Foaming in Hanford River Protection Project
Waste Treatment Plant LAW Evaporation Processes – FY01 Summary Report

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Foaming in Hanford River Protection Project
Waste Treatment Plant LAW Evaporation
Processes – FY01 Summary Report

SAVANNAH RIVER TECHNOLOGY CENTER

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Publication Date: December 2001

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Prepared for the U.S. Department of Energy under Contract No. DE-AC09-96SR18500
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Executive Summary

The LAW evaporation processes currently being designed for the Hanford River Protection Project Waste Treatment Plant (RPP-WTP) are subject to foaming. Experimental simulant studies have been conducted in an effort to achieve an effective antifoam agent suitable to mitigate such foaming. Specifically, nine non-radioactive Hanford simulants were prepared by SRTC and shipped to Illinois Institute of Technology for foaming tests: AN105 (A), AN105 with entrained solids (A), AN107 (C), AN107 with entrained solids (C), AZ101 (B), AZ102 (B), AN104 (A), AN104 with entrained solids (A), and Pretreated AN107 (C). Each simulant went through a small-scale evaporation process (boiling) from which respective degrees of foaminess and foam textures were recorded. The waste simulants that were selected for evaporation testing are representative of the Phase 1 LAW feeds to the WTP. Phase 1 LAW feeds, instead of Phase 2 single-shell tank wastes, were selected because these wastes are better characterized and actual Phase 1 waste samples are available for comparison with the simulants.

Researchers at the Savannah River Technology Center (SRTC) and the Illinois Institute of Technology (IIT) jointly completed testing. A report was issued by the researchers at IIT (Attachment A). This document is designed to summarize the key results and explain the preparation of the simulants used in this testing.

Selected Hanford simulants were boiled and tested at different evaporation flux rates and at both atmospheric pressure and vacuum—760 mmHg and 110 mmHg, respectively. A vacuum of 110 mmHg was selected to simulate the operating temperature range (50-60°C) expected to be encountered by the RPP-WTP evaporators. As the simulants’ total solids concentrations increased, severe foaminess (650 to 800% at atmospheric pressure, 100% at vacuum) was observed in three of the nine simulant samples: AN107, AN107 with entrained solids, and Pretreated AN107, which exhibited the greatest amount of foam. All the remaining simulants, AN104, AN104 with entrained solids, AN105, AN105 with entrained solids, AZ101, and AZ102, produced less foam (< 71%) and thus, were not tested under vacuum (see Table 1).

Mechanistically, foaming is particularly caused by precipitating solids and this process is encouraged by the presence of organic complexants. Maximum foaminess occurs when simulants’ total solids concentrations are greater than 50 wt. % (see Table 1), which is well beyond the saturation point for these simulants and beyond the desired concentration endpoint of the RPP evaporators. A review of the experiments conducted with Hanford tank sample 241-AN102 shows that foaming occurred at a very low total solids concentration (approx. 30 wt. %, 5 M Na), thus indicating that the mechanisms for foaming in the real waste may be much different than the mechanisms for the simulants.

A section of study was devoted to documenting the effects of separable organics, namely normal paraffin hydrocarbons (NPH) and tributyl phosphate (TBP), on LAW processes. The addition of 100 ppm of NPH did not relieve foaming, but increased it by 25%. On the contrary, the addition

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1 The designation AN105, AN107, etc., refers to a simulant designed to model the contents of Hanford waste tank 241-AN105, 241-AN107, etc. (A), (B), (C) refers to envelopes A, B, and C.
of 100 ppm of TBP did reduce foaminess by a factor of three. However, adding 300 ppm of TBP to the simulants resulted in an increase in foaminess, suggesting a deminimus concentration for TBP somewhere between 100 and 300 ppm.

Dow-Corning Pulp Aid 3472 Concentrate and Dow-Corning Antifoam 1520-US were both tested as possible antifoam candidates. The effectiveness of 1520-US was poor. In contrast, antifoam agent Pulp Aid 3472 greatly reduced foaminess at low pressures by a factor of three. However, it is suggested that Pulp Aid 3472 be tested at higher evaporation fluxes.2

Future antifoam testing for the RPP will include surveying relevant literature and investigating the effect of trace surfactants known to be present in the 241-AN102 sample. Information found from the literature search concerning organics in Hanford waste simulants will be exploited with the goal of designing a better simulant. Experimentation to determine the impact on foaminess and foam stability when trace quantities of separable organics (TBP and NPH) are added to the simulants will also be performed and thus allow the RPP to determine the deminimus concentration of separable organics for the LAW evaporator. Modifications to the small-scale experimental rigs used by IIT will be made to increase the vapor flux during testing. Additionally, the consultants will continue testing using three commercial antifoams and final recommendation of the most effective antifoam will be given. Testing with the recommended antifoam will be conducted using actual radioactive, pilot scale Hanford tank samples.

Based upon IIT testing, selected LAW simulants with the antifoam agent will undergo pre- and post-irradiation testing to determine whether or not such a process is destructive to the antifoam. Testing using selected simulants and recommended antifoams will be conducted at the RPP design-basis evaporation flux rate.3

Introduction and Background

The River Protection Project Waste Treatment Plant (RPP-WTP) has identified a reference process for the treatment of low activity waste (LAW) solutions and high-level waste (HLW) slurries.4 The reference process includes evaporation of LAW solutions and RPP-WTP recycle streams using forced circulation evaporators. The RPP-WTP is currently considering changing the design to also evaporate Envelope D HLW sludge stream and the associated supernate fraction. Since this design change is not yet finalized, the scope of design change has not been fully evaluated with respect to this task and will be considered in future efforts as requested by RPP-WTP R&T.

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Excessive foaming during the evaporation of Envelope C (241-AN102) has been identified by SRTC.\(^5\) Excessive foaming in waste evaporators can cause carryover of radionuclides and other waste constituents, which could result in the process condensate exceeding waste acceptance criteria for the 200 Area Effluent Treatment Plant. The antifoams used at Hanford and tested by SRTC in Part B1 are known to degrade and become inactive in high pH solutions.\(^6\) Hanford wastes have been known to foam during operation of the 242-A evaporator causing excessive down time and processing delays.\(^7\)

The DWPF and the SRS Alternative Salt Disposition programs have an ongoing antifoam research program.\(^8\) Researchers at SRTC in collaboration with Illinois Institute of Technology (IIT) have developed several antifoams for the DWPF\(^9\) and SRS Alternative Salt Disposition\(^10\) programs. The DWPF plans to deploy one of these antifoams in FY01. It is desirable for the RPP-WTP to leverage data and resources from the existing DOE antifoam development programs.

The RPP-WTP R&T organization has requested SRTC to establish a contract with Illinois Institute of Technology to characterize the foaminess of the Hanford waste and to develop and optimize the existing commercial antifoam technology for the River Protection Project Waste Treatment Plant LAW Evaporators. The test specification/task plan for this task is contained in references.\(^11,3\) This report discusses the results and conclusions of the FY01 testing conducted at the SRTC and IIT laboratories. Specifically, this report partially completes item III.B in the referenced test specification. Item III.B was not completed because a simulant for AW101 and AN102 was not available by the end of FY01.

The overall objectives of the RPP-WTP antifoam program are:

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**Evaluate the foaming tendencies of LAW Feed, LAW Melter Feed, and Eluate streams in the WTP conditions.** In these tests, SRTC/Illinois Institute of Technology (IIT) will determine the foaming tendencies in the LAW evaporator systems.

**Identify, through testing and analysis, a suitable antifoam reagent for use in plant**

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\(^5\) M. L. Crowder, C. L. Crawford, H. H. Saito and T. B. Calloway, Jr., Bench-Scale Evaporation of Large Hanford Envelope C Sample (Tank 241-AN102), WSRC-TR-2000-00469 (Draft), Westinghouse Savannah River Company, 12/15/00.


\(^7\) M. D. Guthrie, 95-1 242-A Evaporator Campaign Post Run Document, WHC-SD-WM-PE-055 Rev. 0, Westinghouse Hanford Company, 2/7/96. The 242-A Evaporator was shutdown over four times during the campaign. Antifoam addition was increased from 300 to 600 ppm to compensate for the excessive foaming.


evaporators. In these tests, SRTC/IIT will make an initial antifoam recommendation for the LAW evaporators.

Determine if tributyl phosphate (TBP) and normal paraffin hydrocarbons (NPH) create problems in evaporator operation and if so, at what concentration do these problems occur. In these tests, SRTC/IIT will determine the concentration of TBP and NPH that effect foaming in the evaporator.

To achieve these objectives, the RPP-WTP R&T organization via SRTC plan to conduct testing at beaker scale with simulants (this task and report), bench scale with simulants (with and without irradiation), bench scale with actual waste samples, and pilot scale testing.

The specific objectives of this portion of the work are:

- Identify the mechanism of foam formation and characterize the foamingness of selected Hanford waste simulants at various pressures. The specific intent will be to reduce the number of simulants (most of which will involve simulants of AN102) to be tested by future experiments.

- Characterize foamingness as a function of water evaporation flux rate.

- Characterize the foaminess of TBP and NPH with the intent to establish a deminimus concentration for future testing by SRTC.

- Characterize the performance of industry-recommended and available commercial antifoams.

**Experimental**

Nine Hanford simulants, which were previously developed and approved by the project, were prepared at SRS in accordance with the recipes outlined in reference.\(^{12}\) Pretreated AN107 was formulated from the AN107 recipe listed in reference 11 and was subjected to the Sr/Tru precipitation/filtration process by the SRTC Thermal Fluid Laboratory. All nine simulants underwent analytical testing at the SRTC Laboratory for cation/anion concentrations. The instrument used for cation testing was the Varian Vista AX Inductively Coupled Plasma Atomic Emission Spectrometer (ICP-AES). The samples were aqueous so no sample preparation was necessary. A calibration was performed using 1 mg/L, 5, mg/L and 10 mg/L standards with the cations of interest. A 10X, 100X, 1000X, and 10,000X dilution was performed by volume on the original sample to bring the analytes of interest into the calibration range. The instrument used for anion determination was the Dionex DX-500 Ion Chromatograph. No sample prep was necessary. A calibration of 1, 5, and 10 mg/L was performed using the anions of interest. Dilutions of 100X, 1000X, and 10,000X were performed by volume on the original sample to bring the analytes of interest into the calibration range.

The predicted and actual chemical composition values of the nine Hanford simulants were recorded and percent target values were plotted in Microsoft® Excel. The percent target range Na (ICP-ES) concentration in all simulants was between 98.0 and 104.4% of target. However, in some simulants, the concentrations of Al, Ca, and Si were less than 50% of target. Also, in some simulants, concentrations of F, PO₄, K, and Cr were over 100% of target. It is suspected that the cause of such error in ion concentrations (e.g. Al) is due to misbatching. Low Ca concentrations in the simulants are due to the lack of complexants in several of the simulants. For example it is now known that the trace amounts of complexants are present in the Envelope A and B waste tanks. It is likely that the high dilution factors used for the analyses increased the analytical error for these elements (Appendix B).

Simulant samples were tested for foaminess during boiling. Several experiments were conducted in a laboratory set-up as shown in Figure 1. A specially designed heating jacket was installed around the beaker, which helped to increase the water evaporation rate of boiling. A digital camera was installed to monitor the foaming characteristics at varied temperatures and pressures. The foaming tests were conducted to identify the mechanism of foam formation so that a more effective antifoam agent could be selected. Additionally, these tests were conducted to reduce the number of simulants to be tested by future experiments.

In order to understand the effects of TBP and NPH on foaminess, small concentrations of normal paraffin hydrocarbons (NPH) and tributyl phosphate (TBP) were separately added to simulant samples. Adding 25 ppm of NPH or TBP to the samples did not result in any notable effects on foaminess; therefore, research focused on using 100-ppm concentrations of NPH and TBP.

Using AN107, Pretreated AN107, and AN107 with entrained solids, the more severely foaming Hanford simulants, two commercial antifoams were selected for testing: Dow-Corning Antifoam 1520-US and Dow-Corning Pulp Aid 3472 Concentrate. 1400 ppm of the 1520-US antifoam was added to the samples prior to boiling as was Pulp Aid 3472 antifoam and the effectiveness of each was recorded (figures 28-33, Appendix A).

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13 Pretreated AN107 was excluded from the spreadsheet since no predicted values existed for this simulant.
14 1520-US antifoam is used at the Hanford 242-A evaporator. Pulp Aid 3472 antifoam, suggested by Dow, is used by paper industries for processes of high alkaline conditions.
Figure 1 – Experimental Set-Up To Study Foaminess Of Sludge Simulants During Boiling Using Vacuum
Results

There were essentially four main experimental studies carried out by IIT that focused upon the Hanford simulants AN105, AN105 with entrained solids, AN107, Pretreated AN107, AN107 with entrained solids, AZ101, AZ102 envelope B, AN104, and AN104 with entrained solids. These were (1) foaminess at atmospheric pressure and vacuum, (2) foaminess at various evaporation flux rates, (3) foaminess as a function of TBP and NPH concentrations, and (4) foaminess in the presence of a commercial antifoam. Table 1 gives a summary of the simulants’ characteristics and chemistry.

1) Less foam is produced at vacuum rather than at atmospheric pressure. For samples AN107 with entrained solids, AN107, and Pretreated AN107, Figures 16-18 in Appendix A show how lowering the pressure to which a simulant is exposed while boiling can significantly decrease foaminess. Experimental testing at standard atmospheric pressure also indicated that the amount of foam produced correlated to the amount of water evaporated during boiling. As sample volumes decreased over time from the evaporation of water, its total solids concentration increased linearly (Appendix A, figures 3a-11a) and in turn, foaming potential greatly increased (see figures 3b-11b in Appendix A).

2) Higher evaporation flux will result in higher foaminess. For tested simulants AN107 with entrained solids, AN107, and Pretreated AN107, data clearly indicate that a high water evaporation flux rate (0.09 ml/min cm²) will result in a greater amount of foaminess. For instance, when the flux rate was increased from 0.05 to 0.09 ml/min cm², the amount of foam drastically increased by an average factor of nearly 15 (see Appendix A, figures 19-21).

3) Studies concerning the concentration of TBP and NPH in Hanford simulants yielded contrasting results. The presence of NPH led to more foam while TBP acted as an antifoamer at low concentration and lead to more foam at high concentrations. Consequently, the greatest reductions in foaminess were observed with the addition of 100 ppm of TBP to simulants that were at a low pressure of 110 mmHg.

4) Use of the 1520-US antifoam did not result in a relatively significant foam reduction (see Appendix A, figures 28-30) whereas the Pulp Aid 3472 antifoam did indeed reduce the amount of foam produced during boiling by a factor of 3-4 (see Appendix A, figures 31-33). The evaporation flux rate was 0.04 ml/min cm² for experimental tests concerning 1520-US antifoam whereas the evaporation flux rate was 0.09 ml/min cm² for tests concerning Pulp Aid 3472 antifoam.
Table 1: Summary of Hanford Simulant Characteristics and Chemistry

<table>
<thead>
<tr>
<th>Name of Simulant</th>
<th>Weight of solids (g)</th>
<th>% TOC</th>
<th>% foaminess as a function of pressure(^b, c)</th>
<th>% foaminess as function of total solids conc. and Na molarity (at 0.04 ml/min cm(^2) and 760 mmHg)(^a)</th>
<th>pH(^d)</th>
<th>% NaNO(_3) / NaNO(_2) (^d)</th>
<th>Color(^d)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pretreated AN107</td>
<td>2.49</td>
<td>800</td>
<td>100</td>
<td>60 &gt; 9.7 800 14 3.5/1 Blackish Red</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AN107 with Entrained Solids</td>
<td>2.49</td>
<td>650</td>
<td>100</td>
<td>80 &gt; 11.7 633 12 3.5/1 Brown</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AN107 3M</td>
<td>2.49</td>
<td>700</td>
<td>100</td>
<td>70 &gt; 10.3 660 12 8/2.7 Brick Red</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AN104 with Entrained Solids</td>
<td>0.48</td>
<td>N/A</td>
<td>N/A</td>
<td>55 &gt; 8.6 70 14 4.4/3.3 Green</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AN104 3M</td>
<td>0.48</td>
<td>N/A</td>
<td>N/A</td>
<td>60 &gt; 10.1 35 14 6.3/4.3 Green</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AN105 3M</td>
<td>0.50</td>
<td>N/A</td>
<td>N/A</td>
<td>60 &gt; 9.8 50 14 6.3/4.3 Green</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AN105 3M with Entrained solids</td>
<td>0.50</td>
<td>N/A</td>
<td>N/A</td>
<td>60 &gt; 9.0 25 14 6.3/4.3 Yellow</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AZ101</td>
<td>0</td>
<td>N/A</td>
<td>N/A</td>
<td>65 &gt; 10.3 30 14 3.5/3.8 Yellow</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AZ102</td>
<td>0.30</td>
<td>N/A</td>
<td>N/A</td>
<td>55 &gt; 9.7 55 14 1.0/3.0 Yellow</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\(^a\) Data transcribed from graphs in Appendix A (figures 3b-11b)
\(^b\) At evaporation flux rate of 0.05 ml/min cm\(^2\)
\(^c\) “N/A” signifies that test was not conducted with simulant
\(^d\) Data copied from Table 1 in Appendix A
Mechanistically, it is suggested that foam stability is more dependent on physical (formation due to precipitating solids) rather than chemical attributes of the foam.\textsuperscript{15} For instance, foam collapsed shortly after a simulant was removed from the hot plate. Had surfactants been the primary constituent in a simulant’s foaming capability, the foam should have remained for a longer period of time. Moreover, maximum foaminess occurs when simulants’ total solids concentrations are greater than 50 wt. % (see Table 1), which is well beyond the saturation point for these simulants and beyond the desired concentration endpoint of the RPP evaporators. However, Pretreated AN107, AN107 with entrained solids, and AN107, all of which exhibited the greatest amount of foaming, contained a significantly greater concentration of organic complexants\textsuperscript{12} (see Table 1) than the other simulants, thus undermining a purely physical-based mechanism. Furthermore, a review of the experiments conducted with Hanford tank sample 241-AN102 shows that foaming occurred at a very low solids concentration (approx. 30 wt. %, 5 M Na), thus indicating that the mechanisms for foaming in the real waste may be very different than with simulants. Examination of the video results for the evaporation of Hanford tank sample 241-AN102 leads to the hypothesis that foaming in this sample may be due to surfactants that are present in the waste.

**Conclusion**

Studies were completed by SRTC/IIT that characterized the foaminess of Hanford simulants AN105 (A), AN105 with entrained solids (A), AN107 (C), AN107 with entrained solids (C), AZ101 (B), AZ102 (B), AN104 (A), AN104 with entrained solids (A), and Pretreated AN107 (C) at atmospheric and vacuum conditions. The following can be summarized from experimental results:

- Foaminess in all the Hanford waste simulants occurs due to the formation of particles (precipitated salt species) that occur when the solution reaches saturation. The precipitated salt particles stabilize the foam by preventing the liquid from draining through each foam lamella. The particles attach themselves to the bubble and create a stabilization barrier creating stable foam surfaces. Foam occurred in the 241-AN102 sample at a relatively low total solids content (31.5 wt. % total solids, <0.002 wt. % insoluble solids). Foams in the Hanford waste simulants only occurred after bulk saturation was reached (~ 40 wt. %) and significant quantities of salt particles began to form. Therefore, the mechanism for foam formation in the Hanford simulant is very different from the actual 241-AN102 tested by SRTC in the B1 phase. It is known that various salt species (e.g. sodium carbonate) experience retrograde solubility when the temperature is increased. Therefore, it is likely that the foaminess of Hanford waste simulants is related to the chemical composition and morphology of the particle formed in the solution.

- Lowering the pressure during the evaporation process can reduce foaminess. Table 1 compares the amount of foaminess at both atmospheric pressure and at a vacuum pressure of 110 mmHg and clearly shows that such a lower pressure will greatly reduce the amount of foaminess during boiling of the simulant.

\textsuperscript{15} See Appendix A (p. 13) for proposed mechanism of foam stability.
• The foaminess of all the simulants decreased significantly when evaporated under vacuum conditions and even more so with a low vapor flux rate of 0.05 ml/min cm² (see Table 1). Essentially, a higher vapor flux rates always increases foaming potential at a given solids concentration. Testing at the design basis vapor flux is needed to determine the foaminess under actual evaporator conditions.

• The addition of 25 ppm of normal paraffin hydrocarbons (NPH) and tributyl phosphate (TBP) did not affect the foaminess of the AN107 with entrained solids or the AN107 pretreated simulants. Higher additions (100 ppm) of TBP resulted in a decrease in foaminess by a factor of three. However, with additions of 300 ppm of TBP, the foaminess was increased. Foaminess may have increased due to the decomposition of TBP into organic and inorganic components. The decomposed organic component (dibutyl phosphate-DBP) may lead to an increase in foaminess of the simulant. Addition of 100 ppm of NPH increases the foaminess by 25%. Since the effects of TBP (and DBP) and NPH on foaminess are different and non-linear in nature, further work needs to be conducted to define a deminimus level for separable organics.

• Currently, Dow-Corning Pulp Aid 3472 Concentrate seems to be a much more effective antifoam agent than Dow-Corning Antifoam 1520-US. Even though Pulp Aid 3472 antifoam was tested at a higher flux rate than 1520-US, it still outperformed 1520-US and decreased foaminess by a factor of 3-4.

Future antifoam testing for the RPP will include surveying relevant literature and investigating the effect of trace surfactants known to be present in the 241-AN102 sample. Information found from the literature search concerning organics in Hanford waste simulants will be exploited with the goal of designing a better simulant. Experimentation to determine the impact on foaminess and foam stability when trace quantities of separable organics (TBP and NPH) are added to the simulants will also be performed and thus allow the RPP to determine the deminimus concentration of separable organics for the LAW evaporator. Modifications to the small-scale experimental rigs used by IIT will be made to increase the vapor flux during testing. Additionally, the consultants will continue testing using three commercial antifoams and final recommendation of the most effective antifoam will be given. Testing with the recommended antifoam will be conducted using actual radioactive, pilot scale Hanford tank samples.

Based upon IIT testing, selected LAW simulants with the antifoam agent will undergo pre- and post-irradiation testing to determine whether or not such a process is destructive to the antifoam. Testing using selected simulants and recommended antifoams will be conducted at the RPP design-basis evaporation flux rate.16

References

See footnotes.

Appendixes

Appendix A – IIT Final Report for FY01

Appendix B – Simulant Analytical Data
FINAL REPORT ON SUBCONTRACT NO: AC19845S

FOAMING IN HANFORD RPP-WTP LAW
EVAPORATION PROCESSES

By

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September 13, 2001
EXECUTIVE SUMMARY

A program was implemented at IIT to develop a basic understanding of the severe foaming experienced in Hanford River Protection Project Waste Treatment Plant (RPP-WTP) evaporation processes. A study, to understand foaminess, was carried out using simple experiments with nine Hanford sludge simulants. Foaminess and foam textures were monitored during boiling of the simulants. Results of foaminess of nine non-radioactive Hanford simulants are summarized as follows:

Foaminess during boiling of Hanford sludge simulants was tested at both atmospheric pressure (760 mm Hg) and at low pressure (110 mm Hg) (only for simulants showing severe foaminess). The foaminess of all the simulants goes to a maximum at higher solid concentration (>50 %). At atmospheric pressure (760 mm Hg) severe foaminess (>600%) was observed in sludge samples AN-107 with entrained solids, AN-107 3M and pretreated AN-107 at the maximum in foaminess. Less Foaminess at atmospheric pressure (760 mm Hg) was observed in sludge samples AN-104 3M, AN-104 with entrained solids, AN-105 3M with entrained solids, AN-105 3M, AZ-101 and AZ-102.

The increase in water evaporation flux rate enhances foaminess. Experiments were carried out at both atmospheric (760 mm Hg) and low pressure (110 mm Hg) to understand its effect on foaminess. Significantly less foaminess occurred at low pressure (110 mm Hg) and low water evaporation flux (0.05
ml/min sq.cm). However, at low pressure and high evaporation flux rate (0.09 ml/min sq.cm.) the maximum in foaminess was about 1200%. This shows that foaminess increases with increasing water evaporation flux. Extrapolated data for water evaporation flux (0.56 ml/min sq.cm) simulated at Hanford plant show severe foaminess (>1500%) at low solid concentration (<40 %).

The addition of 25 ppm of normal paraffin hydrocarbon (NPH) and tributyl phosphate (TBP) to the sludge samples with severe foaminess did not show reduction in foaminess. However, addition of 100 ppm of NPH increases foaminess at maximum by 25 % and addition of 100 ppm of TBP reduces foaminess by a factor of three at maximum. A detailed study of effect of TBP on foaminess needs to be done at higher water evaporation fluxes.

The antifoaming efficiency of two antifoamers (DOW PULPAID CONCENTRATE 3472 and DOW 1520 US) was tested. The antifoaming performance of DOW 1520 US was poor. Antifoamer DOW PULPAID CONCENTRATE 3472 reduces the foaminess at low pressures by a factor of three (at foaminess corresponding to maximum). However its antifoaming efficiency needs to be studied at higher evaporation fluxes.
INTRODUCTION

Hanford wastes have been known to foam during operation of the 242-A Evaporator causing excessive down time and processing delays\(^1\). The Savannah River Technology Center (SRTC) found that severe foaminess occurred during the continuous bench scale evaporation of the Envelope C tank (sludge sample 241-AN102) and during the small beaker evaporation of simulated 241-AN107 without SR/TRU pretreatment\(^2,3\). Foaming in waste evaporators can cause excessive carryover of radionuclides and non-radioactive waste constituents to the condensate system. The antifoamer (DOW-1520 US) used at Hanford and tested by SRTC is known to decompose and become inactive in high pH solutions\(^1\). Therefore, a fundamental research program was initiated by the IIT researchers (Nikolov and Wasan) with the three main objectives: 1) To develop a basic understanding of the mechanisms of foaminess in the RPP Evaporation process; 2) To identify the key phenomena which aggravate foaming; and 3) To identify a more effective means to eliminate or mitigate foaming.

\(^2\) M.L.Crowder, C.L.Crawford, H.H.Saito and T.B.Calloway, Jr., Bench Scale Evaporation of Large Hanford Envelope C (Tank 241-AN102), WSRC-TR- 2000-00469 (Draft), Westinghouse Savannah River Company, 12/15/00.
In the initial phase of the research program, nine non-radioactive simulants of Hanford salt solution prepared by SRTC were supplied to the IIT researchers to conduct foaming tests during boiling.

- Laboratory tests were conducted at both atmospheric (760 mm Hg) and below atmospheric pressure (110 mm Hg).
- Foaming tests were also conducted when trace quantities of separable organics such as tributyl phosphate (TBP) and normal paraffin hydrocarbons (NPH) were added to the simulants with entrained solids.
- The performance of the commercially available antifoamers DOW PULPAID CONCENTRATE 3472 and DOW-1520 US was tested (the Hanford 242-A evaporator currently uses DOW-1520 US antifoamer).

**EXPERIMENTAL**

Nine sludge samples simulating Hanford nuclear waste sludges were tested for foaminess during boiling. The foaming experiments were conducted in a laboratory set-up as shown in Figure 1. A specially designed heating jacket was installed around the beaker, which helped in increasing the water evaporation rate of the boiling sludge samples. The arrangement included specially designed fiber optics and a digital camera to monitor the degree of foaminess and the foam lamella texture at different temperatures and pressures (see Figure 1). The foaming tests were conducted to understand the foaming mechanisms, and to select the potential antifoamers. The foaming
tests were conducted at both atmospheric (760 mm Hg) and below atmospheric pressure (110 mm Hg). The set up for the experiments conducted below atmospheric pressure (110 mm Hg) is shown in Figure 2.

A 300 ml of the sludge sample was placed in a beaker with a diameter of about 9 cm and height 22 cm. The boiling temperature of the sludge at atmospheric pressure (760 mm Hg) was approximately 98 °C. Several minutes after the beaker was placed on the hot plate the sludge sample began to boil, typically after 10-12 minutes. To eliminate overheating of the sludge sample and creation of local hot spot areas, small amount of boiling chips were added to the sludge in the beaker. To homogenize the sludge sample in the beaker a magnetic stirrer was placed inside the beaker. The pH of the simulants was measured using a pH paper and the results are presented in Table 1.

In order to study the effect of the water evaporation flux on foaminess, the hot plate temperature and the efficiency of heating were controlled. The boiling water evaporation flux for each of the sludge sample was calculated from the measurement of the volume of water evaporated from the sludge in the known time interval. It is important to note that after stopping the boiling the foam collapses in 10-15 seconds. So foaminess was observed only during boiling of the simulant. The volume of liquid remaining in the sludge was measured by removing the sludge from the hot plate and allowing foam to collapse. The change in the liquid level during the known interval of time divided by the
evaporating area of the beaker gave the water evaporation rate for that simulant. During boiling the volume of foam starts increasing which is measured at regular interval. Without boiling the foam collapses and sludge volume was measured after the foam collapsed by removing the beaker from the hot plate. The difference in the volume with and without boiling gives the volume of foam during boiling. Foaminess was calculated as the percentage of foam volume to the stationary liquid volume.

To investigate the effect of sludge particle concentration on foaminess, approximately 15 g of sludge sample was taken in a plastic cup and weighed to measure the soluble solid concentration. The simulant contained both the insoluble as well as the soluble solids, which could be summed up as the total solids in the simulant. The cups were then kept at room temperature for evaporation and sludge samples were weighed after every 6 hrs. After about 48-60 hours the dried sludge samples were weighed and the percentage of total solid concentration of the simulant was calculated. The initial solid concentration was measured by taking the weight fraction of solids in the sludge sample to the total weight of the sludge sample containing both the liquid as well as the solids.

**Foaminess at atmospheric pressure (760 mm Hg)**

Foaming of sludge samples was studied at the atmospheric pressure (760 mm Hg). The heat input to the water evaporation rate was monitored by measuring
the temperature of the hot plate using a thermocouple. The heating temperature of the hot plate was kept at 400 ± 10°C with a flux rate of 0.04 ml/min sq.cm. During the boiling, water evaporated and the concentration of the total solids (soluble and insoluble solids) in the sludge increased. The rate of water evaporation and the foaminess were monitored for each sludge samples. Figures 3a to 11b show the effect of solid concentration on the foaminess and the evaporation rate of the simulant for experiments carried out at 400 °C. All the results reported in the figures 3a to 11b were conducted with the first batch of simulants. In order to investigate the effect of flux on foaminess, the temperature of the hot plate was increased from 400 to 450 °C. The plot showing the effect of water evaporation flux on foaminess is shown in figure 12. The evaporation flux rate for experiments carried out at 450 °C with the heating jacket was 0.08 ml/min sq. cm for the tested sludge samples. The comparative plots for the water evaporation rate from 0.04 to 0.08 ml/min sq.cm is shown in figures 13-15 for the simulants AN-107 with entrained solids, AN-107 3M and Pretreated AN-107. The results clearly show that as the water evaporation rate increases the foaminess also increases.

**Foaminess at low pressure (110 mm Hg)**

The effect of low pressure (110 mm Hg) on the foaminess of the Hanford simulants was also investigated. Figure 2 shows the experimental set-up used to achieve the low pressure. The experiments were carried out at below the
atmospheric pressure of 110 mm Hg, which was obtained by using a water pump to remove vapor from the headspace above the boiling surface. The beaker containing the sludge sample was covered with a specially made rubber enclosure to eliminate any leaks and a small pipe was connected to the water pump in order to achieve the below atmospheric pressure. Mercury manometers were used to calibrate the pressure transducers, which was then used to measure pressure below atmospheric. The foaminess and the foam texture of the sludge sample during boiling were monitored using the video camera. The water evaporation rate for the experiments carried out at 400 °C was 0.05 ml/min sq.cm for the tested sludge samples. Figures 16-18 show the effect of low pressure on foaminess for the simulants AN-107 with entrained solids, AN-107 3M and Pretreated AN-107. The experiments were carried out with the first batch of simulants. The result of all the three samples depicts that foaminess decreased significantly with decreasing of pressure e.g. for the simulant Pretreated AN-107 the maximum foaminess decreases by about 10 times. (See the movie clip).

**Foaminess at higher flux and low pressure**

A heating jacket was specially designed so that higher water evaporation flux value could be obtained. Using a camera and fiber optics the foaminess of the boiling sludge sample was monitored. Applying a low pressure (110 mm Hg) in conjunction with the heating jacket the water evaporation flux value could be increased from 0.05 ml/min sq.cm to 0.09 ml/min sq.cm for the tested
sludge samples. Figures 19-21 shows the effect of higher water evaporation rate on foaminess for the samples AN-107 with entrained solids, AN-107 3M and Pretreated AN-107.

In order to evaluate the foaminess at conditions consistent with Hanford water evaporation rate at about 0.56 ml/min sq.cm, we extrapolated the value of foaminess at a higher water evaporation rate value. Figures 22-24 show the extrapolated values of foaminess for the simulants AN-107 with Entrained Solids, AN-107 3M and Pretreated AN-107. Extrapolated values of foaminess show that for the conditions consistent with Hanford, the foaminess values are about 1500%.

**Foaminess with addition of TBP and NPH**

It is known in the literature that tributyl phosphate (TBP) helps in reducing the foaminess and acts as an antifoamer. So in order to understand the effect of TBP on foaminess, experiments were carried out using 25 ppm and 100 ppm of TBP with the sludge samples. By using 25 ppm of the TBP no notable effect on foam reduction was found and so our research focused on using 100 ppm of TBP in order to study the effect on foaminess (figures 25-27).

Experiments were also carried out to study the effect of normal paraffin hydrocarbons (NPH) on the foaminess of the boiling sludge sample. Trace quantities (25 ppm) of NPH were added to the boiling sludge samples. It was found that 25 ppm of the NPH did not show notable effect on foaminess and it
was decided that a minimum amount of NPH would be required in order to have any effect on the foaminess. So after a number of experiments 100 ppm of the NPH was found to show some effect on foaminess (Figures 25-27). In summary it was observed that low pressure (110 mm Hg) and 100 ppm TBP sufficiently reduces the maximum foaminess by a factor of three. From the figures we can conclude that TBP shifts the maximum in foaminess towards the higher concentration.

**Foaminess in the presence of antifoamer**

In the study of the foaminess of the boiling sludge samples, our observations clearly showed that at higher flux for both low pressure (110 mm Hg) and atmospheric pressure (760 mm Hg) foaminess of the simulants was very severe. To reduce foaminess two commercial antifoamers provided by SRTC were tested. The two commercial antifoamers studied were DOW-1520 US and the DOW PULPAID CONCENTRATE 3472. Due to the severe foaminess of the simulant, a large volume of the antifoamer such as 1400 ppm was added to the boiling sludge samples. In the experimental study we first added 1400 ppm of DOW-1520 US as an antifoamer to the sludge sample right from the start of the experiment and the effect of the antifoamer was monitored (Figures 28-30). The value of the water evaporation rate was 0.05 ml/min sq.cm. for these experiments. The antifoamer DOW PULPAID CONCENTRATE 3472 was tested at a higher vapor flux value of 0.09 ml/min sq.cm. (Figures 31-33). The antifoamer DOW PULPAID
CONCENTRATE 3472 reduces the maximum foaminess by a factor of about 3-4. (See figures 31-33, which compares the foaminess with and without the antifoamer)

**RESULTS AND DISCUSSION**

With the evaporation of water the solid concentration in the simulant increases, which was monitored along with the foaminess of the sludge samples. Figures 3a, 4a, 5a, 6a, 7a, 8a, 9a, 10a and 11a show the rate of change of sludge volume at a hot plate temperature of 400 °C and an atmospheric pressure of 760 mm Hg (water evaporation flux 0.04 ml/min sq.cm) for the sludge samples Pretreated AN-107, AN-107 with entrained solids, AN-107 3M, AN-104 with entrained solids, AN-104 3M, AN-105 3M, AN-105 3M with entrained solids, AZ-101 and AZ-102. The figures show that the rate of water evaporation is constant e.g. the sludge volume decreases linearly with time. As a result of the linear decrease in volume of the sludge the concentration of solids increases linearly with time. Figures 3b, 4b, 5b, 6b, 7b, 8b, 9b, 10b and 11b show the dependence of foaminess on the concentration of solids for the sludge samples pretreated AN-107, AN-107 with entrained solids, AN-107 3M, AN-104 with entrained solids, AN-104 3M, AN-105 3M, and AN-105 3M with entrained solids, AZ-101 and AZ-102.
For all the Hanford simulants during boiling we observed a maximum in foaminess with concentration. The data shown in the above figures clearly indicate that for the Hanford sludge samples, the maximum foaminess occurs at a more than 50 wt % solid concentration. Multiple tests were conducted to ensure the reproducibility of the trends in foaminess as a function of solid concentration (Figures 3b, 4b, 5b, 6b, 7b, 8b, 9b, 10b and 11b). We have observed that during the boiling experiments at low solid concentrations (less than 30 wt %) the foam cells are spherical in shape. The foam lamella was only a few millimeters in size and the lamella increases in size as the solid concentration increases. At higher solid concentrations, the foam lamella was several centimeters in size and its texture changed dramatically. It is important to note that the foam collapses shortly after the boiling sludge sample was removed from the hot plate, thereby suggesting that the foam stability is not due to the presence of surfactants, because in the presence of surface active materials the foam will be stable. Table 1 shows the Hanford Low Activity Waste chemistry, the type of the simulants, the compositions and the maximum of foaminess of the simulants that were studied.

Based on the results for foaminess during boiling, the Hanford sludge samples are categorized into two major categories i.e. sludge samples showing a higher foaminess (700-800 %) and sludge samples showing a lower foaminess (<100 %). The Hanford simulants pretreated AN-107, AN-107 with entrained solids and AN-107 3M showed a higher degree of foaminess (Figures 3b, 4b, 5b) while AN-104 with entrained solids, AN-104 3M, AN-
105 3M, and AN-105 3M with entrained solids, AZ-101 and AZ-102 showed a lower degree of foaminess. Our subsequent tests focussed on the three sludge samples, which showed higher foaminess.

The mechanism of stability of the foam lamella in sludge samples with severe foaming is discussed below.

During boiling of the sludge sample, the water evaporates and the sludge is concentrated. At a certain temperature and concentration, the sludge becomes even over saturated and fine crystals or nuclei are formed. These fine crystals (solids) attach to the bubble surfaces generated during boiling. The process of crystal formation was also observed on the walls/surface of the glass beaker. The crystals (solids attached to the bubble surface) provide a structural stabilization barrier, which prevents adjacent bubbles from coalescing, and, thereby, stabilizes the foam lamella and increases the foaminess. When the boiling is stopped the particles drain from the foam lamella and the foam lamella collapses. This mechanism of foam stabilization formed by the structural stabilization barrier was discussed in our previous report using the acidified SRTC sludge with PHA⁴. The main difference between the Hanford and the STRE sludge samples is that the Hanford simulants before boiling contain much less insoluble solids (≦< 1 wt %) while the SRTC acidified sludge contained 7-10 wt % insoluble solids. As a result of the high concentration of insoluble solids in the SRTC acidified sludge, the foaminess started when the sludge sample was concentrated to about 15 wt % solids
during boiling, and had a maximum at about 18-23 wt % solids. Based on this difference in the initial concentration of insoluble solids, one can expect the foaminess for the Hanford simulants to begin when the concentration of the insoluble solids (crystals) becomes higher than 15 wt %. So the concentration of the total solids (crystals and soluble solids) becomes higher than about 40 wt %. The maximum in foaminess was visually observed when the solid concentration is about 50-60 %, e.g. for the case AN-107 with entrained solids the maximum occurs at solid concentration at 75% while for the AN-107 3M it occurs 70 % and for pretreated AN-107 it occurs at 60 %. The reason for maximum in foaminess is due to the competition of two major phenomena: the structural stabilization barrier and the flocculation (attraction-depletion) mechanisms. At higher concentration of particles (crystals) due to the flocculation (aggregation), particles cannot form the structural stabilization barrier.

In our future study, we need to investigate the factors like boiling temperature and solid concentration of the sludge, which affect the nucleation and crystal growth processes (particle formation), as well as factors such as crystal size and morphology, surface active agents and sludge composition on foam lamella stability and foaminess.
Effect of Water evaporation flux on foaminess

To study the effect of water evaporation flux on foaminess of the sludge samples AN-107 with entrained solids, experiments were carried out at two different temperatures i.e. 400 °C and 450 °C. The flux rate for the experiment where the temperature was increased from 400 to 450 °C was 0.06 ml/min sq.cm. From Figure 12 it is clear that at higher water evaporation flux the foaminess at a fixed solid concentration increases significantly. The maximum in foaminess at higher water evaporation flux occurs at a slightly lower solid concentration. Therefore, we can conclude higher foaminess is observed at higher water evaporation rate. In the following section we discuss the effect of pressure on foaminess, because the water evaporation flux also depends on pressure.

Effect of water evaporation rate on foaminess at atmospheric pressure (760 mm Hg)

The experiments were carried out over a range of 0.04 ml/min sq.cm to 0.08 ml/min sq.cm in order to study the effect of water evaporation flux on foaminess at atmospheric pressure. The water evaporation flux rate at 400 °C was 0.04 ml/min sq.cm at atmospheric pressure conditions (760 mm Hg) and increased to 0.08 ml/min sq.cm at 450 °C with the heating jacket. Figures 13, 14 and 15 show the effect of the water evaporation rate on the foaminess
for the sludge AN-107 with Entrained solids, AN-107 3M and Pretreated AN-107 for the experiments carried out at 760 mm Hg. The three sludge samples show the increase in foaminess with the increase in water evaporation rate at atmospheric pressure (boiling point 98 °C). The reason for this increase is the decrease in surface tension with the increase in the temperature and more bubbles are generated, and consequently foaminess increases.

**Effect of water evaporation rate on foaminess at low pressure (110 mm Hg)**

The effect of low pressure (110 mm Hg) on foaminess was studied for the three sludge samples which showed severe foaminess namely AN-107 with entrained solids, AN-107 3M and pretreated AN-107 (Figures 16-18). The water evaporation rate was at a constant value of 0.04 ml/min sq.cm for these experiments. It is evident that pressure below the atmospheric conditions (110 mm Hg) reduces the foaminess for AN-107 with entrained solids, AN-107 3M and pretreated AN-107 to a large extent (figures 34-36). For the case of AN-107 3M the low pressure reduces the foaminess by a factor of 10 (e.g., see the movie clip for the effect of pressure on foaminess for the simulant pretreated AN-107). At low pressure the solubility of the sludge solids (particles) is higher. Due to this the concentration of sludge solids in the simulant is lower and hence foaminess of the simulant is much lower.
Our study also investigated the foam lamella structure and its size in these experiments. As the pressure is decreased the foaminess decreases dramatically (Figures 16-18). We observed that in the case when initial boiling experiment of the sludge was conducted at atmospheric pressure, and then the pressure was decreased for a while, the foaminess increased dramatically, and this effect is due to the gas solubility in sludge simulant (see video clip). It was seen that at low pressure the foam lamella was of the order of a few centimeters, was highly unstable and it collapsed very quickly. At higher pressure the gas solubility is high, more gas is contained in the sludge and when the pressure is reduced from high to low the gas solubility decreases and for a short period of time many bubbles are generated.

**Effect of low pressure (110 mm Hg) and higher vapor flux on foaminess**

The experiments were carried out over a range of 0.05 ml/min sq.cm to 0.09 ml/min sq.cm in order to characterize the effect of water evaporation rate on foaminess. The water evaporation rate at 400 °C and low pressure was 0.05 ml/min sq.cm and it increases to 0.09 ml/min sq.cm at 450 °C with the heating jacket at low pressures. Figures 19-21 shows the effect of the water evaporation rate on the foaminess for the sludge AN-107 with entrained solids, AN-107 3M and pretreated AN-107 for the conditions of 110 mm Hg. It is evident that with the increase in the water evaporation rate the foaminess also increased. At low pressure and low evaporation flux the solubility of the
sludge solids (less particles are formed) is higher and so foaminess of the sludge simulant is lower. But by increasing the water evaporation rate (increasing the boiling temperature) and the solid particles, more gas is generated and more bubbles are formed. Due to this, the particles attach themselves to the bubble and create the stabilization barrier creating stable foam and one can expect higher foaminess with the increase in water evaporation rate. In summary, at low pressures and high evaporation fluxes foaminess is higher.

In order to simulate the operation conditions (low pressure and high water evaporation flux) at Hanford it was necessary to extrapolate the data for the water evaporation rate value of 0.56 ml/min sq.cm. and at total solid concentration level of 40-50%. We extrapolated our data for foaminess as a function of evaporation flux (0.56 ml/min sq.cm.) for the case of 40 wt % solids using a LINEAR relation (Figures 22-24). From the calculations the foaminess was in the range of 1500%.

**Effect of Hydrocarbons and tributyl phosphate on foaminess**

A study was carried out to check the impact of normal paraffin hydrocarbon (NPH) and tributyl phosphate (TBP) on foaminess of the Hanford sludge samples. It was noted that addition of 25 ppm concentration of the NPH and TBP each did not affect the foaminess (Figures 34-36). To see the effect of 100 ppm of TBP and NPH each on foaminess experimental runs were carried
out with the simulant. The results for the degree of foaminess during boiling of the sludge samples with and without the NPH and TBP each are shown in the figures 25-27. It is evident that in the presence of the hydrocarbons the foaminess increases by about 25%. From the figures 25-27 it is evident that by adding 100 ppm of TBP the foaminess decreases by a factor of three for the simulants AN-107 with entrained solids, AN-107 3M and Pretreated AN-107.

In summary, with the addition of 100 ppm of NPH the foaminess increases by 25% at the concentration corresponding to a maximum in foaminess. However, with the addition of 100 ppm of TBP the maximum foaminess decreases by a factor of three and shifts the maximum in the foaminess to a higher solid concentration.

In order to understand the effect of TBP on foaminess, three different levels (50, 100 and 300 ppm) of TBP were added. The results (Figure 37) show that with 50 and 100 ppm concentration of TBP foaminess decreases, however with the addition of 300 ppm foaminess increases. This kind of effect on foaminess as a function of solid concentration needs a further study. A possible explanation of the increase in foaminess with the concentration of TBP is due to the decomposition of TBP into organic and inorganic components. The decomposed organic component of TBP leads to an increase in the foaminess of the simulant.
Effect of Antifoamer on foaminess

The efficiency of two commercial antifoamers (DOW-1520 US and DOW PULPAID CONCENTRATE 3472) on the Hanford higher foaminess sludge samples were tested. Due to the severe foaminess of the simulants a large volume of antifoamer (1400 ppm) was added to the simulants before boiling. The results for the antifoamer DOW 1520 US performance are as shown in the Figures 28-30 for the sludge samples AN-107 with entrained solids, AN-107 3M and pretreated AN-107, respectively. It was observed that for the sludge samples AN-107 with entrained solids and AN-107 3M, the antifoamer did show a small change in reducing the degree of foaminess. The maximum of foaminess occurred at a lower solid concentration, than for experiments performed without the antifoamer. The antifoamer did have some effect on the pretreated AN-107 sludge sample. The maximum of foaminess decreased from 800% to 700% for the pretreated AN-107 sludge sample and it showed a shift towards the higher solid concentration. Overall, it can be concluded that the antifoamer did not show good antifoaming performance.

The other commercial antifoamer (DOW PULPAID CONCENTRATE 3472) provided by SRTC was tested. Experiments were conducted by adding 1400 ppm of the antifoamer to the sludge simulant before boiling. Figures 31-33 show the effect of the antifoamer on the foaminess. The maximum in foaminess is about 300% for the case of the AN-107 with entrained solids
(Figure 31). Although the antifoamer reduced the foaminess of the simulant sufficiently further testing has to be done at higher water evaporation flux value where the foaminess is expected to be higher. So, either the concentration of the antifoamer may have to be increased or a new commercial antifoamer may have to be tested.

RECOMMENDATIONS

The following additional experiments need to be performed to better understand foaminess in the Hanford simulants.

- It was observed that, at a higher water evaporation rate, the foaminess increased. Studies have to be carried out for the Hanford simulants at a rate at least twice or three times the present water evaporation rate of 0.09 ml/min sq.cm. This study requires a new design of the sludge heating elements and use of a special heating jacket in order to achieve higher evaporation rates.

- It is seen that 1400 ppm of the commercial antifoamer DOW PULPAID CONCENTRATE 3472 reduces the maximum foaminess upto three times. However the maximum in foaminess for the boiling sludge with the antifoamer is 300 %. Studies need to be carried out at higher concentration
levels of antifoamer to efficiently reduce foaminess. However it has to be determined whether this antifoamer is efficient at higher evaporation fluxes.

- In the case of sludge simulant the foaminess is due to the particle structural stabilization barrier. To reduce the foaminess the particle surface properties (interactions between particles) have to be modified as was done for the case of acidified SRTC sludge with PHA.

- The commercial antifoamers, which have been tested, normally perform only when the foaming is caused by surfactants. Foaming in the Hanford simulants is caused by sludge solids (particles). So we need to select a commercial antifoamer which will reduce structural barrier caused by the particles and not by the surfactants.

- Microscopic properties of the particles (surface size and biphilicity) in the simulant under low pressure conditions need to be studied. It is necessary to understand the effects of nucleation, crystal growth, size and shape of the particles on foam lamella stability and foaminess.

- Due to the unavailability of an AN102 simulant, testing was not completed. Since the radioactive AN102 sludge sample tested by SRTC shows a higher degree of foaminess, studies with an AN102 simulant need to be completed before recommending final antifoam to the RPP-WTP.
Studies need to be performed using the mixed simulants such as the mixture of envelope A and envelope C simulants and mixtures of RPP recycles with LAW simulants.
Table 1: Hanford Low Activity Waste Chemistry

<table>
<thead>
<tr>
<th>Name of simulant</th>
<th>Weight of solids (g)</th>
<th>Total weight of solids (g)</th>
<th>Foaminess %</th>
<th>pH</th>
<th>Color</th>
<th>Hydroxide %</th>
<th>Acids %</th>
<th>% NaNO$_3$ / NaNO$_2$</th>
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<tbody>
<tr>
<td>Pretreated AN-107</td>
<td>115</td>
<td>857</td>
<td>4620</td>
<td>825</td>
<td>14 Blackish Red</td>
<td>1.5/0</td>
<td>---</td>
<td>3.5/1</td>
</tr>
<tr>
<td>AN-107 with Entrained Solids</td>
<td>115</td>
<td>857</td>
<td>4620</td>
<td>750</td>
<td>12 Brown</td>
<td>1.5/0</td>
<td>---</td>
<td>3.5/1</td>
</tr>
<tr>
<td>AN-107 3M</td>
<td>115</td>
<td>857</td>
<td>4620</td>
<td>725</td>
<td>12 Brick Red</td>
<td>1.0/0</td>
<td>2-Glycolic acid 1.2 Citric acid</td>
<td>8/2.7</td>
</tr>
<tr>
<td>AN-104 with Entrained Solids</td>
<td>22</td>
<td>845</td>
<td>4612</td>
<td>75</td>
<td>14 Green</td>
<td>4/2.3</td>
<td>---</td>
<td>4.4/3.3</td>
</tr>
<tr>
<td>AN-104 3M</td>
<td>22</td>
<td>845</td>
<td>4612</td>
<td>75</td>
<td>14 Green</td>
<td>4/2.3</td>
<td>---</td>
<td>6.3/4.3</td>
</tr>
<tr>
<td>AN-105 3M</td>
<td>23</td>
<td>892</td>
<td>4612</td>
<td>55</td>
<td>14 Green</td>
<td>3.2/2.2</td>
<td>---</td>
<td>6.3/4.3</td>
</tr>
<tr>
<td>AN-105 3M with Entrained solids</td>
<td>23</td>
<td>892</td>
<td>4612</td>
<td>35</td>
<td>14 Yellow</td>
<td>3.2/2.2</td>
<td>---</td>
<td>6.3/4.3</td>
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<tr>
<td>AZ-101</td>
<td>0</td>
<td>592</td>
<td>4396</td>
<td>40</td>
<td>14 Yellow</td>
<td>1.5/1.2</td>
<td>---</td>
<td>3.5/3.8</td>
</tr>
<tr>
<td>AZ-102</td>
<td>13</td>
<td>517</td>
<td>4396</td>
<td>45</td>
<td>14 Yellow</td>
<td>0.8/0.1</td>
<td>---</td>
<td>1.0/3.0</td>
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Figure 1  EXPERIMENTAL SET-UP TO MONITOR LAMELLA TEXTUREDURING SLUDGE BOILING
Figure 2 EXPERIMENTAL SET-UP TO STUDY FOAMINESS OF SLUDGE SIMULANTS DURING BOILING USING VACUUM
PRE TREATED AN-107 (ENVELOPE C)  
(FIRST BATCH OF SIMULANTS)

Hot Plate temperature 400 °C  
Flux-0.04 ml/min sq.cm  
Atmospheric pressure (760 mm Hg)

Figure 3a: Water evaporation flux with time

Initial Volume 300 ml

Run 1

Run 2
Figure 3b: % Foaminess vs Solid Concentration

Flux-0.04 ml/min sq.cm
Atmospheric pressure (760 mm Hg)

Run 1  Run 2
AN-107 with Entrained Solids (ENVELOPE C)
(FIRST BATCH OF SIMULANTS)

Initial Volume
300ml

Flux 0.04 ml/min sq.cm
Atmospheric pressure (760 mm Hg)

Figure 4a: Water evaporation flux with time
AN-107 with Entrained Solids (ENVELOPE C)  
(FIRST BATCH OF SIMULANTS)

Flux-0.04 ml/min sq.cm.  
Atmospheric pressure (760 mm Hg)

Figure 4b: % Foaminess vs Solid Concentration
AN-107 3M (ENVELOPE C)  
(FIRST BATCH OF SIMULANTS)

Flux-0.04 ml/min sq.cm.  
Atmospheric pressure (760 mm Hg)

Initial Volume 300 ml

Figure 5a: Water evaporation flux with time
AN-107 3M (ENVELOPE C)  
(FIRST BATCH OF SIMULANTS)

Flux-0.04 ml/min sq.cm  
Atmospheric pressure (760 mm Hg)

Figure 5b: % Foaminess vs Solid Concentration
AN-104 with Entrained Solids
(FIRST BATCH OF SIMULANTS)

Flux-0.04 ml/min sq.cm
Atmospheric pressure (760 mm Hg)

Initial Volume
300 ml

Run 1
Run 2

Figure 6a: Water evaporation flux with time
AN-104 with Entrained Solids
(FIRST BATCH OF SIMULANTS)

Flux-0.04 ml/min sq.cm.
Atmospheric pressure (760 mm Hg)

Figure 6b: % Foaminess vs Solid Concentration
Figure 7a: Water evaporation flux with time

AN-104 3M

(FIRST BATCH OF SIMULANTS)

Initial Volume
300 ml

Flux-0.04 ml/min sq.cm.
Atmospheric pressure (760 mm Hg)

Run 1
Run 2

Volume of sample (ml)

Time (min)

0 20 40 60 80 100 120 140
AN-104 3M
(FIRST BATCH OF SIMULANTS)

Flux-0.04 ml/min sq.cm.
Atmospheric pressure (760 mm Hg)

Figure 7b: % Foaminess vs Solid Concentration
AN-105 3M (ENVELOPE A)
(FIRST BATCH OF SIMULANTS)

Flux-0.04 ml/min sq.cm.
Atmospheric pressure (760 mm Hg)

Initial volume
300 ml

Run 1

Run 2

Figure 8a: Water evaporation flux with time
AN-105 3M (ENVELOPE A)
(FIRST BATCH OF SIMULANTS)

Flux-0.04 ml/min sq.cm.
Atmospheric pressure (760 mm Hg)

Solid Concentration (wt %)

% Foaminess

Figure 8b: % Foaminess vs Solid Concentration
AN-105 3M with Entrained Solids (ENVELOPE A)  
(FIRST BATCH OF SIMULANTS)

Flux-0.04 ml/min sq.cm.  
Atmospheric pressure (760 mm Hg)

Initial volume  
300 ml

Run 1
Run 2

Figure 9a: Water evaporation flux with time
AN-105 3M with ENTRAINED SOLIDS (ENVELOPE A)  
(FIRST BATCH OF SIMULANTS)

Flux-0.04 ml/min sq.cm.  
Atmospheric pressure (760 mm Hg)

Figure 9b: % Foaminess vs Solid Concentration
AZ-101 (ENVELOPE B)
(FIRST BATCH OF SIMULANTS)

Flux-0.04 ml/min sq.cm.
Atmospheric pressure (760 mm Hg)

Initial Volume
300ml

Run 1
Run 2

Figure 10a: Water evaporation flux with time
AZ-101 (ENVELOPE B)  
(FIRST BATCH OF SIMULANTS)

Flux-0.04 ml/min sq.cm.  
Atmospheric pressure (760 mm Hg)

% Foaminess vs Solid Concentration (wt%)

Figure 10b: % Foaminess vs Solid Concentration
Figure 11a: Water evaporation flux with time

Initial volume: 300 ml
Atmospheric pressure: 760 mm Hg
Flux: 0.04 ml/min sq.cm.
Figure 11b: % Foaminess vs Solid Concentration

AZ-102 (ENVELOPE B)
(FIRST BATCH OF SIMULANTS)

 Flux-0.04 ml/min sq.cm.
Atmospheric pressure (760 mm Hg)

% Foaminess

Solid Concentration (wt%)

Run 1
Run 2
AN-107 with Entrained Solids (ENVELOPE C)
(FIRST BATCH OF SIMULANTS)

Atmospheric pressure (760 mm Hg)

% Foaminess

Solid Concentration (wt%)

Figure 12: Effect of water evaporation flux on foaminess
at atmospheric pressure (760 mm Hg)
Figure 13: Effect of water evaporation flux on foaminess at atmospheric pressure (760 mm Hg)
Figure 14: Effect of water evaporation flux on foaminess at atmospheric pressure (760 mm Hg)
Figure 15: Effect of water evaporation flux on foaminess at atmospheric pressure (760 mm Hg)
AN-107 with Entrained Solids (ENVELOPE C)
(FIRST BATCH OF SIMULANTS)

![Graph showing the effect of pressure on foaminess at water evaporation flux (0.05 ml/min sq.cm)]

Atmospheric Pressure
(760 mm Hg)

Low pressure
(110 mm Hg)

Figure 16: Effect of pressure on foaminess at water evaporation flux (0.05 ml/min sq.cm)
Figure 17: Effect of pressure on foaminess at water evaporation flux (0.05 ml/min sq.cm.)
Figure 18: Effect of pressure on foaminess at water evaporation flux (0.05 ml/min sq.cm)
AN-107 with Entrained Solids (ENVELOPE C)
(SECOND BATCH OF SIMULANTS)

Figure 19: Effect of water evaporation flux on foaminess at low pressure (110 mm Hg)
Figure 20: Effect of water evaporation flux on foaminess at low pressure (110 mm Hg)
Figure 21: Effect of water evaporation flux on foaminess at low pressure (110 mm Hg)
AN-107 with Entrained Solids (ENVELOPE C)
(SECOND BATCH OF SIMULANTS)

Figure 22: Predicted foaminess at SRTC's water evaporation flux pressure 110 mm Hg)
Figure 23: Predicted foaminess at SRTC's water evaporation flux (pressure 110 mm Hg)
Figure 24: Predicted foaminess at SRTC's water evaporation flux (pressure 110 mm Hg)
Figure 25: Effect of NPH (100 ppm) and TBP (100 ppm) on foaminess at pressure (110 mm Hg) and water evaporation flux (0.09 ml/min sq.cm)
Figure 26: Effect of NPH (100 ppm) and TBP (100 ppm) on foaminess at pressure (110 mm Hg) and water evaporation flux (0.09 ml/min sq.cm)
PRE TREATED AN-107 (ENVELOPE C)  
(SECOND BATCH OF SIMULANTS)

Figure 27: Effect of NPH (100 ppm) and TBP (100 ppm) on foaminess at pressure (110 mm Hg) and water evaporation flux (0.09 ml/min)
Figure 28: Effect of antifoamer (DOW 1520 US, 1400 ppm) on foaminess at atmospheric pressure (760 mm Hg) and water evaporation flux (0.04 ml/min sq.cm.)
Figure 29: Effect of antifoamer (DOW 1520 US, 1400 ppm) on foaminess at atmospheric pressure (760 mm Hg) and water evaporation flux (0.04 ml/min sq.cm)
Figure 30: Effect of antifoamer (DOW 1520 US, 1400 ppm) on foaminess at atmospheric pressure (760 mm Hg) and water evaporation flux (0.04 ml/min sq.cm.)
Figure 31: Effect of antifoamer (DOW PULPAID CONCENTRATE 3472, 1400 ppm) on foaminess at pressure (110 mm Hg) and water evaporation flux (0.09 ml/min sq.cm)
Figure 32: Effect of antifoamer (DOW PULPAID CONCENTRATE 3472, 1400 ppm) on foaminess at pressure (110 mm Hg) and water evaporation flux (0.09 ml/min sq.cm)
PRE TREATED AN-107 (ENVELOPE C)  
(SECOND BATCH OF SIMULANTS)

Figure 33: Effect of antifoamer (DOW PULPAID CONCENTRATE 3472, 1400 ppm) on foaminess at pressure (110 mm Hg) and water evaporation flux (0.09 ml/min sq.cm)
Figure 34: Effect of TBP concentration on foaminess at low pressure (110 mm Hg) and water evaporation flux (0.09 ml/min sq.cm)
Figure 35: Effect of NPH (25 ppm) and TBP (25 ppm) on foaminess at atmospheric pressure (760mm Hg) and water evaporation flux (0.04 ml/min sq.cm)
Figure 36: Effect of NPH (25 ppm) and TBP (25 ppm) on foaminess at atmospheric pressure (760 mm Hg) and water evaporation flux (0.04 ml/min sq.cm)
Figure 37: Effect of NPH (25 ppm) and TBP (25 ppm) on foaminess at pressure (760 mm Hg) and water evaporation flux (0.04 ml/min sq.cm)
## Appendix B. Hanford Simulants Analytical Spreadsheet

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- Reported values that are inaccurate because they are below the instrument's detection limit. Thus, no % target values given for these.
- For predicted values, this color indicates the analyte was not added.
- Significant (> 50%) discrepancies between reported analytical value and predicted values.
### Reported Data, Predicted Data, and % Target Values (continued)

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- **Reported values that are inaccurate because they are below the instrument's detection limit. Thus, no % target values given for these.**

- **For predicted values, this color indicates the analyte was not added.**

- **Significant (> 50%) discrepancies between reported analytical value and predicted values.**