Apparent Benzene Solubility in Tetraphenylborate Slurries

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Apparent Benzene Solubility in Tetraphenylborate Slurries (U)

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SUMMARY

Personnel conducted testing to determine the apparent solubility of benzene in potassium tetraphenylborate (KTPB) slurries. The lack of benzene vapor pressure suppression in these tests indicate that for a 6.5 wt % solids KTPB slurry in 4.65 M Na⁺ salt solution at approximately 25 °C, no significant difference exists between the solubility of benzene in the slurry and the solubility of benzene in salt solution without KTPB solids. The work showed similar results in slurry with 6,000 mg/L sludge and 2,000 mg/L monosodium titanate added. Slurries containing tetraphenylborate decomposition intermediates (i.e., 4,200 mg/L triphenylboron (3PB), 510 mg/L diphenylborinic acid (2PB) and 1,500 mg/L phenylboric acid (1PB)) or 100 mg/L tri-n-butyl phosphate (TBP) also showed no significant difference in benzene solubility from filtrate containing no KTPB solids. Slurry containing 2,000 mg/L Surfynol™420 did exhibit significant additional benzene solubility, as did irradiated slurries. The vapor pressure depression in the irradiated slurries presumably results from dissolution of biphenyl and other tetraphenylborate irradiation products in the benzene.

INTRODUCTION

During operation of the In-Tank Precipitation (ITP) facility, benzene is formed as a result of the decomposition of sodium tetraphenylborate (NaTPB). The benzene released during tank stirring is retained by the tetraphenylborate (TPB) solids in the tank during quiescent, unstirred periods. An understanding of retention of the benzene by solids in the tank is important to understanding and predicting releases from the tank. Generation and release calculations use benzene vapor-equilibrium (Henry's Law) constants as key parameters. Walker reported vapor-liquid equilibrium information for benzene in simulated ITP salt solutions. However, until recently, no work determined the amount of benzene held by benzene solids. Crawford began work to determine the benzene capacity of the slurry. Crawford’s work indicated that slurries could hold more benzene than salt solution. High Level Waste Engineering (HLWE) requested further information about the benzene capacity of the slurries to resolve questions on TPB slurry benzene retention and release as described in the Defense Nuclear Facility Safety Board (DNFSB) Recommendation 96-1 Implementation Plan. This report documents the results of tests to determine the apparent solubility of benzene in KTPB slurries.

DISCUSSION

Crawford’s Tests

During 1996, Crawford conducted several tests which indicated that the solubility of benzene in KTPB slurries containing ≥3 wt % solids was increased over that of filtrate containing no solids or ≤1 wt % solids. The apparent solubility of 3 wt % KTPB slurry at 35 °C was estimated to be ~1800 mg/L as opposed to ~200 mg/L for filtrate or slurry containing ≤1 wt % KTPB solids. This apparent solubility of benzene in slurry is defined as the lowest benzene concentration in the slurry that produced a saturated vapor phase. The suppression of vapor pressure was thought to result from adsorption of benzene on the solids, increasing the benzene capacity for the slurry above that which would be dissolved in the filtrate. Crawford’s method used a magnetic stirrer to agitate slurries kept in a constant temperature in a water bath. Figure 1 gives a diagram of Crawford’s Vessel. Data from Crawford’s tests are given in Figures 2 (25 °C) and 3 (35 °C). The results at both temperatures indicate that for slurries containing ≤1 wt % solids, there is no apparent solubility effects. For slurries containing ≥3 wt % solids, an apparent solubility effect is evident. Tests with benzene dyed red to enhance the visibility indicated that this method did not consistently give good mixing and therefore contact between slurry phase and the vapor phase might not be adequate to obtain equilibrium within a reasonable time. Task technical and QA plans were issued to further study this effect. Because Crawford’s work and a variety of previous work conducted by SRTC personnel indicated that benzene was difficult to completely contain, an initial method was developed to examine the benzene solubility in KTPB slurries by measuring both the slurry and vapor phase benzene concentrations.
Figure 1. Crawford Test Method Vessel

Sample Ports with Teflon Stopcocks and Teflon-lined Septa

Magnetic Stir Bar

Figure 2. Results of Crawford’s Tests at 25 °C

<table>
<thead>
<tr>
<th>Vapor Phase Benzene Concentration (g/L)</th>
<th>Liquid Phase Benzene Concentration (g/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.45</td>
<td>0.1</td>
</tr>
<tr>
<td>0.4</td>
<td>0.2</td>
</tr>
<tr>
<td>0.35</td>
<td>0.3</td>
</tr>
<tr>
<td>0.3</td>
<td>0.4</td>
</tr>
<tr>
<td>0.25</td>
<td>0.5</td>
</tr>
<tr>
<td>0.2</td>
<td>0.6</td>
</tr>
<tr>
<td>0.15</td>
<td>0.7</td>
</tr>
<tr>
<td>0.1</td>
<td>0.8</td>
</tr>
<tr>
<td>0.05</td>
<td>0.9</td>
</tr>
<tr>
<td>0.0</td>
<td>1.0</td>
</tr>
</tbody>
</table>

- □ 3 wt % Solids
- △ 5 M Na+ Salt Soln
- ○ 1 wt % Solids
Figure 3. Results of Crawford's Tests at 35 °C

Initial Test Method

Researchers prepared KTPB slurries. An aliquot of slurry was placed in each test vessel and the vessel closed. Figure 4 is a diagram of the initial test method vessel. An amount of benzene specified by the researcher was injected into the slurry through a septum. The test vessel was placed in a temperature controlled shaker/water bath. The slurry was allowed to agitate until vapor and liquid concentrations stabilize. The vessels were checked visually to verify effectiveness of agitation. The test vessels were vented periodically during the first few hours of equilibration to avoid pressurization of the test vessel. After allowing the vapor and liquid to equilibrate, samples were drawn from both slurry and vapor phases and analyzed for benzene using gas chromatography. Tests of this method using dyed benzene indicated satisfactory mixing.

Figure 4. Initial Test Method Vessel
Tests of this method seemed to indicate no apparent solubility effect in the slurries tested. The difference between these results and previous results from Crawford was attributed to a lack of adequate mixing in the Crawford tests. However, the slurry phase benzene concentration results proved erratic and therefore cast doubts on the interpretation of the results. Figure 5 gives an example of the results using this method for a 6.5 wt % solids slurry containing 5 M Na⁺ at 35 °C.

Figure 5. Example of Slurry and Vapor Phase Benzene Concentrations Using Initial Test Method

![Graph showing slurry and vapor phase benzene concentrations](attachment:graph.png)

Second Test Method

The authors decided to reconsider use of a mass balance method. In this method a known amount of benzene is injected into a sealed test vessel containing slurry. Figure 6 contains a diagram of the vessel. The vessel was agitated in the shaker water bath overnight. A sample of the vapor phase was taken and analyzed for benzene using gas chromatography, and the slurry phase benzene calculated by subtracting the benzene in the vapor phase from the total benzene added.⁹
Preliminary tests of this method with vapor phase benzene indicated good, but incomplete containment of the benzene (80 - 100% found) and insignificant benzene losses over time (Figures 7 and 8). Preliminary tests with benzene in water gave good agreement with referenced data for water (Figure 9), and the data improved when calculated liquid phase benzene concentrations were used instead of measured (Figure 10) as done for the slurry tests.
Figure 8. Vapor Phase Benzene Results with Respect to Time

Figure 9. Benzene Solubility Results in Water at Room Temperature with Measured Liquid Benzene Concentrations
Initial Test Method Results

As stated above, the initial tests used a method that measured both the slurry and vapor phase benzene concentrations. The original scope of tests included 5 test sequences with the majority of the tests run at 35°C. The first of these test sequences examined the impact of salt solution contents (i.e., Na⁺ and OH⁻ concentrations on the apparent solubility of benzene in KTPB slurries). The effect of sludge and titanate solids on the solubility of benzene in filtrate and slurry was investigated in Test Sequence #2. Researchers planned to examine the impact of slurry irradiation and high temperatures in the third test sequence; this sequence was to look at apparent benzene solubility in the Late Wash process and therefore, a low sodium concentration slurry was to be used. Researchers investigated the impact of tetraphenylborate degradation intermediates (i.e., triphenylboron, diphenylborinic acid and phenylboric acid) and surfactants used in the tank farm (e.g., tri-n-butyl phosphate, Surfynol™420 and Do-Corning 544, H-10 and Emulsion B) on the apparent solubility of benzene in KTPB slurry in the fourth test sequence. The effect of solids contents between 1 and 12 wt % solids and temperatures between 25 °C and 50 °C were to have been examined in Test Sequence #5.

Several problems exist with the initial test method. The tests at ~35 °C, above the ambient temperature (~21 °C) for the lab module. Since the method initially used syringes at room temperature to extract samples, the temperature difference led to some condensation of benzene in syringes for samples with high benzene concentrations. In some cases this additional benzene condensate increased the measured benzene concentration to levels well in excess of the saturation concentration. This problem was recognized, and the syringes later warmed to 55 °C (i.e., the maximum temperature allowed for handling without protection) to eliminate this problem. Vapor phase benzene measurements agreed much better with expected values with this change, but the problem was not entirely eliminated. A second problem with the initial test method involved inconsistency in the measured slurry benzene concentration results. Unfortunately, no immediately available way of combating this problem existed. A final problem with the initial test method involved the potential for leakage from the vessels. In this method, only a single Teflon-lined septum separated the slurry vapor space and the atmosphere. Due to multiple piercing of the septum during each test, the potential for leakage of the benzene increased. Large leakage could reduce the apparent equilibrium vapor phase benzene concentration with respect to the liquid phase benzene concentration. The second test method controlled this problem by adding a Teflon stopcock between the slurry and the septum. The stopcock was only open when adding benzene or sampling the vapor space.
Test Sequence #1 from Reference 7 examined the effect of salt solution composition on benzene retention in slurry. The tests used slurry containing ~6.5 wt % KTPB solids with ~4.65 M Na⁺ and an average OH⁻ content, with 5 M Na⁺ and a high OH⁻ content and with 5 M Na⁺ with a low OH⁻ content. Slurries with 2 M Na⁺ and an average OH⁻ content and 0.2 M Na⁺ and an average OH⁻ content were also tested. Table 1 gives the salt contents of these slurries. The tests occurred at a temperature of ~35 °C. Figures 11A and 11B graph the results. Previous work estimates the benzene saturation for 4.65 M and 5 M Na⁺ salt solution as approximately 200 mg/kg; saturation for the 0.2 M Na⁺ salt solution would approach the benzene saturation in water of approximately 2,000 mg/kg. Benzene saturation for the 2 M Na⁺ salt solution would fall somewhere between these two values. Though the results contain considerable scatter at high benzene concentrations, the 4.65 M Na⁺, average OH⁻ slurry and the 5 M Na⁺, high hydroxide slurry do appear to give benzene saturation results close to 200 mg/kg, and the 0.2 M Na⁺ slurry appears to give a benzene saturation value close to 1800 mg/kg (Figure 11A). Unfortunately, the results for the other two slurries tested prove inconclusive (Figure 11B).

Table 1. Final Dissolved Salt Composition of Slurries to be Tested

<table>
<thead>
<tr>
<th>Component</th>
<th>Concentration in 4.65 M Na⁺ Salt Solution</th>
<th>Concentration in 5 M Na⁺ Salt Solution</th>
<th>Concentration in 5 M Na⁺ Salt Solution</th>
<th>Concentration in 2 M Na⁺ Salt Solution</th>
<th>Concentration in 0.1 M Na⁺ Salt Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>NaOH</td>
<td>1.32</td>
<td>0.67</td>
<td>2.50</td>
<td>0.57</td>
<td>0.066</td>
</tr>
<tr>
<td>NaNO₃</td>
<td>1.89</td>
<td>2.58</td>
<td>1.50</td>
<td>0.80</td>
<td>0.094</td>
</tr>
<tr>
<td>NaNO₂</td>
<td>0.59</td>
<td>0.29</td>
<td>0.43</td>
<td>0.24</td>
<td>0.023</td>
</tr>
<tr>
<td>NaAlO₂</td>
<td>0.31</td>
<td>0.17</td>
<td>0.50</td>
<td>0.13</td>
<td>0.016</td>
</tr>
<tr>
<td>Na₂SO₄</td>
<td>0.13</td>
<td>0.30</td>
<td>0.02</td>
<td>0.06</td>
<td>-</td>
</tr>
<tr>
<td>Na₂CO₃</td>
<td>0.14</td>
<td>0.35</td>
<td>0.02</td>
<td>0.07</td>
<td>-</td>
</tr>
<tr>
<td>NaTPB</td>
<td>0.001</td>
<td>0.001</td>
<td>0.001</td>
<td>0.0001</td>
<td>0.0002</td>
</tr>
<tr>
<td>Na⁺</td>
<td>4.65</td>
<td>5.0</td>
<td>5.0</td>
<td>2.0</td>
<td>0.20</td>
</tr>
</tbody>
</table>
Figure 11A. Results of Sequence #1 Tests from Initial Test Method with Regression of Low Concentration Data

Figure 11B. Results of Sequence #1 Tests from Initial Test Method with No Regression
Test Sequence #2 using the initial method examined the effect of sludge and titanate solids on both 4.65 M Na⁺ salt solution alone and on salt solution containing 6.5 wt % KTPB solids. Figure 12 gives the results. Again, though noisy, the results indicate no appreciable difference in benzene solubility between salt solution containing no solids and salt solution containing either 6,000 mg/L simulated Purex sludge or 2,000 mg/L monosodium titanate. Additionally, 6.5 wt % solids KTPB slurry in 4.65 M Na⁺ salt solution with either sludge or titanate gives a benzene solubility not appreciably different from that of salt solution alone.

Figure 12. Results of Sequence #2 Tests from the Initial Test Method

Test Sequence #4 of the initial test method examined the variance in the solubility of benzene in 6.5 wt % solids KTPB slurry in the presence of tetraphenylborate degradation intermediates and several surfactants used in the High Level Waste process. The slurry used to examine the effects of TPB intermediates contained 4,200 mg/L of triphenylboron (3PB), 510 mg/L of diphenylborinic acid (2PB) and 1,500 mg/L of phenylboric acid (1PB). The slurries used to examine the effects of surfactants contained either 100 mg/L of tri-n-butyl phosphate, 2,000 mg/L of Surlynol™420 or 100 mg/L each of Dow-Corning 544, Dow-Corning H-10 and Dow-Corning Emulsion B. Figure 13 provides the results. The results proved too noisy to draw any conclusion.
Test Sequence #3 was planned to examine the effect of radiation and high temperature on the solubility of benzene in KTPB slurry. Test Sequence #5 was planned to examine the effect of KTPB solids content and temperature on the solubility of benzene. However, as a result of the problems encountered with the initial test sequences, the authors decided to discontinue use of the experimental method.

Results of Second Test Method

The authors adopted a second test method using the vessel shown in Figure 6. Also, the scope was modified due to knowledge gained from the first tests. Since the results from the original method indicated no apparent suppression of benzene vapor pressure over KTPB slurry, the scope of testing was reduced to verify no vapor pressure suppression effect with the addition of KTPB solids and to examine the effect of the various additives. More expansive testing of the effects of solids content and other variables remained in Reference 9 as an optional addition should further detail be desired. Testing at higher temperatures was also eliminated because lack of any apparent solubility effect would imply that the temperature effect is predictable (i.e., it would be the same as for benzene in salt solution alone) and because of the difficulties encountered with getting accurate vapor phase measurements at those conditions.

For the second test method, Test Sequence #1 established a baseline by examining the solubility of benzene in 4.65 M Na⁺ salt solution. Table 1 gives the composition of this salt solution. For each of the filtrate samples, the benzene concentration was measured on the two consecutive days to verify equilibrium. Figure 14 shows the results. The dashed line represents a regression of measured vapor and liquid phases. The discrete points represent measured vapor phase concentrations and implied liquid phase concentrations (i.e., total benzene added minus the amount found in the vapor phase divided by the liquid/slurry mass). The implied results shown represent the
average of the two values for each sample. The shaded area was derived from the difference between the measured and implied liquid concentration results and represents the area where benzene solubility data for a solution or slurry with a benzene solubility the same as that of our 4.65 M Na+ filtrate might be expected to fall with a 95% confidence level. For comparison, this data is included in the graphs of all the other results. The saturation vapor phase benzene concentration and Henry’s Law liquid-vapor phase benzene concentration relationship inferred from previous work conducted by Wake are also given. The difference between the Walker Henry’s Law line and our data result from small differences in the salt solutions.

Test Sequence #2 examined the effect of the addition of KTPB solids on the solubility of benzene in salt solution. Figure 14 shows the results. For each benzene concentration, two test vials were prepared. The vapor phase benzene concentration in each vial was measured on three consecutive days to verify equilibrium. The average standard deviation of each of those sets of three measurements was 6.5%. There was no uniform trend in the changing concentrations. The three measurements for each test vial were averaged; the averages are given in Figure 14. These results indicate no apparent benzene solubility effect in KTPB slurry.

Test Sequence #3 of the second test method examined the effect of 6,000 mg/L sludge and 2,000 mg/L titanate solids on the solubility of benzene in the same KTPB slurry as used above. In this sequence, five test vials were prepared containing different amounts of benzene. The benzene concentration in each test vial was measured one
day and three days after preparation of the vials. The vials showed an average of 9% decrease in vapor phase benzene concentration from the first to the second sample. The results from the two measurements for each of the test vials were averaged and are presented in Figure 15. Again the benzene solubility results show no significant deviation from the solubility of benzene in salt solution.

Figure 15. Results Sequence #3 Tests Using the Second Test Method

The shaded area was derived from a comparison of the measured liquid phase and implied liquid phase benzene concentrations. It represents the region where implied data would fall and still be considered to represent the measured (Henry's Law) data with a 95% confidence level.

The effect of TPB intermediates was examined in Test Sequence #6. Researchers mixed the standard 6.5 wt % KTPB slurry with 4,200 mg/L of 3PB, 510 mg/L of 2PB and 1,500 mg/L of 1PB in five test vials along with varying amounts of benzene. The amounts of intermediates used reflected the maximum concentrations of these intermediates expected in the process. As with previous tests, the benzene concentration in each of the three vials was measured on consecutive days starting one day after the test vials were prepared. The values for the two measurements for each of the test vials were averaged with the results given in Figure 16. As with previous results, the TPB intermediates appear to have negligible effect on benzene solubility in the slurry.
Test Sequence #5 examined the effect of 100 mg/L tri-n-butyl phosphate (TBP) on the solubility of benzene in KTPB slurry. This amount of tri-n-butyl phosphate approximates the concentration which will be added to filtrate upstream of the ITP stripper to control foaming. Again, the same 6.5 wt % KTPB slurry was used and five test vessels with varying benzene concentrations were prepared. Benzene concentration measurements were taken on consecutive days starting one day after preparation of the test vials. The results of the two measurements for each test vial were averaged with the averages given in Figure 17. The results appear to indicate little if any vapor pressure suppression of benzene in the slurry. However, previous work indicated a significant suppression of benzene vapor pressure in the presence of large quantities of tri-n-butyl phosphate.\textsuperscript{10} Because TBP is relatively insoluble in salt solution and would float on salt solution or slurry, large quantities could accumulate in tanks downstream of the ITP stripper. Further testing would help to quantify the magnitude of this effect.
The shaded area was derived from a comparison of the measured liquid phase and implied liquid phase benzene concentrations. It represents the region where implied data would fall and still be considered to represent the measured (Henry's Law) data with a 95% confidence level.

The effect of 2,000 mg/L of Surfynol™420 on the solubility of benzene in 6.5 wt % KTPB slurry was investigated in Test Sequence #7. This concentration of Surfynol™420 reflects the concentration expected to be added to control foaming in the Late Wash process; the concentrations expected to reach ITP as part of the recycle stream are much smaller. Five tests were conducted with varying benzene concentrations. The vapor phase benzene concentrations in the vessels were measured at 1 and 2 days after preparation of the vessels. The average results of the two analyses for each vessel are shown in Figure 18. The results of this test show a strong benzene vapor pressure suppression effect. This effect likely results from the dilution of benzene by the Surfynol™420. If this effect causes the benzene vapor pressure suppression, the benzene vapor-liquid equilibrium in a benzene-Surfynol™420 mixture would obey the following relationship.

\[
PP_{Bz} = VP_{Bz} \left( \frac{x_{Bz}}{x_{Bz} + x_S} \right),
\]

where \( PP_{Bz} \) = the partial pressure of benzene, 
\( VP_{Bz} \) = the saturation pressure of benzene, 
\( x_{Bz} \) = the moles of benzene in the benzene-Surfynol™420 solution and 
\( x_S \) = the moles of Surfynol™420 in the benzene-Surfynol™420 solution.
This treatment considers only the benzene-Surfynol™420 system and assumes that that system follows Raoult's Law. Any deviation from Raoult's Law would cause a difference between the predicted and measured concentration values. Researchers also ignored any benzene or Surfynol™420 that might be dissolved in the aqueous phase, and any other effects of the slurry phase. Figure 19 plots the benzene vapor pressure from this relationship along with the measured benzene concentrations. The shapes of the curves in the area studied are about the same, but the theoretical vapor concentrations are offset low with respect to the measured values. This may be the result from the assumptions discussed above. A example calculation of the theoretical vapor pressure of benzene in the Surfynol™420-benzene system is provided in the Appendix.

Figure 18. Results of Sequence #7 Tests Using the Second Test Method

The shaded area was derived from a comparison of the measured liquid phase and implied liquid phase benzene concentrations. It represents the region where implied data would fall and still be considered to represent the measured (Henry's Law) data with a 95% confidence level.
The Sequence #4 tests used slurry irradiated to 95 Mrad in the Gamma Cell. Similar amounts of benzene were added to the five test vessels as added in the previous tests. The benzene concentration was measured both one day and four days after the test vessels were prepared. The vapor phase concentrations increased an average of 14% during that period. The results of the Sequence #4 tests are shown in Figure 20. As with the Surfynol™420 tests, there is obvious vapor pressure suppression. This suppression likely results from dilution of the benzene with biphenyl that accumulates during the irradiation. The slurry was analyzed for biphenyl concentration with the result given in Table 2.

Because of the vapor pressure suppression seen in Sequence #4, it was decided to initiate Test Sequences # 8, 9 and 10 with the slurry irradiated to 30, 60 and 120 Mrad, respectively. Also, because of the vapor pressure suppression noted in Sequence #4, larger amounts of benzene were used in the later test sequences. Figure 20 gives the results of all four sequences. The results of these tests indicate the existence of vapor pressure suppression that increases with irradiation exposure. Because the vapor pressure suppression in the irradiated slurries presumably results from the same mechanism as vapor pressure suppression in slurry with Surfynol™420, it should also obey Equation (1). Figure 21 shows the results of these test sequences along with the predicted theoretical values. The calculations used to produce the predicted curves are given in the Appendix. The shapes of the curves are similar, but again there appears to be an offset. The offset is not as consistent as the offset in the slurries containing Surfynol™420. Since the biphenyl is insoluble in aqueous solutions, the difference between the predicted and measured vapor pressure of benzene probably results from nonideality of the system with respect to Raoult's Law, from experimental variability, and from any benzene which might be dissolved in the aqueous phase.
Table 2: Biphenyl Concentrations in the Irradiated Slurries

<table>
<thead>
<tr>
<th>Slurry Exposure (Mrad)</th>
<th>Biphenyl Concentration (mg/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>30</td>
<td>1,903</td>
</tr>
<tr>
<td>60</td>
<td>2,411</td>
</tr>
<tr>
<td>95</td>
<td>3,000</td>
</tr>
<tr>
<td>120</td>
<td>3,365</td>
</tr>
</tbody>
</table>

Figure 20. Results of the Sequence #’s 4, 8, 9 and 10 Tests Using the Second Test Method
Figure 21. Theoretical and Measured Vapor Phase Benzene Concentrations Over Irradiated 6.5 wt % KTPB Slurry

CONCLUSIONS

From these tests, it appears that KTPB solids alone do not appreciably affect the solubility of benzene in salt solution. The same is true of sludge solids, monosodium titanate and TPB decomposition intermediates. In low concentrations the same appears true for tri-n-butyl phosphate. However, large quantities of materials soluble in benzene, such as Surfynol™420, biphenyl and other radiolytic decay products of KTPB can significantly affect the solubility of benzene in KTPB slurry and significantly reduce the vapor pressure.

QUALITY ASSURANCE

The work documented in this report was performed under Task Technical Plan WSRC-RP-96-775, Revisions 0 and 179 and Task QA Plan WSRC-RP-96-776, Revision 0. Instructions and results are documented in Lab Notebooks WSRC-NB-95-308, WSRC-NB-96-611, WSRC-NB-97-00494 and WSRC-NB-97-493.

REFERENCES


APPROVALS

M. J. Barnes, Design Check per Manual E7, Procedure 2.40
Waste Processing Technology
Date: 11/13/97

S. D. Fink, Manager
Liquid Waste Processing
Date: 11/17/97

W. B. Van Pelt, Manager
Waste Handling and Mechanical Processing
Date: 11/18/97

J. L. Barnes, Deputy Manager
Waste Pre-Treatment Engineering
Date: 11/19/97

J. T. Carter, ITP Flow-Sheet Task Team
Date: 11/19/97

W. L. Tamosaitis, Manager
Waste Processing Technology
Date: 12/1/97
APPENDIX

Prediction of Vapor Pressure of Benzene Containing Dissolved Organics

This calculation assumes that any benzene not dissolved in the aqueous phase exists in a solution with the dissolved organic in question (e.g., Surfynol™420 or biphenyl). The vapor pressure of the benzene above the mixture remains proportional to the mole fraction of the benzene in the benzene-organic solution according to equation (1) restated below.

\[
PP_{Bz} = VP_{Bz} \left( \frac{x_{Bz}}{x_{Bz} + x_D} \right),
\]

where

- \( PP_{Bz} \) = the partial pressure of benzene,
- \( VP_{Bz} \) = the saturation pressure of benzene,
- \( x_{Bz} \) = the moles of benzene in solution with the dissolved organic and
- \( x_D \) = the moles of the dissolved organic in the solution with the benzene.

This calculation assumes the dissolved organics discussed have negligible solubility in the aqueous phase with respect to the total amount added. Previous work shows the solubility of Surfynol™420 as less than 25 ppm in salt solutions.\(^\text{12}\) Lange's Handbook of Chemistry lists biphenyl as insoluble in water.\(^\text{13}\)

Surfynol™420

Given the following information:

- Molecular Weight for Surfynol™420: 283.63\(^\text{12}\)
- Molecular Weight for Benzene: 78.11\(^\text{13}\)

Implied Benzene Concentration in Liquid (assumed all in Benzene-Surfynol™420 phase): 57.2 mg/kg

Added Concentration of Surfynol™420 in liquid (assumed all in Benzene-Surfynol™420 phase): 2,000 mg/L

Amount of Slurry/Liquid Phase: 30 mL with a density of 1.2 g/mL (i.e., 36 g)

Vapor Pressure of Benzene at 25 °C: 0.126 atm\(^\text{14}\) (403 mg/L).

The total number of moles of Surfynol™420 may calculated by:

Moles = Concentration(mass/volume) * Liquid Volume / Molecular Weight with appropriate unitary conversions.

Moles Surfynol™420 = (2,000)*(30/1000)/1000/283.63 = 0.00021

The total number of moles of Benzene may be calculated by:

Moles = Concentration(mass/mass) * Liquid Weight / Molecular Weight with appropriate unitary conversions.

Moles Benzene = (57.2)*(36/1000)/1000/78.11 = 0.0000264
Using Equation (1) to calculate the vapor phase benzene concentration gives

\[
\text{Vapor Phase Benzene Concentration} = \frac{(403)(0.0000264)}{(0.0000264 + 0.000211)} = 44.8 \text{ mg/L}
\]

**Biphenyl**

Given the above information plus the following additional information:

Molecular Weight of Biphenyl: 154.20^{13}

Implied Benzene Concentration in Liquid (assumed all in Benzene-Surfynol^{TM}420 phase): 128 mg/kg

Measured Biphenyl Concentration in Liquid (Irradiated to 30 Mrad): 1903 mg/L

The total number of moles of benzene calculated as above is

\[
\text{Moles Benzene} = \frac{(128)(36/1000)}{1000/78.11} = 0.0000590
\]

The total number of moles of biphenyl is calculated by:

\[
\text{Moles Biphenyl} = \frac{(1903)(30/1000)}{1000/154.20} = 0.000370
\]

Using Equation (1) to calculate the vapor phase benzene concentration gives

\[
\text{Vapor Phase Benzene Concentration} = \frac{(403)(0.0000590)}{(0.0000590 + 0.000370)} = 55.4 \text{ mg/L}
\]
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