilutor to dispense 250μL of mixed volatile solution and 2000μL of internal standard (ISTD) into labeled headspace vial.

4.1.8.4.5.3 Seal **immediately** with crimp cap as illustrated in figure 4.



4.1.8.4.4

Figure 4. Crimp cap assembly

# 4.1.8.5 <u>Preparation Calibration Standards</u>

4.1.8.5.1 Label vials for standards.

4.1.8.5.2 Use Pipetter/Dilutor to dispense 250μL of appropriate ethanol concentration and 2000μL of

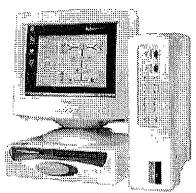
internal standard (ISTD) into each labeled headspace vial.
Seal immediately with crimp cap.
Establish ethanol calibration plot with a minimum of three calibration points.

Sing of Specimens
Open the sample submittal kit and remove the specimen's inner compartment. After inspecting

4.1.8.6 <u>Initial Processing of Specimens</u>

4.1.8.5.3 4.1.8.5.4

- 4.1.8.6.1 Open the sample submittal kit and remove the specimen's inner compartment. After inspecting and noting the condition of seals, open inner compartment (plastic tray or biohazard bag) and place laboratory number on each blood/urine/vitreous humor specimen.
- 4.1.8.6.2 When two blood/fluid samples are present, the samples should be labeled "A" and "B" or equivalent. Utilize sample "A" for analysis unless it contains insufficient sample.
- 4.1.8.7 <u>Preparation of Samples for Analysis</u>
  - 4.1.8.7.1 Label two headspace vials with the laboratory number without the prefix.
  - 4.1.8.7.2 Place one of the sample tubes or urine specimen bottle on tube rocker for at least two minutes.
- 4.1.8.8 Addition of blood, wrine or vitreous humor sample to headspace vials.
  - 4.1.8.8 1 Use Pipetter/Dilutor dispense 250µL of sample and 2000µL of internal standard (ISTD) to a labeled beadspace vial.
  - 4.1.8.8.2 Seal headspace vials **immediately** with crimp caps as illustrated in figure 4.
- 4.1.8.9 Preparation for Run
  - 4.1.8.9.1 Open Sequence Editor
  - 4.1.8.9.2 Into Sequence log table, enter the sample case numbers, ethanol standards, other volatiles mix, blanks and controls.



Issued: 05/02 tox-man-PE-HSA.doc 4.1.8.9.3 Load samples, calibration standards, blank and controls into the carousel of the headspace sampler as noted in the sequence table.

# 4.1.8.9.4 Active headspace sampler

- Click on the **Setup** button to open the setup instrument dialog box.
- Select sequence as the setup type, and select the desired sequence file.
- On **Setup Instrument** dialog box, designate starting and ending row.
- Verify that the paths for raw and result data files specified in the sequence indicate the desired destinations.
- Select OK in the **Setup instrument** dialog box to initialize the instrument.

# 4.1.8.10 Gas Chromatography Parameters Q

4.1.8.10.1 Refer to instrument METHOD printout for oven program and zone temperatures. Temperature program must provide for baseline separation of volatile compounds of interest as indicated by analysis of multicomponent mixtures.

# 4.1.8.11 <u>Calibration</u>

4.1.8.11.1 Ethanol calibrators should be analyzed in order of increasing concentration.

4.1.8.11.2 The least squares line resulting from the analysis of the ethanol calibrators must have a coefficient of correlation of ≥0.995.

# 1.1.8.12 Acceptance Criteria

4.1.8.12.1 Accuracy

# 4.1.8.12.1.1 Qualitative

The presence of ethanol can be established if there are no significant differences in the retention time between sample and standards. The relative retention times specimen must be within  $\pm$  0.10 minutes of the relative retention time for the compound in question. This criterion should rejection be designated in the TotalChrom analysis method.

# 4.1.8.12.1.2 Quantitative

The quantitative results for a batch of samples can be accepted if the values obtained for control samples fall within 10% of their target value range.

### 4.1.8.12.2 **Precision**

The results obtained from duplicate analysis must agree within 0.015g/100mL. If this precision requirement is not met, the sample is reanalyzed.

# 4.1.8.13 Reporting of Results

### 4.1.8.13.1 **Blood**

Samples are quantitated to three significant figures. Report truncated mean value, of grams of ethanol per 100cc of whole blood, to two significant figures.

### 4.1.8.13.2 Urine

Samples are quantitated to three significant figures. Result obtained from blood alcohol curve should be multiplied by 0.67. Report truncated mean value, as grams of ethanol per 67 mL of urine, to two significant figures. A warning statement such as Urine results may be of questionable value, must be included in the report.

Vitreous Human

# 4.1.8.13.3 Vitreous Humor

Samples are quantitated to three significant figures. Report truncated mean value, as grams of ethanol per 100mL of vitreous humor, to two significant figures.

# 4.1.9 QUALTTY ASSURANCE

- 4.1.91 Blood or vitreous samples are to be refrigerated while at the laboratory. Urine samples can be either refrigerated or frozen.
- 4.1.9.2 Refer to toxicology manual section 5.1 for pipette calibration options.
- 4.1.9.3 Refer to toxicology manual section 5.2 for balance calibration requirements.
- 4.1.9.4 Refer to toxicology manual section 5.3.2 for GC-HS maintenance schedule.
- 4.1.9.5 Blood calibrators should be ordered prior to the current supply running out. This will allow for the analysis of new lots against existing calibrators.

### 4.1.10 REFERENCES

- 4.1.10.1 Stafford, D.T., *Chromatography. in:* Principles of Forensic Toxicology, edited by Barry Levin, pp. 93-101, 103-114, AACC Press, 1999.
- 4.1.10.2 Levine, B., *Alcohol. in:* Principles of Forensic Toxicology, edited by Barry Levin, pp. 170-184, AACC Press, 1999.
- 4.1.10.3 Caplan, Y.H., *The Determination of Alcohol in Blood and Breath. in:* Forensic Science Handbook, edited by Richard Saferstein, pp. 594-648, Prentice-Hall New Jersey, 1982.
- 4.1.10.4 Julien, R.M., Central Nervous System Depressants: Alcohol and the Inhalants of Abuse, in: Primer of Drug Action, pp. 64-92, Freeman-New York, 1998.
- 4.1.10.5 Saker, E.G., Screening and Quantitation by Head Space Technique of Some of the Vapors Most Commonly Found in Forensic Toxicology, in: Current Approaches in Forensic Toxicology, Chapter 11, SOFT Meeting, 1994.
- 4.1.10.6. Perrine, D.M., Depressants: Alcohol, Benzodiazepines, Barbiturates, in: The Chemistry of Mind-Altering Drugs, pp. 113-129, ACS, Washington, DC, 1996.
- 4.1.10.7 Hobbs, W.R., Rall, T.W. and Verdoorn, T.A., Drugs Acting on the Central Nervous System Hypnotics and Sedatives; Ethanol, in: Goodman and Gilman's The Pharmacological Basis of Therapeutics, pp. 361, 386-393, McGraw-Hill, 1996.
- 4.1.10.8 Idaho Administration Code, IDAPA 11.03.01, Rules Governing Alcohol Testing.
- 4.1.10.9 Christmore, D.S., Kelly, R.C. and Doshier, L.A. Improved Recovery and Stability of Ethanol in Automated Headspace Analysis, J. Forensic Sci. 29(4): 1038-1044; 1984.
- 4.1.10.10 Restek Applications Note #59598, Dual-Column Confirmational GC Analysis of Blood Alcohols Using the Rtx<sup>®</sup>-BAC1 and Rtx<sup>®</sup>-BAC2 Columns Optimized for the Perkin-Elmer HS-40 Headspace Autosampler, 1999.