Impact of Precipitation Parameters on the Specific Surface Area of PuO₂

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ABSTRACT: Controlling the properties of PuO₂ through processing is of vital importance to environmental transport and fate, production of nuclear fuels, nuclear forensic analyses, stockpile stewardship, and storage of nuclear wastes applications. A number of processing conditions have been identified to control final product properties, including specific surface area (SSA), residual carbon content, adsorption of volatile species, morphology, and particle size. In this paper, a novel approach is developed for the prediction of PuO₂ SSA via the synthetic route of Pu(IV) oxalate precipitation followed by calcination. The proposed model utilizes multivariate regression methodology and leave one out formalism to link Savannah River Site (SRS) precipitation and calcination production data to the SSA of the final product. A comparison among the models provides insight into the accuracy and ability to identify variations amongst the processing data. Additionally, the models may also be used to fit new data outside of the parameters explored in a production facility. Finally, the trained model was compared to a similarly trained conventional model form to illustrate the influence of precipitation parameters on the prediction of the final SSA. The models presented here attempt to provide new methods for more accurate prediction of the PuO₂ product properties in a production scale environment for key environmental and nuclear applications.

INTRODUCTION

For decades, plutonium(IV) oxide has been studied in both laboratory and industrial scale settings to understand how the physical characteristics and chemical stability are controlled by changes in process parameters of the production process.¹ ⁷ Several dependencies have been linked to the different chemical preparation routes, for example, peroxide, oxalate, hydroxide precipitation, or metal oxidation. However, the most commonly studied synthetic route via Pu(III/IV) oxalate intermediate precipitation produces diverse final oxide characteristics depending on the processing conditions.⁸ In general, the mean particle size, shape, and morphology are controlled by the precipitation process parameters, whereas the specific surface area (SSA), tap density, residual moisture content, and carbon content are functions of the calcination process.⁹ ¹⁰ While these are the dominant processing conditions affecting the final oxide product, insights into the interplay between the two remain incomplete due to the wide parameter space among the processing units. For example, a variance in the SSA of 5–15 m²/g is seen at a calcination temperature of 650 °C, with wider window occurring at lower calcination temperatures. Indeed, when analyzing processing conditions outside of comparing calcination temperature to the SSA specifically, it appears that no dependency exists in either the literature or using known data.⁵ ⁷ ⁸ ¹⁰

Recent work which explored process dependencies between the PuO₂ product characteristics and the preparation method has analyzed the effect of adding calcination time to known temperature data to produce a complex correlation using the two variables.¹ ² The data were curve fit using a non-linear regression function over a wide temperature range and suggested a time dependency with the SSA around 2.5–3 h. A subset of the data was further analyzed using a narrower temperature range more in-line with industrial-scale processing parameters and showed minimal dependence on the time to predict the SSA. Orr et al. then developed an exponential function to fit known experimental SSA data based on temperature alone with 95% predictive accuracy.¹ However, the SSA distribution is still wide especially at temperatures below 650 °C, suggesting that additional process-dependent conditions could exist and be traced to the final PuO₂ product. In other words, perhaps there are unidentified interdependencies that can be extracted beyond evaluating processing conditions associated with one unit operation (such as calcination).

Received: September 8, 2021
Accepted: October 27, 2021
Published: December 29, 2021
Since our basic premise is that other processing parameters may account for variability in the SSA within a small calcination temperature window, we turned to data-driven regression modeling as a starting point for analysis. Although less accurate compared to machine learning models, data-driven regression models provide an interpretability not available with some machine learning algorithms.\textsuperscript{11} Furthermore, data-driven regression models utilize some amount of domain knowledge in their creation because the parameters tend to be defined in physical terms versus machine learning models.\textsuperscript{12,13} Due to this interpretability, it is possible to use data-driven models to see the influence of individual parameters by comparing their associated weighting values. The largest difference between the two model types exists in the amount of training data required to obtain an accurate prediction. For good machine learning algorithms, the minimum amount of training data required can vary from hundreds to thousands of data points while for more simple regression models only require tens of data.\textsuperscript{14} It can be easily stated that the required minimum amount of training data will change based on application, scope, and/or modeling algorithm. Ultimately though, the trend that more data lead to a more accurate model exists for both model types.

Herein, we develop a data-driven model capable of utilizing PuO$_2$ processing parameters to predict the SSA. Our particular interest is to evaluate industrial-scale processing data during the precipitation and calcination of Pu(IV) oxalate, since the correlation may be more complex than at the laboratory scale. Results suggest that while the PuO$_2$ SSA is strongly correlated to the calcination processing conditions, other parameters can cause variations in the SSA from the value predicted using the conventional quasi-exponential temperature model. These variations can provide insights into the large variability of SSA at a small calcination temperature window, thus increasing the predictive capability of calcination models.

## MOTIVATION AND METHODOLOGY

From 2014 to 2018, a mission at SRS was to process plutonium metal to produce PuO$_2$ for feed to the MOX Fuel Fabrication Facility (MFFF) with a targeted production rate of 1 metric ton per yr.\textsuperscript{15,16} The flowsheet proceeded as follows: After dissolution of the Pu metal in 8 M nitric acid, the solution was then purified via anion exchange, precipitated as Pu(IV) oxalate, and calcined at $\sim$650 °C for at least 4.5 h to form PuO$_2$. The Pu(IV) oxalate was precipitated using the direct strike batch method or adding 0.9 M oxalic acid to a Pu(IV) nitrate solution at 55 °C. An excess of 0.1 M oxalic acid and target nitric acid concentration between 1.5 and 3 M was necessary to minimize Pu losses and to produce a more dense PuO$_2$ product.\textsuperscript{15,16} In addition to tracking trace element impurities throughout the process, the physical characteristics of the oxide powders, such as bulk and tap density, SSA, particle size, and moisture content were also measured. Table 1 shows the range of each processing parameter explored in this study, specifically the target nitric acid and Pu(IV) nitrate concentrations during precipitation, and the calcination temperature and time. The SSA range during the PuO$_2$ production campaign was $4-11$ m$^2$/g. It is important to note that other processing variables were also analyzed (e.g., agitation time, cake mass, and air flow), but these four parameters were found to be the most relevant when correlating the SSA to the processing conditions (see below). Also, several parameters remained constant during the process and while they could have an effect (e.g. oxalic acid concentration, agitation rate), there was not enough variability in the data to draw conclusions at this time.

Several studies have analyzed the SSA of the final PuO$_2$ product with respect to the calcination temperature because of its importance in understanding radiolytic decomposition of water on the PuO$_2$ surface.\textsuperscript{1−4,7,8,16} Since the 1960s, the quasi-exponential decrease in the SSA as a function of increasing calcination temperature is thought to be the result of crystalline rearrangement occurring at high temperatures, leading to the disappearance of pores and irregularities on the surface.\textsuperscript{1−4} A plot of the SRS SSA versus calcination temperature data (Figure 1) illustrates that an exponential relationship does not describe the trend over a narrow temperature range. Another interesting observation is that higher calcining temperatures in general led to higher SSA in the data set, which is counterintuitive to higher temperatures leading to decreasing surface areas. During production runs, two furnaces were used, one calcining at $\sim$665 °C and the other at $\sim$690 °C. In many cases, the time required to heat the oxalate product in the lower temperature furnace to the set temperature exceeded the facility threshold requirements, thereby necessitating an additional $\sim$4.5 h of calcination time.

Preliminary investigation of the SRS data showed an interesting relationship between the nitric acid molarity and SSA not previously identified in the literature. This subset represents data where variation occurred only in the nitric acid concentration; the Pu concentration, calculation temperature, calculation time, and cake size remained near constant. Figure 2 shows that the effect of the nitric acid molarity on the SSA can be characterized by a negative linear correlation. This newfound result acted as the basis for analyzing several different process parameters for potential correlations to the PuO$_2$ physical characteristics, which were ultimately narrowed down to the nitric acid molarity, Pu(IV) nitrate concentration, and calcination temperature and time.

Of the 39 batch runs, our analysis only considered 27 as the others were identified as outliers falling outside normal processing conditions; SSA range for those data is $6-11$ m$^2$/g. The resultant SRS data set was further divided into training and testing sets using an iterative leave-one-out (LOO) approach\textsuperscript{27} to provide an unbiased identification of a model’s performance. Figure 3 describes a flowchart of the iterative LOO approach utilized here.

In this paper, 26 data points acted as the training data and 1 data point for the testing data set, wherein the model was iterated resulting in total 27 different models. After each training and testing stage, the testing data point was changed and the model was reanalyzed utilizing the new data sets. This was repeated until each data point was utilized as a testing data point. This methodology provides an avenue for identifying the true capabilities of the model without bias as some data points would result in a high error being very different from the

### Table 1. Range of Processing Parameters at SRS during the PuO$_2$ Production Campaign Used in This Study

| parameter                        | minimum value | maximum value |
|----------------------------------|---------------|---------------|
| nitric acid concentration (M)    | 1.7           | 2.7           |
| Pu(IV) nitrate concentration (g/L)| 25            | 45            |
| calcination temperature (°C)    | 655           | 695           |
| calcination time (min)          | 275           | 655           |
training data set while other data points would result in a low error being very similar to the training data set. The training data set was used to train a linear multi-variable model to predict the SSA after calcination. Equation 1 shows the generalized form of the where the variables, $A_{ij}$, correspond to the weighting values which will be calculated using the training data set.

$$A_1 \times T + A_2 \times t + A_3 \times M + A_4 \times C = SSA$$

Equation 1 was fit for three independent model forms each utilizing different processing input parameters. The first form analyzes the calcination temperature ($T$) and time ($t$) as the variable of interest (i.e., $A_1 = A_4 = 0$). The second form examines relationship between the nitric acid ($M$) and Pu(IV) nitrate ($C$) concentrations only as a model variable (i.e., $A_1 = A_2 = 0$). The final model form utilizes all four process parameters of interest to illustrate if remnant precipitation conditions have bearing on the material properties after calcination. Comparison of the four process parameters and the yield confirms that the process parameters are not intercorrelated, and thus, multicollinearity is avoided (Figure S1 in Supporting Information). This final model form attempts to provide information on processing history to more accurately model product characteristics. Finally, statistical analyses were performed on each of the model forms to illustrate the predictability of each form.

**Figure 1.** Comparison of SSA vs calcination temperature for all SRS data.

**Figure 2.** Effect of nitric acid molarity on SSA based on a subset of the SRS PuO$_2$ data. The trend line corresponds to the equation $y = -1.8796x + 11.58$ and had a $R^2$ value of 0.7503.
RESULTS AND DISCUSSION

Initial evaluation of the PuO2 SSA with the processing data considered only calcination temperature, and those results are described in the Supporting Information. Using a linear relationship, the average SSA among all the runs is 8.292 m²/g with an average error of 14%. Because the temperature-only prediction to the experiment is fairly vertical, and the data range between 6 and 11 m²/g over a small temperature window, our next consideration added calcination time to the model since roughly half of the process runs included an extended calcination period. Equation 2 describes the simplified forms of equation 1 for the prediction of SSA using this new model form with the coefficients of $A_1$ and $A_2$ being 0.0142 and $-0.00235$, respectively.

$$0.0142^T - 0.00235^t = SSA$$ (2)

For each simulation, the error in predicting the training data set and the test data point was calculated and plotted (Figure 6). The data points lying above (below) the line indicate that the linear regression predictive model has overpredicted (underpredicted) the SSA. The best line fit for the data points is generated, and the $R^2$ is calculated which indicates the goodness of fit of the predictive model on the training data. When incorporating time to the model, the average error lowers to 12.9% and the $R^2$ coefficient is 0.34 (Table 2). This is a marked improvement from using temperature data alone, and the results better describe the SSA variation occurring over a temperature difference of 50 °C. However, the model is not accurate over all data points even though the average error is low, as shown in Figure 6. In other words, the model is accurate for some but not all the data. This discrepancy can be quantified using the $R^2$ value, indicating the amount of data variability in the model, with roughly a 3× increase from the temperature-only model which had a value of 0.12.

Our next model follows the same LOO methodology but evaluates how the nitric acid and plutonium concentrations affected the SSA without inclusion of calcination conditions. Equation 3 shows the form with the average coefficients across the simulation runs which are 0.03231 and 0.2332 for $A_3$ and

Figure 3. Flowchart of the iterative LOO methodology. The loop is repeated until $i$ is equal to the number of data points in the data array.

Figure 4. Comparison of SSA and calcination temperature across the SRS data set.

Figure 5. Plot of residuals for each data point using equation 4 to fit the data.
respectively. Of the two coefficients, there is larger variability with $A_3$ suggesting the data are not representative of the nitric acid concentration effect. However, the model performed better than evaluating calcination temperature alone and had a lower average prediction error and $R^2$ value of 10% and 0.32, respectively (Table 2).

Equation 3:

$$0.03231^*M + 0.2332^*C = SSA$$

These results were unexpected since precipitation parameters alone have not been attributed to affecting the PuO$_2$ SSA. From Figure 6 and Table 2, equation 3 results in lower values for the average error and statistically the same $R^2$ coefficient when compared to equation 2. Comparison of the

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leading to decreases in the SSA, but instead is reflective of the data set used to generate the model. Figure 4 compares the calcination temperature with the SSA for all data points used from the SRS data set and shows a positive trend with increasing temperature corresponding to an increase in SSA.

This trend is believed to be artificial as the data points were obtained from samples calcined in two different furnaces. The first furnace at low temperature was calcined for almost twice the time that the high temperature furnace was held at temperature. Therefore, this positive trend in the temperature can actually be associated with the hold time parameter. As such it is expected that with a larger data set, the accepted decreasing trend with increasing temperature will be identified by the model. Furthermore, in both equations 3 and 4, the Pu(IV) nitrate concentration coefficient is positive, while the nitric acid concentration coefficient fluctuates between positive to negative values, predicting the SSA tends to increase with increasing Pu(IV) nitrate concentration. However, the same conclusion cannot be gathered from the nitric acid concentration. This could be due to the lack of data available as the nitric acid concentration values were obtained for only a relatively small range. The scatter of the residuals from equation 4 versus the predicted SSA (Figure 5) illustrates that a simple model achieves moderately high accuracy in terms of R. The scatter of the residuals is random, indicating no ties in the data. Moreover, the random underprediction and over-prediction, that is, data points lying randomly under and over the identity line, indicate that the model is not biased in a certain direction. Comparison of the $R^2$ for the three predictive models reveals that including precipitation parameters and calcination time to the general correlations of temperature has the best estimation power. Thus, inclusion of the precipitation parameters may be preferred for the predictive modeling of the PuO$_2$ SSA. This is a significant revelation for industrial-scale PuO$_2$ process optimization to reduce the SSA within tight controls defined by the facility requirements.

### CONCLUSIONS

Multiple linear regression models based on precipitation and calcination parameters were developed to predict the SSA from the PuO$_2$ production data, specifically the target nitric acid and Pu(IV) nitrate concentrations during precipitation, and the calcination temperature and time. Our analysis showed that individual unit operations (e.g., only calcination or precipitation parameters) have an average error similar to the conventional temperature-only model but a nearly three-times increase in the description of the variability of the data. However, the final model which utilized both precipitation parameters and calcination parameters to predict PuO$_2$ SSA was not only more accurate than the conventional model with a decrease in average error from 13.90 to 8.41% but also showed an increase in the $R^2$ coefficient from 0.1202 to 0.7376, thereby showing that both precipitation and calcination provides the best predictive capability out of the models shown here. Therefore, inclusion of precipitation parameters to a calcination time–temperature model could allow for more accurate prediction of the PuO$_2$ SSA, which has not been identified by previous studies. Furthermore, although data are limited, an exhaustive analysis of all possible process parameter combinations may be needed to identify further parameters of interest, for example, oxalate concentration, impurity concentration and carbon content. This justifies the need for advanced machine learning algorithms to provide model

### Table 2. Comparison of Model Statistics Across All Reported SSA Models

| parameters of interest | percent error  | $R^2$ coefficient |
|------------------------|----------------|------------------|
|                        | average (%)    | standard deviation | average | standard deviation |
| calcination parameters only ($T$ and $t$) | 12.90 | 9.3002 | 0.3437 | 0.0356 |
| precipitation parameters only ($M$ and $C$) | 10.65 | 7.8573 | 0.3236 | 0.0491 |
| precipitation and calcination parameters ($T$, $t$, $M$, and $C$) | 8.41 | 5.0744 | 0.7376 | 0.0151 |

Experimental and predicted SSA (Figure 6) shows two distinct clustering of values at ~7 m$^2$/g and again ~9.5 m$^2$/g. During the middle of the campaign, an effort was made to increase the Pu concentration from ~30 to ~45 g/L, and in general, higher SSAs were seen with higher Pu concentration. Although there was some variation in the nitric acid concentration early on (Figure 2), the concentration remained steady at 2.6 M, well within industrial-scale processing parameters. While the results are interesting, the model is only capturing about 32% of the data variability and therefore very limited in its predictive capabilities.

The last model defined in equation 4 describes the specific form of this four-parameter model for the prediction of SSA. An important note is that the value of the $A_3$ coefficient, which corresponds to the nitric acid molarity parameter, has a similar value to the linear regression in Figure 2, corroborating the preliminary hypothesis of a negative correlation between the nitric acid molarity and SSA. Also, the coefficient is stabilized with respect to the average value and sign. The average error and $R^2$ value in the prediction of SSA were found to be 8.41% and 0.74, respectively. Interestingly, results from simulation 15 show that most of the data now reside to ~10% error compared to the experiment (Figure 6). Also, while the model improves the predictability overall, simulation 21 is a significant outlier. When the process parameters of the test data point associated with simulation 21 were compared to the training data set, the nitric acid molarity was outside the training model range and hence the predicted value must be extrapolated rather than interpolated. The error associated with simulation 21 shows that the regression model developed here is highly accurate for simulations involving interpolation of data while it is less accurate for simulations involving extrapolation. Therefore, our analysis is highly accurate for the given processing parameter range and can be extended to encompass the full range of processing parameters with more experimental data.

$$0.01162*T - 0.0013*t - 1.7299*M + 0.15604*C = SSA$$

(4)

While a direct comparison of the parameter coefficients across all three models would be desired, it is often not possible due to the nature of regression modeling, wherein the coefficient values are defined by the data that is chosen as the training data set. However, an indirect comparison of the coefficient signs can provide insights into the trends of the data. For example, in equations 2 and 4, the calcination temperature coefficient is positive while the calcination time coefficient is negative. Therefore, the trend shows that SSA tends to increase with an increase in temperature but decrease with an increase in time. This is a counter description to the known correlations of increasing calcination temperature
comparisons to quantify the model predictive capabilities as a function of process parameters.

**ASSOCIATED CONTENT**

* Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsomega.1c04964.

Discussion on the regression analyses of PuO2 SSA versus calcination temperature and coefficient values for each question across all 27 simulations performed (PDF)

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Notes
The authors declare no competing financial interest.

**ACKNOWLEDGMENTS**

We would like to thank Steve Hensel and Chip McClard for helpful discussions. This work was funded by the U.S. Department of Homeland Security (DHS), Countering Weapons of Mass Destruction (CWMD), under a competitively awarded Interagency agreement no. HSHQDN-16-X-00054. This work was produced by Battelle Savannah River Laboratory-Site: Savannah River National Laboratory: Aiken, SC, 2012.

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