Sandia National Laboratories Waste Isolation Pilot Plant

Uncertainty Analysis of Actinide Solubilities for the WIPP CRA-2014 PA, Rev. 1 Supersedes ERMS 559278

Work Carried Out under Task 4 of the Analysis Plan for WIPP Near-Field Geochemical Process Modeling, AP 153, Rev. 1. To be included in the AP-153 records package

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WIPP: 1.4.2.2: PA: QA-L: 557522 **Information**

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TABLE OF CONTENTS

1 INTRODUCTION	7
2 METHODS	
3 RESULTS	
3.1 Th(IV) Solubility Model3.2 Am(III) Solubility Model	
4 CONCLUSIONS	
5 REFERENCES	

LIST OF FIGURES

Figure 1.	Histogram of the difference $D = \log_{10} S_m - \log_{10} S_p$ for Th(IV) for the CRA-2014 PA (this revised analysis).	38
Figure 2.	Plot of the CDF of the differences between the measured and predicted Th(IV) solubilities for the CRA-2014 PA (this revised analysis).	41
Figure 3.	Histogram of the difference $D = \log_{10} S_m - \log_{10} S_p$ for Nd(III) and Am(III) for the CRA-2014 PA (this revised analysis).	58
Figure 4.	Plot of the CDF of the differences between the measured and predicted Nd(III) and Am(III) solubilities for the CRA-2014 PA (this revised analysis)	64

LIST OF TABLES

Table 1.	Abbreviations, Acronyms, Initialisms, etc.	8
Table 2.	Run-Control Information for the Calculations Carried Out for the Original Analysis.	17
Table 3.	Final Evaluations of Lab Studies for Comparisons of Measured and Predicted Th(IV) Solubilities.	21
Table 4.	Results from Rai et al. (1997) Excluded from Th(IV) Comparisons Because of Complete Dissolution of Solids.	36
Table 5.	Statistical Properties of the Distribution of $D = log10(Sm) - log10(Sp)$ for Th(IV) for the CRA-2014 PA (This Revised Analysis).	38
Table 6.	CDF of the Differences between Measured and Predicted Th(IV) Solubilities for the CRA-2014 PA (This Revised Analysis). This CDF Defines the Solubility Multiplier SOLVAR for the Oxidation-State IV Model Material SOLMOD4 in the WIPP PA Parameter Database.	39
Table 7.	Final Evaluations of Laboratory Studies for Comparisons of Measured and Predicted Nd(III) and Am(III) Solubilities.	42
Table 8.	Statistical Properties of the Distribution of $D = log10(Sm) - log10(Sp)$ for Nd(III) and Am(III) for the CRA-2014 PA (This Revised Analysis).	58
Table 9.	CDF of the Differences between the Measured and Predicted Nd(III) and Am(III) Solubilities for the CRA-2014 PA (This Revised Analysis). This CDF Defines the Solubility Multiplier SOLVAR for the Oxidation-State III Model Material SOLMOD3 in the WIPP PA Parameter Database.	59

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1 INTRODUCTION

This analysis report revises the results of Brush and Domski (2013b). For this revised analysis, we established an additional general criterion (G11) (see Section 2) that we applied to the solubility experiments that had already satisfied criteria G1 through G10 and the specific criterion S1. We established criterion G11 so that the approach used to quantify the uncertainties in the predictions of Th(IV) and Am(III) solubilities in Waste Isolation Pilot Plant (WIPP) disposal rooms is consistent with that used to develop the uncertainties in the other parameters sampled by WIPP performance assessment (PA).

Brush and Domski (2013b), the original version of this analysis report, provided the results of their updated uncertainty analysis of the solubilities of Th(IV) and Am(III). Actinide solubilities (the sums of the concentrations of all dissolved actinide species in chemical equilibrium with actinide-bearing solid phases under the conditions expected in WIPP disposal rooms) and the concentrations of colloidal (suspended) actinides (calculated from the baseline solubilities) together constitute the actinide source term used in WIPP PA calculations. These solubilities are often referred to as the "baseline solubilities" because they comprise unique values predicted using thermodynamic models for each actinide element and brine under expected near-field chemical conditions. However, PA codes actually use actinide solubilities that are the products of the baseline solubilities and sampled uncertainty factors that describe possible deviations of the predicted Th(IV) and Am(III) solubilities from experimentally measured solubilities. The reason for using these uncertainty factors is to adjust the baseline solubilities up if the model underpredicts the experimentally measured solubilities, or to adjust the baseline solubilities down if the model overpredicts the measured solubilities. The baseline solubilities have been established in a separate analysis (Brush and Domski, 2013a).

Brush and Domski (2013b) did not include U(VI) in their analysis because a thermodynamic speciation-and-solubility model has not been developed for U(VI). Instead, the EPA specified that an estimate of 1×10^{-3} M be used for the solubility of U(VI) in Generic Weep Brine (GWB) and Energy Research and Development Administration (WIPP Well) 6 (ERDA-6). GWB is a synthetic brine representative of intergranular Salado Formation (Fm.) brines at or near the stratigraphic horizon of the repository (Krumhansl et al., 1991; Snider, 2003). ERDA-6 (Popielak et al., 1983) is a synthetic brine representative of fluids in brine reservoirs in the Castile Fm., which underlies the Salado Fm. An estimate of 1×10^{-3} M has been used for U(VI) since 2005. Furthermore, the EPA specified a fixed value for its estimate of the solubility of U(VI) in GWB and ERDA-6 (U.S. EPA, 2006).

Furthermore, Brush and Domski (2013b) did not include Np(V) because PA does not sample the uncertainties in the solubilities of Np in view of the fact that releases of this element do not affect the long-term performance of the WIPP (Brush and Garner, 2005).

SNL PA personnel will use the results of this revised uncertainty analysis in the PA for the third WIPP recertification application to the U.S. Environmental Protection Agency (EPA)

(the 2014 Compliance Recertification Application, or CRA-2014 PA). Brush et al. (2005) described the PA implementation of these results in detail.

Brush and Domski (2013b) used EQ3/6, Version 8.0a, (Wolery and Jarek, 2003; Wolery, 2008; Wolery et al., 2010; Xiong, 2011b) and the thermodynamic database (DB) DATA0.FMT.R2, also known as DATA0.FM1, (Xiong, 2011a) for their analysis. Wolery (2008), Wolery et al. (2010) and Xiong (2011b) completed the qualification of Version 8.0a of EQ3/6 according to Sandia National Laboratories' (SNL's) WIPP quality assurance (QA) procedures for WIPP compliance-related actinide-solubility calculations. We did not carry out any additional EQ3/6 runs for this revised analysis. Instead, we used the EQ3/6 input and output (I/O) files and associated spreadsheets from Brush and Domski (2013b) (see Section 2 below) to implement criterion G11 in this revision.

Brush and Domski (2013b) conducted their analysis under Task 4 of AP-153, Rev. 1, the current analysis plan (AP) for WIPP near-field geochemical process modeling (Brush et al., 2012, Subsection 4.4). We also performed this revised analysis under Task 4 of AP-153, Rev. 1.

Table 1 defines the abbreviations, acronyms, initialisms, etc., used in this report.

Abbreviation, Acronym, or Initialism	Definition
Am, Am(III)	americium, americium in the +III oxidation state
am	amorphous
AP	analysis plan
aq	aqueous (dissolved in an aqueous solution)
atm	atmosphere(s)
C, c	carbon, crystalline
Ca, Ca^{2+}	calcium, calcium ion
CAP	(SNL/WIPP QA) corrective action plan
CAR	(SNL/WIPP QA) corrective action request
CDF	cumulative distribution function
citrate	$(CH_2COO)_2C(OH)(COO)^{3-}$ or $(CH_2CO_2)_2C(OH)(CO_2)^{3-}$
Cl, Cl⁻	chloride, chloride ion
ClO ₄ , ClO ₄ ⁻	perchlorate, perchlorate ion
Cm, Cm(III)	curium, curium in the +III oxidation state
CMS	(Sandia/WIPP software) Configuration Management System
$\text{CO}_3, \text{CO}_3^{2-}$	carbonate, carbonate ion
col	colloidal (suspended in an aqueous or other solution)

Table 1. Abbreviations, Acronyms, Initialisms, etc.

Table 1 continued on next page

Abbreviation, Acronym, or Initialism	Definition
CPR	cellulosic, plastic and rubber (materials)
cr	crystalline
CRA-2004	the first WIPP Compliance Recertification Application, submitted to the EPA in March 2004
CRA-2009	the second WIPP Compliance Recertification Application, submitted to the EPA in March 2009
CRA-2014	the third WIPP Compliance Recertification Application, to be submitted to the EPA in March 2014
D	the difference(s) between logs (base 10) of the measured and predicted actinide solubilities (Sm and Sp)
EPA	(U.S.) Environmental Protection Agency
EQ3/6	a geochemical software package for speciation and solubility calculations (EQ3NR) and reaction-path calculations (EQ6)
ERDA-6	Energy Research and Development Administration (WIPP Well) 6, a synthetic brine representative of fluids in Castile brine reservoirs
f_{CO_2}	fugacity (similar to the partial pressure) of CO_2
Fm.	Formation
FMT G-Seep	Fracture-Matrix Transport, a geochemical speciation and solubility code a naturally-occurring brine formerly collected from G Drift in the WIPP underground workings and used for laboratory and modeling studies
GWB	Generic Weep Brine, a synthetic brine representative of intergranular Salado brines
H, H^+	hydrogen, hydrogen ion
HCO ₃ , HCO ₃ ⁻	bicarbonate, bicarbonate ion
H_2O	water
Ι	ionic strength, defined by $I = \frac{1}{2} \times \sum_{i} (M_i \times z_i^2)$, in which M_i and z_i are the molarity and charge of species <i>i</i>
I/O	(EQ3/6) input and output files
K, K ⁺	potassium, potassium ion
LÍB	Library
log	the common logarithm or logarithm (base 10)
М	molar
m	molal
mcr	microcrystalline
Mg, Mg ²⁺ mM	magnesium, magnesium ion millimolar

Table 1. Abbreviations, Acronyms, Initialisms, etc. (Cont.).

Table 1 continued on next page

Abbreviation, Acronym, or					
Initialism	Definition				
N	bin number (see Subsection 5.2 for explanation)				
Na, Na $^+$	sodium, sodium ion				
NaTr	sodium trifluoromethanesulfonate (also abbreviated as sodium triflate)				
Nd, Nd(III)	neodymium, neodymium in the +III oxidation state				
NO_3, NO_3	nitrate, nitrate ion				
Np, Np(IV), Np(V)	neptunium, neptunium in the $+IV$ oxidation state , neptunium in the $+V$ oxidation state				
0	oxygen				
OH, OH ⁻	hydroxide, hydroxide ion				
P	probability				
PA	performance assessment				
PABC	(WIPP) Performance Assessment Baseline Calculations, carried out in 2005 and 2009				
рсН	the negative log (base 10) of the molar concentration of H^+				
p _{CO2}	partial pressure (similar to the fugacity) of CO_2				
pmH	the negative log (base 10) of the molal concentration of H^+				
ppm	parts per million				
pH	the negative log (base 10) of the activity of H^+				
Pu, Pu(III), Pu(IV)	plutonium, plutonium in the +III oxidation state, plutonium in the +VI oxidation state				
QA	quality assurance				
R	gas constant				
Rev.	Revision				
S _m	measured solubility				
S _p SNL	predicted solubility Sandia National Laboratories				
SO_4 , SO_4^{2-}	sulfate, sulfate ion				
504, 504 Т	temperature				
Th, Th(IV)	thorium, thorium in the +IV oxidation state				
TIC	total inorganic C (the sum of the dissolved species of inorganic C)				
U, U(IV), U(VI)	uranium, uranium in the +IV oxidation state, uranium in the +VI oxidation state				
μm	micrometer				
WIPP	Waste Isolation Pilot Plant				
XRD	X-ray diffraction				
Zr, Zr(IV)	zirconium, zirconium in the +IV oxidation state				
μ ⁰ , μ ⁰ /RT	standard chemical potential, dimensionless standard chemical potential				

Table 1. Abbreviations, Acronyms, Initialisms, etc. (Cont.).

2 METHODS

This analysis report revises the results of Brush and Domski (2013b). For this revised analysis, we established an additional general criterion (G11) (see below) that we applied to the Th(IV), Nd(III), Am(III), and Cm(III) solubility experiments that had already satisfied criteria G1 through G10 and the specific criterion S1. We established criterion G11 so that the approach used to quantify the uncertainties (errors) in the predictions of Th(IV) and Am(III) solubilities in WIPP disposal rooms is consistent with that used to develop the uncertainties in the other parameters sampled by WIPP PA.

The methods used by Brush and Domski (2013b) for their original analysis were similar to those used by Xiong et al. (2005, 2010, 2011). They compared experimentally measured solubilities of Th(IV), Nd(III), and Am(III) with the Th(IV) and Am(III) solubilities predicted using EQ3/6, Version 8.0a, (Wolery and Jarek, 2003; Wolery, 2008; Wolery et al., 2010; Xiong, 2011b) and the thermodynamic DB DATA0.FM1 (Xiong, 2011a). Brush and Domski (2013b) used EQ3/6, Version 8.0a, and DATA0.FM1 to recalculate all of the previous comparisons by Xiong et al. (2005, 2010, and 2011) using FMT (Babb and Novak, 1997 and addenda; Wang, 1998). They then constructed histograms and cumulative distribution functions (CDFs) of the differences between the logarithms (base 10) of the measured and predicted solubilities. They constructed separate histograms and CDFs for the Th(IV) and Am(III) models. However, these CDFs will not be used for the CRA-2014 PA; instead, the CDFs from this revised analysis will be used.

Brush and Domski (2013b) used ten general criteria for including results of experimentally measured solubilities in their comparisons of measured Th(IV), Nd(III), Am(III), and Cm(III) solubilities and predicted Th(IV) and Am(III) solubilities. The general criteria were:

G1. Include only results from experimental studies published from January 1, 1990, through October 31, 2011. We deviated from this criterion by including the results of Borkowski et al. (2012) and Borkowski (2012). We included these results because we were directed to do so by our management.

G2. Include results from both papers published in peer-reviewed journals and unpublished reports (e.g., officially released reports from government laboratories such as Los Alamos National Laboratory, Lawrence Livermore National Laboratory, Pacific Northwest National Laboratory, etc.).

G3. Include only results from solubility studies. Exclude other studies that do not provide solubilities (e.g., studies of corrosion, leaching, sorption, or transport).

G4. Include only results from studies in which water was the solvent. Exclude studies in which other solvents were used (e.g., solids, molten metal or salts, or organic liquids).

G5. Include only results obtained from studies at pressures at or close to atmospheric, at temperatures at or close to those expected in the WIPP (i.e., 20-30 °C), and with posttest phase-separation methods similar to those used for the WIPP, because these were the conditions and methods used to parameterize the Th(IV) and Am(III) speciation and solubility models for WIPP compliance-related PA calculations.

G6. Include only results from studies of Th(IV); and Nd(III), Am(III), and Cm(III); because these are the elements used to parameterize the WIPP Th(IV) and Am(III) solubility models. Exclude studies of U(IV), Np(IV), and Pu(IV); and Pu(III) because there could be systematic differences between the solubilities of these elements and those of their oxidation-state analogs used to parameterize the models; and because the difficulties inherent in maintaining these elements in these oxidation states could introduce experimental artifacts in the results obtained with these elements.

G7. Include only results from studies with a characterized solubility-controlling solid for which the value of the dimensionless standard chemical potential (μ^0/RT) is in the WIPP Th(IV) or Am(III) model (i.e., in the EPA-certified thermodynamic database), and in which the quantity of solid initially present was sufficient to prevent complete dissolution of this solid during the experiments.

G8. Include only results from studies with aqueous solutions of known composition. Exclude studies performed with groundwaters, sedimentary pore waters, and soil solutions that may contain unknown quantities of species that can be complexants or adsorbents (e.g., humic acids or other dissolved organic compounds, microbial colloids, or pseudocolloids).

G9. Include only results from studies with dissolved elements or species for which values of μ^0/RT and Pitzer ion-interaction parameters are in our models. Exclude studies with dissolved elements or species for which our models do not include values of μ^0/RT or Pitzer parameters.

G10. Include only results from studies for which the investigators provided a complete description of their experiments and the original solubilities. Exclude literature reviews and summaries, and studies in which the authors only provided average dissolved concentrations or solubility products, thus necessitating backcalculation of the solubilities.

Brush and Domski (2013b) also continued to use the specific criterion established by Xiong et al. (2005) for the experimentally measured solubilities of Th(IV):

S1. Include only results from experiments carried out with solutions with ionic strength (I) \geq 3 M or m.

Xiong et al. (2005, Subsection 5.2.3) described the reason for establishing S1.

Xiong et al. (2005, 2010, 2011) did not establish any specific criteria for experimentally measured solubilities of Nd(III), Am(III), or Cm(III). Furthermore, Brush and Domski (2013b) did not establish any additional specific criteria for their comparison.

For this revised analysis, however, we established an additional general criterion (G11) and applied it to the Th(IV), Nd(III), Am(III), and Cm(III) solubility experiments that had already satisfied criteria G1 through G10 and S1. This additional criterion is:

G11. Include only results from experimental studies carried out under conditions at or close to those predicted for WIPP disposal rooms. Specifically, include only results from experiments in which: (1) I \geq 3 m or M, (2) pcH = 8.0–11.2, and (3) total inorganic C (TIC) = 0-1.4 × 10⁻² M. (In WIPP brines after equilibration with the important solids in the repository, the TIC concentration is essentially the sum of the concentrations of the dissolved species HCO₃⁻ and CO₃²⁻ and is closely related thermodynamically to f_{CO₂} or p_{CO₂}.)

We established criterion G11 so that the approach used to quantify the uncertainties (errors) in the predictions of Th(IV) and Am(III) solubilities in WIPP disposal rooms is consistent with that used to develop the uncertainties in the other parameters sampled by WIPP PA. The objective of WIPP PA is to predict the performance of the repository during the 10,000-yr regulatory period. Therefore, the future conditions expected in the repository must inform and guide the establishment of the parameter uncertainty distributions used in WIPP PA. Criteria G1 through G10 and S1, used by Xiong et al. (2005, 2011) to quantify the previous errors in the predictions of actinide solubilities, allowed for the inclusion of experimental results obtained under conditions quite different from the future conditions expected in WIPP disposal rooms (see below). Although potentially useful for validating the Th(IV) and Am(III) speciation and solubility models for other possible applications, the previous uncertainties established by Xiong et al. (2005, 2011) were inconsistent with the conditions expected in the repository.

When Xiong et al. (2005) specified criterion S1, they noted that "Exclusion of 114 solubilities measured in intermediate-ionic-strength solutions from the new An(IV) comparison is reasonable, given the intended use of FMT for predicting [Th(IV)] solubilities in high-ionic-strength brines." This argument also applies to the uncertainties in the predictions of Am(III) solubilities, given that the objective is to establish a distribution of prediction errors for the conditions expected in WIPP disposal rooms. Furthermore, Xiong et al. (2005) noted that the prediction error is dependent on pcH, even after the exclusion of the Felmy et al. (1991) data for pcH \leq 3.6. Therefore, the pcH expected in the repository must be taken into account when developing the uncertainty distributions for the Th(IV) and Am(III) models. Finally, the TIC concentration is also expected to affect the prediction error of these models, so the TIC expected in the WIPP must also be specified. Consequently, the approach used to develop actinide-solubility uncertainties was inconsistent with that used for the other uncertain parameters in WIPP PA. Thus, the addition of criterion G11 is necessary to quantify the uncertainties in the predictions of Th(IV) and Am(III) solubilities in a manner consistent with the other uncertainties in the predictions of Th(IV) and Am(III) solubilities in a manner consistent with the other uncertainties in the predictions of Th(IV) and Am(III) solubilities in a manner consistent with the other uncertainties in the predictions of Th(IV) and Am(III) solubilities in a manner consistent with the other uncertainties in the prediction parameters in WIPP PA.

Therefore, criterion G11 specifies narrow ranges of I, pcH, and TIC to simultaneously include: (1) measured Th(IV), Nd(III), Am(III), and Cm(III) solubilities from experiments performed under conditions close to those predicted for WIPP disposal rooms, and (2) enough results to establish ranges and probability distributions for the uncertainties in our predictions of solubility predictions for those conditions. We stress that our specifications of I \geq 3 m or M, pcH = 8.0–11.2, and TIC = 0-1.4 × 10⁻² M do not imply that we expect I, pcH, or TIC to deviate from the values predicted by Brush and Domski (2013a, Tables 5 and 6) for the minimum volumes of WIPP brines required for a release from the repository (I = 6.44 M, pcH = 9.54 or pH = 8.82, and TIC = 3.79×10^{-4} M in GWB; I = 5.99 M, pcH = 9.69 or pH = 8.99, and TIC = 4.55×10^{-4} M in ERDA-6). Indeed, we refer to these combinations of conditions for GWB and ERDA-6 as "invariant points," because they are buffered (maintained) by reversible reactions (see Brush et al., 2012, Subsection 2.1.2). Rather, criterion G11 specifies narrow ranges of these parameters to avoid screening out all or nearly all of the results that had already met criteria G1 through G10 and S1.

We established the specification that I be ≥ 3 m or M so as to apply the same criterion to the solubility experiments carried out with Nd(III), Am(III) or Cm(III) as that previously applied to experiments with Th(IV). We did not specify an upper limit of I because all of the results included after applying criterion G11 had values of I within a factor of two greater than those predicted for the WIPP brines GWB and ERDA-6 (see above). We retained criterion S1 for this revision of Brush and Domski (2013c) to minimize the changes required in the text of Subsections 3.1 and 3.2, and in Tables 3 and 7, even though this specific criterion is now included in the general criterion G11.

We established a pcH range of 8.0–11.2 by rounding the pcH of 9.54 predicted for GWB (see above) to 9.5 and setting the lower limit of the range 1.5 units of pcH below this value. Similarly, we rounded the pcH of 9.69 predicted for EERDA-6 to 9.7 and set the upper limit 1.5 units above it. We reiterate that we do not expect the pcH or pH to deviate from the values predicted for GWB and ERDA-6 at the invariant points.

Criterion G11 specifies that the lower limit of the TIC range be 0 to ensure that this revised analysis includes CO₂-free Th(IV) solubility measurements, and CO₂-free Nd(III), Am(III), and Cm(III) solubilities. Inclusion of CO₂-free conditions is appropriate because the speciation and solubilities predicted by the Th(IV) and Am(III) models for the invariant points for GWB and ERDA-6 described above are more like those in CO₂-free systems than high- f_{CO_2} or high-HCO₃ and/or high-CO₃²⁻ systems. This is because: (1) the dominant dissolved species predicted for Th(IV) and Am(III) under these conditions are the Th(OH)4(aq) and various Am-organic or $Am(OH)_x^{3-x}$ complexes, respectively; and (2) the solids predicted to control the solubilities of Th(IV) (hydrous, amorphous ThO₂) and Am(III) (Am(OH)₃) are both CO₂-free phases. The reason that Th(IV) and Am(III) will behave in this manner is that the MgO emplaced in the WIPP to satisfy the EPA requirement for multiple engineered and natural barriers will consume essentially all of the CO₂ that would be produced should microbial activity consume all of the cellulosic, plastic, and rubber (CPR) materials in the TRU waste, waste containers, and waste-emplacement materials in the repository. Although MgO will consume essentially all of the CO₂, minute quantities (relative to the quantity that would be produced by microbial consumption of all of the CPR materials) would persist in the aqueous



and gaseous phases. This residual CO₂ and the MgO carbonation reaction would maintain f_{CO_2} at 3.14×10^{-6} atm and the TIC concentrations at the values provided above for the invariant points predicted for GWB and ERDA-6 (Brush et al., 2012, Subsection 2.1.2).

Finally, we specified the upper limit of the TIC range by multiplying the TIC content of ERDA-6, the higher of the two brines, by $10^{1.5}$ or 31.6. This is consistent with the lower and upper limits of the pcH range, which are 1.5 pcH units below and above the values in GWB and ERDA-6, respectively, rounded to two significant figures (see above).

The use of criterion G11 to restrict the experimentally measured solubilities included in this revised analysis to conditions at or close to those predicted for WIPP disposal rooms is a deviation from the methods specified for this uncertainty analysis in AP-153, Rev. 1 (Brush et al., 2012, Subsection 4.4). This is because AP-153, Rev. 1, stated that only criteria G1–G10 and S1 would be used.

Domski carried out a literature search for published papers and unpublished reports on laboratory studies of Th(IV), Nd(III), Am(III), and Cm(III) solubilities for the original version of this analysis. He used Search Plus, Version 2.4, operated by Los Alamos National Laboratory (LANL), Web of Science (including the Science Citation Index and INSPEC), and the DOE Energy Citations Database.

This literature search identified hundreds of papers and reports on actinide chemistry. Therefore, Domski conducted preliminary evaluations of these studies by applying the criteria described above to the information provided in the abstracts of these papers and reports. Because of the large number of papers and reports identified, it was not practical to document which criteria were used to include these studies in or exclude them from the final evaluations described below. Domski excluded all papers and reports that, in his judgment, obviously failed to meet the criteria. In those cases in which it was not obvious that a paper or report should be included or excluded, Domski retained them for the final evaluations. Both Brush and Domski performed the final evaluations of the laboratory studies of actinide solubilities for this analysis.

We did not obtain or evaluate any additional papers or reports for this revised analysis.

Most of the newly acquired papers and reports did not include data tables. Therefore, Brush and Domski (2013b) attempted to obtain numerical values for all of the experimentally measured solubilities included in their analysis, as well as the parameters required to predict these solubilities with EQ3/6. This was possible in some, but not all cases. Consequently, Domski used the open source program Engauge Digitizer (http://digitizer.sourceforge.net/) to obtain numerical values of experimental variables such as pcH or pmH, and the resulting solubilities, from the figures in the papers and reports.

Domski then set up EQ3/6 input files to predict the solubilities of Th(IV), Nd(III), and Am(III) for the conditions reported for each of the measured solubilities in the laboratory studies selected by the final evaluations during the original analysis. (We did not carry out any additional EQ3/6 runs for this revised analysis.) Simulations for most of the experimental data were performed using the EQ3NR code. In these runs, the pH was specified;

the composition of the supporting solution was used as input, either in molar or molal units; and the radionuclide composition was calculated based on equilibrium with the solubility-controlling phase in the experiment. However, in some cases it was necessary to use both EQ3NR and EQ6 to simulate more complex experiments. In these cases, the experiments were conducted with a carbon dioxide (CO_2) atmosphere, and/or they were titrated to specific values of pH values at which the solubility measurements were made.

In many of the experiments, ionic concentrations are reported in units of moles per liter, or molarity (M); and the hydrogen ion (H^+) content of the solutions was measured in terms of the molar concentration of H^+ , also known as the pcH. When using these data in EQ3/6, which uses the mass-based unit moles per kg, or molality (m); and the negative log of the hydrogen ion activity (pH), additional steps were required.

In instances in which the ionic strength was high and data are reported in molarity, it was necessary to input the following EQ3 input parameters: total dissolved solids (tdspl), solution density (rho), and the aqueous-phase scale factor (scamas). These parameters are calculated based on the fluid composition in the MS-Excel spreadsheet "Conc_density_calcs_brines.xls." This spreadsheet will be submitted to the SNL/WIPP Records Center.

For those experiments that measured the pcH, a two-step approach was used to match the measured pcH. To arrive at the measured value of pcH, an initial EQ3/6 run was performed in which the input pH was referred to as pH_{init} . This resulted in an output value of pcH referred to as pcH_{init} . The difference between pcH_{init} and the measured value pcH_{exp} was then calculated and used to calculate a new pH_{init} , and a second EQ3/6 run was performed to obtain a simulated pcH equal to pcH_{exp} .

We implemented criterion G11 by postprocessing the results of Brush and Domski (2013b) with progressive sorting of the EQ3/6 output data using the Microsoft Excel sort function. Only data that met all three of the G11 subcriteria were retained for the final statistical analysis. We did not conduct out any additional EQ3/6 runs for this revised analysis; instead, we used the EQ3/6 I/O files and associated spreadsheets from Brush and Domski (2013b) (see below) to implement criterion G11 in this revision.

Table 2 (see next page) provides the location and run-control information for the thermodynamic database, Excel spreadsheets, and EQ3/6 I/O files for the calculations for the original analysis.

Table 2. Run-Control Information for the Calculations Carried Out for the Original Analysis.

Description or Title of File(s)	Location of File(s)		
EQ3/6 DB DATA0.FMT.R2 also called	In zip file DATA0_FMT.ZIP,		
DATA0.FM1	library LIBEQ36, class DATABASES		
Excel macro GetEQData_v101e.xls and	In zip file EQ36_IO_files.zip,		
GetEQData_v101f.xls	library LIBEQ36, class AP153		
EQ3/6 I/O files and Excel spreadsheets with input and extracted data	In zip file EQ36_IO_files.zip, library LIBEQ36, class AP153		

Finally, Domski generated frequency distributions of differences (D) between the logs of the measured and predicted actinide solubilities (S_m and S_p , respectively) and displayed them as histograms in Microsoft Excel spreadsheets using Excel's frequency function in this commercial spreadsheet software. One other deviation was implemented. AP153, Rev. 1 (Brush et al., 2012) specifies that histograms of the data will be produced by binning the data and implies that the data will be used in this form. However, the Latin Hypercube Sampling (LHS) code uses a cumulative distribution function (CDF) rather than probability density functions such as the histograms (see Subsection 3.1 and 3.2 below). The CDF must be constructed using the unbinned data so that those data can be submitted directly to PA for entry into their parameter database.

Negative values of D indicate that the Th(IV) or Am(III) model implemented in EQ3/6 predicted a solubility greater than the corresponding measured value (overprediction of solubility by the model); positive values of D indicate that the model predicted a solubility less than the measured value (underprediction). The Excel files "Thorium_Uncertainty_Results_2014_PA.xls" and "Americium_Uncertainty_Results_2014_PA.xls" and "Americium_Uncertainty_Results_2014_PA.xls" contain spreadsheets with S_m or S_p values (or logs thereof), frequency distributions, and histograms for this analysis. The bin numbers (N) in the histograms are defined as follows:

Bin N contains values of D greater than N - 0.15 and less than or equal to N, where $D = \log_{10}(S_m) - \log_{10}(S_p)$.

Examples of the definitions of bins follow:

Bin -0.3 contains values of D greater than -0.45 and less than or equal to -0.3; Bin -0.15 contains values of D greater than -0.30 and less than or equal to -0.15; Bin 0 contains values of D greater than -0.15 and less than or equal to 0; and Bin 0.15 contains values of D greater than 0 and less than or equal to 0.15.

We will submit electronic copies of the files "Thorium_Uncertainty_Results_2014_PA.xls" "Americium_Uncertainty_Results_2014_PA.xls," and all of the other spreadsheets to

the archived file EQ36_IO_files.ZIP, which is stored in the CMS in class ANALYSIS of library LIBPABC09_FMT.

3 RESULTS

This section provides the results of our revisions of the uncertainty analysis by Brush and Domski (2013b) of predictions of the solubilities of Th(IV) (Subsection 3.1) and Am(III) (Subsection 3.2).

Brush and Domski (2013b) included all of the measured solubilities included by Xiong et al. (2011) in their uncertainty analysis. However, application of criterion G11 (see Section 2 above) to the measured solubilities included by Brush and Domski (2013b) resulted in the exclusion of some of these solubilities. Brush and Domski also excluded all of the measured solubilities excluded by Xiong et al. (2011). Therefore, Brush and Domski (2013c) retained for ease of reference all of the text from Xiong et al. (2011) that described their final evaluations of the papers and reports that they considered. However, Brush and Domski (2013b) updated their text as necessary to include comparisons to their most recent predictions of the baseline solubilities of Th(IV) and Am(III), and the compositions of WIPP brines (Brush and Domski, 2013a).

Each of the following subsections provides separate lists of the published papers and unpublished reports identified and evaluated previously by Xiong et al. (2005) and/or Xiong et al. (2011), and the papers and reports evaluated by Brush and Domski (2013b). These lists appear at the beginning of each subsection.

3.1 Th(IV) Solubility Model

This subsection describes the final evaluations by Brush and Domski (2013b) of the experimentally measured Th(IV) solubilities from the published papers and unpublished reports that they identified for their analysis. It documents which measured solubilities they included and excluded, and why they included or excluded them. This subsection then applies criterion G11 to the measured solubilities included by Brush and Domski (2011b), compares the measured Th(IV) solubilities included in this revision with those predicted using the WIPP Th(IV) speciation and solubility model in EQ3/6, and presents our revised results. Finally, it establishes a revised CDF for the differences between our measured and predicted Th(IV) solubilities for the CRA-2014 PA.

The measured Th(IV) solubilities from published papers and unpublished reports evaluated previously by Xiong et al. (2005) and/or Xiong et al. (2011) include: Altmaier et al. (2004, 2005, 2006, 2008), Baston et al. (1996), Felmy et al. (1991, 1997, 2006), Neck et al. (2002, 2003), Östhols et al. (1994), and Rai et al. (1995, 1997, 2000) (see Table 3 and the explanatory text below).

The measured Th(IV) solubilities that Brush and Domski (2013b) evaluated for their analysis include: Borkowski (2012), Borkowski et al. (2012), Brendebach et al. (2007),

Colàs et al. (2011), Fellhauer et al. (2010), Kim et al. (2010), Rai et al. (2005, 2008, 2009, 2011), Vandenborre et al. (2008, 2010), Wood (2005), and Xia et al. (2003) (Table 3 and text).

Altmaier et al. (2004) measured the solubilities of $ThO_2(cr)$ and $ThO_n(OH)_{4-2n} \cdot xH_2O(am)$ in NaCl and MgCl₂ solutions with various ionic strengths (see Table 3 for additional information on these experiments). Xiong et al. (2005) did not consider the results of Altmaier et al. (2004) in their uncertainty analyses of Th(IV) solubilities for the CRA-2004 PABC because this paper came to their attention after they completed their analysis. For the CRA-2009 PABC, Xiong et al. (2011) included all six of the uncentrifuged results from solutions with $I \ge 3$ M from Altmaier et al. (2004, Figure 2) according to criterion S1 (see Section 2 above): two from their 5 M NaCl solutions, two from their 2.5 M MgCl₂ solutions, and two from their 4.5 M MgCl₂ solutions (see Table 3 for additional information on these results). Brush and Domski (2013b) also included these results. All six of these results satisfied criterion G11 and were also included in this revised analysis. However, Xiong et al. (2011) excluded the ultracentrifuged results of Altmaier et al. (2004) because this posttest phase-separation method was significantly different from those employed in the studies used to parameterize the WIPP Th(IV) solubility model (e.g., centrifugation or filtration) and appeared to support a significantly different solubility that established for the model than WIPP (i.e. ultracentrifugation showed that 99% or more of what had been interpreted as dissolved Th(IV) present as the Th(OH)₄(aq) species appeared to be present as the eigencolloid $Th(OH)_4(col)).$ The WIPP Th(IV) model was parameterized in the mid-1990s, before it was realized that the phase-separation methods used then might not have removed all of the $Th(OH)_4(col)$. Thus, Xiong et al. (2011) excluded the ultracentrifuged results of Altmaier et al. (2004) according to Criterion G5 (Section 2). Brush and Domski (2013b) also excluded these results.

Although our model significantly overpredicts the solubility of Th(IV) from about pH = 6 to 13.5 in bicarbonate- (HCO₃⁻-) or carbonate-free (CO₃²⁻-free) solutions, it is still adequate for WIPP compliance-related calculations because both the Th(IV) solubility and the total mobile concentration (dissolved plus colloidal) of Th(IV) predicted under these conditions are higher than they would be based on the ultracentrifuged results of Altmaier et al. (2004). This is because the concentration factors used to calculate the colloidal Th(IV) concentration (solubility) that is about 100 times higher than its actual value, based on the ultracentrifuged results of Altmaier et al. (2004).



Reference	Solution(s) and lonic Strength(s)	pH, pcH, or pmH	Solubility-Controlling Solid	Evaluation and EQ3/6 Run Numbers for Results Included in This Analysis
Altmaier et al. (2004)	(A) 0.5 M NaCl (B) 5 M NaCl (C) 0.25 M MgCl ₂ (I = 0.75 M) (D) 2.5 M MgCl ₂ (I = 7.5 M) (E) 4.5 M MgCl ₂ (I = 13.5 M)	 (A) pcH = 10.4 & 11-13.5 (B) pcH = 10.8 (C) pcH = 9.0 (D) pcH = 8.9 (E) pcH = 8.8 	ThO ₂ (cr) & ThO _n (OH) _{4-2n} xH ₂ O(am), which they referred to as "Th(OH) ₄ (am) or ThO ₂ ·xH ₂ O(am)"	Not considered by Xiong et al. (2005) because this paper came to their attention after completion of their analysis. Six uncentrifuged results (no replicates) obtained at $I \ge 3$ M included by Xiong et al. (2011) and Brush and Domski (2013b): 01ALT04.3i – 06ALT04.3i. All 6 results satisfied criterion G11 and were also included in this revised analysis. Ultracentrifuged results excluded by Xiong et al. (2011) and Brush and Domski (2013b) because the posttest phase-separation method was different than those employed in the studies used to parameterize the Th(IV) model and appeared to support a different model (see text).
Altmaier et al. (2005)	(A) Mostly NaHCO ₃ - Na ₂ CO ₃ -NaCl-NaOH with I = 0.5 M (B) Some with 0.25-2 M Na ₂ CO ₃ & 0.1 M NaOH	(A) pcH = 4.5-7.5 & 8.5-13.5 (B) pcH = 13	Th(OH)₄(am) or ThO₂·xH2O(am)	Not considered by Xiong et al. (2005) because this paper came to their attention after completion of their analysis. Most results excluded by Xiong et al. (2011) and Brush and Domski (2013b) because I < 3 M. All other results excluded because the Th(OH) _y (CO ₃) _z ^{4-y-2z} complex(es) found to be important by Altmaier et al. (2005) are not in the WIPP Th(IV) model. Four results (no duplicates) obtained at I \geq 3 M modeled but excluded by Xiong et al. (2011) and Brush and Domski (2013b).
Altmaier et al. (2006)	NaHCO ₃ -Na ₂ CO ₃ -NaCl with $I = 0.1-4$ M.	pcH = 8–11	Th(OH)₄(am) or ThO₂·xH₂O(am)	Most results excluded by Xiong et al. (2011) and from this analysis because $I < 3$ M. All other results excluded because the Th(OH)(CO ₃) ₄ ⁵⁻ complex found to be important by Altmaier et al. (2006) is not in the WIPP Th(IV) model. Twelve results (no duplicates) obtained at $I \ge 3$ M modeled but excluded by Xiong et al. (2011) and Brush and Domski (2013b).

Reference	Solution(s) and Ionic Strength(s)	pH, pcH, or pmH	Solubility-Controlling Solid	Evaluation and EQ3/6 Run Numbers for Results Included in This Analysis
Altmaier et al. (2008)	(A) $0.2-5.26 \text{ m CaCl}_2$ (I = 0.6-15.78 m) (experiments with Th(IV)) (B) 2.40 m Ca(ClO ₄) ₂ (I = 7.20 m) (Th(IV))	pmH = 9.3-12.1 (all Th(IV))	Solutions with $CaCl_2$ < 1 M (entire pH range) or $CaCl_2 > 1$ M (pH < 11.5): Th(OH) ₄ (am) or ThO ₂ ·xH ₂ O(am). Solutions with $CaCl_2$ > 1 M (pH > 11.5): unidentified, possibly a Ca-Th solid(cr).	Most results excluded by Xiong et al. (2011) and Brush and Domski (2013b) because I < 3 M. Some results excluded because of ultracentrifugation. All other results excluded because the $Ca_4(Th(OH)_8)^{4+}$ complex found to be important by Altmaier et al. (2008) is not in the WIPP Th(IV) model.
Baston et al. (1996)	Nirex (0.02 M)	pH = 10-12	ThO ₂ (am)	Excluded by Xiong et al. (2005, 2011) and Brush and Domski (2013b) because $I < 3$ M.
Borkowski et al. (2012)	 (A) HCO₃⁻/CO₃²⁻-free & HCO₃⁻/CO₃²⁻-bearing GWB, diluted to 95% of its specified concentrations (B) HCO₃⁻/CO₃²⁻-free & HCO₃⁻/CO₃²⁻-bearing ERDA-6, diluted to 95% of its specified concentrations 	(A) pcH = $6.8-10.13$ (GWB: carbonate- free, or 10^{-2} or 10^{-3} M carbonate; & over- & undersaturation) (B) pcH = $7.86-11.04$ (ERDA-6: carbonate- free, or 10^{-2} or 10^{-3} M carbonate; & over- & undersaturation)	Solids precipitated in oversaturation runs not reported. "Th oxyhydroxide" (composition not reported) used in undersaturation runs	Brush and Domski (2013b) included the final results and only the final results from every solubility experiment with both GWB and ERDA-6, because the Th(IV) concentrations generally decreased with time during oversaturation experiments, and increased during undersaturation runs. However, the final results obtained from runs with ERDA-6 appear to be metastable or perhaps more metastable than those from GWB (see text). After application of Criterion G11, 12 results from GWB (7 replicates) and 28 from ERDA-6 (15 replicates) were included in this revised analysis. Used numerical values from Borkowski (2012) (see below). EQ3 files: 02GWB_12.3i - 05GWB_12.3i, and 08GWB_12.3i and 09GWB_12.3i and 01ERD_12.3i - 14ERD_12.3i.

Table 3. Final Evaluations of Lab Studies for Comparisons of Measured and Predicted Th(IV) Solubilities (Cont.).

Reference	Solution(s) and Ionic Strength(s)	pH, pcH, or pmH	Solubility-Controlling Solid	Evaluation and EQ3/6 Run Numbers for Results Included in This Analysis
Borkowski (2012)	Same as above	Same as above	Same as above	Same as above
Brendebach et al. (2007)	0.2-4.5 M CaCl ₂ (I = 0.6–13.5 M) (experiments with Th(IV))	pcH ≈ 9.3-12.1 (all Th(IV)).	Solutions with $CaCl_2$ < 1 M (entire pH range) or $CaCl_2 > 1$ M (pH < 11.5): Th(OH) ₄ (am) or ThO ₂ ·xH ₂ O(am). Solutions with $CaCl_2$ > I M (pH > 11.5): unidentified, possibly a Ca-Th solid(cr).	Not considered by Xiong et al. (2011) because this paper came to their attention after completion of their analysis. The results of Brendebach et al. (2007) were superseded by those of Altmaier et al. (2008) (see above). Therefore, Brush and Domski (2013b) excluded the results of Brendebach et al. (2007) from this analysis for the same reasons that Xiong et al. (2011) excluded those of Altmaier et al. (2008).
Colàs et al. (2011)	0.5 M NaClO ₄ with 10^{-6} to 10^{-1} M gluconate	pcH ≈ 9-13	Th hydroxide (ThO₂·xH₂O)	Not considered by Xiong et al. (2011) because this paper came to their attention after completion of their analysis. All results excluded by Brush and Domski (2013b) because gluconate is not in the WIPP Th(IV) model.
Fellhauer et al. (2010)	1.02–5.26 m CaCl ₂ (I = 3.06–15.78 m) (experiments with Np(IV) and Pu(IV))	pcH ≈ 11-12 (Np(IV)) pcH ≈ 10-12 (Pu(IV))	NpO ₂ ·xH ₂ O(am) or PuO ₂ ·xH ₂ O(am)	Not considered by Xiong et al. (2011) because this paper came to their attention after completion of their analysis. All results excluded by Brush and Domski (2013b) for the same reasons used to exclude those of Brendebach et al. (2007) and Altmaier et al. (2008). Furthermore, there could be systematic differences between the solubilities of Np(IV) and Pu(IV and Th(IV), which was used to parameterize the models; and because the difficulties inherent in maintaining Np and Pu in the +IV oxidation state could have introduced experimental artifacts in these results.

Reference	Solution(s) and Ionic Strength(s)	pH, pcH, or pmH	Solubility-Controlling Solid	Evaluation and EQ3/6 Run Numbers for Results Included in This Analysis
Felmy et al. (1991)	 (A) 0.1 M NaClO₄ (B) 0.6 M NaCl (C) 1.2 M NaCl (D) 3.0 M NaCl (E) 0.6 M KCl 	(A) pcH = 4.9-5.9 (B) pcH = 2.91-10.24 (C) pcH = 2.83-10.19 (D) pcH = 3.82-10.56 (E) pcH = 3.75-7.95	Th(IV) hydrous oxide (all)	Forty-five results (no duplicates) obtained at I = 3 M included in Xiong et al. (2005, 2011) and Brush and Domski (2013b): 01FEL91.3i – 46FEL91.3i. I = 3 M results obtained at pcH \leq 3.6 excluded by Xiong et al. (2005, 2011) and Brush and Domski (2013b) because the solids dissolved before saturation was attained. After application of criterion G11, 18 results (no replicates) were included in this revised analysis. EQ3 files: 15FEL91.3i, 23FEL91.3i – 31FEL91.3i, 39FEL91.3i – 46FEL91.3i, File 16FEL91.3i not included because experimental pH \leq 3.6.
Felmy et al. (1997)	Information provided in previous publications	Information provided in previous publications	Information provided in previous publications	Excluded by Xiong et al. (2005, 2011) and Domski and Brush (2013b) because this paper contains only previously published results (e.g., Felmy et al., 1991).
Felmy et al. (2006)	(A) 0.5 M NaNO ₃ & 0.01 M citrate (B) 3.0 M NaNO ₃ & 0.01 M citrate (C) 6.0 M NaNO ₃ & 0.01 M citrate (D) 0.5 M NaNO ₃ & 10 ⁻⁴ to 10^{-2} M citrate (E) 0.5 M NaNO ₃ & 10 ⁻⁴ to 10^{-2} M citrate	 (A) pcH = 5.2–12.2 (B) pcH = 5.0–11.8 (C) pcH = 4.7–12.1 (D) pcH = 6-8 (E) pcH = 9.5-10.1 	Th(IV) hydrous oxide (ThO ₂ (am)) (all)	All results excluded by Xiong et al. (2011) and Brush and Domski (2013b) because NO_3^- is not in the WIPP Th(IV) model.

Table 3 continued on next page

Reference	Solution(s) and Ionic Strength(s)	pH, pcH, or pmH	Solubility-Controlling Solid	Evaluation and EQ3/6 Run Numbers for Results Included in This Analysis		
Kim et al. (2010)	Distilled H ₂ O; dilute solutions with various combinations of HClO ₄ , Na ₂ CO ₃ , NaClO ₄ and CaCl ₂ ; and naturally occurring, dilute granitic groundwater; 1 = 0-225 mM (all solutions)	pH = 8.6–11.2 (all solutions)	ThO ₂ (cr) (all solutions)	Not considered by Xiong et al. (2011) because this pape came to their attention after completion of their analysis All results excluded by Brush and Domski (2013b) because I < 3 M.		
Neck et al. (2002)	0.5 M NaCl	pcH = 1.11–13.7	Th(OH) ₄ (am) or ThO ₂ ·xH ₂ O(am)	Excluded by Xiong et al. (2005, 2011) and Brush and Domski (2013b) because I < 3 M.		
Neck et al. (2003)	(A) 0.1 M NaCl (B) 0.5 M NaCl	(A) pcH = 1.00-1.75 (B) pcH = 1.03-1.28 & 11.03-13.43	ThO₂(cr) or ThO₂·xH₂O(mcr)	Excluded by Xiong et al. (2005, 2011) and Brush and Domski (2013b) because I < 3 M.		
Östhols et al. (1994)	0.5 NaClO ₄ & 0, 0.1, & 0.97 atm CO ₂	pH = 3.31–7.36 (continuous titrations) & 8.21-10.45 (batch experiments)	ThO ₂ (mcr)	Excluded by Xiong et al. (2005, 2011) and Brush and Domski (2013b) because I < 3 M.		

Reference	Solution(s) and Ionic Strength(s)	pH, pcH, or pmH	Solubility-Controlling Solid	Evaluation and EQ3/6 Run Numbers for Results Included in This Analysis
Rai et al. (1995)	 (A) 0.005-0.25 M NaHCO₃ (B) 0.1-2.0 M Na₂CO₃ (C) 0.1-2.0 M Na₂CO₃ & 0.1 M NaOH (D) 0.1-0.5 M NaOH & 1.0 M Na₂CO₃ 	Not provided	ThO ₂ (am)	Not considered by Xiong et al. (2005) because this paper came to their attention after completion of their analysis. All results excluded by Xiong et al. (2011) and Brush and Domski (2013b) because the Th(OH) _y (CO ₃) _z ^{$4-y-2z$} complexes found to be important by Altmaier et al. (2005) under the conditions used by Rai et al. (1995) are not in the WIPP Th(IV) model.
Rai et al. (1997)	 (A) 0.2 m NaCl (used only for U solubilities) (B) 1.0 m NaCl (used only for U solubilities) (C) 3.2 m NaCl (data from Felmy et al., 1991) (D) 4.0 m NaCl, (E) 6.0 m NaCl, (F) 1.0 m MgCl₂ (I = 3 m), (G) 1.82 m MgCl₂ (I = 5.46 m), & (H) 3.0 m MgCl₂ (I = 9 m) 	 (A) U not included in this analysis (B) U not included in this analysis (C) See entry for 3.0 M NaCl for Felmy et al. (1991) in this table (D) pcH = 3.02-5.16 (E) pcH = 3.32-5.28 (F) pcH = 3.84-4.94 (G) pcH = 3.17-5.46 (H) pcH = 3.52-5.65 	ThO ₂ (am)	Not considered by Xiong et al. (2005) because this paper came to their attention after completion of their analysis. Eighty-nine results (8 duplicates) obtained at $I \ge 3$ M included by Xiong (2011) and Brush and Domski (2013b): 01RAI97.3i - 80RAI97.3i. However, one EQ3 run (29RAI97.3i) did not converge, so only 88 results used. Results obtained from the most acidic solutions in each set of experiments (see Table 6) excluded by Xiong et al. (2011) and Brush and Domski (2013b) because the solids dissolved before saturation was attained. These data do not satisfy criterion G11 and were not included in this revised analysis.

Reference	Solution(s) and Ionic Strength(s)	pH, pcH, or pmH	Solubility-Controlling Solid	Evaluation and EQ3/6 Run Numbers for Results Included in This Analysis		
Rai et al. (2000)	(A) 0.1 M NaCl (B) 0.1 M NaCl (C) 0.1 M NaCl	(A) pH = 2.0-4.7 & 1.5-2.2 (B) pH= 4.2-5.1 & 2.0-2.1 (C) pH = 1.9-4.2 &	(A) ThO ₂ (am) & ThO ₂ (am to cr) (B) ThO ₂ (am) & ThO ₂ (am to cr) (C) ThO ₂ (am) &	Not considered by Xiong et al. 2005) because this paper came to their attention after completion of their analysis Most results excluded by Xiong et al. (2011) and Brush and Domski (2013b) because $I < 3$ M. All other results excluded because $T = 90$ °C.		
	(D) 0.1 M NaCl	1.9-3.0 (D) $pH = 1.3-3.6$	ThO ₂ (am to cr) (D) ThO ₂ (cr)			
	(E) 0.1 M NaC1	(E) $pH = 1.4-3.5$	(E) $ThO_2(cr)$			
Rai et al. (2005)	$HCO_3^{-}/CO_3^{2^-}$ -free & $HCO_3^{-}/CO_3^{2^-}$ -bearing (up to 6.2 m) solutions with numerous other solutes dissolved from borosilicate glass	pH ≈ 0.5–12	Not determined. May have included a Th- & Si-bearing solid(s)	Not considered by Xiong et al. 2005, 2011) because this paper came to their attention after completion of their analyses. Excluded by Brush and Domski (2013c) because: (1) several components of the glass, especially SiO ₂ and Al ₂ O ₃ , are not in the WIPP model; (2) the solu- bility-controlling solid(s) were not identified; and (3) many results were obtained from solutions with I < 3 M or m.		

Reference	Solution(s) andReferenceIonic Strength(s)		Solubility-Controlling Solid	Evaluation and EQ3/6 Run Numbers for Results Included in This Analysis			
Rai et al. (2008)	(A) 0.0025-0.1 M Na ₂ SiO ₃ , with "carbonate impurities"	(A) pH 11, 12, or 13	ThO ₂ (am) containing NO_3^- impurities form preparation and SiO ₂ ,	Not considered by Xiong et al. 2011) because this paper came to their attention after completion of their analysis. Excluded by Brush and Domski (2013b) because			
	(B) 0.0025-0.14 M Na ₂ SiO ₃ , HCO ₃ ^{-/} CO ₃ ²⁻ -free	(B) pH = 10, 12, or 13	probably sorbed to the surface (all experiments)	the Th(OH) ₃ · (H ₃ SiO ₄) ₃ ²⁻ complex found to be important by Rai et al. (2008) is not in the WIPP Th(IV) model.			
	(C) 0.0356 M Na ₂ SiO ₃ , (fixed)	(C) pH = 10–13.3 (variable)					
	(D) 0.0157 M Na ₂ SiO ₃ , (fixed)	(D) pH = 10–13.3 (variable)					
Rai et al. (2009)	(A) 0.0002–0.2 m ISA	(A) pH ≈ 6 or 12 (fixed)	ThO ₂ (am) (all experiments)	Not considered by Xiong et al. 2011) because this paper came to their attention after completion of their analysis Excluded by Brush and Domski (2013b) because			
	(B) 0.008 or 0.08 m ISA	(B) pH = 4.5–12		the Th(OH)ISA ²⁺ , Th(OH) ₃ (ISA) ₂ ⁻ , and Th(OH) ₄ (ISA) ₂ ²⁺ complexes found to be important by Rai et al. (2009) are not in the WIPP Th(IV) model.			
Rai et al. (2011)	Information provided in previous publications	Information provided in previous publications	Information provided in previous publications	Not considered by Xiong et al. (2011) because this paper came to their attention after completion of their analysis. Excluded by Brush and Domski (2013b) because this paper contains only previously published results.			
Vandenborre et al. (2008)	0.01 M NaCl	pH = 3.0 or 4.0	ThO ₂ (cr) spheres with an average diameter of 500 μ m	Not considered by Xiong et al. (2011) because this paper came to their attention after completion of their analyses. Excluded by Brush and Domski (201b) because $I < 3$ M.			

Reference	Solution(s) and Ionic Strength(s)	pH, pcH, or pmH	Solubility-Controlling Solid	Evaluation and EQ3/6 Run Numbers for Results Included in This Analysis
Vandenborre et al. (2010)	0.01 M NaCl	pH = 0.0 to 5.2	ThO ₂ (cr) spheres with an average diameter of 500 μm	Not considered by Xiong et al. (2011) because this paper came to their attention after completion of their analysis. Excluded by Brush and Domski (2013b) because $I < 3$ M
Wood (2005)	(A) 0.1, 1.0, & 3.0 M NaNO ₃ ; 100–500 μM citrate, EDTA; & oxalate	(A) pH ≈ 2–10.5	(A) CeO ₂ (crystallinity and hydration state unspecified)	Not considered by Xiong et al. (2005, 2011) because this paper came to their attention after completion of their analyses. Excluded by Brush and Domski (2013b) because NO_3^- is not in the WIPP Th(IV) model.
	(B) 0.1 M NaNO ₃ ; 100–500 μM citrate, EDTA; & oxalate	(B) pH ≈ 3–12	(B) "A relatively poorly crystalline form of ThO ₂ "	
Xia et al. (2003)	(A) 0.5 M NaNO ₃ & 0.01 M EDTA (B)) 3.0 M NaNO ₃ & 0.01 M EDTA (C)) 6.0 M NaNO ₃ & 0.01 M EDTA (C)) 0.5 M NaNO ₃ & 0.01 M EDTA (D)) 0.5 M NaNO ₃ & 10 ⁻⁵ -10 ⁻² M EDTA (E) 0.5 M NaNO ₃ & 10 ⁻⁵ -10 ⁻² M EDTA (F) 0.5 M NaNO ₃ & 10 ⁻⁵ -10 ⁻² M EDTA (G) 0.5 M NaNO ₃ & 10 ⁻⁵ -10 ⁻² M EDTA	pcH = 4.3-11.8 pcH = 6.7-12.8 pcH = 6.8-13.2 pcH = 8 pcH = 10 pcH = 8 pcH = 10	Th(IV) hydrous oxide (ThO ₂ (am)) (all)	Not considered by Xiong et al. (2005, 2011) because this paper came to their attention after completion of their analyses. Excluded by Brush and Domski (2013b) because NO_3^- is not in the WIPP Th(IV) model.

Altmaier et al. (2005) measured the solubilities of Th(OH)₄(am) and ThO₂·xH₂O(am), mostly with NaHCO₃-Na₂CO₃-NaCl solutions with I = 0.5 M. They also carried out some experiments with Na₂CO₃ solutions with various ionic strengths (Table 3). Xiong et al. (2005) did not consider this paper because it came to their attention after they completed their analysis. The results in Altmaier et al. (2005, Figure 4b) led Xiong et al. (2011) to suspect that, in some of their experiments, the ionic strength might have equaled or exceeded 3 M. Therefore, they used Data Thief to extract the numerical values of the plotted CO_3^{2-} concentrations from their Figure 4b, and calculated the ionic strengths of the solutions. Xiong et al. (2011) found that four of these solutions had I \geq 3 M. According to criterion S1 (see Section 2), Xiong et al. (2005) would have included these results in their Th(IV) uncertainty range and probability distribution. They excluded them, however, because Altmaier et al. (2005) concluded that, at high CO_3^{2-} concentrations (log[CO_3^{2-}] greater than about -0.5) in their Figure 4b, the dominant aqueous species is (are) ThOH(CO_3)4⁵⁻ (or ThOH(CO_3)4⁵⁻ and Th(OH)₂(CO_3)4⁶⁻), neither of which is included in the WIPP Th(IV) model. Thus, Xiong et al. (2011) used criterion G9 to exclude these results. Brush and Domski (2013b) also excluded these results.

The WIPP Th(IV) model includes the following $Th(OH)_y(CO_3)_z^{4-y-2z}$ complexes: Th(CO₃)₅⁶⁻, Th(OH)₃CO₃, and Th(OH)₄(aq). It does not include ThOH(CO₃)₄⁵⁻ and Th(OH)₂(CO₃)₂²⁻, both of which were identified by Altmaier et al. (2005, Figure 5) as important at high CO_3^{2-} concentrations. Furthermore, it does not include Th(OH)₂CO₃(aq) and Th(OH)₄CO₃²⁻, which – although not identified as important in their Figure 5 – were quantified by Altmaier et al. (2005): Nevertheless, the WIPP Th(IV) model is still adequate for WIPP compliance-related calculations. This is because none of the important $Th(OH)_v(CO_3)_z^{4-y-2z}$ complexes with z > 0 approaches the Th(OH)₄(aq) concentrations predicted by Brush and Domski (2013a, Tables 7 and 8) for the minimum brine volume for the CRA-2014 PA $(4.55 \times 10^{-8} \text{ M for GWB and } 4.75 \times 10^{-8} \text{ M for ERDA-6})$ until the log of the CO₃²⁻ concentration exceeds about -1 (see Altmaier et al., 2005, Figure 4b). By contrast, the (TIC) concentrations (essentially the sum of the concentrations of HCO_3^- and CO_3^{2-}) predicted by Brush and Domski (2013a, Tables 5 and 6), 0.379 mM for GWB and 0.455 mM for ERDA-6, for the minimum brine volume were more than two orders of magnitude lower than that threshold. The reason why the TIC concentrations predicted for these WIPP brines are too low to form important Th(OH)_v(CO₃)_z^{4-y-2z} complexes with z > 0 is because the brucite-hydromagnesite carbonation reaction will buffer the TIC at the values given above.

Altmaier et al. (2006) determined the solubilities of Th(OH)₄(am) and ThO₂·xH₂O(am) in NaHCO₃-Na₂CO₃-NaCl solutions with various ionic strengths (Table 3). Xiong et al. (2011) excluded most of their results according to criterion S1 (Section 2) because they were obtained using NaHCO₃-Na₂CO₃-NaCl solutions with ionic strengths less than 3 M. However, they found 12 results from experiments with I \geq 3 M (see Altmaier et al., 2006, Figure 2). Xiong et al. (2011) used criterion G9 to exclude these data because the WIPP Th(IV) model does not include the ThOH(CO₃)₄⁵⁻ complex, which Altmaier et al. (2006) found to be the dominant aqueous species. Brush and Domski (2013b) also excluded these results.

Nevertheless, the WIPP Th(IV) model is still adequate for WIPP compliance-related calculations for the reasons described above in the discussion of the data from Altmaier et al. (2005) (see above).

Altmaier et al. (2008) measured the solubilities of Th(OH)₄(am) and ThO₂·xH₂O(am), a (possible) Ca-Th solid, and Zr(IV) or Pu(IV) solids in CaCl₂ with various ionic strengths and in 2.40 m Ca(ClO₄)₂ (Table 3). Xiong et al. (2011) used criterion G6 (Section 2) to exclude most of their results because they were obtained from experiments with Zr(IV) or Pu(IV). They used criterion S1 to exclude the Th(IV) solubilities from 0.20 and 0.51 m CaCl₂ solutions. Finally, Xiong et al. (2011) used criterion G9 to exclude the Th(IV) solubilities from 1.02, 2.11, and 5.26 m CaCl₂ solutions (I = 3.06, 6.33, and 15.78 m, respectively) because Altmaier et al. (2008) concluded that, at high CaCl₂ concentrations (> 0.5 m for pcH = 11-12 according to their abstract, but > 0.2 m at pH = 12 from their Figure 5), the dominant aqueous species was Ca₄(Th(OH)₈)⁴⁺, which is not included in the WIPP Th(IV) model. Brush and Domski (2013b) excluded these results for the same reasons.

Nonetheless, the Th(IV) model is still adequate for WIPP compliance-related calculations because the Ca concentrations predicted by Brush and Domski (2013a, Tables 5 and 6) for the minimum volumes of GWB and ERDA-6 are 0.0111 and 0.0116 M, respectively. Figure 5 in Altmaier et al. (2008) demonstrates that both the total dissolved Ca concentration and the pcH must be much higher than the values expected in WIPP brines to increase the concentration of $Ca_4(Th(OH)_8)^{4+}$ to values greater than those expected when Th(OH)_4(aq) or Th(OH)_4(col) is the dominant species. For example, if the CaCl₂ concentration is 0.51 m, the pcH must be 12 or higher to increase the Ca₄(Th(OH)₈)⁴⁺ concentration to values greater than those expected when Th(OH)_4(col) is the dominant mobile species. If the CaCl₂ concentration is 1.02 m, the pcH must be 11-11.5 or higher to increase the Ca₄(Th(OH)₈)⁴⁺ concentrations greater than those expected when Th(OH)₄(col) is the dominant mobile species. Therefore, the absence of $Ca_4(Th(OH)_8)^{4+}$ from the Th(IV) model does not affect its validity under expected WIPP conditions.

Although $Ca_4(Th(OH)_8)^{4+}$ and the analogous Am(III) and Cm(III) complexes described below will not be important in the WIPP, these complexes could be important in salt repositories with brines that have higher $CaCl_2$ concentrations. Brendebach et al. (2007) (see below) were, to the best of our knowledge, the first investigators to report the formation of, characterize, and assess the importance of these ternary Ca-M(IV)-OH complexes, in which the metal M can be Th(IV) or Zr(IV). Subsequently, Rabung et al. (2008) and Neck et al. (2009) reported, characterized, and evaluated analogous Ca-M(III)-OH complexes, in which M can be Am(III) or Cm(III). Subsection 3.2 and Table 8 (see below) describe the results of Rabung et al. (2008) and Neck et al. (2009).

Baston et al. (1996) studied the solubility of $ThO_2(am)$ in a low-ionic-strength solution (Table 3). Xiong et al. (2005) excluded the results of Baston et al. (1996) from their analysis, because they were obtained from solutions with I < 3 M (see criterion S1 in Section 2). Xiong et al. (2011) and Brush and Domski (2013b) also used criterion S1 to exclude them.

Borkowski et al. (2012) measured the solubilities of Th(IV) in the synthetic WIPP brines GWB and ERDA-6, diluted to 95% of their specified concentrations to avoid coprecipitation of Th(IV) by evaporite minerals (Table 3). (Small changes in conditions, especially temperature, during solubility experiments can cause precipitation of solids such as evaporite minerals and coprecipitation of actinide elements if the solutions are saturated with these solids under

the conditions used for these tests.) Borkowski et al. (2012) used carbonate-free GWB and ERDA-6, and GWB and ERDA-6 with 10⁻³ or 10⁻² M TIC. Brush and Domski (2013b) considered all of the results of Borkowski et al. (2012) for inclusion in their uncertainty analysis of Th(IV) for the CRA-2014 PA according to criterion G6 (see Section 2 above). These results demonstrated that the Th(IV) concentrations generally decreased with time during their oversaturation experiments, and increased during their undersaturation runs. Therefore, Brush and Domski (2013b) included all of the results obtained from their final samples from each experiment, and excluded those from samples taken prior to their final samples. The final results obtained from the oversaturation runs with ERDA-6 are generally greater than those from the oversaturation runs with GWB, probably because the former were carried out for 2 years and the latter for 4 years. Nevertheless, Brush and Domski (2013b) included all of the final results from experiments with both GWB and ERDA-6. Subsequent to the release of Borkowski et al. (2012), Borkowski (2012) provided numerical values of the brine compositions, pcH, Th(IV) concentrations, and experimental durations shown graphically by Borkowski et al. (2012). Brush and Domski (2013b) used these numerical values from Borkowski (2012) for the EQ3/6 runs for this analysis. After application of criterion G11, 12 results from GWB and 28 results from ERDA-6 were included in this revised analysis.

Brendebach et al. (2007) carried out solubility experiments with $Th(OH)_4(am)$ and $ThO_2 xH_2O(am)$, a (possible) Ca-Th solid, and Zr(IV) solids in CaCl₂ solutions with various ionic strengths (Table 3). As noted above, this was the first paper to identify Ca-M(IV)-OH complexes, in which the metal M can be Th(IV) or Zr(IV). Xiong et al. (2011) did not consider the results of Brendebach et al. (2007) because this paper came to their attention after they completed their analysis. Altmaier et al. (2008) (see above) expanded the study of Brendebach et al. (2007) were superseded by those of Altmaier et al. (2008), Brush and Domski (2013b) excluded the results of Brendebach et al. (2008) (see above).

Colàs et al. (2011) conducted solubility experiments with Th hydroxide (for which they provided the composition $ThO_2 \cdot xH_2O$) in 0.5 M NaClO₄ solutions with 10^{-6} to 10^{-1} M gluconate and pcH \approx 9–13 (Table 3). Xiong et al. (2011) did not consider this paper because it came to their attention after they completed their analysis. Brush and Domski (2013b) used criterion G9 (Section 2) to exclude their results because gluconate is not in the WIPP Th(IV) model.

Fellhauer et al. (2010) measured the solubilities of Np(IV) and Pu(IV) solids in CaCl₂ solutions with various ionic strengths (Table 3). Fellhauer used 2 mM Na₂S₂O₄ or metallic Fe powder to establish reducing conditions and maintain Np and Pu in the +IV oxidation state. Thus, they extended the oxidation-state-analog approach of Brendebach et al. (2007) and Altmaier et al. (2008), both of whom used Th(IV), to a direct evaluation of Ca-M(IV)-OH complexes, in which the metal M was Np(IV) or Pu(IV). Xiong et al. (2011) did not consider these results because they found this paper after they completed their analysis. Brush and Domski (2013b) excluded these results from our uncertainty analysis for the same reasons that they excluded those of Brendebach et al. (2007) and Altmaier et al. (2008). Furthermore, Brush

and Domski (2013b) used criterion G6 (Section 2) to exclude the results of Fellhauer et al. (2010), because they were obtained from experiments with Np(IV) or Pu(IV).

Felmy et al. (1991) measured the solubility of Th(IV) hydrous oxide in 0.1 M NaClO₄, in NaCl solutions of various ionic strengths, and in 0.6 M KCl (Table 3). Xiong et al. (2005, Subsection 5.2.3) established criterion S1 based on their comparison of the solubilities measured by Felmy et al. (1991) and those predicted using the WIPP Th(IV) solubility model, and excluded most of them because they were obtained using solutions with I < 3 M. Furthermore, inspection of the remaining results showed that those obtained at the most acidic values of pH were essentially independent of pcH. Xiong et al. (2005, p. 12) noted that:

[T]his behavior can be explained by noting that in ... the experiments at $pcH \le \sim 3.6$..., the Th(IV)-bearing solids were consumed before saturation was attained (see Felmy et al., 1991, p. 298). Therefore, the constant Th(IV) concentration of slightly less than 10^{-2} M in the experiments reflects the fact that the constant quantity of Th(IV) added at the start of these experiments was insufficient to saturate these solutions under these conditions... Thus, the concentrations in the experiments at $pH \le \sim 3.6$... are not solubilities. Because the concentrations measured by Felmy et al. (1991) at $pcH \le \sim 3.6$ are not solubilities, we excluded them from the revised An(IV) comparison.

Therefore, Xiong et al. (2005) excluded these acidic results, and Xiong et al. (2011) incorporated the reason for excluding these and similar results from other studies into criterion G7 (see Section 2; the descriptions of the results of Rai et al., 1997, in Subsection 3.1 and Tables 4 and 5; and the discussions of the results of Rao et al., 1996a, in Subsection 3.2 and Table 8). After applying criteria G7 and S1, Xiong et al. (2005) included 45 results from Felmy et al. (1991) in their analysis; Xiong et al. (2011) and Brush and Domski (2013b) included the same 45 results in their analyses. After application of criterion G11, 18 results were included in this analysis.

Felmy et al. (1997) reviewed the development of models for the solubility of Th(IV) solids in high-ionic-strength solutions. Xiong et al. (2005, 2011) excluded all of the results in Felmy et al. (1997) from their analyses according to criterion G10, because this paper is a literature review and does not conduct any new data. Brush and Domski (2013b) also excluded these results.

Felmy et al. (2006) studied the solubility of $ThO_2(am)$ in NaNO₃ solutions with various ionic strengths and various concentrations of the organic ligand citrate (Table 3). Xiong et al. (2011) used criterion G9 (Section 2) to exclude their results because NO₃⁻ is not in the WIPP Th(IV) model. Brush and Domski (2013b) excluded these results too.

Kim et al. (2010) measured the solubility of $ThO_2(cr)$ in dilute solutions with various combinations of HClO₄, Na₂CO₃, NaClO₄, and CaCl₂ (Table 3). They also samples of a naturally occurring, dilute granitic groundwater. Xiong et al. (2011) did not consider the results of Kim et al. (2010) because this paper came to their attention after they completed

their analysis. Brush and Domski (2013b) excluded their results according to criterion S1 (Section 2), because they were all obtained using solutions with I < 3 M.

Neck at al. (2002) measured the solubilities of $Th(OH)_4(am)$ and $ThO_2 \cdot xH_2O(am)$ in 0.5 M NaCl solutions (Table 3). Xiong et al. (2005) excluded their results according to criterion S1 (Section 2), because they were obtained using solutions with I < 3 M. Xiong et al. (2011) also used criterion S1 to exclude them. Brush and Domski (2013b) excluded them too.

Neck et al. (2003) performed a solubility study with $Th(OH)_4(am)$ and $ThO_2 \cdot xH_2O(am)$ in 0,1 and 0.5 M NaCl solutions (Table 3). Xiong et al. (2005, 2011) used criterion S1 (Section 2) to exclude their results. Brush and Domski (2013b) also excluded their results.

Östhols et al. (1994) studied the solubility of $ThO_2(mcr)$ in 0.5 M NaClO₄ under atmospheres with various concentrations of CO₂ (Table 3). Xiong et al. (2005, 2011) used criterion S1 (Section 2) to exclude their results. Brush and Domski (2013b) also excluded them.

Rai et al. (1995) measured the solubilities of ThO₂(am) and UO₂(am) in a variety of NaHCO₃, Na₂CO₃, and Na₂CO₃ and NaOH solutions with various ionic strengths (Table 3). Xiong et al. (2005) did not consider the results of Rai et al. (1995) because this paper came to their attention after they completed their analysis. Based on their previous work and that of others, Rai et al. (1995, p. 1146) expected that "... it is probably that [sic] the dominant aqueous species of Th and U in these high-carbonate solutions are Th(CO₃)₅⁶ and U(CO₃)₅⁶, respectively." However, the response of the Th and U solubilities to concentration variations of OH⁻ in their experiments led them to speculate that the complexes U(OH)₃CO₃⁻ or U(OH)₄CO₃²⁻ or, from the oxidation-state analogy, Th(OH)₃CO₃⁻ or Th(OH)₄CO₃²⁻ could be dominant (Rai et al., 1995, p. 1148). Subsequently, Altmaier et al. (2005, 2006) demonstrated that, under the conditions of Rai et al. (1995), the important Th(OH)_y(CO₃)_z^{4-y-2z} complexes are actually Th(CO₃)₅⁶⁻, Th(OH)(CO₃)₄⁻⁵, and Th(OH)₂(CO₃)₂⁻². Although the WIPP Th(IV) model includes Th(CO₃)₅⁶⁻, it does not include Th(OH)(CO₃)₄⁻⁵ and Th(OH)₂(CO₃)₂⁻². Therefore, Rai et al. (1995) used conditions under which there are two important Th(OH)_y(CO₃)_z^{4-y-2z} species that are not in the WIPP Th(IV) model and Xiong et al. (2011) used criterion G9 (Section 2) to exclude them. Brush and Domski (2013b) also excluded these results.

Rai et al. (1997) carried out solubility experiments with $Th(OH)_2(am)$ in 4.0 and 6.0 M NaCl solutions and in MgCl₂ solutions with various ionic strengths (Table 3). Xiong et al. (2005) did not consider this paper because it came to their attention after they completed their analysis. Initially, Xiong et al. (2011) decided to include all of their results obtained at $I \ge 3$ M according to criterion S1 (Section 2). However, many of the results of Rai et al. (1997) displayed the same feature observed by Xiong et al. (2005) in the results of Felmy et al. (1991) (see above): the results obtained at the most acidic pH values were independent of pcH. Therefore, Xiong et al. (2011) used criterion G7 to exclude the most acidic results (see Table 4 on the next page), which left 88 results for inclusion in their analysis. Brush and Domski (2013b) also used these results in their analysis. These data do not satisfy criterion G11 and are not included in this analysis.

Rai et al. (2000) conducted solubility experiments with $ThO_2(am)$ and $ThO_2(cr)$ in 0.1 M NaCl solutions (Table 3). Xiong et al. (2005) did not consider these results because they found this paper later. Xiong et al. (2011) used criteria S1 and G5 (Section 2) to exclude them. Criterion G5 also applied to some of these results because they were obtained at 90 °C, a temperature higher than that expected in the WIPP. Brush and Domski (2013b) excluded these results for the same reasons.

Rai et al. (2005) determined the concentration of Th(IV) dissolved from powdered, phosphate-free borosilicate glass in various solutions (Table 3) at 23 ± 2 °C "to determine whether the observed thorium concentrations exhibit an equilibrium phenomenon and to ascertain whether existing thermodynamic data can be used to interpret these results." The glass contained 24 components, most of which are not in the WIPP Th(IV) model. The most important of these were Al₂O₃ (5.70 wt %) and SiO₂ (48.60 wt %). The KHCO₃ and/or K₂CO₄ concentrations of these solutions varied from 0 to 6.2 m; the pH from about 05 to 12. Xiong et al. (2005, 2011) did not consider the results of Rai et al. (2005) because this paper came to their attention after they completed their analysis. Brush and Domski (2013b) used criterion G9 (Section 2) to exclude the results of this study because SiO₂, Al₂O₃, and numerous other components of the glass are not in the WIPP Th(IV). Brush and Domski (2013b) also used criterion G7, because Rai et al. (2005) were unable to identify the solubility-controlling solid(s) in their experiments. Criterion S1 also applies in many cases, because many of the results of this study were obtained from solutions with I < 3 M or m.

Rai et al. (2008) studied the solubility of Th(IV) in basic solutions of 0.0004 to 0.14 M Na₂SiO₃ under basic conditions at 22 ± 2 °C. (Note that the range of Na₂SiO₃ concentrations given in their abstract, from which Brush and Domski (2013b) obtained the preceding range, does not agree with that in their Table 1, from which Brush and Domski (2013b) obtained the ranges in their Table 3). Xiong et al. (2011) did not consider this paper because it came to their attention after they completed their analysis. Brush and Domski (2013b) used criterion G9 to exclude these data because the WIPP Th(IV) model does not include the Th(OH)₃· (H₃SiO₄)₃²⁻ complex, which Altmaier et al. (2006) found to be the dominant aqueous species. Criterion G7 also applies because the solubility-controlling solid(s) in their experiments seems to contain impurities from preparation and sorbed species from their solutions that could alter its behavior from that of the ThO₂(am) used to start their runs.

Rai et al. (2009) measured the solubility of ThO₂(am) in 0.0002 to 0.2 m isosaccharinate (ISA) solutions under acidic, neutral, and basic conditions at 22 ± 2 °C (Table 3). Xiong et al. (2011) did not consider these results because they found this paper after they completed their analysis. Brush and Domski (2013b) used criterion G9 to exclude these results because the WIPP Th(IV) model does not include the Th(OH)ISA²⁺, Th(OH)₃(ISA)₂⁻, and Th(OH)₄(ISA)₂²⁻ complexes, which Rai et al. (2009) found to be the dominant in their study.

Run Time (days)	Solution	Maximum pcH	Number of Results Excluded	Figure in Rai et al. (1997) with Excluded Data
7	4.0 m NaCl	3.9	7	Figure 6
51	4.0 m NaCl	4.0	5	Figure 6
91	4.0 m NaCl	4.0	5	Figure 6
8	6.0 m NaCl	3.9	3	Figure 2
125	6.0 m NaCl	3.85	3	Figure 2
159	6.0 m NaCl	3.8	5	Figure 2
7	1.8 m MgCl ₂	4.0	5	Figure 3
73	1.8 m MgCl ₂	4.0	4	Figure 3
8	3.0 m MgCl ₂	3.9	2-	Figure 4

Table 4.	Results from	Rai et	al.	(1997)	Excluded	from	Th(IV)	Comparisons	Because	of
	Complete Disa	solutior	n of S	Solids.						

Rai et al. (2011) reviewed the literature review to determine if using the solubilities of radioelements based on knowledge of the important solubility-controlling solids provides a feasible alternative to the use of numerical predictions of these solubilities based on the results of empirical leaching experiments. Xiong et al. (2011) did not consider these results because they found this paper later. Brush and Domski (2013b) used criterion G10 to exclude these results, because Rai et al. (2011) carried out a literature review and did not conduct any new experiments.

Vandenborre et al. (2008) measured the solubility of well crystallized ²³²ThO₂ with reproducible surface properties in 0.01 M NaCl at a pH 3.0 or 4.0. They spiked their solutions with ²²⁹Th to determine the rate of attachment of this isotope to the solid surfaces, and measured the dissolved ²³²Th concentration to establish the rate at which this isotope dissolved. Vandenborre et al. (2010) expanded the range of pH to 0.0 < pH < 5.2. Perhaps the most interesting results of this work are: (1) Figure 1 of Vandenborre et al. (2010), which shows the extent of the scatter in Th(IV) solubilities reported in the literature from 1964 through 2006; (2) the results of Vandenborre et al. (2010, Figure 1), which are lower than most of the values form the literature but show almost no dependence on pH over the range 0 < pH < 4; and



(3) the values of the logs of the solubility products of Th(IV) solids, which vary from -56.9 for ThO₂(cr) to -45.5 for ThO_x(OH)_y(H₂O)_z(s), a range of over 11 orders of magnitude. Xiong et al. (2011) did not consider the results of Vandenborre et al. (2008, 2010) because these papers came to their attention after they completed their analysis. Brush and Domski (2013b) used criterion S1 (Section 2) to exclude the results of Vandenborre et al. (2008) and Vandenborre et al. (2010), because they were obtained from experiments carried out in solutions with I < 3 M.

Wood (2005) measured the solubilities of CeO_2 and ThO_2 in NaNO₃ solutions with and without the organic ligands citrate, desferrioxamine B, EDTA, and oxalate at 22 or 25 °C (Table 3). Xiong et al. (2005, 2011) did not consider this paper because it came to their attention after they completed their analysis. Brush and Domski (2013b) used criterion G9 (Section 2) to exclude their results because NO₃ is not in the WIPP Th(IV) model.

Finally, Xia et al. (2003) studied the solubility of $ThO_2(am)$ in NaNO₃ solutions with various ionic strengths and various concentrations of the organic ligand EDTA (Table 3). Xiong et al. (2005, 2011) did not consider these results because they found this paper later. Brush and Domski (2013b) used criterion G9 (Section 2) to exclude their results because NO₃⁻ is not in the WIPP Th(IV) model.

Figure 1 provides the histogram for our comparisons of experimentally measured and predicted Th(IV) solubilities for the CRA-2014 PA. The current comparison includes a total of 64 measured and predicted solubilities. The current distribution peaks at Bin 1.05 with a frequency of 17 comparisons, with a secondary peak at Bin -1.2 with 10 comparisons. Table 5 summarizes the statistical parameters of the current frequency distribution.

Inspection of Figure 1 shows that the WIPP Th(IV) model, the EQ3/6 code, and the DB Data0.FM1 overpredicted 14 of the 64 measured solubilities included in the current analysis and underpredicted 50. Therefore, Figure 1 contains fewer negative values of the difference between the measured and predicted solubility ($D = \log_{10}(S_m) - \log_{10}(S_p)$) than positive values. The mean and median values of the log of the sampled uncertainty factor used to adjust the baseline Th(IV) solubilities are 0.673 and 1.029, respectively (Table 6), and the mean and median values of the sampled uncertainty factor are $10^{0.673} = 4.708$ and $10^{1.029} = 10.703$. (The WIPP PA codes use actinide solubilities that are the products of the baseline solubilities and the sampled uncertainty factors).

Table 6 provides values of cumulative distribution function (CDF) for Th(IV) for the CRA-2014 PA. PA will use this CDF to define the solubility multiplier SOLVAR for the oxidation-state IV model, material SOLMOD4 in the WIPP PA Parameter Database for the CRA-2014 PA. Brush et al. (2005) described the PA implementation of these results in detail.

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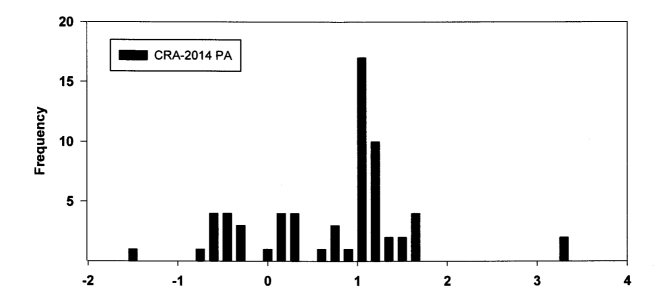


Figure 1. Histogram of the difference $D = \log_{10} S_{\rm m} - \log_{10} S_{\rm p}$ for Th(IV) for the CRA-2014 PA (this revised analysis).

Table 5.Statistical Properties of the Distribution of D = log10(Sm) - log10(Sp) for Th(IV) for
the CRA-2014 PA (This Revised Analysis).

Analy	rsis	Mean	Median	Standard Deviation	Minimum	Maximum
CRA-20	14 PA	0.673	1.029	0.869	-1.52	3.19



Table 6.	CDF of the Differences between Measured and Predicted Th(IV) Solubilities for
	the CRA-2014 PA (This Revised Analysis). This CDF Defines the Solubility
	Multiplier SOLVAR for the Oxidation-State IV Model Material SOLMOD4 in
	the WIPP PA Parameter Database.

-1.51771	0.0156
-0.80033	0.0313
-0.72033	0.0469
-0.71032	0.0625
-0.61033	0.0781
-0.61033	0.0938
-0.56033	0.1094
-0.54032	0.1250
-0.52033	0.1406
-0.47768	0.1563
-0.37699	0.1719
-0.33032	0.1875
-0.30327	0.2031
-0.08032	0.2188
0.02967	0.2344
0.08509	0.2500
0.11565	0.2656
0.14961	0.2813
0.15967	0.2969
0.17237	0.3125
0.26967	0.3281
0.28967	0.3438
0.45967	0.3594
0.67967	0.3750
0.68967	0.3906
0.74967	0.4063
0.77783	0.4219
0.97783	0.4375
1.02567	0.4531
1.02571	0.4688
1.02653	0.4844
1.02863	0.5000

Table 6 continued on next page

Informa³⁹6f⁷³on Only

D	<i>P(D<x)< i="">CRA-2014 PA</x)<></i>
1.03036	0.5156
1.03252	0.5313
1.03516	0.5469
1.03730	0.5625
1.03736	0.5781
1.03798	0.5938
1.04386	0.6094
1.04576	0.6250
1.04779	0.6406
1.04922	0.6563
1.04986	0.6719
1.04993	0.6875
1.05263	0.7031
1.05670	0.7188
1.07033	0.7344
1.07537	0.7500
1.08572	0.7656
1.11229	0.7813
1.12607	0.7969
1.12659	0.8125
1.13123	0.8281
1.13919	0.8438
1.22809	0.8594
1.23038	0.8750
1.39185	0.8906
1.48279	0.9063
1.50341	0.9219
1.59187	0.9375
1.60342	0.9531
1.61913	0.9688
3.19246	0.9844
3.19268	1.0000

Table 6.CDF of the Differences between the Measured and Predicted Th(IV) Solubilities for
the CRA-2014 PA (This Revised Analysis) (Cont.).

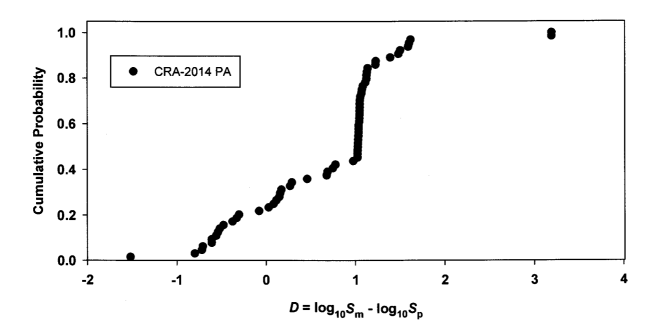


Figure 2. Plot of the CDF of the differences between the measured and predicted Th(IV) solubilities for the CRA-2014 PA (this revised analysis).

3.2 Am(III) Solubility Model

This subsection describes the comparisons by Brush and Domski (2013b) of the experimentally measured Am(III) solubilities with those predicted using the WIPP Am(III) speciation and solubility model in EQ3/6. It includes information similar to that provided for Th(IV) in Subsection 3.1 (see above). In particular, it applies criterion G11 to the measured solubilities included by Brush and Domski (2013b), presents our revised results, and provides a revised CDF for the differences between our measured and predicted Am(III) solubilities for the CRA-2014 PA,

The measured Am(III) solubilities from published papers and unpublished reports evaluated previously (Xiong et al., 2005; and/or Xiong et al., 2011) include: Borkowski et al. (2009), Borkowski (2010), Bunyakina and Storozhenko (2004), Khalili et al. (1994), Lucchini et al. (2007), Makino et al. (1993), Meinrath and Kim (1991), Meinrath and Takeishi (1993), Peretrukhin et al. (1996), Rao et al. (1996a, 1996b, 1999), Runde et al. (1992), Runde and Kim (1995), Silva (1982), and Wood (2002) (see Table 7 and the text below).

The measured Am(III) solubilities from published papers and unpublished reports that Brush and Domski (2013b) evaluated for their analysis include: Borkowski et al. (2010), Borkowski (2012), Migdisov and Williams-Jones (2007), Neck et al. (2009), Rabung et al. (2008) and Vitorge and Tran The (1991) (see Table 7 and text).

Reference	Solution(s) and Ionic Strength(s)	pcH, pmH, or pH	Solubility-Controlling Solid	Evaluation and EQ3/6 Run Numbers for Results Included in This Analysis
Borkowski et al. (2009)	 (A) 5 M NaCl (B) GWB, diluted to 95% of its specified concentrations (C) ERDA-6, diluted to 95% of its specified concentrations 	(A) pcH = 8.2–13.0 (B) pcH = 6.6–8.7 (C) pcH = 7.9–10.3	Presumably Nd(OH) ₃ , based on their thermodynamic predictions.	Superseded by Borkowski (2010) (see below).
Borkowski (2010)	 (A) 5 M NaCl (B) GWB, diluted to 95% of its specified concentrations of solutes to avoid coprecipitation of Nd by evaporite minerals (C) ERDA-6, diluted to 95% of its specified concentrations of solutes to avoid coprecipitation of Nd by evaporite minerals 	(A) pcH = 8.14–13.17 (B) pcH = 6.54–8.64 (C) pcH = 7.55–10.62	Presumably Nd(OH) ₃ , based on their thermodynamic predictions.	Superseded Borkowski et al. (2009). All 120 results from 5 M NaCl included by Xiong et al. (2011) and Brush and Domski (2013b). After application of criterion G11, 100 results (20 replicates) were included in this revised analysis. EQ3 files: 01-21Bor10.3i and 37-115Bor10.3i. Twenty-eight results with pcH \leq 7.4 from GWB included by Xiong et al. (2011) and Brush and Domski (2013b) because Borkowski concluded that they were not affected significantly by Nd(III)-borate complexation (see the description of Borkowski et al., 2009, in text). These results do not satisfy Criterion G11 and were not included in this revised analysis. Fourteen results with pcH \leq 8.1 included by Xiong et al. (2011) and Brush and Domski (2013b) because Borkowski concluded that these results were unaffected by borate. After application of criterion G11, 6 results (3 replicates) were included in this revisior 05 & 06BorERD.3i and 11-14BorERD.3i. A total of 106 results (23 replicates) included.

Table 7. Final Evaluations of Laboratory Studies for Comparisons of Measured and Predicted Nd(III) and Am(III) Solubilities.

Reference	Solution(s) and Ionic Strength(s)	pcH, pmH, or pH	Solubility-Controlling Solid	Evaluation and EQ3/6 Run Numbers for Results Included in This Analysis
Borkowski et al. (2010)	1, 2, 3, 4, or 5 M NaCl with 0.00. 0.01, 0.02, 0.03, or 0.04 M Na ₂ B ₄ O ₇ ; all runs from undersaturation	pcH = 8.59 ± 0.04 (constant, all runs)	Presumably Nd(OH) ₃ , which was used to start these runs	Not considered by Xiong et al. (2011) because this paper came to their attention after completion of their analysis. Results obtained with 1, 2, 3, 4, or 5 M NaCl & 0.00 M Na ₂ B ₄ O ₇ included by Brush and Domski (2013b), but they used numerical values from Borkowski (2012) (see below). Following the application of criterion G11, 3 results (no replicates) were included in this revised analysis: 03bor12.3i through 05bor12.3i. All other results excluded from this analysis because the NdHB ₄ O ₇ ²⁺ species found to be important by Borkowski et al. (2010) is not in the WIPP Am(III) model.
Borkowski (2012)	Same as above	Same as above	Same as above	Same as above
Bunyakina & Storozhenko (2004)	Solutions saturated with MgSO ₄ and/or Nd ₂ (SO ₄) ₃ . Ionic strengths not provided.	Not provided	Nd₂(SO₄)₃·8H₂O	Not considered by Xiong et al. (2005) because this paper came to their attention after completion of their analysis. Excluded by Xiong et al. (2011) and Brush and Domski (2013b) because $Nd_2(SO_4)3\cdot 8H_2O$ is not in the WIPP Am(III) model.

Reference	Solution(s) and Ionic Strength(s)	pcH, pmH, or pH	Solubility-Controlling Solid	Evaluation and EQ3/6 Run Numbers for Results Included in This Analysis
Khalili et al. (1994)	Synthetic brine similar to GWB	pcH = 6.4, 8.4, 10.4, & 12.4	Nd(OH) ₃ ·nH ₂ O(am) at pcH = $6.4 \& 8.4$, Nd ₂ (CO ₃) ₃ ·8H ₂ O or "uncertain" at pcH = 10.4 , NdO(OH) & Nd(OH) ₃ or "uncertain" at pcH = 12.4	Twenty-nine results for $pcH = 6.4 \& 8.4$ included by Xiong et al. (2005) because $Am(OH)_3$ is in the WIPP Am(III) model. Results for $pcH = 10.4 & 12.4$ excluded by Xiong et al. (2005) because $Nd_2(CO_3)_3 \cdot 8H_2O$ and NdO(OH) are not in the model, and because it is unclear that all of the solids were identified. Twenty-four of the 29 results included by Xiong et al. (2005) excluded by Xiong et al. (2011) and Brush and Domski (2013b) because the pcH in these runs exceeded 7.4, the pcH above which Borkowski concluded that complexation of Nd(III) by borate in GWB is significant (see text). Five results (5 duplicates) included by Brush and Domski (2013b). These results do not satisfy criterion G11 and were not included in this revised analysis.
Lucchini et al. (2007)	(A) GWB (B) ERDA-6	(A) pcH = 6.6-8.7 (B) pcH = 7.9-10.3	Nd(OH) ₃ (both solutions), based on their thermodynamic predictions.	Excluded by Xiong et al. (2011) and Brush and Domski (2013b) because Borkowski et al. (2009) and Borkowski (2010) superseded this paper.
Makino et al. (1993)	0.01 M NaClO ₄ & 0.1 ppm CO ₂	pcH = 7–13	Nd(OH) ₃ (cr) with lesser amounts of Nd(OH)CO ₃ (cr) (unclear whether both of these solids were in equilibrium one was replacing another).	Not considered by Xiong et al. (2005) because this paper came to their attention after completion of their analysis. Excluded from by Xiong et al. (2011) and Brush and Domski (2013b) because neither the TIC concentration of the aqueous phase nor the pCO_2 of the gaseous phase was provided, thus precluding predictions of Nd(III) solubilities for their experiments.

Reference	Solution(s) and Ionic Strength(s)	pcH, pmH, or pH	Solubility-Controlling Solid	Evaluation and EQ3/6 Run Numbers for Results Included in This Analysis
Meinrath & Kim (1991)	(A) 0.1 M NaClO ₄ & 0.00031 atm CO ₂	(A) pH = 4.1-4.3	(A) Nd ₂ (CO ₃) ₃ & Nd(OH)CO ₃ initially, Nd(OH)CO ₃ after 2 weeks	Not considered by Xiong et al. (2005) because this paper came to their attention after completion of their analyses. All nine results (no duplicates) from (A) included by Xiong et al. (2011) and Brush and Domski (2013b)
	(B) 0.1 M NaClO ₄ & 0.01 atm CO ₂	(B) $pH = 5.2-5.4$	(B) $Nd_2(CO_3)_3$	because $Nd(OH)CO_3$ is in the WIPP Am(III) model. These results do not satisfy Criterion G11 and were not
	(C) 0.1 M NaClO ₄ & 1 atm CO ₂	(C) pH = 5.7-6.2	(C) Nd ₂ (CO ₃) ₃	included in this revised analysis. Results from (B) and (C) excluded by Xiong et al. (2011 and Brush and Domski (2013b) because $Nd_2(CO_3)_3$ is not in the model.
Meinrath & Takeishi (1993)	(A) 0.1 M NaClO ₄ & 0.0003 atm CO ₂	(A) pH = 5.6-6.6	(A) Nd ₂ (CO ₃) ₃ & Nd(OH)CO ₃ initially, Nd(OH)CO ₃ (mcr) after 5-7 days	Not considered by Xiong et al. (2005) because this paper came to their attention after completion of their analysis. All 17 results (no duplicates) from (A) and (B) included by Xiong et al. (2011) and Brush and Domski (2013b)
	(B) 0.1 M NaClO ₄ & 0.003 atm CO ₂	(B) pH = 5.4-6.3	(B) Nd ₂ (CO ₃) ₃ & Nd(OH)CO ₃ initially, Nd(OH)CO ₃ (mcr) after 30-40 days	because Nd(OH)CO ₃ , is in the WIPP Am(III) model. These results do not satisfy Criterion G11 and were not included in this revision .Results from (C), (D), and (E) excluded because Nd ₂ (CO ₃) ₃ is not in the model.
	(C) 0.1 M NaClO ₄ & 0.01 atm CO ₂	(C) $pH = 5.2-6.2$	(C) Nd ₂ (CO ₃) ₃ ·(4.5- 6 H ₂ O)	excluded because $\operatorname{INd}_2(\operatorname{CO}_3)_3$ is not in the model.
	(D) 0.1 M NaClO ₄ & 0.08 atm CO ₂	(D) pH = 4.7-5.9	(D) Nd ₂ (CO ₃) ₃ ·(4.5- 6 H ₂ O)	
	(E) 0.1 M NaClO ₄ & 1 atm CO ₂	(E) pH = 4.2-5.2	(E) Nd ₂ (CO ₃) ₃ ·(4.5- 6 H ₂ O)	

Reference	Solution(s) and Ionic Strength(s)	pcH, pmH, or pH	Solubility-Controlling Solid	Evaluation and EQ3/6 Run Numbers for Results Included in This Analysis
Migdisov and Williams-Jones (2007)	HClO ₄ =NaHClO ₄ (total ClO ₄ ⁻ = 0.057- 1.97 M	pH = 1.04-1.65 (at 25 °C)	NdF3	Not considered by Xiong et al. (2005, 2011) because this paper came to their attention after completion of their analysis. All results excluded by Brush and Domski (2013b) because Migdisov and Williams-Jones (2007) carried out their experiments at 150, 200, and 250 °C, and because F is not in the WPP Am(IIII) model.
Neck et al. (2009)	 (A) 0.1, 0.5, 2.5, & 5.0 M NaCl; 0.1 M NaClO₄ (B) 0.25, 1.0, 2.5, & 3.5 M MgCl₂ (C) 0.25, 1.0, 2.5, & 3.5 M CaCl₂ 	 (A) pcH ≈ 7 to 13, 14, or even 15 (B) pcH ≈ 7 or 7.5 to 9 (C) pcH ≈ 7-9 to 12 	(A) Nd(OH) ₃ or Cm(OH) ₃ (5 M NaCl &1.0, 2.5, & 3.5 M CaCl ₂)	Not considered by Xiong et al. (2011) because this paper came to their attention after completion of their analysis. All 161 results from NaCl and MgCl ₂ solutions included by Brush and Domski (2013b). After application of criterion G11, 8 results from NaCl solutions (no replicates) were included in this revised analysis: 95NeckNa.3i through 102NeckNa.3i, and 01NeckMg.3i through 42NeckMg.3i. After application of criterion G11 16 (no replicates) results from MgCl ₂ solutions were included in this revision: 17-21NeckMg.3i, 27- 29NeckMg.3i, 35-42NeckMg.3i. All 46 results from CaCl ₂ with pcH < 10 included by Brush and Domski (2013b). Following the application of criterion G11, 32 (2 replicates) results from CaCl ₂ solutions were included in this revision: 15NeckCa.3i through 46NeckCa.3i. Results with pcH> 10 excluded because the complexes Ca(Cm(OH) ₃) ²⁺ , Ca ₂ (Cm(OH) ₄) ³⁺ , and Ca ₃ (Cm(OH) ₆) ³⁺ found to be important by Neck et al. (2009) are not in the WIPP Am(III) model.

Reference	Solution(s) and Ionic Strength(s)	pcH, pmH, or pH	Solubility-Controlling Solid	Evaluation and EQ3/6 Run Numbers for Results Included in This Analysis
Peretrukhin et al. (1996)	1 & 5 M NaOH (experiments with Am(III))	Not provided	Am(OH) ₃ aged by boiling for 3 hours (undersaturation runs) or Am(OH) ₃ , crystallinity unspecified (oversaturation runs)	Not considered by Xiong et al. (2005) because this paper came to their attention after completion of their analyses. Four results (two duplicates) modeled by Xiong et al. (2011) but excluded from their comparison because they predicted solubilities four to six orders of magnitude less than their measured values, possibly due to the presence of one or more $Am(OH)_x^{3-x}$ complexes with x > 4 in their experiments, but not in the WIPP Am(III) model. These results also excluded by Brush and Domsk (2013b).
Rabung et al. 2008	(A) 0.1–3.5 M CaCl ₂	pcH ≈ 11.7	Cm(OH) ₃ (am) (all)	Not considered by Xiong et al. (2011) because this paper
	(B) 1.0 M CaCl ₂	pcH = 10.8 - 11.9		came to their attention after completion of their analysis. All results excluded by Brush and Domski (2013b) because the $Ca(Cm(OH)_3)^{2+}$, $Ca_2(Cm(OH)_4)^{3+}$, and
	(C) 2.5 M CaCl ₂	pcH = 10.8 - 11.9		$Ca_3(Cm(OH)_6)^{3+}$ complexes found to be important by Rabung et al. (2008) are not in the WIPP Am(III) model.
	(D) 3.5 M CaCl ₂	pcH = 10.8 - 11.9		Rabung et al. (2000) are not in the wirt i run(in) model.
Rao et al. (1996a)	0.1 M NaCl	pH = 4.8-9.0 (filtered samples in their Fig. 1) & 5.1-9.0 (41-day samples in their Fig. 2.	Nd(OH)₃(cr)	Not considered by Xiong et al. (2005) because this paper came to their attention after completion of their analysis. Thirty-two of the 25 °C results (no duplicates) included b Xiong et al, (2011) and by Brush and Domski (2013b). These data do not satisfy criterion G11 and were not included in this revised analysis. Results obtained at pH 5.8 excluded by Xiong et al. (2011) and by Brush and Domski (2013b) because the solids dissolved before saturation was attained.

Reference	Solution(s) and Ionic Strength(s)	pcH, pmH, or pH	Solubility-Controlling Solid	Evaluation and EQ3/6 Run Numbers for Results Included in This Analysis
Rao et al. (1996b)	(A) 0.1-1.0 M NaHCO ₃ (B) 0.1-2.0 M Na ₂ CO ₃	Not provided	NaNd(CO ₃) ₂ ·6H ₂ O(cr) (all)	Not considered by Xiong et al. (2005) because this paper came to their attention after completion of their analyses. Sixty results included by Xiong et al. (2011) and Brush and Domski (2013), but only 53 of these EQ3/6 runs attained equilibrium. These data do not satisfy criterion G11 and were not included in this revision.
Rao et al. (1999)	(A) 0.1-1.1 m NaHCO ₃	(A) pH = 7.9-8.3	NaNd(CO ₃) ₂ ·6H ₂ O (all)	One-hundred-and-five results included in Xiong et al. (2005). Twenty-eight of the 105 results included by Xiong et al. (2005) evaluated by Xiong et al. (2011) and
	(B) 0.1-2 m Na ₂ CO ₃	(B) pH = 10.4-10.8		Xiong et al. (2005) excluded by Xiong et al. (2011) and Brush and Domski (2013b) because the pcH in these runs
	(C) 2 M NaCl & 0.1-0.5 m NaHCO ₃	(C) pH = 7.9-8.3		exceeded 7.4 or 8.1, the pcH above which Borkowski concluded that complexation of Nd(111) by borate in GWB or ERDA-6, respectively, is significant (see description of
	(D) 2 M NaCl & 0.1-2 m Na ₂ CO ₃	(D) pH = 10.4-10.8		Borkowski et al., 2009, in text). Seventy-seven results (25 replicates) included by Brush and Domski (2013b), but only 71 of the EQ3/6 runs attained equilibrium. After
	(E) 4 M NaCl & 0.1-2 m Na2CO3	(E) $pcH = 10.4-10.8$		application of criterion G11, one of these results was included in this revised analysis: 46Rao99.3i.
	(F) G Seep, a synthetic brine similar to GWB	(F) pcH = 5.6-7.8		
	(G) ERDA-6	(G) $pcH = 6.2-10.6$		

Reference	Solution(s) and lonic Strength(s)	pcH, pmH, or pH	Solubility-Controlling Solid	Evaluation and EQ3/6 Run Numbers for Results Included in This Analysis
Runde et al. (1992)	0.1 M NaClO ₄ & 0.0003, 0.01, & 1 atm CO ₂	pH = 5.5 - 6.3	Nd(OH)CO ₃ at 0.0003 atm CO ₂ , Nd ₂ (CO ₃) ₃ at 0.01 & 1 atm CO ₂	Not considered by Xiong et al. (2005), because this paper came to their attention after completion of their analysis. Excluded by Xiong et al. (2011) and Brush and Domski (2013b) because only K_{sp} 's reported, not the actual solubility data.
Runde & Kim (1995)	(A) 5.0 M NaCl (B) 5.0 m NaCl & 0.01 atm CO ₂	(A) pcH = 6.7–13.8 (B) pcH = 6.6-9.3	(A) $Am(OH)_3(cr)$ (B) $NaAm(CO_3)_3$ $\cdot xH_2O(cr)$	Seventy one results (18 duplicates) included by Xiong et al. (2005, 2011) and by Brush and Domski (2013b). Following the application of criterion G11, 6 results (no replicates) were included in this revision: 10Run95.3i – 15Run95.3i.
Silva (1982)	(A) 0.1 M NaClO ₄ (B) 0.1 M NaClO ₄	(A) pH = 5.7-9.5 (B) pH = 7.0-9.4	(A) Nd(OH) ₃ (cr) (B) Am(OH) ₃ (cr)	Included in Xiong et al. (2005). Excluded by Xiong et al. (2011) and Brush and Domski (2013b) because this paper was published prior to 1990.
Vitorge and Tran The (1991)	 (A) "Lime water" (composition unspecified) with 0-10 M KOH (B) "Lime water" (composition unspecified) (C) "Cement water" (composition unspecified) with 0-10 M KOH 	Unspecified for (A), (B), and (C)	Am(OH) ₃ ?	Not considered by Xiong et al. (2005, 2011) because this paper came to their attention after completion of their analyses. All results excluded Brush and Domski (2013b) because the compositions of the solutions were not specified.

Reference	Solution(s) and Ionic Strength(s)	pcH, pmH, or pH	Solubility-Controlling Solid	Evaluation and EQ3/6 Run Numbers for Results Included in This Analysis
Wood et al. (2002)	(A) 0.03 m NaTr ^A (30 ℃) (B) 0.03, 0.10, & 1.0 m NaTr ^A (50 ℃)	(A) pmH = 7.01-9.13 (B) pmH = 6.494– 9.548 (I = 0.03 m) pmH = 6.653-6.711 (I = 0.10 m) pmH = 6.681-6.809	Nd(OH)3(cr)	Not considered by Xiong et al. (2005) because this paper came to their attention after completion of their analyses. Most results excluded by Xiong et al. (2011) and Brush and Domski (2013b) because $T > 30$ °C. Fifteen results (no duplicates) obtained at 30 °C included by Xiong et al. (2011) and Brush and Domski (20113b). These results do not satisfy criterion G11 and were not
	(C) 0.03, 0.10, & 1.0 m NaTr ^A (100 °C)	(I = 1.0 m) (C) pmH = 5.362– 8.421 (I = 0.03 m) pmH = 5.739-5.744 (I = 0.10 m) pmH = 5.857-5.859		included in this revision.
	(D) 0.03, 0.010, & 1.0 m NaTr ^A (150 °C)	(I = 1.0 m) (D) pmH = 4.791- 5.890 (I = 0.03 m) pmH = 5.017-5.021 (I = 0.10 m) pmH = 5.139-5.141 (I = 1.0 m)		

Reference	Solution(s) and Ionic Strength(s)	pcH, pmH, or pH	Solubility-Controlling Solid	Evaluation and EQ3/6 Run Numbers for Results Included in This Analysis
Wood et al. (2002) (continued)	(E) 0.03, 0.10, & 1.0 m NaTr ^A (200 °C)	(E) $pmH = 4.419$ - 5.088 (I = 0.03 m) pmH = 4.472-4.473 (I = 0.10 m) pmH = 4.558-4.560	See Wood et al. (2002) on previous page.	See Wood et al. (2002) on previous page.
	(F) 0.03, 0.10, & 1.0 m NaTr ^A (250 ℃)	(I = 1.0 m) (F) pmH = 3.873- 4.873 (I = 0.03 m) pmH = 4.050-4.068 (I = 0.10 m) pmH = 4.180-4.182 (I = 1.0 m)		
	(G) 0.03 & 0.10 m (290 °C)	(G) pmH = $3.493-$ 6.898 (I = 0.03 m) pH = $3.406-5.221$ (I = 0.10 m)		

- A. Sodium trifluoromethanesulfonate, abbreviated as "sodium triflate" or "NaTr," is a noncomplexing salt used to set the ionic strength. In EQ3/6 calculations, the noncomplexing medium, sodium perchlorate, is used as the supporting solution.
- B. Sodium trifluoromethanesulfonate, abbreviated as "sodium triflate" or "NaTr," is a noncomplexing salt used to set the ionic strength.

Borkowski et al. (2009) measured the solubilities of various Nd(III) solids in 5 M NaCl and the synthetic WIPP brines GWB and ERDA-6, diluted to 95% of their specified concentrations to avoid coprecipitation of Nd(III) by evaporite minerals (see Table 7). (Small changes in conditions, especially temperature, during solubility experiments can cause precipitation of solids such as evaporite minerals and coprecipitation of actinide elements if the solutions are saturated with these solids under the conditions used for these tests.) Nd(III), an oxidation-state analog of Am(III), was used along with Am(III) and Cm(III) to parameterize the WIPP Am(III) speciation and solubility model. However, Borkowski et al. (2009) stated that, in many of their experiments in GWB or ERDA-6, complexation of Nd(III) by dissolved borate probably increased the solubility of Nd(III). Because there are no Nd(III)-borate species in the WIPP Am(III) solubility model, SNL personnel asked Borkowski to select those experiments that had values of pcH low enough to preclude significant complexation of Nd(III) by borate, thereby satisfying criterion G9. Borkowski selected 18 runs with GWB (pcH \leq 7.4) and 10 with ERDA-6 (pcH \leq 8.1) and e-mailed them the numerical values of the results. This e-mail is in the SNL/WIPP Records Center in the records package entitled "Calculation of Actinide Solubilities for the WIPP CRA-2009 PABC," ERMS 548686. Borkowski also sent them the results of 54 runs with 5 M NaCl, because these solutions contained no borate. This e-mail is in the same records package.

Borkowski (2010) issued a report containing tables of numerical values of the final values of pcH and Nd(III) solubilities presented graphically by Borkowski et al. (2009). From the standpoint of the analysis by Xiong et al. (2011) and Brush and Domski (2013b), Borkowski (2010) superseded Borkowski et al. (2009). However, comparisons of the numerical values in Borkowski (2010) and those e-mailed by Borkowski (see above) revealed that: (1) Borkowski (2010) contained more results than those e-mailed by Borkowski, and (2) the numerical values e-mailed by Borkowski could not be correlated with any of those in Borkowski (2010). Therefore, Xiong et al. (2011) and Brush and Domski (2013b) used the numerical values from Borkowski (2010). However, they used the compositional and pcH criteria e-mailed by Borkowski (see the description of Borkowski et al., 2009, in the text above) to select these numerical values. Thus, Xiong et al. (2011) and Brush and Domski (2013b) used the following 162 results reported by Borkowski (2010): all 120 results from the 5 M NaCl solutions, 28 results from GWB with pcH \leq 7.4, and 14 from ERDA-6 with pcH \leq 8.1. After application of criterion G11, 100 results from the 5 M NaCl solutions and 6 from ERDA-6 were included in this revised analysis.

Borkowski et al. (2010) carried out solubility experiments with Nd(OH)₃ in 1, 2, 3, 4, or 5 M NaCl with 0.00, 0.01, 0.02, 0.03, or 0.04 M Na₂B₄O₇ at a constant pcH of 8.59 \pm 0.04. Xiong et al. (2011) did not consider the results of Borkowski et al. (2010) in their analysis because this paper came to their attention after they completed their analysis. Because all of these experiments were undersaturation runs and because the solutions were carbonate-free, the solubility-controlling solid in these experiments was presumably Nd(OH)₃. Brush and Domski (2013b) included the results obtained from solutions with 1, 2, 3, 4, or 5 M NaCl and 0.00 M Na₂B₄O₇. However, they excluded all of the other results, because the NdHB₄O₇²⁺ complex found to be important by Borkowski et al. (2010) is not in the WIPP Am(III) model. After the release of Borkowski et al. (2010), Borkowski et al. (2010). Brush and Domski (2013b) included the results obtained form solutions in the WIPP Am(III) model.

(2013b) used these numerical values from Borkowski (2012) for their analysis. Following the application of criterion G11, three results (no replicates) were included in this revised analysis.

Bunyakina & Storozhenko (2004) conducted solubility experiments with $Nd_2(SO_4)_3 \cdot 8H_2O$ in solutions saturated with $MgSO_4$ and/or $Nd_2(SO_4)_3$ (see Table 7). Xiong et al. (2005) did not consider the results of Bunyakina & Storozhenko (2004) because this paper came to their attention after they completed their analysis. Xiong et al. (2011) excluded these results from their analysis according to criterion G7 (see Section 2) because $Nd_2(SO_4)_3 \cdot 8H_2O$ is not in the WIPP Am(III) model. Domski and Brush (2013b) excluded these results for the same reason.

Khalili et al. (1994) determined the solubilities of various Nd(III) solids in a synthetic brine similar to GWB (Table 7). According to criterion G7 (Section 2), Xiong et al. (2005) included 29 measured solubilities from solutions with pcH = 6.4 and 8.4 in their analysis because Am(OH)₃, the solubility-controlling solid in these experiments, is in the WIPP Am(III) model. However, they used the same criterion to exclude the results obtained from solutions with pcH = 10.4 and 12.4 because $Nd_2(CO_3)_3 \cdot 8H_2O$ and NdO(OH), the solubility-controlling solids in these runs, are not in the model, and because it is unclear whether all of the solids were identified at these values of pcH. After considering the results of Borkowski et al. (2009) (see text and Table 7 above), Xiong et al. (2011) used criterion G9 to exclude 24 of the 29 results included by Xiong et al. (2005), because the pcH of the solutions in these experiments exceeded 7.4, the pcH above which Borkowski concluded that complexation of Nd(III) by dissolved borate in GWB is significant. Therefore, Xiong et al. (2011) included five of the results of Khalili et al. (1994) in their analysis. Brush and Domski (2013b) included the same five results in their analysis. These data do not satisfy criterion G11 and were not included in this revised analysis.

Lucchini et al. (2007) included preliminary results from Borkowski's study of the solubility of Nd(III) (see above and Table 7). Borkowski et al. (2009) and Borkowski (2010) superseded Lucchini et al. (2007). Therefore, Xiong et al. (2011) evaluated the results of Borkowski (2010) and included many of them in their analysis. Brush and Domski (2013b) included the same results included by Xiong et al. (2011) in this analysis. See the text and the entries in Table 7 for Borkowski et al. (2009) and Borkowski (2010) for a detailed explanation of why these results were included or excluded.

Makino et al. (1993) measured the solubilities of Nd(III) solids in 0.01 M NaClO₄ (Table 7). Xiong et al. (2005) did not consider this paper because it came to their attention after they completed their analysis. Makino et al. (1993) stated that "the solid phase was identified as Nd(OH)₃(c) by X-ray diffraction [XRD] before and after the experiments." However, both of the posttest XRD patterns in their Figure 1 show that Nd(OH)CO₃(cr) was also present. Clearly, sufficient CO_3^{2} was present in their solutions to convert at least some of the Nd(OH)₃(cr) to Nd(OH)CO₃(cr) during their undersaturation and/or oversaturation experiments, but it is unclear whether both of these solids were in equilibrium with the solutions or whether one solid was replacing another. It is also unclear what the TIC concentration of their solutions was, or even what the p_{CO₂} of the atmosphere in the headspace above their solutions was, because

the p_{CO_2} of 0.1 ppm provided in the English summary of their paper was a detection limit (upper limit) for p_{CO_2} , not the actual concentration of CO₂ in the gaseous phase. Therefore, Xiong et al. (2011) excluded all 22 measured solubilities of Makino et al. (1993) according to criterion G8. Brush and Domski (2013b) also excluded these results from this analysis.

Meinrath and Kim (1991) measured the solubilities of $Nd_2(CO_3)_3$ and $Nd(OH)CO_3$ in 1 M NaClO₄ under atmospheres with various concentrations of CO₂ (Table 7). Xiong et al. (2005) did not consider the results of Meinrath and Kim (1991) because this paper did not come to their attention until after they completed their analysis. Xiong et al. (2011) included all nine results obtained with 0.1 M NaClO₄ under a gaseous phase that contained 0.00031 atm CO₂ according to criterion G7 (Section 2), because Nd(OH)CO₃, — the solubility-controlling solid in these experiments — is in the WIPP Am(III) model. However, they used the same criterion to exclude the results obtained from 0.1 M NaClO₄ under 0.01 and 1 atm CO₂, because Nd₂(CO₃)₃ — the solubility-controlling solid in these runs — is not in the model. Brush and Domski (2013b) included and excluded the same results from this analysis. These results do not satisfy Criterion G11 and were not included in this revised analysis.

Meinrath and Takeishi (1993) also determined the solubilities of $Nd_2(CO_3)_3$ and $Nd(OH)CO_3$ in 1 M NaClO₄ under atmospheres with various concentrations of CO_2 (Table 7). Xiong et al. (2005) did not consider this paper because it came to their attention after they completed their analysis. Xiong et al. (2011) included all 17 results obtained with 0.1 M NaClO₄ under 0.0003 and 0.003 atm CO₂ according to criterion G7 (Section 2), because Nd(OH)CO₃ — the solubility-controlling solid in these experiments (Table 7) — is in the WIPP Am(III) model. However, they used the same criterion to exclude the results obtained from 0.1 M NaClO₄ under 0.01, 0.08, and 1 atm CO₂ because Nd₂(CO₃)₃ — the solubility-controlling solid in these runs — is not in the model. Brush and Domski (2013b) included and excluded the same results from their analysis. These data do not satisfy Criterion G11 and were not included in this revision.

Migdisov and Williams-Jones (2007) performed a solubility study of NdF₃ in HHClO₄-NaHClO₄ solutions (total ClO₄⁻ = 0.057-1.97 M) at 150, 200, and 250 °C. Xiong et al. (2005, 2011) did not consider these results because they found this paper came to their attention after they completed their analyses. Brush and Domski (2013b) used criterion G5 to exclude their results, because they carried out their experiments at 150, 200, and 250 °C. However, Criteria G7 and G9 also apply, because NdF₃ — the solubility-controlling solid in their experiments — and NdF²⁺ — one of the important complexes in their runs — are not in the WIPP Am(III) model.

Neck et al. (2009) measured the solubilities of Nd(OH)₃ and Cm(OH)₃ in NaCl, MgCl₂, and CaCl₂ solutions of various ionic strengths (Table 7). Xiong et al. (2011) did not consider the results of Neck et al. (2009) because this paper came to their attention after they completed their analyses. Brush and Domski (2013b) included all of their results from experiments with NaCl solutions, because these results are consistent with the dominance of Nd(OH)_x^{3-x} or Cm(OH)_x^{3-x} complexes with $0 \le x \le 3$. Furthermore, their data from solutions with very high values of pcH (13 ≤ pcH ≤ 15) showed no evidence for Nd(OH)_x^{3-x} or Cm(OH)_x^{3-x} complexes with x > 3. (The WIPP Am(III) model contains Am(OH)_x^{3-x} complexes with $0 \le x \le 3$, but not

x > 3.) Brush and Domski (2013b) also included all of their results from experiments with MgCl₂ solutions, because the pcH in these runs (\leq 9) was not high enough to stabilize possible Nd(OH)_x^{3-x} or Cm(OH)_x^{3-x} complexes with x > 3. Finally, Brush and Domski (2013b) included all of their results obtained with CaCl₂ solutions with pcH < 10, but excluded those from CaCl₂ solutions with pcH > 10. The reason why they excluded the latter results is that Neck et al. (2009) provided solubility data (negative slopes of solubility versus pcH) and time-resolved laser fluorescence spectroscopic (TRLFS) results that imply that Ca[Cm(OH)₃]²⁺, Ca₂[Cm(OH)₄]³⁺, and Ca₃[Cm(OH)₆]³⁺ and — by application of the oxidation-state analogy — Ca[Nd(OH)₃]²⁺, Ca₂[Nd(OH)₄]³⁺, and Ca₃[Nd(OH)₆]³⁺ are the dominant species in CaCl₂ solutions with pcH > 10. (The WIPP Am(III) model does not contain any of the complexes Ca[Am(OH)₃]²⁺, Ca₂[Am(OH)₄]³⁺, or Ca₃[Am(OH)₆]³⁺.) After application of criterion G11, 8 results from NaCl solutions (no replicates, 16 results from MgCl₂ solutions (no replicates) were included in this revised analysis.

Nevertheless, the WIPP Am(III) model is still adequate for WIPP compliance-related calculations. This is because Figures 4a through 4d of Neck et al. (2009) show that the concentrations of these $Ca_x[Am(OH)_y]^{2x+3-y}$ complexes at a pcH of 9.54 or 9.69, the values predicted by Brush and Domski (2013a) for the minimum volume of GWB or ERDA-6, respectively, for the CRA-2014 PA, do not exceed ~10^{-9.5} M until the CaCl₂ concentration exceeds ~0.25 M. This concentration of ~10^{-9.5} M is several orders of magnitude less than the predicted concentrations of AmEDTA⁻ and Am(OH)₂⁺ (the most important Am(III) complexes in the WIPP), which are ~10⁻⁶ or 10⁻⁷ m. (Note that the numerical values of these concentrations predicted by Brush and Domski (2013a) for the minimum volumes of GWB and ERDA-6 are 0.0111 and 0.0116 M. Therefore, the Ca concentrations of WIPP brines are far too low to stabilize the Ca_x[Am(OH)_y]^{2x + 3- y} complexes characterized by Neck et al. (2009), even after reacting with the solids in WIPP disposal rooms.

Peretrukhin et al. (1996) studied the solubilities of Tc(IV), Tc(V), Np(IV), Np(VI), Pu(III), Pu(IV), Am(III), and Am(VI) in strongly basic solutions. They carried out four experiments with Am(III), two in 1 M NaOH and two in 5 M NaOH (Table 7). Xiong et al. (2005) did not consider this paper because it came to their attention after they completed their analysis. Xiong et al. (2011) predicted Am(III) solubilities about four to six orders of magnitude less than their measured values (see their Table 9). They concluded that these differences were caused by one or more $Am(OH)_x^{3-x}$ complexes with x > 3 that were dominant in the experiments of Peretrukhin et al. (1996), but are not in the WIPP Am(III) model. Therefore, they used criterion G9 (Section 2) to exclude these measured Am(III) solubilities. Brush and Domski (2013b) also excluded these results. Note, however, that other results from solutions with $13 \le pcH \le 15$ showed no evidence for Nd(OH)_x^{3-x} or Cm(OH)_x^{3-x} complexes with x > 3 (see Neck et al, 2009).

Rabung et al. (2008) measured the solubility of Cm in 0.1, 1.0, 2.5, and 3.5 M CaCl₂ solutions under basic conditions (Table 7). Xiong et al. (2011) did not consider these results because they found this paper later. Brush and Domski (2013b) suspected that Rabung et al. (2008) was superseded by Neck et al. (2009). Brush and Domski's (2013b) evaluation of the results of Neck et al. (2009) led them to conclude that all of their results obtained with

CaCl₂ solutions with pcH < 10 are consistent with the dominance of Cm(OH)_x^{3-x} complexes with $0 \le x \le 3$, which are included in the WIPP Am(III) model. However, their results obtained with CaCl₂ solutions with pcH > 10 imply that Ca_x[Am(OH)_y]^{2x + 3- y} complexes dominate. Domski and Brush (2013b) excluded the results of Neck et al. (2009) from CaCl₂ solutions with pcH > 10 and they excluded all of the results of Rabung et al. (2008) with pcH > 10, because the WIPP Am(III) model does not contain any of these Ca_x[Am(OH)_y]^{2x + 3- y} complexes.

Rao et al. (1996a) carried out solubility experiments with Nd(OH)₃(cr) in 0.1 M NaCl at 25 and 90 °C. Xiong et al. (2005) did not consider the results of Rao et al. (1996a) because this paper came to their attention after they completed their analysis. Initially, Xiong et al. (2011) decided to include the 25 °C results of Rao et al. (1996a) and exclude their 90 °C results because the latter temperature is too high to be relevant to the WIPP. However, inspection of the 25 °C results showed that those with pH values < 5.8 were essentially independent of pH. This feature is similar to that observed by Xiong et al. (2005) in the results of Felmy et al. (1991) (see the discussion of these results in Subsection 3.1 above). Therefore, according to criterion G7 (Section 2), Xiong et al. (2011) included only the 32 solubilities measured by Rao et al. (1996a) at pH values > 5.8. Brush and Domski (2013b) included the same results in this analysis. These data do not satisfy criterion G11 and were not included in this revised analysis.

Rao et al. (1996b) conducted solubility experiments with NaNd(CO₃)₂·6H₂O(cr) in NaHCO₃ and Na₂CO₃ solutions of various ionic strengths (Table 7). Xiong et al. (2005) did not consider this paper because it came to their attention after they completed their analysis. Xiong et al. (2011) included all 60 results in their analysis. Brush and Domski (2013) also included these results, but only 53 of them attained equilibrium in their EQ3/6 runs. These results do not satisfy criterion G11 and were not included in this revision.

Rao et al. (1999) studied the solubilities of NaNd(CO₃)₂·6H₂O in NaHCO₃, Na₂CO₃, NaCl-NaHCO₃, and NaCl-Na₂CO₃ solutions of various ionic strengths; and in synthetic G Seep (a WIPP brine similar to GWB) and ERDA-6 (Table 7). Xiong et al. (2005) included all 104 results of this study in their analyses. Xiong et al. (2011), however, excluded 28 of the 104 results included by Xiong et al. (2005) according to criterion G9 because the pcH of the solutions in these experiments exceeded 7.4 or 8.1, the pcH above which Borkowski concluded that complexation of Nd(III) by dissolved borate in GWB or ERDA-6, respectively, is significant. Therefore, Xiong et al. (2011) included 77 of the results of Rao et al. (1999) in their analysis. Brush and Domski (2013b) included these results too, but only 71 attained equilibrium in their EQ3/6 runs. After application of criterion G11, one of these results was included in this revised analysis.

Runde et al. (1992) measured the solubilities of Nd(OH)CO₃ and Nd₂(CO₃)₃ under atmospheres with various concentrations of CO₂ (Table 7). Xiong et al. (2005) did not consider the results of Runde et al. (1992) because this paper came to their attention after they completed their analyses. Runde et al. (1992) reported values of $K_{sp}s$ instead of solubilities expressed as concentrations of dissolved Nd(III). Therefore, according to criterion G10 (Section 2), Xiong et al. (2011) excluded their results. Brush and Domski (2013b) excluded them for the same reason.

Runde and Kim (1995) determined the solubility of $Am(OH)_3(cr)$ in 5.0 M NaCl and that of $NaAm(CO_3)_3 \cdot xH_2O(cr)$ in 5.0 m NaCl under an atmosphere with 0.01 atm CO_2 (Table 7). Xiong et al. (2005, 2011) included all 71 results of this work in their analyses. Brush and Domski (2013b) also included all 71 results in their analysis. Following the application of criterion G11, six results (no replicates) were included in this revision.

Silva (1982) performed solubility experiments with $Nd(OH)_3$ and $Am(OH)_3$ in 0.1 M NaClO₄ (Table 7). Xiong et al. (2005) included all 37 results of this study in their analyses. Xiong et al. (2011), however, included only measured solubilities from papers published or unpublished reports issued from January 1, 1990, through December 31, 2008 (criterion G1 in Section 2). Therefore, they excluded the results of Silva (1982) from their analysis. Brush and Domski (2013b) excluded them too.

Vitorge and Tran The (1991) measured the solubilities of $Am(OH)_3$ and Am(V)-bearing solids in solutions that they referred to as "lime water" and "cement water." They did not provide enough information on the compositions of their solutions to predict the solubilities of Am(III) in the absence of assumptions as to what solid(s) were present when samples were taken for analysis of Am concentrations, what the TIC concentration was, etc. Xiong et al. (2005, 2011) did not consider this paper because it came to their attention after they completed their analyses. Brush and Domski (2013b) used criterion G8 (Section 2) to exclude all of their results.

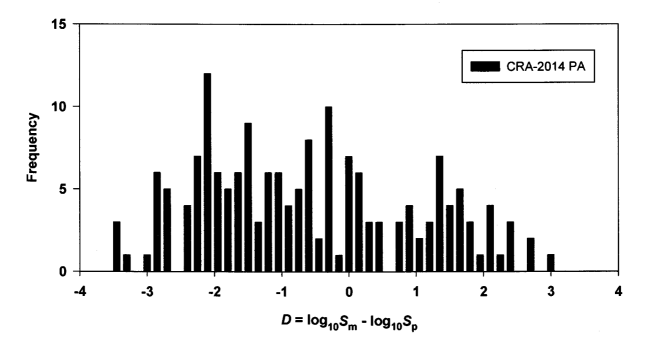
Wood et al. (2002) studied the solubility of Nd(OH)₃(cr) in 0.03, 0.10, and 1.0 m sodium trifluoromethanesulfonate (NaCF₃SO₃) solutions at 30, 50, 100, 150, 200, 250, and 290 °C. (Sodium trifluoromethanesulfonate is often abbreviated as "sodium triflate" or "NaTr.") Wood et al. (2002) used NaTr to set the ionic strength in their experiments because it dissolves to form species that do not complex Nd(III) in aqueous solutions. Xiong et al. (2005) did not consider these results because they found this paper later. Xiong et al. (2011) included the 15 results obtained at 30 °C in this revision, but excluded all results obtained at higher temperatures because they were above the acceptable temperature range of 20–30 °C specified in criterion G5 (Section 2). Brush and Domski (2013b) included and excluded the same results. These results do not satisfy criterion G11 and were not included in this revision.

Figure 3 provides the histogram for our comparison of experimentally measured and predicted Nd(III) and Am(III) solubilities for the CRA-2014 PA.. The current comparison includes a total of 172 measured and predicted solubilities. The current distribution peaks at Bin -2.10 with a frequency of 12, comparisons, with a secondary peak at Bin -0.30 with 10 comparisons. Table 8 summarizes the statistical parameters of the current frequency distribution.

Figure 3 shows that the WIPP Am(III) model, the EQ3/6 code, and the DB data0.FM1 overpredicted 117 of the 172 measured Nd(III) and Am(III) solubilities in this analysis, and underpredicted 55. The mean and median values of the log of the sampled uncertainty factor used to adjust the baseline Am(III) solubilities are -0.678 and -0.866, respectively (Table 8), and the mean and median values of the sampled uncertainty factor are $10^{-0.678} = 0.210$ and $10^{-0.866} = 0.136$. WIPP PA uses actinide solubilities that are the products of the baseline solubilities and the sampled uncertainty factors.



Table 9 provides values of the CDF for Nd(III) and Am(III) for the CRA-2014. PA will use this CDF to define the solubility multiplier SOLVAR for the oxidation-state III model, material SOLMOD3 in the WIPP PA Parameter Database for the CRA-2014 PA. Brush et al. (2005) described the PA implementation of these results in detail.



- Figure 3. Histogram of the difference $D = \log_{10} S_m \log_{10} S_p$ for Nd(III) and Am(III) for the CRA-2014 PA (this revised analysis).
- Table 8. Statistical Properties of the Distribution of D = log10(Sm) log10(Sp) for Nd(III) and Am(III) for the CRA-2014 PA (This Revised Analysis).

Analysis	Mean	Median	Standard Deviation	Minimum	Maximum
CRA-2014 PA	-0.678	-0.866	1.57	-3.55	2.97

Table 9. CDF of the Differences between the Measured and Predicted Nd(III) and Am(III) Solubilities for the CRA-2014 PA (This Revised Analysis). This CDF Defines the Solubility Multiplier SOLVAR for the Oxidation-State III Model Material SOLMOD3 in the WIPP PA Parameter Database.

D	<i>P(D<x)< i="">CRA-2014 PA</x)<></i>	
-3.54668	0.005814	
-3.53804	0.01163	
-3.48624	0.01744	
-3.39484	0.02326	
-3.12625	0.02907	
-2.99787	0.03488	
-2.92819	0.04070	
-2.90889	0.04651	
-2.89789	0.05233	
-2.89417	0.05814	
-2.88167	0.06395	
-2.81960	0.06977	
-2.80340	0.07558	
-2.77909	0.08140	
-2.75392	0.08721	
-2.71310	0.09302	
-2.50363	0.09884	
-2.44562	0.1047	
-2.42524	0.1105	
-2.40328	0.1163	
-2.38528	0.1221	
-2.38125	0.1279	
-2.36584	0.1337	
-2.34077	0.1395	
-2.30728	0.1453	
-2.30370	0.1512	
-2.28221	0.1570	
-2.22527	0.1628	
-2.21692	0.1686	
-2.21193	0.1744	
-2.21057	0.1802	

Table 9 continued on next page

D	<i>P(D<x)< i="">CRA-2014 PA</x)<></i>
-2.21035	0.1860
-2.19890	0.1919
-2.17941	0.1977
-2.14466	0.2035
-2.14017	0.2093
-2.11004	0.2151
-2.10131	0.2209
-2.10044	0.2267
-2.08652	0.2326
-2.06574	0.2384
-2.02624	0.2442
-2.02078	0.2500
-1.99435	0.2558
-1.99094	0.2616
-1.89183	0.2674
-1.88602	0.2733
-1.88082	0.2791
-1.87526	0.2849
-1.81645	0.2907
-1.79857	0.2965
-1.79072	0.3023
-1.77062	0.3081
-1.73992	0.3140
-1.71789	0.3198
-1.65342	0.3256
-1.64679	0.3314
-1.60274	0.3372
-1.58665	0.3430
-1.57875	0.3488
-1.57686	0.3547
-1.55805	0.3605
-1.54605	0.3663
-1.53335	0.3721
-1.50332	0.3779

Table 9 continued on next page

D	<i>P(D<x)< i="">CRA-2014 PA</x)<></i>
-1.47121	0.3837
-1.43567	0.3895
-1.35594	0.3953
-1.32260	0.4012
-1.29978	0.4070
-1.28158	0.4128
-1.27820	0.4186
-1.23176	0.4244
-1.20372	0.4302
-1.17824	0.4360
-1.17469	0.4419
-1.13616	0.4477
-1.08408	0.4535
-1.07590	0.4593
-1.06288	0.4651
-1.01740	0.4709
-0.96962	0.4767
-0.94413	0.4826
-0.90486	0.4884
-0.87890	0.4942
-0.87434	0.5000
-0.85831	0.5058
-0.84284	0.5116
-0.75515	0.5174
-0.74714	0.5233
-0.74650	0.5291
-0.71044	0.5349
-0.69913	0.5407
-0.69468	0.5465
-0.66058	0.5523
-0.63582	0.5581
-0.61706	0.5640
-0.59335	0.5698
-0.48697	0.5756

Table 9 continued on next page

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D	<i>P(D<x)< i="">CRA-2014 PA</x)<></i>
-0.42447	0.5814
-0.41645	0.5872
-0.41048	0.5930
-0.39289	0.5988
-0.37683	0.6047
-0.36274	0.6105
-0.35914	0.6163
-0.33371	0.6221
-0.32341	0.6279
-0.30807	0.6337
-0.16053	0.6395
-0.12886	0.6453
-0.10835	0.6512
-0.07919	0.6570
-0.06936	0.6628
-0.06422	0.6686
-0.05862	0.6744
-0.02520	0.6802
0.02195	0.6860
0.04981	0.6919
0.07414	0.6977
0.07414	0.7035
0.08615	0.7093
0.10609	0.7151
0.19880	0.7209
0.25433	0.7267
0.26417	0.7326
0.31303	0.7384
0.34636	0.7442
0.36008	0.7500
0.68528	0.7558
0.70158	0.7616
0.74939	0.7674
0.77821	0.7733

Table 9 continued on next page

D	<i>P(D<x)< i="">CRA-2014 PA</x)<></i>
0.80209	0.7791
0.85027	0.7849
0.88995	0.7907
0.95297	0.7965
1.02872	0.8023
1.06857	0.8081
1.09543	0.8140
1.17080	0.8198
1.21546	0.8256
1.21768	0.8314
1.23265	0.8372
1.23608	0.8430
1.23617	0.8488
1.28595	0.8547
1.29523	0.8605
1.41886	0.8663
1.41967	0.8721
1.45195	0.8779
1.47603	0.8837
1.51897	0.8895
1.53738	0.8953
1.55726	0.9012
1.57030	0.9070
1.64380	0.9128
1.67223	0.9186
1.70369	0.9244
1.79257	0.9302
1.91679	0.9360
1.96394	0.9419
2.02303	0.9477
2.05965	0.9535
2.06862	0.9593
2.17544	0.9651
2.25179	0.9709

Table 9 continued on next page

Table 9.CDF of the Differences between the Measured and Predicted Nd(III) and Am(III)Solubilities for the CRA-2014 PA (This Revised Analysis) (Cont.).

D	<i>P(D<x)< i="">CRA-2014 PA</x)<></i>	
2.34368	0.9767	
2.37300	0.9826	
2.62793	0.9884	
2.68349	0.9942	
2.97147	1.0000	

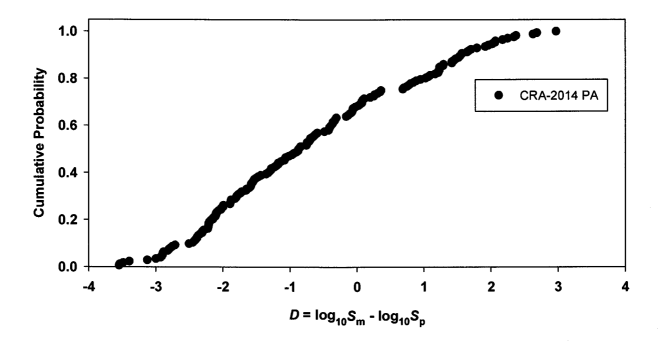


Figure 4. Plot of the CDF of the differences between the measured and predicted Nd(III) and Am(III) solubilities for the CRA-2014 PA (this revised analysis).

4 CONCLUSIONS

This revised analysis report provides the results of the new uncertainty analysis of the solubilities of Th(IV) and Am(III). SNL PA personnel will use the results of this uncertainty analysis in the CRA-2014 PA.

We used EQ3/6, Version 8.0a, and the thermodynamic database DB DATA0.FM1 for this analysis. We carried out this analysis under Task 4 of AP-153, Rev. 1, the current analysis plan (AP) for WIPP near-field geochemical process modeling.

Our Th(IV) comparison included a total of 64 measured and predicted solubilities. The WIPP Th(IV) model overpredicted the majority of the measured solubilities in the current analysis. The mean and median values of the log of the sampled uncertainty factor used to adjust the baseline Th(IV) solubilities are 0.673 and 1.029, respectively. The mean and median values of the sampled uncertainty factor are $10^{0.673} = 4.708$ and $10^{1.029} = 10.703$. (The PA codes use actinide solubilities that are the products of the baseline solubilities and the sampled uncertainty factors).

Our Nd(III) and Am(III) comparison included a total of 172 measured and predicted solubilities. The WIPP Am(III) model underpredicted the majority of the measured solubilities. The mean and median values of the log of the sampled uncertainty factor used to adjust the baseline Am(III) solubilities are -0.678 and -0.866, respectively. The mean and median values of the sampled uncertainty factor are $10^{-0.678} = 0.210$ and $10^{-0.866} = 0.136$.

Tables 6 and 9 provide the CDFs of the uncertainties for the predicted Th(IV) and Am(III) solubilities for the CRA-2014 PA.

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