ION EXCHANGE IN THE NUCLEAR INDUSTRY (U)

by

Jane P. Bibler

Westinghouse Savannah River Company
Savannah River Site
Aiken, SC, USA 29808

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ION EXCHANGE IN THE NUCLEAR INDUSTRY

JANE P. BIBLER
Westinghouse Savannah River Company
Aiken, South Carolina 29808 USA

ABSTRACT

Ion exchange is used in nearly every part of the nuclear fuel cycle - from the purification of uranium from its ore to the final recovery of uranium and transmutation products. Ion exchange also plays a valuable role in the management of nuclear wastes generated in the fuel cycle.

INTRODUCTION

The nuclear industry encompasses a variety of operations and ion exchange processes play an important role in nearly every operation. Uranium, recovered from its ore, must be purified and concentrated. Enriched uranium is fabricated into fuel elements and placed in nuclear reactors where it produces energy, fission products, and transmutation products. If the fuel cycle is completed, uranium and useable transmutation products are separated from the fission products and from each other, to be recycled or used elsewhere. Finally, new waste disposal requirements have prompted studies for the use of ion exchange in remediating wastes generated in each of the above operations.

URANIUM PURIFICATION AND CONCENTRATION

Triggered by the U.S. nuclear defense efforts in the late 1940's and early 1950's, extensive pioneering research and development efforts [1-8] laid down the relevant process chemistry of anion exchange purification of uranium. The first actual plant-scale recovery, concentration, and purification of uranium from H2SO4 ore leach liquors by anion exchange was demonstrated in South Africa in 1952. [9] This demonstration was quickly followed in many countries by construction and operation of similar ion exchange processes for clarified and unclarified (resin-in-pulp) ore leach liquors, both acidic and alkaline carbonate. Subsequent investigations of improvements to this earliest
technology have been concerned with the combination of ion exchange with solvent extraction and with development and testing of new resins and new equipment for countercurrent applications. Recently, ion exchange technology has been directed at the recovery of uranium from mine waters [10] and mill waste streams.[11]

**Important Ion Exchange Resins: Types and Properties**

Strong base Type I resins were the first ion exchange resins used for uranium recovery [6] and still remain as resins of choice in the industry.[12] Typically, the exchange site on a Type I resin is a symmetrical quaternary ammonium ion. Strong base Type II resins use an exchange site similar to Type I resins except that the quaternary ammonium ion is no longer symmetrical, one methyl group having been substituted by an ethanolic group. The advantage in sorption offered by Type II resins is due to their lower affinity for Cl" or NO₃" compared to Type I resins.[12] Vinyl pyridine strong base resins are effective in removing uranium species from solutions but allow much easier stripping of uranium by Cl" or NO₃" than either Type I or Type II resins. Weak base resins used for uranium recovery have tertiary amine functional groups. New waste disposal regulations have precipitated studies of the use of ion exchange for removal of very small concentrations (2 to 10 ppm) of uranium from mine waters and waste streams. The resins that have been used in these studies are conventional strong base anion exchangers, weak acid (carboxylic) cation exchangers, and those containing chelating functional groups (e.g. iminodiacetate, amidoxime, picotylamine, aminophosphonic).

The earliest ion exchange uranium recovery operations used gel-type 16-50 mesh anion resin beads in conventional fixed-bed columns. Modern continuous or countercurrent ion exchange (CIX) processes employ macroporous beads of 12-20 mesh because of their greater resistance to attrition.

**Ion Exchange Chemistry**

Dilute H₂SO₄ is generally used to leach uranium from ground ores. Acidic sulfate leach liquors contain UO₂²⁺, UO₂SO₄, [UO₂(SO₄)₂]²⁻, and [UO₂(SO₄)₃]⁴⁻ in equilibrium.

\[
\text{UO}_2^{2+} + n\text{SO}_4^{2-} \leftrightarrow \text{UO}_2(\text{SO}_4)_n^{2-2n} \quad (1)
\]

\(n = 1, 2, \text{ or } 3\)

Ion exchange of uranium from these solutions is represented on either strong or weak base anion resins by the following general reaction:

\[
4 \text{R-X} + \text{UO}_2(\text{SO}_4)_3^{4-} \leftrightarrow (\text{R})_4(\text{SO}_4)_3 + 4 \text{X}^- \quad (2)
\]

where \(X = \text{Cl} \text{ or NO}_3\) and \(R = \text{the resin support network}\).
For mined or ground ores that are high-lime ores (~15% CaCO₃), leaching with H₂SO₄ would consume very large quantities of the acid and carbonate solutions are, instead, used to leach uranium. (Some in situ leach operations have used ammonium carbonate/bicarbonate mixtures.)[13] Carbonate leach liquors contain the UO₂(CO₃)₃⁴⁻ ion as the predominating species. This ion is sorbed on the weak and strong base ion exchange resins according to equation (3). The symbols R and X have the same meaning as in equation (2).

\[
4 \text{R-X} + \text{UO}_2(\text{CO}_3)_3^{4-} \leftrightarrow (\text{R})_4\text{UO}_2(\text{CO}_3)_3 + 4 \text{X}^-
\]  

(3)

Process variables such as pH, concentrations of sulfate or carbonate, competing anions, temperature, and resin poisons have been extensively documented for the reactions shown in equations (2) and (3).[5]

When a strong base anion exchange resin has been loaded with uranyl sulfate, it can be efficiently eluted using acidic chloride (0.9 M NH₄Cl or NaCl and 0.1 M HCl or H₂SO₄) or acidic nitrate (1.0 M NaNO₃ and 0.1 M H⁺) solutions.[2] Elution can also be effected using dilute (~1.0 M) H₂SO₄ although with less efficiency than with chloride or nitrate. Sulfuric acid elution is useful in conjunction with a subsequent solvent extraction process because the eluate can be directly used as feed for that process and the step of converting the resin from the nitrate or chloride form to the sulfate form before the next adsorption cycle can be eliminated.[2]

Acid elution of sorbed UO₂(CO₃)₃⁴⁻ from anion resins would lead to gas generation and consequent disruption of the resin bed. Thus, elution of this species is accomplished with NaCl or NH₄NO₃ solutions.

After elution of uranium, common resin poisons such as thiocyanate ion, S₂O₆²⁻, molybdenum, and iron are removed by treatment with NaOH or NH₄OH solutions. [2,5]

**Important Ion Exchange Equipment and Methods**

The traditional ion exchange systems designed for collecting and concentrating uranium used fixed-bed columns arranged in a "merry-go-round" array. These systems require clarified feeds for smooth operation. Clarification of feeds after leaching is very expensive, accounting for about half of the operating costs of a uranium milling plant.[14] Costs of the clarification process are driven up by the presence of very small particles (ore slimes) of the ore. One way to lower costs is to first separate the ore slimes from the easily washed and clarified sands fraction of the leachate. Sands fraction wash water and ore slimes are then combined to produce a slurry of up to 30% suspended solids. Such slurries are used as feedstock for
resin-in-pulp (RIP) or countercurrent ion exchange processes which have been successful processes for treating unclarified feedstock.

In all CIX systems, the resin bed is fluidized by an upflow stream of feedstock. Continuous ion exchange processes have been described as a major development in ion exchange technology. The success of CIX processes is directly related to the careful experimental and theoretical work of a variety of investigators.[15-22] Three particular fluidized column systems have been used commercially with great success:

NIMCIX System—This system was developed from Cloete-Streat concepts by the National Institute for Metallurgy in South Africa.[23,24]

USBM-MCIX System—This system was developed by the U.S. Bureau of Mines. It is used for in situ uranium mining processes.[17-19]

HIMSLEY COLUMN—The most recently proven fluidized bed contacting system, the Himsley column, has seen use in South Africa, Canada, and the U.S.[21,22]

Porter has developed an upflow, multi-tank fluidized bed ion exchange system for Namibia.[25,26] A downflow pulsed-bed ion exchange method (Higgins Loop CIX) has been used to recover uranium from copper dump leach liquors at Bingham Canyon, Utah.[11]

Continuous RIP processes mix resin beads with desanded pulp, separate the uranium-rich beads by a screening step, and then elute uranium from the beads in a separate system, recycling the regenerated beads for reuse in uranium sorption. One or more of these operations may be used in any given process. An upflow RIP process has been used to recover uranium in France.[27] Several RIP processes have been used successfully over the past 25 years in uranium mills.[27] All use screen-mix systems, which are the only technology capable of handling medium or high density slurries. No basket RIP systems, developed in the 1980's, are in operation. Development of high specific gravity (≥ 1.25) ion exchange resins would significantly extend the scope of RIP applications.

A combination of ion exchange and solvent extraction, based on pioneering work done at Oak Ridge National Laboratory [28], represents the most widely used method for uranium recovery from sulfate ore leach liquors. [22,29-31] The uranium is eluted from ion exchange resins using H₂SO₄ and the eluate is used as feed for an extraction by a tertiary amine.

**SEPARATION PROCESSES**

Uranium, concentrated in fuel assemblies, undergoes fission, after capturing a neutron in nuclear reactors, to generate fission products, energy, and more neutrons. In power reactors,
energy is the desired product; in production reactors, the additional neutrons are the desired product because they react to produce new isotopes. The main isotope of the neutron irradiation of natural uranium is \(^{239}\text{Pu}\), produced when a neutron is captured by \(^{238}\text{U}\) (accompanied by two beta decay reactions). The \(^{237}\text{Np}\) isotope is also produced in much lower yields.

After the irradiated fuel elements are removed from the reactor they are stored in cooling basins to allow short lived fission product isotopes to decay. The last step in the nuclear fuel cycle is the separation of uranium, plutonium, and useful byproducts from each other and from unwanted fission products. Ion exchange cannot compete with solvent extraction in the primary process of uranium recovery, but it is used in several important secondary recovery steps.

**Nuclear Fuel Reprocessing**

Throughout the world, reprocessing of nuclear fuels employs some form of the Purex or Thorex processes, solvent extraction processes using tri-n-butylphosphate (TBP) as the extracting agent. Although no reprocessing of power reactor fuels is planned in the U.S., the Savannah River, Hanford, and Idaho Falls defense sites use TBP-based solvent extraction. European, Japanese, and Indian reprocessing facilities use Purex or modified Purex processes. The Thorex process was developed to process thorium-based fuels.

In Purex reprocessing, such as at the Savannah River Site, irradiated uranium is first dissolved in nitric acid and then separated from plutonium and fission products during solvent extraction with TBP. The high activity waste concentrate from the first Purex cycle contains most of the \(^{237}\text{Np}\) and some of the \(^{239}\text{Pu}\). It also contains aluminum and most of the fission products. As shown in Figure 1 [32], the neptunium and plutonium are separated from other ions, except for thorium, by an agitated bed anion exchange step using a strong base resin. This is possible because Pu(IV) and Np(IV) both form stable anionic hexanitratooxy species in the concentrated (7-9M) nitric acid solution.

The Pu and Np are then eluted with dilute (0.35M) nitric acid as the Pu(IV) and Np(IV) ions and reconverted after elution to the hexanitratooxy ions with 8M nitric acid. A second anion column is used to resorb Pu(NO\(_3\))\(_6^{2-}\) and Np(NO\(_3\))\(_6^{2-}\). Plutonium is separated from neptunium by elution as Pu(III) with 5.5M nitric acid, ferrous sulfamate, and hydrazine. Neptunium ion is not reduced under these conditions and remains sorbed on the resin.

A strong acid cation resin is used next to separate \(^{234}\text{Th}\) that is often found with the \(^{237}\text{Np}\). The neptunium is oxidized to Np(V) in 1-2M nitric acid and, in that oxidation state, is not strongly sorbed by the cation exchanger. Thorium, Th\(^{4+}\), is retained on the resin and is periodically eluted with NaHSO\(_4\). Any \(^{239}\text{Pu}\) is then recycled back to the rest of the process and
FIGURE 1

ION EXCHANGE USED IN REPROCESSING URANIUM

uranium in nitric acid

First cycle Purex process

Second uranium cycle

Pu

Second plutonium cycle

Pu Cation Exchange

PuO₂ - PuF₄

Conversion to

Precipitation

Reduction

Dissolution of residue

Anion exchange

Pu

FissioN products, Al

Aqueous waste

Fission products

Np, Al, trace Pu

Evaporation

Anion exchange Agitated bed

Waste

Anion Exchange Fixed bed Two Stages

Pu

Np

Th cation exchange

Np

Oxalate precipitation Calcination

NpO₂

Pu metal
the $^{237}$Np is used in targets for irradiation to make $^{238}$Pu.

The second cycle of the Purex process produces a dilute (1-3 g/L) plutonium stream. Plutonium is concentrated (Figure 1) on a strong acid cation resin after reduction to Pu(III) with hydroxylamine. Sorbed plutonium is eluted with 5M nitric acid to yield a solution (40-60 g/L) that is concentrated enough for subsequent Pu recovery by precipitation.

When $^{239}$Pu metal is produced by reduction using Ca metal in the recovery process, unreacted plutonium compounds and metal fragments remain in the reaction residue. The residue is treated with concentrated nitric acid to dissolve the plutonium, filtered, and the Pu is sorbed as Pu(IV) on a strong base anion resin. Plutonium collected in this way is recycled to the main process after elution with dilute HNO$_3$.

ION EXCHANGE IN NUCLEAR POWER PLANTS

A wide variety of reactor designs have been used throughout the world to produce power. Regardless of the design, the basic operation employs the use of a controlled nuclear reaction to supply energy to produce steam to drive turbine generators. Ion exchange is used extensively to purify water entering and leaving the reactor.[33,34] These applications are briefly noted below:

- Plant makeup water demineralizers usually are strong base or strong acid nuclear grade resins. Nuclear grade resins are distinguished from regular grade resins by their uniform particle size, their purity relative to leachable ions, and their high degree of conversion to the desired ionic form. These very pure resins generate effluent water of the high quality needed to minimize neutron activation product formation and corrosion.

- Reactor coolant water is purified by mixed bed cation and anion exchange systems, normally found in the H$^+$ and OH$^-$ forms, respectively. In pressurized water reactors (PWR) the anion resin also serves to remove borate ion, a nuclear poison.

- Steam condensates are polished using mixed bed, powdered resins that act as filters as well as ion exchangers. Often, deep bed strong acid/strong base systems are used to treat condensate. Evaporator condensates are usually discharged or recycled as makeup water after treatment.

- Contaminated water from fuel pools and other dilute miscellaneous wastes can be treated by ion exchange to yield high purity water.

DECONTAMINATION OF NUCLEAR WASTES

Waste streams from the reprocessing of nuclear fuels contain
essentially all of the fission products found in irradiated fuel. The number of variations of the Purex or Thorex reprocessing processes used throughout the world leads to an equal number of nuclear wastes, each having its own characteristic properties. It is not possible to consider them all here. Instead, the uses of ion exchange in treatment of some general categories of nuclear wastes are described. In addition, specific waste treatments of interest involving ion exchange technology are noted.

**High Activity Waste Streams**

Waste streams from fuel reprocessing are acidic in nature. At Savannah River and Hanford the pH of the waste is adjusted to 12 or greater with NaOH and the waste is stored in carbon steel tanks. During aging, insoluble materials precipitate and the primary radionuclide left in the supernate, as the soluble portion of the waste is called, is $^{137}\text{Cs}$.

At Hanford [35], cesium ion was recovered and purified by cation exchange. The first ion exchanger used was a zeolite but this was later changed to a phenolic resin, Duolite® ARC-359.[36] Feed for the process consisted of supernate blended with the product from a phosphotungstate precipitation performed on acid waste. These solutions contained as much as 250 Ci of $^{137}\text{Cs}$ per gallon and were about 5M in sodium ion. After sorption, the sodium ion was preferentially eluted from the resin using 0.25M ammonium carbonate. Cesium was eluted with a 3M (NH$_4$)$_2$CO$_3$/2M NaOH mixture at 60°C. Ammonium carbonate in the eluate was thermally decomposed and the resulting solution was concentrated by evaporation and sent to storage.

At the Savannah River Site [37], a resorcinol/formaldehyde condensation resin has been developed that is capable of selectively sorbing 2.5 E-4M Cs$^+$ from supernate containing up to 6M Na$^+$. The cesium can be recovered by elution with 1M HCOOH or 0.5M HNO$_3$. Two modes of final disposal of the waste cesium have been proposed:

- Incorporate the eluate containing cesium ion with feed to the glass melter for final encapsulation in glass and then reuse the resin.
- Incorporate spent resin with the feed to the glass melter for final vitrification.

At West Valley Nuclear Services Co., a subsidiary of Westinghouse Electric Corp., cleanup has begun on about 550,000 gallons of neutralized Purex waste. The initial treatment step involves passing waste supernate through zeolite ion exchange columns to remove Cs$^+$ and Sr$^{2+}$. The loaded zeolite will be combined with washed hydroxide sludge and about 8,000 gallons of acidic Thorex waste, generated during processing of thorium based fuel, for incorporation into glass.[38]
Low Activity Waste

High activity wastes are usually concentrated by evaporation to minimize the storage volume required. Condensate from waste evaporators usually contains radioactive cesium due to entrainment. Removal of the cesium can be effected using a zeolite or an organic ion exchange resin, depending on the pH of the waste stream and the fate of the decontaminated waste stream. For example, evaporator condensate from nuclear power plants is decontaminated using mixed bed ion exchange in order to generate very pure effluent that can be recycled as makeup water.

At Savannah River Site, condensate is decontaminated first by passage through a zeolite bed and then treatment in an effluent treatment facility. A final polishing step in the effluent treatment uses strong base cation exchange to remove the small quantity of cesium ion that remains in the waste after a reverse osmosis step that precedes the ion exchange. The cation resin is eluted with 2M NaNO₃ to remove Cs⁺ and the eluate is incorporated in concrete for storage in vaults.[39]

At Oak Ridge National Laboratory, cesium ion is separated from dilute waste by passage through a column of a phenolic/carboxylic acid resin. It is subsequently eluted with 0.5M HNO₃.[40]

After the accident at Three Mile Island Unit 2 Nuclear Power Station (TMI), large volumes of contaminated water remained in the facility that required decontamination. The Containment Building sump held approximately 600,000 gallons of contaminated water; about 500,000 gallons were stored in the Auxiliary and Fuel Handling building and another 90,000 gallons were in the Reactor Coolant system. The primary radionuclides of concern were ¹³⁷Cs and ⁹⁰Sr in each case but the activity level in the Auxiliary and Fuel Handling building was less by a factor of ten than that in the other two facilities. A slightly different ion exchange treatment was used for the two streams.

Waste water from the Auxiliary and Fuel Handling building was decontaminated by the Epicor II system [41] that was composed of a primary series of carbon steel ion exchange liners (columns) followed by two polishing steps. The liners contained organic ion exchange resins, although some also contained zeolite. The Epicor II system removed approximately 55,000 Ci of gamma emitters, generating 50 highly loaded liners that were disposed of at the commercial waste burial ground at Richland, Washington.

A submerged demineralizer system was used to decontaminate the waste held in the Containment Building sump and the Reactor Core system. Two parallel trains of four ion exchange columns in series were operated in carousel mode while submerged in the reactor fuel pool, which provided shielding. The sorbing medium was a 60/40 mixture of Linde IONSIV™ IE-96 and A-51 zeolites, designed to remove Cs⁺ and Sr²⁺, respectively, with nearly
simultaneous breakthrough of the two radionuclides. The effluent from the zeolite beds was further polished with the Epicor system and stored in tanks for reuse at the TMI facility.

Ion exchange technology has been proposed as a method of removing other fission product radionuclides from nuclear wastes. The pertechnetate ion, TcO₄⁻, is readily sorbed on a strong base resin. Large quantities of 4M nitric acid are required to elute the ⁹⁹Tc ion from the resin and the process by itself has not been cost effective. Recent work has shown that the pertechnetate ion can be removed from the eluate by sorption onto a weak base resin, allowing recycle of the nitric acid as a cost saving measure. Elution from the weak base resin with a small volume of 1M NaOH completes the isolation of this long-lived isotope from the waste stream.[42]

Chelating resins can be used to complex and retain many cationic radionuclides such as ⁹⁰Sr, ⁶⁰Co, ⁵¹Cr, and ⁹²Zr that may be found in trace quantities in low activity wastes. The ¹⁰⁶Ru (nitrosyl) species found in waste streams have historically resisted efficient isolation by ion exchange. Recent work with somewhat flexible sulfur-containing copolymers [43] gives hope that specially designed functional groups may be able to effect removal of nitrosylruthenium ions from nuclear waste.

On a much smaller scale, except for the case of zeolite ion exchangers, several inorganic exchangers have been used for the sorption and separation of radionuclides. These include the zeolites, synthetic zeolites, hexacyanoferrates, oxides and hydrous oxides, and acidic salts of multivalent metals.[44]

Non-Radioactive Waste-Hg

Fuel elements containing alloys of aluminum with enriched uranium are reprocessed by dissolution in nitric acid using mercuric nitrate as a catalyst.[45] The mercury introduced at this point ultimately finds its way into both high- and low-activity wastes. Strict regulation of dispensing and storing waste streams containing toxic materials such as mercury has led to studies of ways to isolate and immobilize dissolved mercury from the waste to protect the environment. Duolite® GT-73 cation exchange resin is used extensively at the Savannah River Site [39] to remove Hg²⁺, Hg₂²⁺, and dissolved Hg⁰ from dilute waste streams.

OTHER ACTINIDE SEPARATIONS OF INTEREST

Ion exchange has been used in several applications to separate actinide isotopes produced in special irradiation campaigns.

Transplutonium Elements

A decade and a half ago, kilogram quantities of ²⁴³Am and ²⁴⁴Cm and milligram amounts of ²⁵²Cf were produced at Savannah River.[46] At about the same time, Oak Ridge produced those
radionuclides as well as $^{249}$Bk, $^{253}$Es, and $^{257}$Fm in their High Flux Isotope Reactor.\cite{47} The heavy transplutonium elements are made by neutron irradiation of a mixture of Pu isotopes \cite{46,48} or a mixture of Pu, Am, and Cm.\cite{47} The ion exchange separations of actinides at Oak Ridge and Savannah River were accomplished using high pressure ion exchange.\cite{46,48-51} There are two major advantages in operating high pressure ion exchange for this particular application. First, radiation damage to the resin is minimized because very small resin beads are used, allowing for rapid flow rates. Second, high pressure keeps radiolytic gases, that otherwise would disrupt the bed, in solution.

At Oak Ridge, the lanthanides and actinides were separated from each other on a strong base ion exchanger. Lanthanides were eluted with 10M LiCl and most of the Am and Cm were then eluted with 9M LiCl. The rest of the americium and curium and the heavier actinides were eluted with alpha-hydroxyisobutyrate (AHIB).\cite{50} The heavier nuclides elute first because they form more stable complexes with AHIB.

The Savannah River process used displacement development, similar to that used at Hanford \cite{52} to purify Am, on cation exchange resin to effect separation and purification. The lanthanides and actinides were first sorbed on the resin column behind a barrier band of zinc ion. Development of the band with diethylenetriaminepentaacetic acid (DTPA) moved the more stable actinide complexes to the front of the band, leaving the lanthanides behind. The actinides were resorbed on the resin when the band reached the zinc because zinc forms an even more stable complex with DTPA. The net result was that the actinides separated into discrete bands on the resin. They were eluted, resorbed and developed on a series of smaller columns, and finally eluted in very pure form.

Concentration of $^{233}$U from $^{232}$Th Irradiation Products

Neutron irradiation of $^{232}$Th produces $^{233}$U.\cite{53,54} The two isotopes are separated from each other by the Thorex solvent extraction process using TBP. The uranium stream is concentrated by sorption of $^{233}$U on a strong acid cation resin followed by elution with a NH$_4$NO$_3$/HNO$_3$ mixture.

$^{241}$Am from Aged $^{241}$Pu

A product of the beta decay of $^{241}$Pu is $^{241}$Am, which is used in commercial applications such as smoke detectors, thickness gauging, and oil well logging.\cite{55} The separation of plutonium and americium is possible by solvent extraction \cite{56,57}, molten salt extraction \cite{55}, or precipitation.\cite{58} Each of these separation processes, however, must be coupled with at least one ion exchange process step. Cation exchange allows a preliminary separation of Pu(III) and Am(III) from ionic impurities of +1 or +2 charge. Also, anion exchange of Pu(NO$_3$)$_6^{2-}$ from concentrated nitric acid solution effects its separation from Am(III).
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