Capillary GC Analysis of Fusel Oils and Other Components of Interest

Scope and Application

Methanol, acetaldehyde, ethyl acetate and fusel oils (F.O.) are natural fermentation products. Fusel Oils consist of n-propanol, iso-butanol, n-butanol, and amyl alcohol (2-methyl-1-butanol and 3-methyl-1-butanol). Absence of these compounds in products suggests either non-fermented products or the use of neutral spirits. Acetic acid is indicative of spoilage. Benzaldehyde and propylene glycol (greater than 0.01%) are indicative of flavoring/adulteration. Glycerol (1,000-2,000 ppm) is present naturally in wines.

This method may be used for the analysis of most alcohol beverages and nonbeverage alcohol (NBA) products with the following caveats:

1. NBA products which are NOT miscible with water cannot be analyzed by this method.
2. NBA products containing >10% solids must be diluted prior to analysis for nonbeverage analytes.
3. Distilled spirit products containing >10% solids are diluted (or distilled) prior to analysis. Liqueur products are distilled prior to analysis.
4. Acetic acid, benzaldehyde, propylene glycol, and glycerol cannot be determined after distillation.

Regulatory Tolerances:

Methanol—0.1 % by volume max. in wine (Industry Circular IC-93-3)(CPG 7119.09 Section 510.200).

Methanol—0.35 % by volume max. in brandy (FDA Administrative Guides 7401.01 and 1701.01)(Topical Digest 1710.41-43)(CPG7119.09)

Fusel Oil—less than 20 ppm indicates neutral spirits (Commodity Classification Branch 4/4/1983)

Volatile Acidity/acetic acid (27CFR4.21) —

0.14 % by volume max. in red wine when starting brix ≤ 28
0.17 % by volume max. in red wine when starting brix is >28
0.12 % by volume max. in white wine when starting brix ≤ 28
0.15 % by volume max. in white wine when starting brix is >28

Propylene Glycol – For NBP’s, ±5% of the stated value. The finished alcohol beverage may not contain more than 5% PG (21 CFR 184.166).

Acetic Acid – For NBP’s, ±5% of the stated value. The finished alcohol beverage may not contain more than 0.15% acetic acid (21 CFR 184.1005).
## Levels and Limitations

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Detection Limit</th>
<th>Quantitation Limit</th>
<th>Calibration Range</th>
<th>Validated Linear Range</th>
<th>Interferences</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ethyl Acetate</td>
<td>1.8 mg/L (0.18 g/100L)</td>
<td>5.9 mg/L (0.59 g/100L)</td>
<td>9-900 mg/L (0.9-90.0g/100L)</td>
<td>5.9-20000 mg/L (0.59-2000 g/100L)</td>
<td>None</td>
</tr>
<tr>
<td>Methanol</td>
<td>0.0004 %</td>
<td>0.005 %</td>
<td>0.015-1.5 %</td>
<td>0.01-20%</td>
<td>None</td>
</tr>
<tr>
<td>n-Propanol</td>
<td>0.5 mg/L (0.05 g/100L)</td>
<td>1.6 mg/L (0.16 g/100L)</td>
<td>3-300 mg/L(0.3-30g/100L)</td>
<td>1.6-10000 mg/L (0.16-1000 g/100L)</td>
<td>None</td>
</tr>
<tr>
<td>iso-Butanol</td>
<td>0.5 mg/L (0.05 g/100L)</td>
<td>1.7 mg/L (0.17 g/100L)</td>
<td>6-600 mg/L(0.6-60g/100L)</td>
<td>1.7-20000 mg/L (0.17-2000 g/100L)</td>
<td>None</td>
</tr>
<tr>
<td>n-Butanol</td>
<td>0.3 mg/L (0.03 g/100L)</td>
<td>0.9 mg/L (0.09 g/100L)</td>
<td>3-300 mg/L(0.3-30g/100L)</td>
<td>1.25-2000 mg/L (0.125–200 g/100L)</td>
<td>None</td>
</tr>
<tr>
<td>Amyl Alcohol</td>
<td>0.3 mg/L (0.03 g/100L)</td>
<td>1.0 mg/L (0.10 g/100L)</td>
<td>12-1200 mg/L(1.2-120g/100L)</td>
<td>2-40000 mg/L (0.2-4000 g/100L)</td>
<td>None</td>
</tr>
<tr>
<td>Acetic Acid **</td>
<td>0.001 g/100mL</td>
<td>0.0032 g/100mL</td>
<td>0.01-0.20 g/100mL</td>
<td>0.002-0.2 g/100mL</td>
<td>Furfural</td>
</tr>
<tr>
<td>Benzaldehyde</td>
<td>0.0014 g/100mL</td>
<td>0.0046 g/100mL</td>
<td>0.01-0.20 g/100mL</td>
<td>0.002-0.2g/100mL</td>
<td>None</td>
</tr>
<tr>
<td>Propylene Glycol</td>
<td>0.0010% by Vol</td>
<td>0.0033 % by Vol</td>
<td>0.01-0.20 % by Vol</td>
<td>0.002-0.2% by Vol</td>
<td>None</td>
</tr>
<tr>
<td>Glycerol</td>
<td>0.006 % by Vol</td>
<td>0.019 % by Vol</td>
<td>0.05-1.00 % by Vol</td>
<td>0.01-1% by Vol</td>
<td>None</td>
</tr>
<tr>
<td>Acetaldehyde</td>
<td>4.9 ppm</td>
<td>16 ppm</td>
<td></td>
<td>40-2000 ppm</td>
<td>None</td>
</tr>
</tbody>
</table>

** When distilled, Acetic Acid in the presence of ethanol may react to form ethyl acetate.
Equipment

Instrumentation:

GC: Hewlett Packard 6890 with 7673 Autosampler and Chemstation Software, or equivalent
Column: DB-WAXETR, 30m x 0.53mm x 1μm film thickness
Carrier Gas: Hydrogen, from generator, Proton Model G600 or equivalent, constant flow, 5.9 ml/min
Temperature: 40°C initial, hold 5 min, ramp at 10°/min to 215°C, hold 2.5 min.
Injector: 220°C, 5:1 split
Detector: FID 250°C, Hydrogen flow 40 ml/min, air flow 450 ml/min
Injection Volume: 1 μL

Glassware and Supplies:

Class A pipets / Micropipettes
Class A volumetric flasks

Reagent and Sample Preparation and Handling

**Reagents:** (All chemicals for standards are 99.0+% pure.)

- 40% Ethanol/Water
- Ethyl Acetate
- n-Propanol
- iso-Butanol
- n-Butanol
- 2-methyl-1-butanol (active Amyl alcohol)
- Methanol
- Propylene Glycol
- Glycerol
- Acetic Acid
- Benzaldehyde
- Acetaldehyde
- 200 Proof Ethanol

**Preparation of Fusel Oil (F.O.) stock, working, and second source standards:**

1. Prepare a **F.O. stock standard solution**. Weigh the following into a 100 mL volumetric flask and quantitate to volume with >95% Ethanol by volume. Stock solution is stable in the refrigerator for up to 12 months.
   a. 3.00 g ethyl acetate ± 1.0%
   b. 1.00 g n-propanol ± 1.0%
   c. 2.00 g iso-butanol ± 1.0%
   d. 1.00 g n-butanol ± 1.0%
   e. 4.00 g active amyl alcohol ± 1.0%
   f. 50.00 mL methanol ± 1.0%
2. Prepare F.O. working standards by transferring stock or diluted standard as outlined below. Use glass volumetric pipettes or positive displacement pipettes. The standards may be stored in the refrigerator for up to 3 months.

**F.O. Level 4:** Transfer 6 mL standard stock solution into 200 mL volumetric flask or equivalent concentration. Q.S. with 40% ethanol by volume.

**F.O. Level 3:** Transfer 1 mL standard stock solution into 200 mL volumetric flask or equivalent concentration. Q.S. with 40% ethanol by volume.

**F.O. Level 2:** Transfer 5 mL Level 4 into 200 mL volumetric flask or equivalent concentration. Q.S. with 40% ethanol by volume.

**F.O. Level 1:** Transfer 2 mL Level 4 into 200 mL volumetric flask or equivalent concentration. Q.S. with 40% ethanol by volume.

3. Prepare F.O. second source stock and working solution (F.O. Level 3 concentration) in the same manner as step 1 and 2.

This preparation results in F.O. standards with the following concentrations:

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Stock Std. ppm</th>
<th>FO4 ppm</th>
<th>FO3 and 2nd source ppm</th>
<th>FO2 ppm</th>
<th>FO1 ppm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ethyl Acetate</td>
<td>30,000</td>
<td>900</td>
<td>150</td>
<td>22.5</td>
<td>9</td>
</tr>
<tr>
<td>n-propanol</td>
<td>10,000</td>
<td>300</td>
<td>50</td>
<td>7.5</td>
<td>3</td>
</tr>
<tr>
<td>Iso-butanol</td>
<td>20,000</td>
<td>600</td>
<td>100</td>
<td>15</td>
<td>6</td>
</tr>
<tr>
<td>n-butanol</td>
<td>10,000</td>
<td>300</td>
<td>50</td>
<td>7.5</td>
<td>3</td>
</tr>
<tr>
<td>Amyl alcohol</td>
<td>40,000</td>
<td>1200</td>
<td>200</td>
<td>30</td>
<td>12</td>
</tr>
<tr>
<td>Methanol</td>
<td>500,000 (1.5%)</td>
<td>15000</td>
<td>2500 (0.25%)</td>
<td>375 (0.037%)</td>
<td>150 (0.015%)</td>
</tr>
</tbody>
</table>

**Preparation for NBA stock and working standards:**

1. Prepare a **NBA standard stock solution** to proper concentration. Place the amounts below into a 1 L volumetric flask. Q.S. with 40% Ethanol by volume. Store stock solution in the refrigerator for up to 12 months.
   a. 2.07 g Propylene Glycol ± 1.0%
   b. 12.50 g Glycerol ± 1.0%
   c. 2.00 g Acetic Acid ± 1.0%
   d. 2.00 g Benzaldehyde ± 1.0%
2. Prepare the following **NBA working standards**:

   NBA Level 3: NBA Stock Standard, as is.

   NBA Level 2: Pipet 5 mL of NBA standard stock solution into a 10 mL volumetric flask. Q.S. with water. Prepare daily.

   NBA Level 1: Pipet 0.5 mL of NBA standard stock solution into a 10 mL volumetric flask. Q.S. with water. Prepare daily.

   This preparation results in NBA standards of the following concentrations:

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Units</th>
<th>Stock/Level 3</th>
<th>Level 2</th>
<th>Level 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>propylene glycol</td>
<td>% by Vol</td>
<td>0.2</td>
<td>0.1</td>
<td>0.01</td>
</tr>
<tr>
<td>glycerol</td>
<td>% by Vol</td>
<td>1.0</td>
<td>0.5</td>
<td>0.05</td>
</tr>
<tr>
<td>acetic acid</td>
<td>g/100mL</td>
<td>0.2</td>
<td>0.1</td>
<td>0.01</td>
</tr>
<tr>
<td>benzaldehyde</td>
<td>g/100mL</td>
<td>0.2</td>
<td>0.1</td>
<td>0.01</td>
</tr>
</tbody>
</table>

**Preparation for Acetaldehyde standard**:

1. Prepare an **Acetaldehyde Standard** by pipeting 1 mL of acetaldehyde into a 1000 mL volumetric flask. Q.S. with 40% Ethanol by volume. Prepare daily.

**Procedures**

1. Check to see if potential high solid samples (e.g., NBA, enforcement, salted wine samples) were run with the current liner. If so, replace the liner.

2. Run a wash and a blank at beginning of sequence. A wash is used to condition the system and a blank is used to evaluate the cleanliness of the system.

3. Run standards on the GC. The choice of standards depends on the sample type and is typically as follows:

   - For **beer and DSP**, use Fusel Oil Standards 1-4
   - For **wine**, use Fusel Oil Standards 1–4. NBA Level 1–3 is used when an acetic acid value is required.
   - For **nonbeverage products**, use NBA Level 1-3 and/or Fusel Oil Standards 1-4.
   - For **moonshine**, use Fusel Oil Standards 1-4 and the Acetaldehyde Standard.

4. Calibrate based on area using external standards on all levels for each component. The correlation coefficient (r) shall be > 0.99.

5. Run a blank after the highest standard to assure the system is free from carryover.

6. Run a second source check to verify the system, then inject samples. Bracket the samples with a second source check after every ten (10) samples and at the end
of the sequence as a control to assure the system is free from carryover and as a control to check for drift, respectively.

7. Inject a wash (e.g., 40% ethanol) after potentially “dirty” sample (i.e., high solids, high contaminant) to assure the system cleans out properly prior to next injection.

8. If sample results are above the calibration range, dilute into range and re-inject.

Quality Control

1. The correlation coefficient of the external standards is to be greater than 0.99. If the correlation coefficient is not >0.99, change the liner and/or septa and rerun. If the correlation coefficient is still not >0.99, re-run using fresh working standards. If the correlation coefficient remains out of spec, contact the principal analyst.

2. Run an LCS check in duplicate as appropriate for the sample type (i.e., neat LCS with neat samples, distillate LCS with distillate samples). The accuracy and precision values are to be within the prescribed limits.

3. Wash injections are allowed to contain carryover >LOQ. Blank injections are acceptable if results are less than LOQ for all analytes being reported. If unacceptable, do not report affected results and inspect liner, syringe/plunger and replace or clean if necessary.

4. If the second source standard differs by >15% of the expected value, repeat the sample injections performed since the last passable control result.

If any second source analytes exceeds the tolerance limits, inspect the liner and syringe and clean or replace as necessary. Remake second standard(s) if results exceed the tolerance limits on more than one instrument.

5. Violative results are confirmed as follows:
   - Propylene Glycol and Glycerol - confirm using LC Acids Method.

6. For precision quality control failures, inspect the liner and syringe and replace if necessary.

7. If neat LCS values fall outside the acceptable accuracy range for at least two out of three results, re-run the calibration curve using a fresh aliquot of the calibration standards.

Sources of Uncertainty

1. Weighing errors for standards
2. Preparation of working standards (e.g., Dilution, pipet, etc.)
3. Dirty injection liner
4. Problem with GC syringe (e.g., dirty syringe or bad plunger)
5. Change in analyte retention time
Calculations

GC is operated in external standard mode with calculations using peak areas. Total Fusel Oil is the sum of n-propanol, iso-butanol, n-butanol and amyl alcohols.

Reporting Results

Report the results as follows:

<table>
<thead>
<tr>
<th>Component</th>
<th>Sample Type</th>
<th>Units</th>
<th>Precision</th>
<th>Format</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetaldehyde</td>
<td>All</td>
<td>mg/L (ppm)</td>
<td>No decimal</td>
<td>XX</td>
</tr>
<tr>
<td>Acetic Acid</td>
<td>All</td>
<td>g/100mL</td>
<td>2 decimals</td>
<td>X.xx</td>
</tr>
<tr>
<td>Benzaldehyde</td>
<td>All</td>
<td>g/100mL</td>
<td>1 decimals</td>
<td>X.x</td>
</tr>
<tr>
<td>Ethyl Acetate</td>
<td>DSP</td>
<td>g/100L</td>
<td>1 decimal</td>
<td>X.x</td>
</tr>
<tr>
<td>Ethyl AcetateDSP</td>
<td>Wine, Beer, NBA</td>
<td>mg/L (ppm)</td>
<td>No decimal</td>
<td>XX</td>
</tr>
<tr>
<td>Fusel Oils (Total and indiv. components)</td>
<td>DSP</td>
<td>g/100L</td>
<td>1 decimal</td>
<td>X.x</td>
</tr>
<tr>
<td>Fusel Oils (Total and indiv. components)</td>
<td>Wine, Beer, NBA</td>
<td>mg/L (ppm)</td>
<td>No decimal</td>
<td>XX</td>
</tr>
<tr>
<td>Glycerol</td>
<td>All</td>
<td>% by volume</td>
<td>2 decimals</td>
<td>X.xx</td>
</tr>
<tr>
<td>Methanol</td>
<td>All</td>
<td>% by volume</td>
<td>2 decimals</td>
<td>X.xx</td>
</tr>
<tr>
<td>Propylene Glycol</td>
<td>All</td>
<td>% by volume</td>
<td>2 decimals</td>
<td>X.xx</td>
</tr>
</tbody>
</table>

Safety Notes

Consult the MSDS for any chemicals used that are unfamiliar. All chemicals shall be considered hazardous - avoid direct physical contact.

Hydrogen is explosive and is used as a carrier gas. Extreme caution shall be used when working with the GC hardware.

If the GC is not equipped with a Hydrogen leak sensor to automatically shut down the GC, this method shall not be used.

References

(Note: Similarities – Method 968.09 is the basis for SSD:TM:200 to separate higher alcohols in distilled liquors using GC-FID; Differences – Method 968.09 has been modified with different standard mixtures, and instrumentation/column).

The colored ink stamp indicates this is a controlled document. Absence of color indicates this copy is not controlled and will not receive revision updates.

(Note: Similarities – Method 972.11 prepared methanol standard solutions in 40% ethanol without the use of internal standard; Differences – Method 972.11 differed in the standard mixture, quantifying using peak height instead of peak area, and GC column and operating parameters).

Title: Gas-Liquid Chromatographic Determination of Congeners in Alcoholic Products with Confirmation by Gas Chromatography/Mass Spectrometry.
(Note: Similarities – standard mixture in 40% ethanol; Differences – SSD:TM:200 was modified to updated instrumentation/column).

Kelly, J., et al. (1999) J. AOAC Int. 82, 6, 1375-1388.
Title: Gas Chromatographic Determination of Volatile Congeners in Spirit Drinks: Interlaboratory Study.
(Note: Similarities – calculation using peak area; Differences – differed from SSD:TM:200 with use of internal standard, and modification of instrument parameters/column dimension)

Location of Validation Package.

Quality System Files

Required Training, Certification and Re-certification.

1. In-house training by a certified chemist in GC and chemstation operation. Training on GC (in-house or vendor provided).
2. Periodically, chemists are re-tested for competency (e.g., every 5 years) and/or given proficiency testing.

Revision History.

Revision 4 – changes as a result of a document review to clarify and harmonize units used in the test method – 11/14/2008

Revision 5 – Change reporting of Propylene Glycol to 2 decimal places from 1 – 9/1/2009

Revision 6 – changed DL, QL and linear range units to match reporting units; added values to DL, QL and linear range to cover both DS and wine units.

Revision 7 – changes to calibrant levels used; changes to LOD and LOQ, edits for clarity and to better reflect lab practices; addition of Sources of Uncertainty; addition of what to try for precision QC failures (Quality control section)

NOTE: Revision 7 had errors in units in the standard concentrations. This has been revised and issued prior to implementation date. 10/29/2014

Revision 8 – addition of second source solution instructions; added instructions to Quality Control; added Blanks to the procedure; added requirement for running a second source check, as well as bracketing samples with a second source check to the procedure.
Revision 9 – additions to procedure for quality items; differentiation between ‘blank’ and ‘wash’ injection; added instruction to quality control for second source and LCS; update references listed.