Validation of Styrene using SKC Passive Sampler 575-003



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Contents

Abstract		1		
Importance of	f Validation of Passive Samplers	2		
Summary of M	NIOSH Validation Protocol	4		
Bi-Level Vali	dation	6		
Comments on	the Relationship Between the NIOSH and			
CEN Diffusi	ve Sampler Evaluation Protocols	8		
Scope of the I	Method	10		
Background		11		
Analytical Re	covery	12		
Sampling Rate and Capacity 1				
Reverse Diffusion				
Storage Stabil	lity	15		
Factorial Resu	ılts	16		
Factorial Sum	ımary	17		
Temperature l	Effects	18		
Accuracy and	Precision	19		
Appendix A.	Atmosphere Generation Apparatus	20		
Figure 1.	Atmosphere Generation Apparatus	21		
Figure 2.	Analytical Instrument	22		
Abbreviations	s, Trademarks	23		
Deferences		24		

Research Report

Validation of Styrene using SKC Passive Sampler 575-003

Abstract

A sampling method for Styrene in air has been validated for concentration levels from 5 to 250 ppm and for exposure times from 7.5 minutes to 12 hours. The 575-003 passive sampler used has a sample medium of Anasorb® 727. Desorption was with carbon disulfide and analysis by gas chromatography with flame ionization detection.

The analytical recovery over the range of 5 ppm for 8 hours to 250 ppm for four hours (0.13 to 3.3 mg) was estimated to be 100%. There was no effect of humidity on recovery.

The sampling rate is 13.7 ml/min which was confirmed by the precision and accuracy calculations using 101 results (see Background; Sampling Rate Determination). Samples can be taken up to 40° C.

Minimum recommended sampling time is 15 minutes. Maximum recommended sampling time is 8 hours.

Samples were stable for up to three weeks when stored at room temperature, or in a refrigerator.

A full validation of Styrene was done according to NIOSH Protocol.¹

Two samplers were validated simultaneously. The 575-002 sampler has greater capacity, and can be used where more volatile compounds (e.g. acetone) are present. The 575-003 has the better recovery, and is therefore more useful at low concentrations or for measuring STELs. The 575-003 should not be used to sample compounds with boiling points below that of Styrene.

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Styrene (003)

Importance of Validation of Passive Samplers

There are distinct differences between a passive sampler and a sample tube.

The most important difference is that a passive sampler does not have a foolproof back up section that guarantees that all the chemical hazard has been collected and there is a true and total measure of the worker exposure.

Secondly, the sorbent media is exposed to the external environment and this poses problems not associated with a sample tube where the air sample passes into the sample tube directly contacting the sorbent media. That is why it is critical to use a strong sorbent medium in passive samplers to assure complete capture and retention.

Therefore, for compliance purposes a passive sampler must be laboratory tested and validated under worst case field conditions for all factors that affect sampling accuracy as well as interaction between affects.

NIOSH has laid out a rigorous and complete validation protocol to assure that the sample collected is a complete and true measure of worker exposure. The following are the factors that the NIOSH protocol addresses:

Factors That Affect Complete Sample Uptake & Retention

Chemical Hazard Concentration Temperature

Time of Exposure Humidity

Sorbent Capacity Interfering Chemicals

Sorbent Strength Reverse Diffusion from Sorbent Surface

Wind Velocity Sampler Orientation

Interaction of Any of the Above Factors

Validation by NIOSH protocol assures that the sample results are a true and total measure of worker exposure.

SKC Validation follows the NIOSH Validation Protocol. Certain experiments may have been modified for practical reasons, or to provide more rigorous tests.

User Responsibility

The sampler manager should be a professional trained in air sampling and aware of the limitations and advantages of the method being used. It is also very helpful if they have a working relationship with the analytical techniques being used and the requirements of record keeping.

In accordance with ASTM D6346-98 and ANSI 104-1998 standards, use of samplers outside the range of conditions used in these validation tests does not assure accurate results and is not recommended. It is the user's responsibility to determine whether the conditions of the sampling site fall within the range tested. For bi-level validations it can be assumed that the applicable range is that used for testing the lower member of the homologous series.

Workers should be trained in the use of the equipment. In collecting the sample, care should be taken in the location of the sampler on the worker. It is to be openly exposed near the breathing zone. Exact times of exposure must be recorded. No moisture condensation should occur on the sampler. Workers should not be allowed to touch the sampler as they may transfer contamination. Particular attention must be paid to environments where liquid aerosols may be present, since droplets of liquid solvent on the sampler face will invalidate the sample. Any other field conditions outside of the limits used in the NIOSH protocol, such as extreme temperatures or stagnant air conditions which might affect the sampler operation should be recorded.

Good laboratory practice must be followed. Follow the operating instructions for the desorption time needed for complete desorption. Use only the correct desorption instrument (SKC Cat. No. 226D-03-01). If gas chromatography is used as the analysis method, base line separation should occur with the chemical hazard of interest and proper instrument calibration procedures used.

NIOSH or OSHA analytical methods should be used.

Publication No. 1315 Rev 0510

Summary of NIOSH Validation Protocol¹

Characteristic	Experimental Design		Interpretation of Results
1. Analytical Recovery	Spike 16 samplers, 4 at ea levels (0.1, 0.5, 1.0 & 2.0 about 12 h and analyze.		For the higher 3 levels require \geq 75% recoveries with $S_r \leq 0.1$.
2. Sampling Rate and Capacity	Expose samplers (4 per tin 1/2, 1, 2, 4, 6, 8, 10 & 12 and 20 cm/s face velocity. time exposed. Determine	n to 2 x STD, 80% RH Plot concentration vs.	Verify sampling rate. State useful range at 80% RH & 2 x STD. Capacity - sample loading corresponding to the downward break in conc. vs time curve from constant concentration. SRST - time linear uptake rate achieved. MRST-0.67 x capacity (1 analyte) MRST-0.33 x capacity (Multi-analyte)
3. Reverse Diffusion	Expose 20 samplers to 2 x MRST. Remove and and Expose others to 80% RH remainder of MRST.	alyze 10 samplers.	Require \leq 10% difference between means of the two sampler sets at the 95% CL.
4. Storage Stability	Expose 3 sets of samplers RH, 1 x STD, and 0.5 x M within 1 day, second set a about 25° C, third set after about 5° C.	IRST. Analyze first set fter 2 weeks storage at	Require ≤ 10% difference at the 95% CL between means of stored sampler sets and set analyzed within 1 day.
5. Factor Effects	Test the following factors Use a 16 -run fractional fa samplers per exposure) to factors.	ctorial design (4	Indicate any factor that causes a statistically significant difference in recovery at the 95% CL. Investigate further to characterize its effect.
	Factor analyte concentration exposure time face velocity relative humidity interferant sampler orientation	Test Levels 0.1 & 2 x STD SRST & MRST 10 & 150 cm/s 10 & 80% RH 0 & 1 x STD parallel & perpendicular (to air flow)	
6. Temperature Effects	Expose samplers (10 per t 10, 25, & 40° C for 0.5 x 1		Define temperature effect and verify correction factor, if provided.
7. Accuracy and Precision	Calculate precision and bi conc. level) exposed to 0. 80% RH for ≥ MRST. Us experiments.	1, 0.5, 1 & 2 x STD at	Require bias within \pm 25% of true value at 95% CL with precision $S_r \le 10.5\%$ for 0.5, 1, & 2 x STD levels.

Summary of NIOSH Validation Protocol (cont.)

Characteristic	Experimental Design	Interpretation of Results		
8. Shelf Life	Observe samplers throughout evaluation for changes in blank values, physical appearance, etc. Test samplers from more than one lot, if possible.	Note shelf storage time at which changes begin to occur. Indicate whether correctable or not.		
9. Behavior in the Field	Consider problems not predictable from laboratory experiments.	Record temperature, humidity, air velocity, other contaminants, etc.		
Area Sampling:	Expose passive samplers and independent method samplers (13 each) to the same environment.	Calculate precision and bias. Compare with laboratory results.		
Personal Sampling:	Conduct personal sampling with ≥ 25 sampler pairs. Place pairs of passive samplers and independent samplers on the same lapel of each worker.	Calculate bias. Compare with area sampling and laboratory results		

Bi-Level Validation (previously designated by SKC as 5B)

Validation of passive samplers is essential to ensure accurate determination of airborne chemical levels. To assist manufacturers and users, the National Institute for Occupational Safety and Health (NIOSH), the Health and Safety Executive (HSE)², and the Comité Européen de Normalisation (CEN)^{3,4} have developed comprehensive protocols for the validation of passive samplers.

Bi-level validation can also be used to assure a sample that gives the total and complete exposure to a chemical hazard.

Bi-level validation is only for a series of chemically related compounds, i.e., members of a homologous series. Bi-level validation includes a full protocol validation on key compounds followed by a partial validation on other members of the series.

The concept of a bi-level validation of chemically related compounds for a given sorbent and sampler design is based on the following premises and has been studied by Guild et al.⁵

- Full validation by NIOSH, HSE, or CEN Protocol of a lower member of the series is essential to assure accurate, routine sampling under all field conditions without the need for error-corrective measures.
- 2. Capacity and retentivity are directly related to the affinity of a sorbent for a specific chemical. For a series of chemically related compounds, the affinity of a sorbent for a particular member compound will increase with the molecular weight and boiling point of the member. If a sorbent is suitable for collecting a low molecular weight member of the series, it will be suitable for the higher molecular weight members of the series as well.
- 3. For chemically stable compounds, sample loss by reverse diffusion and loss during storage are inversely related to the affinity of the sorbent for the adsorbate. Therefore, compounds with higher molecular weights and boiling points will exhibit less loss by reverse diffusion and storage. Again, if a sorbent is suitable for a member with a lower molecular weight and boiling point, it will be suitable for the higher members.
- 4. The linearity of uptake with time is also a function of sorbent affinity and capacity. Uptake becomes increasingly linear as the molecular weight and boiling point increases and the sample load decreases. (Protocol validation requires study of concentrations ranging from 0.1 to 2.0 x the permissible exposure limit.)

Bi-Level Validation (cont.)

- 5. Temperature affects the accuracy of passive samplers in two different ways; the relation of temperature to adsorption affinity and the relation of the molecular diffusion of the sample to the sampler.
 - a. It is well known that the affinity of a sorbent for a chemical decreases with increasing temperature. If the sorbent has adequate affinity for a low molecular weight member of the series at 40° C (the maximum temperature tested under protocol), it will also be adequate at lower temperatures, and for higher molecular weight members of the series.
 - b. The effects of temperature on sample uptake follow established mathematical relationships and are not significant compared to other random sampling errors.
- 6. The effects of humidity because of competition or modification of sorbent affinity will be most pronounced for lower members of the series.
- 7. Adsorption affinity decreases with the mass adsorbed. Therefore, the "key" member chosen for full validation should have a high PEL relative to the other members of the series.
- 8. Air velocity and sampler-orientation effects are functions of sampler design and will be similar for all compounds.
- 9. If all the factors affecting sampling accuracy improve with increasing molecular weight and boiling point and there are no interacting effects of these parameters with a lower member of the series, then there will be no interacting effects with higher members.
- 10. The accuracy of a sampler is determined by its bias and precision. For most passive samplers, the bias is the result of the deviation of the calculated sample rate from the actual rate. By determining the sample rate under known conditions at 1 PEL, the bias is reduced to zero. Therefore, measured sample rates should be determined for all compounds.
- 11. The precision of a sampler is a function of the consistency of sampler manufacture and the analytical procedures in the laboratory.
- 12. Analytical recovery tends to decrease with increased sorbent affinity and is a function of the chemical compound, the concentration, and the sorbent. Therefore, analytical recovery should be determined for every compound over the concentration range of 0.1 to 2.0 PEL, as recommended by protocol.

Conclusion: The above premises have been verified, peer reviewed and published.⁵ Therefore, Bi-Level validation (5B) is an excellent way to assure accurate performance of a passive sampler for higher members of a homologous series.

Comments on the Relationship Between the NIOSH and CEN Diffusive Sampler Evaluation Protocols

The Comité Européen de Normalisation (CEN) is engaged in writing standards for air sampling equipment which include the limitations on precision and accuracy (EN 482) and the required performance tests. In the case of passive samplers the relevant performance test standard is yet to be published, but draft copies are available (EN 838).

The precision and accuracy requirements in EN 482 are based on the use that will be made of the results, principally either for problem identification or compliance purposes. The standard for compliance purposes is a combined precision and accuracy of less than 30%, which is a looser standard than the 25% in the NIOSH protocol.

The performance tests are closely related to those in the NIOSH protocol, as might be expected, since they are trying to confirm the performance of the samplers over a similar range of environmental conditions. As in the NIOSH protocol there are tests for desorption efficiency, uptake rate at different concentrations and for different time-periods, reverse diffusion, storage stability, wind velocity and orientation, humidity, temperature and the presence or absence of interferences. As in the NIOSH protocol these factors are normally tested using a "high" and a "low" measure, whether alone or in combination. Since there is little difference between workplace conditions in the U.S.A. and Europe, these "high" and "low" conditions are very similar in the two protocols. In general, the NIOSH test provides the more stringent conditions (e.g. 7.5 minutes up to 12 hours in the NIOSH uptake rate experiment versus 30 minutes and 8 hours in the CEN equivalent). In addition, for the majority of the experiments, the NIOSH protocol requires more samples to be taken for each data point (typically 10 rather than 6). The reverse diffusion test is one test that might be considered significantly different, and a paper showing that the results of the tests are actually comparable has been submitted for publication.⁶

In addition, the CEN protocol requires tests for shelf-life and packaging integrity that have been carried out for one analyte (n-Hexane) only. The 575 Series passive sampler successfully passed these tests.

For the reasons given above, SKC considers the validations presented in these research reports to be at least sufficient to meet the requirements of the European Standards prEN 838 and EN 482 for compliance monitoring. This conclusion is supported by a detailed comparison which has been submitted for publication.⁷

The CEN protocol supports the Bi-level theory of validation.

SHELF-LIFE STUDY ON 575 SERIES PASSIVE SAMPLERS

Protocol: 4 expired and 2 unexpired 575-001 samplers were exposed to an atmosphere 100 ppm n-Hexane (2 X PEL) at 80% relative humidity (25° C) for 30 minutes, and then analyzed. Study was conducted August 1995.

Results:

Calculated atmosphere concentration: 106 ppm

Gas sample analysis concentration: 102 ppm (RSD = 7.0%)Sorbent tube analysis concentration: 115 ppm (RSD = 3.2%)

Sampler analysis concentration:[◊]

Sampler expired 12/92: 106 ppm

Sampler expired 4/94: 106 ppm

Sampler expired 10/94: 108 ppm

Sampler expired 10/94: 110 ppm

Sampler unexpired (7/96): 100 ppm

Sampler unexpired (7/96): 100 ppm

Conclusion: Samplers will perform as expected up to their expiration date.

PACKAGING INTEGRITY STUDY ON 575 SERIES SAMPLERS

Protocol: 6 575-001 samplers in unopened Tedlar® pouches were exposed to an atmosphere of 100 ppm n-Hexane (2 X PEL) at 80% relative humidity (25° C) for four hours, and then opened and analyzed.

Results:

Calculated atmosphere concentration: 103 ppm

Gas sample analysis concentration: 104 ppm (RSD = 8.7%)Sorbent tube analysis concentration: 103 ppm (RSD = 2.7%)

Sampler analysis: No detectable n-Hexane in any sampler.

(estimated LOD = 1.5 micrograms, equivalent to 0.125 ppm)

Conclusion: Packaging will prevent contamination of stored samplers.

[⋄] Based on 111.6% desorption efficiency

Scope of the Method

Analyte: Styrene Matrix: Air **Procedure:** Adsorption on a 575-003 SKC passive sampler, desorption with 2 ml of CS₂, and analysis by GC-FID. **Exposure Guidelines:** ACGIH-TLV (1994/95) 50 ppm TWA, 100 ppm STEL 50 ppm TWA, 100 ppm STEL OSHA (1995) 50 ppm TWA, 100 ppm STEL NIOSH (1995) Validation Range, Recovery: Compound Validation Range ppm in air Mean % Recovery Styrene 5-250 100% **Detection Limits:** Depending on the instrumentation, it is possible to determine 13 µg/sampler with a relative standard deviation of less than 10%. This corresponds to an air concentration of 0.5 ppm (v/v) based on an 8 hour sample at the validated sampling rate of 13.7 ml/min. **Temperature Effects:** Samples could be taken up to 40° C. **Factorial:** No significant effects were found due to the interaction of factors that affect sampling accuracy. High humidity conditions (80% RH at 25° C) did not affect the **Humidity Effects:** recovery of Styrene on the 575-003 passive sampler or the uptake rate. **Storage Effects:** The passive sampler can store for at least 21 days at room temperature or in a refrigerator with no loss in recovery. **Interferences:** Any compound that has the same retention time as Styrene will interfere with the analysis. A study was also conducted where passive samplers were exposed to 200 ppm toluene and 100 ppm Styrene and no significant loss in recovery was observed. **Validation Completion Date: April** 1995 **Physical Properties:**

Mol. Weight (g/mole) Boiling Pt. at 760 mm Hg Density (g/ml) 104.15 142.2° C 0.9060

Background

History of Methodology

Previous methodologies have used activated charcoal SKC Lot 120 in a sample tube, or there is a newer method which uses a sample tube containing charcoal coated with tertiary butyl catechol.

Research Purpose

The present work was to evaluate and validate the SKC 575 Series passive sampler containing Anasorb 727 as a method for sampling Styrene. The passive sampler was validated over a concentration range of 0.1 to 5 x PEL. Critical parameters such as analytical recovery, concentration, relative humidity, reverse diffusion, storage stability, temperature, sampling time, wind speed and orientation, and the presence of interfering compounds were addressed.

Experimental

99+% Styrene (Aldrich Chemical Co.) was used. The HPLC-grade carbon disulfide (99.9%) was obtained from Aldrich Chemical Company. The 575 passive sampler containing Anasorb 727 (SKC Cat. No. 575-003) and the Anasorb 747 tubes used for atmosphere calibrations (SKC Cat. No. 226-81) are available from SKC, Inc.

A dynamic atmosphere generation apparatus was used to generate precise concentrations of Styrene in air for exposure of the passive samplers. The system is described in Appendix A and Figure 1. The atmosphere was fed into an exposure test chamber. The passive samplers were exposed on a rotating bracket inside the test chamber to simulate wind velocity and orientation.

Analytical recoveries for the passive samplers were conducted by injecting a known amount of Styrene (as a CS₂ solution) into the back of each sampler. The passive samplers were capped, allowed to equilibrate overnight, and analyzed the next day to determine analytical recovery or desorption efficiency. The tests were conducted at mass loadings equivalent to an 8-hour time weighted average sample (6.48 L at the expected sampling rate of 13.5 ml/min) at 0.1, 0.5 and 2.0 PEL under dry conditions.

The sampling rate, reverse diffusion and storage stability experiments on the passive sampler were conducted under dynamic conditions in the test chamber described above. In the storage stability study, recovery is referred back to the reference samples analyzed on Day 1.

The passive samplers were desorbed (in situ) with 2 ml of CS₂ and shaken on a flatbed shaker for 30 minutes. All extracts were transferred to autosampler vials and analyzed by flame ionization gas chromatography. A chromatogram with analytical conditions is shown in Figure 2.

Sampling Rate Determination

Sampling rates can be determined by one of several statistical methods from the experimental data and they differ by only a small amount. Any bias taken is toward the protection of the worker.

We use the time-weighted average from one to eight hours where results fall within NIOSH criteria.

We constantly review our data and conduct experimental work to provide the most precise sampling rate. This rate may differ slightly from previously published sampling rates. Use the rate listed in this report.

Analytical Recovery

NIOSH Requirements

Experimental Design

Interpretation of Results

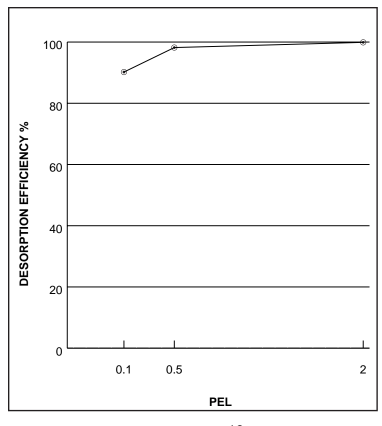
Spike 16 samplers, 4 at each of 4 concentration levels (0.1, 0.5, 1.0 & 2.0 x STD) Equilibrate about 12 h and analyze.

For the 3 higher levels require $\geq 75\%$ recoveries with $S_r \leq 0.1$.

Results

.8
.6
.3

The results from these experiments confirmed observations with sampling tubes containing the same adsorbent indicating a 100% desorption efficiency around the loading of the PEL. Therefore, 100% was used as the desorption efficiency correction throughout. The use of this figure was confirmed by consistency with the results from 575-002 samplers exposed in the same experiments.



Sampling Rate and Capacity

NIOSH Requirements

Experimental Design

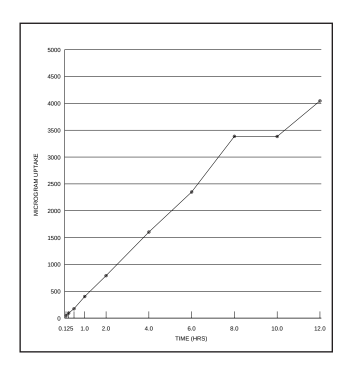
Expose samplers (4 per time period) for 1/8, 1/4, 1/2, 1, 2, 4, 6, 8, 10 and 12 h to 2 x STD, 80% RH and 20 cm/s face velocity. Plot concentration vs. time exposed. Determine MRST and SRST.

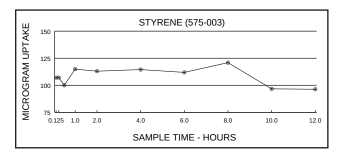
Interpretation of Results

Verify sampling rate. State useful range at 80% RH and 2 x STD. Capacity - sample loading corresponding to the downward break in conc. vs time curve from constant concentration. SRST-time linear uptake rate achieved. MRST - 0.67 x capacity (1 analyte)
MRST-0.33 x capacity (Multi-analyte)

Results

Time (hrs)	Uptake (µg)	Mean (µg)	RSD%	DE Corr	Concn. (ppm)
0.125	44.1	4.9		4.6	(FF)
	45.0				
	49.1				
	49.2	46.8	5.7	46.8	107.0
0.25	97.0				
	84.9				
	102.5				
	90.3	93.7	8.2	93.7	107.1
0.5	179.9				
	180.6				
	158.2				
_	181.1	175.0	6.4	175.0	100.0
1	397.0				
	405.6				
	386.7	402.2	2.4	102.2	1110
2	418.9	402.2	3.4	402.2	114.9
2	785.3				
	816.4 739.8				
	822.1	790.9	4.1	790.9	113.0
4	1444	790.9	4.1	790.9	113.0
4	1587				
	1676				
	1705	1603	7.3	1603	114.5
6	2354	1003	7.5	1003	114.5
O	2422				
	2254				
	2367	2349	3.0	2349	111.9
8	3295	20.7	5.0	20.7	111.,
	3418				
	3445				
	*	3386	2.4	3386	120.9
10	3391				
	3336				
	3495				
	3310	3383	2.4	3383	96.7
12	4052				
	3986				
	3994				
	4150	4046	1.9	4046	96.3





Concentration values are calculated using the sampling rate of 13.7 ml/min obtained from the 575-002 validation, since the 8-hour result documented here is an outlier which would bias the results high. This uptake rate is confirmed by the overall precision and accuracy result (13.9 ml/min \pm 1.2 mL/min).

^{*} Sampler lost.

Reverse Diffusion

NIOSH Requirements

Experimental Design

Expose 20 samplers to 2 x STD 80% RH for 0.5~x MRST. Remove and analyze 10 samplers. Expose others to 80% RH and no analyte for remainder of MRST.

Interpretation of Results

Require \leq 10% difference between means of the two sampler sets at the 95% CL.

Results (in micrograms)

Exposed 4 hours to analyte

Exposed 4 hours to analyte plus 4 hours at zero analyte concentration

Uptake	DE Corr	Uptake	DE Corr.
<u>(µg)</u>	$(\mu \mathbf{g})$	$(\mu \mathbf{g})$	<u>(µg)</u>
3264	3264	2914	2914
3245	3245	2787	2787
2951	2951	2660	2660
3049	3049	2957	2957
Mean:	3127		2830
SD:	152.5		134.1
RSD:	4.9%		4.7%

The difference between the two sets of results is less than 10%.

Results normalized to 240 minutes from 243 minutes.

Study carried out at 4 x PEL (205.7 ppm) as a more rigorous test.

Storage Stability

NIOSH Requirements

Experimental Design

Expose 3 sets of samplers (10 per set) at 80% RH, 1 x STD, and 0.5 x MRST. Analyze first set within 1 day, second set after 2 weeks storage at about 25° C, third set after 2 weeks storage at about 5° C.

Interpretation of Results

Require \leq 10% difference at the 95% CL between means of stored sampler sets and set analyzed within 1 day.

Results (in micrograms)

	Room Ter	np	Refrigerator		
	Uptake	DE Corr.	Uptake	DE Corr	
	<u>(µg)</u>	<u>(µg)</u>	$(\mu \mathbf{g})$	$(\mu \mathbf{g})$	
Day 1 for Day	7 (52.8 ppm)				
	732.9	732.9	768.1	768.1	
Day 1 for Day	14 (54.6 ppm)				
	678.1	678.1	743.7	743.7	
Day 1 for Day	21 (52.4 ppm)				
	720.6	720.6	762.5	762.5	
Day 7					
	758.7	758.7	794.1	794.1	
	707.6	707.6	773.1	773.1	
	715.2	715.2	787.3	787.3	
Day 14					
	734.7	734.7	745.1	745.1	
	723.3	723.3	745.2	745.2	
	690.0	690.0	716.9	716.9	
Day 21					
	795.4	795.4	765.7	765.7	
	762.2	762.2	776.8	776.8	
	731.6	731.6	775.4	775.4	
Mean: Day 7 - 21		735.4		764.4	
SD:		32.1		24.4	
RSD:		4.4%		3.2%	

There is no significant loss of sample on storage.

Experiment altered to track storage closely and to extend storage to 21 days. Day 14 results normalized to 240 minutes from 250 minutes.

Factorial Results

NIOSH Requirements

Experimental Design

Test the following factors at the levels shown. Use a 16 run fractional factorial design (4 samplers per exposure) to determine significant factors.

Factor Test Levels
analyte concentration 0.1 & 2 x STD
exposure time SRST & MRST
face velocity 10 & 150 cm/s
relative humidity 10 & 80% RH
interferant 0 & 1 x STD
sampler orientation parallel &

perpendicular (to air flow)

Interpretation of Results

Indicate any factor that causes a statistically significant difference in recovery at the 95% CL. Investigate further to characterize its effect.

Results (All results in micrograms per ppm per minute).

<u>Run #</u>		Individual M	onitor Results		<u>Average</u>	%RSD
1	3.8580	3.7394	3.8793	3.6430	3.7799	2.9
2	3.2954	3.3959	3.4673	3.6087	3.4418	3.8
3	3.3093	3.1641	3.0097	3.3184	3.2004	4.5
4	3.9346	3.6929	3.8234	3.6400	3.7727	3.5
5	3.1592	3.2074	3.2683	3.3849	3.2550	3.0
6	2.4490	2.5941	2.5114	2.3960	2.4876	3.4
7	3.9648	4.0248	3.7215	4.2464	3.9894	5.4
8	3.3878	3.3469	3.3701	3.2291	3.3335	2.1
9	3.2627	3.3701	3.3804	3.2711	3.3211	1.9
10	3.3624	3.5157	3.4453	*	3.4411	2.2
11	3.3299	3.4115	3.2391	3.1660	3.2866	3.3
12	3.8123	3.7452	3.7644	3.6877	3.7524	1.4
13	4.3246	4.1551	4.0882	3.9824	4.1377	3.5
14	3.0529	2.9844	3.1285	3.1947	3.0901	3.0
15	3.7983	3.6361	3.7235	*	3.7193	2.2
16	3.1087	3.3602	3.3400	3.2797	3.2722	3.5

Notes: Low face velocity = 20 cm/sLow concentration = 0.1 PEL

Minimum sample time = 2 hours

200 ppm Toluene used in the interference experiments.

^{*} Sampler lost

Factorial Summary

Run Num	<u>ıber</u>	μg/j	<u>ppm/hour</u>	
Run#	1	=	3.7799	
Run#	2	=	3.4418	
Run#	3	=	3.2004	
Run#	4	=	3.7727	
Run#	5	=	3.2550	
Run#	6	=	2.4876	
Run#	7	=	3.9894	
Run#	8	=	3.3335	
Run#	9	=	3.3211	
Run#	10	=	3.4411	
Run#	11	=	3.2866	
Run#	12	=	3.7524	
Run#	13	=	4.1377	
Run#	14	=	3.0901	
Run#	15	=	3.7193	
Run#	16	=	3.2722	
Avera	ge	=	3.4550 = 13.5 r	nl/

Average = 3.4550 = 13.5 ml/min

Factor		Effect	Percent	Significance
A -	Concentration	-0.04	1.0%	N.S.
В -	Relative Humidity	0.16	4.7%	N.S.
C -	Interferants	-0.19	5.6%	N.S.
D -	Time	0.01	0.3%	N.S.
E -	Face Velocity	0.05	1.5%	N.S.
F -	Orientation	0.29	8.5%	N.S.
E1 -	ABC	-0.13	3.8%	N.S.
E2 -	ABD	-0.10	2.8%	N.S.
E3 -	AB + EF	0.25	7.1%	N.S.
E4 -	AC + DF	-0.47	13.5%	N.S.
E5 -	AD + CF	0.17	5.0%	N.S.
E6 -	AE + BF	-0.09	2.6%	N.S.
E7 -	CD + BE	-0.07	2.0%	N.S.
E8 -	BC + DE	0.16	4.8%	N.S.
E9 -	BD + CE	-0.26	7.6%	N.S.

 $\label{eq:minimum} \mbox{Minimum Significant Effect (MSE)} = \pm \ 0.50$ No significant effect of factors or their tested interactions

Temperature Effects

NIOSH Requirements

Experimental Design

Expose samplers (10 per temp) to 0.5 x STD at 10, 25, & 40° C for 0.5 x MRST.

40° C

Interpretation of Results

Define temperature effect and verify correction factor, if provided.

40° C

Results (in micrograms)

40)° C	40	° C
@ 0.5	x PEL	@ 5 x	R PEL
Sample	DE Corr.	Sample	DE Corr
$(\mu \mathbf{g})$	$(\mu \mathbf{g})$	<u>(µg)</u>	$(\mu \mathbf{g})$
356.4	356.4	3243	3243
351.4	351.4	3458	3458
341.3	341.3	3286	3286
365.1	365.1	*	*
356.4	356.4	*	*
355.5	355.5	3285	3285
353.7	353.7	3380	3380
363.2	363.2	3253	3253
354.1	354.1	3165	3165
333.3	333.3	3285	3285
Mean:	353.0		3294
RSD:	2.7%		2.7%
Concentration ¹ :	25.8		254
Uptake ² :	3.421		3.242
Theoretical:	3.589		3.589

Uptake is within 10% of theoretical for both low and high concentrations.

^{*} Sampler lost, mean of group substituted for statistical calculations.

¹ In ppm at the sampling temperature.

² Uptake rate measured as micrograms/ppm (sampling temperature)/hour.

Accuracy and Precision

NIOSH Requirements

Experimental Design

Calculate precision and bias for samplers (10 per conc. level) exposed to 0.1, 0.5, 1 & 2 x STD at 80% RH for \geq MRST. Use data from previous experiments.

Interpretation of Results

Requires bias within \pm 25% of true value at 95% CL with precision S_r \leq 10.5% for 0.5, 1 & 2 x STD levels.

All Values in µg/ppm/hr

	Sar	nplers run :	at 5.0 x PEL			Sar	nplers rur	n at 1.0 x	PEL	
Values fo	r individua	l monitors f	or the		Values for individual monitors for the					
Temperat	ure Effects	Experiment	t		Storage St	tability Ex	periment			
3.1121	3.3185	_	3.1524		Day 7 -	3.4702	3.5923	3.3504	3.3864	
3.2436	3.1217	3.0373	3.1524		•	3.6368	3.7599	3.6605	3.7277	
					Day 14 -	3.1049	3.3640	3.3118	3.2670	
	Sar	nplers run a	at 4.0 x PEL		·	3.4052	3.4179	3.4121	3.2825	
Values fo	r individua	l monitors f	or the		Day 21 -	3.4380	3.7948	3.6364	3.4905	
Reverse I	Diffusion E	xperiment			·	3.6379	3.6531	3.7061	3.6994	
3.9669	3.9439	3.5865 3	.7056							
3.5416	3.3872	3.2329 3	.5938			Sar	nplers rur	at 0.5 x	PEL PEL	
					Values for	r individua	l monitors	for the		
	Samplers run at 2.0 X PEL			Temperati	ure Effects	Experime	nt			
Values fo	Values for individual monitors for the				3.3672	3.3199	3.2245	3.4493	3.3672	
Rate/Capa	acity Expen	riment				3.3586	3.3416	3.4314	3.3454	3.1489
4 Hour -	3.6050	3.9620	4.1841	4.2565		<u>Sar</u>	<u>nplers rur</u>	at 0.1 x	PEL	
6 Hour -	3.9178	4.0310	3.7514	3.9395	Values for	r individua	l monitors	for the		
8 Hour -	3.2700	3.2170	3.3703	3.1919	Factorial l	Experimen	t			
					Run #1 -	3.8580	3.7394	3.8793	3.6430	
Values fo	r individua	l monitors f	or the		Run #3 -	3.3093	3.1641	3.0097	3.3184	
Factorial	Experimen	t			Run #14 -	3.0529	2.9844	3.1285	3.1947	
					Run #16 -	3.1087	3.3602	3.3400	3.2797	
Run #2 -	3.2954	3.3959	3.4673	3.6087						
Run #4 -	3.9346	3.6929	3.8234	3.6400						
Run #13 -	4.3246	4.1554	4.0882	3.9824						
Run #15 -	3.7983	3.6351	3.7235							

Summary			Average Values in µg/ppm/hr		
	Relative Standard	Degrees of	Experiment	Average	RSD
PEL	Deviation	Freedom	Rate/Capacity	3.7049	5.9%
			Factorial, 2 PEL	3.6632	3.4%
0.1	6.7%	12	Storage Stability	3.5086	3.6%
0.5	2.7%	9	Temperature 0.5 PEL	3.3354	2.7%
1.0	3.6%	21	Temperature 5.0 PEL	3.1614	2.7%
2.0	3.3%	20	Reverse diffusion, 4.0 PEL	3.6198	6.7%
4.0	6.7%	7	Factorial 0.1 PEL	3.3356	3.5%
5.0	2.7%	7			
			Overall average	3.5395	4.3%
			Overall sampling rate = 13.9 ml/min \pm 1.2 ml/min		

Appendix A

Atmosphere Generation Apparatus

The instrument is designed to expose a known concentration of a chemical hazard to a passive sampler under controlled conditions of: 1. Concentration, 2. Temperature, 3. Humidity, 4. Wind Velocity Effect, 5. Time, and 6. Up to four multicomponent hazards.

Description

The instrument consists of:

- 1. an exposure chamber in which the wind velocity effects are controlled by internal rotating holders,
- 2. an air supply and purification train such that dry air is blended with saturated air under desired temperature conditions so as to provide air at a known flow and selectable humidity,
- 3. an injection system composed of precision motor driven syringes in which 1 to 4 chemical hazards can be injected into the flow system and in which the temperature of the injectors is closely controlled,
- 4. an electrical control system that controls the entire instrument operation,
- 5. the chamber concentration can be verified by either solid sorbent sampling tubes actively sampled or by gas analysis of the gas phase. The particular verification method used will depend on the analyte of interest.

Means are also included to check the relative humidity.

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Styrene (003)

Figure 1 Atmosphere Generation Apparatus

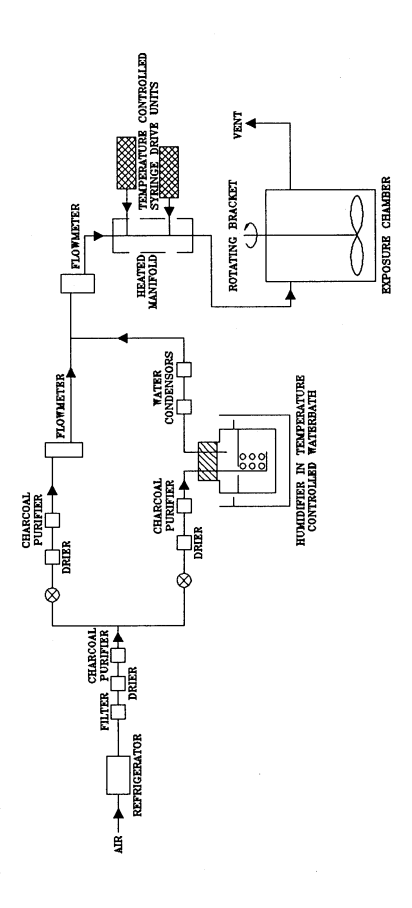
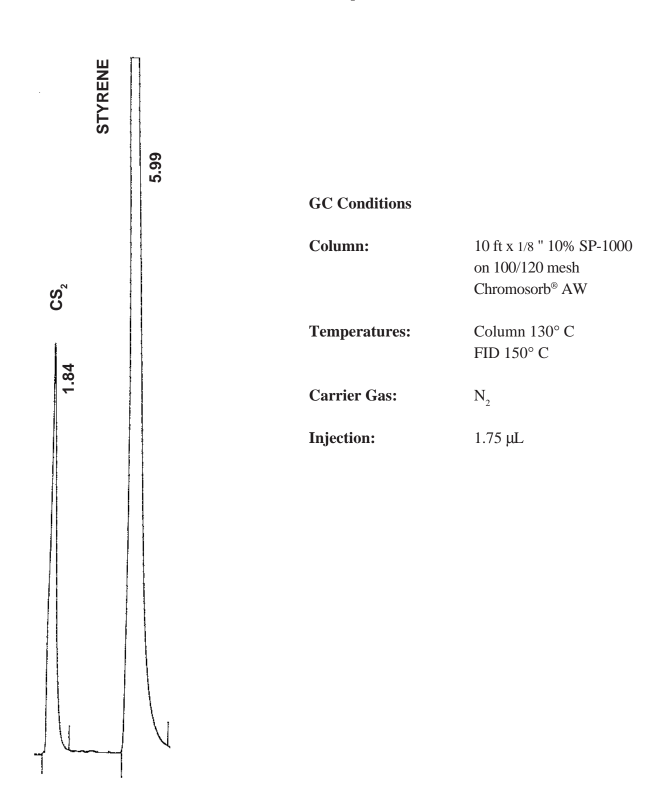


Figure 2 Analytical Instrument

Sample Chromatogram Styrene in CS₂



Abbreviations

C Celsius

CL confidence level

cm centimeter
ml milliliter
min minute
g gram

GC-FID gas chromotography - flame ionization detector

h hourL liter

LOD limit of detection

MRST maximum recommended sampling time

N.S. not significant

PEL permissible exposure limit

RH relative humidity
TLV threshold limit value
TWA time-weighted average
RSD relative standard deviation

SD standard deviation

SRST shortest recommended sampling time

STD the appropriate exposure standard (OSHA PEL, ACGIH TVA, or NIOSH recommended

standard)

S_r Pooled relative standard deviation

S second V volume

Trademarks

Anasorb is a registered trademark of SKC Inc.

Chromosorb is a registered trademark of Manville Corp.

Tedlar is a registered trademarik of DuPont Corporation.

Porapak is a registered trademark of Waters Associates, Inc.

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