STANDARDS IN THE NUCLEAR ENERGY PROGRAM

C. J. RODDEN

United States Atomic Energy Commission, New Brunswick Laboratory, New Brunswick, New Jersey

Summary—The types of standards used in the nuclear energy field will be considered under the heading of chemical, spectrochemical and isotopic standards. The various fuel materials used as standards, such as uranium, plutonium, thorium and their compounds, are discussed. In addition, standards of certain non-fuel materials of special interest in certain operations, such as zirconium and boron, are considered.

According to Webster, a standard is defined as "that which is established by authority, custom or general consent, as a model or example, criterion, test in general." To the analyst, a standard is usually considered as a standard sample and is defined as a homogeneous material, which resembles as closely as possible, in chemical and physical nature, the material with which the analyst expects to deal, and which has been analyzed by a sufficient number of methods and analysts to establish its composition with considerable certainty.

Theoretically, the analyst should be provided with as many standard samples as he has different kinds of material to analyze. Such a program would require thousands of samples and is quite out of the question. For many important groups of materials the National Bureau of Standards issues standard analyzed samples.

In addition to standard analyzed samples, the analyst also uses primary standards. To the analyst, the term "primary standard" generally means an element or compound of very high purity which is usually certified as to composition by an organization such as the National Bureau of Standards. From the above, you will note that results on standard analyzed samples are given in terms of composition.

There is a widely prevalent misconception, even among analysts themselves, as to the accuracy of the results obtained in analytical work, and many confuse precision with accuracy. Accuracy is a measure of the degree of correctness. Precision is a measure of reproducibility of results in the hands of a given operator or operators. The precision of a result does not necessarily guarantee its accuracy. It is a comparatively simple matter to establish the precision that can be expected if a certain method of analysis is applied by an analyst or group or analysts to a given material. On the other hand, it may be extremely difficult to judge the probable accuracy of a result, for all results are matters of opinion rather than fact, and so the true value is never known. This is one of the most unsatisfactory aspects of chemical analysis. When an analyst scans a result, he never can be sure whether the determinator provided for all interfering compounds, hoped that they were absent, or was too ignorant to worry. This brings up the question of the definitions of determinators and analysts which was so ably discussed by Dr. G. E. F. Lundell of the National Bureau of Standards in his famous lecture given in 1933 and entitled "The Chemical Analysis of Things as They Are" (Ind. Eng. Chem. Anal. Ed. 1933, 5, 221).

"So we have two classes of workers in the analytical field, the determinator and the analyst. The determinators, who by far are the more numerous, may in turn be divided into two general classes: first, the common determinators who follow a method explicitly, without knowledge or concern as to the reactions involved; and the second, the educated determinators who can handle perhaps two variables or who, like determinators of pH, are chiefly interested in group effects and make no effort to go beyond this. The first are laborers. They need not be chemists and they require constant supervision. The second are the white-collar workers who exhibit the usual extremes of workers in this classification. The determinator's salvation lies in the development of truly selective methods of analysis, and his final resting place will be a heaven in which he has a shelf containing ninety-two reagents (this number has changed), one for each element, where No. 13 is the infallible specific for aluminum, No. 26 the sure shot for iron, No. 39 the unfailing relief for yttrium, and so on to uranium. As for the analyst, he is a comparatively rare bird and is often referred to as a disappearing species, like the old family practitioner who does the best he can, guided by theory and experience, of which the most comforting is experience. One of the reasons for the scarcity of analysts is perhaps the nature of the special characteristics, some inherent and some acquired, that he must possess. He must have the inquisitive habit of a detective, for oftentimes the composition of the material under test is absolutely unknown. He must be an expert manufacturer of pure chemicals, for on this ability the success of his gravimetric determinations depends. He must be an efficient dispenser of liquids of which he knows the exact effect. And above all, he must serve a long apprenticeship and keep in constant practice.

As for estimating the accuracy of his work, it can be said that the determinator is usually an optimist in thought and expression, while the analyst is a confirmed pessimist. The determinator reports silicon in glass as 71.61, if not 71.611. The analyst, who knows that he is doing nicely to insure results in the first decimal place, reports 71.6 and thus is honest with himself and deceives no one as to his power."

I think we all are in agreement that standard analyzed samples should be analyzed by analysts and the reported results should not imply an accuracy that is greater than can be obtained by the methods used. How many of you have seen results reported out to the second or third decimal, indicating a precision of 1 part in 8000, when you know the analysis was done with a buret where the reading was 26.15 ml which might just as easily have been 26.17 or 26.13 ml? In certain other reported values, you will also see a precision given which only indicates the precision obtained with the instrument used to make the final reading. Many times this is not stated, however, and the recipient of the data believes they are better than they actually are. In such a case, if the results had been referred to a standard of similar composition carried through all stages of the analysis, one would be on much safer ground as to the accuracy achieved.

The advantage of a standard sample is that it behaves like the unknown material which the chemist intends to analyze; consequently, errors arising from factors such as solubility of precipitates, varying concentrations and so forth will affect both sample and standard alike. If the standard and the unknown sample are analyzed at the same time and under practically identical conditions, and if results are obtained for the standard corresponding to those that are certified, the presumption is strong that the values obtained for the unknown material are equally accurate, whereas wide divergence from the certificate results shows that something is radically wrong.

Let us now consider what is desirable in a standard.

It should be a metal, alloy, compound or ore which is homogeneous and is stable under all normal atmospheric conditions. Since we are dealing with materials which do not always follow this criterion, it is desirable that, if any change does occur—such as the absorption of water—the material can be brought to a certain composition by a standard heat treatment (the usual condition is to heat to constant weight at 105° C). In the case of the very reactive metals with which one deals in the nuclear energy field, one may be forced to use a chemical treatment, such as the pickling of uranium with nitric acid or the electropolishing of plutonium to remove an oxide coating. This is not ideal, but it may be the best that can be done. In certain other instances, materials of a hygroscopic or oxidizable nature are packaged into bottles in a dry box and sealed. Needless to say, this type of sample is not the most reliable. Human fallibility must also be considered in the preparation of standards. There is always the possibility of mislabeling, of mixing samples or using unclean containers in the handling of the materials. All these must be guarded against. The amount of work that goes into the preparation of a reliable standard does not make this a very promising business venture. In addition, the fact that many users want assurance that a financially disinterested, as well as a technically competent, party is concerned has resulted in the greatest reliance being placed on standards certified by the National Bureau of Standards.

In many respects the nuclear energy field is at a disadvantage since there are not enough nationally certified standards available. This has resulted in many companies setting up their own standards. This works very well in running a plant but may cause some problem when inter-company or international transfers occur. It is highly desirable that certified standards acceptable to all parties concerned be available.

URANIUM

Probably the most wanted standards at the present time are those containing varying amounts of U^{235} . Since the accountability of this material is of prime importance, especially with increasing U^{235} content, it is highly desirable that the same standard materials be used by all as reference materials. As far back as 1948, the Atomic Energy Commission constituted the Fissionable Standards Committee and requested it to frame recommendations and to handle the miscellaneous details incidental to providing all standard fissionable materials needed by the Commission laboratories and contractors. This work was taken over in 1956 by the Committee for Uranium Isotopic Standards, whose scope was broadened in 1958 under the Advisory Committee for Standard Reference Materials and Methods of Measurement.

Under these various committees, the Union Carbide Nuclear Company at Oak Ridge prepared two lots of very high U^{235} concentration (99.82 per cent) and one of very high U^{238} concentration (99.9996 per cent). These materials are the Primary Generative Materials and are now at the National Bureau of Standards where they are to be used only for better characterization of their composition, or to prepare Primary Uranium Isotopic Standards at some future date. These Primary Uranium Isotopic Standards will be used only for isotopic certification of relatively large quantities of material which will then be designated as Secondary Generative Materials to be known as Reference Uranium Isotopic Standards. These Reference Uranium Isotopic Standards will be used in the certification of "Uranium Isotopic Standards." A series of such "Uranium Isotopic Standards" is now being distributed by the National Bureau of Standards. They consist of highly purified U_3O_8 of varying U^{235} content.

Table I lists the various isotopic standards now available for distribution.

In the interest of resolving discrepancies, it would be extremely useful if all parties concerned with the determination of U^{235} used these standards of the N.B.S. as

reference materials. Any existing bias could be resolved and all concerned would be referring to the same yardstick and not to their own individual standards.

While the determination of uranium may be done with N.B.S. standard potassium dichromate, 136a, in many instances it is highly desirable to standardize procedures with a uranium-containing material. Certain installations have used uranium metal but the greatest use has been made of a purified U_3O_8 . Material of this type was

Standard No.		Isotopic abi	Indances, %	
Standard No	U ²³⁴	U ²³⁵	U ²³⁶	U ²³⁸
U005	0.0023	0.483	0.0046	99.51
U010	0.0054	0.991	0.0067	98.99
U015	0.009	1.51	0.016	98.47
U020	0.012	2.01	0.016	97.96
U030	0.018	3.01	0.020	96.95
U050	0.028	4.95	0.048	94.98
U100	0.068	10.19	0.038	89.70
U150	0.100	15.30	0.066	84.53
U200	0.125	19.80	0.209	79.86
U350	0.249	34.89	0.170	64.69
U750	0.593	75.12	0.252	24.03
U800	0.660	80.07	0.246	19.02
U850	0.64	84.99	0.37	14.00
U900	0.77	90.10	0.33	8.80
U930	1.08	93.27	0.205	5.44

TABLE I.--- URANIUM ISOTOPIC STANDARDS

originally distributed to the Manhattan Project contractors during the war, and this sample was renewed by the New Brunswick Laboratory under the designation of No. 15. This material has been transferred to the National Bureau of Standards where it is sold as their No. 950. This material, when ignited to 900° C for 1 hr, contains 99.94 per cent U_3O_8 . It was made from UO_3 obtained by thermal decomposition of the nitrate. It is well to point out here that the formation of U_3O_8 , or of a material corresponding to this analysis, is very temperature-sensitive. If we consider that heating in a furnace is usually practiced, followed by removing and cooling in a desiccator, then 850–900° C appears to be the temperature range necessary. This applies only at sea level. At lower temperatures, there is an excess of oxygen and somewhat above 900° C there is a deficiency of oxygen. The history of the material appears to have some effect; it is quite difficult to obtain complete conversion to U_3O_8 from a nitrate by thermal decomposition. On the other hand, the peroxide or oxalate converts to essentially 100 per cent U_3O_8 quite readily under the above heat treatment.

It is well to point out at this time that the atomic weight of uranium must be questioned. The group at Oak Ridge propose that this be changed to 238.037 for normal uranium instead of 238.07. This question will no doubt be considered at the next international meeting of the Committee on Atomic Weights. It may be inadvisable to change at this time since the question of using C^{12} in place of oxygen as the basis for atomic weights is also under consideration.

As shown in Table II, the New Brunswick Laboratory has a group of naturally occurring minerals which were prepared for use in the chemical analysis of uraniumbearing materials. They have been distributed mainly to plants and private analytical laboratories, but some use of these has been made by prospectors. Sample No. 3A was prepared by ball-milling pitchblende ore with dunite. Admittedly, the way that sample 3A was prepared is not the best way to prepare a standard. For use in chemical analysis where a 5-g sample is usually taken, it has proven satisfactory.

Material hosphate rock	U_3O_8 content, %
hosphate rock	0.020
	0.029
itchblende mixture*	4.3
itchblende ore	53.5
Carnotite	0.18
Carnotite	0.11
	Carnotite

TABLE II.—NATURALLY OCCURRING MINERALS FOR USE IN THE ANALYSIS OF URANIUM-BEARING MATERIALS

 $\boldsymbol{\ast}$ Sample No. 3A was prepared by ball-milling pitchblende ore with dunite.

A series of samples, as shown in Table III, was prepared by the New Brunswick Laboratory for use in calibrating counting apparatus to be used in prospecting work. All samples were analyzed after mixing.

NBL No.	BL No. Material U conter		
73	Pitchblende*	1	
74	Pitchblende*	0.1	
75	Pitchblende*	0.05	
76	Pitchblende*	0.01	

TABLE III.—STANDARDS FOR USE IN THE CALIBRATION OF COUNTING APPARATUS USED IN PROSPECTING

* Pitchblende diluted with dunite.

For the analysis of impurities in uranium, a sample of uranium metal chips has been prepared by the New Brunswick Laboratory. This is shown in Table IV.

Other materials available from the same source are UF_4 (Tables V and Va) and UO_3 (Table VI).

For the spectrographic analysis of impurities in uranium and its compounds, two series of samples were prepared by the New Brunswick Laboratory. One of these, for impurities in U_3O_8 , is given in Table VII.

The U_3O_8 samples 65-1 to 65-5 are synthetic materials prepared from a low-levelimpurity U_3O_8 by adding to it solutions of the impurity elements in volumes just sufficient to wet the oxide. Sample No. 65-5 is the matrix material to which the impurities were added. After drying, the material was ball-milled in a plastic container and then ignited at 900° C. The above samples are suitable for the preparation of spectrochemical working curves for the analysis of uranium and its compounds after conversion to U_3O_8 at 900° C. All concentrations listed are the values obtained by chemical analysis, with the exception of cadmium in 65-2 to 65-5, which values are the concentrations added to the U_3O_8 .

Element	Values, <i>p.p.m.</i>
Aluminum	10
Boron	0.23
Carbon	410
Copper	2
Iron	45
Lead	1
Magnesium	2
Manganese	6
Nickel	45
Nitrogen	40
Potassium	<1
Silicon	50
Silver	<0.1
Sodium	2
Thorium	0.3

TABLE IV.---ANALYZED SAMPLE NO. 16. URANIUM METAL

Note: < = less than.

TABLE V.—ANALYZED SAMPLE NO. 17-A. UF₄. Chemical values on UF₄ basis

75.8 %
74.2 %
0.31 %
2.0 %
97.6 %
20 p.p.m.
16 p.p.m.
3 p.p.m.

Table Va.—Analyzed sample No. 17-A. UF₄. Spectrographic on uranium basis (p.p.m.)

<u>`````</u>	
Cu 4	P <10.0
Fe 26 d	Pb <1
Mg 25	Si <20
Mn <4	V <20
Mo <4	Sn <1
Na 2	Zn <20
Ni <20	
	Fe 26 d Mg 25 Mn <4

Note: a = about

d = densitometer.

Since the rare earths are analyzed on a concentrate, a series was prepared for this purpose, to be used with a certain spectrographic procedure. This series is shown in Table VIII.

Several companies use uranium metal as a main standard. From our experience, this is quite satisfactory providing a pure uranium is available. The New Brunswick Laboratory has used "Dingot" metal from the Mallinckrodt Chemical Works for

U HCl, insoluble NO ₃ H_2O Fe Ni Cu Cd Cr Mo	82.10 % 0.6 % 0.9 % 0.6 % 12 p.p.m. 2 p.p.m. <1 p.p.m. <0.2 p.p.m. 3 p.p.m. <1 p.p.m.
Мо	<1 p.p.m.
Bulk density Tap density	$\begin{array}{c} \textbf{2.6} \pm \textbf{0.2} \text{ g/ml} \\ \textbf{4.0} \pm \textbf{0.2} \text{ g/ml} \end{array}$

TABLE VI.—ANALYZED SAMPLE NO. 18. UO₃

TABLE VII.—ANALYZED SAMPLE NO. 65 (1 TO 5).	U_3O_8 samples for spectroscopic
ANALYSIS	

Element	65–1	65–2	65-3	65–4	655
Boron	2.0	1.0	0.5	0.25	0
Cadmium	1	0.5	0.2	0.1	0
Chromium	45	25	12	8	
Copper	53	23	12	6	about 1
Iron	100	55	28	20	10
Manganese	24	12	6	3	about 0.5
Molybdenum	4	1.5	1.0	0.5	
Nickel	100	48	22	13	6
Phosphorus	200	100	50	20	0
Silicon	125	60	25	17	6
Silver	2.3	1.1	0.6	0.2	
Vanadium	100	50	20	10	1

Values in parts per million (based on metal)

TABLE VIII,—RARE EARTH MIXTURES FOR THE SPECTROCHEMICAL ANALYSIS OF RARE EARTH CONCENTRATES FROM URANIUM

Analyzed	sample	No.	28 - 1	to	28-5	
----------	--------	-----	--------	----	------	--

Element	28-1	28-2	28-3	28-4	28-5
Gadolinium	20	10	4	2	0
Dysprosium	20	10	4	2	0
Erbium	20	10	4	2	0
Samarium	20	10	4	2	0
Europium	20	10	4	2	0

Note: Concentrations are in micrograms per milligram of lanthanum.

this purpose. The oxide coating usually present is readily removed with nitric acid, and it does not re-oxidize at an excessive rate. Tables IX and IXa show the values obtained on this material as well as the analysis for the impurities.

While the above samples are available for distribution, many installations use their own working standards. These may be synthetic samples made from reagent grade chemicals, or on the other hand, they may be production materials suitably prepared as to be homogeneous. The value assigned to this material may be from

analyses referred to primary standards such as potassium dichromate or to one of the available uranium analyzed samples. Some plants prefer to prepare U_3O_8 which they use as a standard. The question of composition is all important in these instances. In most of these companies, standard local certification is relied upon.

Analyst	Uranium, %	Variability
NBL	99.974	$\sigma = 0.0032$
NBS	99.975	$\sigma = 0.0071$
K-25	99,976	±0.009 %

TABLE IX .- ANALYSIS OF DINGOT METAL

Note: Above corrected for 30 p.p.m. Fe present in sample, and using 238.037 for atomic weight of uranium.

TABLE IXa.—I	MPURITIES IN DINGOT METAL
Ag 0.03	Мо —
Al 10	Na <5
As —	Ni <8
B <0.1	P
Bi —	РЬ 1
Ca 10	Sb
Cd —	Si 70
Co 2	Sn
Cr 6	v
Cu 1	Zn —
Fe 30	O 10
Li <1	C 10
Mg 8	N 8
Mn 5	

Note: -, not detected. Results are in parts per million.

For impurity analyses in uranium, many rely on synthetic spectrographic standards made from analytical reagent grade chemicals in a purified base. The question, of course, of homogeneity of the final mixture, of the chemicals used and of the impurities in the purified base must be resolved in all cases.

Proposed standards

Certain installations have indicated that a uranium compound such as U_3O_8 , containing enriched uranium with certified uranium and isotopic content, would be useful.

The use of U^{233} in the analysis of highly radioactive solutions of spent enriched uranium fuel elements indicates that possibly a U^{233} certified as to isotopic content might be useful. This would be used chiefly in mass spectrographic measurements.

In this connection, some interest has also been expressed in standards of known U^{236} content.

PLUTONIUM

Standards for plutonium have been a problem for the past decade or more. Up to the present time, a compound suitable for a standard has not been found. The oxide, PuO_2 , as in the case of uranium, is difficult to obtain. One reference¹ states that it is necessary to ignite the metal to 1200° C when preparing PuO_2 from it. When

the oxide is made from plutonium salts at 870° C in air, it has a higher molecular weight. The authors suggest that the formula ranges from $PuO_{2.00}$ to $PuO_{2.00}$, depending upon starting material and temperature. From the above, it would seem that a usable oxide could be made by heating to 1200° C. Unfortunately, plutonium dioxide is very difficult to dissolve in acids. It becomes less soluble when heated to a higher temperature. That formed at 1200° C is exceptionally difficult to dissolve. In some instances, even fusing in bisulfate does not effect solution. For the above reasons, PuO_2 has not been used to a great extent.

At the present time, most installations use plutonium metal of known purity as a standard, the atom per cent of plutonium-239, 240, 241 and 242 being determined by the mass spectrometer. There are several drawbacks to the use of plutonium metal. It is readily oxidized in a moist atmosphere, forming an oxide coating which, unlike that of uranium, is not readily dissolved by acids. Also, the question of obtaining pure plutonium metal has not been entirely solved. Iron is a common impurity and is an undesirable contaminant because after reduction it will titrate like plutonium.

In spite of the fact that plutonium metal is not entirely satisfactory, the Advisory Committee for Standard Reference Materials and Methods of Measurement have authorized the preparation of a provisional plutonium metal standard which is expected to be available during the coming year. The samples are to be prepared at the Los Alamos Laboratory and will consist of weighed pieces of plutonium metal of 0.5-1 g, which will be individually sealed in glass tubes in an atmosphere of argon. The isotopic composition will be given since this is necessary for the calculation of the atomic weight of the plutonium. It is hoped that the impurities will be less than 100 p.p.m. and that the Pu²³⁹ content will be greater than 96 per cent. Distribution will no doubt be handled by the National Bureau of Standards.

As for preparing a plutonium compound of known composition, little has been realized along these lines in spite of a considerable amount of work on various compounds. The New Brunswick Laboratory has prepared $Pu(SO_4)_2 \cdot 4H_2O$. This compound was prepared by dissolving high-purity plutonium metal in 4 N sulfuric acid and heating to about 130°C until red crystals formed. The sulfate was washed ten times with 95% methanol and then air-dried for 3 days. The analysis is shown in Table X.

	Theoretical, %	Found, %
Pu	47.51	47.61
SO4	38.17	38.19
H ₂ O*	14.32	14.20

TABLE X.—ANALYS	S OF	Pu(SO ₄) ₂ ·4H ₂ O
-----------------	------	--

* By difference.

Because there is a question as to the stability of this compound—especially since it has four molecules of water and since it has been shown that the α -particles from the plutonium will decompose water in an aqueous solution—a series of weights was taken on one sample over a period of about 2 months. Table XI shows the change in weight during this period. A different preparation of 1 g weight shows similar stability.

Date, 1959	Weight of sample, mg	Temperature, °F	Relative humidity, %
June 18	666.31		
19	666.29		
22	666.33		
24	666.33	_	
25	666.33	76	78
26	666.42	84	76
29	666.44	85	80
30	666.43	89	68
July 1	666.40	78	79
2	666.40	79	79
6	666.33	81	68
10	666.36	82	76
22	666.37	86	77
August 3	666.29	78	53
4	666.30	79	64
7	666.36	72	77
10	666.35	_	

TABLE XI.—STABILITY OF PLUTONIUM SULFATE TETRAHYDRATE

At present, while the above work on the sulfate looks promising, it is too early to state that this will be a satisfactory compound to use as a standard.

THORIUM

We now come to a prospective fuel in which the interest is like the appearance of the moon. It waxes and wanes. Several years ago there was considerable interest in thorium, and the New Brunswick Laboratory prepared a sample of thorium metal for impurity analysis as given in Table XII.

Element	Values, p.p.m.
Aluminum	a 20
Boron	a 1
Beryllium	a 150
Calcium	<10
Copper	<5
Iron	140
Magnesium	<10
Manganese	a 2
Nickel	a 10
Lead	a 4
Silicon	a 20
Uranium	a 2
Nitrogen	60
Carbon	230
HCl, insoluble	1%

TABLE XII.—ANALYZED SAMPLE NO. 19. THORIUM METAL

Note: a = about

Spectrographic standards of ThO_2 were also prepared, in the same fashion as with the U_3O_8 , for use in the carrier-distillation method of analysis. These are shown in Table XIII.

NBL	26-1	26–2	26–3	26–4	26–5	26-6
Element	Α	В	с	D	Е	F
Al	500	200	100	50	20	10
Fe	500	200	100	50	20	10
Mo	500	200	100	50	20	10
P	500	200	100	50	20	10
Si	500	200	100	50	20	10
v	500	200	100	50	20	10
Zn	500	200	100	50	20	10
Be	170	70	35	17	7	4
Bi	50	20	10	5	2	1
Со	50	20	10	5	2	1
Cu	50	20	10	5	2	1
Mg	50	20	10	5	2	1
Mn	50	20	10	5	2	1
Ni	50	20	10	5	2	1
Pb	50	20	10	5	2	1
Sn	50	20	10	5	2	1
Ag	5	2	1	0.5	0.2	0.1
B		7*	5*	4*	3*	2*
Cd	5.	2	1	0.5	0.2	0.1

TABLE XIII.—SAMPLE NO. 26-1 TO 26-6. ThO₂ FOR SPECTROSCOPIC ANALYSIS Values are in parts per million (based on metal)

Note: Figures given are amounts added except as asterisked. This material is to be used for visual estimation. Cooperative laboratory results indicated that the data given is within ± 25 per cent of the stated value.

* Determined values.

The present program, in which a breeder reactor using thorium is to be constructed under the Oak Ridge Operations Office, will no doubt result in increased interest in this element and possibly new standards may be desired.

Many years ago a monazite ore was prepared at the New Brunswick Laboratory for chemical analysis. The composition is as shown in Table XIV.

MPLE NO. 7. MONAZITE SAND		
Percentage		
9.65		
0.38		

This sample is about exhausted and consideration is being given to renewing it. A candidate material is on hand at the New Brunswick Laboratory and may be transferred to the National Bureau of Standards for certification.

A series of thorium standards made from monazite by mixing with dunite was prepared many years ago by the National Bureau of Standards. These standards

Sample No.	Thorium, %	Uranium, %
79	1.01	0.04
80	0.101	0.004
81	0.051	0.002
82	0.020	0.0008
83	0.0102	0.0004
84	0.0011	0.00004

were exhausted and a new series was prepared by the New Brunswick Laboratory, as shown in Table XV.

These	standards	have	been	distributed	internationally.	All	were	analyzed
chemically	after mixin	ng. If-	—as ha	as happened-	-a very small san	nple,	in the	milligram
range, is u	sed, it is do	oubtful	whet	her these san	nples are sufficien	tly ho	omoge	neous.

A series of standards of radium in radium-bearing sludge and of radium in pitchblende ore has been prepared under the New Brunswick Laboratory auspices, but since these have been used chiefly in international agreements, and since the

amounts prepared are not very great, they are not available for public distribution.

RADIOACTIVITY STANDARDS

For certain work, the National Bureau of Standards has prepared a series of α , β and γ standards, as shown in Table XVI.

Radiation	Nuclide	Nominal activity
α	Polonium-210	200 d.p.s.
α	Polonium-210	500 d.p.s.
α	Polonium-210	1000 d.p.s.
α	U_3O_8	15 d.p.s.
β (γ)	Sodium-22	10 ⁴ d.p.s./ml
γ (β)	Sodium-22	10 ⁶ d.p.s.
β	Carbon-14	10 ⁸ d.p.s./ml
β	Carbon-14	104 d.p.s./ml
β	Hydrogen-3	10 ⁴ d.p.s./ml
β	Hydrogen-3	10 ⁶ d.p.s./ml
K	Iron-55	10 ⁵ d.p.s./ml
$K(\gamma)$	Zinc-65	10 ⁵ d.p.s./ml
$\gamma(\vec{\beta})$	Mercury-203	10 ⁶ d.p.s./ml
γ (β)	Krypton-85	10 ⁷ d.p.s./gmc

TABLE XVI.—ALPHA	BETA	AND	GAMMA	STANDARDS
------------------	------	-----	-------	-----------

Note: d.p.s. = disintegrations per second.

The Bureau is preparing less of these materials than in the past, as several commercial sources of reference materials are now available. In most instances, the companies interested in nuclear fuels prepare their own reference materials, such as, for example, plutonium deposited on stainless-steel discs by vacuum distillation as an α -standard or by evaporating aliquots of a cesium-137 solution for use as a γ -standard.

Radium standards for use in radon analysis, as well as radium γ -ray standards,

Sample No.	Radium content, g	Volume, ml
4950	10-9	100
4951	10-11	100
4952	Blank Solution	100
4955	0.1	5
4956	0.2	5
4957	0.5	5
4958	1.0	5
4959	2.0	5
4960	5.0	5
4961	10	5
4962	20	5
4963	50	5
4964	100	5

are also available from the National Bureau of Standards. These are shown in Table XVII. TABLE XVII.-RADIUM STANDARDS

As you may have noted, nothing has been said of alloy standards. For certain classified work, a program for the preparation of Zircaloy-uranium standards is in progress. The amounts prepared at present, as well as the classification restriction, will not allow these alloys to be made available for general use. It is hoped that in the not-too-distant future, such alloy standards will be available for sale.

The preparation of alloy standards of all types is difficult chiefly because of the lack of homogeneity of such preparations. For certain work-such as the determination of the U²³⁵ content of fuel elements themselves without destruction of the element—the possibility has been brought up of having a library, so to speak, of fuel elements of known U²³⁵ composition. The types of fuel elements now in use in the A.E.C. complex are so varied, and so many of them are of an experimental nature, that it would be impractical to attempt an A.E.C.-wide program to provide standardized fuel elements. The Argonne National Laboratory has been asked to collect specific data as to costs, estimated number of fuel elements required, and details of evaluation of elements to be placed in such a library. I am sure many here would be pleased to see such a program put into effect.

Up to the present time, no consideration has been given to the preparation of standards for the various types of ceramic fuel elements which have been proposed. No doubt, before very long, it may be necessary to consider ceramic or cermet fuel elements containing the oxides and/or carbides of uranium, plutonium and thorium; many other fuel elements have been proposed, such as ones in which the above compounds may be used in a metallic or graphitic matrix as well as in a matrix in which carbides of titanium, zirconium, tantalum and niobium may be used.

NON-FUEL MATERIALS

In addition to fuel materials, there are many other materials used which may require standards. Some of these, such as steel and other common materials, will not be considered.

In the analysis of hafnium, niobium, zirconium and uranium base alloys, synthetic standards are made by making known additions to a high-purity matrix material. High purity matrix materials for hafnium, niobium and zirconium are usually difficult to obtain. At present, there are no available primary standards for hafnium, niobium or uranium base alloys. The A.S.T.M. is actively engaged in work on zirconium metal and Zircaloy-2 standards, and have just initiated a program for niobium metal.

Zirconium–Zircaloy

At the present time, only for zirconium and Zircaloy-2 have standards which approach primary standard quality been developed.

Table XVIII shows the zirconium standards values kindly submitted by G. W. Goward of the Bettis Division of Westinghouse.

	ZAS-1	ZAS-2	ZAS-3	ZAS-4	ZAS-5	ZAS-6
Al*	42	56	79	125	174	252
В	Not agre	ed upon				
Cr†	124	110	82	83	108	245
Co*	< 0.1	1.6	0.8			<0.
Cu*	16	479	197	28	97	10
Hf	Not agre	ed upon				
Fe†	730	820	540	450	630	1120
Mg†	<15	<15	<15	<15	<15	<15
Mn*	26	27	24	20	26	17
Mo*	<0.1	1.2	0.5			0.
Ni	Not agre	ed upon				
РЬ	Not agre					
Si†	52	58	109	106	82	110
Sn	Not agree	ed upon				
Ti*	10	-	4			13
V	Not agree	ed upon				
W†	312	76	233	290	483	19

TABLE XVIII.-ZIRCONIUM STANDARDS

* = Accepted values.

 $\dagger =$ Provisional values.

Table XIX gives the values for the only Zircaloy-2 standard now available. These zirconium and Zircaloy-2 standards are reserved for contractors working on the Naval Reactor Program.

	TABLE XIX5	^Y 303 ZIRCALOY-2
Sn	1.34%	Hf 70
Fe	0.125%	Pb 90
Cr	0.106%	Mg 12
Ni	0.051%	Mn 26
С	45	Mo <5
Al	55	Si 46
В	0.3	Ti <10
Cd	<0.1	V <5
Co	<2	W 108
Cu	13	
		1

Note: All values in parts per million except where otherwise indicated.

Certain secondary standards for specific elements, such as oxygen, hydrogen, nitrogen, tin, iron, chromium and nickel, are used.

Spectrographic standards for zirconium are made by grinding mixtures of zirconium oxide and the oxides of the elements to be added.

At the present time, the National Bureau of Standards has in preparation two samples of Zircaloy-2 and two samples of reactor-grade zirconium to be used for impurity analysis by spectrographic means. In addition, there is one sample of Zircaloy-2 which has been prepared for chemical analysis. It is expected that this last sample will be available in the not-too-distant future.

Beryllium

In the past, several types of beryllium metal standards were prepared. These materials were of electrolytic flake, pebble and vacuum-cast grade. Since, in the past few years, the interest has been chiefly in vacuum-cast, a series of vacuum-cast beryllium metal standards was prepared under the auspices of the New Brunswick Laboratory. These samples are for use in chemical and spectrographic analysis. The list of these materials with their composition is given in Table XX.

Element	New Brunswick Laboratory standard No. 85	New Brunswick Laboratory standard No. 86	New Brunswick Laboratory standard No. 87	New Brunswick Laboratory standard No. 88
Cu	30	300	210	250
Мо	5	25	32	51
Ni	80	200	210	370
Со	10	30	10	10
Fe	400	1400	1700	2500
Mn	80	120	215	250
Si	580	650	1300	1450
Cr	70	200	250	430
Al	100	1400	1300	1400
Mg	90	60	170	50
В	1.1	>50	1.3	1.0
N	55	70	120	110
С	500	400	500	400
BeO	0.5%	0.5%	0.5%	0.5%
Be	99.0%		99.25%	99.5%

TABLE XX.—TENTATIVE	VALUES	OF	BERYLLIUM	METAL	STANDARDS
(Base	d on che	mic	al analyses)		

Note: Values in parts per million except where noted otherwise.

In addition, a series of beryllium oxide standards for spectrographic work with the carrier-distillation technique was prepared by the New Brunswick Laboratory. They are shown in Table XXI.

Since the lowest value given on some elements, in the beryllium metal samples, is rather high, and since the beryllium metal of commerce is of better grade, it may be necessary to prepare a sample of lower impurity content.

In addition to the above, interest has also been expressed in the preparation of standards of beryllium alloys.

Element	72–1	72–2	72–3	72-4	725
В	5.3	2.3	1.3	0.8	0.3
Fe	2070	1070	570	170	70
Al	a2200	a1150	700	290	180
Mg	500*	200*	100*	50*	a60
Ni	500	200	100	50	<2
Mn	540	200	100	50	<10
Cr	450	180	90	50	4
Cd	6	2*	1.0*	0.5*	<3
Li	5*	2*	1*	0.5*	<0.1
Ca	1000*	500*	200*	100*	a60
Со	20	10	5*	2*	<2
Cu	100	50	20	10	<4
Zn	460	210	115	70	20
Ag	10	5	2	1	<0.1
РЬ	50	20	10*	5*	<4
Si	2120	1180	750	480	230
Мо	50	18	8	5	<2

TABLE XXI.—ANALYZED SAMPLE NO. 72 (1 TO 5). BERYLLIUM OXIDE SAMPLES FOR SPECTROSCOPIC ANALYSIS Values in parts per million (based on metal)

* amount added to blank material 72-5

a, about

<, less than

Boron

Since boron in elemental or carbide form is used in certain fuel elements, it has become increasingly important that isotopic standards for this work be made available. The B^{10} content of naturally occurring boron is still open to question. Values ranging from 18.5 to 20.0 per cent have been reported. In addition, reactorgrade elemental boron of naturally occurring composition appears to give a higher value for B^{10} than is found in naturally occurring sodium tetraborate or boric acid.

It is proposed that three standards be prepared on these materials: (1) a salt or oxide of naturally occurring boron such as the tetraborate or boric acid, (2) a sample of elemental boron of natural origin, (3) a highly enriched elemental boron of about 93 per cent B^{10} content.

OTHER MATERIALS

There appears to be interest in standards of the following materials, listed in Table XXII.

1 2 3 4 5 6 7 8	B ₄ C B ₄ C in a Zircaloy matrix Niobium Niobium-uranium Niobium-zirconium-tin Niobium-uranium-tin-zirconium Molybdenum Hafnium
8	Hafnium

Standards in the nuclear energy program

The above discussion on standards is naturally incomplete. Changes in the fuel-elements program and in materials of construction will no doubt result in needs for new types of standards. The Advisory Committee for Standard Reference Materials and Methods of Measurement would be pleased to receive comments on desirable standards for use in the nuclear energy field. We would appreciate the forwarding of your suggestions to: Dr. Samuel C. T. McDowell, Division of Nuclear Materials Management, U.S. Atomic Energy Commission, Washington 25, D.C., U.S.A.

REFERENCE

¹ J. L. Drummond and G. A. Welch, J. Chem. Soc., 1957, 4781.

THE HIGH-RADIATION-LEVEL ANALYTICAL FACILITY AT THE OAK RIDGE NATIONAL LABORATORY*†

CALVIN E. LAMB

Oak Ridge National Laboratory, Oak Ridge, Tennessee

Summary—Unusual techniques and remotely controlled manipulations are required in carrying out analytical work on highly radioactive materials. In order to isolate highly radioactive materials for analysis and to centralize remotely controlled operations in one area, a concrete structure, called the High-Radiation-Level Analytical Facility (HRLAF), was built at Oak Ridge National Laboratory (O.R.N.L.) in 1955.

This facility is used for the analysis of samples of radioactivity greater than 1 r/hr at contact; the samples are received from the Power Reactor Fuel Reprocessing Pilot Plant as well as from many other sources. It consists of a sample-storage cell, seven work cells, a "cold" preparation area, a decontamination area, a receiving dock and an office. Barytes concrete, in addition to concrete of normal composition, is used in the cell walls to meet different shielding requirements. Zinc bromide solutions are used for shielding in the work-cell windows, and high-density lead glass is used for shielding in the storage-cell window.

The facility is provided with Master Slave Manipulators, analytical instruments designed for use by remote control, and special equipment for transporting samples, for continuously monitoring air-borne and background radioactivity, for disposing of solid and liquid wastes, and for carrying out decontamination procedures.

THE various studies at the O.R.N.L. which are related to isotope production, reactor experiments and reactor fuel processing require analytical chemistry on highly radioactive samples. Improvised shielding of stacked lead bricks and other barricades, in use, proved to be inadequate in preventing excessive exposure to personnel who worked for extended periods of time on the high-level radioactive solutions. Each individual laboratory unit was confronted with the problem of trying to work on the hazardous materials with limited space, limited shielding and limited equipment for remotely controlled operations. Therefore, in order to isolate the highly radioactive samples for analysis and to centralize remotely controlled analytical operations into one area, a concrete structure, called the High-Radiation-Level Analytical Facility (HRLAF) was designed and constructed¹ at O.R.N.L. It was completed in 1955 at a cost of approximately \$500,000. The cost included the purchase of fifteen Master Slave Manipulators.

The purpose of this paper is to familiarize the reader with some of the highlights of the facility, its equipment and some of the operations involved.

The High-Radiation-Level Analytical Facility was built as an annex (Fig. 1) to an existing building already housing a pilot plant and laboratories engaged in chemical processing and analytical work on highly radioactive materials.

RECEIVING SAMPLES

Groups in other buildings must transport their samples in carriers with adequate shielding to protect the personnel handling these containers. Sample carriers are

^{*} Work performed under contract with the U.S. Atomic Energy Commission. † This paper was also presented at the Southeastern Regional Meeting of the American Chemical Society, Richmond, Virginia, 5–7 November 1959. The Society has generously released the paper in order that it may be included in these Proceedings.



FIG. 1.-Radiochemical Processing Pilot Plant and HRLAF.



FIG. 2.-Unloading platform and tunnel cart.

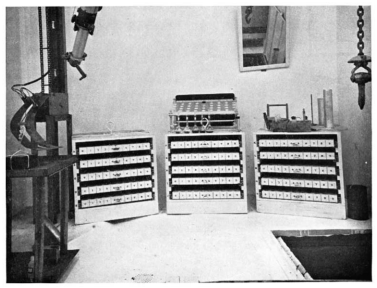


FIG. 3.—Sample storage cell.

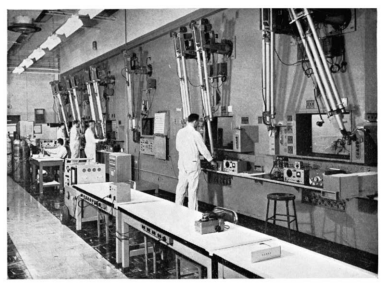


FIG. 6.—Front view of cells.

made in different sizes and shapes to meet varying shielding requirements. The carrier shown in Fig. 2 is a stainless-steel shell filled with lead, and weighs approximately 1000 lb. It is used to transport samples from the Homogeneous Reactor Project. The carrier is designed for a graduated 10 ml centrifuge cone fitted with a screw cap usually containing approximately 5 ml of solution.

In Fig. 2, the carrier is in a position for placement on the target plate of the cart in the tunnel. After loading, the cart is driven electrically 22 ft to the other end of the tunnel. A hydraulic lift then raises the carrier $5\frac{1}{2}$ ft which places it inside the sample storage cell through the opening in the cell floor shown in Fig. 3.

The sample is removed from the carrier in the storage cell and is sent to an adjoining work-cell where the analyses are to be made. It is transferred from one cell to another on a motorized conveyor track. Approximately 10 min are required to send or deliver a sample through the tunnel and to return the carrier to the unloading platform.

Samples received from the Power Reactor Fuel Reprocessing Pilot Plant are loaded onto a chain conveyor in the sampling gallery at the other end of the building. They are carried overhead through lead shielding and down into the sample storage cell where they are unloaded automatically onto a rack. The chain conveyor carries four sets of metal cups (ten cups per set) for holding the sample bottles in transit.

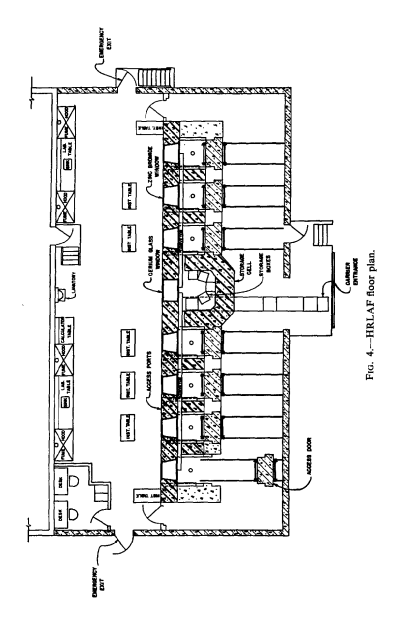
In the storage cell, there are three boxes for the retention of samples after the analytical work is complete. Each box has five drawers wherein 100 standard sample bottles can be stored in each drawer.

BUILDING LAYOUT

On the floor plan of the HRLAF in Fig. 4, the storage cell is centrally located with three work-cells on one side and four on the other. At the present time, there are three analytical chemistry groups using the HRLAF. The spectrochemical laboratory group is using Cell No. 6, the Reactor Analyses Laboratory is using Cells No. 2 and 4, the Pilot Plant Control Laboratory is using Cells No. 1, 3 and 5, and all groups are jointly using Cell No. 7 and the storage cell. The Cold Preparation Zone is the area in front of the cells and is used for the operation of the equipment controlling the work being done inside the cells. It is also used for the preparation of reagents and glassware to be used in the cells. Supporting laboratories which handle low and intermediate level (less than 1 r/hr at contact) radioactive samples are adjacent to the Cold Preparation Zone. The area behind the cells is the Decontamination Zone.

A sectional view of a work-cell is shown in Fig. 5. The walls of the work-cells are 3 ft thick and are made of standard poured concrete. The storage cell walls are made of barytes concrete. Each work-cell has a rear access door made of stacked concrete blocks. The door weighs approximately 9 tons. A steel grill covered with a stainless-steel pan is mounted approximately 3 ft above the actual floor of each work-cell. The pan forms the working surface for the manipulations required in making the analyses. The space beneath the pan is used for the storage of extra equipment. There are four mercury vapor lamps in each work-cell and they are wired so that any two, but only two, can be used at a time. There are also two incandescent work lights in each work-cell.

A view of the front of the cells is shown in Fig. 6. The Master Slave Manipulators



were manufactured by the American Machine and Foundry Company and were designed from the Argonne National Laboratory Model 8. They cost approximately \$7000 a pair. The slaves have been modified with a thumb-activated micro-switch for canting the arm in the cell either forward or backward.

The storage cell window consists of six 6 in. thick sections of lead glass sandwiched together for shielding and cost approximately \$10,000. Zinc bromide solution is used for shielding in the work-cell windows at a cost of approximately \$3000 per window.

The area opposite the front of the cells is shown in Fig. 7. The work benches and

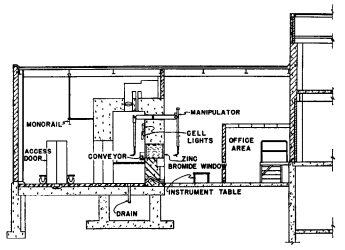


FIG. 5.-Section through work cell.

hoods are used for the preparation of reagents and glassware to be used during the analyses in the work-cells. The non-radioactive work relative to an analysis in the cell is done in this area to increase the cell time available for work involving radiation.

SAFETY DEVICES

Safety features of the facility include a constant air monitor in the Cold Preparation Zone and one in the Decontamination Zone. It continuously monitors and records any radioactivity which has been airborne in its vicinity.

There is a pressure sensitive device in the off-gas system exhausting the cells. It is activated as an alarm if a change of pressure in the system is outside the limits of safety.

There are two probes in the Cold Preparation Zone, two in the Decontamination Zone, and one in each work-cell wired into the console (Fig. 8) of a Remote Area Monitor. It continuously indicates the background radiation at the location of each probe. An individual setting for each probe can be made so that it will act as an audible alarm at any radiation level up to 100 r/hr. A red light flashes on at the console to indicate the position of the probe where the radiation is greater than the preset limit.

Figure 9 lists some of the analyses made in the HRLAF by the Pilot Plant Control Laboratory. Generally, standard "bench-top" methods are used with modification or adaptation to remotely controlled techniques. In many of the methods, the radioactivity level of the material being determined is reduced during the analysis to less than 1 r/hr at contact and the analysis completed in a supporting laboratory.

Most of the equipment failures encountered in the operation of the HRLAF have been the same as those found in a normal laboratory, e.g., a burned-out exciter bulb on a filter photometer, a faulty salt bridge on a coulometer or a polarograph, a broken buret or electrode on a titration unit, etc.

Frequent breakage of the tongs and inner-finger assemblies (Fig. 10) on the Master Slave Manipulators made it necessary to increase the diameter of the tension

Analysis	Method		
U (macro)	Spectrophotometric ammonium thiocyanate		
Th` ĺ	Spectrophotometric thoron		
Zr	Spectrophotometric thoron		
AI	Spectrophotometric ammonium aurintricarboxylate (aluminon)		
P	Spectrophotometric phosphomolybdate		
Hg	Spectrophotometric dithizone		
NH4	Spectrophotometric Nessler's reagent		
Fe	Spectrophotometric o-phenanthroline		
Cr	Spectrophotometric diphenylcarbazide		
Ni	Spectrophotometric dimethylglyoxime		
gross β	Dilution		
gross γ	Dilution		
рН	Direct measurement		
Ċi l	Volumetric titration		
Free H+	Volumetric titration		
Free H+	Potentiometric titration		
Density	Falling drop		
U (micro)	TBP-Amsco extraction—Fluorometric		
U ²³³	Hexone extraction		
Ρυα	Thenoyltrifluoroacetone extraction		
Np	Thenoyltrifluoroacetone extraction		

Fig. 9 —Some methods used by the Pilot Plant Control Laboratory in the HRLAF. Pretreatments include: dilution, fusion, weighing, resin column separation, precipitation.

cables, and to use tension cables and torsion springs made from stainless steel on the inner-finger assembly. The "tru-arcs" used on the tongs were replaced with stainless-steel washers and the pins braded to hold them in place.

A broken tong can be replaced in less than 5 min with the manipulator left in place; however, a broken inner-finger assembly necessitates removal of the manipulator from the cell as shown in Fig. 11. Approximately $1-l_2$ hr is required for two men to remove a manipulator, repair a broken inner-finger assembly and return the slave to its normal position in the cell. The frame used in removing a manipulator from a cell is movable and can be positioned for the removal of any one of the fifteen manipulators. If the proper precautions are not taken, floor contamination often results from particles of radioactivity falling from the inner arm of the manipulator while it is outside the cell for repair.

CELL EQUIPMENT

Cell No. 1 in Fig. 12 is used for non-routine analytical work requiring a temporary set-up of equipment which is disassembled and removed when the work is finished. The equipment which remains in the cell consists of a densimeter mounted on the pan and a sample disposal unit mounted on the wall. The densimeter is used for the measurement of density by the Falling Drop Method.²

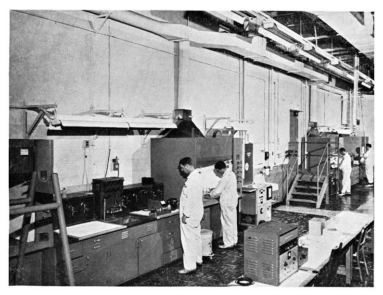


FIG. 7.-Cold Preparation Zone-across from cells.

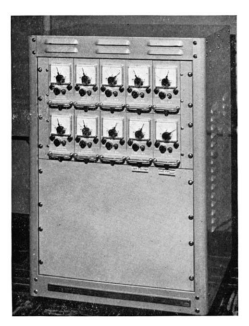


FIG. 8.—Remote area monitoring console.

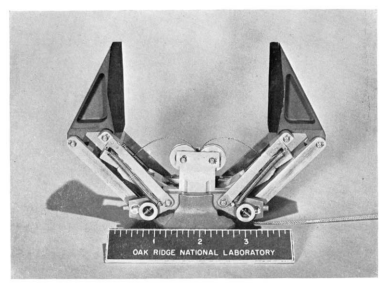


FIG. 10.-Master Slave Manipulator tong and inner-finger assembly.



FIG. 11.-Manipulator being removed from work cell.

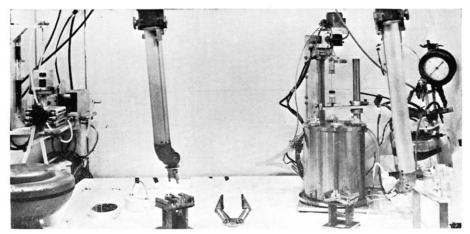


FIG. 12.-Cell No. 1-sample disposal unit and densimeter.

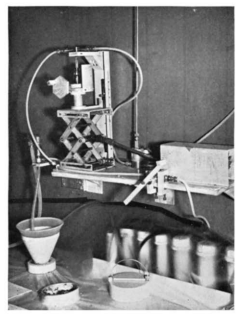


FIG. 13.--Sample disposal unit.



FIG. 14.—Waste materials receptacle.

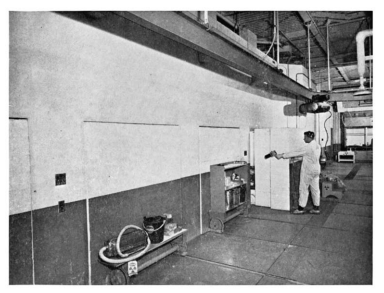


FIG. 15.—Rear of cells-decontamination area.

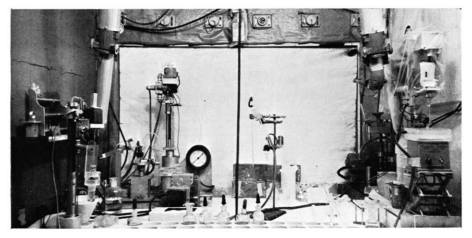


FIG. 16.-Cell No. 2-microelectrodes for pH.

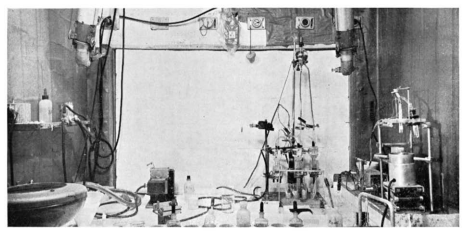


FIG. 17.--Cell No. 4--coulometric titration unit and polarograph cells.

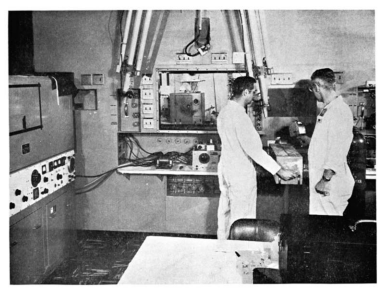


FIG. 18.-Cell No. 6-exterior view showing Jaco-varisource Spectrograph.

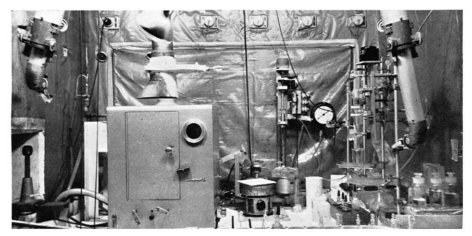


FIG. 19.-Cell No. 6-interior view.

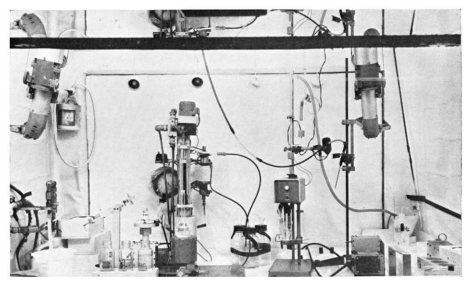


FIG. 20.--Cell No. 3-showing filter photometer and automatic free H⁺ titration unit.

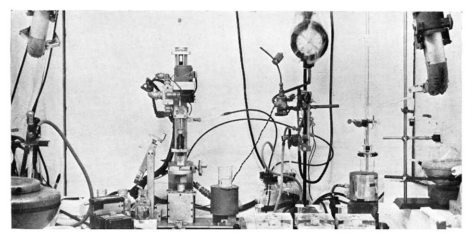


FIG. 21.-Cell No. 5-equipment for liquid-liquid extractions.

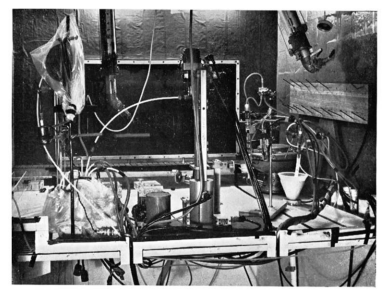


FIG. 22.-Cell No. 5-rear view.

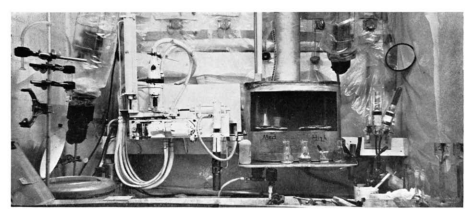


FIG. 23.-Cell No. 7-showing fume hood.



Fig. 24.—Laboratory for receiving, sorting and pretreating samples measuring 1 r/hr at contact.



FIG. 25.—Laboratory for analyzing samples containing intermediate and low-level radioactivity.

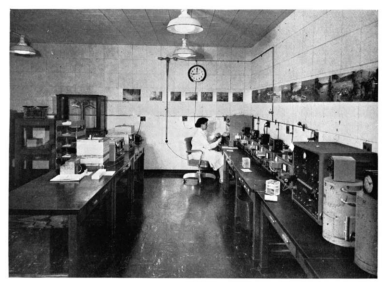


FIG. 26.—Counting room.

The sample disposal unit (Fig. 13) is used for removing the solutions from the sample bottles after the analytical work on them has been completed. The unit is designed for sample bottles which are $2\frac{1}{2}$ in. tall and 1 in. in diameter. Each bottle has a Bakelite top which has been drilled open and sealed with a rubber diaphragm. On disposal, the bottle is clamped onto a motorized lab jack and raised until the two needles have punctured through the rubber diaphragm and extend to within $\frac{1}{8}$ in. of the bottom. One of the needles is connected to a stainless steel water aspirator which draws the sample solution out of the bottle and down the drain to a storage tank. The other needle is connected to a solenoid which allows air or water to be drawn through the bottle. After the bottle is rinsed several times, it is removed and dropped through a chute in the cell pan.

A receptacle (Fig. 14) for catching the empty sample bottles is mounted on a bracket attached to the access door of Cell No. 1. It consists of a cheap metal can with a polyethylene bag fitted inside for a liner. The can is in a lead shielded container. As the cell door is opened into the decontamination area (Fig. 15), the receptacle moves out of the cell and into a position under an overhead hoist. The hoist is on a monorail and is used to carry the receptacle to a vehicle for transfer to a burial ground.

The equipment in Cells No. 2 and 4 is used by the Reactor Analyses Laboratory primarily for analyzing materials from the Homogeneous Reactor. Cell No. 2 (Fig. 16) contains a free-acid titration unit; a temporary pipettor (made from a mohr pipet, rubber tube and syringe for pipetting solutions containing halides), a stainless-steel remote pipettor and a microelectrode unit (for measuring the pH of one drop of solution).

Cell No. 4 (Fig. 17) contains a coulometric titration unit, three dropping mercury polarograph cells, a filter photometer and a centrifuge.

Cell No. 6 is used by the Spectrochemical Laboratory, and a Jaco-Varisource Spectrograph is shown on the left in the exterior view shown in Fig. 18. An access drawer is located at the front of each cell for passing glassware and reagents into the cell. The drawer is also used for removing flasks containing dilutions or pretreated radioactive materials but which must measure less than 1 r/hr before they can be safely handled outside of the cell.

Figure 19 shows the interior of Cell No. 6. The equipment is used for chemical separation and preparation of the sample for mounting onto a graphite porous cup. The prepared material is sparked in the housing in the center of the cell. A duct is located directly over the housing which carries the radioactive gases through a filter and empties them into the exhaust system. The duct can be moved by remote control to a position over a drain in the cell and then flushed with water to remove the collected radioactive materials from the filter.

Cells No. 1, 3 and 5 are used by the Pilot-plant Control Laboratory for analyzing samples from the Power Reactor Fuel Reprocessing Pilot Plant, Volatility Pilot Plant and Fission Products Pilot Plant.

Cell No. 3 (Fig. 20) is equipped with a Beckman Model K automatic titrator, a filter photometer and a stainless-steel remote pipettor. The titrator utilizes an electromagnetic buret which has a fill and delivery valve activated by solenoids. The filter photometer is used for colorimetric analyses. The vacuum system is protected by three bottles used as traps for the collection of waste solutions. The bottles are emptied through a line connected to a stainless-steel water aspirator.

The interior of Cell No. 5 is shown in Fig. 21. The cell is used for making liquidliquid extractions, chloride distillations and colorimetric analyses. By using a magnetic stirrer, an extraction can be made in an enclosed container which prevents the spread of contamination to the cell, cross contamination between extractions, and any change of volume through evaporation during the extraction period. Lucite holders, with holes drilled to support the extraction vessels, are mounted on top of the magnetic stirrers.

A rear view of Cell No. 5 is shown in Fig. 22. The cables attached to the instruments are brought into the cell through conduit embedded in the concrete cell walls and through specially designed plugs in the front wall of the cells.

Polyethylene coverings are used extensively in the cells for protection of the equipment from acids, solvents and radioactive contamination. It is easily and rapidly removed when decontamination of the cells is necessary. It is also inexpensive and can be discarded without the expense of cleaning it.

Cell No. 7 (Fig. 23) is used for analytical work requiring heat. A small fume hood containing two hot plates is used for dissolution and the fuming of samples. The cell is exhausted through the hood. The duct of the hood has a continuous flow of water down the interior walls to dilute or absorb corrosive fumes.

SUPPORTING LABORATORIES

The materials and dilutions removed from the work-cells which measure less than 1 r/hr at contact are sent into a supporting "warm" laboratory such as the one shown in Fig. 24.

Shadow shielding of stacked lead bricks is used for shielding the materials during additional chemical treatment or measurement. The groups submitting radioactive samples also require analytical chemistry on related nonradioactive samples which are analyzed in the supporting laboratories. Open racks are used for storage of the nonradioactive samples after aliquots have been removed for analysis. Samples containing high concentrations of α -emitters, but which are low in Beta activity, and Gamma activity, are analyzed in a glove box.

Another room of the supporting laboratories is shown in Fig. 25. In the Pilotplant Control Laboratory an average of over 4000 analyses per month have been completed for the past year using the work-cells and supporting laboratories. It is staffed with 19 persons, 12 of whom are on rotating shifts operating 24 hr a day, 7 days a week.

Since many of the analyses require measurement of the α -, β - or γ -activity content, a counting room is necessary and is shown in Fig. 26. The equipment consists of a gamma ionization chamber, scintillation gamma counters, alpha proportional counters and end-window beta counters.

CONCLUSION

Since the initial operation of the High-Radiation-Level Analytical Facility in 1955, analytical chemistry has been applied to thousands of samples containing hundreds of curies each. The personnel operating it have averaged bodily radiation exposures of less than a third of the maximum permissible limits as set by the International Commission of Radiological Protection. Acknowledgement—The author gratefully acknowledges his indebtedness to S. A. Reynolds and his Radiochemical Methods Development group, and P. F. Thomason and his Ionic Methods Development group for providing methods for many of the analyses, and also to D. J. Fisher and his Analytical Instrumentation group for providing most of the remote-controlled instruments.

REFERENCES

¹ E. J. Frederick, Nucleonics, 1954, 12, 36.

² R. B. Quincy, Jr., Method Nos. 1 100 and 9 0060 (2 November, 1956) O.R.N.L. Master Analytical Manual, TID-7015, Section 1.

THE DETERMINATION OF IRON IN YTTRIUM*

J. O. HIBBITS, W. F. DAVIS and M. R. MENKE General Electric Company, Aircraft Nuclear Propulsion Department,

Cincinnati, Ohio

Summary—An improved method for determining Fe in Y metal or Y_2O_3 was investigated further in order to determine its applicability to a wider range of samples. The method consists essentially of the extraction of Fe^{III} from a 6 *M* HCl solution into a solution of tri-*n*-octylphosphine oxide (TOPO) in cyclohexane. The Fe is re-extracted into 3 *M* H₂SO₄, an equal volume of HCl is added and the Fe is again extracted with TOPO. An aliquot of the extract is used for development of the 1,10-phenanthroline complex. The method is almost specific for Fe. Only a few elements interfere, even when present in large quantities, and of these elements all but Ga, Te and Si produce a warning turbidity in the colored solution. No interference with the determination of 100 μ g of Fe was caused by 10 mg of Al, Sb^{III}, As^{III}, Ba, Be, Bi, B, Cd, Ca, Ce^{III}, Ce^{IV}, Cr^{III}, Cr^{VI}, Cs, Co, Cu, Dy, Er, Eu^{III}, Gd, Ge, Au, Hf, Ho, In, I, Ir, La, Pb, Li, Lu, Mg, Mn^{II}, Mn^{VII}, Hg^{II}, Nd, Ni, Nb, Pd, P, Pt, K, Pr, Re^{VII}, Rh, Rb, Sm, Sc, Se^{IV}, Na, Sr, Ta, Tb, Tl, Th, Tm, Sn, Ti, V, Yb, Zn; 5 mg of Si, Zr; 1 mg of Mo, W, Ga; 0.1 mg of Te, or U, or 100 mg of Br or F.

PROBABLY the most popular method for determining iron colorimetrically is by means of 1,10-phenanthroline. A number of elements interfere, however, with this method of determination, and a preliminary separation is desirable. Dr. White of the Oak Ridge National Laboratory has published data concerning the extraction of the ferric chloride complex with tri-*n*-octylphosphine oxide (TOPO) which can be used to make a preliminary separation. At the 1958 Southeastern Regional Meeting of the American Chemical Society, he and his co-workers reported on a spectrophotometric method for determining iron after separating it in this manner. A number of elements still interfered, however, such as cadmium, cobalt, chromium VI, copper, nickel, etc. Based on the extraction data published by White, we have developed a method for determining iron which is nearly specific.

The sample is dissolved in hydrochloric acid, and the concentration of this acid is adjusted to 6 M. The iron is oxidized with bromine water, after which it is extracted with 0.01 M TOPO in cyclohexane. The organic extract is shaken with 3 M sulfuric acid, wherein the iron reverts to the aqueous phase. When large amounts of indium, hafnium or zirconium are present, the sulfuric acid solution is washed several times with a solution of 0.1 M TOPO in chloroform, and these wash solutions are discarded. An equal volume of hydrochloric acid is added to the sulfuric acid solution and the iron is again extracted with TOPO. An aliquot of the organic phase is taken for the development of the 1,10-phenanthroline complex. The absorbance of the complex is measured at 510 m μ . Despite the number of manipulations involved, the procedure is relatively rapid and very precise. The method is accurate to ± 3 per cent or 3 p.p.m., whichever is greater, on 1 g samples.

In order to ensure that the method was indeed applicable to the samples to be analyzed, spectrographic analyses of material produced over a 2 year period were

^{*} Work performed under contract with the U.S. Atomic Energy Commission.

tabulated. The method was expected to be accurate when applied to material containing at least twice the maximum amount of each element found by spectrographic analysis to be present in any sample. Furthermore, the analysis was expected to be accurate in the presence of at least 100 p.p.m. of any element normally determined spectrographically. These requirements were met, as can be seen by the data given in Table I.

Added	Fe added, μ g	Fe found, μg	Bias, µg
Y, 1 g	0 100	0 100	
Y, 1 g; Zn, Ca, Mg, Sn ^{II} , Al, Mn ^{II} , V ^V , Pb, Ba, Cd, B, Si, Li, K, Be, Dy, Ho, Tb, Yb, Co, Ni, Mo ^{VI} , Cr ^{VI} , W ^{VI} , Nb, 100 μ g; Cu, 200 μ g; Ta, 400 μ g; Ti ^{IV} , 4 mg; F, 5 mg; Zr, 10 mg.	100 100	97 98	-3 -2

TABLE I.-RECOVERY OF IRON IN THE PRESENCE OF OTHER ELEMENTS

Further investigation of the effects of sixty-eight elements in 10 mg quantities has indicated interference only from silicon, antimony, gallium, molybdenum, tungsten, tellurium and uranium at this level. No interference is caused by 5 mg of silicon, 1 mg of antimony, gallium, molybdenum or tungsten, or 0.1 mg of tellurium or uranium. When interfering amounts of antimony, molybdenum or tungsten are present, the colored solution whose absorbance is to be measured becomes turbid. Only gallium, tellurium and silicon interfere without producing this warning turbidity.

SOME APPLICATIONS OF HIGH-FREQUENCY TITRIMETRY

D. L. MANNING and OSCAR MENIS Oak Ridge National Laboratory, Oak Ridge, Tennessee

Summary—A parallel transmission line oscillator operating at 190 Mc/s was demonstrated to be a useful instrument for the performance of high-frequency titrations. The end point is established by graphically recording either the oscillator grid current or the IR drop, which is produced by passing this current through a load resistor, versus the volume of titrant and extrapolating the two segments of the titration graph to a point of intersection. The instrument has been utilized for the titration of Th, free acid and sulfate in solutions of high ionic strength. The method is of particular value in titrating solutions containing substances which mask the color change of indicators or otherwise interfere with visual titrations. Furthermore, it appears to be adaptable for use in the titration of radioactive materials by remote control.

For the performance of high-frequency titrations, a vessel containing the solution to be titrated is placed in the field of a high-frequency oscillator. Then, as the reaction between titrate and titrant proceeds, changes occur in plate or grid currents or voltages, or in the frequency of the oscillator which result from variations in the composition of the solution. A plot of any one of these quantities against the volume of titrant results in a curve with a break at the end point, similar to that of a conductometric titration graph.

To locate the end point in a high-frequency titration, electrodes, indicators and physical contact of any nature with the solution are not required. The potentiality of the method, therefore, becomes apparent for titrations in which no suitable color indicators or electrodes for end point detection are available. For high-frequency titrimetry to be applicable, however, a difference must exist between the "loading characteristics" of the reactants and the products. This loading is caused by the time lag of dipoles and ionic atmospheres in reorienting themselves in response to the rapid changes in field strength which are caused by the high-frequency oscillations. When the frequency approximates the relaxation time of the dipole or ionic atmosphere, a maximum amount of energy is removed from the oscillator circuit.

A limitation of the high-frequency titration method at the present time is the inability of the commercially available oscillators, most of which operate in the range of 5–50 Mc/s, to respond to changes in fairly concentrated solutions. Debye and Faulkenhagen¹ in their work on the absorption of radio-frequency energy by electrolytes showed that, for absorption of energy to occur, the frequency of the source must increase in proportion to the concentration of the solution. Thus, whereas a 30 Mc/s instrument may respond to changes in approximately $10^{-3}-10^{-2}M$ solutions, an oscillator operating at a frequency of hundreds of megacycles per second is required to detect changes in solutions where the electrolyte concentration is of the order of 0.1*M* or greater. The feasibility of applying high-frequency titrimetry, however, to solutions of high ionic strength was first demonstrated by Blaedel and Malmstadt² who reported on a 350 Mc/s titrimeter and later by Lane³ who presented several

examples of the use of a 250 Mc/s instrument in carrying out conventional titrimetric procedures.

In this work, a study was made of the application of a parallel transmission line oscillator which operates at 190 Mc/s to chemical analyses. This type of oscillator was investigated by Stelzner and Kelley⁴ who made use of an in-line arrangement for the measurement of varying concentrations of alkali in aqueous solutions. As pointed out by Stelzner and Kelley, whenever radio-frequency energy is removed from the circuit, the plate current of the oscillator tube increases and the grid current decreases. Either of the two variables, plate current or grid current, may be measured. In this application, the energy loss which occurs as a consequence of the titration was

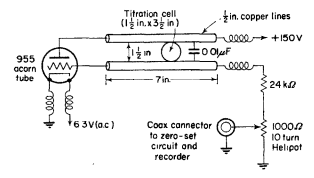


FIG. 1.—Schematic diagram of 190 Mc/s parallel transmission line oscillator.

observed by following the change in grid current. This was accomplished by passing the grid current through a known resistance and recording the voltage drop across the resistance with a 50 mV Brown recorder. The end point is established by extrapolating the two segments of the titration graph to a point of intersection.

In this study, the instrument was applied to the detection of the end point in acidbase titrations, and also the titration of free acid in solutions of uranyl sulfate, sulfate with barium, and thorium with oxalic acid and ethylenediaminetetraacetic acid (EDTA).

Description of oscillator

A simple schematic diagram of the 190 Mc/s parallel transmission line oscillator is shown in Fig. 1. The design is based on a parallel transmission line oscillator in which a type 955 acorn vacuum tube is used in a triode circuit. This type of instrument is described in detail by Stelzner and Kelley⁴. The transmission lines are 8 in. sections of copper tubing which are $\frac{1}{2}$ in. in diameter and spaced $1\frac{1}{2}$ in. between centers. Radio-frequency chokes are utilized to isolate the cathode and filament from r-f ground and to keep stray voltages from the power supply. The calculated Q for this circuit is approximately 4000. A holder for the titration vessel is placed between the parallel transmission lines near the shorted ends of the lines.

Description of titrimeter

A photograph of the high-frequency titrimeter, in which a 190 Mc/s oscillator is used, is shown in Fig. 2. The components are, reading from left to right, 190 Mc/s

oscillator, motor-driven syringe buret, power supply for the oscillator, zero-set circuit and a 50 mV Brown recorder with a 4 sec response.

The oscillator is housed in a 6 in. by 5 in. by 9 in. metal cabinet. The parallel transmission lines run horizontally and mounted underneath them is a plastic well-type holder for the titration cell. The glass cell, which measures $1\frac{1}{2}$ in. in diameter and $3\frac{1}{2}$ in. in height, is inserted between the copper lines through a circular opening in the top of the housing. The volume of solution in the cell for a typical titration is about 30 ml.

The output of the oscillator is measured in terms of the voltage drop produced when the grid current flows through a known resistance. This resistance is composed of a 1000-ohm, 10-turn helipot which is incorporated into the instrument. The sensitivity of the instrument can be changed by means of the variable resistor. In the majority of cases, if the resistor is set at 500 Ω , the voltage change during a titration will be of the order of 30 mV which is adequate for recording purposes.

The delivery of titrant at a constant rate is accomplished through the use of a 900-S single-speed infusion-withdrawal pump equipped with a 2 r.p.m. synchronous motor. This device delivers 1.304 ± 0.004 ml of titrant per minute. For some of the titrations, however, an alternate motor-driven buret was utilized.

The Fluke power supply, Model 407 d.c., furnishes +150 V for the plate and 6.3 V a.c. for the filament of the 955 tube.

The output of the oscillator is connected to a zero-set circuit which is also housed in a small metal cabinet. This type of zero-set circuit is the same as described by Stelzner and Kelley⁴ with the exception that a 6 V "A" battery is used instead of the 1.5 V telephone battery. With this device it is possible to position the pen of the Brown recorder at the start of a titration so that, in recording the titration graph, it will not go off scale, provided the voltage change during the course of the reaction does not exceed the full-scale deflection of the recorder.

Response curves

An idea of the response of the instrument to changes in electrolyte concentrations can be gained from Fig. 3 in which is plotted the response of the oscillator to progressively increasing concentrations of hydrochloric acid and sodium chloride. From a plot of this type, the range of electrolyte concentration over which the instrument is applicable can be ascertained. For the 190 Mc/s oscillator, it can be seen from the steep portion of the curves in Fig. 3 that the optimum response of the instrument to changes in concentration occurs when the electrolyte content of the test solutions falls within the approximate range of 10^{-2} –1*M*. This instrument can, therefore, be used on a wide range of concentrations. Outside of these limits, however, the response decreases markedly.

TITRATIONS PERFORMED

Strong acid–strong base

The response of the instrument to a typical titration of a strong acid with a strong base is illustrated in Fig. 4. The recorded titration curve for this system is very well defined and the end point can be fixed precisely by the break in the curve. When titrating approximately 0.2 mequiv. of a strong acid in a volume of 30 ml with 0.1N NaOH, the end point is reproducible with a coefficient of variation of about 0.6 per cent.

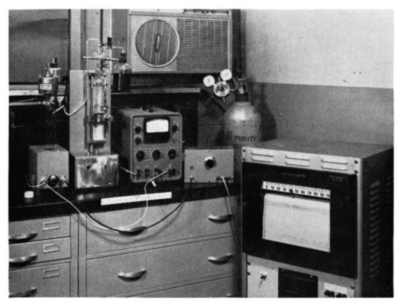


FIG. 2.—Assembled 190 Mc/s high-frequency titrimeter.

The effect of added electrolyte on the response of the oscillator to this system was evaluated by repeating the titrations after making the test solutions 0.1 M and 1 M in sodium chloride, respectively. Essentially the same shape of titration graph was

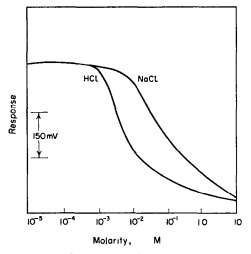


FIG. 3.-Response curves for the 190 Mc/s parallel transmission line oscillator.

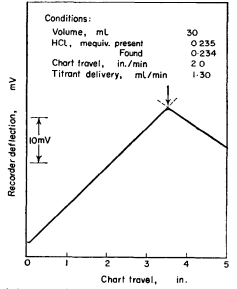


FIG. 4.—High-frequency titration of hydrochloric acid with sodium hydroxide.

obtained in the presence of 0.1M chloride and no difficulty was experienced in locating the end point. When the solution was 1M in sodium chloride, however, the slopes of the two segments of the titration curve were appreciably decreased and the break in the graph indicating the end point could not be located with any degree of certainty. Thus, from a practical point of view, when the 190 Mc/s oscillator is used in titrating a strong acid with a strong base, the concentration of electrolyte in the test portion should be less than 1M.

Determination of free acid in the presence of uranium

The applicability of the instrument to the determination of free acid in the presence of hydrolyzable substances, in this case uranyl sulfate, is illustrated by the data presented in Table I and also graphically in Fig. 5.

The first end point corresponds to the neutralization of the free acid while the second end point is observed when the hydrolysis of the uranyl ion is complete. The volume of titrant consumed between the two end points is, qualitatively, or roughly quantitatively, a measure of the uranium present. Rodden⁵ has also pointed

Uranium,	Chart to	Free ac	id, <i>mequiv</i> .	Deserve
mg	end point, in.	Present	Found	Percent
None	2.30	0.210	0.209	
	2.32		0.210	100
	2.10	0.187	0.190	101
	2.10		0.190	101
14	2.00		0.181	97
	2.05		0.186	100
35	2.00		0.181	97
	2.10		0.190	101
165	2.15		0.195	104
	2.10		0.190	101
	2.15		0.195	104
	2.10		0.190	101
350	2.10		0.190	101
	2.10		0.190	101

TABLE I.-HIGH-FREQUENCY TITRATION OF FREE ACID. EFFECT OF URANIUM Conditions: NaOH, N

0.100

out the feasibility of estimating uranium by a sodium hydroxide titration; but he further emphasized the necessity of utilizing empirical factors inasmuch as the reaction is not a truly stoichiometric one. Since the preliminary results indicated that the titration serves only as an approximation of the uranium present, further tests for the purpose of determining uranium by this technique were suspended. Relatively large amounts of uranium can be present without introducing a significant error in the free acid measurement. The coefficient of variation is of the order of 2 per cent. The uranium does not have to be removed or complexed in any way prior to the acid titration.

Titration of sulfate with barium chloride

The results from the high-frequency titration of sulfate with barium chloride are presented in Table II and in the form of typical titration curves in Fig. 6.

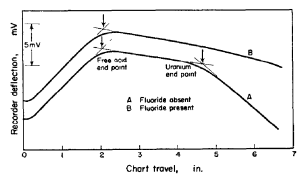


FIG. 5.-High-frequency titration of free acid and uranium with sodium hydroxide.

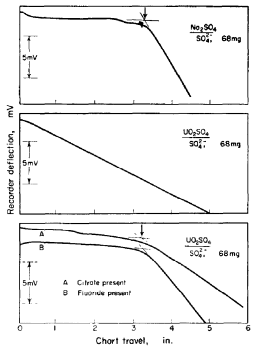


FIG. 6.-High-frequency titration of sulfate with barium chloride.

No difficulty was encountered in titrating aqueous solutions of Na_2SO_4 . Welldefined titration curves were recorded as shown by the upper graph in Fig. 6. The coefficient of variation for a limited amount of data is of the order of 1 per cent.

When an aqueous solution of uranyl sulfate was titrated with barium chloride, the resulting high-frequency titration curve (middle graph, Fig. 6) essentially consisted of one continuous straight line instead of the customary two segments of different slopes. The same type of curve was obtained when attempts were made to titrate sulfate as Na_2SO_4 in the presence of uranyl nitrate. In this situation, where the reaction is

$$UO_2SO_4 + BaCl_2 \rightarrow UO_2Cl_2 + BaSO_4$$

the change in the response of the oscillator at the start of the titration is undoubtedly due to the difference between the "loading characteristics" of UO_2SO_4 and UO_2Cl_2 . Beyond the end point no change in slope is observed as excess $BaCl_2$ is added. The response of the oscillator consists of one continuous straight line.

Substance	Madium	Chart to end		Sulfate	
Substance	Medium	point, <i>in</i> .	Present, mg	For	und
			·	mg	%
Na ₂ SO ₄	Aqueous	3.8	82.3	81.6	99
	1	2.5	54	54.7	101
		2.45	54	53.8	100
	I	İ	1	Aver	age 100
			Coef	ficient of variat	ion 1
UO₂SO₄	Aqueous	No end point	68.4		
	Sodium citrate,	3.2	68.4	68.5	100
	0.03 <i>M</i>	2.10	45.6	46	101
	NaF, 0.03 <i>M</i>	2.16	45.6	47	103
	NaF, 0.1 <i>M</i>	2.12	45.6	46.2	102
		3.17	68.4	69.4	101
]	2.10	45.6	46.0	101
		2.10	45.6	46.0	101
		2.45	54	53.8	100
		2.45	54	53.8	100
l	-	1	Ļ	Avera	ige 101

TABLE II.—HIGH-FREQUENCY TITRATION OF SULFAT	E WITH BARIUM CHLORIDE
Conditions: Barium chloride, M	0.251
Delivery, ml/min	0.906
Volume, ml	30

It is possible, however, to titrate the sulfate in a uranyl sulfate solution if a complexing agent such as sodium citrate or fluoride is added. When this is done, the uranyl ion is masked and remains inactivated throughout the titration of the sulfate which is now present as Na_2SO_4 . In fact, the titration graphs (curves A and B, Fig. 6) resemble those obtained with solutions of sodium sulfate. Sodium fluoride was found to be preferable to the citrate as a masking agent because, with the fluoride, the titration curves are better defined. Also, the concentration of the fluoride is less critical than citrate. Whereas 50 mg of sulfate as uranyl sulfate can be titrated satisfactorily in sodium fluoride solutions ranging from 0.03 to 0.1*M* (optimum concentration, 0.1*M*), when citrate is used, the citrate concentration must be carefully

36

controlled within the range of 0.03-0.05M, to attain satisfactory results. When masking agents are used, sulfate can be determined in uranyl sulfate solutions by this method with a coefficient of variation of 2 per cent or less.

Titration of thorium with oxalic acid

High-frequency titrations of thorium with oxalic acid were carried out both manually and automatically. A representative titration curve which was recorded manually is presented in Fig. 7. The oscillator response was measured directly as a voltage drop across a 500 Ω resistor in the grid circuit after the addition of each increment of titrant. It is of interest to note when thorium is reacted with oxalic acid, in

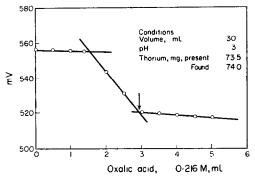


FIG. 7.-Titration of thorium with oxalic acid.

an unbuffered solution having a pH of approximately 3, two breaks are obtained in the high-frequency titration graph as illustrated in Fig. 7. It appears probable from these observations that the formation of the insoluble thorium oxalate occurs stepwise, according to the reactions:

$$Th^{4+} + H_2C_2O_4 \rightarrow Th(C_2O_4)^{2+} + 2H^+ \qquad (first break)$$

$$Th(C_2O_4)^{2+} + H_2C_2O_4 \rightarrow Th(C_2O_4)_2 + 2H^+.$$
 (first break)

The second break is taken as the end point because the first break is not reproducible enough for quantitative purposes. It was also noted that, in order to obtain two breaks, the pH of the test solution must be at least 3 at the beginning of the titration. When the titrations were performed in more acidic solutions, only one break corresponding to the second break of Fig. 7 was obtained, as illustrated graphically by curve B of the titration graphs in Fig. 8. Other than obscuring the first break in the titration curve, the pH of the solutions to be titrated is not a critical factor. The titrations can be successfully carried out at any pH within the range of 1.5 to 3.5. When determining thorium in amounts of the order of 50–100 mg, the coefficient of variation is approximately 2 per cent.

Titration of thorium with ethylenediaminetetraacetic acid (EDTA)

The applicability of the high-frequency titrimeter to the complexometric titration of thorium with EDTA was studied. In view of recent interest in the determination of thorium in the presence of moderate quantities of uranium, aluminum and corrosion products of stainless steel, the effect of these substances (with the exception of nickel and chromium) on the high-frequency titration of thorium was ascertained. The results of these tests are presented in Table III and graphically in the form of titration curves in Figs. 9, 10 and 11. Finally, in view of the fact that conventional EDTA titrations of thorium with indicators are usually performed in acetate-buffered solutions, it was of interest to evaluate the effect of acetate, i.e. moderately buffered solutions, on the high-frequency titrations. The results of this study are shown graphically in Fig. 12.

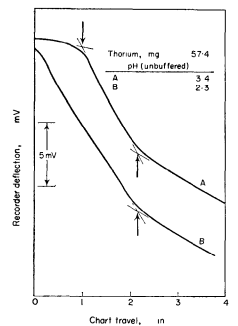


FIG. 8 .--- High-frequency titration of thorium with oxalic acid. Conditions: oxalic acid, M, 0.236; delivery, 0.98 ml/min; chart travel, 2.0 in./min.

TABLE III.—TITRATION O Conditions: EDTA, <i>I</i> Delivery, Chart tra pH (unbu	M 0.1 , ml/min 1.3 , wel, in./min 2.0	44 10 10
Inches of chart to	Thorium,	mg
end point	Present	Found
2.66	58	58.2
2.60		57.0
2.62		57.4
2.64		57.8
2.67		58.5
2.61		57.2
2.60		57.0
2.62		57.4
	Average	e 57.6

It is evident from the results listed in Table III that the 190 Mc/s oscillator provides an accurate means of locating the end point when thorium is titrated with EDTA. A quantitative titration is realized with a coefficient of variation of 1 per cent or less. This compares favorably with visual titrimetric methods for thorium. The governing factor in obtaining reproducible results is the location of the end points from an extrapolation of the two segments of the graph to a point of intersection. In many

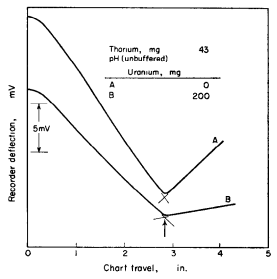


FIG. 9.—Effect of uranium on the high-frequency titration of thorium with EDTA. Conditions: EDTA, M, 0.0712; delivery 1.3 ml/min; chart travel, 2.0 in./min.

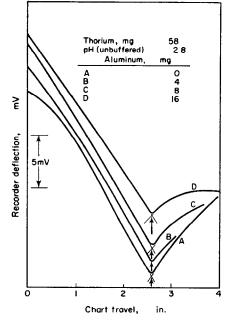


FIG. 10.—Effect of aluminum on the high-frequency titration of thorium with EDTA. Conditions: EDTA, M, 0.144; delivery, 1.3 ml./min; chart travel, 2.0 in./min.

instances, however, a very sharp break in the curve is obtained, and an extrapolation is unnecessary.

As is indicated by Figs. 9, 10 and 11, relatively large quantities of uranium and moderate quantities of aluminum and iron^{II} can be tolerated without introducing a significant error in the thorium measurement. Iron^{III} cannot be tolerated as this substance reacts with EDTA along with the thorium. Larger quantities of these interfering substances were not tested since the upper limit already explored far

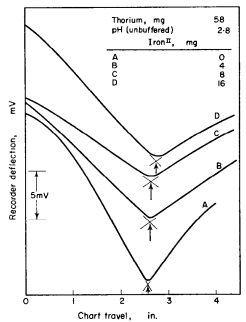


FIG. 11.—Titration of thorium with EDTA. Effect of iron^{II}. Conditions: EDTA, M, 0.148; NH₂OH.HCl, mg 200; delivery, 1.3 ml/min; chart travel, 2.0 in./min.

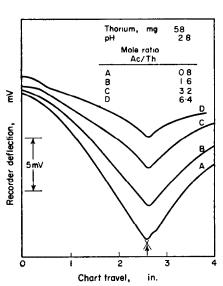


FIG. 12.—Titration of thorium with EDTA. Effect of acetate. Conditions: EDTA, M, 0.144; delivery, 1.3 ml/min; chart travel, 2.0 in./min.

exceeded the amount that would be present in a typical test portion to be analyzed for thorium. Nickel and chromium remain yet to be studied; however, it is expected that moderate quantities of these elements will likewise not interfere.

The effect of acetate on the high-frequency titration curves is illustrated in Fig. 12. In general, as long as the mole ratio of acetate to thorium does not exceed 3, it makes no difference whether the solution titrated is moderately buffered or not. Nothing is gained, however, by the addition of acetate and, in fact, the titration curves recorded from unbuffered solutions are generally better defined. In the presence of large quantities of acetate, the curve becomes poorly defined although it is still possible to locate the end point.

It is to be concluded from these studies that the high-frequency titrimetric procedure appears to be a relatively precise method for determining thorium under conditions where the only other alternatives would be to use a lengthy gravimetric procedure, a potentiometric titration using the Hg–EDTA electrode, or a less precise colorimetric method. The time required for a high-frequency titration, including aliquoting and pH adjustment, is less than 10 min.

CONCLUSION

It is concluded that the parallel transmission line oscillator is a useful instrument that can be utilized to follow chemical changes in rather complex solutions. The simplicity of the oscillator circuit combined with good stability qualities make it attractive as a high-frequency titration instrument.

REFERENCES

- ¹ H. Falkenhagen, *Electrolytes*, Oxford University Press, London, 1934, p. 213.
- ² W. J. Blaedel and H. V. Malmstadt, Anal. Chem., 1950, 22, 1413.
- ⁸ E. S. Lane, Analyst, 1957, 82, 406.
- ⁴ R. W. Stelzner and M. T. Kelley, *Radio-frequency Oscillators for Determination of Alkali*, U.S. Atomic Energy Commission, ORNL 1742, 1955.
- ⁵ C. J. Rodden, Analytical Chemistry of the Manhattan Project, McGraw-Hill, 1950, p. 74.
- ⁶ L. P. Pepkowitz, W. W. Sabol and D. Dutina, Anal. Chem., 1952, 24, 1956.

X-RAY ABSORPTION EDGE ANALYSIS*

H. W. DUNN

Oak Ridge National Laboratory, Oak Ridge, Tennessee

Summary—Very few industrial laboratories are using the X-ray absorption edge method of analysis at the present time. Without a doubt, it will be used much more in the future, for it has several advantages over other methods. Probably the greatest advantage is the relative freedom from interferences, both direct and matrix. Most elements will have little or no interference, but a few elements will have a direct interference from the absorption edge of one or two elements. Secondly, a wide range of elements can be determined; it should be possible to determine all elements from chromium on, in the table of atomic numbers. In addition, the precision and the accuracy of the method are fairly good. For Zr a relative standard deviation of 1 per cent has been obtained for concentrations down to 1 mg/ml, 3 per cent for 0.2 mg/ml, and 50 per cent for 0.05 mg/ml. Means to improve the precision and accuracy of the method are being planned. The method is also fairly fast. If the sample is in solution, the determination of each element requires from 15 min to 1 hr. Other advantages are that any number of elements can be determined in the same solution, the solution is not consumed or contaminated during the analysis, and only a small amount of solution (0.2–2 ml) is required.

Disadvantages of the X-ray absorption edge method are that elements of low atomic number cannot be determined by this method, and the equipment required is expensive. A satisfactory instrument will cost \$15,000 to \$20,000.

A NUMBER of chemical determinations are difficult or slow by ordinary methods. It has been shown that some of these determinations can be made quickly and accurately by the use of X-ray fluorescence. When X-ray fluorescence was tried at the Oak Ridge National Laboratory, however, it was found to be unsatisfactory for the accurate quantitative analysis of the variety of samples we receive. By the time we could work out a good set of working curves and correction factors, the people had completed the experiment and had gone to another phase of the project and were submitting a different type of sample. We just could not keep up with the changes that were taking place at the laboratory.

While in another laboratory, the author had done some work on the analysis of uranium by the X-ray absorption edge method. This method was found to be accurate and essentially free from interferences. In fact, it has been used satisfactorily for routine analyses in a service laboratory for some time.

The X-ray absorption edge method uses white X-ray radiation. At a certain wavelength in the white X-ray radiation (the excitation potential of the element to be determined), there is an abrupt change in the absorption coefficient of that element. This abrupt change is referred to as the absorption edge. The position of the absorption edge is different for each element and the magnitude of the break is a function of the amount of the element present. By taking readings on both sides of the absorption edge (Fig. 1), it is possible to make a quantitative analysis for the element. Such analyses are usually performed on solutions.

There are three L absorption edges and also one K absorption edge for each

^{*} Work performed under contract with the U.S. Atomic Energy Commission.

element. In a few cases the absorption edge of one element will interfere with an edge of another element, but most elements will have no interferences. The direct overlapping of absorption edges is the only type of interference that will give trouble with the X-ray absorption edge method.

This method is applicable to a wide range of elements. It should be possible to determine all elements from chromium up, in the table of atomic numbers. In addition, the precision and the accuracy are fairly good. If the sample is in solution, the determination of each element requires from 15 min to 1 hr. Other advantages



FIG .1.

are that any number of elements can be determined in the same solution, the solution is not consumed or contaminated during analysis, and only a small amount (0.2-2 ml) is required.

Disadvantages of the X-ray absorption edge method are that the elements of low atomic number cannot be determined by this method, and the equipment required is expensive. A satisfactory instrument will cost \$15,000 to \$20,000. Since the equipment is fairly complicated and must be kept in excellent condition, the services of a competent electronic man will be required occasionally.

The equipment required is the X-ray diffractometer with a single crystal replacing the powder sample and the absorption cell mounted between the collimator and the crystal. The X-ray source must be well regulated. Due to the high counting rates (up to 20,000 counts/sec) a low resolving time detection system is needed. In order to correct for resolving time losses, it is necessary to have a constant potential filter. The absorption cell must be mounted in a very reproducible position, perpendicular to the beam.

A consideration of the advantages and disadvantages of the X-ray absorption edge method seemed to indicate that this method would be quite satisfactory for analysis at the Oak Ridge National Laboratory. We had a GE XRD-5 with a constant potential filter and special detection equipment. The equipment seemed adequate, so we decided to try this method.

The basic equation for X-ray absorption is

$$I_{\lambda} = I_{0\lambda} e^{-\sum_{i} \mu_{m_{i}} lC_{i}}$$

where I_{λ} is the observed intensity, $I_{0\lambda}$ is the incident intensity, e is the base of natural logarithms, μ_{m_i} is the absorption coefficient of an element at one wavelength, l is the cell length in centimeters, and c is the concentration in grams of element per gram of sample. The total absorption is the summation of all the individual absorptions. To obtain the absorption coefficient for an element, the equation may be written in the form

$$\mu_m = \frac{2.303}{lC} \log_{10} \frac{T_\lambda}{T_{b1}}$$

where T_{λ} is the time for a specified number of counts at wavelength λ , and $T_{b_{\lambda}}$ is the

time for the same number of counts at the same wavelength for a blank containing everything except the element for which the absorption coefficient is being obtained. By obtaining the absorption coefficient, μ_m , at two wavelengths, λ_1 and λ_2 , the equation may be written in the form

$$C = \frac{2.303}{\Delta \mu_m \ell} \log_{10} \frac{T_{\lambda_1}}{T_{\lambda_2}} \frac{T_{b\lambda_2}}{T_{b\lambda_1}}$$

where T_{λ_1} is the time for a specified number of counts on the short wavelength side of the absorption edge, and T_{λ_2} is the time for the same number of counts on the long wavelength side of the absorption edge. $T_{b\lambda_1}$ is the time for a specified number of counts for a blank on the short wavelength side of the absorption edge, and $T_{b\lambda_2}$ is the time for the same number of counts for the blank on the long wavelength side of the absorption edge. $\Delta \mu_m$ is the difference in the absorption coefficient at the two wavelengths.

By taking the time for a specified number of counts at λ_1 and λ_2 with a series of Al foils, it has been found that the slope of the white radiation curve changes with the amount of absorbing material in the beam. It has also been found, using other types of foils, that the slope at a given wavelength (i.e. $T_{b\lambda_1}/T_{b\lambda_2}$) is almost independent of the atomic number of the absorbing material. This indicates that the $T_{b\lambda_2}/T_{b\lambda_1}$ or blank ratio (BR) could be read from a curve and a blank would not have to be determined for each sample. The curve was set up by plotting the time for a specified number of counts at λ_1 against the blank ratio. The data for this curve were determined by back calculating to the required blank ratio from a set of standard samples.

It has also been found that there is a change in the slope of the white radiation curve near the absorption edge. This may account for the fact that the concentration of the element has a slight effect on the calculated concentration of that element. This effect may be corrected for by applying a concentration correction,

$$CK \frac{T_{\lambda_1}}{T_{\lambda_2}},$$

where CK is the concentration constant.

The equation now takes the form

$$C = \frac{2.303}{\Delta \mu_m l} \log_{10} \frac{T_{\lambda_1}}{T_{\lambda_2}} BR - CK \frac{T_{\lambda_1}}{T_{\lambda_2}}$$

where BR is the blank ratio and CK is the concentration constant.

The solutions used for standards for the determination of Zr were made from pure Zr metal. The calculated concentrations were obtained using the above equation. The percentage error, both observed and theoretical, was based on a single determination.

Recovery of zirconium was tested for both pure and contaminated Zr solutions. The results are shown in Table I.

It is evident that recovery of zirconium was satisfactory over a wide range of concentration, regardless of the presence of impurities.

It will be noted from the data presented in Table I that the observed error was well below the expected limit of error in a few cases. Of course, this could not be done consistently. The theoretical limit of error is based on statistical considerations only and assumes that all other errors are quite small.

With X-ray absorption edge analysis, it makes no difference whether the impurity is in the solution in the cell or in the beam outside the cell. The components of stainless steel, Y, and U were in the solution in the cell and the Al was outside the cell. The elements selected as impurities were chosen because they have absorption

Actual Zr concentration, mg/ml	Actual impurity concentration, mg/ml	Observed Zr concentration, mg/ml	Error observed, %	Theoretical limit of error, %
25.04	0.00	25.04	<0.1	0.1
5,007	0.00	5.006	<0.1	0.2
1.001	0.00	1.009	0.8	0.9
0.200	0.00	0.203	1.5	4.6
2.003	4.12 NiMo	2.011	0.4	0.8
	Alloy*			
2.003	1.00 Y	2.008	0.2	0.8
2.003	3.01 U	2.017	0.7	0.8
0.200	~20.00 Al	0.201	0.5	8.0

TABLE I.--RECOVERY OF Zr IN PURE AND CONTAMINATED Zr SOLUTIONS

* Composition: 72 per cent Ni, 5 per cent Cr, 20 per cent Mo, 3 per cent Fe.

5

edges near the Zr K absorption edge and were the most likely elements to cause trouble. Since all the data were within the expected limit of error, the impurities evidently cause little or no interference.

The X-ray absorption edge method has been used for several months for the analysis of Zr in a variety of samples. Zr-containing alloys, solutions from solubility studies and precipitates and residues from chemical reactions have been analyzed. Unknown to us, those submitting samples have put known standards in with their samples. We have consistently checked the standards within our limits of error. Also, the material balances obtained using our data have been good.

For the varied type of work done at the Oak Ridge National Laboratory, the X-ray absorption edge method looks very promising.

I would like to express my appreciation to Mr. Cyrus Feldman for his help and encouragement on this project.

X-RAY FLUORESCENCE SPECTROGRAPHIC DETERMINATION OF IMPURITIES AND ALLOYING ELEMENTS IN TANTALUM CONTAINER MATERIALS*

E. A. HAKKILA and G. R. WATERBURY University of California, Los Alamos Scientific Laboratory, Los Alamos, New Mexico

Summary—X-ray fluorescence methods have been used for determining molybdenum, niobium, thorium, tungsten, yttrium and zirconium as impurities or alloying constituents in tantalum and binary tantalum alloys being investigated as container materials for molten plutonium fuels. Various solvents were tested for dissolving tantalum samples, and hydrofluoric acid was found to be the most suitable. Samples are dissolved in hydrofluoric acid and diluted to a known volume; then the solution is analyzed by means of an X-ray tube with a tungsten target. X-ray tube operating conditions and counting times were selected to give good precision in a minimum time. Effects of impurities and variations in X-ray tube voltage and current, scintillation counter voltage, solvent composition, dilution volume and count rate were determined.

Molybdenum, niobium and zirconium as impurities in tantalum within the concentration range of 25–2000 p.p.m. are determined by means of a line-to-background intensity-ratio method with a precision ranging from 17 p.p.m. at the 25–100 p.p.m. level to 43 p.p.m. at the 2000 p.p.m. level. From 0.5 to 10 per cent of tungsten in tantalum-tungsten alloys is determined with an absolute standard deviation of 0.09 per cent by comparing the intensity ratio of the tungsten L_{α_1} to tungsten $L_{\alpha Compton}$ for the sample with ratios obtained for known standards. Thorium or yttrium in binary alloys is determined in the 0.025 to 10 per cent concentration range, using an internal standard procedure. Thorium is added as an internal standard in the determination of yttrium, and yttrium is added as an internal standard in the determination of thorium. Following sample dissolution, the ThF₄ and YF₃ precipitates are separated from the tantalum, dissolved in H₃SO₄ and diluted to volume, and the intensity ratio of the thorium to yttrium lines is measured. Relative standard deviations of 7 and 2 per cent, respectively, were obtained in the 0.025–0.1 and 0.1–10 per cent concentration ranges for yttrium, and 12 and 3 per cent for thorium in the 0.025–0.1 and 0.1–10 per cent concentration ranges.

INTRODUCTION

HIGH-PURITY tantalum metal and various tantalum alloys are being investigated as container materials for molten plutonium reactor fuels. The concentration of numerous impurities as well as alloying constituents in a sample must be accurately known in order to evaluate its applicability under reactor conditions. Molybdenum, niobium and zirconium are expected to occur in high-purity tantalum metal as impurities in amounts up to 2000 p.p.m. Tungsten in the concentration range of 0.5–10 per cent, and thorium and yttrium in the range of 0.025 to 10 per cent were considered as possible alloying constituents with tantalum. Because X-ray fluorescence spectroscopy offers a rapid method of analysis for most metallic elements, its application to the determination of these elements in tantalum was investigated.

APPARATUS

A North American Philips three-position-head X-ray spectrograph with a Philips FA60 tungsten-target X-ray tube was used in this work. The optical arrangement

* Work performed under the auspices of the U.S. Atomic Energy Commission.

consisted of a $\frac{1}{8}$ in. source collimator, lithium fluoride crystal with a 2d spacing of 4.0276 Å, Philips Type 52189 receiving collimator, and a sodium iodide-thallium iodide scintillation counter. Voltage to the counter was supplied from a Baird Atomic Model 312A Super Stable High Voltage Power Supply. In the determination of tungsten, an Atomic Instrument Company Model 510 single-channel pulse-height analyzer was also used. The inside of the stainless-steel sample cell provided with the instrument was coated with paraffin to prevent dissolution by the concentrated hydrofluoric acid used in the determinations.

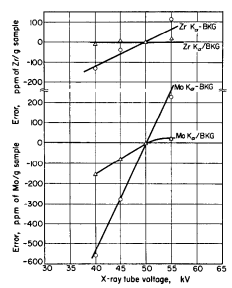


FIG. 1.—Effect of X-ray tube voltage on the X-ray spectrographic determination of molybdenum and zirconium in tantalum.

SAMPLE PREPARATION

Various methods were investigated for obtaining either solid or liquid solutions of the tantalum samples. Fusion with borax, pyrosulfate or sodium hydroxide proved time consuming and unsatisfactory. Tantalum is slowly dissolved by concentrated hydrofluoric acid, and the reaction can be hastened with a drop or two of nitric acid. Because 2 or 3 ml of 35% hydrofluoric acid will hold 1 g of tantalum in solution, this acid was selected as the solvent. Weighed samples are dissolved overnight in 48% hydrofluoric acid, evaporated to 2 or 3 ml on a hot plate, transferred to a quartz 5-ml volumetric flask, and diluted to volume with 35% hydrofluoric acid.

DETERMINATION OF 25-2000 p.p.m. OF MOLYBDENUM, NIOBIUM AND ZIRCONIUM IN TANTALUM

In the determination of 25–2000 p.p.m. of molybdenum, niobium and zirconium in 1-g tantalum samples, scattered background radiation was used as an internal standard to reduce errors caused by daily fluctuations in line voltage, variations in X-ray tube voltage and current settings, fluctuations in noise of the electronic circuitry, and errors in sample preparation. The use of scattered radiation as an internal standard has been discussed by Andermann and Kemp¹. As an example, Fig. 1 shows that, in the determination of molybdenum and zirconium, the effect of varying the X-ray tube voltage is less on line-to-background intensity ratio than on line intensity corrected for background. Variations of X-ray tube current and scintillation counter voltage also show a lesser effect on the line-background intensity ratio than on line intensity corrected for background. Likewise, errors due to dilution and variations of hydrofluoric acid concentration are minimized by using the ratio method.

A total of 128,000 counts are accumulated at the K_{α} lines for molybdenum, niobium and zirconium, and at the background 24.00° 2 θ . To eliminate bubble formation on the Mylar window of the sample cell during the long counting times

Concentration	Stand	lard deviation, p.	p.m.
range, p.p.m.	Мо	Nb	Zı
0-125	16	24	19
300-800	-	22	25
900-1100	31	35	30
1400-1600	_	37	36
1800-2000	54	34	36

TABLE I.—STANDARD DEVIATIONS FOR X-RAY SPECTROGRAPHIC DETERMINATION OF MO, ND AND Zr IN Ta

involved, the window is coated with a nonwetting agent such as Desicote. The total of 128,000 counts is required in order to obtain a statistical counting error sufficiently small to yield a sensitivity approaching 25 p.p.m.

The standard deviation of the procedure was determined at various concentration levels for each of the elements studied. Fourteen determinations were made at each point and values obtained are shown in Table 1.

In Table II, niobium concentrations, as determined by the X-ray spectrographic procedure, are compared with values obtained by a spectrophotometric procedure in which the niobium is separated by ion exchange and determined as the thiocyanate complex. Each value shown is the result of one determination. Assuming the chemical values to be correct, only one X-ray value exceeds the 95 per cent confidence limits of two standard deviations.

DETERMINATION OF 0.5–10 PER CENT OF TUNGSTEN IN TANTALUM

The L_{α_1} line for tunsgten was selected for this work because it is the most intense line available and it is not subject to errors from overlap of lines of other elements expected to be present. To correct for errors caused by fluctuations in X-ray tube voltage and current, incoherent or Compton scattered radiation was selected as an internal standard. By comparison of line-Compton intensity ratio and line intensity corrected for background, it was indicated that the internal standard procedure is more precise. Coefficients of variation of 7 and 1 per cent were determined for the ratio method at the 1 and 7 per cent tungsten concentration levels compared to 12 and 3 per cent, respectively, using line intensity corrected for background intensity.

Because niobium in varying concentrations was expected in most samples, the

48

use of a pulse-height analyzer was investigated for eliminating interference of the second order niobium K_{α} line with the Compton line for tungsten. Although each 0.1 per cent of niobium caused a negative absolute error of 0.06 per cent in tungsten values when no discrimination is used, 3 per cent of niobium causes no error when a pulse-height analyzer is used.

Because the intense L_{α_1} line of tantalum at 44.41° 2 θ slightly overlaps the Compton line at 43.75° 2 θ , care must be exercised in setting the goniometer. An absolute error of 0.28 per cent in tungsten concentration is caused by a 0.01° 2 θ error in goniometer setting. The goniometer setting is approached from the high-energy side each time to

Nb found, <i>p.p.m</i> .		
X-ray	Spectrophotometric	
120	123	
45	78	
90	76	
55	66	
395	420	
285	309	
1055	1095	
95	97	
65	60	
695	694	
50	57	
80	67	
600	671	

 TABLE II.—COMPARISON OF X-RAY AND SPECTROPHOTOMETRIC METHODS

 FOR DETERMINATION OF Nb in Ta

avoid backlash in the dial, and standards are analyzed daily to observe any drift in the goniometer calibration.

For samples containing more than 7 per cent of tungsten, a portion of sample containing 20–70 mg of tungsten is diluted to 1.00 g with high-purity tantalum. This dilution is necessary because it is difficult to maintain more than 70 mg of tungsten in solution during preparation of the sample.

The standard deviation of the procedure, as calculated from twelve to fourteen determinations at each of the concentrations, 1, 5 and 7 per cent of tungsten in 1.00 g samples, was 0.07, 0.10 and 0.09 absolute per cent, respectively. One analyst can perform twenty to thirty determinations per day.

DETERMINATION OF 0.025-10 PER CENT OF YTTRIUM AND THORIUM IN TANTALUM

Samples containing yttrium or thorium are dissolved in hydrofluoric acid as previously described. However, because thorium and yttrium are insoluble in hydrofluoric acid which is the best solvent for tantalum, a separation from tantalum is necessary.

For the determination of yttrium in tantalum, a sample containing 0.25–5 mg of yttrium is dissolved with 48 per cent hydrofluoric acid and a few drops of nitric acid

and 3.00 mg of thorium as the nitrate solution are added. The sample is evaporated to a volume of 3-5 ml, transferred to a 15-ml Pyrex centrifuge cone, diluted to 10 ml with water, and centrifuged for 5 min. The precipitate is washed with two 10-ml portions of dilute hydrofluoric acid, four drops of concentrated sulfuric acid are added, and the mixture is evaporated to fumes in a platinum dish. The soluble sulfate residue is transferred to a 5-ml volumetric flask and diluted to volume with water. The time to accumulate 64,000 counts is measured at the K_{α} line for yttrium and the L_{α_1} line for thorium, and the intensity ratio is compared to ratios obtained for known standards. Standards are prepared by diluting 3.00 mg of thorium as the

Interfering	Th/Y	ratio	Th $L_{\alpha 1}$ line intensity	
substance	Th found, mg	Error, %	Th found, mg	Error, %
Al, 5 mg	2.98	-0.7	3.03	+ 1.0
10 mg	3.01	+0.3	3.00	0
La, 5 mg	2.94	-2.0	2.98	- 0.7
10 mg	2.92	-2.7	2.88	- 4.0
15 mg	2.92	2.7	2.83	- 5.7
Ta, 1 mg	3.05	+1.7	3.07	+ 2.7
2 mg	2.96	-1.3	2.92	- 2.3
H_2SO_4 , 0 drops	3.19	+6.3	3.43	+14.3
4 drops*	3.01	+0.3	3.00	0
6 drops	2.96	-1.3	2.81	- 6.3
10 drops	2.90	-3.3	2.57	-14.3

TABLE IIIINTERFERENCE OF Al, La, Ta AND H2SO4 WITH THE X-RAY
SPECTROGRAPHIC DETERMINATION OF THORIUM
Thorium 3.00 mg

* Standard procedure.

nitrate solution, four drops of sulfuric acid and various amounts of yttrium to 5 ml with water. If the intensity of the internal standard line for the samples varies by more than 6 per cent from intensities obtained for standards, a poor separation is indicated and the sample is discarded.

The procedure for determining thorium is similar to that described for yttrium except the sample size is adjusted to contain 0.25–7 mg of thorium, and 1.00 mg of yttrium is added as an internal standard.

Errors in the determination of thorium attributable to various amounts of aluminum, lanthanum, tantalum and sulfuric acid were studied. Solutions were prepared in the same manner as standards, with no precipitation or centrifugation, and the amount of impurity indicated in Table III was included. These results show that interferences are minimized or eliminated by using the internal standard rather than the line-intensity procedure.

Because no standard alloys of yttrium or thorium with tantalum are available, the reliability of the method is based upon samples prepared by adding known amounts of yttrium or thorium as nitrate solution to weighed amounts of tantalum metal dissolved in 48 per cent hydrofluoric acid. In the concentration range of 0.025-0.075 per cent, 0.25-0.75 mg of yttrium or thorium was added to 1-g samples of high-purity tantalum. In the concentration range of 0.10-10 per cent, samples were taken which contained 1-5 mg of yttrium or 1-7 mg of thorium. Average recoveries and coefficients of variation are shown in Table IV.

Element determined	Concentration range, % in Ta	Number of samples	Average recovery, %	Coefficient of variation, %
Y	0.025 0.075	9	103	7
Y	0.10 -10	33	100.1	2
Th	0.026- 0.075	6	98.7	12
Th	0.1010	18	100.2	3

TABLE IV .-- RECOVERY OF 0.025-10 PER CENT OF Y OR TH FROM TA

Element determined	Concentration range	Coefficient of variation
Nb	100 p.p.m.	24
	1000 p.p.m.	3.5-4
	2000 p.p.m.	1.7–2
Мо	100 p.p.m.	16
	1000 p.p.m.	3.1-3
	2000 p.p.m.	2.7-3
Zr	100 p.p.m.	19
	1000 p.p.m.	3
	2000 p.p.m.	1.8-2
W	1%	7
	5%	2
	7%	2 2
Y	0.025-0.075%	7
	0.10 -10	2
Th	0.205-0.075%	12
	0.10 -10	3

TABLE V.—PRECISION FOR DETERMINING MO, Nb, Zr, W, Y AND Th IN Ta

By using the internal-standard technique for determining thorium or yttrium in tantalum, corrections are made for variations in sample composition as well as for small sample losses during sample preparation. The procedure shows no bias in the recovery of yttrium or thorium from tantalum, whereas calculations based on line intensity show a negative bias of 3.2 per cent for the recovery of thorium.

REFERENCES

¹ G. Andermann and J. W. Kemp, Anal. Chem., 1958, 30, 1306.

THE DETERMINATION OF MODERATOR TURBIDITY PARTICLE SIZE BY AN ANGULAR-DEPENDENT LIGHT-SCATTERING TECHNIQUE*

R. H. GADDY

E. I. du Pont de Nemours and Company, Savannah River Plant, Aiken, South Carolina

Summary—In the characterization of the moderator in heavy-water reactors, information is frequently required concerning the size of the suspended particulate matter. A rapid analytical procedure adaptable to routine control has been developed whereby the particle-size distribution may be derived from angular-dependent light-scattering measurements. A commercial instrument, the Meeco Scattermaster, designed originally for measurements on transparent films, has been adapted for use with liquid samples. Measurements are made of the intensity of the light scattered by the sample over a wide angular range, and converted to a special graphical form in which log $I\theta^2$ is plotted against log θ , where I is the light intensity and θ is the angle of scattering.

This method is based on the assumption that a light-scattering curve for a distribution of size groups is a composite curve in which the individual contributions of the size groups appear as steps in the decline of intensity with angle.

Although not quantitative, the method is, under controlled conditions, capable of providing the following semiquantitative information: uniformity of particle size, presence of particles in a specific range, maximum particle size and most frequent particle size.

This analytical procedure is applicable to particle matter ranging from 0.2–50 μ in diameter.

Particle-size distributions determined by this method on mono- and poly-disperse systems of polystyrene spheres and on representative moderator samples are in satisfactory agreement with the distribution data obtained with the electron microscope.

* Work performed under contract with the U.S. Atomic Energy Commission.

THE ANALYSIS OF A LIQUID METAL REACTOR FUEL*

L. NEWMAN, T. PRACH and H. L. FINSTON Brookhaven National Laboratory, Upton, New York, N.Y., U.S.A.

Summary—A procedure has been developed for the analysis of a uranium–molten bismuth fuel containing 1400 p.p.m. of uranium-235, 350 p.p.m. of magnesium and 250 p.p.m. of zirconium. Analyses are performed for the above elements and for the corrosion products, iron, nickel, chromium and manganese. The activity of the bismuth is approximately 0.2 r hr⁻¹ g⁻¹ at one foot and contains an approximate alpha activity of 10⁹ d min⁻¹ g⁻¹ in the form of polonium-210.

The sample is dissolved in 50% nitric acid, whereupon zirconium and iron can be extracted directly by 0.5M 2-thenoyltrifluoroacetone in xylene. The absorbance due to the iron complex is measured directly on the organic phase. The zirconium is stripped into a fluoride solution and determined spectrographically.

For magnesium and the remaining corrosion products, an aliquot of the original solution is diluted tenfold. The polonium along with the bismuth is extracted by a 0.5M solution of the acid chloride form of tri-iso-octylamine in methyl isobutyl ketone. Three contacts are sufficient to reduce a solution originally containing alpha activity of 10° to less than 10° d min⁻¹ ml⁻¹.

To decontaminate the aqueous solution from rare earth elements, a praseodymium fluoride precipitation is used. Finally, to decontaminate from the alkali and alkaline earth elements, bismuth hydroxide is used to coprecipitate the magnesium and corrosion products. This precipitate is then dissolved in such a way that the usual matrix for spectrographic analysis is obtained.

Uranium is determined in a separate aliquot by a modified dibenzoyl methane method; 1,2diaminocyclohexane-tetra-acetic acid is used to chelate the bismuth and other interfering materials. After adjustment of the pH to between 6 and 7, the uranium dibenzoyl methane complex is extracted into amyl acetate. An adequate decontamination is thus obtained.

THE use of liquid reactor fuels is being investigated at a number of laboratories. Extensive studies have been devoted to the use of uranium in bismuth as a liquid metal reactor fuel at Brookhaven National Laboratory. Most of the chemical investigations have centered around metallurgical studies of corrosion, mass transfer and stability of fuel systems.

Elements in the fuel, for which analyses are required, are uranium, zirconium, magnesium, iron, nickel, chromium and manganese. Approximately 250 p.p.m. of zirconium and 350 p.p.m. of magnesium are added to the fuel as stabilizers and corrosion inhibitors. The presence of iron, nickel, chromium and manganese, in the parts-per-million range, is due to the corrosion of the container material.

In the past, most of the samples submitted for analysis have been "cold". In order to cope with the large number of samples, a spectrographic technique has been used. The sample is dissolved in 50% nitric acid with a resulting bismuth concentration of 10 per cent by weight. Analyses for zirconium, magnesium, iron, nickel, chromium and manganese can be obtained simultaneously, utilizing a solution technique with a rotating disc and a high voltage spark as the source.¹ The instrument used is a Baird 3-m spectrograph with a direct reader attachment. The uranium analysis is done colorimetrically by a modified dibenzoyl methane (DBM) method.^{2,3,4}

In order to test the effects of radiation on corrosion and fuel stability, it has been decided to circulate a bismuth fuel containing approximately 1400 p.p.m. of uranium-235 in the Brookhaven Reactor. It became necessary to devise a method to analyze

* Research performed under the auspices of the U.S. Atomic Energy Commission.

samples containing an α -activity of 10⁹ d min⁻¹ g⁻¹ in the form of polonium-210 and a fission-product β , γ -activity of 0.2 r hr⁻¹ g⁻¹ at one foot. Ideally, this problem could have been solved by putting an arc-spark stand in a shielded α facility, and indeed this approach was used for samples containing one thousandth of this β , γ -activity and one tenth of this α -activity. In order to handle samples of the higher activity, it was thought advisable, because of the limitations of our facilities, to use prior chemical separation of the polonium and a major part of the fission-product activities. The approach pursued was to make use of, as much as possible, our existing spectrographic and chemical techniques.

The procedure is divided into three sections, each of which is described below.

1. THE DETERMINATION OF URANIUM

The main feature of this method is the formation of the DBM complex in an aqueous phase at a pH of 6–7, followed by an extraction into amyl acetate, and a spectrophotometric determination. The use of 1,2-diaminocyclohexane-tetra-acetic acid (DCTA) prevents interference from bismuth and other elements. The amyl acetate extraction serves as a means of obtaining the uranium free of polonium and fission-product activity. The resulting polonium activity in the organic phase is less than 100 d min⁻¹ ml⁻¹ with negligible fission-product activity. The polonium activity can be reduced still further by washing the organic phase with a blank solution of the appropriate pH. Identical results are obtained in the presence or absence of bismuth. A detailed procedure for the determination of uranium is presented.

Reagents

Combine in order:

2 mg Brom cresol green

 $600 \text{ ml } H_2O$

50 ml NH₄OH (conc.)

200 g NH₄Cl

100 g 1,2-diaminocyclohexane-tetra-acetic acid (available as Chel 600 from Geigy Industrial Chemical Co., Saw Mill River Road, Ardsley, N.Y.).

Dilute to 1 liter in a graduated cylinder.

Dibenzoyl methane solution (1,3-diphenyl-3-propanedione), 1 g in 125 ml of acetone.

All other chemicals are of standard laboratory grade.

Procedure

Add to centrifuge tube:

10 ml of DCTA buffer solution.

A sample aliquot of appropriate size containing 10 per cent of Bi in 50% HNO₃.

Adjust the pH with 2M NH₄OH until a blue color is obtained. (If already blue, omit addition of NH₄OH.)

Immediately return to a faint yellow color by the dropwise addition of 1M HCl.

Add 2 ml NH_4OH (2*M*), and 1 ml of dibenzoyl methane solution.

Wait 10 min, and extract with 10.0 ml of amyl acetate.

Determine the absorbancy at 412 or 426 m μ . (For less than 10 μ g of U, extract with 5.0 ml of amyl acetate and measure the absorbancy at 404 m μ .)

Process appropriate standards and a blank through the same procedure as the sample.

2. THE DETERMINATION OF IRON AND ZIRCONIUM

Iron and zirconium are extracted directly from an aliquot of the 10% bismuth-50% nitric acid solution into 0.5M 2-thenoyltrifluoroacetone (TTA) in xylene. If the amount of sample is limited, the aqueous phase after this extraction can be used for the uranium determination.

Direct, p.p.m.	Extract and strip, p.p.m.			
254	250			
248	240			
240	231			
258	249			
251	242			
245	236			
·				
Average 249 \pm 5	241 ± 7			

TABLE I.—Spectrographic zirconium analysis

Moore has adequately demonstrated that zirconium is quantitatively extracted from nitric acid solutions⁵, and we have found no deleterious effect due to the presence of high concentrations of bismuth. The zirconium is re-extracted from the organic phase (TTA in xylene) into 1*M* fluoride solution containing, as the standard matrix for the subsequent spectrographic analysis, 50% nitric acid and 10% bismuth. An internal standard consisting of 0.05 mg of yttrium per milliliter is added to the solution. The results obtained for the direct spectrographic determination of zirconium are compared in Table I with those obtained after the TTA extraction and fluoride stripping steps. There appears to be a negative bias of approximately 3 per cent; however, if one compares the averages they are found to be statistically the same. If desired, this effect can be adequately compensated for by using a calibration curve obtained by carrying standards through the procedure.

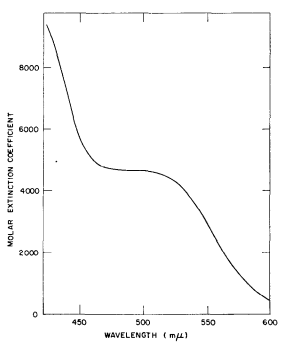
Bolomey and Wish⁶ have recommended that the pH of the aqueous phase be within the range of 2-4 for the analytical extraction of iron by TTA. However, in a review article by Moore it was reported that iron can be extracted efficiently from perchloric and nitric acid solutions.⁷ Furthermore, iron invariably appears as a contaminant in TTA extractions of metals from acidic media. It appeared that it might be feasible to extract iron quantitatively from nitric acid solutions, and indeed, this approach has been utilized in a recent publication.⁸ We have investigated the percentage of iron extracted from 50% nitric acid by 0.5M TTA-xylene as a function of shaking, number of contacts and the organic/aqueous volume ratios. The results are presented in Table II. As can be seen, the 10 min shaking time is sufficiently long to yield almost equilibrium extraction. It was calculated that an organic/aqueous ratio of 4 should yield essentially quantitative extraction of the iron from solutions containing 10 per cent bismuth and 50 per cent nitric acid.

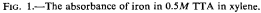
Experience has shown that it is difficult to obtain meaningful iron analyses in the low p.p.m. range if the chemical and manipulative steps are not kept to a minimum. It was, therefore, decided to ascertain whether a spectrophotometric determination of

Shaking time, min		10		30		60	
Number of contac	ts	1	2	1	2	1	2
Organic/Aqueous,	1/1	81	96	84	96	86	
(Volume ratio)	2/1	90		93		93	-
Used Fe ⁵⁹ Tracer		concentration of Fe = 1 μ g/ml					

concentration of Fe = 1 μ g/ml

TABLE II.—PERCENTAGE OF IRON EXTRACTED FROM 8M HNO3 BY 0.5M TTA





the iron could be made directly in the organic phase. The absorbance of the iron-TTA complex versus an appropriate blank is shown in Fig. 1. A plateau is observed between 470 and 510 m μ . A Beer's law plot at 490 m μ is presented in Fig. 2. Blanks were found to be of the order of 0.01 absorbance units. If the lower limit of measurement is considered to be twice that of the blank, 5 p.p.m. of iron can be determined in bismuth on the basis of a 10% bismuth solution and an organic/aqueous volume ratio of 4. In practice, it has been found that this lower limit is adequate for our needs. If a still lower limit is desired, various manipulative techniques can be resorted to.

3. THE DETERMINATION OF MAGNESIUM, CHROMIUM, NICKEL AND MANGANESE

Considering the chemical properties of these elements, it seemed most feasible to decontaminate from α -activity by the removal of the polonium. Using bismuth as a model, it would appear that a chloride anion-exchange approach could be used.^{9,10} On the basis of ease of handling, speed, versatility and maintaining small solution volumes, it is desirable to separate the polonium by means of solvent extraction. Consequently, the use of tertiary amines as liquid anion exchangers was investigated. Tri-iso-octylamine (TIOA) has been used as an extractant for uranium from sulfate¹¹ and chloride¹² media.

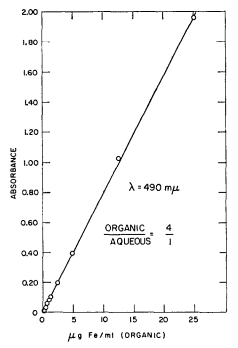


FIG. 2.—Beer's law plot of iron-TTA complex in xylene.

A study was made of the extraction of bismuth by TIOA* as a function of the molarity of hydrochloric acid in the aqueous phase. A 0.5M solution of TIOA in methyl isobutyl ketone (MIBK) was pretreated with 2M hydrochloric acid to put the amine in the hydrochloride form. This solution was then contacted with an equal volume of aqueous solution containing 100 mg Bi/ml and hydrochloric acid in concentrations varying between 2 and 8M. The results for the extraction of bismuth after one and two amine contacts are shown in Fig. 3. The relatively low distribution coefficient after the first contact is due to the fact that the amount of bismuth extracted approaches the capacity of the amine solution. The results obtained for the amine liquids anion exchange are in good qualitative agreement with those obtained by Nelson and Kraus⁹ for resin anion exchange.

Fronaeus has shown, for the case of resin anion exchange, that a maximum in the

^{*} Purchased from Union Carbide Chemical Company, New York 17, N.Y.

distribution coefficient should appear when the concentration of the neutral complex in the aqueous phase is at its maximum.¹³ The mathematical details were later explored by Marcus and Coryell.¹⁴ The same mathematical relationships apply to the liquid anion exchange situation and, indeed, the interpretation might be simplified due to a smaller "resin invasion" by the hydrochloric acid.

On the basis of the theory, the distribution coefficient increases until the aqueous concentration of the neutral complex attains its maximum value, in this case BiCl₃. When the formation of anionic complexes becomes more important, the distribution

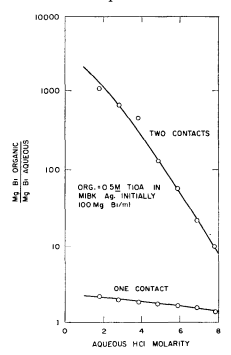


FIG. 3.--Extraction of bismuth by tri-iso-octyl amine hydrochloride.

coefficient decreases. According to the data of Newman and Hume¹⁰, the maximum in the distribution coefficient should appear at a chloride concentration of approximately 0.1M. This low chloride concentration could not be used due to the precipitation of BiOCl. However, we have observed that, if the TIOA in the hydrochloride form is contacted with a nitric acid solution, the amine is distributed according to equation (1):

$$R_3NH^+Cl^-(\text{org.}) + NO_3^- = R_3NH^+NO_3^-(\text{org.}) + Cl^-.$$
 (1)

Now, if bismuth is present in the aqueous solution, the following two equilibria would predominate:

$$3 \operatorname{Cl}^{-} + \operatorname{Bi}^{3+} = \operatorname{Bi}^{3+} \operatorname{Cl}_{3}$$

$$R_{3}NH^{+}Cl^{-} (\text{org.}) + BiCl_{3} = R_{3}NH^{+}BiCl_{4}^{-} (\text{org.})$$
(3)

Therefore, bismuth can be extracted from a nitric acid solution as a chloride complex by contact with an amine solution in the hydrochloride form. Furthermore, considering that the bismuth samples are made up in nitric acid, it is the most convenient medium for the extraction. If the nitric acid concentration is made too low, then BiOCl can once again precipitate upon contact of the aqueous solution with the amine. In order to avoid precipitation, the maximum permissible dilution of the original 10% bismuth-50% nitric acid solution is approximately ten to fifteen-fold.

It then remained necessary to test whether the decontamination from polonium was sufficient at the optimum conditions found for bismuth extraction. The original 10% bismuth-50% nitric acid solution was diluted ten-fold and "spiked" with some neutron irradiated bismuth to yield a resulting α -activity of 1.4×10^6 d min⁻¹ ml⁻¹.

TABLE III.—RESIDUAL PO ACTIVITY AFTER EXTRACTION WITH 0.5M TIOA IN MIBK

Initial activity =	1.4	Х	106	d	min	¹ m	[-1
--------------------	-----	---	-----	---	-----------------	----------------	-----

Number of extractions	d min ⁻¹ ml ⁻¹		
3	203, 97		
4	96		
3, aqueous transferred to clean vial for third	60		
3, aqueous transferred to clean vial for second and third	34		

TABLE IV.—ANALYSIS OF THE BISMUTH HYDROXIDE FRACTION

	Mg	Cr	Ni	Mn	
	244	148	88	107	
	245	150	86	112	
	233	128	67	108	
	242	153	116	104	
	240	149	82	111	
	235	146	107	103	
	238	118	74	102	
	238	129	78	105	
Found	233 ± 4	140 ± 13	87 ± 16	107 ± 4	
Added	236	145	82	105	

A 0.5*M* solution of TIOA in MIBK pretreated with 2*M* hydrochloric acid was used for the extraction which was performed in vials. The results for the polonium decontamination are presented in Table III. The polonium activity is reduced to approximately 100 to 200 d min⁻¹ ml⁻¹ after three contacts with the amine; however, high results were sometimes encountered. Although spurious high results were not encountered after a fourth contact, the decontamination was not significantly increased. The organic phase was removed from the vial between contacts by applying suction to a transfer pipet. It was reasoned that droplets of the organic phase adhering to the sides and top of the vial could contaminate the pipet used in taking the aliquot for residual polonium determination. The simple expedient of pouring the aqueous phase into a clean vial after contact with the organic liquid was found to yield better and more consistent polonium decontaminations. Decontamination was next achieved from rare-earth fission products by a fluoride precipitation, using praseodymium as a carrier. Finally, to separate the magnesium, chromium, manganese and nickel from the alkaline and alkaline-earth fission products, a hydroxide precipitation was performed using bismuth as a carrier. The precipitate was redissolved and analyzed by the standard spectrographic technique. The results after this decontamination procedure are presented in Table IV. They are in agreement with the amount added and the precision compares favourably with the spectrographic analyses obtained on a direct aliquot of a cold sample.

REFERENCES

- ¹ J. Forrest and H. L. Finston, *Third Conference on Analytical Chemistry in Nuclear Reactor Technology*, Paper 11, Gatlinburg, Tenn., 1959.
- ² J. H. Yoe, F. Will and R. A. Black, Anal. Chem. 1953, 25, 1200.
- ^a R. Pribil and M. Jelinek, Chem. listy 1953, 47, 1326.
- ⁴ R. W. Stoenner, unpublished results, Brookhaven National Laboratory, Upton, New York.
- ⁵ F. L. Moore, Anal. Chem. 1956, 28, 997.
- ⁶ R. A. Bolomey and L. Wish, J. Am. Chem. Soc. 1950, 72, 4483.
- ⁷ F. L. Moore, Solvent Extraction in Analysis of Metals, *Am. Soc. Testing Materials, Spec. Tech. Publ.* 238, December 1958.
- ⁸ F. L. Moore, W. D. Fairman, J. G. Ganchoff and John G. Surak, Anal. Chem. 1959, 31, 1148.
- ⁹ F. Nelson and K. A. Kraus., J. Am. Chem. Soc. 1954, 76, 5916.
- ¹⁰ L. Newman and D. N. Hume, J. Am. Chem. Soc. 1957 79, 4576.
- ¹¹ D. J. Crouse, K. B. Brown and W. D. Arnold, U.S. Atomic Energy Commission Report, O.R.N.L. 2173, January 1957.
- ¹² F. L. Moore, Anal. Chem. 1958, 30, 908.
- ¹³ S. Fronaeus, Svensk Kem. Tidskr. 1953, 65, 1.
- ¹⁴ Y. Marcus and C. D. Coryell, Bull. Research Council Israel 1954, 4, 90; Abstracts 132nd Meeting American Chemical Society, New York, September 1957.

A METHOD OF URANIUM ISOTOPE ANALYSIS BY DIRECT READING EMISSION SPECTROSCOPY

JASON L. SAUNDERSON

Baird-Atomic Inc., Cambridge, Massachusetts, U.S.A.

Summary—A special-purpose direct reading spectrometer has been developed for the determination of U²³⁵, U³³⁶ and U²³⁸, using the spectral lines at 4244.12, 4244.22, and 4244.37 Å, respectively. The parallel-channel integrating method is used, in which the ratios of U²³⁵ and U²³⁸ to U²³⁸ are determined simultaneously. A 5 A 2400 Å a.c. arc is used, the sample (uranium oxide) being held in a graphite crater. The three spectral lines are separated by optical means; exact optical alignment is maintained automatically. At 3 per cent concentration of U²³⁵, a coefficient of variation of 1 per cent of the amount present is achieved. The sensitivity for U²³⁶ is better than 0.1 per cent. Sample preparation is simple and straightforward. Less than 3 min is required for an analysis.

INTRODUCTION

EMISSION spectroscopy has been recognized as an extremely promising method for the analysis of uranium isotopes.^{1,2,3} In measuring the relative intensities of the shifted isotope components of a single spectrum line, emission spectroscopy should approach its theoretical limit of precision. This is due to the fact that all the isotope components are homologous and have identical self-reversal or absorption properties. An additional advantage of emission spectroscopy is that it is free from memory effects.

Emission spectrographic techniques fall into two major categories: photographic methods and direct reading methods, in which spectrum line intensities are converted into electronic signals by photomultiplier tubes. Photographic emission spectroscopy has the disadvantage that errors may be introduced in plate development and densitometry. An additional disadvantage, which is applicable in the case of isotope analysis, is that intense spectrum lines may cause halation within the photographic emulsion and consequent broadening of the spectrum line image on the photographic plate. Such broadening may interfere with the accurate densitometry of adjacent lines. These disadvantages point toward the use of direct reading techniques, of which there are several. In choosing between the alternative direct reading techniques, a scanning technique is attractive in the separation of closely adjacent lines. However, a scanning method requires exceptional source stability, and invites the use of such cumbersome sample excitation devices as cathode lamps. A parallel channel technique offers the advantage of allowing the simultaneous integration of the intensities of all the spectrum lines measured. This somewhat relieves the requirements on source stability and consequently on sample preparation. The problems of a parallel channel technique in isotope analysis are principally problems of optical alignment. They concern the optical manipulation necessary to bring light from closely spaced spectrum lines to separate photomultipliers, and the rather stringent requirements for stability placed on such a system.

This paper describes a method of analyzing the 235, 236 and 238 uranium isotope components in the vicinity of 4244 Å (Figs. 1 and 2). The method employs parallel channel, direct reading spectrographic equipment of standard design. To this equipment has been added a special device to obtain the required optical manipulation.

JASON L. SAUNDERSON

APPARATUS

The instrument, a standard Baird-Atomic three-meter concave grating eaglemounting spectrograph is used in the third order with a 25,000 line/in. grating blazed for this application. An interference filter transmitting about 75 per cent at 4244 Å is used in front of the entrance slit, preventing higher orders from reaching the focal curve. The entrance slit is 15μ , and the dispersion of the instrument under these conditions is about 0.8 A/mm.

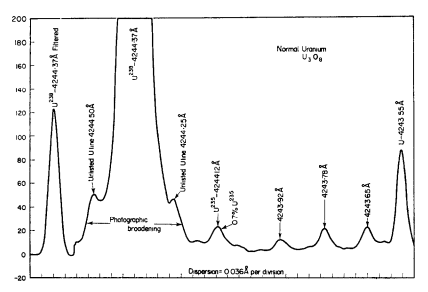


FIG. 1.—Densitometer recording of the spectrum (obtained by photographic spectroscopy) of normal uranium in the 4244 Å region. The 4243.55 Å line at the extreme right is the nearest line listed in Harrison's tables to the short wavelength side of the 4244.37 Å line. Between these two lines are weak unlisted uranium lines at about 4243.65 Å, 4243.78 Å, 423.92 Å and 4244.25 Å, as well as the isotope-shifted components of the 4244.37 Å line. The line at 4244.25 Å is only about 0.03 Å longer wavelength than the position of the 236 uranium istope line. The line at 4244.50 Å on the long wavelength shoulder of the broadened 4244.37 Å line is also an unlisted uranium line. Considerable work was done to establish the validity of the conclusion that the 4244.25 Å and 4244.50 Å lines are indeed uranium line, nor any impurity in the uranium. This conclusion was also reached earlier by personnel of Westinghouse Atomic Power Division. The 4244.25 Å uranium line has frequently been mistaken for the 236 isotope line by other workers.

The densitometer trace of the 4244.37 Å uranium 238 isotope line reduced in intensity by a filter appears at the extreme left. It is evident that in the absence of photographic broadening this line is adequately sharp to permit optical separation from other isotopic lines in direct reading work.

The portion of the apparatus described above can be used for photographic spectrographic analysis of uranium isotopes. Photographic plates with a Spectrum Analysis No. 1 emulsion are used. This emulsion is insensitive to the second and third order wavelengths.

The spectrograph is equipped with devices which enable it to be used for direct reading work. The plateholder is removed and a direct reading head inserted in its place. The direct reading head contains a focal curve, upon which exit slits can be mounted, and means for mounting photomultiplier tubes. The exit slits can be precisely positioned on spectrum lines by manual controls on the front of the spectrograph. The separation of the three spectrum lines of interest, the 235, 236 and 238 isotope lines at 4244 Å constitutes a special problem. Whereas the 235 and 238 isotope lines are about 300 μ apart on the focal curve, since the focal curve is inclined at about 45° to the direction of the light, the actual physical separation of these two extreme lines is only about 210 μ . The 236 isotope line is about 84 μ from the 235 isotope line. Although a 15 μ entrance slit was used, the lines are believed to be about 30 μ wide. Some of this width is presumably Doppler broadening.

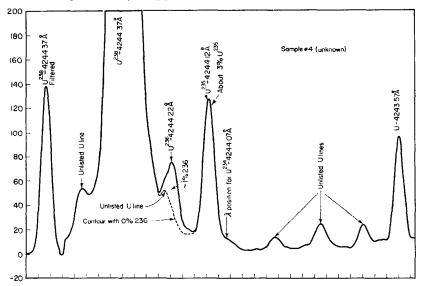


FIG. 2.—Densitometer recording similar to Fig. 1 except that the exposure has been made with an enriched sample. Note that the 236 isotope line is very nearly superimposed on the unlisted uranium line at 4244.25 Å. Since this interfering line is a uranium line, its contribution to the 236 signal is constant and its effect on the 236 calibration curve is exactly the same as background.

The line separation problem is solved by a beam-splitter (Fig. 3). The beam splitter consists of three glass prisms. The coplanar face of the three prisms is placed on the focal curve facing the grating. The prism is adjusted so that the U^{236} isotope line at 4244.22 Å is accepted by the narrow leading edge of the central prism, while the U^{235} and U^{238} components fall on the side prisms and are reflected to the right and left by the aluminized interfaces between prisms. The front surfaces of the side prisms are masked to exclude unwanted portions of the spectrum.

From consideration of the geometry of the apparatus (the long focal distance required for adequate dispersion, and the small and critical dimensions of the beam splitter), it is obvious that some extraordinary means of maintaining optical alignment is required. This is supplied by the automatic servo monitor (Fig. 4) which is a standard component, only slightly modified for this application.

Briefly described, the servo monitor is a servo system which adjusts the apparent position of the entrance slit in response to any change in spectrum position on the focal curve. The sensing device is a tilted slit and two photomultiplier systems. The slit is aligned on a mercury spectrum line produced by a mercury light source which is projected through the entrance slit by a quartz plate. Any lateral misalignment of the mercury line causes an unbalance in the photomultiplier outputs, and this unbalance is corrected by the automatic rotation of a plane parallel quartz deflection plate located between the entrance slit and the grating. The correction is made on mercury light but, because the sample light from the analytical gap follows the same path, all corrections are valid as well for the uranium spectrum. This device holds the spectrum

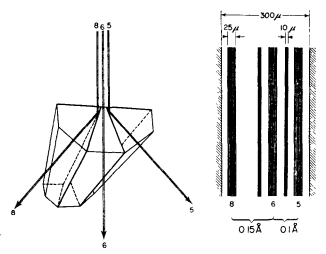


FIG. 3.—Optical device for separating spectrum lines.

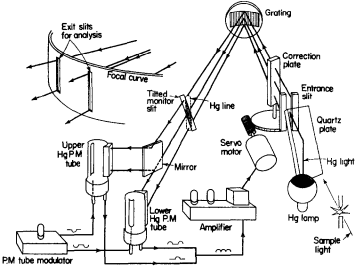


FIG. 4.-Baird-Atomic automatic servo monitor.

to within 1 μ of the desired position, and is absolutely necessary in overcoming shifts due to temperature and pressure variations as well as mechanical dislocations transferred from external sources to the inside of the spectrograph. Although the lateral stability required of the spectrograph for uranium isotope analysis is one order of magnitude greater than the stability required for most direct reading spectrographic applications, no loss of precision due to spectrum shift has been detected even though repeatability runs have been made under unfavorable conditions of laboratory temperature variation.

The readout system is the same as used with a standard Baird-Atomic direct reading spectrometer and Spectromet. During the exposure period, condensers are charged by photomultiplier tube currents simultaneously for all the unknown channels as well as for the internal standard or reference channel. Photomultiplier tube dark currents are automatically cancelled. During the measuring period, all unknowns are read out simultaneously in terms of ratios of voltages of the unknown channels to the voltage of the internal standard. This is accomplished by discharging the internal standard condenser through a resistance, and recording the time taken for the internal standard condenser voltage to decline so as to equal the unknown condenser voltages being measured. Thus, the following equation applies,

$$t = -RC \log_{e} \frac{V_{u}}{V_{s}}$$

where C is the capacity of the internal standard condenser, R is the resistance through which the condenser discharges, V_u is the voltage on an unknown element condenser, V_s is the voltage on the internal standard condenser and t is the time required for V_s to become equal to V_u . If V_s is greater than V_u , t is positive. Percentage concentrations can thus be plotted against t (time or clock readings) on semilog paper, resulting in approximately linear calibration curves, except at concentrations close to the background level. Readout precision is usually better than 0.5 per cent in terms of voltage ratios, but can be increased to almost any extent by using a larger RC constant.

In normal uranium and uranium enriched with low concentrations of U^{235} , the U^{238} line is the internal standard. In highly enriched samples, the U^{235} line is made the internal standard. Yet the change need not be made at the 50 per cent concentration. It is acceptable to use the 235 component as the internal standard from 35 per cent to 95 per cent.

Since this method is based upon the determination of isotope ratios, that is, 235/238 and 236/238, the concentration of the 236 isotope affects the 235 result. If a calibration curve has been established for the 235 isotope based on the 235 to 238 ratio with standards containing no 236 isotope, then as the concentration of 236 increases the true 235 concentration will be smaller than the value read directly from the calibration curve. If the value read from the calibration curve is, say, 3.00 per cent and the sample contains 1.0 per cent of 236, then the actual 235 concentration would be 3.00 per cent less 0.03 per cent, or 2.97 per cent.

BACKGROUND CORRECTION

In addition to measuring the isotope line intensities, the spectrograph is equipped with an exit slit which is positioned to accept background light from a portion of the spectrum near 4244 Å. The ratio of background intensity to reference line intensity is recorded on a clock in the usual manner. The measurement thus obtained may be used to correct for variations in the background light which falls on the isotope photomultipliers. Although such corrections appear to be helpful in some cases, the results obtained so far are contradictory on this matter, and more data must be obtained to prove or disprove the value of background correction.

SAMPLE PREPARATION AND EXCITATION

A weighed quantity of U_3O_8 is mixed with an equal weight of powdered graphite and mixed in a Wig-L-Bug for 2 min. The lower electrode is a $\frac{1}{8}$ in. diameter graphite rod having a 2 mm diameter crated $2\frac{1}{2}$ mm in depth. The electrode is loaded by pushing the rod several times down into the capsule in which the sample was mixed. About 10 mg of uranium oxide are consumed per run. The upper electrode is a flat tipped $\frac{1}{8}$ in. diameter rod. A 2 mm arc gap is used, imaged on the grating of the spectrograph.

The discharge is an automatically ignited a.c. arc, 5 A at 2400 V. A pre-arc period of 80 sec is used. During the pre-arc period, a bead of uranium-containing material,

	0.7%	2.0%	4.0%	6.0%			
	Clock	Clock	Clock	Clock			
	31.0	22.8	14.7	9.5			
	31.0	22.8	14.5	9.8			
	31.0	22.7	14.5	9.6			
	30.9	22.6	14.5	9.9			
Average	30.98	22.73	14.55	9.70			

TABLE I.— U^{235} calibration data

glossy and refractory in appearance, forms on the lower electrode; as the bead forms, the discharge becomes increasingly stable and quiet. During the pre-arc period, the analytical gap widens to such an extent that, at the beginning of the exposure period, the images of the electrode tips no longer fall on the grating. Exposure periods used are 80 sec for photographic analysis, and 40 sec for direct reading analysis.

ANALYTICAL PERFORMANCE

A typical calibration series is given in Table I. When these data are plotted on semilog paper, with the percentage of U^{235} as ordinate and U^{235} clock reading as abscissa, a U^{235} calibration curve is obtained as shown in Fig. 5. The point at 0.1 per cent can be obtained by calculation. The curve at diminishing percentage concentration becomes steeper because 0 per cent is at infinity. Hence the curve approaches a vertical line asymptotically, the position of the vertical line representing the intensity of the background. These properties are easily calculated from the logarithmic equation given previously

Figure 6 is a calibration curve for U^{236} . This curve should be considered as only approximate due to some uncertainties in the actual U^{236} content of the standards used.

Figure 7 is a combination of the calibration curves in Figs. 6 and 7 with a calibration curve for U^{238} in the range from 5.5 per cent to 50 per cent.

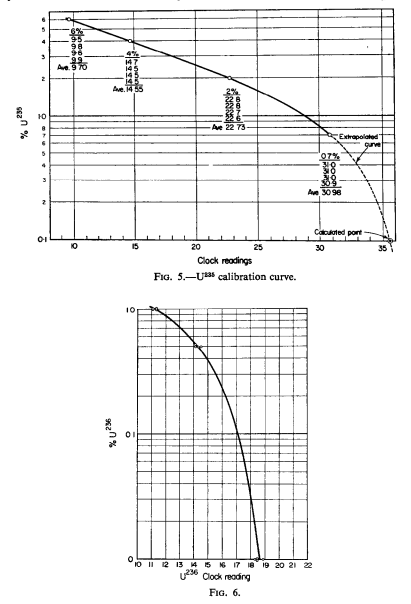
The precision^{*} is close to 1 per cent of the amount present at approximately 3 per cent U^{235} . Work on other samples, for example, nearly equal concentrations of the 235 and 238 isotopes, has indicated possible reproducibility of 0.2 per cent of the ratio.

standard deviation
$$= \left(\frac{\sum d^2}{n}\right)^{1/2}$$
.

^{*} In this discussion, precision is given the same meaning as coefficient of variation, or standard deviation of a single run expressed in terms of percentage of the amount present. Standard deviation is calculated by the usual formula:

Improvement of precision in this case is believed to be due to the fact that the background intensity is an insignificant fraction of the light intensities measured.

At 93 per cent U^{235} , the precision of U^{238} is about 1.2 per cent, and consequently the precision of U^{235} (calculated by difference) is on the order of 0.07 per cent.



The precision for the determination of the 236 isotope is a few per cent of the amount present at 1 per cent concentration. Thus, less than 0.1 per cent should be detectable by suitable calibration. There appears to be a leak of about 5 per cent of the 235 light into the 236 channel. For wide variations in the 235 concentration, a suitable correction could be made to improve the accuracy of the 236 result.

7

CONCLUSIONS

(1) By the use of equipment which automatically maintains optical alignment of a mercury line on an exit slit with a precision of about 1 μ , it has been proved feasible to separate the three closely spaced spectrum lines of the 235, 236 and 238 isotope lines of uranium near 4244 Å for photoelectric measurements. Separation of the three lines is accomplished by a specially constructed optical device which operates as three closely spaced slits.

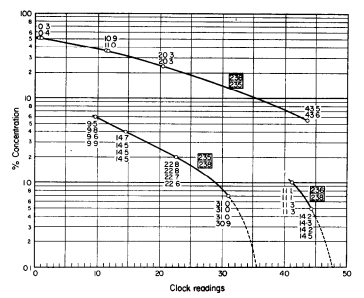


FIG. 7.—For each of the three calibration curves in this Figure, the ordinate is percentage concentration of the isotope listed uppermost in the square box. The abscissa represents the clock reading of the direct reading readout system. In each case, the lower number in the square box represents the isotope component used as the internal standard. The point on the extreme lower right of the top curve represents two runs of a sample containing 5.5 per cent U²⁸³. The U²⁸⁵ concentration for this sample is about 93 per cent, the remainder being other isotopes.

(2) The equipment described in this report allows the simultaneous measurement of the spectrum line intensities of interest over any convenient period of integration, so that stability of the source of excitation is of minor importance. Thus the use of timeconsuming and cumbersome techniques, such as hollow cathode discharge tubes, are avoided.

(3) The fact that the spectrum lines used are perfectly "homologous" and also have identical self-reversal or absorption properties indicates that the method should be capable of high precision. This is borne out by the analytical precision obtained.

REFERENCES

- ¹ L. E. Burkhardt, George Stukenbroeker and Sam Adams, Phys. Rev. 1949 75, 83.
- ² James K. Brody, AECD-2994; ANL-HDY-678, Decl. Oct. 1950, 12.
- ^a G. V. Wheeler, Modern Approaches to Isotopic Analysis or Uranium, P.25-31. A Conference held at Chicago, 5-7, 1957.

THE SPECTROCHEMICAL ANALYSIS OF BISMUTH USING A PHOTOELECTRIC SPECTROMETER

JOSEPH FORREST and H. L. FINSTON

Brookhaven National Laboratory, Upton, N.Y., U.S.A.

Summary—In the Liquid Metal Fuel Reactor Program, a large number of Bi samples must be analyzed rapidly and accurately. A spectrochemical procedure for the determination of Mg, Zr, Fe, Cr, Mn, Ni and Mo, using a rotating-disk technique with a Baird-Atomic spectrograph and direct-reader attachment, is described.

To ensure proper performance of LMFR test loops and related engineering experiments, frequent monitoring is required of molten Bi–U alloy for Mg and Zr additives as well as corrosion products of stainless steel. During the past few years, Mg and Zr in the range of 50–700 p.p.m. were determined spectrographically by spark-excitation of an HNO₃ solution of Bi in a porous-cup electrode. In order to obtain sufficient sensitivity in the analysis of corrosion products at the 10–500 p.p.m. level, a d.c. arc procedure was utilized. This involved time-consuming sample preparation, consisting of the evaporation of Bi solutions followed by ignition, grinding, mixing and electrode packing.

With the present instrument, quicker analyses can be made on a larger number of samples. In order to take full advantage of the instrument, a procedure was developed whereby the concentration of all seven elements of interest is determined by a single exposure. A solution technique, utilizing Y as an internal standard, has been found to yield results which are satisfactory for control of the experimental loops.

Samples are prepared as a 10% (w/v) solution of Bi in 1 : 1 HNO₃. Rapid dissolution is effected by dissolving the sample in the appropriate amount of concentrated HNO₃. The solution is then diluted to constant volume with water. A 10-ml aliquot is then mixed with 1 ml of Y solution containing 0.5 mg of Y per ml. Standard solutions are similarly prepared from pure Bi and solutions of pure metals; in these solutions Y is also added as internal standard.

The prepared solution is placed in a Pyrex glass boat and excited, using a $\frac{1}{2} \times 0.200$ in. rotating graphite counter electrode. Excitation conditions are as follows:

Inductance, µH 1250	
Voltage, V 18,00	Ю
Current, r.f. amp 3	
Number of discharges per second 960	
Analytical gap, mm 4	
Entrance slit, μ 50	
Time of preburn, sec 30	
Time of exposure, sec 60.	

Using the Baird-Atomic direct-reader attachment, condenser discharge time (in clock units) is plotted against the logarithm of concentration to give analytical working curves.

A tabulation will be presented of the spectral lines utilized for the determination of Mg, Zr, Fe, Cr, Mn, Ni and Mo, together with relative standard deviations obtained for each of these elements within specified ranges of concentration.

The method was originally developed by means of a photographic technique; therefore, this technique can also be used, providing some of the intensity levels are reduced with step filters or sectors and the period of exposure is lengthened to 3 min.

DETERMINATION OF U²³⁵ BURN-UP IN IRRADIATED NATURAL URANIUM BY CHEMICAL AND COUNTING METHODS*

D. R. MACKENZIE*

Brookhaven National Laboratory, Upton, Long Island, N.Y., U.S.A.

Summary—A method was desired for determining, with a precision of 1 per cent, the U^{235} content of U samples taken lengthwise of a highly irradiated NRX natural U fuel rod. A mass spectrometric method capable of achieving this accuracy was not available at the time this work was begun.

The method chosen was based on measurements of the fission and α -counting rates of natural U samples and U samples from the fuel rod after chemical purification, in order to compare their U²³⁵ : U²³⁸ ratios.

Due to the thin sources necsssary for the counting, the time required for the direct determination of U^{338} by α -counting was too long to be practical. An indirect measure of the U^{238} content was, therefore, obtained by counting the Pu^{239} produced by neutron irradiation of the sources in a position of high flux in NRX. To ensure a valid basis for comparison of samples, all irradiation and counting was done with a source of natural U and one of depleted U placed back to back. The sources were not disturbed on changing from fission- to α -counting. This type of counting was accomplished by use of an ion chamber with a double grid, located at the opening of a thermal-neutron beam hole. Fission counting was done with the beam hole shutter open; α -counting was done with the shutter closed, a higher gas pressure in the chamber, and a higher electronic gain.

Because of difficulties caused by radiation damage to the sources, it did not appear possible to approach the precision desired for U^{235} burn-up values, and a further complexity had to be introduced into the method.

The new procedure consisted of the following steps:

(1) A set of sources and a set of targets were prepared from natural U and each depleted U sample. The sources were designed for the determination of U^{235} by fission counting; the targets, after processing, were used for the determination of U^{238} via Pu^{239} . Each set of sources and corresponding targets contained Pu^{238} which was added as a tracer. For any given U sample, sources and targets were prepared at the same time and in a manner designed to maintain the Pu^{238} : U^{238} ratio constant for both sets.

(2) Sources were fission- and α -counted in the gridded ion chamber to obtain U²³⁵ : Pu²³⁸ ratios.

(3) Targets were irradiated in the back-to-back arrangement used previously.

(4) After irradiation, the U and Pu were dissolved off the Al target backing and the Pu was carefully purified.

(5) This Pu was used to prepare sources on Pt for α pulse height analysis which gave Pu²³⁹ : Pu²³⁸ ratios directly and thus, indirectly, U²³⁸ : Pu²³⁸ ratios. From these and the U²³⁵ : Pu²³⁸ ratios obtained in (2), U²³⁵ : U²³⁸ ratios were calculated.

The use of an α -active tracer was necessary to link the two sets of counting results. The advantage of using Pu²³⁸ as a tracer was that the method is independent of chemical yields during the processing of irradiated samples.

The precision of the counting methods used for burn-up measurements varied from 1.3 to 2.6 per cent. Before the chemical and counting methods had been fully worked out, a satisfactory mass spectrometric method was developed and put into routine use. The precision of the mass spectrometer measurements, for samples which were analyzed by the counting techniques, varied from 0.5 to 1.4 per cent. Results obtained by analysis with the mass spectrometer and the counting methods agreed within the experimental errors listed above.

* Contribution from A.E.C.L., Chalk River, Ontario, Canada.

INTRODUCTION

The problem

In studying the results of long irradiation of natural uranium fuel, one of the more useful pieces of information which can be obtained is the U^{235} depletion. This gives a direct measure of the integrated irradiation received by a fuel element at any point without the use of external flux monitors.

One NRX fuel rod in particular (Rod. No. 683) has been the object of a very detailed investigation.^{1,2} This rod was cut up in the manner shown by the sketch in

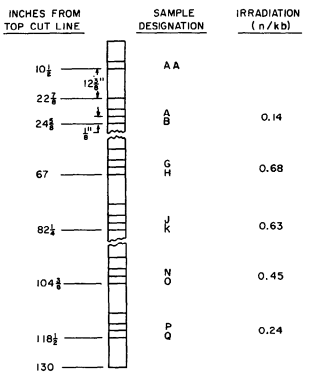


FIG. 1.—Cutting diagram of rod 683 showing location of $\frac{1}{8}$ -inch disc samples.

Fig. 1, and the pieces used for various measurements. The 12 in. lengths were used for reactivity measurements at ANL, and the $\frac{1}{8}$ in. discs for chemical and mass spectrometric analyses, principally at Chalk River. Among the measurements made on the discs were the radial distributions of plutonium isotopes and Cs¹³⁷. Also desired were the values of the U²³⁵ burn-up for as many samples as possible on which other measurements had been made.

At the irradiation levels met with in Rod. No. 683 (shown on the right of the diagram in Fig. 1), burn-up of U^{235} can readily and accurately be determined by use of a sensitive mass spectrometer to measure the ratio U^{235}/U^{238} . At the time work on this problem was started, it was understood that U^{235}/U^{238} ratios obtained with the Chalk River mass spectrometer would be subject to an error roughly 10 times greater than desired. Attempts were, therefore, made to develop a method based on α -particle and fission fragment counting, with a target accuracy of 0.1 per cent so as to give an accuracy of 1 per cent in irradiation at 10 per cent burn-up.

D. R. MACKENZIE

Subsequently, it was found that, with considerable effort, the mass spectrometer was capable of measuring U^{235}/U^{238} ratios to the required accuracy. It was, therefore, decided to use the mass spectrometer routinely for the depletion measurements, and the counting methods were abandoned while still under development. The results obtained had not met with the rather stringent accuracy requirements, and it is not possible to state what accuracy could eventually have been achieved. However, the methods merit description since, apart from any intrinsic interest they may have, the counting and chemical techniques that were developed have formed the basis for a series of precision measurements of fission cross sections³; the apparatus used was, in fact, turned over to these studies.

A detailed description of the apparatus and its use in the burn-up measurements, together with a thorough discussion of the requirements of various methods and the results achieved, are given by Bigham, Hanna and Tunnicliffe⁴. Although the present paper deals with the chemical problems encountered, in order to present an understandable report of the whole investigation, it seemed desirable to include modest descriptions of many of the nuclear physics aspects. For a more detailed consideration of these aspects, the reader is referred to the report of Bigham *et al.*⁴

Basis of counting methods used

Basically, what is required for a depletion measurement is a comparison of the thermal neutron fission rates per unit mass of pure natural uranium and purified depleted uranium in the same neutron flux. The fission rates in this case give a direct measure of the amounts of U^{235} in the samples. A small, easily calculable correction is required for the mass of the depleted sample to take into account the small amount of U^{238} destroyed during irradiation.

Comparison of fission counting rates can be accomplished with the required accuracy, but comparison of the masses of the samples posed several problems. It might be thought, for example, that accurate aliquots of accurately standardized solutions need only be deposited quantitatively as counting sources. However, in the first place, it is extremely difficult to mount samples quantitatively in a form suitable for precise fission counting. And even if this had been feasible in the present study, many of the depleted uranium samples were so small (some as low in weight as 15 mg) that standardization to the required accuracy by gravimetric and volumetric procedures was virtually impossible.

Conventional gravimetric and volumetric methods of uranium analysis were thus of no use, and the possibility of determining U^{238} by α -particle counting was investigated. Counting of U^{238} α -particles directly with the help of a pulse height analyzer did not seem particularly attractive because of low counting rates, and two indirect methods involving α -counting were developed. In all these methods, the α -counting rate actually observed was either that of the U^{238} , or proportional to it. Thus the U^{235}/U^{238} ratio of a given sample was arrived at through a combination of fission- and α -counting.

EXPERIMENTAL APPARATUS

Apparatus used for the chemical work in this study consisted for the most part of conventional equipment and can most conveniently be discussed along with the various procedures. Most of the counting equipment, on the other hand, was developed especially for this work and requires separate description. The individual pieces of experimental apparatus are described in detail elsewhere.⁴ Only the salient feature will be given here for the sake of completeness.

Ionization chamber

In order to be certain that fission counting rates of natural and unknown uranium samples are obtained under identical conditions, the samples should be counted in the same neutron flux—in this case a collimated beam. This can best be done in a double ionization chamber with the samples mounted back-to-back and with as little backing and construction material as possible between the two sources. The effects of attenuation of the neutron beam due to scattering and absorption in the source backings can be eliminated by reversing the sources halfway through the counting. However, it was considered desirable to limit the scattering to that of 1 mm of aluminum, which is about one tenth that of a similar thickness of platinum or stainless steel.

The arrangement of the ionization chamber is shown schematically in Fig. 2. The chamber was filled with an argon-10% methane mixture to a pressure of either 20 cm of Hg, for fission counting, or 40 cm of Hg for α -counting. The gas filling the chamber was continuously purified by convection circulation through a side-tube containing calcium metal chips at 300° C.

The neutron beam

In this work, the neutron spectrum was not important, provided that the fast neutron flux was insufficient to give an appreciable U^{238} fission rate. It was necessary, however, that the beam intensity be rather uniform so that non-uniformity of sample thickness could be tolerated. Of the two beams used for this work, one was uniform to better than ± 1 per cent and the other to better than ± 2 per cent.* One of the experimental arrangements is shown in the diagram of Fig. 3.

Auxiliary equipment

In two of the methods, α pulse-height analysis was required. For this purpose, an alpha grid ionization chamber recording on an eighteen-channel 'kicksorter' was used. The design was similar to that of the system described by Harvey and Jackson⁵.

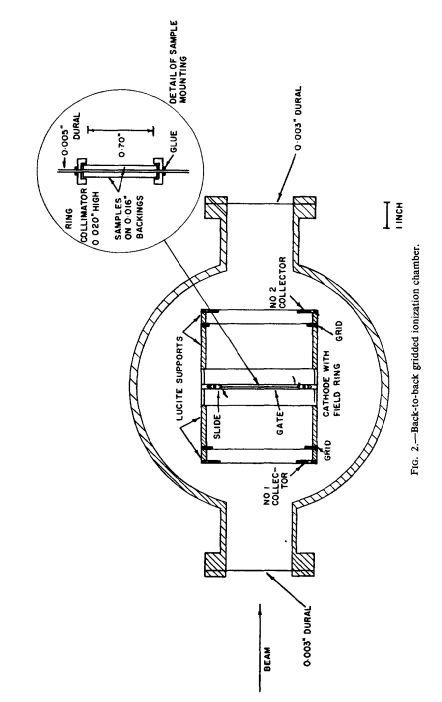
In the third method, the fission counting was done after activation irradiation and it was necessary to correct for the approximately 10 per cent contribution due to Pu²³⁹ fission. This was done by measuring, simultaneously with the main fission counting, the fission rate of a subsidiary Pu²³⁹ source of known α disintegration rate. The simple 'monitor' chamber provided for this purpose was mounted in the neutron beam between the main chamber and the beam catcher. It was very similar to one half of the back-to-back chamber described by Cockcroft⁶.

CHEMICAL TREATMENT OF URANIUM USED FOR SAMPLES

Natural uranium

All samples came from CP-grade starting material which was used without further purification. As a check on purity, samples supplied by two different sources were used. One source was UO_3 powder prepared several years earlier from CP-grade uranyl

^{*} During this work it was necessary to move from the original location (the thermal column) to another part of the reactor (experimental hole S-8).



nitrate hexahydrate; the other was CP-grade uranyl acetate. The former was dissolved in HNO₃ and the latter evaporated with concentrated HNO₃ to form the nitrate starting material. No significant difference was observed in the α /fission ratios of sources prepared from the two different starting materials.

Irradiated uranium

Degree of purity required. Since, in all of the counting methods, it was necessary to determine both fission- and α -counting rates, either all fissile and α -emitting impurities had to be removed from the irradiated uranium samples, or else corrections applied to

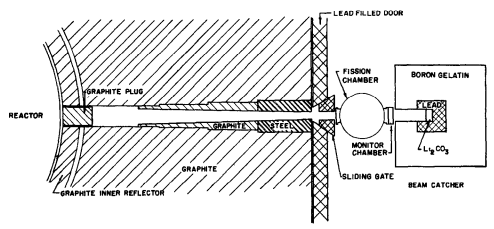


FIG. 3.-Thermal column beam assembly.

account for their presence. Because of the high final accuracy desired, it was evident that, the fewer the corrections which were required, the more hope there would be of achieving it. Consequently, the former course was chosen. The presence of fission products was undesirable only if their β , γ -activity was high enough to cause errors in the fission- or α -counting. In any case, the purification steps used for the heavy elements thoroughly decontaminated uranium from fission products.

It is possible to test for the presence of plutonium and transplutonium elements in uranium by α -pulse analysis—in fact, during the purification procedure, this was done as a check and further decontamination carried out, if required. Three nuclides which could have caused errors due to difficulty of detection in the starting material were Th²³⁰, Np²³⁷ and U²³³, all of whose principal α -particle energies are very nearly the same as that of U²³⁴.

Th²³⁰ was unlikely to be present, and its separation from uranium is, in any case, particularly straightforward. Although neptunium separation is not so simple, only moderate decontamination was required. The maximum Np²³⁷ α -activity originally present in any samples was calculated to be 3 per cent of the uranium activity, and the maximum decontamination factor required for any of the methods was somewhat less than 10³. U²³³ could have been produced from Th²³² impurity in the uranium rod. According to all available evidence, the thorium content was much lower than the 200 p.p.m. which would have been required to cause a significant error due to U²³³ production in Rod. No. 683.

D. R. MACKENZIE

Material for use in the direct counting method required greater decontamination from plutonium, americium and curium than did material for use in the other methods. With these nulides as impurities the α -counting would have been affected more adversely than the fission counting. In order to keep the α -counting rates of plutonium (Pu²³⁹ + Pu²⁴⁰), Am²⁴¹ and Cm²⁴² to 0.1 per cent of the uranium counting rate, the maximum decontamination factors required were 3×10^5 for plutonium, 10^5 for Am²⁴¹ and approximately 10^5 for Cm²⁴².

Purification procedure. Although requirements for the different methods varied somewhat, the most stringent were invariably the objective since starting material purified to such a degree could then be used for any method. The sequence of steps was: (1) ether extraction from saturated $Mg(NO_3)_2$ solution 0.1 M in HNO₃ to separate uranium from Np, Am, Cm, the bulk of the fission products, and some Pu, (2) adsorption on an anion-exchange column (Dowex-1) from 12 M HCl solution to further decontaminate from Pu, Am and Cm, and (3) passage through a Dowex-1 column in 9 M HNO₃ to remove the remainder of the Pu.

Two single volume ether extractions were normally carried out to ensure sufficient decontamination from Np²³⁷, since its detection was not possible by α -pulse analysis. In the second step, after washing Am and Cm through the column, 5–10 column volumes of 1 per cent NH₄I solution in 12 *M* HCl were passed through to elute Pu as Pu^{III}. The uranium was then eluted with distilled water. In the third step, uranium passed rapidly through the column in 9 *M* HNO₃ while the plutonium was strongly adsorbed. Normally, these steps reduced all contaminants to a satisfactorily low level, but in one instance a repetition of the third step was required to give sufficient decontamination from Pu.

METHODS

Three distinct methods were used in this work. As pointed out earlier, they all depended on fission counting for the determination of U^{235} and α -counting for the determination of U^{238} . In Method A* fission and α -counting were done directly on purified uranium samples. In Method B, both fission and α -counting were done on samples which had been neutron activated in the NRX in order to increase their α -counting rates via Pu^{239} production. In Method C, fission counting was done on separate sources, and an even more indirect procedure than in Method B was used to arrive at the U^{238} contents of samples.

Method A

From the standpoint of the number of operations required, this method was the simplest of the three which were tried. It consisted of (1) mounting a source of natural uranium back to back with one of depleted uranium in the double ionization chamber already described, (2) determining the ratio of their fission counting rates in the neutron beam, (3) closing the beam shutter, doubling the gas pressure and determining the ratio of their α -counting rates.

Since the relative amounts of the uranium isotopes are different in the depleted and natural samples, for the determination of the U²³⁸ content of depleted uranium by α counting, a knowledge of its specific α -activity is required. The determined U²³⁸
content of depleted samples must be corrected for U²³⁸ destroyed during irradiation, in

^{*} The methods will be given the same identifying letters as used by Bigham et al.4

order to determine the U²³⁸ initially present; however, this is a small and easily calculable correction. In order to determine the specific activity, on the other hand, an analysis of the α -spectrum is required in order to determine the fraction of total α -counts in the U²³⁸ peak and correct this for the contributions of the other α -particles.

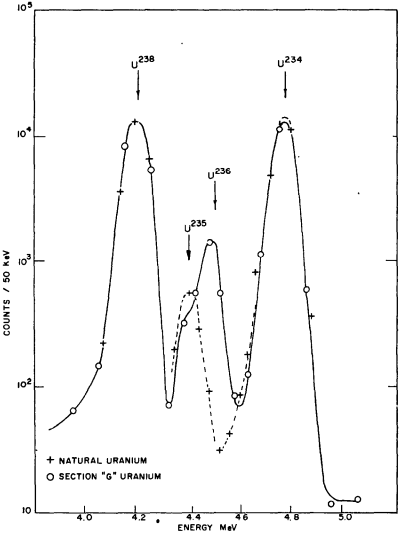


FIG. 4.-Alpha spectra of natural and Section "G" uranium.

The α -spectra were obtained by use of the grid ion chamber and 'kicksorter' described previously. In Fig. 4, the spectra for natural and Section G uranium are shown normalized to give the same intensity in the U²³⁸ peak at 4.20 MeV. Results of the analysis are given in Table I in terms of the relative α -particle specific activities per initial U²³⁸ atom.

Sources used for these subsidiary measurements contained about 100 μ g of U spread over mirror-finish platinum discs $1\frac{1}{8}$ in. in diameter. These were prepared in

two ways—electrodeposition, using the method of McAuliffe⁷, and lacquer spreading, according to the method of Glover⁸ as developed by the author.⁹ Basically, the latter method consists of either pipetting or brushing a lacquer solution containing dissolved $UO_2(NO_3)_2$ over the metal surface to be coated, allowing the solvent to evaporate, and then flaming to burn off the lacquer residue and convert the uranyl nitrate to oxide. For the spectral analysis, the sources prepared by the lacquer technique proved to be the more suitable, and were used to obtain the results given above (Fig. 4, Table I).

For depletion measurements, Method A was very tedious. Because of the low specific α -activity of uranium, a 100 μ g sample requires counting for about 10 days to

Isotope	Energy, MeV	Natural	G
U ²³⁸	4.20	1	0.9975
U ²³⁴	4.76	1	0.9070 ± 0.0043
U ²³⁵	All groups	0.046 ± 0.001	0.0293 ± 0.0007
U^{236}	4.50	0	$\textbf{0.0768} \pm \textbf{0.0008}$
Total		$\textbf{2.046} \pm \textbf{0.001}$	2.0106 ± 0.0045
Ratio: $\frac{\text{natural}}{\text{depleted}}$		1	1.0176 ± 0.0022
U ²³⁵ Burn-up		0	36.3%
value assumed			

Table I.—Relative α -particle specific activities per initial U^{238} atom

achieve the necessary statistical accuracy. At the same time, the background and spurious rates have to be known and held constant to within about 3 counts/hr. Because of this, the method was not considered practical at first; instead, Method B was planned and the double ionization chamber designed with that method in mind. In fact, Method A was not tried until trouble was experienced with Method B. As a consequence, sources prepared by procedures developed for Method B were used for Method A, and these preparative methods are described in a separate section.

Method B

This method was designed to reduce times required for α -counting by a factor of 50 to 100 over those required by Method A. If a thin sample of natural uranium is irradiated for one month, for example, in a high-flux irradiation position of NRX, the resulting Pu²³⁹ α -disintegration rate will be about 100 times greater than that of the natural uranium. Irradiations were carried out with a natural sample back to back with each unknown sample. Since natural and unknown samples were irradiated in the same flux, a comparison of their α -counting rates was equivalent to a comparison of their U²³⁸ contents. Similarly, comparison of their fission rates was equivalent to comparison of their U²³⁵ contents, after applying a suitable correction for the fission rates due to plutonium by the method described previously.

Because of a combination of two effects, the results obtained by Method B were less accurate than had been hoped for. One of these was the deterioration of sources due to irradiation; the other was the persistent and variable contamination of the sources with Po^{210} . The latter affected only the α -counting, the former mainly the fission counting. Although the better sources after irradiation were hardly distinguishable visually from unirradiated sources, marked deterioration of their fission counting plateaus showed that they had, indeed, been thickened by irradiation. This drawback might perhaps have been tolerated but the Po^{210} contamination could not be. Although all materials used in the irradiations (aluminum irradiation capsules, blank discs and spacers, as well as the source disc material) were given various hot acid cleaning treatments to remove traces of surface bismuth, small but significant and uncontrollable amounts of Po^{210} persisted.

Method C

In order to avoid the contamination and irradiation damage problems encountered in Method B, a more complicated method involving the use of Pu²³⁸ as a tracer was investigated. The procedure for this method (Method C) consisted of the following steps:

- (1) A set of sources and also a set of targets were prepared from natural uranium and each depleted uranium sample. The sources were designed for U²³⁵ determination by fission counting; the targets, after processing, were used for U²³⁸ determination via Pu²³⁹. Each set of sources and corresponding targets contained Pu²³⁸ added as a tracer. The amount of tracer was so chosen that its α -activity was about equal to that of the Pu²³⁹ produced in the irradiation.
- (2) For any given uranium sample, sources and targets were prepared at the same time and in a manner designed to maintain the Pu²³⁸ : U²³⁸ ratio constant for both sets.
- (3) Sources were fission and α -counted in the double gridded ion chamber to obtain U²³⁵ : Pu²³⁸ ratios.
- (4) Targets were irradiated in a manner similar to the back-to-back arrangement used previously.
- (5) After irradiation, the uranium and plutonium were dissolved off the aluminum target-backing and the plutonium was carefully purified.
- (6) This plutonium was used to prepare sources on platinum for α-pulse-height analysis, which gave Pu²³⁹ : Pu²³⁸ ratios directly and thus, indirectly, U²³⁸ : Pu²³⁸ ratios. From these and the U²³⁵ : Pu²³⁸ ratios obtained in (3), U²³⁵ : U²³⁸ ratios were calculated.

The use of an α -active tracer was necessary to link the two sets of counting results. As the tracer, Pu²³⁸ had the advantage over a non-plutonium isotope in that the method was independent of chemical yields during target processing.

Although Method C eliminated all the problems it was supposed to, a new and unexpected difficulty arose which prevented the method from achieving the high final accuracy expected on the basis of the counting results. This difficulty will be described during the discussion of the various methods and the results obtained by them.

SOURCE PREPARATION AND CHEMICAL PROCESSING

General requirements

Regardless of method, certain general principles had to be observed throughout the chemical work in an effort to prevent contamination, especially cross contamination, of the samples. The strictness of the precautions required in this work cannot be too greatly stressed. Reagents were checked for α -activity and fissile material. Whenever possible, a separate fume hood was kept for natural uranium and each of the two depleted materials used. All pieces of apparatus, such as electrodeposition cells, were used for only one type of uranium. The one exception to this was the muffle furnace which was used for igniting all sources prepared by the lacquer technique; in this case, it was considered satisfactory to rest each sample on its own insulator during its time in the furnace.

One of the most important requirements for satisfactory fission and α -counting is a uniformly thin source. Coupled with this must be a very smooth backing surface which is reasonably level. Although the neutron beams used were quite uniform, to avoid any adverse effects from even slight nonuniformity, the sources should be reasonably uniform over their whole area. The tolerance was set at ± 10 per cent which was unnecessarily strict in view of the neutron beam uniformity attained. Another requirement of the sources was that they be adherent; this was particularly important for sources which had to undergo relatively long irradiations before being counted.

From the point of view of the counting, it was very desirable to use source backing material which affected the measurement of neutron flux as little as possible. This meant that it had to have rather a low absorption cross section, and also be thin to keep scattering effects small. It was preferable, too, that during in-pile irradiation, either low-level or short-lived activity be produced. From the point of view of source preparation, a material capable of withstanding temperatures of ca. 500° C was desirable in case ignition of the source was required. Aluminum was considered the most suitable backing material on all counts, and pure aluminum sheet (grade 1S) 0.016 in. thick was used.

Source preparation for Method B

At the time the work was started, it was not known how uniform the collimated neutron beam would be. Consequently, much stricter limits were set on source uniformity than later proved to be necessary. For the achievement of such uniformity, it was felt that electrodeposition offered more hope of success than did lacquer painting or pipetting. Also, provided the electrodeposition could be made quantitative, or its rate controlled, the average thickness of the uranium oxide deposit could be closely controlled. Actually, electrodeposition methods were not invariably satisfactory, although a number of suitable sources were obtained; the lacquer method, on the other hand, proved capable of producing at least as good sources, if not slightly better. It should be noted that 'suitable' here is used with reference to sources before irradiation. None of the sources stood up to irradiation quite as well as had been hoped, but again, the lacquer sources were somewhat better.

Electrodeposition directly on aluminum

In applying almost all the methods for electrodepositing uranium oxide adherently on any metal from aqueous solution, a temperature of 70–80° C is required (see, for example, McAuliffe⁷). McAuliffe was not successful in depositing uranium oxide on aluminum, but Wilson and Langer¹⁰ succeeded in depositing thick films of oxide (ca. 2 mg/cm²) at 80° C, using a very thin subcoat of zinc on the aluminum and essentially the same ammonium oxalate electrolyte as recommended by McAuliffe.

The method of Wilson and Langer was found to be of no use in depositing uranium oxide in the range of thickness, $20-50 \ \mu g/cm^2$, required for this work. It has since been shown by the author¹¹ that below a critical average deposition rate no uranium is

deposited, but instead the aluminum cathode surface is attacked and a floc of Al(OH)₃ is formed in the electrolyte. The critical rate varies somewhat for different surface preparations of the aluminum (such as zinc subcoating¹⁰ surface abrading¹² or HNO₃-HF etching¹³) and is such that a minimum thickness of the order of 200 μ g U/cm² can be deposited when aiming for quantitative deposition. Thinner deposits than this can, of course, be obtained by aiming at thick sources and stopping the deposition in its early stages. However, due to variable induction periods and relatively high initial rates of deposition, control of the amount deposited is difficult.

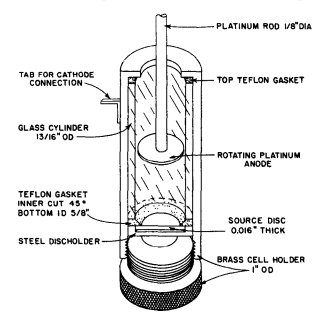


FIG. 5.—Sketch of electrodeposition cell.

In any case, the nature of the deposition process on aluminum appears to be such that roughened surfaces (by etching or abrading) are required for adherent deposits. Since, for precise fission and α -counting, very smooth surfaces are required, no further effort was expended on straight aluminum; instead, the use of thin copper and nickel plates on the aluminum was investigated.

Electrodeposition on copper and nickel plated aluminum

Both copper and nickel can be plated directly on aluminum after the proper pretreatment of the surface, and both elements have low enough neutron absorption cross sections (approximately 4 b) that a plate 0.001 in. thick, for example, can be tolerated by the counting methods. Also, the only activities induced to an appreciable extent (Cu⁶⁴ and Ni⁶⁵) have short enough half-lives to cause no trouble. There appeared to be no doubt that McAuliffe's method⁷ would work well for both metal plates, and the principal problems were to obtain adherent sources on adequately smooth and adherent metal plates.

The cells used for electrodeposition were similar to those used by numerous other investigators.⁷ One arrangement is shown in the sketch of Fig. 5, in which the parts of

the cell were held in place by a brass container with threaded bottom plug. At first, an ordinary washer-shaped gasket had been used at the bottom of the cell and the anode stirrer was moved up and down, rather than rotated, in an effort to prevent an area of light deposition around the center of the cathode which appeared to result from rotational stirring.⁷ When this procedure was used, the uniformity was quite good over most of the area, but the edges were markedly scalloped. The same effect was observed with rotational stirring at speeds of 2-600 r.p.m. Several arrangements were tried with subsidiary copper gaskets of different shapes next to the copper plated cathode source discs; none were satisfactory. The most satisfactory arrangement found was that shown in the sketch, with the Teflon gasket cut at a 45° angle. Although, with this arrangement, quite uniform sources were prepared with a reasonably sharp cut-off at the edge of the Teflon gasket, a fairly large fraction had small spots of heavy deposition or small areas of almost no deposition, apparently due to formation of gas bubbles. These effects were observed even on pure copper discs, and serve to illustrate the fact that phenomena observed during electrodeposition are very dependent on the surface properties*of the base metal, being particularly affected by those irregularities which are not apparent to the eye.

Conditions used for electrodeposition were all within the ranges found most satisfactory by McAuliffe⁷. Those which gave the best results are listed below.

Temperature	80° C
Current density	125 mA/cm ²
$(NH_4)_2C_2O_4$ concentration	0.15 M
Initial pH	8 ± 0.5
Stirring rate	6–700 r.p.m.

Electroplating

Aluminum sheets were prepared for plating by the anodic treatment of Bengston¹⁴ after preliminary cleaning and a 0.5 *M* HF etch at room temperature. After the anodic treatment at 32° C for 10 min, the sheets were immediately transferred to the copper or nickel plating baths. The nickel plating bath was of the high sulfate type used for plating aluminum which has been given a zincate pretreatment. The copper plating process employed was the Unichrome process described by Coyle¹⁵, in which a bath is used of Cu₂P₂O₇ dissolved in a Na₄P₂O₇ solution and the electrodeposition is made at fairly high current densities at 40 to 60° C.

Because of the surface roughening caused by the reported pretreatment of aluminum for direct copper and nickel plating, it did not seem worthwhile to attempt "mirror finish" plating. Instead, after plating, the sheets were buffed to a mirror finish on a high speed buffing wheel and cut into discs. These were thoroughly cleaned by boiling in "Alconox" solution before they were used in the electrodeposition process. Since the original etched aluminum surfaces were rough, the metal plates were also rough, and plates of the order of 0.001 in. in thickness were required to allow mirror polishing without going through to the base metal. Almost invariably, blisters formed on copper plates originally 0.002 in. thick after they were polished and boiled in "Alconox" solution, and nickel plates of 0.001 in. often suffered the same fate. While nickel plates around 0.0005 in. in thickness were usually adherent, they were not easily polished without going through to the aluminum; consequently, most of the sources were prepared on copper plated discs. If the latter survived the boiling treatment and the 30-40 min electrodeposition at 80° C without any signs of blistering, it was observed that they did not blister under irradiation.

Sources by lacquer method

The basic steps in the lacquer method for depositing thin films of uranium oxide (and many other medium and heavy metal oxides) have already been outlined. Lacquer solution containing dissolved $UO_2(NO_3)_2$ was prepared as follows. A strong nitric acid solution of $UO_2(NO_3)_2$ was evaporated just to dryness under an infra-red lamp, avoiding too strong heating. The residue was dissolved in a small volume of acetone and transferred to the required volume of a solution of cellulose nitrate in amyl acetate. The volumes were calculated so that the final solution would have the correct uranium concentration for the particular purpose for which it was required, and the proper cellulose nitrate and acetone concentrations for either painting on with a brush or applying with a pipette. For example, a solution containing 3–3.5 g of cellulose nitrate per 100 ml was most suitable for pipetting, but this was too viscous for painting, becoming tacky before it could be completely brushed out. Solutions containing around 2 g/100 ml were most suitable for painting.

At first, sources were prepared by pipetting onto the counting discs the amounts of uranium solution required to give the desired counting rates. Since there were still some doubts about the uniformity required, it was felt that such sources might not be uniform enough. Accordingly, it was decided to switch to the painting technique. In this procedure, about ten successive coats were applied to a relatively large sheet (2 in. \times 3 in.) and discs were punched out from the areas that appeared by visual inspection to be uniform. For macroscopic uniformity, such inspection was adequate since layers of all thicknesses above about 10 μ g U/cm² were colored (interference effects) and, in the range of thickness of sources used (20–40 μ g/cm²), the color varied rather rapidly with thickness. Thus, it was only necessary to select areas of a sheet where the color did not vary in order to obtain uniform sources. By means of the color, the film thickness could also be estimated quite closely and, hence, the weight of uranium on the disc.

For successful application of the lacquer technique, it was required that the sheets be flat and level during painting. To keep them level they were placed on a small plate with a bubble levelling device. Each polished sheet for painting was heated at 500° C for a few minutes to soften it, then flattened by passing it through a small pair of rolls while sandwiched between two sheets of polythene (to avoid roll marks).

For successful counting, very smooth surfaces were desirable. With a little practice a mirror finish on small areas ($\ll 1 \text{ in.}^2$) can be obtained by electropolishing. Quite a lot of effort was put into developing methods for electropolishing the 0.016 in. 1S aluminum sheet in areas of 5–6 in.², with only modest success. It was found that chemical polishing produced surfaces which were satisfactorily smooth for counting, and 2 in. by 3 in. pieces could easily be handled, so all subsequent polishing was done chemically. The baths used were the R5 polishing bath¹⁶ containing phosphoric, acetic and nitric acids, and one due to Jumer¹⁷ which was largely phosphoric acid with small amounts of Na₂SO₄ and NaNO₃. Both baths were operated at around 100° C or a slightly higher temperature.

Chemical work associated with Method C

Sources were all prepared by the lacquer technique in the same manner as described in the preceding section. Successive coats of lacquer were applied to and burnt off chemically polished aluminum sheets 0.016 in. in thickness, and source discs were punched out in areas where the uranium oxide deposit was uniform.

Target foils were also prepared by lacquer painting; for this purpose, however, 2 in. squares of 1S aluminum foil 0.005 in. in thickness were used. For each irradiation three foils were prepared, one containing approximately $10 \,\mu g/cm^2$ of depleted uranium and Pu²³⁸ tracer, and two containing approximately $5 \,\mu g/cm^2$ of natural uranium and Pu²³⁸. These foils were wrapped in 0.001 in. aluminum foil to prevent cross contamination during irradiation, rolled into cylinders and placed inside an irradiation capsule. The depleted foil was sandwiched between the two natural foils. This arrangement was chosen to minimize differential attenuation of resonance neutron flux by averaging the effect between the natural and depleted samples.

For each irradiation, a set of foils prepared in exactly the same manner, except that the uranium was not 'spiked' with Pu^{238} , was included with the 'spiked' set. These were used to measure the production of Pu^{238} from uranium, since the rates of production were different for the natural and depleted samples due to the presence of U^{236} in the depleted material.

Chemical treatment of irradiated targets

After removal of the samples from the reactor, a waiting period of about 2 weeks was allowed for the bulk of the Pu²³⁹ to grow in from the Np²³⁹ remaining at the end of the irradiation. About 0.3 μ g of plutonium was produced in each irradiated sample. Chemical separations were carried out simultaneously on the natural and depleted material. The steps of the separation procedure outlined below were all given thorough preliminary tests, using unirradiated target foils and (particularly to check on decontamination from Po²¹⁰) irradiated sources from Method B. The steps were:

- Dissolving the deposit off the target foils and their 0.001 in. wrappers with hot 1:1 HNO₃. This dissolved appreciable amounts of aluminum, but weaker acid did not satisfactorily dissolve the deposit.
- (2) Adsorption of plutonium from 9M HNO₃ on 'large' Dowex-1 resin columns and washing uranium, aluminum, neptunium and polonium through with 15 column volumes of 9M HNO₃.
- (3) Elution of plutonium by means of 0.5M NH₂OH·HNO₃.
- (4) Repetition of steps 2 and 3, using 'small' Dowex-1 resin columns.

The 'large' columns were approximately 1 cm in diameter and 5-6 cm in length to allow reasonable flow rates for the relatively large volumes of viscous aluminum nitrate solution. The small columns were approximately $1\frac{1}{2}$ mm in diameter by 4 cm in length. They were designed to give a final decontamination from traces of uranium and aluminum which might have come through the large columns. This was necessary because no visible solid residue could be tolerated in the final plutonium sources.

Plutonium source preparation

The lacquer method of depositing plutonium after the above purification was useless because, in fact, there was invariably a visible residue on the platinum discs after preparation of the sources. This residue was inorganic (since flaming did not remove it) and was definitely associated with the plutonium and not the lacquer solution. Later experience in the preparation of thin β -sources¹⁸ has shown that virtual elimination of solid residues from source materials purified by ion-exchange can only be achieved by special purification of reagents and working in polythene instead of glass to avoid pick-up of inorganic matter.*

The use of electrodeposition promised to overcome the residue problem since the conditions required are not favorable for deposition of cations such as the alkali and

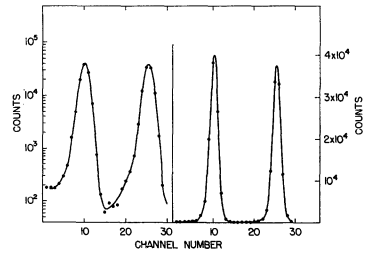


FIG. 6.—Pulse analysis of Pu²³⁸-Pu²³⁹ peaks using source SP-15.

alkaline earth elements which might have come from the glassware and reagents. In the method of Miller and Brouns¹⁹, provision is made for the oxidation of plutonium to the plutonyl state, using ozone to avoid the introduction of interfering cations. Since an ozonator was not available, a method of oxidizing the plutonium with the aid of persulfate was developed.²⁰ Any alkali (or ammonium) persulfate may be used since the Miller and Brouns electrolyte is a 1-2M KOH solution. One ml portions of 0.2M nitric acid solution containing $0.3 \ \mu g$ Pu were placed in a boiling water bath, a few small crystals of an alkali persulfate was added with stirring, and the solutions were heated for about five more minutes. After transfer to the 1.2M NaOH electrolyte in electrodeposition cells of the same type as shown in Fig. 5, plutonium was deposited on mirror-finish platinum discs 1 in. in diameter in a circle 1 cm in diameter at the center of the disc. The sources were invisible until after they had been flamed, at which time a faint brown shadow appeared. This was attributed to the presence of traces of iron in the caustic electrolyte.

Pu²³⁸-Pu²³⁹ pulse-height analysis was performed, using the grid ion chamber and eighteen-channel 'kicksorter'. For the "spiked' samples, in which counting rates of the two isotopes were roughly equal, a correction of about 0.4 per cent had to be applied for the contribution which the tail of the Pu²³⁸ peak made to the Pu²³⁹ peak. Fig. 6 shows an α -spectrum obtained for source SP-15, using a 'kicksorter'-ion chamber

* One exception is the method of Børnholm et al., Nature 1956, 178, 1110 using what might be called microcolumns in glass capillaries and only a very few λ of extremely pure eluting solutions.

combination very similar to that described earlier. Correction for the Pu^{238} tail in this case amounts to about 0.3 per cent, and Po^{210} activity is undetectable (<0.1 per cent of either peak).

EXPERIMENTAL RESULTS

Of the measurements by Method A, three were made on Section G material, using two pairs of electrodeposited sources, and one on Section P material, using lacquer sources (refer to Fig. 1 for location of discs P and G.)

The results are given in Table II, and represent about 800 hr of counting for the G and about 80 hr for P data. Typical rates in the experiments with material from

Samples	Burn-up, %	Deviation from mean
G14/U14	38.47	1.48
	±0.23	
G15/U15	36.68	0.31
	±0.28	
G15/U15	35.81	1.18
	±0.28	
	Mean value 36.99 \pm 0.64*	
P90/U90	15.00	ĺ
	±0.23	

TABLE II.—DEPLETION MEASUREMENTS BY METHOD A

Note: Errors given are statistical errors of counting except for that marked with an asterisk, which is the root mean square deviation.

section G were 30 α counts/min with a background of 2.5 counts/min and about 3000 fission counts/min with a background of about 2 counts/min. The results show a deviation from the mean considerably larger than expected from the statistical errors. The one result for P material was obtained from sources with counting rates about three times those used for the G material, and is in good agreement with the mass spectrometer result (see Table V).

The same sort of scatter obtained for results from the analysis of material from Section G, using Method A, is observed in the results by Methods B and C. These results are summarized in Tables III and IV, respectively. Using Method B, seven pairs of sources were measured, including one natural uranium pair which was included as a check. Of the other six results, one was rejected as an obvious 'flier'. The root mean square deviation of the set of five 'good' determinations was 2.3 per cent, which is 15 times that expected from statistics.

Considering Method C, the fission/ α -count ratio measurements on the unirradiated sources were straightforward, with counting rates of a few thousand per minute. The measurement of Pu²³⁸/Pu²³⁹ activity ratios in the plutonium separated from the irradiated targets also presented no problems. However, the result for material from the G Section was quite wide of the mark, and deviation of the results for material from the P Section was considerably higher than would be expected from the purely statistical errors in the counting.

In Table V, the collected results of depletion measurements by the three counting methods are compared with the mass spectrometer results. Except for the isolated

Capsule number	Sample numbers	Prepared by	(235/238) Depleted (235/238) natural	Statistical error	Deviation from mean
C 79	U41/G41	Painting	0.6606	0.0015	0.0208
C 79	U40/G40	Painting	0.5438	0.0009	Rejected
C 80	U14/G14	Electrolysis	0.6348	0.0006	0.0050
C 80	U42/G44	Electrolysis	0.6526	0.0012	0.0128
D 70	U68/G66	Painting	0.6190	0.0016	0.0208
D 70	U65/G65	Painting	0.6322	0.0016	0.0076
D 70	U66/U67	Painting	1.0014	0.0018	

TABLE III.-DEPLETION MEASUREMENTS BY METHOD B

Notes: Mean of G/U comparisons, 0.6398 \pm 0.0067.

Root mean square deviation of a single measurement, 0.0149 = 2.3 per cent.

Number	Pu source number	$\frac{(\alpha_{9}/\alpha_{8})_{\rm D}}{(\alpha_{9}/\alpha_{8})_{\rm N}}$	Еггог, %	Fission source number	$\frac{(F/\alpha_8)_D}{(F/\alpha_8)_N}$		Error, %	Burn-up, %
I	SG2 SU2	0.9784	0.14	GS1/US1 GS2/US2	0.5872 0.6010	0.6000 0.6134	0.16 0.17	39.5 ± 0.4
п	SP6,7 SU6,7	0.9511	0.14	PS5/US5 PS6/US6	0.8063 0.8151	0.8474 0.8570	0.16 0.18	14.9 ± 0.4
ш	SP12,13 SU12,13	0.9332	0.11	PS10/US10 PS11/US11	0.8004 0.8051	0.8577 0.8628	0.13 0.18	14.1 ± 0.3

TABLE IV .--- DEPLETION MEASUREMENTS BY METHOD C

Note: D = depleted; N = natural.

TABLE V	Collected	RESULTS
---------	-----------	---------

Method	Burn-up, %		
Method	G	Р	
 A. Alpha and fission counting of unirradiated samples B. Alpha and fission counting of irradiated samples 	37.0 ±0.6 36.0 ±0.7	15.0* ±0.2	
C. Pu ²³⁸ tracer method	(39.5)†	14.5‡ ±0.4	
Mass spectrometer analysis (Lounsbury ¹)	36.3 ±0.16	14.96 ±0.21	

* Isolated measurement; accuracy quoted is statistical. † Isolated measurement; probably invalidated by bad mixing of tracer.

‡ Mean of two; error corresponds to the difference between them.

measurement of G material by Method C, the counting results agree with the mass spectrometer results within the accuracy claimed. However, the accuracy achieved with any of the counting methods did not reach the ± 1 per cent desired.

DISCUSSION AND CONCLUSIONS

It was emphasized in the last section that the scatter in the counting results was disappointingly large, considering the relatively small errors arising in the counting itself and in the corrections (mostly small) which were required in the various methods. The reasons for this, insofar as they are known, were different for each method, and thus would require different treatment for their reduction or elimination.

In the case of the results for G material by Method A, the scatter is believed to be due primarily to fluctuations in background during the lengthy periods of α -counting. Trouble from this source can be greatly alleviated simply by increasing the counting rates of the samples used. In fact, the counting rates used in the analysis of P material were three times those used in the analysis of G material. By designing a chamber to take larger source discs, sources of still higher counting rate could be used.

The scatter in Method B results was probably due partly to uncertainties in the counting (particularly fission counting) resulting from irradiation damage to the sources. However, a larger uncertainty was introduced by the random polonium contamination mentioned earlier. Although the presence of Po²¹⁰ on the sources was not directly proven, it was found as a surface deposit on the blank spacing discs which had faced the sources at a distance of about 5 mm during irradiations. Small amounts of plutonium, very variable, but averaging about 0.2 per cent of the sample, were also found on the blank discs, so that it is completely reasonable to expect that the polonium, which is notorious for migrating, had been transferred from the sources to the blank discs during irradiation.

The amounts of bismuth required to give the observed Po^{210} activities were very small, being about 5×10^{-10} g for the activities on the blank discs. A random variation of about 2×10^{-9} g between sources would account for the observed scatter of the results, so that amounts of the order of 10^{-8} g were probably present on the sources. Since surfaces of source discs were cleaned (e.g. by chemical polishing) before source deposition, it appears that the bismuth found its way to the sources during deposition. Indeed, it is not surprising that millimicrogram amounts might have been present in the chemical reagents and in apparatus which could not be given standard acid cleaning treatment such as the brushes used for lacquer painting. It should be feasible to decrease this trace of bismuth contamination by a factor of 10 or so without expending too great an effort, especially once the source or sources of contamination have been discovered. However, in dealing with such small amounts there is always the worry that, having sufficiently cleaned up all reagents and apparatus, contamination will sooner or later creep in from some unexpected source.

One obvious way of solving the problem without preliminary decontamination from traces of bismuth would be to determine the exact amounts of Po²¹⁰ contamination by radiochemical and counting methods after the samples had been counted. This course of action, however, it not too attractive because it adds two more steps to the already lengthy procedure.

In the case of Method C results, the lack of constancy was apparently largely due

to lack of constancy of the Pu²³⁸/U²³⁸ ratio on the targets and foils. Detailed examination of the results showed that those obtained with different pairs of samples did not agree; for example, in measurement I there was a discrepancy of 2.2 per cent between the F/ α ratios from GS1 and US1 and those from GS2 and US2. This implies that the Pu²³⁸/U ratio was not constant over the aluminum sheets from which source discs were punched.

Although it was felt that during preparation of 'spiked' lacquer solutions the Pu^{238} tracer was thoroughly mixed with the uranium, some doubt existed because of the high viscosity of the solutions. Special precautions were, therefore, taken with the mixing of the tracer in the third series of measurements. Although the discrepancy between results from the two sets of sources was only 0.6 per cent, the burn-up value obtained still differed considerably from the mass spectrometer value. This leads one to suspect that, regardless of constancy of Pu^{238}/U ratio on the source sheets, this ratio for the sources was different from that of the targets.

Another possibility is that, regardless of the constancy of the Pu^{238}/U ratio on both the sources and targets, the Pu^{238}/Pu^{239} ratio was altered during treatment of the irradiated targets. During the dissolution of the target material off the aluminum backings, this could happen only if the target material were incompletely dissolved and one of the isotopes (presumably Pu^{239} because of recoil from neutron capture in the parent uranium) tended to be imbedded in the backing to a greater extent than the other. Certainly no large effect could be attributed to this cause since a fair depth of aluminum was dissolved from the backing at the same time as the target material. However, it is possible that a small error could have arisen from this cause, especially since attack of the aluminum backing was not uniform on a micro scale.

If it is assumed that any such error was too small to be significant, we are left with the alternative that the Pu²³⁸/U ratios had not in fact been maintained constant. This effect must then have been due to some phenomenon occurring in the lacquer solutions themselves, or in the process of target preparation.

Regarding the former possibility, it is well known that trace amounts of inorganic ions are adsorbed from aqueous solutions on the walls of glass containers. For example, in standardization of solutions of radioisotopes, small amounts of inactive carrier are added and solutions are kept acidic to minimize adsorption. In our situation, it is conceivable that slow adsorption of Pu²³⁸ tracer on the glass container walls was occurring during the course of source and target preparation. A similar type of occurrence, involving Pu²³⁸ tracer, was observed by Hart²¹ in nitric acid solution. He found that, after thorough mixing, considerable time had to elapse before the ratio of Pu²³⁸ tracer to other plutonium isotopes would remain constant.

In our procedure of lacquer painting, adsorption of the tracer would probably have caused the most trouble in the mechanics of source and target deposition. For example, adsorption might have occurred on particles of material from the brushes, or even in the brushes themselves, in such a way as to progressively change the Pu^{238}/U ratios. Then, since more coats were required for sources than for target foils, the final ratios on the two would have been different.

If adsorption on the glass were the only problem, it probably could be solved by allowing the adsorption process to come to equilibrium before beginning lacquer application. However, if adsorption in the brushes was involved, probably the only solution would be to throw away the brushes and develop the technique of applying lacquer with a micropipet. This undoubtedly can be done on the size of sheet used in this work, since it was done successfully²² on extremely thin foil over an area roughly half this size during the course of another investigation.²³

Comparison of Methods A, B and C can be made from both the counting and the chemical point of view. The 'chemical' comparison is the critical one in deciding on a method to use, since counting techniques have been mastered or, in the case of the troublesome α -background in Method A, almost certainly can be without to much effort.

From the chemical point of view, Method A presents no problems at all, and in terms of effort is much simpler than either Method B or C. The chemical problems of both Methods B and C could be, with considerable effort, either solved or circumvented, but both are long and C in particular is rather complicated. By comparison, the chemical work required for preparation of mass spectrometer samples is simple, since Pu and other α -activities many times that of the uranium can be tolerated. Of the three counting methods, Method A most nearly corresponds to the mass spectrometric method as far as overall effort is concerned, although the chemical effort is greater for the counting method.

Two more things should be pointed out. The first is that any of the counting methods, even at the stage of development which was attained in this work, would be applicable where accuracies of only a few per cent are needed. Under such conditions, the large total number of counts taken for each determination in the present work would not be required, and Method A would become quite attractive. The other point to be made is that the accuracy achievable by counting methods increases progressively as the burn-up increases, basically because the fission rate is a good measure of U²³⁵ even at very low concentrations. Since this is unlikely to be true of mass spectrometric determinations, counting methods, particularly method A, may be very useful in the concentration region encountered in very high burn-ups.

This paper should give some idea of the complexity of the type of analytical problem which is often encountered in the determination of specific reactor constants and more general nuclear data. The problem discussed in the present paper proved to be more difficult than most, and was not completely solved before work had to be stopped, although it probably could eventually have been solved. However, the description of our attempts at its solution will serve to illustrate the wide variety of techniques and meticulous attention to detail required for successful solution of such complex problems.

ACKNOWLEDGEMENTS

The experiments described in this paper represent a joint effort of several people. The author is responsible only for the chemical aspects. Dr. C. B. Bigham, G. C. Hanna and P. R. Tunnicliffe were responsible for the conception and design of the physical aspects of the experiments and made most of the physical measurements. Their report has been referred to earlier.⁴ I am greatly indebted to them for helpful comments on the content of this paper and for permission to reproduce Figs. 2, 3 and 4.

Dr. J. P. Butler and Mrs. J. S. Merritt kindly supplied the Pu²³⁸ tracer used.

REFERENCES

¹ D. G. Hurst, A. H. Booth, M. Lounsbury and G. C. Hanna, Atomic Energy of Canada Ltd., CRR-622 (Revised) 1957.

U²³⁵ burn-up in irradiated natural uranium by chemical and counting methods

- ² D. S. Craig, G. C. Hanna, D. G. Hurst, S. A. Kushneriuk, W. B. Lewis and A. G. Ward, *Long Irradiation of Natural Uranium*, Second United Nations International Conference on the Peaceful Uses of Atomic Energy, Geneva, 1958, A/Conf. 15/P205.
- ⁸ C. B. Bigham, G. C. Hanna, P. R. Tunnicliffe, P. J. Campion, M. Lounsbury and D. R. MacKenzie, *The Slow Neutron Fission Cross Sections of the Common Fissile Nuclides*, Second United Nations International Conference on the Peaceful Uses of Atomic Energy, Geneva, 1958, A/Conf. 15/P/204.
- ⁴ C. B. Bigham, G. C. Hanna and P. R. Tunnicliffe, Atomic Energy of Canada Ltd., CRR-861 (1959).
- ⁵ B. G. Harvey, H. G. Jackson, T. A. Eastwood and G. C. Hanna Can. J. Phys. 35, 258 (1957).
- ⁸ H. S. Cockroft, United Kingdom Atomic Energy Authority, A.E.R.E. N/R 890 (1952).
- ⁷ C. McAuliffe, U.S. Atomic Energy Commission, A-3626, February 1946.
- ⁸ K. M. Glover and P. Borrell, J. Nuclear Energy 1955, 1, 214.
- ⁹ D. R. MacKenzie, Atomic Energy of Canada Ltd., C.R.D.C.-621, 1960.
- ¹⁰ C. R. Wilson and A. Langer, Nucleonics 1953, 11, No. 8, 48.
- ¹¹ D. R. MacKenzie, unpublished work, 1957.
- ¹² L. Koch, J. Nuclear Energy 1955, 2, 110.
- ¹³ J. M. McKenzie, *Nucleonics* 1959, 17, No. 1, 60.
- ¹⁴ H. Bengston, Trans. Electrochem. Soc. 1945, 88, 307.
- ¹⁵ T. G. Coyle, Proceedings of the American Electroplaters' Society, 1941, p. 113.
- ¹⁶ B. E. Bunce, Metal Finishing J. London 1954, 52, (1) 70.
- ¹⁷ J. F. Jumer, U.S. Pat. 2,705,191, March 29, 1955.
- ¹⁸ J. S. Geiger, G. T. Ewan, R. L. Graham and D. R. MacKenzie, Phys. Rev. 1958 112, 1684.
- ¹⁹ H. W. Miller and R. J. Brouns, Anal. Chem. 1952, 24, 536.
- ²⁰ D. R. MacKenzie, Atomic Energy of Canada Ltd., PR-CM-5A, p. 54, 1956.
- ²¹ R. G. Hart, M. Lounsbury, C. B. Bigham, L. P. V. Corriveau and F. Girardi, Atomic Energy of Canada Limited, CRRP-761, Part B, September 1959.
- ²² D. R. MacKenzie, Atomic Energy of Canada Ltd., PR-CM-10A, p. 49, 1957.
- ²³ C. H. Westcott, W. H. Walker and T. K. Alexander, Effective Cross Sections and Cadmium Ratios for the Neutron Spectra of Thermal Reactors, Second United Nations International Conference on the Peaceful Uses of Atomic Energy, Geneva, 1958, A/Conf. 15/P/202.

ANALYTICAL METHODS USED IN A STUDY OF REACTIVITY CHANGES WITH LONG IRRADIATIONS

R. G. HART, C. B. BIGHAM and M. LOUNSBURY

Atomic Energy of Canada Limited, Chalk River Project, Chalk River, Ontario, Canada

Summary—A number of samples of natural U have been irradiated to various levels, up to 6200 MWD/t, in the NRX reactor at Chalk River for studies of the variation of reactivity with neutron irradiation. Measurements of the reactivity of the samples before and after irradiation have been made in the GLEEP reactor at Harwell. Chemical and mass spectrometric analyses of the samples have been made at Chalk River. A comparison of some of the results with theoretical predictions, based on the latest BNL cross-section data, was presented at the 1959 Annual Meeting of the American Nuclear Society, 15–17 June, by Kushneriuk⁹. A discussion and interpretation of all the results will be given in subsequent papers by D. S. Craig, R. G. Hart, M. Lounsbury, and A. G. Ward. The analytical methods used are discussed in the present paper.

The U was determined gravimetrically by diuranate precipitation and volumetrically by $Ce(SO_4)_a$ titration. These methods are straightforward and are mentioned only briefly. Plutonium was determined by four different methods: spectrophotometric, direct low-geometry α -counting, isotopic dilution with highly irradiated Pu, and isotopic dilution with Pu⁸³⁸. A comparison of these methods is the primary concern of this paper.

Some brief comments are also given on reactivity changes with long periods of irradiation.

INTRODUCTION

THE Canadian approach to atomic power is based on the heavy-water moderated reactor fueled with natural uranium. To achieve economic generation of power in such a reactor, an energy yield of 8000 thermal megawatt days (MWD) or more per ton of uranium must be attained without reprocessing. Consequently, an integral part of the research program at Chalk River has been an examination of the overall credit and debit balance of neutrons over the lifetime of the fuel and a correlation of this balance with theory.

The important changes in the fuel during irradiation are the depletion of U^{235} and the ingrowth of the successive plutonium isotopes 239, 240 and 241, and also the fission products. Such a program, therefore, is inherently dependent upon accurate analytical methods for uranium and its isotopes, and plutonium and its isotopes. In this paper, the determination of uranium will be discussed only briefly because the methods used are well known. The principal topic for discussion will be the analysis of plutonium as the methods we used are unique. The mass spectrometric aspects of the program will not be discussed, outside of outlining some of the results. Those particularly interested in this subject are referred to a paper by Lounsbury in the 1956 issue of the *Canadian Journal of Chemistry*.

URANIUM ANALYSIS

First of all, methods for uranium analysis will be mentioned briefly. These can only be defined as brute force methods as a certain amount of radiation exposure was accepted in order to achieve the accuracy required. The standard gravimetric method¹ involving ammonium diuranate precipitation and ignition to U_3O_8 was used on the lower irradiated samples. The higher irradiated samples were analyzed volumetrically², using ceric sulfate oxidation of U^{IV} to U^{VI} after reduction by means of a Jones reductor. Approximately 500 mg of U was taken for each gravimetric analysis and approximately 50 mg of U was taken for each volumetric analysis. A correction for a small amount of corrosion-product iron was required in both methods. This correction was based on results obtained by colorimetric analysis, using the *o*-phenanthroline method, and normally amounted to a fraction of a per cent.

The precision of a single gravimetric analysis at the 95 per cent confidence level was ± 0.39 per cent, while that of a single volumetric analysis was ± 0.38 per cent at the same confidence level. By following the usual practice of analyzing each sample in duplicate, the mean error was reduced to below 0.3 per cent. Results for some of the samples which were analyzed by both methods are given in Table I. The cross-checks obtained give us confidence that there is no bias in either method.

Slug No.	Total U, mg/ml	Total U, mg/ml	Atom ratio
	(gravimetric)	(volumetric)	U ²³⁵ /U ²³⁸
AGW-1-8 AGW-1-5 AGW-1-9 AGW-1-11 AGW-1-4	$\begin{array}{c} 241.8 \pm 0.7 \\ 238.0 \pm 0.7 \\ 254.8 \pm 0.7 \\ 242.4 \pm 0.7 \\ 277.2 \pm 0.8 \end{array}$	$\begin{array}{c} 241.5 \pm 0.7 \\ 238.9 \pm 0.6 \\ 254.7 \pm 0.7 \\ 242.7 \pm 0.7 \\ 277.7 \pm 0.7 \end{array}$	$\begin{array}{c} (0.7116 \pm 0.0010) \times 10^{-3} \\ (0.6841 \pm 0.0010) \times 10^{-3} \\ (0.6218 \pm 0.0010) \times 10^{-3} \\ (0.5671 \pm 0.0016) \times 10^{-3} \\ (0.4764 \pm 0.0012) \times 10^{-3} \end{array}$

TABLE I.—COMPARISON BETWEEN GRAVIMETRIC AND VOLUMETRIC URANIUM ANALYSES

PLUTONIUM ANALYSIS

All samples were analyzed for plutonium by two different methods, a spectrophotometric method and a direct α -counting method.

The spectrophotometric method was developed by Allison at Chalk River³ and has been used there routinely since that time. An aliquot of dissolver solution containing 1.5-4 mg of plutonium is adjusted to 8M with nitric acid and is then passed through a 2-ml column of 250 to 350 mesh Dowex-1 in the nitrate form. The plutonium is absorbed while the uranium and fission products pass through the column. The column is then washed with 5 column volumes of 10M nitric acid. The 10Mnitric acid is then displaced with 1 column volume of 1M nitric acid and the purified plutonium is eluted with 5 ml of 10-per cent hydroxylamine nitrate. The eluate is is collected in a 5-ml volumetric flask, centrifuged and transferred to a 5-cm spectrophotometer cell. Absorbancy measurements are then made with a Beckman Model DU spectrophotometer at a wavelength of 602 m μ . A schematic diagram of the apparatus is given in Fig. 1. There is always a small loss of plutonium to the aqueous raffinate. This loss is normally of the order of 0.2-0.3 per cent, but can go higher if dissolving conditions are abnormal. The loss is determined by α -counting an Amand Cm-free source and is added to the spectrophotometric result. The precision of a single analysis at the 95 per cent confidence level is ± 1.4 per cent. The usual practice of analyzing each sample in duplicate reduces the error to ± 1.0 per cent. The method is readily adaptable to remote operation. The scheme used at Chalk River is described in the literature.4

The direct α -counting method for plutonium⁵ was developed primarily for the analysis of radial samples taken from a cross section of an irradiated uranium rod.

These samples, taken to provide a knowledge of the variation of cross sections across the diameter of a rod, contain too little plutonium to be analyzed spectrophotometrically. A sample containing approximately 50 μ g of uranium is accurately pipetted onto a stainless-steel source tray. The micropipet used is washed three times with a solution of tetraethylene glycol in dilute nitric acid (four drops of TEG in 10 ml of 1*M* nitric acid). The solution on the source tray is then evaporated to dryness under an infrared lamp and the source tray is ignited to redness. The evaporation must take place very slowly in order to avoid decomposition of TEG by hot nitric acid. Two to

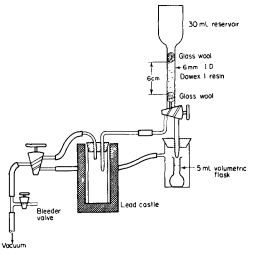


FIG. 1.—Ion exchange apparatus for Method B.

three hours are necessary for the drying. The total α -distintegration rate is measured in a Ward-type low-geometry chamber⁶ and the α -spectrum is measured in a Frisch gridded ion chamber⁷ using an AECL-2230 pulse analyser⁸ and preamplifier described by Harvey *et al.*⁷ A typical spectrum is shown in Fig. 2. From the total count rate, the α -spectrum, a mass analysis of the plutonium and the specific α -activities of Pu²³⁹ and Pu²⁴⁰, the amount of plutonium in the sample can be calculated. The precision of a single analysis at the 95 per cent confidence level is ± 1.9 per cent. The usual practice of analyzing each sample in duplicate and taking the mean reduces the error to ± 1.3 per cent.

The results for a number of samples analyzed by both methods are given in Table II. Again the cross checks obtained give us confidence that there is no bias in either method.

A limited amount of work has been done on two isotopic dilution methods for plutonium; one involves isotopic dilution with Pu^{238} ,⁵ the second isotopic dilution with plutonium rich in $Pu^{240.5}$ The former method is based on an α -count of the purified product while, in the latter method, a mass analysis is made of the product. Since a sufficient number of analyses are not available to establish the precision of these methods in the usual manner, an attempt has been made to arrive at an evaluation of the error by considering all the possible contributing errors. In this manner, it was estimated that the precisions for single analyses at the 95 per cent confidence level is ± 1.8 per cent for the Pu^{238} dilution method and ± 2.2 per cent for the Pu^{240} .

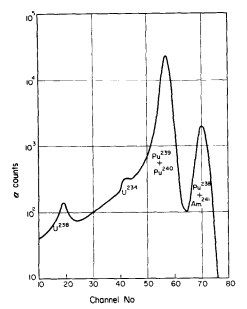


FIG. 2.—Typical a-spectrum of a plutonium sample.

Table II.—Comparison between spectrophotometric and direct α -counting plutonium analyses

Slug No.	Total Pu, mg/l. (spectrophotometric)	Total Pu, mg/l. (direct α-counting)
AGW-1-9	 195 ± 2	194 ± 3
AGW-1-11	295 \pm 3	293 ± 4
AGW-1-4	505 \pm 5	498 ± 6
42-1	455 \pm 5	455 ± 6
42-2	519 \pm 5	529 ± 7
42-3	558 ± 6	567 \pm 7
42-4	517 \pm 5	516 ± 7
42-5	423 ± 4	426 ± 5
42-6	278 ± 3	280 ± 4

dilution method. The cross checks obtained between the four methods on the few samples analyzed by isotopic dilution are shown in Table III. One further point

Table III.—Comparison between spectrophotometric, direct α -counting and isotopic dilution analyses

Slug No.	Total Pu, mg/l. (spectro- photometric)	Total Pu, mg/l. (direct α-counting)	Total Pu, <i>mg/l</i> . (Pu ²³⁸ dilution)	Total Pu, <i>mg/l</i> . (Pu ²⁴⁰ dilution)
AGW-1-5	80.6 ± 0.8		77.3 ± 1.4	79.7 ± 1.6
AGW-1-9	195 \pm 2	194 ± 3	195 \pm 4	(199 \pm 4)
AGW-1-11	295 \pm 3	293 ± 4	301 ± 6	(303 ± 6)
AGW-1-4	505 \pm 5	498 ± 6	514 \pm 10	513 ± 14

Note: ()-isotopic equilibrium not certain.

R. G. HART, C. B. BIGHAM and M. LOUNSBURY

requires mention. We have definite evidence that isotopic equilibrium is not attained immediately after spiking. Sample 4 gave a result of 579 mg/l. when separated shortly after spiking. This factor should be studied in detail before too much faith is put in either of these methods. We have as yet not found time to make this study.

CORRELATION WITH THEORY

Finally, the correlation achieved between experiment and theory will be indicated. Figures 3 and 4 are taken from a paper⁹ presented by Kushneriuk, of our theoretical physics branch, at the 1959 Annual Meeting of the American Nuclear Society, in June of this year. Figure 3 shows the correlation between U²³⁵ burn-up and grams of

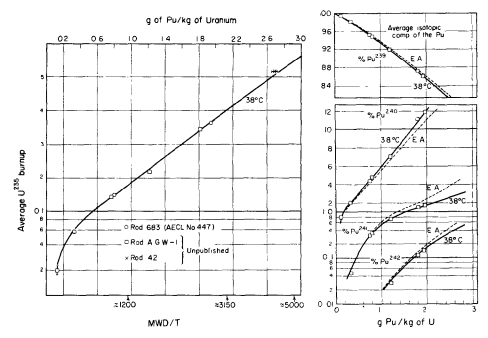


FIG. 3.—Correlation between U²³⁵ burnup and grams of plutonium per kilogram of uranium

FIG. 4.—Correlation between grams of plutonium per kilogram of uranium and the isotopic composition of the plutonium.

plutonium produced per kilogram of uranium. The solid line is calculated, using the latest BNL cross sections for a moderator temperature of 38° C. The slightly high value for the rod 42 point is attributed to a slightly different environment than that assumed in the calculation. Figure 4 shows the correlation between grams of plutonium per kilogram of uranium and the isotopic composition of the plutonium. The dotted curves are elementary approximations based on the assumption of constant cross sections. The solid curves take into account self shielding of the plutonium isotopes. The excellent agreement between experiment and theory gives us confidence that our reactivity predictions contain no very significant unknowns and that our predictions of the burn-up attainable in our power reactors are correct.

REFERENCES

- ¹ N. H. Furman, (Editor) Scott's Standard Methods of Chemical Analysis, Van Nostrand, New York, 1948, pp. 1022, 1024.
- ² R. R. Fraser, Volumetric Methods for the Determination of Uranium, Atomic Energy of Canada, PDB-63, 1952.
- ⁸ G. M. Allison, The Separation of Plutonium by Anion Exchange in 10M Acid Solution-2. Analytical Application to Rod Solution, Atomic Energy of Canada, PDB-87, 1953.
- ⁴ G. M. Allison, A Remote Control Box for the Analytical Separation of Plutonium from Rod Solution, Atomic Energy of Canada, PDB-123, 1954.
- ⁵ R. G. Hart, M. Lounsbury, C. B. Bigham, L. P. V. Corriveau and F. Girardi, *Chemical and Isotopic Analysis of Irradiated Uranium Slug s from Demountable Slug Rod*, Atomic Energy of Canada, CRRP-761, Part. B, 1959.
- ⁶ A. J. Cruikshank, D. J. Littler and A. G. Ward, Atomic Energy of Canada, CRP-378, 1948.
- ⁷ B. G. Harvey, H. G. Jackson, T. A. Eastwood and G. C. Hanna, Can. J. Phys. 1957, 35, 258.
- ⁸ F. S. Goulding, A New Electronic Binary Counter Circuit, Atomic Energy of Canada, 1955, ELI-23.
- ⁹ S. A. Kushneriuk, Calculation of Isotopic Composition of Natural Uranium Fuel Irradiated in the NRX Reactor, Paper Presented at the 1959 Annual Meeting of the American Nuclear Society, 15–17 June, 1959.

THE COLORIMETRIC DETERMINATION OF ZIRCONIUM IN PLUTONIUM-URANIUM-'FISSIUM' ALLOYS*

R. F. BUCHANAN, J. P. HUGHES and C. A. A. BLOOMQUIST, Argonne National Laboratory, Lemont, Illinois, U.S.A.

Summary—This method was developed for the determination of zirconium in plutonium-uraniumfissium' alloys. A sample of the alloy containing $40-100 \ \mu g$ of zirconium in 10N nitric acid is separated from plutonium and from some of the palladium by anion exchange. The effluent is analyzed for plutonium since more than 60 μg will interfere. The ruthenium is removed by fuming with perchloric acid. The uranium and molybdenum do not interfere in reasonable quantities as they exist in this alloy. The alizarin red-S lake is formed in 0.1N nitric acid, and its absorbancy is measured by means of the Beckman Model B spectrophotometer at 520 m μ . The color is stable for several hours and reproducible within 3-5 per cent of the amount present. The interference of the alloying constituents and reagents are given. The time required for duplicate analyses is about $3\frac{1}{2}$ hr, and the equipment and techniques used are standard.

INTRODUCTION

'FISSIUM' (Fs)¹ is the name that has been applied to all the alloys of uranium, plutonium and fission products coming from the work on pyrometallurgical processing.

The problems introduced by starting a reactor cycle on pure uranium or plutonium and then processing and fabricating a slightly different alloy with every cycle are tremendous. As a result, it has been planned to load the reactor with an alloy approaching one of the equilibrium alloys in composition. The change in composition per cycle in this case is slight and changes in properties are expected to be negligible.

A novel feature of the Experimental Breeder Reactor-II (EBR-II) is that the fuel processing is an integral part of the reactor cycle.² The metallurgical refining process and the fabrication methods that will be used involve very short cooling times and incomplete decontamination of the fuel. Chemical methods of processing are much more expensive than the pyrometallurgical processing because they require a longer cooling period, and thus a larger fuel inventory, and much more elaborate processing equipment. After a few cycles in the reactor and through the pyrometallurgical process, the equilibrium alloy will vary according to the original fuel composition. A few typical reference alloys have been set up for test purposes; however, the Pu–U-Fs experimental alloys submitted for analyses have been quite variable, as is shown in Table I.³ The variations in weight per cent of the U–Pu-Fs encountered in our analysis is indicated in the last column of Table I.

Our basic philosophy in analyzing plutonium-bearing samples is to use a method which calls for the smallest sample size as long as the accuracy obtained is adequate for the problem. Consequently, this method with an accuracy of about 3 per cent standard deviation serves the purpose very well for the present state of fuel development with 'fissium' alloys.

It was obvious from some work reported earlier^{4,5} on the analytical application of the nitric acid anion exchange system to plutonium-'fissium' alloys and on the

^{*} Work performed under contract with the U.S. Atomic Energy Commission.

Element	U-'Fissium'	Pu–U-'Fissium'	Variations in Pu–U-'Fissium'
Uranium	95.0	69.2	50-75
Plutonium		20.0	20-40
Zirconium	0.2	0.5	0.05-4.5
Molybdenum	2.5	2.8	1.25-6.0
Ruthenium	1.5	4.3	1.25-6.0
Rhodium	0.3	0.7	0.25-0.9
Palladium	0.5	2.5	0.7 -2.5

TABLE I.— TYPICAL REFERENCE FISSIUM ALLOYS

determination of neptunium, using anion exchange, that the plutonium interference could be removed. In 10N nitric acid, the Pu^{4+} has a distribution coefficient of about 2500 while that for Zr^{4+} is only 0.7.⁵ Some of the palladium is removed and ruthenium, the only other interfering element, can be removed by fuming with perchloric acid after the anion exchange. The complete recovery of the zirconium after fuming was difficult but was found to be quantitative after refluxing with nitric acid.

EXPERIMENTAL

The alizarin lake was found to be stable⁶ in 0.1N nitric acid for several hours if the waiting period after the addition was at least $1\frac{1}{2}$ hr.

If hydrofluoric or sulfuric acid was used in the dissolution of the alloy, complete removal was necessary. If these acids are present, the procedure should be modified. The fuming with perchloric acid after elution from the anion resin column should be continued until the sample has evaporated to dryness. Traces of these acids either inhibited or prevented the formation of the lake. However, the only alloying constituents which did interfere were plutonium and ruthenium. The rhodium, molyb-denum and palladium gave no interference when the ratio of each to zirconium was 1 to 1. The uranium did not interfere when it was 2000 to 1.6

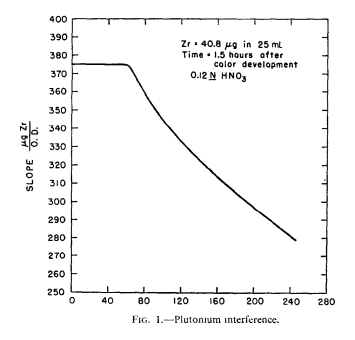
The nitric acid did not affect the slope up to 0.15N but above that the optical density decreased. Mixtures of nitric and hydrochloric acids gave the same slope as long as the total acidity did not exceed 0.15N.

A perchloric acid medium for the development of the alizarin lake would probably work just as well as the nitric acid.⁷

The plutonium content in excess of 60 μ g in a final volume of 25 ml was found to interfere. Excess plutonium caused the slope to decrease as is shown in Fig. 1.

Even though the distribution coefficient for the Pu^{4+} was stated to be about 2500, the decontamination factor varied from 60 to 2500. The erratic decontamination factor may be due to either polymerization or disproportionation in the 10N nitric acid.^{8,9} In all cases, the decontamination was more than adequate to prevent the interference of plutonium (Table II).

The volume of 10N nitric acid necessary for complete elution of the zirconium from a resin column (6 mm \times 6 cm) is less than 25 ml.⁴ An aliquot from a standard solution containing zirconium was eluted from the resin column and compared to a like aliquot from the same solution. Quantitative recovery was indicated by comparison of spectrophotometric measurements.



Some experimental data indicated that the zirconium in 0.1N nitric acid could be accurately determined in the presence of as much as 200 μ g of plutonium if the solution is allowed to stand for at least 12 hr before the addition of sodium alizarin sulfonate.

Sample number	Sample weight in aliquot, mg	Pu, %	Pu in sample, μg	Pu left, μg ,	D.F.
3305-A	1.70	19.4	330	0.7	470
3306	4.90	20.1	985	2.7	365
3307	2.07	20.2	420	5.6	75
3308	1.57	20.7	325	5.3	60
3309-A	1.33	19.3	255	0.1	2550
3310-A	2.58	19.8	510	1.4	365
3311	2.60	41.5	1080	9. 0	120

TABLE II.—Pu DECONTAMINATION BY ANION EXCHANGE—10NHNO3

PROCEDURE

Reagents.

Nitric acid: 10N

Analytical Grade AG 1 × 10 Anion Exchange Resin.* (Processed from Dowex-1 × 10). Moisture 33-39 per cent. Total capacity—3 mequiv./dry gram. Mesh size—200-400. Supplied in the chloride form.

* Supplier: Bio-Rad Laboratories, 800 Delaware, Berkeley, California.

Concentrated perchloric acid (70%). Concentrated nitric acid. Distilled water. Sodium alizarin sulfonate: 0.0125% in H₂O Hydrochloric acid: 6N. Hydrogen peroxide 30%. Hydrofluoric acid: 1N.

The Pu-U-Fs alloy was dissolved in 6N hydrochloric acid, hydrogen peroxide and a drop or two of 1N hydrofluoric acid when necessary. The sample was diluted to a convenient volume and an aliquot containing 40–100 μ g of zirconium was transferred to a small beaker or centrifuge tube. Enough concentrated nitric acid was added to make the solution 10N and it was allowed to stand overnight. The anion column (6 mm × 6 cm) was prepared by washing it alternately with 5 ml portions of 10N nitric acid and water, the final wash solution being 10N nitric acid. The total volume of wash solution was 25–30 ml. The solution of the alloy was transferred to the column by gravity with an elution rate of about 1–2 ml/min. The column was washed with 25–30 ml of 10N nitric acid (using a vacuum) with an elution rate of about 6 ml/ min. The eluant collected was sampled for α -assay to determine the Pu^{239.240} plus Am²⁴¹ content.* If the total plutonium was less than 60 μ g, there was no interference.

The eluant was evaporated to 5–10 ml, cooled, and five drops of concentrated perchloric acid were added. The sample was then fumed in perchloric acid for about 5 min to remove the ruthenium after which it was allowed to cool. Two hundred μ l. of concentrated nitric acid were added and the solution refluxed for about 20–30 min to insure complete dissolution of the zirconium. The solution was cooled, transferred to a 25-ml volumetric flask and diluted to about 15 ml with water. Five millilitres of 0.0125% sodium alizarin sulfonate were added and the solution was mixed. Final dilution to the mark was made with water, following which the contents were mixed again and allowed to stand $1\frac{1}{2}$ hr.

A portion of the solution was transferred to a 1-cm Corex cell, compared to a blank of 0.12N nitric acid containing sodium alizarin sulfonate and the optical density at 520 m μ determined. The concentration was then determined by use of a previously established calibration factor, μg zirconium/O.D. The color remained stable for several hr.⁶

RESULTS AND DISCUSSION

Duplicate test results for a set of four samples ranging from 1.5 to 4.5 per cent zirconium are shown in Table III.

Three standards were prepared from Pu–U-Fs alloys which contained a nominal composition of 0.05–0.15 per cent zirconium. A standard solution of zirconium was added to these alloys to make the percentages equivalent to 2.96, 4.62 and 6.01 zirconium. It was assumed that the nominal composition was correct. The analytical values were biased on the low side (Table IV). Most of the determinations were made about two months after the preparation of the standards. Zirconium is known to hydrolyze readily even in strong acid solutions. In view of this, it is desirable to make the zirconium determinations within a few days of the dissolution of the sample.

• The Am²⁴¹ is the daughter of Pu²⁴¹ which is normally found in the plutonium that we have used.

Sample number	Zr, % (nominal composition)	Zr, % (found)	Average Zr, % (found)
3306	1.5	1.46 1.51	1.48
3307	3.0	2.96 2.88	2.92
3308	4.5	4.51 4.40	4.46
3311*	3.0	2.87 2.93	2.90

TABLE III.-DUPLICATE TEST RESULTS FOR UNKNOWN SAMPLES

*Sample number 3311 contained 40 per cent plutonium in the Pu-U-Fs alloy while the others contained only 20 per cent plutonium.

Sample number	Zr added, %	Zr recovered, %	Average Zr found, %	Std. Dev., %	Error, %
3305A	4.62	4.72 4.50 4.50 4.32 4.49	4.51	3.6	-2.4
3309A	6.01 ?	5.42 5.76 5.64	5.61	3.1	-7.1
3310A	2.96	3.00 2.88 2.88	2.92	2.4	-1.4

TABLE IV.—Zr ANALYSES ON STANDARD Pu–U-FS ALLOYS

REFERENCES

- ¹ L. J. Koch et al., Experimental Breeder Reactor II (EBR-II)Hazard Summary Report, U.S. Atomic Energy Commission, Argonne National Laboratory, ANL-5719, p. 47, May 1957.
- ² L. J. Koch, *The Engineering Design of EBR-II*, American Institute of Chemical Engineers, Chemical Engineering Progress Symposium, Vol. 52, No. 19, pp. 37–47, 1956.
- ³ L. R. Kelman, Fast Reactor Fuel Development at Argonne National Laboratory, U.S. Atomic Energy Commission, Argonne National Laboratory, ANL-FGF-73, p. 13, November 1957.
- ⁴ R. F. Buchanan, J. P. Faris, K. A. Orlandini and J. P. Hughes, The Analytical Application of the Nitric Acid Anion Exchange System to Plutonium-Fissium and Plutonium Binary Alloys Used in Metallurgical Studies, U.S. Atomic Energy Commission, TID 7560, p. 179, January 1959.
- ⁵ F. P. Roberts, An Analytical Method for Neptunium-237 Using Anion Exchange, U.S. Atomic Energy Commission, Hanford Atomic Products Operation, HW-59032, January 1959.
- ⁶ Ruth Gunther and Richard H. Gale, *Determination of Zirconium in the Presence of Uranium:* A *Spectrophotometric Method*, U.S. Atomic Energy Commission, Knolls Atomic Power Laboratory, KAPL-305, March 1950; *Anal. Chem.* 1950, **22**, 1510.
- ⁷ D. L. Manning and J. C. White, Anal. Chem. 1955, 27, 1389.
- 8 Arthur Brunstad, Ind, Eng. Chem. 1959, 51, 38.
- ⁹ J. L. Ryan and E. J. Wheelwright, Ind. Eng. Chem. 1959, 51, 60.

ANALYTICAL PROGRAM FOR PROCESSING STAINLESS STEEL–URANIUM DIOXIDE REACTOR FUEL ELEMENTS*

G. A. HUFF

Phillips Petroleum Company, Atomic Energy Division, Idaho Falls, Idaho, U.S.A.

Summary—Analytical programs have been described for uranium recovery processes for aluminumuranium and zirconium–uranium reactor fuel elements. It is the purpose of this paper to describe the analytical program for the uranium recovery process involving stainless-steel–uranium-reactor fuel elements.

The recovery process consists of a two-step dissolution and the regular liquid-liquid solvent extraction using TBP in kerosene. Sulfuric acid is used to dissolve the stainless steel. As a result of the sulfuric acid dissolution, the uranium is converted to a slurry. The uranium slurry is then dissolved in nitric acid. After the nitric acid dissolution, the feed material is compatible with the extraction columns used at the Idaho Chemical Processing Plant for other type fuel elements previously described.

In order to analyze samples of dissolver solutions for constituents necessary for plant operation, three new analytical methods were developed. These methods are a rapid method for determining uranium, one for nitrate and one for sulfate. Methods are described for uranium, acidity, specific gravity, nitrate and sulphate in the dissolver solution, and for small amounts of uranium in the extraction column raffinates. Samples of dissolver solution and first cycle aqueous raffinate contain large amounts of fission products. These samples require shielded facilities for analytical work. The remote analytical facility described by Dykes and his co-workers was used for this work.

Precision and accuracy data are given for the analytical methods described.

INTRODUCTION

ANALYTICAL programs have been described for uranium recovery processes for aluminum-uranium¹ and zirconium-uranium² reactor fuel elements. It is the purpose of this paper to describe the analytical program for the uranium recovery process of stainless-steel-uranium dioxide reactor fuel elements.

Figure 1 shows a simplified flowsheet of the recovery process used. The scheme consists of a two-step dissolution and the conventional liquid-liquid solvent extraction

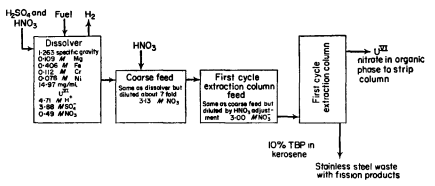


FIG. 1.--Stainless-steel fuel dissolution and extraction column feed preparation.

* Work done under Contract AT(10-1)-205 to the U.S. Atomic Energy Commission.

G. A. HUFF

using tributylphosphate in kerosene. Sulfuric acid is used to dissolve the stainless steel. As a result of the sulfuric acid dissolution, the uranium dioxide is converted to a slurry. The uranium dioxide slurry is then dissolved in nitric acid. After the nitric acid dissolution, the feed material is compatible with the extraction columns used at the Idaho Chemical Processing Plant for other type fuel elements previously described.

SAMPLES ANALYZED

The more important routine samples, constituents determined and analytical techniques are summarized in Table 1.

In order to analyze samples of dissolver solutions for constituents necessary for plant operation, three new analytical methods were developed. These methods are a rapid method for determining uranium, one for nitrate and one for sulfate.

Samples of dissolver solution and first cycle aqueous raffinate contain large amounts of fission products. These samples require shielded facilities for analytical work. The remote analytical facility described by Dykes *et al.*³ was used for this work.

ANALYTICAL METHODS

Uranium in dissolver, coarse feed, first cycle extraction column feed and first cycle extraction column product

From the composition listed in Table 1, it can be seen that these samples, with exception of the first cycle extraction column product, contain constituents which interfere with analytical procedures. Liquid-liquid solvent extraction has been used successfully in separating uranium from fission products and other interfering impurities. After separation, samples readily lend themselves to the isotopic dilution mass spectrometric method. This technique was proposed by Ingram⁴ and further developed for uranium by Duffy and Tingey⁵, and Goris and Tingey.⁶ This is an excellent method since quantitative recovery from the extraction is not required and concentration as well as isotope ratio are obtained at the same time. This method is used for the plant input measurement.

The separation procedure is as follows.

To a test tube is added 0.5 ml of a 1.0 mg/ml uranium-233 standard solution and 4.0 ml of $1N^{B}$, 2.0*M* aluminum nitrate salting solution. The test tube is transferred to the remote facility (Figs. 2 and 3) where 0.5 ml of the sample is added from a remote pipetter (Figs. 4 and 5). This pipetter was discussed at the Second Conference on Analytical Chemistry in Nuclear Reactor Technology in 1958. Details on the apparatus are given by Dykes.⁷ The test tube is then stoppered and the aqueous solution is mixed by shaking to obtain homogeneity. A manipulator (Fig. 6) is used to shake the tube. Two milliliters of organic extractant, hexane, are added and the mixture is stoppered and shaken for 1 min. As much of the organic as possible is poured into a clean centrifuge tube. The tube is transferred from the remote facility to the conventional laboratory. The uranium is simultaneously stripped from the organic phase and precipitated as the peroxide by contacting with 1 ml of 30% hydrogen peroxide.

After centrifuging and decanting, the precipitate is washed with 1 to 1 wateracetone mixture and transferred to the Mass Laboratory.

The peroxide is dissolved in a minimum amount of nitric acid and evaporated on a tantalum filament. Details of the operation of the mass spectrometer and of the

Analytical program for processing fuel elements

Sample	Composition	Constituent determined	Analytical technique
Dissolver	$\begin{array}{cccc} \text{Specific gravity} & 1 \cdot 263 \\ \text{Mg} & 0 \cdot 109M \\ \text{Fe} & 0 \cdot 406M \\ \text{Cr} & 0 \cdot 112M \\ \text{Ni} & 0 \cdot 078M \\ \text{U} & 14 \cdot 97 \text{ g/l. U}^{\text{VI}} \\ \text{H}^+ & 4 \cdot 71M \\ \text{SO}_4^{3-} & 3 \cdot 08M \\ \text{NO}_3^- & 0 \cdot 496M \end{array}$	Uranium Uranium Nitrate	Isotope dilution Mass spectrometer Spectrophotometric Spectrophotometric
Dissolver wash	$H_{2}O < 0.01M \text{ NO}_{3}^{-1}$	Nitrate	Acid-base titration and spectrophotometric
Coarse feed	As dissolver, but diluted about seven-fold and 3.13 <i>M</i> NO₃ ⁻	As dissolver plus Specific gravity Acidity Sulfate Fission products	Falling drop pH titration with oxalate complexing Amperometric Total β - and γ -counting
First cycle extraction column feed	As coarse feed but diluted by nitric acid adjustment NO ₃ - 3.00M	Nitrate	Spectrophotometer
First cycle extraction column raffinate	As first cycle feed but with less than 1×10^{-3} mg/ml of U ^{VI}	Uranium	Pellet fluorophotometric
First cycle extraction column product	1 <i>M</i> U ^{VI} nitrate 0.81 <i>M</i> HNO ₃	Uranium Specific gravity Fission products	Isotope dilution Mass spectrometer Falling drop Total β and γ

TABLE I IMPORTANT ROUTINE SAMPLES	S, CONSTITUENTS DETERMINED AND ANALYTICAL TECHNIQUES
-----------------------------------	--

determination are given by Duffy *et al.*⁸ Results are reported for total uranium concentration and percentages of each isotope.

The isotopic dilution mass spectrometric method is time consuming and is not a method to be used if production schedules are critical. In order to obtain rapid data for uranium concentration, a spectrophotometric method was developed. The procedure is as follows.

To a test tube is added 6 ml of 2.8M aluminum nitrate $(2N^{B}, 0.025M$ in TPAN)

and 3 ml of hexone. The tube is transferred to the remote line where 0.750 ml of sample is added. A polyethylene stopper is used to stopper the tube and it is transferred to the special tumbler (Fig. 7) and mixed for 3 min. As much of the organic as possible is poured off and transferred to the conventional laboratory. The sample is centrifuged and as much of the organic as possible is transferred to a 1-cm Corex absorption cell. The spectrum of the material is scanned from 500 to 400 m μ on a Cary Model 14 recording spectrophotometer at a scan speed of 5 A/sec.

Details on this method are given by Maeck et al.9

Uranium in first cycle extraction column raffinate.

These samples are very low in uranium ($<1.0 \times 10^{-3}$ mg U/ml) and pellet fluorophotometry is used to analyze them. They contain large amounts of fission products which create radiation problems. Also, they contain foreign ions which quench fluorescence. In order to separate the uranium from fission products and other interfering ions, a liquid-liquid extraction is made in the Remote Analytical Facility. The uranium, which is contained in the organic phase, is transferred to the conventional laboratory where pellets are prepared and the fluorescence measured.

The procedure is as follows.

To a test tube is added 0.5 ml of sample and 1.5 ml of solution, 1.5M in aluminum nitrate, 5 per cent in citric acid, 1M in nitric acid and 2 ml of 1 per cent tributylphosphate in n-hexane. After extracting for 1 min, as much of the organic as possible is transferred to the conventional laboratory. Sodium fluoride pellets are prepared in platinum dishes and 0.2 ml aliquots are pipetted onto each pellet. The pellets are dried under infrared lamps, fused in a gas-air flame and the fluorescence is measured in a high-sensitivity fluorophotometer. Uranium concentration is then read from a working curve prepared from processed standards.

Acid in coarse feed

The coarse feed sample contains hydrolyzable ions which require complexing before acidity is measured. A pH titrimetric method is used to determine free acid (or base) in solutions of this nature. Oxalate, as the potassium salt, is used as the complexing agent. The potassium oxalate is adjusted to a reference pH of 5.80. This pH minimizes hydrolysis in a stainless-steel matrix. The method is described by Booman *et al.*¹⁰

The procedure is as follows.

Fifteen ml of an 8% potassium oxalate solution adjusted to a pH of 5.80, is added to a 50-ml beaker. If the sample is known to be nearly neutral or slightly basic, an acid "spike" is added. The beaker is transferred to the Remote Analytical Facility where 0.200 ml of sample is added by the pipetter. The beaker is then transferred to a titration apparatus (Figs. 7 and 8) and allowed to digest for 2 hr. This digestion time is required for chromium to complex. After digestion, the sample is titrated with standard 0.2N sodium hydroxide until the exact reference pH is obtained. Acidity is calculated from the base titer, sample volume and normality of acid "spike".

Specific gravity in coarse feed and first cycle extraction column product

These samples contain fission products; therefore, they require remote handling. Specific gravity is determined by the falling drop method. A small, known volume



FIG. 2.—Operating aisle of remote laboratory.



FIG. 3.-Remote line.

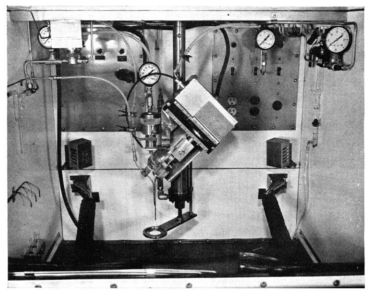


FIG. 4.—Remote analytical facility Model B pipet box installation.



FIG. 5.-Remote analytical facility Model B pipetter.

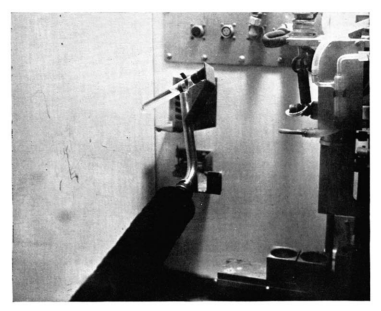


FIG. 6.-Remote analytical facility manual extraction apparatus.

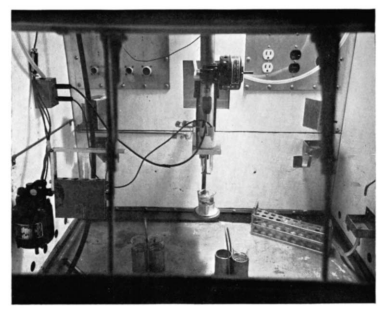


FIG. 7.-Remote analytical facility mechanical extraction apparatus.

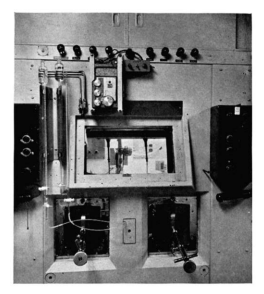


FIG. 8.-Acid titrator.

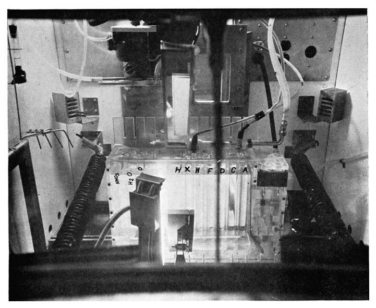


FIG. 9.—Falling drop specific gravity apparatus box installation.

.....

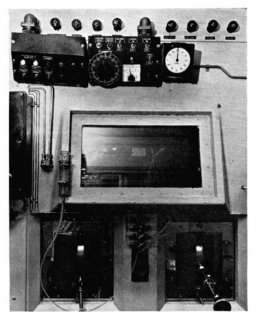


FIG. 10.-Falling drop specific gravity apparatus.

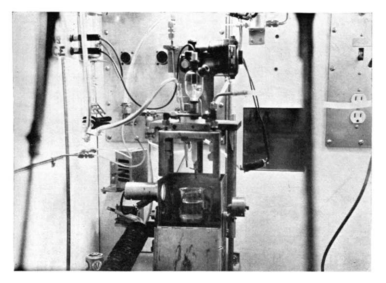


FIG. 11.-Remote sulfate titrator box installation.

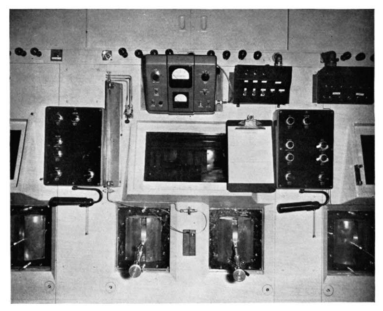


FIG. 12.-Remote sulfate titrator.

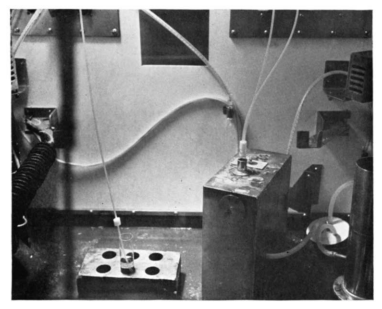


FIG. 13.-Remote colorimetric nitrate box installation.

.

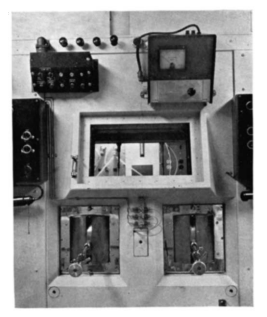


FIG. 14.-Remote colorimetric nitrate apparatus.

of sample is dropped through an immiscible organic liquid (a mixture of o-nitrotoluene, α -chloronaphthalene and α -bromonaphthalene) and the time for the drop to travel a known distance is determined. The remote apparatus used for this measurement is shown in Figs. 9 and 10.

The procedure requires that a 0.005 ml drop be formed on the pipet tip while submerged in the liquid. The drop is released as the pipet tip breaks the surface of the liquid as it is withdrawn. The time for the drops to fall a measured distance is obtained. An average time for five drops is obtained. Specific gravity is then read from a working curve prepared from known standards.

Total β - and γ -activity in coarse feed and first cycle extraction column product

Gross β - and gross γ -activities are reported on these samples as a complete fission product analysis is costly and time consuming. Activities from one stream are compared to the other in order to determine the effectiveness of the decontamination process. Dilutions of these samples from 100 to 10,000 fold are prepared using 0.1N HNO₃ as a diluent. Original dilutions are prepared in the Remote Analytical Facility and transferred to the conventional laboratory for further work.

The gross beta measurement is very simple. An aliquot of the diluted sample is pipetted onto a 1-in. watch glass and evaporated to dryness under infrared lamps. The watch glass is covered with cellophane tape and mounted on a paper card. The sample is counted on a proportional counter at 10 per cent geometry and reported as counts per minute per ml.

The entire dilution is used in the gross gamma measurement. The sample is deposited into the thimble of a high-pressure ion chamber and read at 100 per cent geometry. Since the sample contains a large number of gamma energies and decay schemes are unknown, results are reported in terms of radium equivalents per ml.

Sulfate in coarse feed

Since sulfuric acid is used in the first dissolution step, sulfate data are necessary in obtaining material balance checks over the feed end. A rapid titrimetric method is used. The method involves the amperometric titration of sulfate ion with lead nitrate at -1.2 V versus the saturated calomel electrode with a dropping mercury cathode. The technique was proposed by Elving and Grodzka¹¹ and further developed by the Analytical Development Group at the Idaho Chemical Processing Plant. Remote apparatus for this method is shown in Figs. 11 and 12.

The procedure is as follows.

To a 150-ml beaker is added 10 ml of absolute alcohol. A volume of distilled water is added to give a total aqueous phase of 25 ± 1 ml. This final volume also includes the sample aliquot to be added. Five drops of bromothymol blue indicator and a stirring bar are added. An aliquot of sample containing from 0.25 to 0.65 millimoles of sulfate is added to the beaker. The beaker is placed on a magnetic stirrer and 0.5N sodium hydroxide is added to the bromothymol blue end point (yellow to blue). The sample is scrubbed with nitrogen for 5 min. After scrubbing, the sample is swept with nitrogen during titration. The sample is then titrated with standard lead nitrate solution using 0.5 ml increments. Data are plotted on coordinate graph paper—galvanometer readings versus volume of lead nitrate. The best straight line is constructed through points on either side of the break. The intercept of the two lines is the end point. Molarity of the sample is determined from the lead nitrate titer and sample volume.

Nitrate in dissolver, dissolver wash and first cycle extraction column feed

In order to dissolve stainless-steel-uranium-fuel elements with sulfuric acid, it is necessary to have the dissolver tank free of nitrate. Nitrate renders the elements passive to sulfuric acid. After the nitric acid dissolution step, the dissolver is washed with water until the nitrate concentration is <0.01M. In order to determine the nitrate content of the dissolver, a sample of the wash material is analyzed for total hydrogen ion. A simple, rapid, acid-base titration using phenolphthalein as an indicator is the method involved. If an answer of <0.01M is obtained, the dissolver is assumed to be clean. If an answer of >0.01M is obtained, it may be due to hydrogen ion indicating nitrate content or to hydrolyzable ions being titrated with base. In this case, a more accurate check of nitrate content is obtained by the colorimetric determination of nitrate. The acid-base titration is generally a sufficient check.

The colorimetric determination of nitrate is the method used to determine nitrate concentration on dissolver and first cycle extraction column feed material. This method is dependent upon the purple complex of ferrous nitrosyl sulfate. The technique is one reported by Feigl¹², English¹³, Elving and Grodza¹⁴, and further developed by the Analytical Development Group at the Idaho Chemical Processing Plant. The Remote Colorimeter is shown in Figs. 13 and 14.

The procedure is as follows.

To a 50-ml glass centrifuge tube is added 1 ml of ferrous sulfate solution. Twentyfive ml of 15M sulfuric acid is added to the tube. A volume of distilled water is quantitatively added to give a total aqueous phase of 29.0 ml. This final volume should include the sample aliquot to be added later. The tube is then cooled in an ice bath until its content is less than 5° C. While the tube is still in the ice bath, an aliquot of sample containing from 0.007 to 0.16 millimoles of nitrate is added. The tube is removed from the ice bath and the color is allowed to develop. The absorbance of the sample is read on a spectrophotometer at a wavelength of 530 m μ . Because of the transient nature of the color, the absorbance measurement is made between 35 and 45 min after removal of the tube from the ice bath. Millimoles of nitrate corresponding to the sample absorbance is read from a standard curve.

QUALITY CONTROL AND RELIABILITY OF THE METHODS

A quality control program similar to those described for the analytical programs on aluminum-uranium¹ and zirconium-uranium¹⁶ fuel processing was maintained on the analytical program for stainless-steel-uranium-fuel processing. Also, all analysts were trained and tested on the methods used according to the training and testing program described by Huff and Tingey.¹⁷

Bench standards were used to check performance of apparatus and reagent purity from shift to shift. Control samples for each method were submitted to the laboratory and analyzed at a rate of 8 per method per month. Where practical, bench standards and controls were prepared in matrices similar to process samples.

The reliability data listed in Table II were calculated from the results of control samples.

Analysis	Range	Precision ^(A)	Bias ^(B)
Uranium concentration by isotopic dilution mass spectrometry	1.8 mg/ml	0.04 mg/ml	None
Uranium isotope Distribution	50 % U-235 1–6 % U-236	0.08 % ^(C) 0.02 % ^(C)	None None
Uranium, spectrophotometric	2.20-14.80 mg/ml	0.04-0.18 mg/ml	+0.03-0.04 ±0.03 mg/ml
Uranium extraction, fluorophotometric	$8.0 imes10^{-5}$ - $9.0 imes10^{-2}$ mg/ml	37.48 %	None
Nitrate, spectrophotometric	0.008-3.0 <i>M</i>	0.016 <i>M</i>	None
Specific gravity, falling drop	1.15 to 1.34	0.006	None
Acidity—pH Titration with oxalate complexing	1.44 N ^B -4.44N ^A	0.10 <i>N</i>	$0.05 \pm 0.02N$
Sulfate, amperometric	2.60-5.80M	0.21 <i>M</i>	None

TABLE II .--- PRECISION AND ACCURACY OF METHODS UNDER ROUTINE OPERATING CONDITIONS

Note: (A) Precision is expressed as 95 per cent limits of uncertainty for a single determination.

(B) Precision of the bias is also expressed as 95 per cent limits of uncertainty.

(C) Expressed in absolute percentage units.

REFERENCES

- ¹ R. C. Shank and J. E. Rein, U.S. Atomic Energy Commission, IDO-14412, 1957.
- ² J. E. Rein and R. C. Shank, U.S. Atomic Energy Commission, IDO-14413, 1957.
- ⁸ F. W. Dykes, R. D. Fletcher, E. H. Turk, J. E. Rein and R. C. Shank, Anal. Chem. 1956, 28, 1084.
- ⁴ M. G. Inghram, J. Phys. Chem. 57, 809, 1953.
- ⁸ W. E. Duffy and F. H. Tingey, U.S. Atomic Energy Commission, IDO-14301, 1955.
- ⁶ P. Goris and F. H. Tingey, U.S. Atomic Energy Commission, IDO-14366, 1955.
- ⁷ F. W. Dykes, U.S. Atomic Energy Commission, TID-7568 Part 2, 65, 1959.
- ⁸ W. E. Duffy, G. V. Wheeler and T. D. Morgan, U. S. Atomic Energy Commission, IDO-14318, 1955.
- ⁹ W. J. Maeck, G. L. Booman, M. C. Elliott and J. E. Rein, Anal. Chem. 1959, 31, 1130.
- ¹⁰ G. L. Booman, M. C. Elliott, R. B. Kimball, F. O. Cartan and J. E. Rein, *Anal. Chem.* 1958, 30, 284.
- ¹¹ P. J. Elving and P. Grodzka, Control Indices and Analytical Procedures, University of Michigan Progress Report No. 12, U.S. Atomic Energy Commission Contract No. AT (10-1)-733, Project 2121-1, February 1954.
- ¹² F. Feigl, Spot Tests (3rd Ed.) Elsevier, New York, 1947, p. 243.
- ¹⁸ F. L. English, Anal. Chem. 1947, 19, 850.

¹⁷ G. A. Huff and F. H. Tingey, U.S. Atomic Energy Commission, IDO-14388, 1956; G. A. Huff and F. H. Tingey, *Anal. Chem.* 1957, **29**, 19A-22A.

¹⁴ Ibid¹¹

¹⁸ Ibid¹

¹⁶ Ibid⁹

A SIMPLIFIED METHOD FOR RADIOIODINE ANALYSIS*

H. M. EILAND

General Electric Company, Knolls Atomic Power Laboratory, Schenectady, N.Y., U.S.A.

Summary—A radiochemical method of analysis for gross iodine activity in the primary coolant of pressurized water reactors has been developed at K.A.P.L. It is based upon the rapid isotopic exchange which occurs under suitable conditions between fission product iodine and the iodine atoms in preformed silver iodide. The principal steps in the procedure are the following: (a) about 20 ml of the slightly basic coolant is oxidized with sodium hypochlorite. Lanthanum hydroxide is formed by adding lanthanum nitrate to the solution, (b) the lanthanum hydroxide scavenger is removed by filtration, (c) the solution is contacted with preformed silver iodide, (e) the silver iodide is removed by filtration, washed, mounted and counted. The procedure has been found to give excellent decontamination from fission products and activated corrosion products which occur in the primary coolant of pressurized water reactors. The recovery of iodine has been shown to be 88 ± 5 per cent. During the past year, the procedure has been used extensively at K.A.P.L. for applications where speed and simplicity are of greater importance than highest accuracy.

A RADIOCHEMICAL analysis for total iodine activity in the primary coolant of pressurized water reactors must be performed frequently. This analysis serves two purposes. First, it serves as a basis for determining whether or not the I¹³¹ content of the water exceeds the limit allowed for safe disposal as radioactive waste; Second, it serves as a monitor for detecting fuel element rupture in the reactor.

The choice of a procedure to be used in the iodine analysis was affected by several factors. The use of solvent extraction was severely restricted, for example, because volatile organic solvents, which might constitute a potential hazard in a closed environment, could not be used. Other items common to radiochemical practice, such as an analytical balance and a vacuum line, were not available. The equipment used in the analysis had to be durable and simple, and the procedure itself had to be relatively uncomplicated so that it could be performed by personnel with little training in analytical techniques. Finally, the radioactivity measurement was restricted to β -counting on a G-M counter.

The isotopic exchange approach to this analysis was suggested by Sunderman and Mienke.¹ They found that carrier-free silver-110 could be separated from a large number of other activities by isotopic exchange with silver chloride. It seemed reasonable to expect, therefore, that carrier-free radioiodine would exchange with silver iodide or some other insoluble iodide salt.

In order to test this hypothesis, several tracer experiments were conducted with carrier-free I¹³¹ from Oak Ridge.

In Table I, results are given of an experiment in which solutions of carrier-free I^{131} tracer were contacted with Agl and Cu_2I_2 . The pH of the solution was adjusted with HCl and NaOH. Silver and cuprous iodide were chosen because these salts are easily prepared and they are quite insoluble. It can be seen that the maximum exchange

* Operated for the U.S. Atomic Energy Commission by the General Electric Company under Contract No. W 31 109 Eng 52.

A simplified method for radioiodine analysis

occurred with AgI in strongly acidic solution. Exchange with Cu_2I_2 was only 85 per cent complete at best, and it was quite sensitive to the pH of the solution.

pH of solution	Exchangeable I ¹³¹ on AgI, %	Exchangeable I ¹⁸¹ on Cu ₂ I ₂ , %
1	95	16
3	85	84
5	65	36
9	62	32
11	50	19

TABLE I.-ISOTOPIC EXCHANGE WITH AGI AND CU2I2

Note: exchange time = 3 min.

Table II shows the results of another tracer experiment in which carrier-free solutions of I^{131} were contacted with AgI for varying periods of time. The solutions were 0.1M in HCl. Maximum exchange was attained within the first 3 min.

TABLE II.---EFFECT OF CONTACT TIME ON

Exchangeable [¹³¹ on AgI, %
80
94
100
96
104

The procedure was then tested on freshly prepared solutions of fission products. It was soon evident that any procedure based upon isotopic exchange with AgI would require a step to convert all the radioiodine to a chemical form exchangeable with AgI. Furthermore, a decontamination step would be required to prevent other activities being carried on the AgI during the isotopic exchange step.

The conventional method of converting radioiodine to an exchangeable form consists of hypochlorite oxidation in alkaline solution, followed by bisulfite reduction in acidic solution. This method was found to be satisfactory for our purposes and was adopted for use in the present method.

Scavengers tested for use in the decontamination step were $Fe(OH)_3$, $La(OH)_3$, $Al(OH)_3$ and filter paper pulp. When iron hydroxide was used, the recovery factors for iodine were extremely low. This was attributed to the fact that ferric iron oxidizes iodide ion to molecular iodine. Lanthanum hydroxide appeared to be more effective than either aluminum hydroxide or filter paper pulp as a scavenger. Consequently, a La(OH)₃ scavenging step was combined with the oxidation-reduction cycle in the procedure.

As mentioned earlier, the principal aim of this work was to develop a simple, reliable procedure utilizing a minimum of rugged laboratory equipment. Highest

H. M. EILAND

accuracy and precision were not of paramount importance. The procedure which is diagrammed in Fig. 2 and outlined in detail in the Appendix was also designed to accomplish this with limited laboratory facilities.

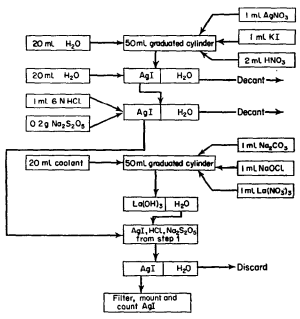


FIG. 2.-Block diagram of procedure.

The first step of the procedure consists of preparing about 50 mg of silver iodide for use in the exchange reaction. This is done by mixing the appropriate amounts of AgNO₃ and KI in a 50-ml graduated cylinder. A few ml of HNO_3 will hasten coagulation of the precipitate. The AgI is washed once with water by decantation. Sodium metabisulfite and HCl are added to the graduated cylinder containing the precipitate and it is set aside for later use.

The second step of the procedule is a combination of the oxidation-reduction cycle and the scavenging step. Twenty milliliters of the primary coolant is measured into a 50 ml graduated cylinder. Sodium carbonate, sodium hypochlorite and lanthanum nitrate are added in that order. Lanthanum hydroxide forms in the solution and the mixture is shaken for 1 min.

In the third step of the procedure, the $La(OH)_3$ scavenger is removed by filtering the mixture from step 2. A short-stem funnel containing a fast filter paper (Whatman No. 41-H) is placed on the graduated cylinder from step 1 which contains the AgI precipitate. The scavenged solution from step 2 is poured into the funnel. The HCl in the funnel acidifies the solution, and the sodium metabisulfite reduces the radioiodine to iodide.

In the fourth step, the isotopic exchange step, the cylinder containing the treated coolant and the AgI precipitate is shaken frequently during a 4-min interval. Finally, the AgI precipitate, which now contains the radioiodine, is collected on a 24-mm glass fiber filter disc, dried under a heat lamp and mounted on a card for counting.

The filtration apparatus used to collect the AgI precipitate is shown in Fig. 1.

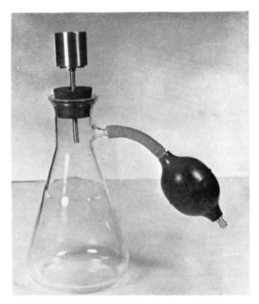


FIG. 1.-Filtration apparatus.

It consists of a stainless-steel filter chimney available from Tracerlab and a suction flask which can be partially evacuated by means of the rubber exhaust bulb.

This procedure has been tested by representatives from K.A.P.L. and the Westinghouse Bettis plant with coolant from a prototype reactor operated by K.A.P.L. Each of 22 coolant samples was analyzed two or more times by the isotopic exchange method and once by a conventional CCL₄ extraction procedure² for which the carrier recovery was measured. Five different analysts were used to perform the isotopic exchange analyses. The efficiency of the isotopic exchange procedure for each analysis was calculated on the basis of the β -count rate of the AgI precipitate obtained by CCl₄ extraction after correction for carrier recovery. The efficiency values ranged from 77–97 per cent in this test. The average value was 88 per cent; the standard deviation was 5 per cent.

It can be said in summary that the method described here is satisfactory for determining gross iodine activity in the primary coolant of a pressurized water reactor. It can be used for radioiodine analyses where simplicity and speed are of greater importance than highest accuracy and precision.

APPENDIX

Stepwise procedure

(1) Prepare a precipitate of AgI by pipetting into a 50-ml graduated cylinder (with ground glass stopper) 1 ml each of $0.3M \text{ AgNO}_3$ and 0.25M KI solutions. Add 20 ml of deionized water and 2 ml of $6M \text{ HNO}_3$. Shake for 10 sec and allow to stand until AgI settles. Pour off the solution and add 20 ml of water to the graduated cylinder containing the AgI. Shake again for 10 sec and allow the AgI to settle. Discard all but about 2 ml of the water. A loss of 10-20 per cent of the AgI during this step of the procedure is allowable.

(2) Add 1 ml of 6N HCl and 0.2 g of Na₂S₂O₅ to the graduated cylinder containing the AgI and set aside for later use.

(3) Measure 20 ml of coolant into a clean 50-ml graduated cylinder and add the following reagents:

1 ml of $2M \operatorname{Na_2CO_3}$ solution 1 ml of NaOCl solution (5 per cent available Cl) 1 ml of $0.1M \operatorname{La}(\operatorname{NO_3})_3$ solution

Shake for 15 secs.

(4) Place a 90-mm filter paper (Whatman No. 41-H) in a short-stemmed funnel and rinse down the sides of the paper once with water. After the water has drained out, place the stem of the funnel in the top of the cylinder containing the AgI so that the mouth of the cylinder supports the funnel.

(5) Carefully pour the treated coolant into the filter paper, allowing the filtrate to drain into the cylinder containing AgI. After all of the solution has been filtered, rinse down the sides of the filter with water. When the filter has drained, remove the filter paper and allow any solution in the stem of the funnel to run into the cylinder.

(6) Place the stopper on the graduated cylinder and shake intermittently for 4 min to assure a good dispersion of the AgI in the solution.

(7) Place a 24-mm fiber glass filter in the stainless-steel filtration apparatus and fill the chimney with the freshly shaken mixture of coolant and AgI. Apply suction by rapidly squeezing and releasing the rubber bulb of the filtration apparatus. Continue H. M. EILAND

refilling the chimney and filtering until all of the AgI has been collected on the filter. A few ml of water can be used to transfer the last traces of AgI from the cylinder into the chimney. The walls of the chimney should be washed down with a few ml of water from the wash bottle. Excessive washing of the precipitate should be avoided.

(8) Maintain suction on the filter for 30 sec after all visible water has run out of the chimney and then remove the chimney with a twisting motion. Remove the filter carefully with the flat tipped tweezers and dry it under the heat lamp for 4 min. Place the filter in a nickel counting planchet, and β -count under the G-M counter.

REFERENCES

¹ D. N. Sunderman and W. W. Mienke, Anal. Chem. 1957, 29, 1578.

^a E. D. Coryell and N. Sugarman, *Radiochemical Studies; The Fission Products*, McGraw-Hill, New York, 1951, pp. 1625–1628.

RADIOCHEMICAL ANALYSIS OF REACTOR EFFLUENT WASTE MATERIALS AT HANFORD*

R. W. PERKINS

Hanford Laboratories Operation, General Electric Company, Richland, Washington, U.S.A.

Summary—Analytical procedures are described which were found to be suitable for the measurement of radioisotopes in effluent water and in erosion products deposited from effluent water. The procedures have been applied in studying the changes in concentration of radioisotopes in effluent water following reactor start-up and in identifying and measuring the radioisotopes deposited on metal surfaces during operation.

The increase in concentration of nine radioisotopes in the effluent water from a chemically cleaned (purged with Turco 4306B) process tube has been measured. The concentrations of all radioisotopes studied in the effluent from the purged tube were, with the exception of Na²⁴, less than those removed from the control tube during the first two days of operation by factors of 5 to 100; several of the radioisotopes, including P³³ and As⁷⁶, were less than 25 per cent of their normal value 2 weeks after the purge. It was demonstrated by means of this study that radioisotope concentrations increased rather slowly in the effluent water during the 1½ month period of the test.

The relative amounts of the sixteen radioisotopes in the effluent water, absorbed during a 35 day exposure on Al, stainless steel and carbon steel surfaces, downstream from both a normal tube and a chemically cleaned tube were measured. The amounts of the majority of the radioisotopes on the Al and stainless-steel surfaces downstream from the purged tube were lower by factors of 2 to 7 than the amounts from the control tube. A smaller difference was observed for the carbon steel samples. The relative affinity of the metal surfaces for the radioisotopes studied was found to vary by a factor of about 3×10^8 ; the most readily adsorbed radioisotopes being the lanthanides and the least readily adsorbed being Np²⁸⁹. Calculations based on the relative amounts of these sixteen radioisotopes indicate that, for a short exposure (35 days), La¹⁴⁰, As⁷⁶ and Cu⁶⁴ are the major short-lived radioisotopes adsorbed on original reactor piping after 15 years of exposure. From these measurements, it was shown that La¹⁴⁰, As⁷⁶ and Cu⁶⁴ were still the major short-lived contributors; however, Eu¹⁵², Co⁶⁰, Tb¹⁸⁰ and Fe⁵⁹, in addition to Zn⁶⁵ and Sc⁴⁶, were important long-lived contributors.

INTRODUCTION

In studying the origin of radioisotopes in effluent water from a reactor, it is of interest to know the effect of the film, which normally forms on fuel element jackets and process tubing, on the observed radioisotopic content of the effluent. It is also important to know the chemical and physical condition of the radioisotopes leaving the film in order to understand and predict their downstream behavior in the process tubing as well as their ultimate behavior in the Columbia River.

Production tests have recently been performed by the Irradiation Processing Department,¹ using the chemical cleaner Turco 4306B on process tubing, with the object of reducing rear face dose rates. These tests have provided an opportunity to study the build-up of radioisotopes in the effluent water after the process tubing and associated fuel element jackets have been cleaned chemically. They have also provided an opportunity to measure the uptake of radioisotopes in effluent water on aluminum, stainless steel and carbon steel surfaces downstream from the reactor.

^{*}Work performed under contract with the U.S. Atomic Energy Commission.

The relative radioisotope concentrations observed on these metal surfaces have been compared with those present on downstream surfaces after 15 years of operation to permit interpolations to other times of interest. This information is useful in understanding the chemical and physical form of the radioisotopes, for designing future studies of in-pile water treatment, and for calculating the contribution of the various radioisotopes to the γ -ray dose on the rear face of the reactor.

An outline of the analytical procedures which were developed or adapted for use in the measurement of the sixteen radioisotopes included in this study is presented.

EXPERIMENTAL

All of the reactor effluent water and metal samples which were analyzed in this study were obtained from reactors during their normal course of operation. The specifications of the reactor process water are given elsewhere.²

Radioisotope build-up in the effluent from a chemically cleaned process tube

This experiment followed the build-up of radioisotopes in the effluent water from a chemically cleaned tube. It was expected that the concentrations of radioisotopes in the effluent water observed would be similar to those from a new reactor. The tube was chemically cleaned while the reactor was down, and the radioisotope concentrations in the effluent water were measured periodically during the following $l\frac{1}{2}$ months. Four sets of

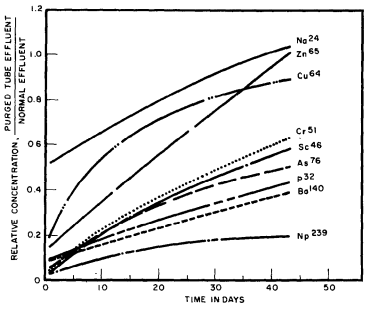


FIG. 1.-Build-up of radioisotope concentrations in effluent water from a Turco purged tube.

samples were collected during this time from the above tube, from a control tube of similar history (that was not purged with Turco), and from the whole reactor effluent. The exposure of metal in the purged and control tube was almost the same.

The first effluent water samples were collected after 26 hr of reactor operation. A continuously running sample line, which permitted sampling without turning on a valve, was used for all sampling. Because of sampling difficulties, only two samples

were taken from the control tube. However, it was found that the control tube had essentially the same concentrations of radioisotopes as the total effluent, so the total effluent was used as a control in following the concentrations of radioisotopes in the purged tube.

The ratios of the effluent radioisotope concentrations in the purged tube to the total effluent are given in Table I and are presented graphically in Fig. 1. From the

TABLE I.—RATIO OF THE RADIOISOTOPE CONCENTRATIONS IN THE CHEMICALLY CLEANED TUBE TO THAT OF THE TOTAL EFFLUENT

Date	Na ²⁴	P ³²	Sc46	Cr ⁵¹	Cu ⁶⁴	Zn ⁶⁵	As ⁷⁶	Ba ¹⁴⁰ *	Np ²³⁹
29 January 1959			0.0106	0.0346	0.222	0.133	0.0540	0.085	0.0130
30 January 1959 2 March 1959	0.716	0.123 0.229	0.0624 0.295	0.0872	0.134 0.673	0.199 0.463	0.0789 0.297	0.23 0.13	0.0295 0.145
12 March 1959	1.05	0.435	0.582	0.629	0.893	0.991	0.501	0.39	0.194

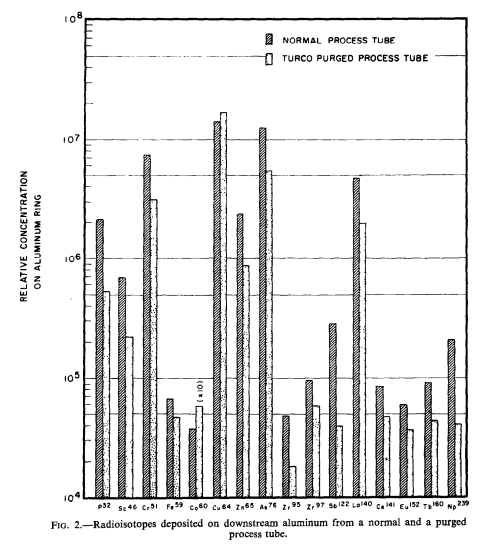
* The precisions of the Ba¹⁴⁰ measurements were relatively poor. Standard deviations were 25-50 per cent, compared with less than 5 per cent for the other radioisotopes.

curves in Fig. 1, it is evident that chemical cleaning reduces the concentrations of most of the radioisotopes in the effluent water by a considerable amount, and that their concentrations increase slowly toward normal operating values over a period of weeks to months. With the exception of Na²⁴, all of the isotope concentrations were lower by at least a factor of five than the total effluent during the first 2 days of operation. The effluent concentrations of P⁸², Sc⁴⁶, Cr⁵¹, As⁷⁶, Ba¹⁴⁰ and Np²³⁹ were reduced by a factor of 10 to 100 during this period. Also, the concentrations of these isotopes were less than 25 per cent of normal for 2 weeks after the purge and less than 65 per cent 6 weeks after the purge. These data support previous observations³ that the process tube film is an important reservoir for radioisotopes and parents of radioisotopes.

It is interesting to note that even the Na²⁴ in the effluent from the purged tube is lower by a factor of two. This is probably due to the lack of good "ion-exchange surface" on the freshly purged tube to hold the parent Na²³.

Radioisotope deposition on metal surfaces during a short exposure

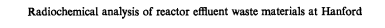
In this experiment, a description is given of the measurement of effluent water radioisotopes adsorbed on metal surfaces downstream from a normal and a chemically cleaned process tube. Dummy rings about 1.4 in. external diameter by 1 in. internal diameter by 0.5 in. long were located in the normal downstream dummy positions of two process tubes in a production reactor and allowed to remain for 35 days. One of these tubes was chemically cleaned by purging with Turco solution prior to positioning the rings. The second tube was used as a control. Dummy rings of aluminum, stainless-steel and carbon steel were studied. The relative amounts of the sixteen radioisotopes measured on these rings are listed in Table II. To better illustrate the relative uptake of these isotopes by the dummy rings in the chemically cleaned and normal tubes, the relative activities of these isotopes are shown on bar graphs as Figs. 2, 3 and 4 for aluminum, stainless steel and carbon steel, respectively. With two exceptions, namely Cu⁶⁴ and Co⁶⁰, the amounts of the radioisotopes found on the aluminum and stainless steel rings were much less for the chemically cleaned tube (a factor of 2 to 7 less) than for the normal tube (Fig. 2 and 3). These differences were smaller for the carbon steel rings (Fig. 4). In this case, the normal tube ring had about



the same or slightly less Co^{60} , Cu^{64} , Zn^{65} , Sb^{122} , La^{140} , Ce^{141} and Eu^{152} (13 years) than the purged tube.

Aluminum, stainless steel and perhaps some carbon steel are the major materials with which effluent water is in contact on the rear face of the reactor. It is, therefore, important to know the relative contribution of each radioisotope to total γ -ray dose from the radioisotope mixtures observed on these surfaces. The relative dose rates for the observed mixtures in Table II have been calculated for a distance of 1 ft from a flat surface. These values were calculated for the control tube only and are normalized to La¹⁴⁰ as 100 (Table III, Fig. 5). It is apparent that La¹⁴⁰, As⁷⁶ and Cu⁶⁴ are the major short-lived contributors to the γ -ray dose, while Zn⁶⁵ and Sc⁴⁶ are the major long-lived contributors.

To illustrate the relation between the radioisotopic content of the effluent water and that on the metal rings, the ratios of the radioisotopes on the rings to the radioisotopes



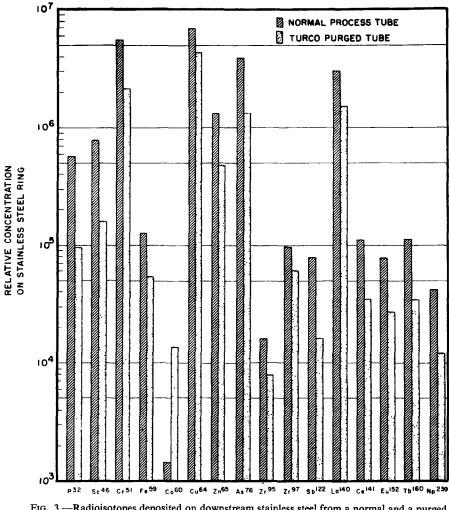
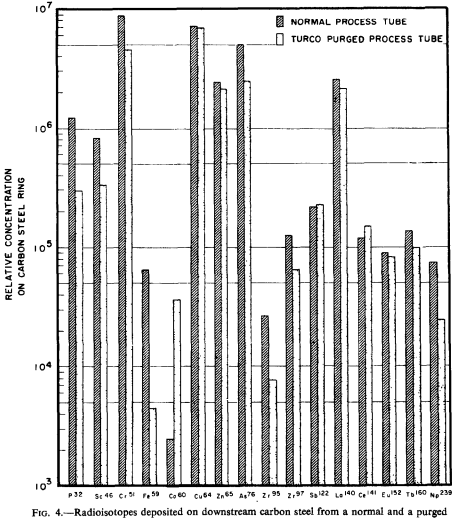


FIG. 3.—Radioisotopes deposited on downstream stainless steel from a normal and a purged process tube.

per ml of effluent are shown as a bar graph in Fig. 6. This graph covers the ring samples for the normal tube only. These values give an indication of the relative affinities of the metal ring surfaces for the various radioisotopes; however, these relationships do not give a complete picture since they are not corrected for radioactive decay or erosion. The relationships are very useful, however, in understanding and interpreting the behavior of these elements in film formation. It is apparent that Sc^{46} and the lanthanons (La¹⁴⁰, Eu¹⁵² and Tb¹⁶⁰) are most readily adsorbed, being present on the ring film at $2-3 \times 10^5$ times that of 1 ml of effluent. Figure 6 also illustrates the relative uptake of each radioisotope by aluminum, stainless steel and carbon steel surfaces. It is apparent that the uptake of most of the isotopes is much lower on stainless-steel surfaces than on aluminum or carbon steel. Sc^{46} , Fe^{59} and the lanthanons are exceptions, being about equally well adsorbed on the three materials.



process tube.

Radioisotope deposition on metal surfaces during long exposures

Since the radioisotope mixtures observed in the above work were the result of a rather short exposure (35 days), it was of interest to compare these relative concentrations with those present after prolonged exposures.

Sections of three stainless-steel pigtails on the rear face and three aluminum process tubes (from a location downstream from the neutron flux region), which had been in a reactor for 15 years, were obtained, the films removed, and their radioisotope compositions measured. Their observed relative concentrations and their corresponding relative dose rates, normalized to Eu^{152} , are presented in Tables IV and V and are shown graphically in Figs. 7 and 8. The difference in relative concentrations of isotopes from these two types of material may be partially due to their positions in the reactor. Unfortunately, the pigtails were from fringe tubes while the tubing samples were more

	Al	uminum	Stair	nless steel	G	arbon steel	
Radioisotope	Normal	Turco cleaned	Normal	Turco cleaned	Normal	Turco cleaned	
P ⁸²	212	· · · · · · · · · · · · · · · · · · ·	55.8	9.51	122	29.1	
Sc46	68.2	21.9	76.8	15.5	81.6	33.1	
Cr51	715	306	560	209	898	455	
Fe ⁵⁹	6.52	4.64	12.7	5.26	6.42	0.434	
Co ^ε	0.365	0.558	0.140	1.35	0.252	3.65	
Cu ⁶⁴	1380	1660	665	439	718	691	
Zn^{65}	237	86.0	128	47.9	244	211	
As ⁷⁶	1200	533	384	128	499	247	
Zr ⁹⁵	4.82	1.79	2.54	0.771	2.65	0.769	
Zr ⁹⁷	9.81	5.92	9.54	5.91	12.3	6.24	
Sb122	28.3	3.91	7.70	1.58	21.7	22.3	
La ¹⁴⁰	472	193	300	148	254	213	
Ce141	8.56	4.55	11.0	3.35	12.1	14.8	
Eu ¹⁵²	5.84	3.59	7.69	2.60	9.24	8.29	
Tb160	9.03	4.29	11.2	3.36	13.6	9.86	
Np ²⁸⁹	20.7	4.05	4.09	1.18	7.39	2.42	

TABLE II.—RADIOISOTOPES	DEPOSITED	ON METAI	. RINGS	DOWNSTREAM	FROM A	a Turco	PURGED A	ND
	A NORMAL	PROCESS	гиве (3:	5 DAY EXPOSUR	E)			
		Relative	D/M/I	Ring				

TABLE III.—FILM MATERIAL FROM METAL RINGS

Relative disintegration rates of the observed radioisotope mixtures and their corresponding γ -ray dose rates⁴ at 1 ft from a flat surface

Isotope	Aluminum		Stainless steel		Carbon steel		Average	
	D/M	Dose	D/M	Dose	D/M	Dose	D/M	Dose
La ¹⁴⁰	100	100	100	100	100	100	100	100
As ⁷⁶	254	89.2	128	44.9	197	69.1	193	67.7
Cu ⁶⁴	293	27.2	221	20.5	283	26.3	266	24.7
Sc46	14.5	14.0	25.6	24.7	32.2	31.0	24.1	23.2
Zn ⁶⁵	50.2	12.5	42.6	10.6	96.1	23.8	63.0	15.6
Cr ⁵¹	152	1.87	187	2.30	354	4.35	231	2.84
Sb122	6.00	1.48	2.56	0.540	8.55	1.80	5.70	1.27
Tb160	1.91	1.05	3.73	2.81	5.36	4.03	3.67	2.63
Fe ⁵⁹	1.38	0.780	4.23	2.39	2.53	1.43	2.71	1.53
Eu ¹⁵⁸	1.24	0.747	2.56	1.55	3.64	2.20	2.48	1.50
Zr ⁹⁷	2.08	0.745	3.18	1.14	4.85	1.74	3.37	1.21
Zr ⁹⁵	1.02	0.349	0.846	0.289	1.04	0.356	0.969	0.331
Np ²³⁹	4.39	~0.2	1.36	~0.1	2.91	~0.2	2.89	~0.17
C0 ⁶⁰	0.0774	0.0929	0.0466	0.0559	0.100	0.120	0.0750	0.0896
Ce141	1.82	0.0371	3.66	0.0747	4.77	0.0972	3.42	0.070

centrally located. Also, some information is lacking in comparing these tables since the tubing (Table V) was not received in time to measure the short-lived isotopes As^{76} , La^{140} and Cu^{64} .

If one compares the relative concentrations of radioisotopes in the film from the

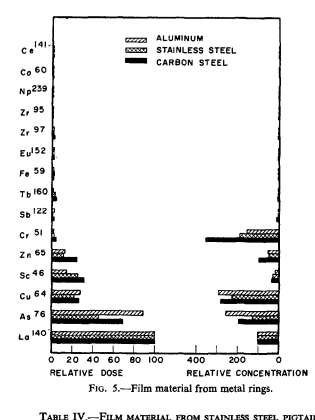


TABLE IV.—FILM MATERIA	. FROM STAINLES	S STEEL PIGTAILS
------------------------	-----------------	------------------

Relative disintegration rates of the observed radioisotope mixtures and their corresponding γ -ray dose rates⁴ at 1 ft from a flat surface

Isotope	Pigtail 1		Pigtail 2		Pigtail 3		Av erage	
	D/M	Dose	D/M	Dose	D/M	Dose	D/M	Dose
Eu ¹⁵²	100	100	100	100	100	100	100	100
As ⁷⁶	351	204	168	97.7	364	211	294	171
Zn ⁶⁵	294	121	162	66.8	216	88.9	224	92.2
La ¹⁴⁰	51.2	85.0	—		22.2	36.8	36.7	60.9
Sc46	35.8	57.2	18.4	29.3	32.1	51.4	28.8	46.0
Cu ⁶⁴	110	16.9	45.6	7.02	121	18.6	92.2	14.2
C0 ⁶⁰	8.08	16.0	2.44	4.84	2.56	5.08	4.36	8.64
Fe ⁵⁸	8.68	7.32	3.98	3.36	16.1	13.6	9.59	8.09
Zr ⁹⁵	6.73	3.82	5.18	2.94	18.8	10.6	10.2	5.79
Cr ⁵¹	345	7.04	51.3	1.04	185	3.74	194	3.94
Np ²³⁹	31.5	2.63	9.81	0.818	24.8	2.06	22.0	1.84
Ce143	5.37	1.56	3.82	1.11	7.19	2.10	5.46	1.59
Ce141	8.10	0.274	4.06	0.137	8.50	0.287	6.89	0.233

long-exposure pigtails (Table IV) with those found on the 35 day dummy ring (Table III) samples, it is obvious that the longer-lived isotopes are relatively much more abundant on the surfaces exposed for long periods of time. This is especially apparent

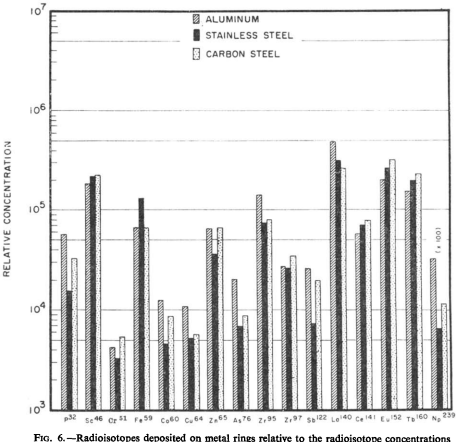


FIG. 6.—Radioisotopes deposited on metal rings relative to the radioisotope concentrations in effluent water.

TABLE V.-FILM MATERIAL FROM ALUMINUM PROCESS TUBING

Relative disintegration rates of the observed radioisotope mixtures and their corresponding γ -ray dose rates⁴ at 1 ft from a flat surface

Isotope	• Tube 1		Tube 2		Tube 3		Average	
	D/M	Dose	D/M	Dose	D/M	Dose	D/M	Dose
Eu ¹⁵²	100	100	100	100	100	100	100	100
Sc46	591	943	384	613	372	593	449	716
Zn ⁶⁵	2200	905	1780	720	1530	629	1840	751
Co ⁶⁰	101	200	126	250	84.8	168	104	206
Tb160	145	133	56.6	52.2	80.8	74.4	94.1	86.5
Fe ⁵⁹	46.0	43.4	11.9	11.3	23.5	22.2	27.1	25.6
Cr ⁵¹	1 770	36.0	897	18.3	1080	22.0	1250	25.4
Np289	378	31.5	167	13.9	190	15.9	245	20.4
Zr ⁹⁵	28.0	15.9	28.8	16.4	22.1	12.6	26.3	15.0
Ce141	52.4	1.77	38.2	1.29	38.1	1.29	42.9	1.45

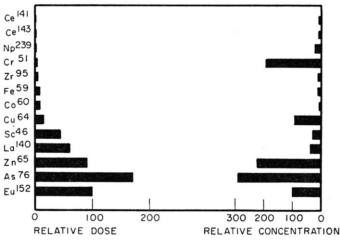


FIG. 7.-Film material from stainless-steel pigtails; average of three samples.

in the case of Co^{60} (5.2 years) and Eu^{152} (13 years). Relative to La^{140} (40 hr), these radioisotopes are about two orders of magnitude greater on the long exposure than on the short exposure surfaces.

As in the case of the short exposure study (Table III), the major short-lived radioisotopes on the stainless-steel surfaces exposed for long periods of time (Table IV) were As⁷⁶, La¹⁴⁰ and Cu⁶⁴. The major long-lived radioisotopes on the long-exposure stainless steel were Eu¹⁵², Zn⁶⁵, Sc⁴⁶, Co⁶⁰ and Fe⁵⁹, with a considerable change in relative importance from that observed on short exposure. These radioisotopes were also major contributors on aluminum exposed for a long period of time (Table V), but they were present in different relative amounts on this material. In addition, the longlived isotope Tb¹⁶⁰ (72 days) was a major contributor on the aluminum surfaces. This isotope could not be detected on the stainless-steel pigtails from their rare-earth γ -ray spectra, and its concentration was determined to be less than five per cent of that of the Eu¹⁵².

It is interesting to note that Eu¹⁵² (13 years), which, prior to this study, had not been measured in effluent water, is responsible for a large fraction of the rear-face

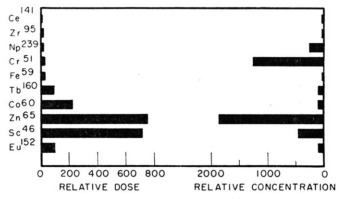


FIG. 8.-Film material from aluminum process tubing; average of three samples.

gamma-ray dose. On stainless steel exposed for a long period of time (Table IV), it is the major long-lived contributor.

DISCUSSION

These measurements have shown the marked difference that exists in the concentrations of radioisotopes in effluent water from a chemically cleaned (or new) process tube and a normal process tube. The slow build-up of the radioisotope concentrations in the effluent from a cleaned tube indicates that film formation in the tube is a slow process, and suggests that any water treatment which would prevent film formation would provide a considerable reduction in the radioisotope output in effluent water. Only nine radioisotopes were included in this phase of the work; therefore, it would be worthwhile in a future study to include several of the radiolanthanons, and also Fe^{59} and Co^{60} , since these isotopes were found to be important in accounting for the rear-face γ -ray dose.

The observed relative affinities of the radioisotopes in the effluent water for metal surfaces are of value in understanding and predicting their effect on rear-face materials of reactors and their fate in the Columbia River. The high relative adsorption for La¹⁴⁰ and the rare-earth elements (Fig. 6) was in agreement with data from a previous study of effluent water, which showed about 80 per cent of the La¹⁴⁰ to be associated with particulate material in the water.⁵ The low adsorption of Np²³⁹ also agreed with observations on effluent water which showed that less than one per cent of the Np²³⁹ was associated with the particulate material.

It is apparent that, since La¹⁴⁰, Eu¹⁵² (13 years) and Tb¹⁶⁰ are major contributors to the γ -ray dose on the rear-face of the reactor, other rare-earth elements will also contribute to this dose. Any subsequent study of this type should include Sm¹⁵³, Eu¹⁵² (9 hr), Ce¹⁴³ and, perhaps, other lanthanons which are known to be present⁶ in effluent water in relatively large concentrations.

Since the relative importance of most of the radioisotopes in effluent water to the γ -ray dose on the rear-face of the reactor has been determined (Tables III and IV), the problem of reducing this dose becomes a problem of removing these isotopes from the rear-face or preventing their accumulation.

ANALYTICAL PROCEDURES

Analytical procedures in which γ -ray spectrometric techniques were used to simplify or eliminate chemical separations were applied wherever possible. Also, the counting of all the γ -ray emitting radioisotopes was done on a multichannel gamma-ray spectrometer since it provided a check on the chemical separation procedures.

Phosphorus-32

The P^{32} is chemically separated by solvent extraction.⁷ The molybdophosphoric acid is extracted from dilute H_2SO_4 with a 10% solution of 1-butanol in diethyl ether.

Scandium-46

The Sc⁴⁶ is measured in the samples without chemical separation by coincidence counting of its cascade photons.⁸ The 0.885 and 1.12 MeV photopeaks are viewed with 90 keV windows by the two 3 in. \times 3 in. NaI(Tl) detectors of the coincidence spectrometer and the coincidence counting rate is measured. Interference from other radioisotopes is negligible. The spectrometer counting efficiency for Sc⁴⁶ is determined by

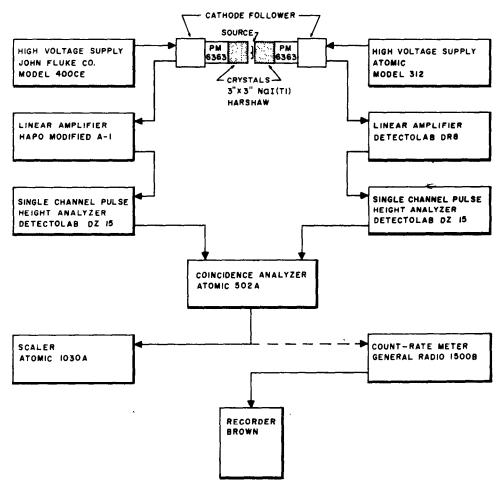


FIG. 9.-Block diagram of coincidence spectrometer system.

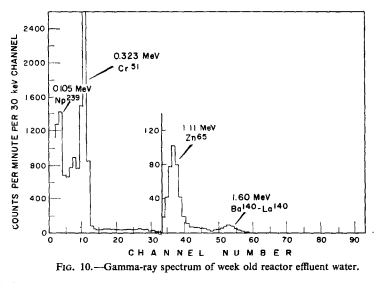
counting a Sc⁴⁶ standard with each group of samples. A block diagram of the coincidence spectrometer is shown in Fig. 9.

Chromium-51

The Cr⁵¹ is determined from the γ -ray spectrum of the sample by measuring the counting rate of its 0.323 MeV photopeak.⁸ Minor Compton corrections for interference from Sc⁴⁶, Zn⁶⁵ and Ba¹⁴⁰ are required. The gamma-ray spectrum of a typical effluent water sample, showing the Cr⁵¹ photopeak, is presented in Fig. 10.

Iron-59

The sample is boiled with concentrated HNO₃, after which the Fe^{59} is separated by a 15-min extraction with 0.45*M* TTA in benzene from a 12*N* HNO₃ solution, each washing being for a period of 5 min, after which it is evaporated on a 1 in. counting dish and plated. A 'spiked' Fe^{59} sample is carried through the procedure to determine the radiochemical yield which is about 90 per cent. The samples are counted on a multi-channel γ -ray spectrometer. If the sample contains HCl, 3 ml of HClO₄ are added, evaporated to 1 ml and the procedure outlined above is then followed.



Cobalt-60

The Co⁶⁰ is uniquely measured in the samples without chemical separation by coincidence counting of its cascade photons⁸ in a manner similar to that described for Sc⁴⁶. The 1.17 and 1.33 MeV photopeaks are viewed with 90 keV windows by the coincidence spectrometer's two 3 in. \times 3 in. NaI(Tl) detectors and the coincidence counting rate is measured. The spectrometer counting efficiency for Co⁶⁰ is determined by counting a Co⁶⁰ standard with each group of samples.

Copper-64

In reactor effluent water, where it is a major radioisotope, Cu^{54} is measured by coincidence counting of its positron annihilation radiation.^{8,9} In metal film samples, where the relative radioisotope concentrations differ considerably from that of effluent water, the Cu⁵⁴ is first removed from the mixture by carrier electrodeposition¹⁰, and then coincidence counted. The positron counting efficiency of the instrument is measured by using the long-lived positron emitter Na²² (2.6 years) as a comparator standard.

Arsenic-76

The As⁷⁶ in effluent water is measured by a β -absorption technique¹¹ in which advantage is taken of the fact that the 2.97 MeV As⁷⁶ β -particle can be counted through an absorber thick enough to shield out other β -radiations from the sample. The As⁷⁶ in the film samples is separated by carrier precipitation from 12N HCl as As₂S₃ after first removing Cu⁶⁴ by carrier electrodeposition. The As⁷⁶ is counted on a multichannel γ -ray spectrometer. The yield for the separation is above 95 per cent.

Zirconium-95-97

Twenty ml of concentrated HNO_3 are added to the sample and the solution is then evaporated to about 5 ml. The sample is diluted to 10 ml with water and extracted

with 10 ml of 0.45*M* TTA in benzene for 15 min. (A 60 ml separatory funnel and a mechanical shaker are used.) The organic phase is washed 5 min each with a 10 ml portion of 8 *N* HCl and three 10 ml portions of 6*N* HCl-2*M* HONH₂·HCl. The organic phase is then washed with 10 ml of 1*N* HNO₃ and evaporated on a 1-in. counting dish. The sample is counted on a multichannel analyzer after 24 hr, during which time the Nb⁹⁷ grows into radioequilibrium with Zr^{97} , and again after 2 weeks when the Zr^{97} -Nb⁹⁷ has decayed.

A Zr^{95} 'spike' is carried through the procedure to determine the yield which is about 95 per cent. Less than 1 per cent of the Nb radioisotopes are carried through the procedure.

Antimony-122

The samples are allowed to stand a week or more to permit the Cu⁶⁴ and most of the As⁷⁶ to decay. Twenty milligrams of As and Sb carrier are added, and also 20 ml of concentrated HCl, after which the solution is evaporated to 5–10 ml. Fifteen ml of concentrated HCl are then added and H₂S is passed through the solution to precipitate the As⁷⁶. The precipitate is discarded. The filtrate is diluted to 60 ml and H₂S is passed through the solution to precipitate the Sb¹²². A Sb¹²⁵ 'spike' is carried through the procedure to determine the yield which is about 95 per cent.

The Sb¹²² is measured from the counting rate of its 0.56 MeV photopeak and a correction is applied for the contribution of Sb¹²⁴. The magnitude of the correction is established from the counting rate of the 1.69 MeV photopeak of Sb¹²⁴.

Cerium-141

The cerium separation involves a TTA extraction similar to that described by Moore.¹²

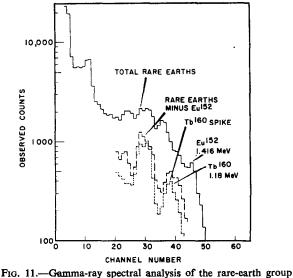
The sample is placed in 50 ml of 4N HNO₃, 40 mg of La carrier and 0.5 g of KBrO₃ are added, and the solution is boiled. It is then cooled, 5 ml of concentrated HF are added and, after standing for 5 minutes, the precipitate is removed by centrifugation and the supernate is discarded. The precipitate is dissolved in 50 ml of 4N HNO₃. A second fluoride precipitation is then made as outlined above, after which HClO₄ is added to the precipitate and the solution is evaporated to dryness.

The residue is dissolved in 10 ml of $1N H_2SO_4-0.1M K_2Cr_2O_7$ and extracted 15 min with 10 ml of 0.45*M* TTA in benzene. The organic phase is washed with two 10-ml portions of $0.1M H_2SO_4-0.1M K_2Cr_2O_7$, and then counted on a multichannel analyzer. The Ce¹⁴¹ is determined from the counting rate of its 0.142 MeV photopeak. The yield, which is about 80 per cent, is determined by carrying a 'spiked' sample through the procedure.

Lanthanum-140, Europium-152 (13 years) and Terbium-160

The initial portion of this separation is the same as the Ce¹⁴¹ procedure. The first aqueous phase from the TTA extraction is diluted to 25 ml, 5 ml of concentrated HF is added and the fluoride precipitate is then separated. The precipitate is evaporated to dryness with HClO₄, dissolved in 5 ml of 0.5N HNO₃, transferred to a separatory funnel with 2–3 ml of water and shaken 5 min with 20 ml of TTA in benzene. The aqueous phase is adjusted to a pH of about 4 by adding 10 ml of 0.5M NaAc and the

mixture is shaken for 15 min. The organic phase is then washed with two 10-ml portions of extremely dilute (about pH 4) HNO₃ and back-extracted by shaking 15 min with 10 ml of 0.1N HNO₈. The aqueous phase is evaporated on a counting dish and counted on a multichannel analyzer. The La¹⁴⁰ is determined from the counting rate of its 1.60 MeV photopeak. The γ -ray spectrum is measured again after a month or more, to permit La¹⁴⁰ and other short-lived rare-earth elements to decay, and the Eu¹⁵² and Tb¹⁶⁰ are determined from the counting rates of their respective 1.416 and 1.18 MeV photopeaks. Since there is overlap in the two spectra at these points, simultaneous equations derived from the spectra of the pure radioisotopes are used in calculating the net counting rates of the photopeaks. The γ -ray spectrum of the elements of the rare-earth group separated from effluent water 3 months old is shown in Fig. 11 with and without the Eu¹⁵² component subtracted. Since the rare-earth



from 3 month old reactor effluent water.

spectrum minus the Eu¹⁵² matches the Tb¹⁶⁰, it is apparent that these are the only rareearth elements which contribute significantly to the spectrum in this energy region.

The yield of rare-earth elements is about 85 per cent and is measured by carrying a 'spike' through the procedure.

Neptunium-239

The sample is placed in 50 ml of 1N HCl, 20 mg of La carrier are added, and the solution is boiled gently and then cooled, after which $\frac{1}{4}$ g of ferrous ammonium sulfate is added, the solution is stirred and let stand 5 min. Five ml of concentrated HF are then added and, after 5 min, the precipitate is removed by centrifugation, dissolved in 50–75 ml of N HCl and the fluoride is precipitate a second time, as outlined above. Three ml of HClO₄ are added to the precipitate and the solution is evaporated to dryness. The residue is dissolved in 8 ml of N HCl and $\frac{1}{4}$ g of ferrous ammonium sulfate is added, after which the solution is transferred to a separatory funnel with the aid of 2–3 ml of N HCl and shaken for 5 min. Ten ml of 0.45M TTA in benzene are

then added and the solution is extracted by shaking for 15 min. The organic phase is washed with two 10-ml portions of 1N HCl, each washing being for a period of 5 min, and then washed for 1 min with 10 ml of water. The Np²³⁹ is back-extracted into 10 ml of 8N HNO₃ and then counted.

The yield, as measured by β -counting, is 96.4 per cent with a standard deviation of 4 per cent.

Acknowledgements—The author wishes to express his appreciation to Mrs. B. L. Bauer for her help on the calculations, and to D. R. Edwards who performed much of the analytical work.

REFERENCES

- ¹ N. R. Miller, R. B. Hall and J. F. Jensen, *Production Test Authorization IP-203-AE 105C*, *Reactor Test Internal Chemical Decontamination*, U.S. Atomic Energy Commission, Hanford Atomic Products Operation, HW-57357, August 10, 1958.
- ² U.S. Atomic Energy Commission, Hanford Atomic Products Operation, HW-27155 Rev. 1., August 1957.
- ³ R. W. Perkins, U.S. Atomic Energy Commission, Hanford Atomic Products Operation, HW-56242, June 3, 1958.
- ⁴ Simon Kinsman *et al., Radiological Health Handbook*, Radiological Health Training Section, Sanitary Engineering Center, Cincinnati, Ohio, 1954, p. 49, Equation J(3).
- ⁵ R. W. Perkins, unpublished work.
- ⁶ W. B. Silker and L. J. Kirby, *Separation and Determination of Radioactive Rare Earths in Reactor Effluent Water*, U.S. Atomic Energy Commission, Hanford Atomic Products Operation, HW-48021, January 7, 1957.
- ⁷ W. B. Silker, Anal. Chem. 1956, 28, 1782.
- ⁸ R. W. Perkins, Gamma-Ray Spectrometric Systems of Analysis, *Proceedings of the Second Inter*national Conference on Peaceful Uses of Atomic Energy, Geneva 1958, P/2377, Vol. 28, p. 445, United Nations, Geneva, 1958.
- ⁹ R. W. Perkins and J. D. McCormack, The Determination of Cu⁸⁴ in Reactor Effluent Water by Coincidence Counting of the Positron Annihilation Radiation, U.S. Atomic Energy Commission, Hanford Atomic Products Operation, HW-45636, October 12, 1956.
- ¹⁰ I. M. Kolthoff and E. B. Sandell, *Textbook of Quantitative Inorganic Analysis*, The Macmillan Company, New York, pp. 420–425, 1947.
- ¹¹ R. W. Perkins, Determination of P³² and As⁷⁶ in Reactor Effluent Water by Beta-ray Absorption Techniques, U.S. Atomic Energy Commission, Hanford Atomic Products Operation, HW-50310, July 2, 1957.
- ¹⁹ Gilbert W. Smith and Fletcher L. Moore, Anal. Chem. 1957, 29, 448.

RADIOCHEMICAL ANALYSIS OF CORROSION PRODUCT MIXTURES FROM HIGH TEMPERATURE PRESSURIZED WATER LOOP SYSTEMS

L. C. BOGAR

Bettis Atomic Power Laboratory, Pittsburgh, Pa.

and

C. J. L. Lock

United Kingdom Atomic Energy Authority Group, Chalk River, Ontario

Summary—This paper describes the various types of samples of activated corrosion products obtained from the high-temperature pressurized water-loop facilities in the NRX reactor and the radiochemical procedures and radioactivity assay methods which are used in analysis of these samples.

Typically, in-pile loops are simulated reactor systems, i.e. they consist of a primary flowing coolant system and auxiliary make-up, purification, sampling and cooling systems. Two of the loops in NRX, namely, the carbon steel English Electric Corporation "Leo" loop and the Bettis Atomic Power Laboratory central thimble CR-V Loop, are described with particular attention to their associated sampling facilities: autoclaves and out-pile test sections for corrosion and corrosion product deposition studies, hot and cold corrosion product filters for filtration of insoluble materials, and ion exchange columns for purification of the loop water and the study of the ion-exchange behaviour of activated water-borne corrosion and fission products.

The types of samples obtained and the method of attack used to separate and isolate specific fractions of the mixed corrosion products are described. In particular, the separation scheme devised is one which permits the isolation of elements of the first transition series and their associated activated impurities from one another. The separation is basically an anion exchange technique involving the successive elution of carrier-equilibrated species.

Resin samples from the loop purification system are attacked in the presence of carriers by either digestion with H_2SO_4 -HNO₃ or by alkaline fusion.

In some cases, corrosion product deposits are assayed directly for activity distribution by means of gamma-scintillation spectrometry (differential pulse height analysis). Application of composite gamma-ray spectrum resolution to deposition coupons and pipe samples is described.

The radionuclides found in loop water and in unfilterable corrosion products are mainly radiative neutron capture products of the corroded in-pile and out-pile materials. The main nuclides analyzed for are $45.1 d \operatorname{Fe^{59}}$, $5.25 y \operatorname{Co^{60}}$, $27.8 d \operatorname{Cr^{51}}$, $12.8 h \operatorname{Cu^{64}}$, $2.58 h \operatorname{Mn^{56}}$, $2.56 h \operatorname{Ni^{65}}$, $245 d \operatorname{Zn^{65}}$ and $65 d \operatorname{Zr^{95}}$. Fast neutron reactions have been also observed through the appearance of $300 d \operatorname{Mn^{54}}$ and $72 d \operatorname{Co^{58}}$. Two radionuclides observed in the water and in solid corrosion products are $60 d \operatorname{Sb^{124}}$ and $2.8 d \operatorname{Sb^{122}}$. The source of these radionuclides has not been positively identified. The corrosion products from the CR-V loop also contain $253 d \operatorname{Ag^{110m}}$, $43 d \operatorname{Cd^{115m}}$ and $50 d \operatorname{In^{114m}}$ arising from the corrosion of an in-pile specimen of a Ag-In-Cd alloy.

Radioactivity assay of the purified fractions from separated samples is mainly by γ -ray scintillation spectrometry. Routine activity assay utilizing comparison with standardized differential pulse height distributions for pure radionuclides has been adopted because of the versatility of its use for both separated and gross samples. Conventional end-window beta-proportional counting techniques are used for those nuclides whose decay modes and energies preclude gamma-ray intensity assay techniques.

The complexity of corrosion product mixtures from in-pile loops requires that stringent control be maintained at all phases of the program from the time of sampling to the time radioactivity assay is completed. Several problem areas are discussed. For example, large amounts of Fe⁵⁹ contaminating Co⁵⁰ samples and Zr⁹⁵ contamination of Mn⁵⁴, are difficult to detect without considerable scrutiny of counting data. Carrier exchange is difficult to effect in some cases: e.g. Zr⁹⁵–Zr^N exchange. The gamma-counting data in particular receive careful scrutiny. The gravimetric assay of carrier yield is in some cases a doubtful and time-consuming operation; therefore, techniques eliminating gravimetric procedures should be explored.

INTRODUCTION

ONE OF the major problems confronting nuclear reactor designers is the selection of structural materials. Neutron economy considerations immediately limit the selections of the designer to those alloys having low macroscopic neutron absorption cross-sections if in-flux utilization is proposed. Metallurgical requirements must also be satisfied with respect to irradiation damage, corrosion behavior and mechanical strength. In addition to these reactor engineering requirements it is necessary that the various components of the reactor system be readily accessible in terms of personnel exposure for repair or refueling. The general requirements and techniques for irradiation testing of pressurized water reactor materials have been described by Wroughton and Cohen.¹

Studies involving reactor spectrum radiation activation of structural and fuel cladding materials, activation of corrosion products (crud) and system contamination with released fission products and with corrosion products, are being carried out in the high-temperature pressurized water-loop facilities in the NRX reactor at Chalk River. These experimental studies are being performed by a cooperative team of Canadian (Atomic Energy of Canada Ltd.), U.S. (Bettis Atomic Power Laboratory) and United Kingdom (U.K. Atomic Energy Research Authority) scientists and engineers. The radiochemical methods described in this paper are the result of evolution and development of techniques to study the corrosion and radioactivation of materials and their subsequent transport and deposition behavior in high-temperature pressurized water systems. Radiochemical analysis of fission products in these systems is described by Lock and Bogar in a complementary paper.²

LOOP FACILITIES IN THE NRX REACTOR

The in-pile NRX loops are simulated pressurized water reactor systems, i.e. they consist of a core (in-pile test section), a primary flowing coolant system and auxiliary make-up, purification, sampling and cooling systems. There are presently six hightemperature pressurized water loops in the NRX reactor which may be classified into three major types. The BAPL CR-IV-X rod loops (X-1, X-2, X-3) are stainless steel throughout and have a common make-up system. The BAPL CR-VI loop and English Electric Corporation "Leo" loop like the CR-IV systems have a water capacity of ~ 60 kg at temperature and pressure; however, they are of carbon steel out-pile with stainless steel in-pile test sections. The maximum empty-hole thermal neutron flux at the in-pile test sections of these loops is $\sim 3 \times 10^{13}$ n/cm²/sec. The BAPL CR-V central thimble (NRX-CT) loop test section occupies a lattice position of maximum thermal neutron flux (ca. 6×10^{13} n/cm²/sec). The CR-V loop is of larger water capacity than the other five loops (~ 100 kg) and has a reentrant in-pile test section; the other loops are of the straight-through type. The CR-V loop in-pile test section is zircaloy to provide maximum utilization of neutron flux. Out-pile sections are of stainless steel with the exception that out-pile runs of inconel and of monel have been used for radiochemical qualification testing of these alloys for use in primary system components.3

All of the loops have operating temperatures and pressures of the order of 600° F and 2000 p.s.i., respectively. Coolant pH $\simeq 10$ is maintained with LiOH formed mixed-bed resin columns (Rohm and Haas type XE-154) in the purification system or in the case of "Leo" and X-2 with KOH formed resin columns (Rohm and Haas

type XE-149). Total gas content is maintained at $30-50 \text{ cm}^3/\text{kg H}_2O$ (measured at STP) by addition of hydrogen to the systems to minimize radiation decomposition of water and to reduce O_2 concentration by radiation induced recombination.

SAMPLING SYSTEMS OF LOOP FACILITIES

Due to the structural material complexity of loop systems, samples with complicated distributions of radioactivities are obtained. Radionuclides which have been observed in NRX loops and their major target nuclide sources are shown in Table I. Close

Radionuclide	Target nuclide source	Probable production reaction
12.4h K-42	КОН	$K^{41}(n,\gamma)K^{42}$
12.4y H-3	LiOH	$Li^{6}(n,\alpha)H^{3}$
1.87 <i>h</i> F-18	LiOH, H₂O	Li ⁶ (n,α)H ³ ,O ¹⁶ (H ³ , n) F ¹⁸ ; O ¹⁸ (p,n)F ¹⁸
5.25y Co-60	CS ⁽¹⁾ , S.S. ⁽²⁾ , Inc ⁽³⁾ , M ⁽⁴⁾	$Co^{59}(n,\gamma)Co^{60+60m}$
72d Co-58	CS ⁽¹⁾ , S.S. ⁽²⁾ , Inc ³⁾ , M ⁽⁴⁾	Ni58(fast n,p)Co58+58m
45.1d Fe-59	CS ⁽¹⁾ , S.S. ⁽²⁾ , Inc ⁽³⁾ , M ⁽⁴⁾	$Fe^{58}(n,\gamma)Fe^{59}$
65d Zr-95	Zircaloy, Uranium Contamination	F.P., $Zr^{94}(n,\gamma)Zr^{95}$
35d Nb-95	Zircaloy, Uranium Contamination	Daughter Zr ⁹⁵
300d Mn-54	CS, SS, INC	Fe ⁵⁴ (fast n,p)Mn ⁵⁴
2.58h Mn-56	CS, SS, INC, M	Mn ⁵⁵ (<i>n</i> ,γ)Mn ⁵⁶
2.56h Ni-56	CS, SS, INC, M	$Ni^{64}(n,\gamma)Ni^{65}$
27.8d Cr-51	CS, SS, INC, M	$Cr^{50}(n,\gamma)Cr^{51}$
12.8h Cu-64	Μ	$Cu^{63}(n,\gamma)Cu^{64}$
245d Zn-65	CS	$Zn^{64}(n,\gamma)Zn^{65}$
60d Sb-124	?	Sb ¹²³ (<i>n</i> , <i>γ</i>)Sb ¹²⁴
2.8d Sb-122	?	$Sb^{121}(n,\gamma)Sb^{122}$
26.8h As-76	CS	$As^{75}(n,\gamma)As^{76}$
253d Ag-110m	CR-V Ag-In-Cd Specimen	Ag109(n, y)Ag110+110m
43d Cd-115m	CR-V Ag-In-Cd Specimen	$Cd^{114}(n,\gamma)Cd^{115+115m}$
50d In-114m	CR-V Ag-In-Cd Specimen	In ¹¹³ (n, y)In ^{114+114m}
Fission Products	U contamination, defected test specimen	U ²³⁵ (<i>n</i> , <i>f</i>)F.P.'s

TABLE I.--RADIONUCLIDES OBSERVED IN LOOP SYSTEMS

(1) Carbon Steel

(2) Stainless Steel

(3) Inconel(4) Monel

(4) Monei

inspection of this tabulation will reveal that few activation products have unique target nuclide sources. Although the fission products and the Ag, In, Cd isotopes have unique sources, Sb¹²² and Sb¹²⁴ are so ubiquitous that it appears conceivable that antimony impurities are in some way routinely introduced to the systems.

These activities may be sampled from many points and in a variety of physical forms. The types of samples currently obtained are:

- 1. Water samples from various points, i.e. before and after purification columns to determine resin bed efficiency.
- 2. Crud (insoluble corrosion products) from hot and cold crud filters. "Hot" and "cold" here imply relative ambient temperatures.

- 3. Deposition coupons in autoclaves or out-of-pile test-sections.
- 4. Pipe sections from various parts of the loops.
- 5. Ion-exchange resins.

Figures 1 and 2 illustrate schematically the Leo and CR-V loops.

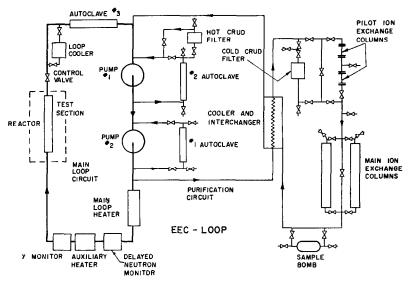


FIG. 1.-Schematic "Leo" loop

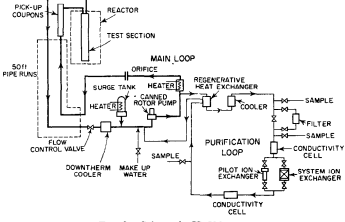


FIG. 2.-Schematic CR-V loop

The "Leo" loop has three out-of-pile autoclaves usable for corrosion product and fission product deposition studies. In both "Leo" and CR-V, the valving may be arranged so that the purification column flow passes through the crud filters. The out-of-pile test section immediately downstream of the in-pile test-section of CR-V is used for deposition and corrosion studies of various alloys suspended in the loop environment. Because several defected fuel tests have been run in "Leo", corrosion product studies are somewhat obscured due to high levels of fission products. The CR-V loop, however, is not intended for defected fuel tests; therefore, fission product background is very low and is accounted for by superficial uranium contamination. Although the initial attack in radiochemical analysis of the samples depends primarily on the physical form of the sample, separation for and decontamination of specific radionuclides is modified, depending on the fission product level in the loop.

GENERAL ASPECTS OF CORROSION PRODUCT SAMPLING AND ATTACK

The major effort in the corrosion product work at the NRX site is directed toward the analysis of samples from the crud filters. A typical cold crud filter is illustrated in Fig. 3. Particulate matter, $>0.5 \mu$ diameter, is filtered from the loop coolant onto 47-mm diameter type HA "Millipore" filter disks. Since this filter is accessible during reactor operation, daily crud levels may be determined. Crud is separable from the filter because of the solubility of "Millipore" filters in acetone. Refiltration of the crud through a weighed "Millipore" disk permits gravimetric assay of the collected crud. Total flow through the crud filter is recorded on a flow integrator; thus, crud concentrations in terms of p.p.m. may be determined. The hot crud filter is similar to the cold filter except that higher temperature operation precludes the use of "Millipore" filters; a sintered platinum disk is used.

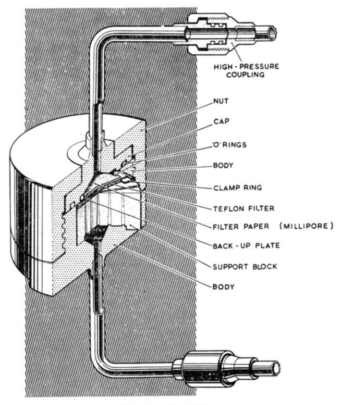


FIG. 3.-Crud filter

It has been found convenient to classify the corrosion product obtained on deposition coupons and pipe specimens into a "Brushed" fraction and an "Etched" fraction. The "Brushed" fraction is that crud which is removable by vigorous mechanical abrasion. Common nylon bristled tooth-brushes have been found to be very convenient for brushing crud. To remove the activities which have diffused into the tightly adherant oxide films of the various metallic samples, more drastic acid etchants such as $HNO_3 + HCl$ are used. This classification system has been developed to minimize dilution of the radionuclides with macro-quantities of etched metal.

Total loss of volatile chlorides such as $SbCl_5$ and loss of radionuclides by deposition on glassware surfaces is minimized by carrying out the dissolution procedures in the presence of known amounts of isotopic carriers.

Dissolution of filtered or "Brushed" crud from CR-V and "Leo" is fairly easy to accomplish by ppeated digestion with aqua regia mixtures. In some cases removal of the activity i.in he "Etched" fraction is more difficult and in extreme cases, long periods of refluxing with strong acids are necessary. Refluxing is avoided where possible because of the large amount of inactive metal also dissolved. After dissolution of the crud has been accomplished, the solutions are reduced to dryness and converted to metal chlorides by dissolving in conc. HCl. The solutions are volumetrically diluted (usually to 25 ml) to provide a stock solution for analysis.

Ion-exchange resins after removal from the loops are dried and stored in vacuo. Alkaline attack is accomplished in the following manner:

100–200 mg of dried resin, 10–20 mg of isotopic carrier for each element to be determined, and two pellets of KOH are reduced to dryness under a heat lamp in a 20-mm diameter, 30-ml Ni crucible. About 2 g of KOH and 1 g of KNO_3 are mixed and added to the crucible and gently heated until the reaction subsides. The temperature is raised until the mass is molten and glassy. Hearing is continued for 10 minutes. The crucible is cooled and the melt is extracted with water (alkaline solution of Zn). Insoluble material is centrifuged off and extracted with conc. HCl (acid solution Fe, Co, Cr, etc.). The acid-insoluble material is centrifuged off and discarded (insoluble sulfates from resin).

Acid attack is accomplished by destruction of the resin in the presence of carriers with red fuming HNO_3 -conc. H_2SO_4 . The carbonaceous organic material is then destroyed with a H_2SO_4 -HClO₄ mixture. This procedure is very time consuming (2-3 days) and its use is limited for determination of long-lived radionuclides.

Water samples (~ 1 L) may be obtained at various points in the purification circuit by sampling into a vessel containing a known volume (and weight) of isotopic carriers. The total volume (or weight) is then measured and the whole reduced to dryness. The residue is taken up in HCl and volumetrically diluted. Samples taken downstream from the crud filter are already filtered; therefore prior filtration is not necessary to yield a sample of the soluble species. Water samples upstream from the crud probe are filtered; the filtrate and precipitate may be combined or analyzed separately. Soluble short-lived nuclides such as Ni⁶⁵, Mn⁵⁶ and Cu⁶⁴ are analyzed from large samples of water as well as in crud samples.

THE SEPARATION SCHEMES FOR CORROSION PRODUCTS

The basic corrosion-product separation scheme is one based on anion-exchange separation of the carrier-equilibrated mixture. This scheme has been described by Lock and Corbett previously.⁴ The scheme is illustrated in Fig. 4. The Dowex-1 anion column is a bed of resin 5 mm in diameter by 15 cm long, 4–8% cross-linked, in the chloride form. Analyses are performed on duplicate aliquots of the sample stock

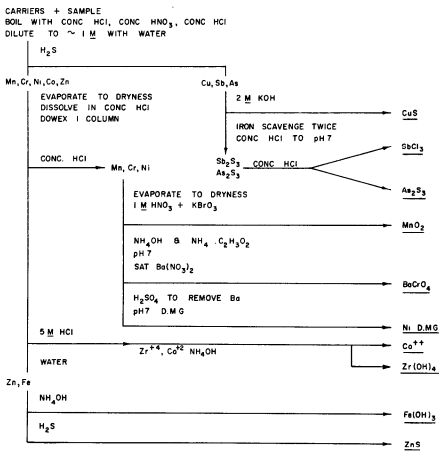


FIG. 4.—Separation scheme for Cu, Sb, As, Mn, Cr, Ni, Zn, Fe, Zr

solutions. The final precipitates, or solutions obtained from the separations, are further decontaminated using the procedures reported by Lock and Corbett⁴ or by the Los Alamos Group.⁵

In CR-V crud the Ag, In, Cd radionuclides are of low specific activity; therefore several crud samples are composited when analysis for these activities is desired. The separation scheme devised (Fig. 5) is a simple and rapid scheme based on basic chemical differences of Ag⁺, Im⁺³ and Cd⁺².

A resume of the separated forms and final precipitates obtained is given in Table II. The precipitates are filtered onto prewashed and weighed 2.4-cm Whatman #42 filter disks; washed, dried and weighed. The filtered precipitates are positioned concentrically with a 1.500 in. o.d. \times 1.020 in. i.d. \times 0.024 in. thick Al ring and are sandwiched between two layers of 1.5 in. wide "Scotch" tape (type 600 M M and M). These sources are then beta-proportional and/or gamma-scintillation counted.

Radionuclide	Separated as	Final precipitate
 Fe ⁵⁹	Fe(OH) ₃	$Fe(C_9H_6ON)_3^{(1)}$
Co ^{58,60}	Co+2	$K_{3}Co(NO_{2})_{6} \cdot H_{2}O$
Mn ^{54,56}	MnO ₂	MnO ₂
Ni ⁶⁵	$Ni(DMG)_{2}^{(1)}$	Ni(DMG)2 ⁽²⁾
Cu ⁶⁴	CuS	CuSCN, or Cu(C ₁₄ H ₁₁ O ₂ N) ⁽³⁾
Cr ⁵¹	BaCrO ₄	BaCrO ₄
Zn ⁶⁵	ZnS	ZnHg(SCN) ₄
Sb ^{122,124}	SbCl ₃	Sb ₂ S ₃
As ⁷⁶	As ₂ S ₃	As ₂ S ₃
Ag ^{110m}	AgCl	AgCl or AgIO ₃
Cd^{115m}	CdS	CdNH4PO4 · H2O
In114m	In(OH)3	$In(C_9H_6ON)_{3}^{(1)}$
Zr ⁹⁵	Zr(OH) ₄	Zr-mandelate

TABLE II.—CORROSION PRODUCT SEPARATED FORMS AND FINAL PRECIPITATES

(1) 8-hydroxyquinolinate or "oxine"

(2) DMG = Dimethylglyoximate

(3) α-benzoin oximate ("Cupron")

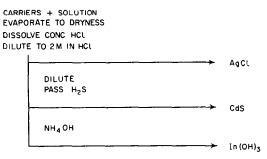


FIG. 5.--Separation scheme for Ag, In, Cd

RADIOACTIVITY ASSAY OF SEPARATED SAMPLES

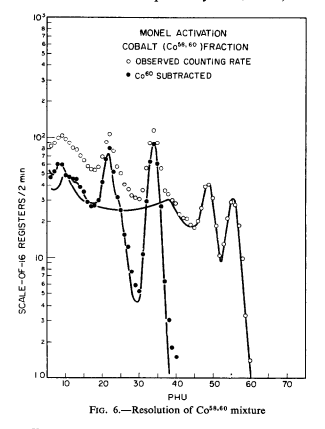
Separated samples are counted on end-window methane gas flow proportional betacounters to check chemical yields. The samples are then counted with a $1\frac{1}{2}$ in. $\times 1$ in. NaI(Tl) crystal multiplier-phototube assembly coupled to a 20-channel "Atomic Instrument Co." Model 520 differential pulse height analyzer. The gamma-scintillation counting apparatus has been described in detail by Hawkings and Edwards.⁶ The NaI(Tl) crystal has been calibrated for photopeak efficiency as a function of gamma-ray energy with samples of standardized nuclides. Total efficiencies, photopeak efficiencies, peak count to total count ratios and a compilation of standard spectra obtained from samples of standardized radionuclides have been reported.⁷

Radioactivity assay for Ni⁶⁵ and Mn⁵⁶ is normally by end-window beta-proportional counting because low activity levels and short half-lives preclude obtaining good gamma-spectra. The samples are beta-counted at frequent intervals to provide a decay curve which is extrapolated to sampling time. Count rates are converted to disintegration rates by means of experimentally determined ($4\pi \beta$ /end-window counter) efficiency factors. Counting data is also corrected for background, counter dead-time (~50 μ sec) and chemical yield. Cd^{115m} and In^{114m} are also beta-proportional counted because the gamma-ray abundances are difficult to establish from the decay scheme. Efficiency factors for beta-counting have been estimated from an empirical curve of beta-efficiency vs. effective beta-energy (+) (\overline{E}_{β}) for several $4\pi \beta$ standardized radio-nuclide samples.

Specific activities thus may be reported as disintegrations/minute-mg crud or dpm/ml of H_2O , dpm/cm² pipe etc.

RADIOACTIVITY ASSAY WITHOUT SEPARATION

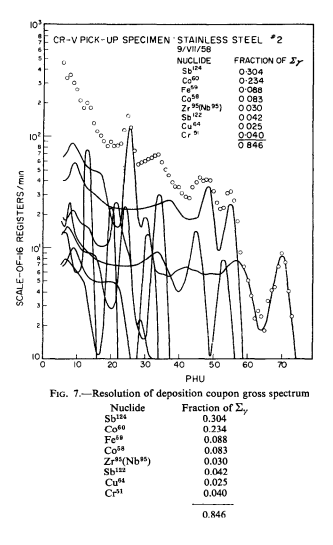
It is sometimes convenient to observe the distribution of radioactivities in certain samples without destroying the sample or expending an excessive amount of analyst time. Utilizing the standard spectra for samples of pure radionuclides, it is possible to resolve a gross or complex spectrum into its components by manual subtraction of spectra. The resolution of a two component system ($Co^{58,60}$) is illustrated in Fig. 6.



Note that the Co^{58} standard spectrum is "fitted" to the points (closed circles) resulting from the subtraction of the Co^{60} standard spectrum from the gross-spectrum (open circles).

This complex spectrum resolution technique has been extended to include the major components in CR-V crud, and is also applicable for examination of loop pipe sections. A typical case, where it was not desirable to destroy the deposition coupon for analysis but it was still possible to determine the activity distribution, is illustrated

 $(+)\overline{E}_{\beta} = \sum_{i} E_{\beta} i\alpha_{i}$ where E_{β} is the maximum beta-energy of the ith group of abundance α_{i}



in Fig. 7. In old CR-V samples (>30 days) $\sim 90\%$ of the total gamma activity may be accounted for by complex spectrum resolution. Absolute activities may also be determined because values for "photopeak count-to-total count ratios" and "total disintegrations per total count factors have been determined for the pure nuclide spectra. Accuracy of complex spectrum resolution compared to radiochemical analysis has proven to be 20–25%. Although this level of accuracy may suffice for some purposes, the resolution technique is tedious, requires absolute reproducility from the counting apparatus, and presumes that the character of the corrosion-product activity distribution is well established.

CONCLUSION

The methods described here have been developed to provide for easy and rapid analysis of activated corrosion product mixtures. These methods have been successfully used for the past two years in loop experiments at Chalk River. Some difficulties and pit-falls have been encountered, e.g. $Zr^{95}-Zr^N$ exchange in some CR-V water samples has been incomplete. In this case prolonged agitation in the presence of HF solved the problem. In other cases Zr^{95} contamination appeared in the Mn fractions in spite of repeated decontamination. Here again, prolonged equilibration of carrier with active material and specific decontamination steps for Zr were required

Scintillation spectrometry data is subjected to careful scrutiny. For example, it is difficult to see fairly large amounts of Fe⁵⁹ contamination in Co⁶⁰ samples. Careful comparison of observed photopeak resolution of a suspected sample with the Co-60 standard spectrum will reveal such contamination.

One of the problem areas requiring further development is the chemical yield determination of separated samples. Precipitates such as the Cu α -benzoin oxime are voluminous, filter with difficulty and are difficult to dry to constant weight. Spectrophotometric techniques for chemical yield assay coupled with well-type scintillation counting would materially decrease analysis time.

Acknowledgements—It is a pleasure to acknowledge the help and discussion of J. A. Corbett in developing many of the methods described here. The assistance of D. McLaughlin (AECL), J. Rees (AECL), D. L. Spate (BAPL) and C. A. Rein (BAPL) in carrying out the bulk of the analyses is also acknowledged. R. F. S. Robertson (AECL), G. M. Allison (AECL) and H. A. Clawson (BAPL) directed various phases of the corrosion product program. We are also indebted to R. C. Hawkings (AECL) and W. J. Edwards (AECL) for their continued assistance in the counter standardization program.

REFERENCES

- ¹ P. Cohen, and D. M. Wroughton, *Requirements and Techniques for Irradiation Testing of Reactor Materials for Pressurized Water Reactors*, Proc. 2nd Int. Conf. at Geneva, 1958, P/2508.
- ² C. J. L. Lock and L. C. Bogar, *The Separation and Radioactive Assay of Fission Products from Samples obtained from High Temperature Pressurized Water Loops*, WAPD-T-1052.
- ³ Y. Solomon and K. H. Vogel, Radiochemical Qualification of Inconel for use in Reactor Primary Coolant Systems, WAPD-T-900 (also issued as WAPD-M-(CDD)-1 Jan. 30, 1959).
- ⁴ C. J. L. Lock and J. A. Corbett, Procedures used in Radiochemical Studies of Activation and Fission Products found in the EEC Loop at Chalk River, UKE-CR-1003.
- ⁵ G.S.S. Radiochemistry Group, Collected Radiochemical Procedures, LA-1721 Sept. 1954.
- ⁶ R. C. Hawkings, and W. J. Edwards, Apparatus for Routine Quantitative Estimation of Radionuclides by Gamma-Scintillation Spectrometry, AECL-819, Nov. 1958.
- ⁷ L. C. Bogar, Routine Gamma-Scintillation Spectrometry: A Complication of Spectra and some Quantitative Analytical Aspects, WAPD report, to be published.

REACTOR FUEL ELEMENT TYPES*

J. W. Ullmann

Oak Ridge National Laboratory, Oak Ridge, Tennessee

Note—This paper is not included in the *Proceedings* because, in essence, the information contained therein can be found in a report by the Civilian Reactor Fuel Element Group, *Report on Civilian Reactor Fuel Elements*, TID-8505.

Summary—The function of a research reactor fuel element is to generate a flux of neutrons, beta rays, and gamma rays for absorption in experiments. The fuel element is usually cooled by water below the boiling point. In reactors of very low power, cooling by natural convection to the air is adequate.

The main function of a power reactor fuel element is to transfer heat to a coolant at a sufficiently high temperature for use in the production of electricity. Power reactor elements are, therefore, cooled by pressurized water, liquid metals, high-boiling organic compounds, and gases. The elevated temperatures introduce problems of corrosion and mechanical failure.

All fuel elements must contain fertile or fissile isotopes, and most elements contain both. In the interest of neutron economy, the quantities of absorbing isotopes are controlled. Analyses in the range of 0.02 to 100 p.p.m. are required for many elements, depending on their nuclear or chemical properties.

Because of the potential hazard and high cost of shutdown from fuel element failures, rigid analytical control is exercised to assure the quality of all materials and components. The concentration of impurities in each part of a fuel element must be controlled. The principal parts of a fuel element are; the fuel material, whose fissile content must fall within predetermined limits; possibly, for added stability, fuel diluent; the gap or bond between the fuel and its cladding, which may be a gas, plated metal, liquid metal, or an intermetallic compound formed by reaction of the fuel with the cladding material; the clad which separates the fuel from the coolant; possibly a moderator or a burnable poison to ease reactor control; miscellaneous hardware, such as a handle, springs, ferrules, insulators and, frequently, braze or weld material.

The two chief types of fuel elements for research reactors are cylindrical rods of natural U clad in Al, and plates of enriched U-Al alloys or tubes clad in Al and assembled into structures containing from 6 to 19 parallel plates or several concentric tubes. In both types the fuel is metallurgically bonded to the cladding.

The most common fuel element in United States power reactors consists of a bundle or cluster of rods containing ceramic fuel pellets in Zircaloy-2 or stainless steel tubes. Most reactors use pellets of slightly enriched UO₂. A few reactors use ThO₂–UO₂ pellets, and in a few reactors, the tube material is an Al-Ni alloy. Helium is used in the space between the ceramic fuel and the tube. The elements usually contain between 7 and 306 rods.

Under development are alternative ceramic fuels such as UC and dispersions of carbides of Th and U in graphite. Other cladding materials such as Be, Nb, graphite, and graphite coated with SiC, are being investigated.

Metallic fuel is also used in power reactors. The British use natural U rods clad in a finned tube composed of a Mg alloy. Rods of U-Mo alloy which are Na-bonded in stainless steel tubes and pins of the same alloy metallurgically bonded to Zr will be used.

Plate elements of highly enriched U-Zr alloy and slightly enriched U-Zr-Nb alloy, both metallurgically bonded to Zircaloy cladding, are in use. Plate elements of highly enriched UO_2 dispersed in stainless steel and clad in stainless steel are also being applied.

* Work performed under contract with the U.S. Atomic Energy Commission.

THE DETERMINATION OF U AND Pu IN U-Pu ALLOYS OF THE FISSION ELEMENTS*

R. P. LARSEN and C. A. SEILS, JR. Argonne National Laboratory, Lemont, Illinois

Note—This paper is not included in the *Proceedings* because it has been submitted for publication in *Analytical Chemistry*.

Summary—A melt-refining process has been developed for reprocessing the enriched U loading(s) of the EBR-II. After a number of cycles, an equilibrium alloy will be formed which contains small percentages of the fission elements Zr, Mo, Ru, Rh and Pd. The rare-earth elements, Sr. Ba, and several per cent of the U will react with the ZrO_2 crucible and be separated as an oxide dross.

Since, in the projected operation of the reactor, Pu bred in the reactor blanket is to replace U³⁸⁵ as the fissile material in the core, it was necessary to ascertain whether or not the melt-refining process was also applicable to alloys of U, Pu, and fission elements. Therefore Pu and U analyses on both the ingot and dross material were required.

Plutonium is determined spectrophotometrically as the nitrate after it is separated from interferences; U is determined by an X-ray spectrometric method after it is separated from the Pu alpha contamination. To remove the interference of the highly colored ions of group VIII metals, the sample solution is converted to a chloride medium and treated with excess Mg. The group VIII metals are precipitated; the Pu and U are reduced to the tri- and tetravalent states, respectively. Uranium(IV) which serves as a reductant to hold the Pu in the trivalent state, is extracted from strong HCl into a CCl_4 solution of tributylphosphate. The Pu is then extracted from the same medium after it is oxidized to the tetravalent state with nitrite. The U and Pu are easily extracted from the organic phase with water or dilute HCl. These solutions are then prepared for X-ray and spectrophotometric assay by converting them to HNO₃ solutions of the proper concentrations.

In an alternate procedure, Pu is separated by coprecipitation with LaF_3 . By dissolution of the precipitate in a $Zr(NO_3)_4$ -HNO₃ mixture, the interference of F⁻ is removed and the Pu is oxidized to the tetravalent state.

The coefficients of variation for the U and Pu analyses are 0.9 and 1.2 per cent, respectively.

* Work performed under contract with the U.S. Atomic Energy Commission.

RADIOCHEMICAL DETERMINATIONS OF Ce AND TOTAL RARE-EARTH ELEMENTS BY LIQUID-LIQUID EXTRACTION*

J. J. McCown and R. P. LARSEN

Argonne National Laboratory, Lemont, Illinois

Note—This paper is not included in the *Proceedings* because it has been submitted for publication in *Analytical Chemistry*.

Summary—A considerable amount of supporting analytical service will be required in the pyrometallurgical processing of EBR-II fuel. In the melt refining of the U 'fissium' alloys used as an equilibrium-type fuel at startup, ingots will be produced containing more than 0.5 per cent each of Mo, Ru, Zr, Pd, and Rh, together with dross samples containing as much as 10 per cent of rare-earth elements. Since the removal of rare-earth elements is of prime importance in the fuel cleanup, it will be necessary to determine Ce and/or total rare-earth elements at several stages in the processing cycle.

Standard radiochemical procedures for determining Ce or total rare-earth elements involve numerous precipitation steps and are somewhat tedious and time consuming. Since a solvent extraction separation would offer several advantages over precipitation methods, several solvent extraction systems for separating Ce and total rare-earth elements were investigated. Data published by Peppard and his co-workers on the use of dioctylphosphoric acid (HDEHP) for extracting carrierfree rare-earth isotopes indicated that such an extraction could be used as the basis for a quantitative radiochemical method.

Using di(2-ethylhexyl) orthophosphoric acid as extractant, rapid and quantitative radiochemical methods were developed for Ce and for total rare-earth elements. Cerium(IV) is extracted from a $10M \text{ HNO}_{s}-1M \text{ KBrO}_{s}$ solution with 0.75M (HDEHP) in n-heptane. After scrubbing the organic phase with a fresh HNO₃-KBrO₃ solution, the Ce is re-extracted into a $10M \text{ HNO}_{3}-3\% \text{ H}_2\text{O}_{s}$ solution. In the procedure for total rare-earth elements, these elements are extracted from a 0.01M HCl solution with 1.5M (HDEHP) in toluene. After the organic phase is scrubbed with fresh 0.01M HCl, the rare-earth elements are re-extracted into 8M HCl. Since each of the separation steps in both procedures is quantitative, it is unnecessary to determine a yield factor. The solutions derived by re-extracting the organic phase are either plated for β^{-} counting or are gamma-counted with a well-type scintillation counter.

The methods were tested on slices of U fuel element and irradiated U 'fissium' alloys. Several samples were also analyzed by standard precipitation procedures. The results compare favorably.

* Work performed under contract with the U.S. Atomic Energy Commission.

ANALYTICAL LABORATORIES FOR THE HANDLING OF PLUTONIUM*

CHARLES F. METZ and GLENN R. WATERBURY

University of California, Los Alamos Scientific Laboratory, Los Alamos, New Mexico

Summary—For the safe handling of Pu in analytical chemical laboratories, equipment is required capable of handling up to 1-g quantities of the radioactive material per analytical sample and the storage of 100-g quantities in the form of unused samples and residues. Special enclosures, known as dryboxes, or glove boxes, have been designed for this purpose. The analytical chemistry laboratories at the Los Alamos Scientific Laboratory that are so equipped are described in this paper. The philosophy of handling Pu is also discussed. These boxes are so designed that all required operations in chemical analyses may be performed safely without contamination of the room and with a minimum of risk to the analyst. Included are operations such as weighing, dissolving, precipitation, filtering, evaporating, centrifuging, and igniting of precipitates and also electrometric, polarographic, spectrophotometric, and spectrochemical techniques. The designs of individual boxes are varied to permit various operations, and at the same time they may be arranged interchangeably in circumstances requiring several boxes in a row. For routine and semiroutine analyses, these boxes are arranged for an assembly-line type of operation. The sample to be analyzed is introduced at one end of the assembly, successive steps in the analysis are performed as the sample is moved progressively through the assembly, and the residue is temporarily stored at the other end of the assembly in a suitable container.

These boxes are fabricated of stainless steel, with windows of Lucite or safety glass. All metal exposed on the inside is painted with a strippable plastic-base paint. Glove ports are located at a convenient height to permit easy use of rubber gloves. Experience has shown that Pu in solution can be safely handled in these enclosures without rubber gloves being on the ports but that for Pu in solid form, rubber gloves must be attached to the glove ports. These enclosures are equipped with all the usual laboratory services. Doors between boxes are of an unusual design, vertically operated by compressed air. Provisions are included for the operation of equipment such as centrifuges and pH meters within the boxes, yet permitting their removal in an uncontaminated condition.

Ventilation is provided through filters for all gloved boxes. It is so regulated by means of dampers in the exhaust ducts that it does not interfere with operations. In case of open front boxes, the flow of air is maintained at a minimum face velocity of 100 ft/min. Stainless steel as an exhaust duct material has been found to be unsatisfactory, especially for those boxes in which acids are fumed. Polyvinyl plastic has been used for the more recently installed exhaust ducts and has been found to be quite satisfactory.

The descriptive material is adequately illustrated by 20 photographs.

THE analytical chemistry laboratories at Los Alamos were designed for the routine and daily handling of plutonium analytical samples in solid or solution form, and of multigram size, if necessary. They were also designed so that a large variety of analyses could be done on samples of many different types, including metal, alloys, compounds and solutions. It was also understood that the plutonium would be free from fission product elements.

The radioactive nature of plutonium gives rise to alpha particles of approximately 6 meV energy, and also of very weak gamma rays which are not penetrating. Due to these facts, radiation shielding is not a factor in the design of laboratories for handling plutonium unless fission product elements are also present. Hence, under these conditions, the problems in designing laboratories for plutonium analysis arise principally

149

^{*} Work performed under contract with the U.S. Atomic Energy Commission.

from its extreme biological hazard. On the basis of a permissible total body tolerance of 0.3 μ g of the element, and in consideration of the ease with which plutoniumcontaining materials may find their way into the laboratory atmosphere, it is clear that special care must be taken if ingestion of plutonium is to be completely eliminated or, under unusual circumstances, kept to a safe level. Partly on the basis of the established permissible body tolerance, and partly on the basis of previous experience which indicated what could reasonably be accomplished, it appeared that a working philosophy in the design of analytical chemistry laboratories for handling plutonium should have the following aims: (1) to do alpha-hot operations in essentially uncontaminated rooms with a minimum of danger to personnel, (2) to keep contamination of exposed surfaces below 20 disintegrations/min per cm², and (3) to keep air counts below 9 disintegrations/min per m³. It was also clear that handling of macro-amounts of the element would be required in daily operations, and facilities must be designed for the safe handling and storage of up to 100-g quantities.

To accomplish these aims, special enclosures, called dryboxes, glove boxes, or enclosed hoods, were designed. These enclosures are essentially boxes with windows for viewing. Two general types are in use, one type being provided with ports to which rubber gloves may be fastened, the other, an "open front" design. They are equipped on the interior with all the usual and necessary laboratory services, such as gas, water, electricity, compressed air, vacuum, etc. They are designed to permit all the usual operations necessary in analytical procedures, such as weighing, filtering, centrifuging. electrolysis, drying, igniting, electrometric titration, etc. They are of various sizes, and may be 32, 36 or 58 in. long, 25 in. deep and 24 to 36 in. high.

The windows are of $\frac{1}{2}$ in. thick safety glass or Lucite. In the glove boxes, the windows are set in a metal frame which, in turn, is bolted to the box, with a gasket providing the seal. In the open front boxes, the windows are framed with an S-shaped neoprene gasket which, in turn, is cemented to the box. These gaskets do not provide a complete seal.

The boxes are made of No. 304 stainless steel, 16 gage, with a satin polish surface. This material, while providing a pleasing appearance, is not completely satisfactory, being easily corroded by halogen acids.

Partly to eliminate corrosion, and partly to provide better lighting, the interior sides, tops, and backs of these boxes are painted with an acid-proof, plastic-base paint, white in color. Adequate illumination is provided by fluorescent fixtures, exterior to the boxes, and resting on the top.

These boxes are supported on metal frames with adjustable legs. Once installed, however, the height of the boxes cannot be altered without corresponding changes in the duct work.

The exhaust ducts, originally of stainless steel, corroded rapidly and are being replaced with polyvinyl chloride. These ducts are equipped with water sprays which permit them to be washed continuously during heavy acid-fuming operations. The exhaust ducts pass downward through the floor into a larger duct which carries the exhaust air to a purification system consisting of a dry filtration system. From this point it is discharged into the atmosphere.

Experience has shown that, in all operations where it can be done, it is advantageous to arrange the dryboxes in such a sequence that assembly-line-type of operations may be carried out. In other words, the sample to be analyzed is introduced at one end of

the drybox series, the necessary sequential operations performed as the sample is passed along, and finally, at the other end, the residues from the analysis are stored in special containers, periodically removed and sent to recovery operations.

Figure 1 is an example of this type of operation. In this case, drybox space and equipment for the potentiometric titration of plutonium is shown. At the right, the plutonium sample is introduced into the weighing box; here it is weighed and then passed to the left where it is fumed with sulfuric acid; from there it is transferred to the titrating area where a potentiometric titration is made. The residue is temporarily stored in 2-liter bottles and later removed to the packaging area. You will note these boxes are either open front or used without gloves; obviously, they are suited for solution work only. As shown in Fig. 1, personnel wear shoe covers, a laboratory coat and rubber gloves. The pH meter seen at the floor of the box is actually under a sealed Lucite cover and supported in such a manner that it can be removed from below for servicing. Extended controls are used to operate the instrument (see Fig. 2). In these boxes, the opening at the front is 7 in. in height, which has been found adequate. It has been found necessary to provide ventilation to the extent of at least 100 linear ft/min across the face of this opening.

In this particular analysis, iron must be determined on a separate portion of the sample, and a correction made in the plutonium determination. To do this, the sample is introduced at the same place as in Fig. 1, but now it is passed to the right, into the dryboxes shown in Fig. 3, where the analysis is made, and the residue stored in the end boxes on the extreme right. In all, 40 linear feet of dryboxes are included in Figs. 1 and 2, and are required for the routine assay of plutonium. Another interesting fact comes to light. Because the equipment becomes contaminated, and also because it takes up considerable space in the dryboxes, once a series of boxes is set up for a particular analysis, it has been found from experience that it is next to impossible to use that particular drybox space for other analytical determinations. This fact simply means that a large number of dryboxes is required to do a variety of analyses.

For the handling of plutonium in solid form, as either metal or compounds, experience has indicated that the boxes should be completely enclosed, and rubber gloves placed on the glove ports. Adequate ventilation is a necessity, with the inlet air being filtered as it enters the box. In the Los Alamos laboratories the exhaust air is not filtered at the outlet of the box but it is realized there are advantages in so doing.

Figure 4 shows an arrangment wherein the boxes are equipped with gloves for work with plutonium-containing solids. Figure 5 shows a combination wherein only parts of the boxes contain gloves, and the remainder are of the open front type. This is permissible in those cases where the sample is a solid and is dissolved at some point in its analysis. Figure 6 shows an arrangement of open front boxes designed for wet chemistry with plutonium solutions. This arrangement can be used to carry out various analyses.

In Figs. 1, 2, 3, 4 and 5 there are boxes of several designs. In addition to the gloved types and the open front types, it is to be noted that some have sloping fronts and others have vertical fronts. In addition, but not noticeable in the photographs, some have reagent shelves, while others do not. Some have exhaust outlets near the bottom, others near the top. At one time it was thought the various designs were necessary for convenience of the workers in performing various kinds of analytical techniques. Experience has shown, however, this is not necessarily true. With two or three

exceptions, it is now thought all analytical techniques can be carried out in a box of one design.

Although boxes of several different designs are in use, these are designed with essentially complete flexibility. It is possible to put together an assembly of any possible combination of all the various types of boxes available; a situation allowed for in the initial flanged construction and endplate design.

It has been found advantageous to isolate regions in an assembly by placing partitions and doors at the required positions. Considerable saving of space was accomplished by using a specially designed, pneumatic-operated door which rises vertically, as indicated in Fig. 7. These doors are made from two $\frac{1}{4}$ -in. stainless steel plates separated by a toggle arrangement which, in turn is connected to the piston. At the end of the closing stroke of the piston, the two plates are expanded laterally against neoprene gaskets, thereby producing an effective seal.

In those cases where all the boxes in an assemblage are equipped with gloves, a problem arose as to how supplies and samples could be introduced without contaminating the room. This was solved by entrance through the bottom. For small items, including samples for analysis, a metal can, Fig. 8, equipped with a bayonetcatch lid and handle for carrying, was designed in size and shape to hold a pint ice cream container. This metal can, with gasketed lid removed, is attached to the underneath side of the box with a bayonet catch as shown in Fig. 9. A circular plate, 4 in. in diameter and gasketed flush with the floor of the box, can then be removed, and the contents of the metal can transferred to the box. The metal transfer can remains fastened to the bottom of the box until it is necessary to remove or transfer additional items. The pint-size ice cream container has proved to be a convenient, inexpensive container for packaging materials for entry into and removal from a box. It helps materially in keeping the inside of the metal can relatively free from contamination. However, these metal cans do become contaminated and, when they do, they are sent to the decontamination room. For larger items, a hole 12 in. in diameter was cut in the bottom of the box. To the bottom of the box on the underneath side a cylinder 6 in. long, designed with a flanged ring at the bottom, was bolted (see Fig. 10). A portable box mounted on a cart was designed with an opening to fit against the bottom of this cylinder, as shown in Fig. 11. This portable box can be raised into position, fitting snugly against the bottom of the cylinder and thereby affording a sealed unit for transferring equipment and supplies into and out of the gloved box. A gasketed plate, flush with the bottom of the glove box, a lid on the bottom of the cylinder, and a lid on the plastic portable box normally remain in place to confine contamination.

Because of the frequent use of centrifuges, and the advisability of keeping them uncontaminated from a servicing standpoint, they were incorporated into the boxes in the following manner. A metal box of suitable dimensions, containing a removable side as indicated in Fig. 12, was bolted and gasketed on to the bottom of the drybox. A hole with the same diameter as the opening in the centrifuge bowl was cut in the bottom of the drybox. By means of levelling screws the centrifuge could then be raised against a gasket on the underneath side of the drybox, thereby effecting a seal as shown in Fig. 13. During normal use, only the inside of the centrifuge bowl becomes contaminated. In case of repair, the instrument can easily be removed by removing the Lucite window.

The question whether sinks should be included in contaminated boxes was finally



FIG. 1.—Drybox assemblage for assembly line-type of operation.

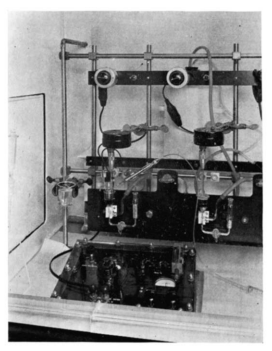


FIG. 2.-Built-in pH meter with extended controls.



FIG. 3.—Drybox assemblage for the determination of iron in plutonium.

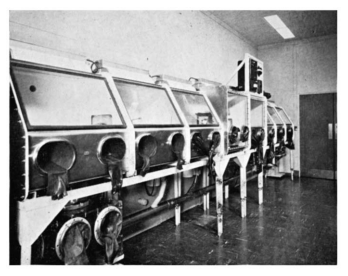


FIG. 4.—Glove box assemblage for work with plutonium-containing solids.

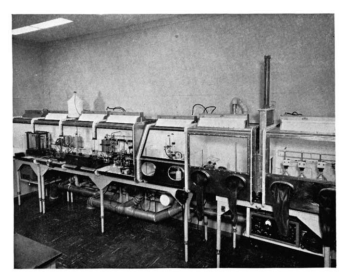


FIG. 5.—Drybox assemblage with open front and glove boxes.

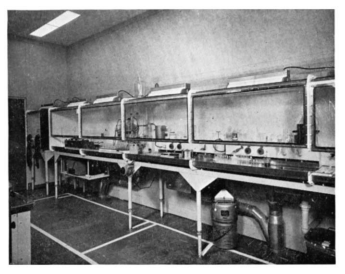


FIG. 6.—Open-front type boxes for general wet chemistry analysis of plutonium.

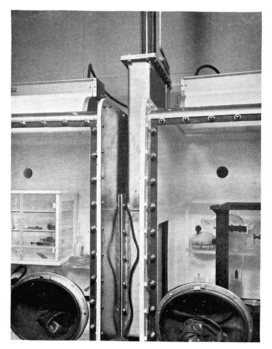


FIG. 7.-Pneumatically operated door between boxes.



FIG. 8.-Small size transfer can.



FIG. 9.--Small transfer can assembled to bottom of drybox.

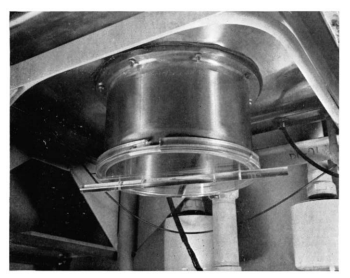


FIG. 10.-Large transfer unit connection.

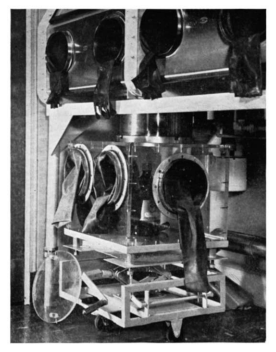


FIG. 11.-Portable transfer unit connected to drybox.

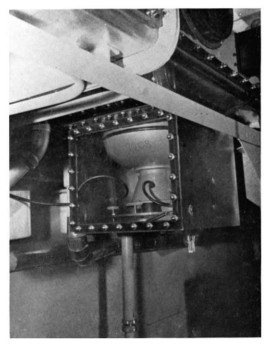


FIG. 12.—Centrifuge and housing on bottom of drybox.

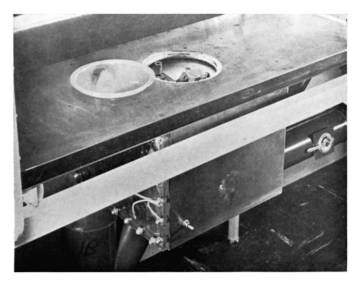


FIG. 13.—Opening in bottom of drybox for centrifuge.



FIG. 14.-Typical plutonium analytical laboratory.



FIG. 15.—Open benches for handling micro-quantities of plutonium.

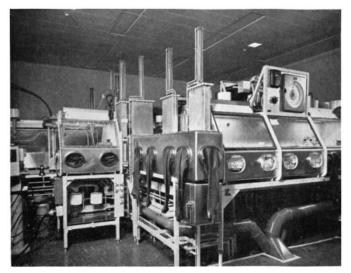


FIG. 16.—Assemblage of dryboxes for spectrochemical operations.

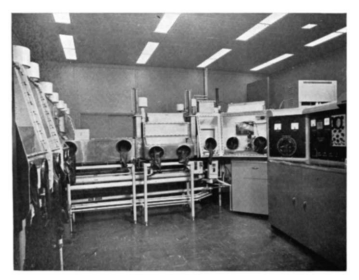
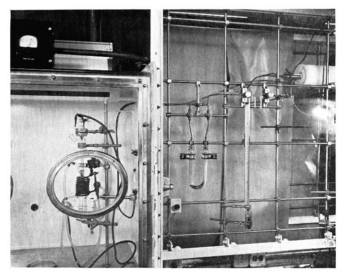


FIG. 17.-Glove box assemblage for spectrochemical operations.



FIG. 18.—Drybox unit for polarography.

ł,



FIG, 19.-Drybox equipment for the determination of oxygen in plutonium.

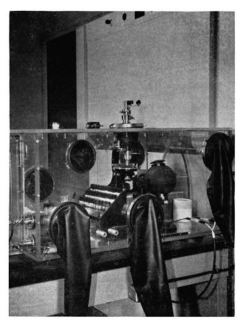


FIG. 20.—Drybox unit for chemical microscopy.

decided in their favor. However, all drains from these sinks are connected to retention tanks. The liquid in these tanks is checked for plutonium content before being released to the industrial treatment plant.

In equipping analytical laboratories with these boxes for handling plutonium, consideration must be given to the fact that a certain amount of "cold" laboratory work is necessary in order to be able to carry on the work in the boxes. For this reason, most of the laboratories for the analysis of plutonium at Los Alamos have been designed with about half the wall space equipped with dry boxes and the remainder occupied with the normal type of laboratory bench as shown in Fig. 14. This arrangement has proven very satisfactory. Experience has shown that drybox assemblages suitable for routine and semi-routine analytical chemistry work are also satisfactory for research on the analytical chemistry of plutonium, and the development of procedures for the analysis of plutonium.

It has been found possible to handle small samples of dilute solutions for radiochemical analysis in specially designed open-front hoods, as shown in Fig. 15. Ventilation is provided through a narrow opening at the bottom and also at the top of the back of the hood. This arrangement has been found to be satisfactory for such microoperations as pipetting a few microliters of plutonium solution into volumetric flasks, diluting, pipetting a few tenths of an ml on to a glass or metal plate, evaporating the solution to dryness and flaming. These working spaces are equipped with centrifuges for separating, by precipitation, a fraction of a microgram of plutonium from other constituents of the solution. In any case, the amount of plutonium removed from the original sample bottle does not exceed 1 or 2 μg . In this work the operator always wears rubber surgical gloves.

Various combinations or arrangements of boxes suitable for special uses, are easily assembled. Figures 16 and 17 show arrangements for assembly-line type of operations in spectroscopy. Figure 18 shows a box specially designed for polarographic work wherein the constant-temperature bath and cells are placed inside the box. The mercury reservoir is also located inside the box with the adjustment extended outside the box. The polarograph itself is located outside the box and is connected to the cells with extended leads. Not shown on the left is an assemblage of boxes for sample preparation. The hinged door which opens into this assemblage may be seen on the left side of the polarographic-cell box. In Fig. 19, a set up is shown for determining oxygen in plutonium. The induction heating coil and enclosed graphite crucible are shown inside the drybox. A glove has been removed for better visibility. The rest of the apparatus is outside the drybox. This is a typical example of adapting apparatus for use inside a drybox. Figure 20 shows a specially designed box for chemical microscopy work. The eye piece of the microscope extends through the top of the box. The actual manipulation of the microscope is done inside the box. An inert, dry atmosphere may be maintained inside the box when necessary.

Nearly 5 years of operations in the boxes have proven the general philosphy to be be sound and the boxes adequate for the work done in them. In retrospect, it now seems that a single design could be made that would be satisfactory for most of the analytical operations involving plutonium that are carried out at Los Alamos.

THE HIGH-ALPHA-RADIATION ANALYTICAL FACILITY OF THE OAK RIDGE NATIONAL LABORATORY*

J. H. COOPER

Oak Ridge National Laboratory Oak Ridge, Tennessee

Summary—The High-Alpha-Radiation Analytical Laboratory has been in operation since 1952. Originally concentrated Pu and U^{233} solutions were assayed in this laboratory and the control analyses associated with the ion-exchange isolation of these solutions were also carried out in this facility. However, during the past few years, the high-alpha-laboratory has been used to provide analytical services for all development programs concerned with the processing of alpha-emitting materials, such as Np²³⁷ and Am²⁴¹.

A brief description of the methods of analysis and techniques is presented. Mention is made of some of the special problems that have arisen concerning the handling of alpha-emitting materials. Future plans for a transplutonic analytical facility are also discussed.

EARLY in 1952 at the Oak Ridge National Laboratory, a special operation was initiated for the purpose of isolating the pure plutonium and uranium-233 product solutions from the metal recovery processes. The operation of collecting product solutions on ion-exchange resins had just been adopted at the recovery process pilot plant. It was the duty of the newly formed Isolation Group to elute these products from the resin and further purify, decontaminate, and concentrate them. Figure 1 shows a diagram

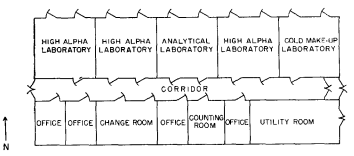


FIG. 1.--Plan of isolation building.

of the isolation laboratory. Briefly, the process involved transporting the loaded ionexchange columns to the isolation laboratory, eluting the plutonium or uranium, and purifying and concentrating the material by precipitation, filtration, and redissolution.

As improvements were made in the pilot plant processes, elution of the essentially pure product material was carried out in the recovery plant itself and the isolation laboratory was no longer needed for its original purpose. Since the initial operation of this laboratory involved the handling of high-alpha radioactive material, all research, development, and small-scale production operations involving this type of material were transferred to this laboratory.

It is the purpose of this paper to present some of the operational techniques of the high-alpha analytical laboratory. The primary duty of this laboratory is to assay the

^{*} Work performed under contract with the U.S. Atomic Energy Commission.

final solutions of the power reactor fuel reprocessing pilot plant. The sampling of these solutions initially was the responsibility of the Analytical Chemistry Group. The original storage and shipping containers were polyethylene bottles of 4-liter capacity from which a sample was withdrawn with a pipet. A later type of container was a stainless steel can, 6 in. in diameter and 6 ft in height. Samples were removed from these containers through Tygon tubing with a siphoning action initiated by suction

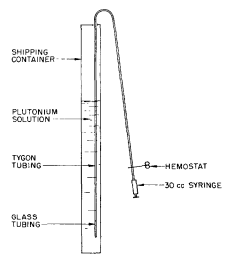


FIG. 2.—Plutonium sampling apparatus.

applied with a 30 cm³ hypodermic syringe. Figure 2 shows a schematic diagram of the apparatus used in sampling. Following is a stepwise description of the sampling procedure:

(1) A glass tube, 8 to 10 in. in length, is fitted into a 3/16-in. i. d. Tygon tube, 10 ft. in length.

(2) The tube, with the glass tube attached, is lowered into the solution container through a 1-in. hole in the top.

(3) The tip of a 30-cm³ syringe is inserted into the end of the Tygon tube.

(4) Gentle suction is applied, allowing the solution to rise in the tube.

(5) When the solution approaches the end of the tube, a hemostat is clamped on the tube and the syringe is removed.

(6) The open end of the tube is placed in a sample bottle and the hemostat is slowly released, allowing the solution to flow into the bottle.

(7) The flow is again stopped with the hemostat and the bottle capped.

(8) The tube, filled with solution, is raised and the hemostat released, allowing the solution to drain back into the product container.

(9) The tube is slowly withdrawn from the container through a compress of absorbent tissue and discarded.

At the present time, samples are taken in the processing plant by recirculating type samplers in glove boxes and sent to the laboratory for assay. Table I lists the determinations made on the plutonium solutions.

Aliquots for plutonium determinations are fumed to near dryness in mineral acids

Determination	Method ¹	
Plutonium	Potentiometric ceric sulfate titration (tentatively, controlled potential coulometric titration)	
Density	Direct weighing	
Iron	Spectrophotometric o-phenanthroline	
Uranium	Fluorophotometric	
Acid	NaOH titration, phenolphthalein	
Gross alpha	Methane proportional counter	
Gross gamma	Scintillation counter	
Gamma spectrum	Single channel recording gamma spectrometer	
Plutonium isotopes	Mass spectrometer	
Trace impurities	Emission spectrograph	

TABLE I.- ASSAY OF PLUTONIUM PRODUCT SOLUTIONS

in the high velocity hoods and diluted to volume from which aliquots are taken for analysis. Plutonium interference in the fluorophotometric determination of uranium is eliminated by precipitating the plutonium as the fluoride and then analyzing the supernatant solution. Aliquots are prepared in this laboratory for mass analysis and emission spectrographic analysis. Samples for the mass spectrometer are evaporated directly on the spectrometer filament and sealed in a polyethylene bag for shipment to the mass spectrograph. This is accomplished by adsorbing plutonium(IV) on an anion resin from a 8M nitric acid solution. The effluent solution from the resin is submitted to the spectrographic laboratory for determination of trace impurities.

A list of the analyses performed on the uranium-233 product solutions is shown in Table II.

Determination	Method ¹	
Uranium	Potentiometric ferric sulfate titration	
Density	Direct weighing	
Acid	NaOH titration—Beckman Model K titrator with ferrocyanide complex of U	
Thorium	Spectrophotometricthoron	
Gross alpha	Methane proportional counter	
Gross gamma	Scintillation counter	
Gamma spectrum	Single channel recording gamma spectrometer	
% U ²³²	Alpha energy analysis of gross alpha mount	
Uranium isotopes	Mass spectrometer	
Trace impurities	Emission spectrograph	

TABLE II.—ASSAY OF URANIUM-233 PRODUCT SOLUTIONS

Interfering uranium is separated from thorium by adsorbing the uranium from 6.5M hydrochloric acid on an anion resin column. The thorium in the effluent is determined spectrophotometrically with thoron. Uranium is removed from sample



FIG. 3.—Solvent extraction apparatus.



FIG. 4.—Plate drying apparatus.

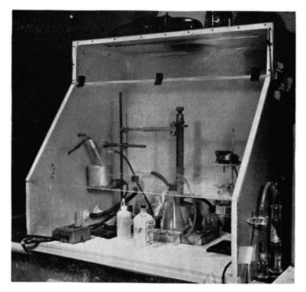


FIG. 6.—Plywood hood used for plutonium titrations.

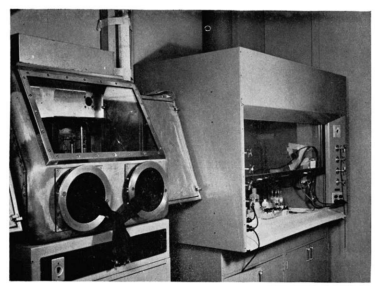


FIG. 7.-Glove box and hood in high-alpha analytical laboratory.

PREFACE

PROGRESS in nuclear reactor technology is intimately related to and contingent on advancements in other disciplines, of which analytical chemistry is by no means the least in importance. In many widely scattered centers of nuclear energy development, new methods, instruments and facilities are continually being devised while established methods and facilities are likewise being improved and adapted to a wide variety of intriguing problems in the analysis of nuclear reactor components, fuels and fission products. As a contribution toward the coordination of these efforts and the dissemination of this information, a series of conferences on analytical chemistry in nuclear reactor technology was initiated in 1957 by the Oak Ridge National Laboratory.

The Third Conference of this series was held at Gatlinburg, Tennessee, on 26–29 October 1959. At this conference forty-one papers were presented on methods, equipment and facilities for carrying out the analytical work which is, or may be, required following the operation of nuclear reactors. Some indication of the scope of the conference is reflected by the subject-titles of the different sessions: *Analytical Chemistry of Fuels; Analytical Chemistry of Plutonium and the Transplutonic Elements; The Analysis of Fission-product Mixtures*, and also one *General* session on related subjects.

In these Proceedings, a compilation is given of the papers which were presented at this Conference, excepting, in a few cases, those papers which are readily available in publications of the U.S. Atomic Energy Commission or have been submitted for publication elsewhere in the open literature. In lieu of reprinting this latter class of papers, a reference to and abstract of each paper is given at the end of each section.

Drafts of the papers in the Proceedings were revised, when required, in the interest of uniformity of presentation or to convert a paper which had been prepared for oral presentation to a form more suitable for publication; consequently, the Editors must assume all responsibility for errors or omissions which have been occasioned by such review.

The success of an effort of the magnitude of this Conference is, most certainly, dependent on the contributions of many persons. First among those who merit our sincere thanks and appreciation are the writers and speakers, and those who served as presiding chairmen over the different sessions. The Editor is also deeply indebted to the members of the program committee: J. H. Cooper, G. W. Leddicotte, J. A. Norris, R. W. Stelzner and J. P. Young, who laid the groundwork for and directed the conference, and also to Dr. M. G. Mellon of Purdue University for his stimulating and factual after-dinner talk, "Analyses and Analysts."

We are also most grateful to D. D. Cowen and his staff for their efficient services in handling the many details of the conference, and to many others, especially H. P. House and Miss M. A. Marler, for their competent assistance in the preparation of the papers for publication in the Proceedings and for numerous other contributions in the conduct of the Conference.

Whereas the proceedings of the two prior conferences of this series were published by the U.S. Atomic Energy Commission, the Proceedings of this Conference is

1

Preface

presented in a special collected issue of *Talanta* through the helpful and understanding cooperation of the staff of Pergamon Press. This departure from prior practice was motivated largely in the knowledge that, because of the extensive experience and contacts of the Pergamon Press in the field of technical and scientific literature, wider dissemination of this useful information could be achieved through the utilization of this service. We are most sincerely grateful to the publishers and staff of *Talanta* for this generous contribution and for their many courtesies in making this compilation possible.

C. D. SUSANO

ü

TALANTA

An International Journal of Analytical Chemistry

VOLUME 6 1960

Editor-in-chief

CECIL L. WILSON

Belfast, N. Ireland

Regional Editors

L. GORDON Cleveland R. PRIBIL Prague T. TAKAHASHI Tokvo

Assistant Editor

M. WILLIAMS

Birmingham

Editorial Advisory Board

Professor F. E. BEAMISH-Toronto Professor W. WAYNE MEINKE-Ann Arbor Professor R. BELCHER-Birmingham (Chairman Michigan of the Board) Professor J. MINCZEWSKI-Warsaw Professor H. BODE-Hanover Dr. A. D. MITCHELL-London Professor G. CHARLOT-Paris Dr. G. H. MORRISON-Bayside, New York Professor C. CIMERMAN-Haifa Professor F. NYDAHL-Uppsala Dr. C. E. CROUTHAMEL-Argonne, Illinois Dr. M. PESEZ-Paris Professor P. DELAHAY-Baton Rouge, Dr. E. RANCKE-MADSEN-Copenhagen Louisiana Professor G. GOPALA RAO-Waltair, S. India Professor H. DIEHL-Ames, Iowa Professor C. N. REILLEY-Chapel Hill, N. Dr. C. Dragulescu-Timisoara Carolina Professor L. ERDEY-Budapest Dr. M. L. SALUTSKY-Clarksville, Maryland Professor E. SCHULEK-Budapest Professor F. FEIGL-Rio de Janeiro Professor G. SEMERANO-Padova Professor H. FLASCHKA-Atlanta, Georgia Professor W. GEILMANN-Mainz Professor G. FREDERICK SMITH-Urbana, Illinois Dr. J. HOSTE-Ghent Professor H. SPECKER-Dortmund-Aplerbeck Mr. H. W. KIRBY-Miamisburg, Ohio Professor E. H. SwIFT-Pasadena, California Professor J. J. LINGANE-Cambridge, Professor P. S. TUTUNDŽIĆ-Belgrade Massachusetts Professor F. L. WARREN-Pietermaritzburg Professor F. LUCENA-CONDE-Salamanca Dr. H. WEISZ-Vienna Dr. S. J. LYLE-Durham Dr. T. S. WEST-Birmingham Dr. R. J. MAGEE-Belfast Dr. JAMES C. WHITE-Oak Ridge, Tennessee Professor H. MALISSA-Vienna Professor H. H. WILLARD-Ann Arbor, Michigan Mr. F. J. WOODMAN-Sellafield



PERGAMON PRESS

LONDON · NEW YORK · PARIS · LOS ANGELES

驇

LIST OF CONTENTS

Volume 6

Preface	•	•		•		•		•	•	•	•		•	•	i
C. D. Sus	ANO:	The r	ole of	the a	nałyte	cal ch	emist	in nu	elear	reacto	r tech	nolog	У	•	iii

SECTION I. GENERAL

C. J. RODDIN: Standards in the nuclear energy program	3
CALVIN E. LAMB: The high-radiation-level analytical facility at the Oak Ridge National Laboratory	20
J. O. HIBBITS, W. F. DAVIS and M. R. MENKE: The determination of iron in yttrium	28
D. L. MANNING and OSCAR MENIS: Some applications of high-frequency fitrimetry .	30
H. W. DUNN: X-ray absorption edge analysis	42
E. A. HAKKILA and G. R. WATERBURY: X-ray fluorescence spectrographic determina- otion of impurities and alloying elements in tantalum container materials	46
R. H. GADDY: The determination of moderator turbidity particle size by an angular- dependent light-scattering technique.	52

SECTION II. THE ANALYTICAL CHEMISTRY OF FUELS

L. NEWMAN, T. PRACH and H. L. FINSTON: The analysis of a liquid metal reactor fuel	55
JASON L. SAUNDERSON: A method of uranium isotope analysis by direct reading emission spectroscopy	63
JOSEPH FORREST and H. L. FINSTON: The spectrochemical analysis of bismuth using a photoelectric spectrometer	71
D. R. MACKENZIE: Determination of U ²³⁵ burn-up in irradiated natural uranium by chemical and counting methods	72
R. G. HART, C. B. BIGHAM and M. LOUNSBURY: Analytical methods used in a study of reactivity changes with long irradiations	94
R. F. BUCHANAN, J. P. HUGHES and C. A. A. BLOOMQUIST: The colorimetric deter- mination of zirconium in plutonium-uranium-fissium' alloys	100
G. A. HUFF: Analytical program for processing stainless steel-uranium dioxide reactor fuel elements .	105
H. M. EILAND: A simplified method for radioiodine analysis	112
R. W. PERKINS: Radiochemical analysis of reactor effluent waste materials at Hanford	117
L. C. BOGAR and C. J. L. LOCK: Radiochemical analysis of corrosion product mixtures from high temperature pressurized water loop systems	133
J. W. ULLMANN: Reactor fuel element types	144
R. P. LARSEN and C. A. SEILS, JR.: The determination of U and Pu in U Pu alloys of the fission elements	145
J. J. McCown and R. P. LARSEN. Radiochemical determinations of Ce and total rare- earth elements by liquid-liquid extraction .	146

.

SECTION III. ANALYTICAL CHEMISTRY OF PLUTONIUM AND THE TRANSPLUTONIC ELEMENTS

CHARLES F. METZ and GLENN R. WATERBURY: Analytical laboratories for the handling of plutonium	149
J. H. COOPER: The high-alpha-radiation analytical facility of the Oak Ridge National Laboratory	154
C. E. PIFTRI and J. A. BAGLIO: The determination of plutonium based on National Bureau of Standards potassium dichromate	159
JOSEPH BUBERNAK, MARION LEW and GEORGE MATLACK: The determination of americiam plutonium alloy fuels for fast reactors	167
R. F. BUCHANAN, J. P. HUGHES, J. J. HINES and C. A. A. BLOOMQUIST: The determina- tion of nitrogen, americium, neptuniu 1, and uranium in p.p.m. quantities in pure plutonium	173
M. T. KELLEY, H. C. JONES and D. J. FISHER: Electronic controlled-potential coulo- metric titrator for plutonium analysis	185
W. S. TURNLEY: X-ray fluorescence analysis of plutonium	189
F. A. SCOTT and R. M. PEEKEMA: The determination of plutonium in irradiated uranium fuel solutions by controlled-potential coulometry .	196
R. W. STROMATT and F. A. SCOTT: Analysis for neptunium by controlled-potential coulometry	197

SECTION IV. THE ANALYSIS OF FISSION-PRODUCT MIXTURES

D. G. OLSON: Quantitative gamma ray spectrometric analysis of nuclide mixtures consecutive standard sources nullification	201
G. W. BOYES, JR., J. B. RAMSAY and R. T. PHELPS: Spectrochemical analysis of 'fissium' for cerium and lanthanum .	209
LYLE L. REED, MARCUS N. MYERS, and WILLIAM W. SABOL: A quantitative deter- mination of several short-lived iodine, barium and strontium fission products in gas-cooled reactor effluents	215
D. G. MULER and M. B. LEBOEUF: A radiochemical technique for determining fissionable material contamination on completed reactor fuel assemblies .	230
GLENN R. WATERBURY and CHARLES F. METZ: The spectrophotometric determination of alloying and fission product elements in nonirradiated plutonium 'fissium' alloys	237
JESSE W. T. MEADOWS, GEORGE M. MATLACK and GILBERT B. NELSON: Fission- product analysis of fast-reactor plutonium fuels	246
H. L. KRIEGER, J. E. GILCHRIST and S. GOLD: Concentration of radioactivity in detection of cobalt-60 and zinc-65 in rainout	254
A. C. LEAF: The determination of zirconium-95 plus niobium-95 by anion-exchange separation from other fission products	265
C. J. L. LOCK and L. C. BOGAR: The separation and radioactive assay of fission products from samples obtained from high-temperature pressurized-water loops	2?3
T. C. RAINS, H. P. HOUSE and OSCAR MENIS: Evaluation of flame photometry for the determination of elements of the rare-earth group	274
W. W. MORGAN, R. G. HART, F. C. MILLER and W. J. OLMSTEAD: Determination of the distribution of fission Xe in irradiated UO ₂ fuel elements	275
DANIEL L. LOVE and ALLEN E. GREENDALE: Radiochemical analyses through polaro- graphic methods—II. Rapid procedure for technetium and ruthenium radionuclides in fission products .	276
in fission products	2.0

THE ROLE OF THE ANALYTICAL CHEMIST IN NUCLEAR REACTOR TECHNOLOGY

C. D. SUSANO*

Associate Director, Analytical Chemistry Division, Oak Ridge National Laboratory[†], Oak Ridge, Tennessee

THE developments which have occurred in nuclear physics during the past 20 years have brought about significant and revolutionary changes in the domain of the analytical chemist; in that, whereas prior to the year 1943, he was concerned primarily with the analysis of non-radioactive materials and only rarely with radioactivity, the emphasis has changed to the extent that he has now become a part of a broad over-lapping discipline in which the utilization of nuclear reactions and the concomitant production of radiation and radioactive materials have become matters of the greatest importance and of real concern. In the development of this nuclear technology, many dilemmas have been encountered in that methods and procedures had to be, and are still being, developed for the production in relatively large quantities of materials which previously had been matters of obscure scientific interest. These developments, particularly in the chemical and metallurgical processing of uncommon metals and also in the fields of ceramics, electronics, and chemical engineering, are now being exploited most extensively to secure better materials for the construction, and the hoped for efficient and competitive operation, of nuclear reactors.

If the prediction of the future utilization of nuclear reactors in the production of electricity and of other forms of energy is, in fact, to be attained, two problems of real import face the analytical chemist: (1) the development of new and better methods of analysis, specifically for those materials which are and which will undoubtedly be developed and utilized in the construction and operation of reactors; and (2) the development of methods, techniques, and instruments for the automatic and remote control of processes and of reactors involving the handling of highly radioactive materials as matters of operations control, safety, by-product utilization and waste disposal.

In this new technology, the analytical chemist is perhaps a bit more fortunate than was his lot in the ferrous metals, and in many other industries, in that his role has been a continuing one since the early days of the nuclear weapons program wherein he has been given the opportunity to make his contributions simultaneously with those of the scientists and engineers engaged in the many revolutionary facets of the technology. With this very close association, the analytical chemist has given invaluable assistance to those engineers, chemists and physicists who have been charged with the responsibilities of processing and refining nuclear and other materials, including the fabrication and utilization of these materials into parts and components

^{*} General Chairman, Conferences on Analytical Chemistry in Nuclear Reactor Technology, held annually at Gatlinburg, Tennessee.

[†] Operated by Union Carbide Corporation for the U.S. Atomic Energy Commission.

of reactors. In this connection, it is worth noting that the analytical chemist has already made many real and significant contributions in the development of process methods for the separation of hafnium and zirconium; in the separation, purification and utilization of rare-earth elements; in the winning of uranium from extremely low-grade raw materials and the large-scale separation of this element and the U^{235} isotope from unwanted, deleterious process materials; in the creation of plutonium and the subsequent development of methods for its processing and refinement; and in many other equally important scientific and technological achievements. In citing these few examples, one is particularly impressed because some of these accomplishments, such as the separation and purification of rare-earth elements, represent the solution of problems which had previously been investigated more or less fruitlessly for more than a century by some of the most able chemists of their time. This is not to imply that the analytical chemist was in the main responsible for these accomplishments, but it cannot be denied that his contribution has been indeed most significant.

Even though remarkable advancements have been made in the accumulation of knowledge and of a better understanding of nuclear concepts, much still remains to be done in the utilization of materials in a manner that the full practical significance of these concepts can be understood and utilized for the eventual conversion of nuclear energy to a whole host of beneficial purposes.

In the further growth of an effective nuclear technology, it is important to note that the analytical chemist will be required to develop methods, procedures, and techniques for use in many lines of endeavor, some of which may appear to be completely unrelated to nuclear technology. In this regard, the analytical chemist working in the ferrous and non-ferrous metals industries will be called upon to provide better methods for the continued development of newer metals and alloys which will be useful in the design, construction and operation of reactors and appurtenances. Analytical chemists in other disciplines, such as biology, public health, ecology, food sterilization, agronomy, agriculture, oceanography and many more, will likewise be similarly involved in the development of improved methods of analysis with manifold beneficial effects on all aspects of human endeavor.

The present state of our knowledge of nuclear reactor technology is such that a considerable portion of the total effort in the field of nuclear energy is being directed toward the development of nuclear reactors which ultimately will produce thermal and electric power at economically acceptable prices. With the advent of second and third generation reactors and the launching of the NS Savannah, it is still quite obvious that considerable work needs to be done in the production, fabrication and evaluation of reactor components, and also of a wide variety of appurtenances and of auxiliary units, such as metals of construction, fuels, coolants, moderators, reflectors, shielding materials, control mechanisms, piping materials, pumping devices, pressure vessels, safety devices, instruments, filters, waste-disposal systems and byproduct recovery installations. Such properties as corrosion and erosion resistance, neutron absorption characteristics, and the effects of temperature, pressure and radiation on the physical properties of various components will require considerable experimentation and testing in prototypes and in smaller test units extending over a period of perhaps many years. In this work, the analytical chemist will be concerned with the analysis of liquids, solids, and gases which will have been subjected to wide varieties of exposure or produced under a multiplicity of conditions and which will vary considerably in the concentrations of many component substances. For example, in the case of coolant materials, he will be called upon to make analyses of lithium, sodium, potassium, or combinations thereof, for purity, extent of contamination, and to assist in the evaluation of certain corrosion tendencies in heat transfer systems and of the degrees of reactivity of these substances with other materials.

Much of the experimental testing of materials prior to the actual construction of prototypes and test models will be conducted under conditions simulating actual plant operation and, consequently, in these early stages, such experimentation involves the analysis of non-radioactive materials. As one proceeds, however, from the conceptual and experimental stages to the development of critical test reactors, and finally to actual operational reactors of increasingly greater capacity with exceptionally high neutron fluxes and concomitantly greater degrees of radioactivity, one soon reaches the other extreme where all reactor components will most surely be contaminated with hazardous radioactive materials which will require the application of methods of analysis under totally different conditions and with a maximum of consideration for the safety of the analyst and others.

In order to make a presentation such as this, it would be helpful if one could isolate as a first approximation those elements (or isotopes) from the periodic table which would appear to be of prime significance in nuclear reactor technology, but if one attempts to make any such simplifying assumption, he is immediately confronted with the incontrovertible fact that this is not a limited technology of a few elements or of simple mixtures of compounds, but is rather a highly complex technology in which he must be ready to evaluate the inter-relationships of all the elements in the periodic table, including thousands of isotopes. It may be possible at some later date, as knowledge is acquired of the relative merits of certain materials, to eliminate some of these from further consideration, but in the present state of the art any such simplification does not now appear, to put it mildly, to be either possible or warranted.

An awareness of the role of the analytical chemist in nuclear reactor technology is immediately obvious from the veritable flood of papers which have appeared on this subject in the technical literature over the past few years and nonetheless equally obvious is this substantiated further by the number of meetings, conferences and symposia which are being held in ever-increasing numbers. The significance of the application and utilization of remote manipulative techniques in this technology can be readily appreciated from the fact that the eighth in a series of Hot Laboratories and Equipment Conferences¹ is now in the planning stage. Even though not engendered specifically for the benefit of analytical chemists, much of the work that is being done in this field of specialized laboratory and equipment design and operation is of vital importance in the analytical chemistry of highly activated materials.

With regard to the many aspects of the analytical chemistry of radioactive materials alone, it is significant to note that Meinke² in a review of recent developments in the analytical chemistry of nucleonics cites no less than 1279 references of papers and reports which appeared in the world-wide scientific and technical literature during the two-year period between late 1955 and late 1957. This review is most extensive and will serve the analyst well in delineating the scope and multiplicity of his interests and responsibilities in the analysis of radioactive materials.

In a recent paper on the role of the analytical chemist in nuclear technology,

Kelley³ spells out the dual functions of the analyst in this field in that one of these is the identification and measurement of radioactive species and the other is the analysis of highly radioactive materials. He also points out the necessity for the development of remote-control instrumentation to effect the safe handling and analysis of substances with ever-increasing degrees of activity which are and will be encountered in the future development and utilization of nuclear reactors.

The historical account, by Rodden,⁴ of the role of the analytical chemist in the preparation of high-purity nuclear materials is a magnificent revelation of the importance of trace analysis in the development of suitable materials, particularly uranium, graphite and beryllium, which ultimately led to the production of "workable" atomic weapons during the war. As indicated by Rodden, this work is by no means complete; even though great progress has been made, to attain greater neutron economy, further improvements in methods will be required, specifically greater sensitivity, precision and accuracy at lower concentrations of all elements and, in a large number of cases, of certain isotopes which possess useful nuclear properties. In this respect, the methods for the analysis of uranium; graphite; beryllium, particularly the highly refractory oxide; and others, which have been examined most extensively, must be continued with even greater emphasis on the precise evaluation of minor components in the microgram and extending into the nanogram (10^{-9}) range.

In order to promote greater interchange of information on this subject, the Oak Ridge National Laboratory, in 1957, inaugurated a series of conferences on Analytical Chemistry in Nuclear Reactor Technology in which it was intended that information would be presented which bears directly on some of the scientific, technical and practical aspects of analytical chemistry as they apply specifically to the technology of nuclear reactors. The first conference⁵ was devoted to a review of recent developments in the analytical chemistry of certain of the more important reactor materials, specifically uranium, thorium, the rare-earth elements, zirconium, titanium, beryllium, graphite and heavy water, including a brief review of some of the recent applications of the methods of X-ray analysis, emission and mass spectroscopy, solvent extraction, and ion-exchange separations. In addition to providing a summary of recent developments in the analytical chemistry of each of the foregoing subjects, the discussions at that conference pointed out the serious need that now exists for the development and establishment of adequate standard samples and precise methods.⁶ This lack of standard authority has been and will continue to be a serious problem until standards of suitable quality become available. The development, design, construction and operation of nuclear reactors will depend in a large measure on the establishment of specification requirements which, most certainly, will be replete with exceedingly strict limitations with regard to the presence of certain substances, particularly those which seriously hamper the operation of reactors from the standpoint of neutron economy or wastage and also with respect to metals possessing high resistivity to corrosion or erosion and with high degrees of structural integrity. This matter of standards development has been investigated for a number of years by the Atomic Energy Commission and its precursor, the Manhattan Engineering District, in cooperation with the Atomic Energy Research Establishment of the United Kingdom, but, even though real progress has been made in the development of some standardization, the job is by no means complete. The recent utilization of newer industrial metals and alloys such as those of zirconium, hafnium, titanium, niobium, yttrium,

molybdenum, the alkali metals, rare-earth elements, such as dysprosium and gadolinium, boron, and special ferrous and non-ferrous alloys gives impetus to the urgency of this problem of securing better standard materials and the development of procedures of unquestioned precision and accuracy for evaluating sample purity and composition, coupled with high speed performance and automation.

Not only is the development of standard samples and procedures essential in the usual sense as applied to analytical chemistry, but even more profoundly is this need apparent in the application of the instrumental techniques such as emission and mass spectroscopy, absorption spectrophotometry, throughout the entire useful part of the electromagnetic spectrum, coulometry, flame photometry, radiochemistry, nuclear magnetic resonance, X-ray absorptiometry, fluorimetry, diffraction methods, gas chromatography, nuclear methods of analysis, polarography, and many other electrical methods based on the concept of electron transfer which can be utilized as a measure of the quality of certain materials.

During the fall of 1958, the Second Conference in the series on Analytical Chemistry in Nuclear Reactor Technology⁷ was again held at Gatlinburg, Tennessee, at which time analytical chemists from Great Britain, Canada and the United States really got down to the business of discussing some of the more important aspects of this work. Whereas the first meeting dealt specifically in generalities on the more recent developments, the emphasis at the second meeting was directed more toward the applications of analytical chemistry in prototype testing, particularly as it applied to the utilization of certain materials in various types of nuclear reactors. Perhaps one of the outstanding features of this Second Conference was the amount of discussion and interest which was shown in the matter of particle-size evaluation. At first glance, it would appear rather strange that a subject of this nature would be given any consideration in discussions on analytical chemistry. It is quite obvious in reading the papers on this subject, however, that the variety of analytical techniques used in this work does indeed make it a proper subject of analytical chemistry. The matter of particle-size analysis is very important in the operation of the thermal breeder reactor where successful operation depends in a large measure on the thorough knowledge of the flow, settling and nuclear characteristics of slurries and other heterogeneous mixtures which contain fertile and fissionable materials. The analytical techniques involved in these analyses are, in a large number of cases, essentially the same as those that are used in the analysis of solutions but are complicated by the one requirement that the estimation of certain constituents need be made on fractional portions of samples rather than on the basis of the total sample. By and large, these analyses are made by first separating the solids into groups of differing particle size by the application of settling techniques, based primarily on Stoke's law. All the known methods of particle-size evaluation were discussed at this meeting, in addition to which some of the newer techniques, such as flying-spot microscopy, and electronic sizing were also discussed. The intense interest shown in these sessions does indeed reveal that much needs to be done in the development of methods for the analysis of subsieve particulate matter.

In the Second Conference, many of the papers continued the discussions of the first session on the utilization of analytical techniques, particularly on X-ray and emission spectroscopy, nuclear methods of analysis, flame photometry, and instrumentation, but in which the preponderance of interest had to do more with unusual mixtures, such as newly developed alloys, fuel materials, boron and specialized preparations.

The Third Conference,⁸ which was held in October 1959, was devoted primarily to the consideration of the analysis of materials and components after having been subjected to different degrees of exposure to neutron activation in operating reactors. The principal point of interest in the analytical chemistry of such materials is the high degree of activity which is encountered and the necessity for carrying out this function in a manner completely foreign to the usual methods of analysis. Much of this work must, of necessity, be carried out in separate installations designed primarily for the remote handling of materials having a high degree of radioactivity. Some of the specially designed laboratories and the details of handling and analyzing extremely "hot" materials in those installations were considered in several of the papers.

In the operation of nuclear reactors, the analytical chemist is called upon to make analyses of mixtures containing a large number of fission products and to continue this assistance through the processing and reprocessing cycles which are intended to separate certain valuable radioisotopes, fertile materials, unburned fuel, plutonium, transplutonic elements and waste products. The successful utilization of by-products and the production of high-purity materials of useful import that will be derived from such reactor operations will constantly rest on the ability of the analytical chemist to develop better methods of control. Typical case histories of some of those more important problems which have been encountered by the analytical chemist were presented in some detail. As was the case at the first meeting, the matter of standardization was again taken up by Dr. Rodden, at which time he also indicated the progress that is being made in this particular field and how different installations are meeting this problem. In order to keep up to date, virtually all of one session was devoted to some of the more recent developments in analytical methodology, even though much of this subject matter did not necessarily follow the main theme of this particular meeting.

At this point, it is interesting to note that, with the advent of nuclear technology, the analytical chemist has again been quick to adapt those instruments and methods of other fields of science, particularly physics, to his repertoire for use in the evaluation of the composition of matter in any form. Even though one ordinarily associates a chemist with the study of chemical reactions, it has always been the attitude of the analytical chemist that if a device or technique can be of any service in establishing the composition of a given sample, the innovation should be used regardless of its source or origin.

In this regard, it is becoming increasingly apparent that mass spectrometry, nuclear magnetic resonance and counting techniques, insofar as they are useful in the evaluation of compositions of matter, to mention only a few of the more recently developed and widely applied methods, are part and parcel of the analytical chemist's "kit of tools" on which he can draw in carrying out his responsibilities. In fact, future workers in this field must of necessity prepare themselves to meet these added responsibilities if they intend to keep pace with the future trends of this profession. There seems to be little, if any, chance that the physicist will ever consider the desirability of developing a new field of endeavor under the heading of, say, analytical physics. If he does, he will find that one of the basic essentials in the performance of this work is a thorough knowledge of the principles and concepts of chemistry and in

the end the field of analysis concerned with the composition of matter would undoubtedly be returned to the chemist. In any case, it is doubtful that the analytical chemist need even be remotely or mildly concerned about this possibility.

In considering the role of analytical chemistry in nuclear reactor technology, the analyst's attention is naturally first drawn to the basic raw materials-the fissionable isotopes of U²³³, U²³⁵ and Pu²³⁹ or, more appropriately, the sources of these isotopes, the naturally occurring mixtures of thorium and uranium isotopes. In view of the limited extent of the known world reserves of conventional fuel materials, of the future potential of nuclear power reactors and of the scarcity of known high-grade uranium ore deposits, particularly of domestic origin, considerable activity in exploratory searches has been going on for a number of years in an effort to find high-grade sources of these materials or to utilize low-grade sources economically. This search has not only been concerned with the usual source of valuable minerals but has been extended to an examination of waste materials and by-products of other industries. Regardless of the source of these potential reserves, the methods of establishing the uranium or thorium content are basically those developed by the analytical chemist, be it by the use of wet-chemical procedures or a variety of instrumental techniques. The extremely wide dispersion of these elements, particularly uranium, in virtually insignificant amounts in all varieties of materials in the earth's crust, has dictated the development of procedures suitable for use on materials of considerable variability and of low uranium or thorium concentration. In this regard, the analytical chemist has been of invaluable assistance in the development of methods for the exploration and evaluation of sources of uranium and thorium-containing materials; such as,

- 1. The monazite sands of India, Brazil and to a lesser extent the low-grade, domestic and Canadian deposits of thorium-containing materials.
- 2. The high-grade deposits of uranium in Africa and Canada.
- 3. Lower-grade deposits of uranium in the mineral deposits of the Colorado plateau and the waste stockpiles of low-grade materials from the mining of vanadium and radium.
- 4. Low-grade sources of shale, lignite, sea water, natural wells and streams, the tailings of the gold mines of South Africa, and the huge piles of waste materials from the phosphate industry.

In all of these and many others, the analytical chemist has been and will continue to be called on to develop methods and procedures, and to make the analyses necessary for the prospecting, mining, milling, processing, and recovery of the mineral values from the so-called ore bodies and control of these operations to provide a product of suitable quality for utilization in the atomic energy industry. Similarly, this activity must be adaptable to the recovery and exploitation of other valuable byproducts. As an example, it may be foreseen, but perhaps not too clearly, how the recovery of uranium from lignite may well provide a margin sufficient for the economical utilization of lignite in supplementing the reserves of fossil fuels essential to the petroleum and petrochemical industries.

In all this analytical work, the important function, of course, is the precise and accurate control of the quality of raw materials and intermediates in a highly complex series of processing steps designed to yield fuel materials in a multiplicity of shapes, forms, concentrations, and configurations. Since, as a condition precedent, the

C. D. SUSANO

feasibility and efficient operability of nuclear reactors is highly dependent on strict limitations of certain impurities in the fuel, it is obvious that methods and procedures for the analysis of fuel components must be inherently sensitive, accurate and precise throughout all the process cycles and recycles of mining, milling, refining, conversion, isotope separation, fuel fabrication and fuel reprocessing. An enumeration of the methods and procedures of analytical control of the fuel cycle is completely outside the scope of this paper; it is sufficient, however, to note that the literature on this subject is voluminous and significantly good but by no means complete. The compilation of Rodden⁹ serves as an excellent starting point on the analytical chemistry of uranium, but with the enforced development of methods and procedures occasioned by the search for domestic fuel sources and the increased emphasis on the peaceful utilization of nuclear energy, a large number of methods and procedures have been developed for application to specific types of materials and process cycles. In order to achieve the highest degrees of precision, accuracy, sensitivity and economy of operations, not only in the analytical chemistry of the fuel cycle but in all aspects of nuclear technology, it is interesting to observe that methods of separation, particularly those based on solvent extraction and the use of ion-exchange resins, have received considerable attention with highly fruitful results. An example of the practical importance of solvent extraction and, incidentally, of the utilization of analytical techniques to process application is best exemplified by the adaptation of the solvent extraction of uranyl nitrate with diethyl ether as an efficient war-time method for the purification of uranium. The first successful refining process for preparing zirconium, free of hafnium, was also similarly accomplished by the adaptation of an analytical method based on the use of hexone and ammonium thiocyanate. In the field of ion-exchange adsorption, the work of W. E. Cohn, F. H. Spedding, G. E. Boyd, K. A. Kraus, Robert Kumin and their associates and many others in the separation and purification of the rare-earth elements was particularly brilliant and opened up a wide field which at present is being explored most successfully for other adaptations both in analytical chemistry and process technology. In this regard, the work of Kraus et al.¹⁰, Wish et al.¹¹ and others has been most significant in providing break-throughs to what previously had been considered difficult and in some cases insoluble problems. As a further step in the development of separative processes of the highest integrity, the successful use of procedures based on combinations of solvent extraction¹² and ion-exchange separations¹³ has given further impetus to the solution of many difficult problems in analytical chemistry, including most particularly the separation and isolation of the components in radioactive mixtures prior to the application of methods of estimation.

In considering further this matter of the analytical chemistry of the fuel cycle, it is to be noted that one of the foremost problems which confront the nuclear technologist is the continued development of in-line instrumentation and automatic devices to measure, evaluate, control and correct the compositional characteristics of process materials as they flow through the many process streams from the mine, through feed preparation, reactor operation, reprocessing, by-product recovery and waste disposal. In carrying out his responsibilities in this regard, the analytical chemist must work in close harmony with the process engineer, the reactor technologist, the safety engineer, the health physicist, the statistician, and the design engineers, interested in the many facets of construction, process development and instrumentation. Even though many operations in the fuel cycle lend themselves to simple methods of control, based on functional and direct measurements of operational variables, much remains to be done in those operational areas wherein the exact parametric relationships have not as yet been established. Until such time as these relationships are evaluated or reasonably understood, the work must be done on an experimental basis with simultaneous consideration being given to the development of these operational methods of control. It goes without saying that much of this work will prove difficult, particularly as it may apply to the control of heterogeneous combinations of materials flowing at high velocities in peculiarly designed and intricate matrices of piping, and heat-transfer units, at high temperatures and at high pressures, exposed to intense neutron fluxes and high degrees of radioactivity behind enormous shields and barricades which rule out the usual, simple methods of control.

REFERENCES

- ¹ The Eighth Conference on Hot Laboratories and Equipment, held in San Francisco, California, December, 1959.
- ² W. W. Meinke, Anal. Chem., 1958, 30, 686.
- ³ M. T. Kelley, Anal. Chem., 1957, 29, 21A.
- ⁴ C. J. Rodden, J. Chem. Educ., 1959, 36, 458.
- ⁵ Proceedings of the First Conference on Analytical Chemistry in Nuclear Reactor Technology, Office of Technical Services, Department of Commerce, Washington 25, D.C., TID-7555.
- ⁶ Anal. Chem., 1957, 29, 33A.
- ⁷ Proceedings of the Second Conference on Analytical Chemistry in Nuclear Reactor Technology, Office of Technical Services, Department of Commerce, Washington 25, D.C., TID-7568.
- ⁸ Third Conference on Analytical Chemistry in Nuclear Reactor Technology held in Gatlinburg, Tennessee, October 26–9, 1959.
- ⁸ C. J. Rodden, Analytical Chemistry of the Manhattan Project, McGraw-Hill, New York (1950).
- ¹⁰ K. A. Kraus, H. O. Phillips, T. A. Carlson and J. S. Johnson, Ion Exchange Properties of Hydrous Oxides, Proceedings of the Second United Nations International Conference on the Peaceful Uses of Atomic Energy, 1958, 28, P/1832, p. 3.
- ¹¹ L. Wish, Anal. Chem., 1959, 31, 326.
- ¹² G. H. Morrison and H. Freiser, *Solvent Extraction in Analytical Chemistry*, John Wiley, New York (1957).
- ¹⁸ L. R. Bunney, N. E. Ballou, J. Pascual and S. Foti, Anal. Chem., 1959, 31, 324.

AUTHOR INDEX

Baglio, J. A. 159 Bigham, C. B. 94 Bloomquist, C. A. A. 100, 173 Bogar, L. C. 133, 273 Boyes, G. W., Jr. 209 Bubernak, J. 167 Buchanan, R. F. 100, 173 Cooper, J. H. 154 Davis, W. F. 28 Dunn, H. W. 42 Eilanu, H. M. 55, 71 Finston, K. L. 55, 71 Fisher, D. J. 185 Forrest, J. 71 Gaddy, R. H. 52 Gilchrist, J. E. 254 Gold, S. 254 Greendale, A. E. 276 Hakkila, E. A. 46 Hart, R. G. 94, 275 Hibbits, J. O. 28 Hines, J. J. 173 House, H. P. 274

Hutf, G. A. 105 Hughes, J. P. 100, 173 Jones, H. C. 185 Kelley, M. T. 185 Krieger, H. L. 254 Lamb, C. E. 20 Larsen, R. P. 145, 146 Leaf, A. C. 265 Leboeuf, M. B. 230 Lew, M. 167 Lock, C. J. L. 133, 273 Lounsbury, M. 94 Love, D. L. 276 MacKenzie, D. R. 72 Manning, D. L. 30 Matlack, G. M. 167, 246 McCown, J. J. 146

Matlack, G. M. 167, 24 McCown, J. J. 146 Meadows, J. W. T. 246 Menis, O. 30, 274 Menke, M. R. 28 Metz, C. F. 149, 237 Miller, F. C. 275 Morgan, W. W. 275 Myers, M. N. 215 Nelson, G. B. 246 Newman, L. 55 Olmstead, W. J. 275 Olson, D. G. 201 Peekema, R. M. 196 Perkins, R. W. 117 Phelps, R. T. 209 Pietri, C. E. 159 Prach, T. 55 Rains, T. C. 274 Ramsay, J. B. 209

Reed, L. L. 215 Rodden, C. J. 3 Sabol, W. W. 215 Saunderson, J. L. 63 Scott, F. A. 196, 197 Seils, C. A., Jr. 145 Stromatt, R. W. 197 Susano, C. D. i, iii Turnley, W. S. 189 Ullmann, J. W. 144

Waterbury, G. R. 46, 149, 237