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# **ICBA**

# Guidance Document to Mitigate the Potential for Benzene Formation in Beverages

Adopted by the ICBA Council on 29 April 2006

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## **ICBA Guidance Document to Mitigate the Potential for Benzene Formation in Beverages**

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### ICBA Guidance Document to Mitigate the Potential for Benzene Formation in Beverages

### 1. Introduction

The International Council of Beverages Associations (ICBA) is a non-governmental organization representing the interests of the worldwide beverage industry. The members of ICBA produce, distribute, and sell a variety of non-alcoholic beverages, including carbonated soft drinks and non-carbonated beverages such as juice-drinks, bottled waters, and ready-to-drink coffees and teas.

### 2. <u>Background</u>

In 1990-1991, the soft drink industry learned that elevated benzene levels could be found in select beverages under certain conditions. Working with the US Food and Drug Administration (FDA), the industry found that when ascorbic acid (Vitamin C) was used as an ingredient along with sodium benzoate (a preservative), benzene formation could occur. This formation was exacerbated when the beverage was stored for extended periods at elevated temperatures.

Although the levels and frequency at which this benzene formation occurred were not considered to pose a public health risk, the industry immediately took proactive steps to reformulate affected products in order to minimize any formation potential, while still ensuring microbiological integrity.

For soft drinks, and other foods and beverages, regulatory authorities e.g. Australia and New Zealand FSANZ, EU JRC, Canada, UK FSA and US FDA use a comprehensive exposure monitoring and evaluation approach to risk assessment. The latest study, Volatile Organic Compounds in Foods: A Five Year Study was conducted by the FDA and published in the Journal of Agriculture and Food Chemistry in 2003. Benzene was found in all foods tested, including fruit and vegetables, apart from in American cheese and vanilla ice cream. These levels ranged from 1-190 ppb (microgrammes per kilo). FDA concluded that data collected during the study demonstrated that the American food supply is comparatively safe and that although there is some oral exposure to volatile organic compounds such as benzene, exposure is actually much higher through inhalation. In fact, according to an article which appeared in the February 27, 2006 Food Chemical News, an official from FDA's Center for Food Science and Applied Nutrition stated that all food products are responsible for only 5% of total exposure to benzene. Other studies which arrive at similar conclusions are: UK-MAFF Food Surveillance No 58 - Benzene and other Aromatic Hydrocarbons in Food-Average UK Dietary Intakes -March 1995; EU Joint Research Centre, HEXPOC, 2005 - Human Exposure Characterisation of chemical substances; quantification of exposure routes; Canada: B.D. Page et al – Journal of AOAC Intl., 1992, 75, (2) 334-340.

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In spite of this, the industry has taken a responsible approach to minimize the presence of benzene in its beverages. Today, as the beverage industry continues to grow and expand, the International Council of Beverages Associations (ICBA) is renewing its commitment to provide guidance on minimizing benzene formation. This guidance will be made available to *all* beverage companies worldwide, regardless of their affiliation with the ICBA. Likewise, the Council will make a concerted effort, through its Member Associations, to provide this information to all beverage producer companies.

### 3. Trigger and Mitigating Factors for Benzene Formation in Beverages

### 3.1 Trigger factors which may lead to the formation of benzene in beverages

\* *Primary Driver:* Benzene formation may occurs at part per billion (microgrammes per kilo) levels in some beverage formulations containing sodium benzoate or potassium benzoate along with ascorbic acid.(<sup>1</sup>) Levels increase with heat and/or light, with heat being the predominant factor.

\* Some studies suggest that erythorbic acid – where permitted - may lead to benzene formation in much the same way as ascorbic acid.

\* Benzene formation may also occur when juices and other ingredients - which naturally, or otherwise - where permitted - contain benzoic acid sources and ascorbic acid - are used in beverage formulations.

# **3.2** Mitigating factors which may mitigate the formation of benzene in beverages containing benzoic acid sources and ascorbic acid

\* Ingredients, such as nutritive sweeteners (sugar, high fructose corn or starch syrup) and calcium disodium ethylenediamenetetraacetic acid (EDTA) - where permitted - or sodium poly (or hexameta) phosphate, may mitigate benzene formation.

\* Evidence indicates that nutritive sweeteners delay the reaction, as the phenomenon seems most noticeable in diet beverages, however the longer a product is in the market (shelf-life), the greater the potential for benzene formation if its precursors are present.

\* Evidence also suggests that EDTA – where permitted – may mitigates the reaction, possibly by complexing metal ions that may act as catalysts. The degree of mitigation may be lessened in products containing calcium or other minerals – especially when used as fortificants - as they may interfere with the mitigating action.

<sup>&</sup>lt;sup>1</sup> L.K. Gardner and G.D. Lawrence, J. Agric. Fd. Chem. 1993, <u>41</u> (5), 693-695

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### 4. <u>ICBA key recommendations to beverage producers to eliminate/minimise benzene</u> <u>formation</u>

Taking into account trigger and mitigating factors for benzene formation in beverages as set out above (section 3), ICBA recommends the following:

### ✓ RECOMMENDATION 1: REVIEW

All beverage companies **review their existing products and new formulations** considering the above information relative to procedures for the prevention / minimization of benzene formation.

### ✓ RECOMMENDATION 2: TEST

All beverage companies **perform analytical sampling of appropriate products** for benzene through accelerated storage tests (*for more detailed guidance on testing, please see section 5 below*).

### ✓ RECOMMENDATION 3: REFORMULATE

Beverage companies **reformulate any affected products** in which benzene may be present to eliminate or reduce benzene formation to the fullest extent possible.

### ✓ RECOMMENDATION 4: MONITOR POST-LAUNCH

Beverage companies include benzene measurement in trade sample analyses.

### 5. <u>Guidance: Testing for the presence of benzene in beverages</u>

### 5.1 Accelerated tests

Accelerated tests should be conducted for product formulations containing benzoic acid sources - including added benzoate - and ascorbic acid. Specific test conditions may vary from producer to producer but should encompass conditions of time and temperature that would cover the normal distribution conditions that the product will experience. As a starting point, producers may want to consider subjecting the product formulations to temperatures of a minimum of 50-70 degrees C for 24 hours, or longer depending on the formulation, eg some product formulations require 14 days of accelerated test exposure to evaluate the reaction potential.

### 5.2 Analytical procedures

Reliable analytical procedures for benzene should be validated through appropriate performance trials or accredited external laboratories, capable of determining <u>at least</u> 5 ppb (microgrammes/kg) of benzene in beverages. (See Annex for examples of methodologies).

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### 6. <u>Guidance: Formulation Control Strategies</u>

As previously noted, the main factors in benzene formation in beverages are generally a combination of benzoic acid sources and ascorbic acid, heat and time. However, other control points (CP) that beverage developers may wish to consider when formulating a product also include:

Product Water

 $\rightarrow$  must meet local regulatory requirements, including benzene levels, for potable water. *In addition, see section below on 'Transition Metals'*. <u>CP</u> – check benzene in water

- ♦ <u>Sugars (Nutritive sweeteners)</u>
  → appears to slow benzene formation, but does not totally inhibit it.
- ♦ <u>Fruit Juices</u>

 $\rightarrow$  can be delivered 'preserved' with benzoate - where permitted - and/or other natural benzoic acid sources

<u>CP</u> – review specifications with supplier to control or eliminate benzoate

 $\rightarrow$  may be a source of ascorbic acid (added or natural) <u>CP</u> – analyse for ascorbate or obtain levels from supplier

• Intense Sweeteners

 $\rightarrow$  Diet / Light products have greatest potential for benzene formation if precursors are present.

<u>Carbon Dioxide</u>

 $\rightarrow$  ensure compliance with local regulatory requirements or International Society of Beverage Technologists (ISBT) standard of 20 ppb (microgrammes/kg) maximum of benzene

<u>CP</u> – supplier specifications and analyses with checks

♦ <u>Acids</u>

 $\rightarrow$  At low pH, ascorbic acid and/or erythorbic acid, in combination with benzoic acid sources, leads to a higher potential of the formation of benzene

• Flavours/Clouding Agents

 $\rightarrow$  Flavours, emulsions and cloudifiers may contain preservatives and antioxidants <u>CP</u> – review specifications with supplier to control or eliminate benzoate

 $\rightarrow$  Benzaldehyde and ascorbic acid can also form benzene

CP - check if benzaldehyde present

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### ♦ <u>Colours</u>

 $\rightarrow$  may contain ascorbate as an antioxidant to prevent fading <u>CP</u> – check with suppliers and re-specify if necessary

### • <u>Preservatives</u>

All manufacture of beverages should take place under strict hygienic conditions, following HACCP principles

 $\rightarrow$  Consider the use of blends of sorbate and benzoate, if there is a technological need (microbiological stability or sorbate solubility).

 $\underline{CP}$  - Consider if benzoate can be removed/reduced/replaced by sorbate or other preservation systems. Note that sorbate may precipitate out in dilutable and post-mix syrups (fountains)

 $\rightarrow$  Dilutables: (Squashes, Cordials – usually 5x concentrated, diluted before consumption) require to be preserved because of their frequent openings during their long shelf-lives.

 $\underline{CP}$  - Consider if benzoate can be removed/reduced/replaced by sorbate or other preservation systems.

Note that sorbate must be used with care to avoid it precipitating. Consider using alternatives to ascorbate, if present.

♦ <u>Antioxidants</u>

 $\rightarrow$  consider the use of ascorbate in relation to overall formula, especially if citrus juices or other natural carriers of ascorbate are present.

<u>CP</u> – Remove/reduce/replace ascorbate as appropriate if a benzoic acid source is present

♦ <u>Light</u>

 $\rightarrow$  UV light may induce free radical formation in products <u>CP</u> – Review storage and shelf-life conditions, and labelling instructions

• <u>Temperature</u>

 $\rightarrow$  accelerates the formation if precursors are present

<u>CP</u> – Review storage and shelf-life conditions, and labelling instructions

<u>Transition Metals</u>

 $\rightarrow$  trace levels of metal ions, such as copper and iron, may act as catalysts in benzene formation in beverages in the presence of benzoic acid sources and ascorbic acid. Sources of transition metals may include product water, sweeteners or other ingredients.

<u>CP</u>- Chelating compounds such as EDTA (where permitted) or sodium polyphosphates may help mitigate formation. Fortification by calcium, or other minerals, may lessen this effect.

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## ANNEX: EXAMPLES OF METHODS OF ANALYSIS

- 1. <u>Purge/Trap</u> GCMS Quantitation of Trace Level Benzene in Carbonated Soft Drinks and Juice Products
- 2. Determination of Benzene in Carbonated and Non-Carbonated Beverages Gas Chromatography Method using <u>Headspace</u>

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### 1. <u>Purge/Trap GCMS Quantitation of Trace Level Benzene in Carbonated</u> <u>Soft Drinks and Juice Products</u>

### Summary:

### A. Instrument Operation Parameters

Three purge and trap GC/MS instruments were used for the analyses:

1. Agilent 6890/5973 #1 GC/MS, Tekmar Velocity purge and trap concentrator and Tekmar Solatek purge and trap autosampler.

2 Agilent 6890/5973 #2 GC/MS, EST Encon purge and trap concentrator and EST Archon 5100 purge and trap autosampler.

3 Agilent 6890/5973 #3 GC/MS, Tekmar Velocity purge and trap concentrator and Archon (OI 4552) autosampler.

1.  $\underline{6890/5973 \#1}$  A 30 m x 0.25 mm x 0.25 µm HP-5MS column (Agilent 19091S-433) was held at 45 °C for 2 minutes, than increased at 10 °C/minute to 65 °C, increased at 25 °C to 250 °C and held at 250 °C for 5 minutes. The injection port was set at 225 °C with a 50:1 split ratio and a 4 mm ID split/splitless liner. Carrier gas regulated by the 6890's EPC is supplied to the Velocity. It is returned to the 6890 via the heated transfer line which is plumbed into the GC inlet.

The column was operated in the constant pressure mode at 12.33 psig with a nominal flow rate of 0.8 mL/minute. The gas saver was set to 20 mL/minute after 2 minutes run time. The column terminated in an Agilent microfluidics splitter which diverted approximately 20% of the column effluent to an Agilent  $\mu$ -ECD. The ECD data was not collected for this analysis. The splitter had a 3.8 psig make up gas supply, making the column output 3.8 psig above ambient pressure.

The MSD transfer line was set at 280 °C, the MS Quad was at 150 °C and the MS Source was at 230 °C. The electron multiplier offset was 106 volts and the multiplier voltage was 1694 volts. A tune file of ATUNE was used. Selected-Ion-Monitoring was employed for the analysis. Ions monitored were m/z 77, 78 for benzene and 82, 84 for d<sup>6</sup>-benzene internal standard in low resolution mode and the dwell time for these ions was 100 ms.

Benzene, target ion m/z 78, qualifier ion 77, and  $d^6$ -benzene, target ion m/z 84, qualifier ion 82, were monitored from 3.23 to 3.65 minutes. Retention times are benzene, 3.45 minutes, and  $d^6$ -benzene, 3.43 minutes.

A Tekmar Velocity XPT purge and trap concentrator with the Solatek autosampler was used in the soils mode. In this mode samples are purged in the 40-mL sample vial (I-Chem Certified 200 series, Fisher Scientific 05-719-102) by the autosampler. Purge gas is carried to the head of the concentrator trap via a heated SilcoSteel transfer line.

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A Supelco type J, or BTEX, trap (Supelco 21064) was used. This is a 25 cm long 1/8" stainless steel tube packed with 7.7 cm of Carbopak C and 1.2 cm of Carbopak B. The analytical conditions for the purge and trap concentrator/autosampler are as follows. These analytical conditions are set up in Tekmar's Teklink software which operates the sampler and starts the GC/MS run when the desorption cycle begins.

Variable	Value	Variable	Value
Rinse Water Temp	90 °C	Pre-Purge Flow	40 mL/min
Sample Cup Temp	40 °C	Sample Heater	Off
Sample Needle Temp	50 °C	Sample Preheat Time	1.00 min
Transfer Line Temp	150 °C	Preheat Temp	40 °C
Soil Valve Temp	125 °C	Purge Temp	0 °C (a default setting)
Sample Sweep Time	0.50 min	Purge Flow	40 mL/min
Needle Rinse Volume	15 mL	Dry Purge Time	0.50 min
Needle Sweep Time	1.00 min	Dry Purge Temp	20 °C
Sample Preheat Time	1.00 min	Dry Purge Flow	50 mL/min
Preheat Stir	Off	GC Start	Start of Desorb
Preheat Stir Mode	Spin	Desorb Preheat Temp	250 °C
Preheat Stir Speed	5	Desorb Drain	On
Purge Time	11.00 min	Desorb Time	2.00 min
Purge Stir	Off	Desorb Temp	250 °C
Purge Stir Mode	Agitate	Desorb Flow	0 mL/min (a default
			setting)
Purge Stir Speed	5	Bake Time	4 min
Valve Oven Temp	150 °C	Bake Temp	260 °C
Transfer Line Temp	150 °C	Dry Flow Bake Temp	300 °C
Sample Mount Temp	90 °C	Bake Flow	400 ml/min
Purge Ready Temp	35 °C	Focus Temp	-150 °C
Dry Flow Standby	200 °C	Inject Time	1.00 min
Temp			
Standby Flow	20 mL/min	Inject Temp	180 °C
Pre-Purge Time	0.50 min	Standby Temp	100 °C

Table 1. Tekmar Velocity and Solatek Settings

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### 2. <u>6890/5973 #2</u>

A 60 m x 0.25 mm x 1.4  $\mu$ m DB-624 column (Agilent 122-1364) was held at 40 °C for 2 minutes, than increased at 10 °C/minute to 180 °C, than increased at 40 °C/minute to 250 °C, and held at 250 °C for 2 minutes. The injection port was set at 200 °C with a 40:1 split ratio. Carrier gas regulated by the 6890's EPC is supplied to the Encon. It is returned to the 6890 via the heated transfer line which is plumbed into the GC inlet.

Some problems were experienced with ghost peaks/sample carryover due to the length of the column. This instrument is configured for another analysis which requires that column. It is recommended to use no longer than a 30 m thin film column for benzene analysis. The column was operated in the constant flow mode at 16.65 psig initial pressure with a flow rate of 1.0 mL/minute. The gas saver was set to 20 mL/minute after 2 minutes run time.

The MSD transfer line was set at 280 °C, the MS Quad was at 150 °C and the MS Source was at 230 °C. The electron multiplier offset was off and the multiplier voltage was 1670 volts. It used a tune file of ATUNE. Solvent delay was 10 minutes.

Selected-Ion-Monitoring was used for the analysis. Ions monitored were m/z 77, 78 for benzene and 84, 82 for d<sup>6</sup>-benzene internal standard. The dwell time for these ions was 85 ms.

Benzene, Rt 11.3 minutes, was detected using target ion m/z 78, qualifier ion 77.  $d^6$ -benzene internal standard was detected using target ion 84 and qualifier ion 82. The mass spectrometer was operated with high resolution.

The Encon purge and trap concentrator with the Archon 5100 autosampler was used in the soils mode. Purge gas is carried to the head of the concentrator trap via a heated SilcoSteel transfer line. The Encon used a Type K, or Vocarb 3000 trap (PTS Catalog E70300-K03). It is a 25 cm long 1/8" stainless steel tube packed with 10 cm Carbopak B, 6 cm Carboxen 1000 and 1 cm Carboxen 1001.

Variable	Value	Variable	Value
Standby Flow	On	Drain	On
Bake Gas Bypass	On	Antifoam	Cont
Total GC Time	0 Minutes	Valve Oven	130 °C
Transfer Line	130 °C	MoRT Ready	40 °C
MoRT Bake	260 °C	Purge Ready	35 °C
Purge Time	11 Minutes	Dry Purge Time	1 Minute
Purge Flow	40 mL/Minute	Desorb Flow	0 mL/Min (a default
			setting)
Preheat Temp	40 °C	Preheat Time	0.5 Minute
Desorb Preheat	245 °C	Desorb Temp	250 °C
Desorb Time	2 Minute	Bake	260 °C
Bake Time	10 Minutes		

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Table 2. Encon and Archon Settings

### 3. <u>6890/5973 #3</u>

A 20 m x 0.18 mm x 1  $\mu$ m DB-624 column (Agilent 122-1324) was held at 40 °C for 3 minutes, than increased at 15 °C/minute to 210 °C and held at 210 °C for 0.33 minutes. The injection port was set at 210 °C with a 50:1 split ratio. Carrier gas regulated by the 6890's EPC is supplied to the Encon. It is returned to the 6890 via the heated transfer line which is plumbed into the GC inlet.

The column was operated in the constant flow mode at 16.53 psig initial pressure with a flow rate of 0.8 mL/minute. The gas saver was off.

The MSD transfer line was set at 190 °C, the MS Quad was at 150 °C and the MS Source was at 230 °C. The electron multiplier offset was off and the multiplier voltage was 1176 volts. It used a tune file of ATUNE. Solvent delay was 4.5 minutes.

Selected Ion Monitoring was used for the analysis. Ions monitored were m/z 77, 78 for benzene and 84, 82 for d<sup>6</sup>-benzene internal standard. The MSD was set to low resolution and the dwell time for these ions was 85 ms.

Benzene, Rt 5.19 minutes, was detected and amount was quantitated using target ion m/z 78.  $d^6$ -benzene, Rt 5.16 minutes, was detected using target ion m/z 84.

The Velocity purge and trap concentrator with the Archon autosampler was used in the soil mode. Purge gas is carried to the head of the concentrator trap via a heated SilcoSteel transfer line. The Velocity used a Type K, or Vocarb 3000 trap. It is a 25 cm long 1/8" stainless steel tube packed with 10 cm Carbopak B, 6 cm Carboxen 1000 and 1 cm Carboxen 1001.

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X7 · 11	X7 1	X7 · 11	X7 1
Variable	Value	Variable	Value
Valve Oven Temp	150	Dry Purge Temp	40
Transfer Line	150	Dry Purge Flow	200 mL/Min
Temp			
Sample Mount	90	GC Start	Start of Desorb
Temp			
Purge Ready Temp	40	Desorb Preheat	245
		Temp	
Dry Flow Standby	175	Desorb Drain	ON
Temp			
Standby Flow	0 mL/min	Desorb Time	2 min
Pre-Purge Time	0.5 min	Desorb Temp	250
Pre-Purge Flow	40 mL/min	Desorb Flow	0 mL/min (a default
C C			setting)
Sample Heater	OFF	Bake Time	2 min
Sample Preheat	1 min	Bake Temp	270
Time		L.	
Preheat Temp	40	DryFlow Bake	300
1		Temp	
Purge Time	11 min	Bake Flow	400 mL/min
Purge Temp	0 (a default	Focus Temp	-150
	setting)	1	
Purge Flow	40 mL/min	Inject Time	1 min
Dry Purge Time	1 min	Inject Temp	180
Sample Type	Soil	Sample Vol	5 mL
Dilution Factor	0	Rinse Vol	0 (a default setting)
# Rinses	0	Standard 1	YES
Standard 2	NO	Sample Preheat Stir	NO
Stir	NO	W. Stir Time	0 (a default setting)
W. Settle Time	0	Syringe Flushes	0 (a default setting)
Opr. Mode	Remote	Cycle Timer	0 (a default setting)
Aux Timer	0	Link To Method #	#0

Table 3. Velocity and Archon Settings

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### **B.** Instrument Calibration

### **Reagents and standards (d<sup>6</sup> benzene ISTD)**

The calibration information is for GC/MS #1 with the Tekmar Velocity and Solatek.

0.9251 g of 5.023 mg/mL benzene in methanol was weighed into a 50 mL volumetric flask and filled to the mark with methanol (117,448 µg/L benzene). This solution was serially diluted 1:10 using 5 mL Class A glass pipettes and 50 mL volumetric flasks to make solutions of 11,745 µg/L benzene, 1,174 µg/L benzene and 117 µg/L benzene. Aliquots of these solutions were added to 10 mL of purged, deionized water in 40 mL EPA vials according to Table 5 to make calibration standards.

Hamilton 800 Series microliter syringes were used to add the calibration standards to the EPA vials containing water or sample. These syringes are listed in Table 4.

Syringe Volume	Model Number	Fisher Catalog Number
10 µ	801RNW	14-815-300
25 μL	802RNW	14-815-301
50 µL	805RNW	14-815-302
100 µL	810RNW	14-815-303

Table 4. Syringes Used for Addition of Standard Solutions to Vials

0.9712 g of 2.015 mg/mL d<sup>6</sup>-benzene in methanol was weighed into a 50 mL volumetric flask and filled to the mark with methanol (49,460  $\mu$ g/L d<sup>6</sup>-benzene). This solution was diluted 1:5 with a 20 mL Class A glass pipet and a 100 mL volumetric flask to make a solution of 9892  $\mu$ g/L d<sup>6</sup>-benzene for use as internal standard. The purge and trap autosampler method was instructed to inject 5  $\mu$ L of the internal standard to each EPA vial prior to purging.

This d<sup>6</sup>-benzene solution was only used for GC/MS #1. The Solatek autosampler was able to inject 5  $\mu$ L of internal standard solution. The other (Archon) autosamplers used for these analyses were capable of injecting 1 and 2  $\mu$ L aliquots, respectively, of internal standard solution and required d<sup>6</sup>-benzene solutions of different concentrations. The Archons did wash the ISTD into the sample with 10 mL of rinse water.

One of the standards reservoir vessels on the Solatek was filled with a portion of the internal standard solution. Prior to purging the sample, an injection valve loaded 5  $\mu$ L of the d<sup>6</sup>-benzene solution which was than added to the EPA vial with 10 mL of rinse water.

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	:	Standard			Standar	b		Standard	
μL of 117 μg/L		Conc	Cal	μL of 1,174 μg/L	Conc	Cal	μL of 11,745 μg/L	Conc	Cal
Benzene		µg/L	Level	Benzene	μg/L	Level	Benzene	µg/L	Level
	10	0.117	1	10	1.	17	4 10	) 11.74	. 7
	20	0.235	2	20	2.	35	5		
	40	0.470	3	40	4.	70	6		
μL of 9892 μg/L d <sup>6</sup> -Benzene	-								
	5	4.90	All						

Table 5. Standards Preparation for 10 mL Calibration Standards and Samples

In addition to these calibration standards 5 and 50  $\mu$ L of the 1,174  $\mu$ g/L benzene solution were added to 10 mL aliquots of purged, deionized water. These are used to make calibration verification samples at 0.59 and 5.87  $\mu$ g/L benzene.

Finally a check sample was prepared from an independent benzene source to verify that the instrument gave consistent data. A 100  $\mu$ L aliquot of a solution containing 2000  $\mu$ g/mL benzene in methanol was added to 100 mL of purged, deionized water.

A 5  $\mu$ L aliquot of the 2000 $\mu$ g/L benzene solution was added to 10 mL purged, deionized water in an EPA vial giving 1.0  $\mu$ g/L benzene. This check standard was treated as an unknown sample for calibration verification and to ensure that all the instruments used for the project gave consistent results.

The Agilent 6890/5973 #1 GC/MS, Tekmar Velocity purge and trap concentrator and Tekmar Solatek purge and trap autosampler was calibrated with these standards. The ChemStation was calibrated using a linear fit of response ratio to amount ratio. The ChemStation was "told" that the d<sup>6</sup>-benzene internal standard concentration was 1.0  $\mu$ g/L for all samples. This caused the Amount Ratio for each calibration level to display as being the same as the benzene concentration for that level.

A list of the purchased calibration standards is included. All are 1 mL of solution in a sealed ampoule.

- 1- Benzene standard: Supelco 40004, 5023 µg/mL benzene in methanol
- 2- d<sup>6</sup>-benzene standard: Supelco 48940-U, 2015 µg/mL benzene in methanol
- 3- Check sample benzene standard: Restek 30249, 2000 µg/mL benzene

### **Instrument Calibration**

Calibration and verification samples are made by adding microliter aliquots of the standards from Table 5 to 10 mL of purged, deionized water in 40 mL EPA vials. Internal standard solution aliquots of 5  $\mu$ L are added by the Solatek. For other instruments the concentration of

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the d<sup>6</sup>-benzene standard will have to be altered to accommodate the size of the aliquot of ISTD solution the autosampler is capable of injecting into the samples. Table 5 will still be applicable for dilution of the benzene calibration standards.

Table 6 shows the calibration data for the 6890/5973/Velocity/Solatek purge and trap GC/MS. Recall that the actual d<sup>6</sup>-benzene concentration is 4.9  $\mu$ g/L and the ChemStation was told it was 1  $\mu$ g/L to adjust the amount ratio to equal the benzene concentration.

The first and last samples in the calibration runs were purged, deionized water used to check the background of the instrument. A typical sequence in these analyses starts with a water blank, a  $5.87 \mu g/L$  benzene verification sample and than the unknown samples with either a 0.59 or a  $5.87 \mu g/L$  verification sample between every 8 - 10 unknown samples. Figure 1 is the ChemStation calibration chart.

Sample Name	Benzene	Benzene	d6 Benzene	d6 Benzene	Benzene
	Area	Conc µg/L	Area	Area	Found µg/L
0 μg/L	10942	0.00	1	818157	0.0
0.12 μg/L	32807	0.12	1	817882	0.1
0.24 μg/L	57300	0.24	1	827709	0.3
0.47 μg/L	101968	0.47	1	828011	0.5
1.17 μg/L	221872	1.17	1	818283	1.2
2.35 μg/L	419894	2.35	1	818604	2.2
4.70 μg/L	906480	4.70	1	831742	4.8
11.74 μg/L	2192056	11.74	1	835169	11.7
0 μg/L	17214	0.00	1	825647	0.0
0.59 µg/L Check Sample	123141	0.59	1	833539	0.6
5.87 µg/L Check Sample	1153599	5.87	1	837021	6.1

Following chart in Excel using the ChemStation data was created.

Table 6. Calibration Data for 0.12 to 11.74 µg/L Benzene Standards

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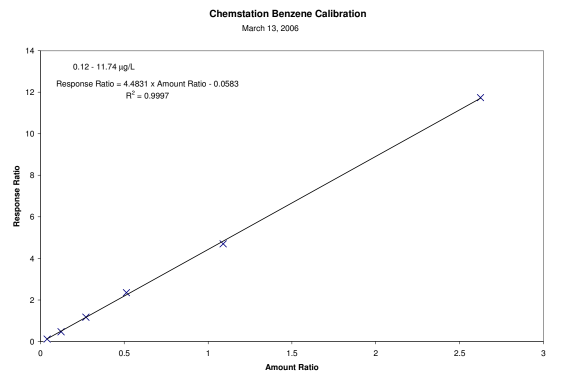


Figure 1. ChemStation calibration curve for 0.12 to  $11.74 \mu g/L$  benzene standards

The manufacturer of the d<sup>6</sup>-benzene standard solution claims 99.9 % purity. The 4.9  $\mu$ g/L d<sup>6</sup>-benzene internal standard could contain as much as 0.004  $\mu$ g/L of benzene. In addition the mass spectrum of d<sup>6</sup>-benzene in the NIST library has a small amount of m/z 78 and 77.

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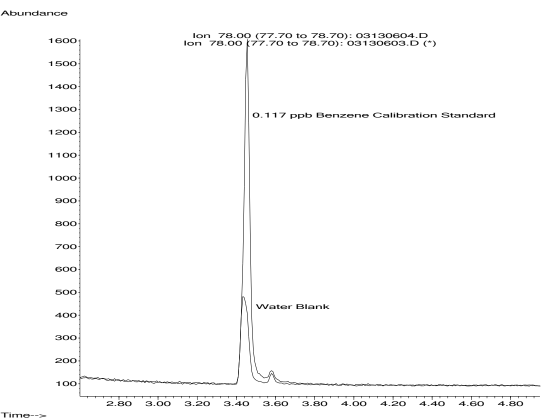


Figure 2. Superimposed ion chromatograms of blank run and 0.12 µg/L benzene

#### **Sample Preparation**

Ten mL sample aliquots (plus 10 mL rinse water) are purged directly in the 40-mL EPA vials. The samples are pipetted into the vials and the vials capped for analysis.

Carbonated beverages (CSDs) tend to experience  $CO_2$  breakout when opened and during sampling by pipet. This is unavoidable and a certain (unknown) of loss of volatiles does occur. The CO<sub>2</sub> breakout can be minimized by cooling the CSDs in a refrigerator overnight or in bucket of ice for several hours before opening the container. Avoid agitating the container before opening. Pre-wetting the interior of the pipet with an aliquot of sample and disposing of it will help to minimize losses due to this bubbling. Fill and drain the pipet slowly to minimize  $CO_2$ escape. Allow the sample to run down the interior wall of the 40-mL flask to reduce  $CO_2$  and volatiles losses.

#### Remark:

Juices containing pulp and other high solids cannot be analyzed using this purge and trap method since they plug sample lines and valves.

These samples have to be analyzed by head-space – GC/MS.

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# 2. <u>Determination of benzene in carbonated and non-carbonated beverages – gas chromatography method using headspace</u>

### **1. SCOPE AND PURPOSE**

This method specifies a gas chromatography method using headspace for the determination of benzene in carbonated and non-carbonated beverages.

### 2. RESPONSIBILITIES

2.1. Trained lab associates are responsible for preparing the reagents and for conducting the analyses. They are also responsible for recording the results of the samples and check samples properly in the proper raw data forms. They need to evaluate the obtained results critically and apply the means necessary to control themselves by calibrations and checks.

2.2. The Laboratory Operation Manager is responsible for the effectiveness of the procedure. He/she also provides expertise assistance in evaluating methods and results.

2.3. The Laboratory Manager is responsible for making sure the laboratory has the necessary resources(equipment and personnel) to perform the tests, checking and calibration activities properly.

### **3. PRINCIPLE**

The sample is being heated in a closed headspace vial, in order to obtain equilibrium between the concentration of the benzene in the headspace above the sample and the concentration in the sample. Prior to analysis, carbonated samples are being treated with sodium hydroxide in order to neutralize the carbon dioxide. In order to increase the headspace efficiency of benzene, a matrix modifier (sodium chloride) is being added to the vial. After establishment of the equilibrium, an aliquot of the gaseous phase is injected on a gas chromatograph. In order to obtain higher sensitivity the benzene is trapped on a cold trap prior to injection on the capillary column. The components are detected by mass spectrometry detector (MS) in full scan mode. Identification is based on retention time and library mass spectra comparison. Quantification is based on the intensity of the main ion using a four-point calibration and the internal standard method.

### 4. EQUIPMENT AND GLASSWARE

- 4.1. Appropriate laboratory equipment
- 4.2. Volumetric flasks with glass stoppers

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4.3. 20 ml headspace vials with magnetic crimp caps

4.4. Finnigan Trace GC- DSQ- mass spectrometer with Combipal auto sampler with headspace configuration (GC210 or 212)

4.5. Capillary column: Restek RTX-1 60m \* 1µm df \* 0.25 mm ID

4.6. Ultra pure water generator: Millipore MilliQ (PW202 or 203)

4.7. Graduated Hamilton syringes (10, 25 and 50  $\mu$ l)

4.8. Hamilton digital syringe 25  $\mu$ l

4.9. Analytical balance (AB201, 202 or 203)

### **5. REAGENTS AND SOLUTIONS**

### **5.1. REAGENTS**

- 5.1.1. Ultra pure water cooled at 4 °C
- 5.1.2. Benzene (CHBEN92)
- 5.1.3. Benzene-d6 (CHBEN91)
- 5.1.4. Methanol, purge and trap grade (CHMET05)
- 5.1.5. Sodium hydroxide (CHSOH04)
- 5.1.6. Sodium chloride (CHSOC05)

### **5.2. SOLUTIONS**

Only use volumetric flasks with glass stoppers (4.2)

### 5.2.1. Benzene stock solution 1000 mg/l (solution code SL-045-01)

Weigh a dry, empty volumetric flask of 50 ml with stopper to an accuracy of 0.1 mg. (**m1**) on the analytical balance (AB201). Place 25 ml methanol (5.1.4) in the volumetric flask, place the flask on the analytical balance and tare. Using a pasteur pipette, add +/- 50 mg of benzene (5.1.2) and determine the exact weight. (**m2**). Make sure the benzene is directly added to the methanol (no contact with the inside surface). Make up to the mark with methanol.

Determine the weight of the flask with the solution with stopper to an accuracy of 0.1 mg (m3). The exact concentration of the working standard is calculated as follows:

C1 = (m2/((m3-m1)/0.7915)) \* 1000 (mg/l)

Where

C1 is the concentration of the benzene stock solution (mg/l)

m1 is the mass of the empty flask (g)

m2 is the mass of the benzene (mg)

m3 is the mass of the full flask (g)

0.7915 is the density of methanol  $(20^{\circ}C)$ 

This solution is stable for three months if stored at -18  $^{\circ}$ C in the dark. Store the solution in an amber glass

### 5.2.2. Benzene control stock solution 1000 mg/l (solution code SL-045-02)

Prepare the independent benzene control stock solution 1000 mg/l using the same procedure as described in 5.2.1. This solution is stable for three months if stored at -18 °C in the dark. Store the solution in an amber glass flask.

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### 5.2.3. Benzene working spike solution 2 ppm (solution code SL-045-03)

Weigh a dry empty volumetric flask of 50 ml with stopper to an accuracy of 0.1 mg (**m4**). Place +/- 45 ml of methanol in the volumetric flask. Using a 0.25 ml syringe add 0.1 ml of the benzene stock solution 1000 mg/l (5.2.1). Record the weight of the added solution (**m5**). Make up to the mark with methanol and determine the weight of the full flask with stopper. (**m6**). The exact concentration of the working spike solution is calculated as follows:

The exact concentration of the working spike solution is calculated as follows:

C2 = ((m5)/(m6-m4)) \* C1 (mg/l)

### Where

C2 is the concentration of the benzene working spike solution

m4 is the mass of the empty flask (g)

m5 is the mass of the added benzene control stock solution 1000 mg/l (mg)

m6 is the mass of the full flask (g)

C1 is the concentration of the benzene stock solution 1000 mg/l (5.2.1)

This solution is stable for one month if stored at -18 °C in the dark. Store the solution in an amber glass flask.

### 5.2.4. Benzene control spike solution 2 ppm (solution code SL-045-04)

Weigh a dry empty volumetric flask of 50 ml with stopper to an accuracy of 0.1 mg (**m7**). Place +/- 45 ml of methanol in the volumetric flask. Using a 0.25 ml syringe add 0.1 ml of the benzene control stock solution 1000 mg/l (5.2.2). Record the weight of the added solution (**m8**). Make up to the mark with methanol and determine the weight of the full flask with stopper. (**m9**). The quart concentration of the working spike solution is calculated as follows:

The exact concentration of the working spike solution is calculated as follows:

C3 = ((m8)/(m9-m7)) \* Ccontr (mg/l)

Where

C3 is the concentration of the benzene control spike solution

m7 is the mass of the empty flask (g)

m8 is the mass of the added benzene control stock solution 1000 mg/l (mg)

m9 is the mass of the full flask (g)

Ccontr is the concentration of the benzene control stock solution 1000 mg/l 5.2.2.

This solution is stable for one month if stored at -18 °C in the dark. Store the solution in an amber glass flask.

### 5.2.5. Internal standard stock solution 1000 ppm (solution code SL-045-05)

Weigh a dry, empty volumetric flask of 50 ml with stopper to an accuracy of 0.1 mg. (**m10**) on the analytical balance. Place 25 ml methanol (5.1.5) in the volumetric flask.

Weigh  $\pm$  50 mg of benzene-d6 (5.1.3) into the flask and note the exact weight (**m11**). Make up to the mark with methanol and close. Make sure the benzene is directly added to the methanol (no contact with the inside surface).

Determine the weight of the flask with the solution with stopper to an accuracy of 0.1 mg (**m12**). The exact concentration for each internal standard is calculated as follows:

C4 = (m10/((m11-m9)/0.7915))\*1000 (mg/l)

Where

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C4 is the concentration of the internal standard stock solution m9 is the mass of the empty volumetric flask with stopper (g) m11 is the mass of the volumetric flask with solution and stopper (g) m10 is the mass of the internal standard (mg) 0.7915 is the density of methanol ( $20^{\circ}$ C) This solution is stable for six months if stored cooled in the dark. Store the solution in an amber glass flask at – 18 °C in the dark.

### 5.2.6. Internal standard spike solution 1 ppm (solution code SL-045-06)

Place  $\pm 40$  ml of methanol (5.1.5) into a 50 ml volumetric flask. Inject 50 µl of the internal standard stock solution (5.2.5) using a 0.25 ml syringe under the surface of the methanol. Make up to the mark with methanol. This solution is stable for one month if stored cooled in the dark. Store the solution in an amber glass flask at -18 °C in the dark.

### 5.2.7. VOC calibration standards (0.4 – 4 ppb) and control standard 1 ppb

All the calibration standards have to be prepared with cooled ultra pure water (5.1.1) Prepare the calibration standards as follows:

**5.2.7.1. 0.4 ppb**: Weigh +/- 3.5 gr of NaCl (5.1.6) into a 20 ml headspace vial. Pipette 10 ml of cooled ultra pure water. Add 10  $\mu$ l of the internal standard spike solution (5.2.6) using a 25  $\mu$ l digital syringe (4.8). Add 2  $\mu$ l of benzene working spike solution 2 ppm (5.2.3) using a 10  $\mu$ l syringe. Close the vial immediately with a magnetic crimp cap.

**5.2.7.2. 1 ppb**: Weigh +/- 3.5 gr of NaCl (5.1.6) into a 20 ml headspace vial. Pipette 10 ml of cooled ultra pure water. Add 10  $\mu$ l of the internal standard spike solution (5.2.6) using a 25  $\mu$ l digital syringe (4.8). Add 5  $\mu$ l of benzene working spike solution 2 ppm (5.2.3) using a 25  $\mu$ l syringe. Close the vial immediately with a magnetic crimp cap.

**5.2.7.3. 2 ppb:** Weigh +/- 3.5 gr of NaCl (5.1.6) into a 20 ml headspace vial. Pipette 10 ml of cooled ultra pure water. Add 10  $\mu$ l of the internal standard spike solution (5.2.6) using a 25  $\mu$ l digital syringe (4.8). Add 10  $\mu$ l of benzene working spike solution 2 ppm (5.2.3) using a 25  $\mu$ l syringe. Close the vial immediately with a magnetic crimp cap.

**5.2.7.4. 4 ppb:** Weigh +/- 3.5 gr of NaCl (5.1.6) into a 20 ml headspace vial. Pipette 10 ml of cooled ultra pure water. Add 10  $\mu$ l of the internal standard spike solution (5.2.6) using a 25  $\mu$ l digital syringe (4.8). Add 20  $\mu$ l of benzene working spike solution 2 ppm (5.2.3) using a 25  $\mu$ l syringe. Close the vial immediately with a magnetic crimp cap.

**5.2.7.5.** Contr 1 ppb: Weigh +/- 3.5 gr of NaCl (5.1.6) into a 20 ml headspace vial. Pipette 10 ml of cooled ultra pure water. Add 10  $\mu$ l of the internal standard spike solution (5.2.6) using a 25  $\mu$ l digital syringe (4.8). Add 5  $\mu$ l of benzene control spike solution 2 ppm

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(5.2.4) using a 25 µl syringe. Close the vial immediately with a magnetic crimp cap.

### 5.2.8. Sodium hydroxide 30 %

Weigh +/- 60 g of sodium hydroxide (5.1.5) into a clean glass beaker and add 200 ml of ultrapure water.

Dissolve the sodium hydroxide and allow cooling to room temperature.

### 6. SAMPLE STORAGE

Samples have to be stored at 4 °C in the dark.

### 7. PROCEDURE

### 7.1. Pretreatment of samples

For carbonated samples add 1 ml of sodium hydroxide 30% to a 40 ml vial and fill the vial to the top with the sample. Close the vial with a PTFE septum containing screwcap. Non carbonated samples are not pretreated.

### 7.2 Sample and blank preparation

Weigh +/- 3.5 g NaCl (5.1.6) into a 20 ml headspace vial. Pipette 10 ml of the sample into the 20 ml headspace vial (4.3). Add 10  $\mu$ l of the internal standard spike solution (5.2.6) using the digital syringe of 25  $\mu$ l. The vial is closed with a magnetic crimp cap with a silicon/PTFE septum. For blank preparation, 10 ml of ultrapure water is treated the same way as a sample.

### 7.2. Preparing the headspace auto sampler.

Place the vials on the tray of the Combipal headspace auto sampler. The vials have to be analyzed in the following order:

blank - calibration standards - blank – control standard 1 ppb – samples – control standard 1 ppb After every 10 samples a control standard of 1 ppb has to be analysed. Indicate in the Cycle Composer software which vials have to be analysed and start the autosampler. Start the cryo-cooling trap once the first thermostatization time has almost expired.

### 7.3. Excalibur software

Fill in the sample table according to the samples to be analysed.

Mark the blanks and samples as unknown and the calibration standards as standard. Use appropriate filenames to save the data. Use the sample name field to describe the sample clearly. Select the correct method for the gas chromatograph. The parameters for the gas chromatograph are described in appendix 2. Save and start the sample table.

### 7.4. Calibration

### 7.5.1. Calibration Frequency

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A full calibration curve has to be analyzed at least once a week. Each day a calibration standard of 1 ppb has to be analyzed.

### 7.5.2. Calibration type

A linear calibration curve using internal standard is used for the calculation. Y=aX+b where Y= relative response factor calculated by As/Ai Where As = area benzene in the sample Ai = area internal standard in the sample X= concentration of benzene in the sample a= slope of the linear regression curve (dY/dX) b= intercept

### 8. EXPRESSION OF RESULTS

### 8.1. Identification and quantification

The identification of the components is based on their retention times and mass spectra library comparison. Table 1 contains the indicative retention times of the components. These retention times slightly vary due to aging of the column and are to be corrected. The quantification of the components is being done automatically by the software using linear regression with internal standard compensation. For carbonated samples, a correction factor of 1,025 is used to calculate the final result (takes into account the addition of NaOH). Subtracting blank values from the sample results is not permitted.

Table 1: Indicative retention times of the components to be quantified

Component name	Retention time	Main ion
Benzene-d6	14,1	84
Benzene	14,17	78

### **8.2. Reporting of results**

Results are expressed as  $\mu g/l$ . Results below 0.5  $\mu g/l$  are to be reported as '<0.5  $\mu g/l$ '.

### 9. REFERENCES

- EPA method 5021A. Volatile organic compounds in various sample matrices using equilibrium headspace analysis.

- EPA method 8260B Volatile organic compounds by gas chromatography/mass spectrometry.

- EPA method 524.2 Measurement of purgeable organic compounds in water by capillary column gas chromatography/mass spectrometry.

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### **Appendix 1: Combipal headspace parameters**

Method name: VOC-25min-benz Incubation temperature: 60 °C Incubation time: 25 min Syringe temperature: 90 °C Agitator speed: 500 rpm Agitator on time: 2 sec Agitator off time: 5 sec Fill speed: 250 µl/sec Injection speed: 15µl/sec GC runtime: 30 min

### **Appendix 2: GC/MS parameters**

Method name: VOC-20-benzene MS parameters: Acquisition time: 16 min Source temperature : 280 °C Scan mode : Full scal Scan rate : 750 amu/s Scan range : 40-250 amu GC parameters: Oven method: Initial temp: 40 °C Initial time: 5 min Rate 1: 5 °C/min Temp 1: 90 °C Hold time 1:0 min Rate 2: 80 °C/min Temp 2: 300 °C Hold time 2: 5 min Injection method: Injection temp: 230 °C Mode: Splitless Splitless time: 6 min Split flow: 25 ml/min Flow method: Mode: Constant flow Flow: 1 ml/min

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### **Appendix 3: Example chromatogram**

