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(54) PROCESS FOR RECOVERING MOLYBDENUM-99 FROM A MATRIX CONTAINING NEUTRON IRRADIATED FISSIONABLE MATERIALS AND FISSION PRODUCTS

KERNFORSCHUNGSZENTRUM KARLSRUHE GESELLSCHAFT MIT BESCHRAENKTER HAFTUNG, formerly Gesellschaft Fuer Kernforschung M.B.H., of 5 Weberstrasse, D-7500 Karlsruhe 1, Federal for which we pray that a patent may be granted to us, and the method by which it is for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:-

The present invention relates to a process for the recovery of molybdenum-99 from a matrix containing neutron irradiated, fissionable materials and fission products, in which the matrix is decomposed in an aqueous alkali hydroxide solution and the molybdenum-99 and part of the fission products are dissolved, the solution containing the molybdenum-99 is separated from a residue of solid particles containing at least actinides and lanthanides and is treated with thiocyanate ions in order to form a molybdenum-99 complex.

In nuclear medicine, the significance of Tc-99 is continuously at an increase as an indicator in the diagnosis of tumors. Since, however, technetium has a relatively short halflife ($T_{1/2}$ =6.0 h), the mother nuclide Mo-99 is eluted when required. Thus, a technetium generator is used to provide the technetium. The technetium generators generally comprise a chromatographic column having Mo-99 bearing molybdate ion absorbed thereon. Radioactive decay of the relatively long-lived Mo-99 produces Tc-99. Elution of the chromatographic column provides an onthe-spot source of the technetium.

Previously, natural molybdenum which had been activated in reactors was used in the generators to produce the technetium. The drawbacks of this natural molybdenum material are that large columns are required for small specific activities, large injection volumes are required in order to retain the required activity, and there is a very limited availability of the generator due to the low

Fission molybdenum has been found to be useful in technetium generators to produce the technetium and has been used to a greater degree in recent times. This, however, requires much refined processing technology in order to obtain the required degrees of molybdenum purity. The required radionuclide purity of the fission molybdenum for use in a technetium generator is:

I-131<0.05 μ C/mC Mo-99 Ru-103<0.05 μC/mC Mo-99 Total γ contamination <0.1 μ C/mC Mo-99

no more than 1 nanocurie total α activity per curie Mo-99 Sr-89<6×10⁻⁴ μC/mC Mo-99 Sr-90<6×10⁻⁵ μC/mC Mo-99

40 A long known method for proving the presence of molybdenum where the 40 molybdenum is present in solution as molybdate comprises reducing the molybdate with SnCl₂ to Mo(III), then binding the Mo(III) to SCN⁻ ions to form a thiocyanate complex, and thereafter extracting the thiocyanate complex with the aid of diethyl ether. This method is completely useless in the recovery of Mo-99 because of the great volatility and combustibility of the diethyl ether inasmuch as the risk of a fire 45 45

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	of explosion must be completely eliminated when working with radioactive substances.	
5	A number of publications discuss methods which use thiocyanate ions, but these methods operate principally with the use of additional extracting agents in the organic phase, such as, for example, with tributyl phosphate (TBP) (Gorlach, V. F., Marchenko, L. M. (Kiev State University), <i>Ukr. Khim. Zh.</i> ; 40: No. 9, 983—985, September, 1974 (in Russian)); or with tribenzylamine (Yatirajam, V., Ram, Jaswant	5
10	(University of Kurukshetra, India), Anal. Chim. Acta, 59; No. 3, 381—387, May 1972); or with 2-furaldehyde (Spaccamela Marchetti, Elena, Cereti Mazza, Maria Teresa (Politecnico, Turin) Ann. Chim., Rome, 59: 902—911, 1969 (in Italian)).	10
	These methods have the drawback that the additional organic extraction agents may lead to contamination of the final product which could result in behavioural malfunctions of the molybdenum on the generators. Furthermore, these	
15	contaminants may have a pyrogenic effect. German Offenlegungsschrift No. 23 49 804 (laid-open on April 17th, 1975) discloses a process for the recovery of molybdenum-99 from fission products in which the matrix containing irradiated uranium, in the case of the use of uranyl nitrate as target, is dissolved in water and treated, before the separation of Mo.	15
20	with 5 N HNO ₃ , or, in the case of the use of an uranium aluminum matrix, the matrix is dissolved within 30 minutes in a 5 M sodium solution which is 2 molar nitrite. In both cases, the solution is then filtered from the residue. The molybdenum is then separated out of the respective solutions from the other fission products by two dividing operations which can be combined and repeated in a	20
25	certain sequence. When an alkali solution is used as the starting solution, it is necessary to pretreat the solution before the first dividing operation. This pretreatment includes an addition of potassium iodide solution (as carrier for iodine), neutralization and acidification of the alkali solution with silver nitrate containing 5 molar HNO ₃ , a standing for 20 minutes to precipitate AgI, and a separation of the precipitation deposit from the solution.	25
30	Then the solution is treated according to the first dividing operation. In the first dividing operation, the solution containing the molybdenum as molybdate, which solution is about 2 molar HNO ₃ , is fed to a column charged with aluminum oxide. This causes practically all fissionable substances and fission products to be sorbed onto the column. The aluminum oxide column is then subjected to a	30
35	number of elution steps. Firstly, the column is eluted with 1 N HNO ₃ . The time required for this first elution, for a column having a diameter of 2 cm and a 8 cm fill level, is about 100 minutes. (The time periods disclosed hereafter for the further steps of the first dividing operation are with respect to a column having a diameter of 2 cm and a 8 cm fill level). Thereafter, the aluminum oxide column is treated	35
40	with water. The time required for the water treatment is about 60 minutes. The aluminum oxide column is then eluted with a 0.1 N NH ₄ OH solution for a required time of about 30 minutes and subsequently with a 25 ml, 1 N NH ₄ OH solution for a required time of about 8 minutes. The elution with nitric acid and diluted	40
45	ammonium hydroxide solution separates the uranium and all fission products, except for tellurium, from the molybdenum. The molybdenum is then supposed to be elutable with a further 1 N NH ₄ OH solution to provide an 80% yield. The time required for the molybdenum elution is about 40 minutes. In the second dividing operation, the eluate containing Mo-99 and	45
50	contaminants, such as, for example, iodine, is acidified with HNO ₃ to a pH of 1 to 2. Potassium iodate and sulfurous acid are added, and thereafter potassium iodate and potassium iodide are again added, as carriers for iodine. The additions of carrier in this sequence are to assure complete conversion of the fission product iodine to iodide. Then the solution is sucked slowly through a layer of fine particles	50
55	of freshly-made silver chloride. The time required for this treatment with silver chloride is about 15 minutes. The molybdate yield in this operation is supposed to be about 95%. The drawbacks of this process can easily be discerned and include time consuming and complicated mode of operation, sorption of all actinides and almost	55
60	all fission products together with molybdenum on the aluminum oxide, carrying along of contaminants with the molybdenum during the elution and low total yield of approximately 76%. In addition, elemental iodine is carried along throughout the entire process due to the use of HNO ₃ . It is an object of the present invention to overcome the drawbacks of the	60
65	known processes and provide a simple, easily-practised and safe process which ensures a very high yield of an extremely pure Mo-99 product.	65

	A further object of the present invention is to provide such a process which has	
	a minimum of process stages, such as, for example, sorption stages and elution stages, and results in a reduction of the quantity of contaminated, organic waste.	
5	Another object of the present invention is to provide such a process which is easily handled as a matter of routine and with remote control.	-
-	According to the present invention there is provided a process for recovering	5
	molybdenum-99 from a matrix which has been irradiated with neutrons and contains fissionable materials and fission products, wherein the matrix is	
10	decomposed in an aqueous alkali hydroxide solution and the molybdenum-99 and	
10	part of the fission products are dissolved, the solution containing the molybdenum- 99 is separated from a residue of particles containing at least actinides and	10
	lanthanides and is treated with thiocyanate ions to form a molybdenum complex	
	comprising the steps of: a) conditioning the alkali solution containing molybdenum in the form of	
15	molybdate ($MoO_A^{}$) with an iodine reduction agent in a quantity corresponding to	15
	a concentration range of from 10 ⁻⁴ Mol to 0.2 Mol per liter alkali solution; b) adding mineral acid to the alkali solution until a hydronium ion	
	concentration in the range from 0.1 to 6 Mol/l has been reached.	
20	c) reducing the molybdenum contained in the acidified solution of step b) to form three-valent molybdenum Mo(III) and complexing the Mo(III) with SCN-	20
	lons to form [Mo(SCN) ₈] ³ ions, said SCN ⁻ ions being present in an ion	20
	subjected to the reduction:	÷
25	d) treating the $[Mo(SCN)_e]^{3-}$ jon-containing acid solution from step c) with a	
23	previously conditioned, organic ion exchanger of the time of a chelate-forming synthetic resin consisting of a styrene divinyl benzone copolymer containing	25
	methylene nitrilo diacetate groups as functional groups and having a particle size in	
	the range of from 35 μ to 840 μ for selectively sorbing the molybdenum; e) separating the ion exchanger from step d), which is charged with	
30	molybdenum, from the solution now free of molybdenum; f) washing the separated molybdenum-charged ion exchanger with a wash	30
	solution of diluted mineral acid containing a weak concentration of an iodine	
	reduction agent, the volume of the wash solution corresponding to 5 to 10 times the volume of ion exchanger employed, in order to remove residual quantities of the	
35	molybdenum-free solution;	35
	g) eluting the molybdenum from the washed ion exchanger with a liquor at an elution temperature in the range from 20°C to 70°C.	33
	In one embodiment of the present invention, the reduction to form the Mo(III)	
40	ions and the complexing of the Mo(III) ions is effected by mixing the acidified solution from step b) with an aqueous thiocyanate ion solution containing metallic	40
	Zinc or metallic aluminum to form a solution which contains the metallic zinc or	40
	aluminum in a concentration range of from 10 mg/l to 2000 mg/l and a concentration of thiocyanate ions in the range of from 0.1 Mol/l to 3 Mol/l of	
45	solution; and reducing the molybdenum contained in the acidified solution of step	
	b) with the aid of the hydrogen produced from a reaction between the hydronium ions and the metallic zinc or the metallic aluminum to form the three-valent	45
	molybdenum Mo(III) which then complexes with the SCN- ions to form [Mo(SCN) ₈] ³⁻ ions.	
50	In another embodiment of the present invention, the reduction to form the	
50	Mo(III) ions and the subsequent complexing is effected by initially subjecting the acidified solution from step b) to a cathodic reduction whereby the molybdenum is	50
	reduced to molybdenum (III), and bringing the resulting Mo(III) into contact with	
	thiocyanate ions to form complexes, the concentration of the thiocyanate ions in the solution being subjected to the cathode reduction being in the range of from 0.1	
55	MOVI to 3 MOVI.	55
	In the process of the present invention, an alkali solution which contains molybdenum as molybdate (MoO ₄ -) is treated. This alkali solution generally is	
	one which is formed during the recovery of Mo-99 from targets which have been	
60	subjected to an enrichment process. Customarily, Mo-99 is recovered and isolated from plates or cylinders in the form of uranium-aluminum targets which have been	60
	enficied with dramum-233. In order to realize a high Mo-99 yield or fission rate	00
	with the same quantity of uranium, the target material contains about 93°, of enriched uranium-235	
65	In view of the short cooling periods for the target to be processed, iodine isotopes, such as I-131, I-132 and I-133, are very much involved in the iodine	
	and 1-133, are very much involved in the lodine	65

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	emission rate. In order to assure that this emission rate is reduced to values below those set by safety councils, agencies and laws of various countries, the uranium-	
	aluminum targets are decomposed in an alkali solution and iodine reduction agents	
_	are added so that elemental iodine is converted to iodide ions. Thus, and due to the	
5	avoidance of the presence of NO ₃ ⁻ or NO ₂ ⁻ ions in the process according to the	5
	present invention, the possible danger of fire in the activated carbon filter beds	
	being used to retard the fission gases Xenon-133 and 135 is prevented. If a U/Al alloy containing uranium which has been enriched to 93% is selected	
	to produce Mo-99, the yield from a U/Al sample, for example, containing 1 g U-235	
10	without consideration of the flux depression of the sample, and with a neutron flux	10
	of 5·10 ¹³ cm ⁻² sec ⁻¹ , after a period of irradiation and a period of cooling and a	10
	period of processing of three days each, is 30 C Mo-99.	
	It is also possible, however, to subject other matrices containing uranium as	
	the fissionable material or other fissionable materials, respectively, to neutron	
15	irradiation in order to produce Mo-99.	15
	For an alkali decomposition, the matrix is treated with caustic soda solution at	
	about 120°C. Thus, the aluminum and the fission products molybdenum, tellurium and iodine, as well as the alkali and earth alkali metals are quantitatively dissolved.	
	Part of the resulting fission products zirconium and ruthenium are also dissolved,	
20	while the lanthanides and actinides as well as the major portion of Ru and Zr	20
	remain undissolved in the form of mud.	. 20
	After filtering the mud, the alkali solution is used as the starting solution to	
	obtain a highly pure molybdenum-99 according to the process of the invention.	
2.5	In the practice of the present invention, the starting alkali solution is	
25	conditioned with an iodine reduction agent. Exemplary of suitable iodine reduction agents are sulfite ions in aqueous solution, such as, for example, sodium sulfite or	25
	potassium sulfate or a mixture thereof. In addition, hydroxyl ammonium sulfate or	
	hydrazine sulfate can be used as iodine reduction agents. The iodine reduction	
	agent is used in a quantity corresponding to a concentration range of from 10 ⁻⁴ Mol	
30	to 0.2 Mol per liter of alkali solution. In order to condition the alkali starting	30
	solution with Na ₂ SO ₃ as the iodine reduction agent, it is sufficient, for example, to	
	have a sulfite concentration in the alkali solution of 0.1 Mol to 0.05 Mol per liter.	
	In the practice of the present invention, a mineral acid is added to the alkali	
35	solution until a hydronium ion concentration (H ₃ O ⁺ ion concentration) in the range between 0.1 and 6 Mol/l has been reached. The mineral acid which is added to the	35
55	alkali solution advantageously can be hydrochloric acid or sulfuric acid. The acid	33
	concentration after addition of the acid preferably can be, for example, in the	
	range from 0.5 to 3 Mol/l.	
40	In the practice of the present invention, the molybdenum contained in the	
40	alkali solution is reduced to form three-valent molybdenum Mo(III) and the	40
	Mo(III) is complexed with SCN ⁻ ions to form [Mo(SCN) ₈] ⁻³ ions. The valency of	
	the Mo ⁺⁺⁺ formed by the reduction is thus stabilized by the creation of the stable thiocyanate complex [Mo(SCN) ₆] ⁻³ . This reduction and complexing can be	
	performed in either of two techniques.	
45	In the first reduction and complexing technique, an aqueous thiocyanate-	45
-	containing solution is added to the alkali solution to form a conditioned alkali	73
	solution which contains mineral acid and thiocyanate ions. The thiocyanate ion	
	containing solution which is added to the alkali solution can be a solution of	
50	NH ₄ SCN, NaSCN or KSCN or mixtures of these thiocyanate ion forming	
50	compounds. In addition to containing these thiocyanate ion forming compounds,	50
	the thiocyanate solution which is added to the alkali solution contains metallic zinc or metallic aluminum. The solution which is formed upon addition of the	
	thiocyanate containing solution to the alkali solution contains the metallic zinc or	
	aluminum in a concentration range between 10 mg/l and 2000 mg/l and a	
55	concentration of thiocyanate ions in the range between 0.1 Mol/l and 3 Mol/l.	55
	The thiocyanate concentration sufficient for complex formation generally is	00
	about 1 to 2 Mol/1 of solution. In the presence of larger quantities of aluminum or	
	foreign cations, respectively, the thiocyanate concentration may possibly have to	
60	be increased to 3 Mol/l The molyhdanum in this conditioned saidified and this events containing	
60	The molybdenum in this conditioned, acidified, and thiocyanate-containing solution is reduced with the aid of the hydrogen produced from a reaction between	60
	the hydronium ions and the metallic zinc or the metallic aluminum to form three-	
	valent molybdenum Mo(III). The Mo(III) is then complexed with the SCN ⁻ ions in	
	the solution to form $[Mo(SCN)_6]^{-3}$ ions.	
65	The conditioning and acidification of the alkali solution and addition of the	65
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thiocyanate solution to the alkali solution can take place in the sequence of firstly conditioning, secondly acidifying, and thirdly adding the thiocyanate solution. The conditioning, acidification and addition of the thiocyanate solution, however, need not necessarily take place in the above sequence, and can be effected in a different 5 sequence such as adding SCN- to the alkali solution and subsequent acidifying. In the second technique for reducing and complexing the molybdenum in the alkali solution, the molybdenum is reduced by a cathode reduction. In this technique, a conditioned and acidified alkali solution is initially subjected to a cathodic reduction to reduce the molybdenum to Mo(III). The resulting Mo(III) is 10 brought into contact with thiocyanate ions to form the [Mo(SCN)₆]⁻³ complexes. 10 The concentration of the thiocyanate ions in the acidified alkali solution to effect this complexing is in the range between 0.1 Mol/l and 3 Mol/l. To achieve the desired thiocyanate ion concentration in the alkali solution, a thiocyanatecontaining solution of NH4SCN, NaSCN, or KSCN or mixtures thereof can be 15 added to the alkali solution. As in the first technique, the various additions to the 15 alkali solution can be in the sequence of conditioning, acidification and addition of thiocyanate containing solution, but other sequences can be employed.

The cathodic reduction of the molybdate to Mo(III) results in the great advantage that no further quantities of ions which would increase the proportion of 20 solids in the solution and thus the waste volume are introduced into the solution 20 and that a lower concentration of thiocyanate ions is required. In the practice of the present invention, the acid solution containing the [Mo(SCN)₆]³⁻ ions is treated with a previously-conditioned, organic ion exchanger. The organic ion exchanger is of the type of a chelate-forming, synthetic resin consisting of a styrene divinyl benzene copolymer containing methylene nitrilo 25 25 diacetate groups as functional groups and have a particle size in the range between 35 μ and 840 μ for the selective sorption of the molybdenum. The organic ion exchanger could consist of e.g. CH,COOH СН,СООН 30 or 30 CH,COONa The [Mo(SCN)₆]³⁻ complex is selectively and quantitatively retained by the pretreated or conditioned organic ion exchanger which may be contained, for example, in a short, thick-walled glass column. All fission products considered to 35 be contaminants in the production of Mo-99 and other constituents of the treated 35 solution flow through this glass column. Conditioning of the organic ion exchanger can be effected with a mineral acid, such as hydrochloric acid or sulfuric acid, as a mobile phase which acts on the ion exchanger which serves as a stationary phase. Conditioning of the organic ion exchanger is best effected with a diluted thiocyanate ion containing mineral acid, 40 40 such as diluted hydrochloric acid or diluted sulfuric acid both containing a small amount of about 0.01 percent by volume of H₂SO₃. The quantity of the mobile conditioning phase that is used is preferably from 5 to 20 ml per gram of ion exchanger. Suitable concentrations of the diluted hydrochloric and sulfuric acids are 2 to 4 N HCl or H₂SO₄ and the concentration of sulfite ions and thiocyanate 45 45 ions is 0.01 N sulfite and 0.1 N SCN-. The treatment of the acid solution containing the [Mo(SCN)₆]⁻³ ions with the ion exchanger charges the ion exchanger with the molybdenum complex and the charged exchanger is then separated from the remaining solution which is now free 50 of molybdenum. 50

The separated molybdenum-charged ion exchanger is then washed with a wash solution of diluted mineral acid which contains a low concentration of an iodine reduction agent in order to remove residual quantities of the molybdenum free solution. The quantity of wash solution corresponds to 5 to 10 times the volume of the quantity of ion exchanger employed. The wash solution can be, for example,

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5	diluted hydrochloric or sulfuric acid. Thus, the wash solution for the ion exchanger charged with $[Mo(SCN)_6]^{3-}$ can be, for example, 0.1—0.001 M hydrochloric acid or 0.1—0.001 M sulfuric acid which is 0.01—0.0001 molar in H_2SO_3 . After washing, the molybdenum from the washed ion exchanger is then eluted with a liquor at an elution temperature in the range from about 20°C to	5
10	about 70°C. The elution of Mo-99 from the charged and washed ion exchanger can be effected, for example, with a caustic soda solution or a caustic potash solution of a concentration in the range from 0.1 Mol/l to 10 Mol/l under normal (atmospheric) pressure or with aqueous ammonium hydroxide solution of a concentration in the range from 1 Mol/l to 10 Mol/l at increased (above atmospheric) pressure up to 10 atmospheres. The elution can also be effected with a liquor of a higher concentration, but higher concentrations generally are not necessary and are not appropriate.	10
15	The elution of Mo-99 is preferably effected with a 0.5 molar to 6 molar caustic soda solution in a quantity of about 50 ml per mg Mo-99 at 60°C or with 2 molar to 6 molar NH ₄ OH solution in a quantity of about 50 ml per mg Mo99 at 60°C and a pressure of about 3 atmospheres.	15
20	For an ion exchanger particle size between 75 μ and 150 μ , the most favorable elution temperature lies in a range from 50° to 60°C, and, for a particle size between 150 μ and 300 μ , the most favorable elution temperature lies at about 20°C. No contaminants can be found in the eluted molybdenum. The	20
25	decontamination factor of the process lies above 10°. The resulting degree of purity for the Mo-99 produced according to the process of the invention is greater than that set for use in medicine. As shown by experiments, the yield of Mo-99 is more than 90% and, under optimum process conditions, even more than 99.5% of the Mo-99 originally present in the alkali starting solution.	25
30	The process according to the present invention has a number of further advantages. For example, volatile, easily-combustible, organic substances, such as, for example, organic solvents, are not present in this process. Thus, a danger of fire is eliminated. This is of particular significance for the handling of freshly-irradiated nuclear fuels, in view of the still present particularly dangerous volatile nuclides, such as, for example, I-131, I-132 and I-133. Additionally, the danger of formation	30
35	of volatile, difficultly-adsorbable organic iodine compounds, such as, for example, CH ₃ I, is minimized. The retention capability of a column with the said ion exchange resin for molybdenum is independent of the aluminum concentration which may still remain in the solution with the use of aluminum as reducing agent in step c).	35
40	The present invention will be explained below with the aid of a few experiments. Experiment 1	40
45	Experiment 1 provides proof of the efficacy of the process of the present invention relative to the separation of Mo-99 from other fission product nuclides. To determine the decontamination factor, an alkali solution containing, in addition to Mo-99, the radioactive fission products Cs-137, Ba-140, Ce-143, La-140, Nd-147, Ru-106, Te-132, I-131, I-132 and I-133, was conditioned with Na ₂ SO ₃ and acidified with sulfuric acid. Thiocyanate ions and zinc granules were added to the	45
50	solution. The decontamination factors were more than 10 ⁵ for Cs, Ba, Ce, La, Nd and more than 10 ⁵ for I-131, Te and Ru. Separation was effected after the preparation of the solution (stages a, b and c in stage d) in which Mo-99 is selectively absorbed on the ion exchanger. All other fission product nuclides remain in the solution.	50
55	Experiment 2 This experiment serves to establish the yield of Mo-99 as a ratio of the amount of Mo-99 which was treated in the acidified solution before treatment with the ion-exchange resin. Experiment 2 therefore begins with the solution obtained from step c) and is concerned with steps d) to g) of the present invention. To determine the yield of Mo-99 after treatment with the ion exchange resin in	55
60	relation to the quantity of Mo-99 present in the acidified solution in the form of the [Mo(SCN) _e] ³⁻ complex before treatment with the ion exchange resin, 100 ml of a sulfuric acid solution containing approximately 3 Mol/l H_3O^+ ions and 4.3 mg Mo present as a thiocyanate complex were added during approximately 10 minutes to approximately 18 ml of a conditioned ion exchange resin of a grain size of 75 μ to	60

Even with the use of larger ion exchange sorption columns, such as those having a volume of greater than 20 cm³, the flowthrough speed during addition of the acidified solution containing the [Mo(SCN)₆]³⁻ ions and during addition of the wash solution can be one-half the ion exchange column volume per minute, and, during addition of the elution liquor (alkali elution solution) to elute the Mo-99, the flowthrough speed can be 1/10 of the column volume per minute. Fluctuation in the flowthrough speed by a factor in the range of 1/2 to 2 of the given values is possible, however, without adverse influence on the process.

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WHAT WE CLAIM IS:-

1. Process for recovering molybdenum-99 from a matrix which has been irradiated with neutrons and contains fissionable materials and fission products, wherein the matrix is decomposed in an aqueous alkali hydroxide solution and the molybdenum-99 and part of the fission products are dissolved, the solution containing the molybdenum-99 is separated from a residue of particles containing at least actinides and lanthanides and is treated with thiocyanate ions to form a

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molybdenum complex, comprising the steps of:

a) conditioning the alkali solution containing molybdenum in the form of molybdate (MoO₄⁻) with an iodine reduction agent in a quantity corresponding to a concentration range of from 10⁻⁴ Mol to 0.2 Mol per liter alkali solution; b) adding mineral acid to the alkali solution until a hydronium ion

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concentration in the range from 0.1 to 6 Mol/l has been reached.

c) reducing the molybdenum contained in the acidified solution of step b) to form three-valent molybdenum Mo(III) and complexing the Mo(III) with SCN ions to form [Mo(SCN)₆]³ ions, said SCN ions being present in an ion concentration in the range of from 0.1 Mol/l to 3 Mol/l of the solution being subjected to the reduction;

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d) treating the [Mo(SCN)₆]³⁻ ion-containing acid solution from step c) with a

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	previously conditioned, organic ion exchanger of the type of a chelate-forming synthetic resin consisting of a styrene divinyl benzene copolymer containing methylene nitrilo diacetate groups as functional groups and having a particle size in the representations of from 25 may 240 m for calculatively corbins the malybdonum.	
5	the range of from 35 μ to 840 μ for selectively sorbing the molybdenum; e) separating the ion exchanger from step d), which is charged with molybdenum, from the solution now free of molybdenum; f) washing the separated molybdenum-charged ion exchanger with a wash	5
	solution of diluted mineral acid containing a weak concentration of an iodine reduction agent, the volume of the wash solution corresponding to 5 to 10 times the	
10	volume of ion exchanger employed, in order to remove residual quantities of the	10
	molybdenum-free solution; g) eluting the molybdenum from the washed ion exchanger with a liquor at an elution temperature in the range from 20°C to 70°C.	
	2. A process as claimed in claim 1, wherein the reduction to form the Mo(III)	
15	ions and the complexing of the Mo(III) ions is effected by mixing the acidified solution from step b) with an aqueous thiocyanate ion solution containing metallic	15
	zinc or metallic aluminum, to form a solution which contains the metallic zinc or aluminum in a concentration range of from 10 mg/l to 2000 mg/l and a	
20	concentration of thiocyanate ions in the range of from 0.1 Mol/l to 3 Mol/l, and reducing the molybdenum contained in the acidified solution of step b) with the aid	20
	of the hydrogen produced from a reaction between the hydronium ions and the metallic zinc or the metallic aluminum to form the three-valent molybdenum	20
	Mo(III) which then complexes with the SCN ⁻ ions to form [Mo(SCN) ₆] ³⁻ ions. 3. A process as claimed in claim 1, wherein the reduction to form the Mo(III)	
25	ions and the subsequent complexing is effected by initially subjecting the acidified solution from step b) to a cathodic reduction whereby the molybdenum is reduced	25
	to molybdenum (III), and bringing the resulting Mo(III) into contact with	
	thiocyanate ions to form complexes, the concentration of the thiocyanate ions in the solution subjected to the cathode reduction being in the range of from 0.1 Mol/l	
30	to 3 Mol/l.	30
	4. A process as claimed in claim 1, 2 or 3, wherein sulfite ions in aqueous solution are used as the iodine reduction agent.	-
	5. A process as claimed in claim 4, wherein the sulfite ions are supplied by	
35	sodium sulfite, potassium sulfite or a mixture thereof. 6. A process as claimed in claim 1, 2 or 3, wherein hydroxyl ammonium sulfate	35
	is used as the iodine reduction agent. 7. A process as claimed in claim 1, 2 or 3, wherein hydrazine sulfate is used as	33
	the iodine reduction agent.	
40	8. A process as claimed in any preceding claim, wherein hydrochloric acid or sulfuric acid is employed for the addition of mineral acid in step b) and for the wash	40
	solution in step f).	40
	9. A process as claimed in any preceding claim, wherein a solution of NH ₄ SCN, NaSCN, KSCN or a mixture thereof is used to provide the thiocyanate	
45	ions for step c). 10. A process as claimed in any preceding claim, wherein the conditioning of	
7.5	the organic ion exchanger is effected with dilute hydrochloric acid or dilute sulfuric	45
	acid. 11. A process as claimed in claim 10, wherein the conditioning of the organic	
50	ion exchanger is effected with diluted hydrochloric acid or dilute sulfuric acid in a quantity of 5 to 20 ml/g ion exchanger.	50
	12. A process as claimed in claim 11, wherein the conditioning of the organic	50
	ion exchanger is effected with a thiocyanate-containing dilute hydrochloric acid or a thiocyanate-containing dilute sulfuric acid containing a small proportion of about	
55	0.01 percent by volume of H ₂ SO ₃ . 13. A process as claimed in any preceding claim, wherein the elution of the	
55	Mo-99 in step g) is effected with caustic soda solution or caustic potash solution of	55
	a concentration in the range of from 0.1 Mol/l to 10 Mol/l under normal pressure. 14. A process as claimed in claim 13, wherein the elution of the Mo-99 is	
60	effected with 0.5 molar to 6 molar caustic soda solution in a quantity of about 50 ml	
60	per mg Mo-99 at 60°C. 15. A process as claimed in any one of claims 1 to 12, wherein the elution of the	60
	Mo-99 in step g) is effected with aqueous ammonium hydroxide solution of a	
	concentration in the range of from 1 Mol/l to 10 Mol/l at increased pressure up to 10 atmospheres.	
65	16. A process as claimed in claim 15, wherein the elution of the Mo-99 is	6.

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effected with 2 molar to 6 molar NH₄OH solution in a quantity of about 50 ml per mg Mo-99 at 60°C under a pressure of about 3 atmospheres.

17. A process as claimed in any preceding claim, wherein the elution temperature for an ion exchanger particle size of from 75 μ to 150 μ lies in a range of from 50° to 60°C.

18. A process as claimed in any one of claims 1 to 16, wherein the elution temperature for an ion exchanger particle size of from 150 μ to 300 μ lies approximately at 20°C.

19. A process for recovering molybdenum-99 from a matrix containing neutron irradiated fissionable materials and fission products as claimed in any preceding claim, substantially as hereinbefore described.

20. Molybdenum-99 wherever recovered by a process as claimed in any one of claims 1 to 19.

POTTS, KERR & CO., Chartered Patent Agents, 15, Hamilton Square, Birkenhead, Merseyside, L41 6BR. and 27 Sheet Street, Windsor, Berkshire, SL4 1BY.

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