BASIS FOR SUBCRITICAL LIMITS IN PROPOSED CRITICALITY SAFETY STANDARD FOR MIXED OXIDES

J. H. CHALMERS,[†] GORDON WALKER,[‡] N. KETZLACH,** Richland, Washington 99352

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E. D. CLAYTON, H. K. CLARK,* D. W. MAGNUSON.* RYOHEI KIYOSE,⁺⁺ C. L. BROWN, D. R. SMITH,^{††} and R. ARTIGAS^{‡‡} Battelle Northwest Laboratories, Battelle Boulevard

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Subcommittee 8 of the Standards Committee of the American Nuclear Society has proposed a standard providing subcritical limits for operations with mixed oxides of plutonium and uranium. The subcritical limit is the limiting value assigned to a controlled parameter that results in a system known to be subcritical, provided the limiting value of no other controlled parameter of the system is violated. The proposed standard includes subcritical limits for mixed oxides containing up to 30 wt% plutonium in Pu + U. A review was made of the available experimental data and validations undertaken that serve as the basis of the limits, and the assertion that they are, indeed, subcritical as given.

INTRODUCTION

The American National Standard for Nuclear Criticality Safety in Operations with Fissionable Materials Outside Reactors (N16.1-1975) provides single-parameter limits for operations with ²³⁵U, ²¹³U, and ²³⁹Pu (Ref. 1). Subcommittee 8 of the Standards Committee of the American Nuclear Society is proposing a standard for operations with mixed oxides, extending N16.1 chiefly through the inclusion of additional subcritical limits. These limits may prove valuable for operations with mixed oxides of plutonium and uranium encountered in light water reactor, liquid-metal last breeder reactor, and gas-cooled fast reactor fuel cycle operations. As defined in N16.1, a subcritical limit is the limiting value assigned to a controlled parameter that results in a system known to be subcritical provided the limiting value of no other controlled parameter of the system is Violated. This limit contains margins designed to

CHEMICAL PROCESSING

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be sufficiently large to allow for uncertainties in calculations and experimental data used in its derivation, but, simultaneously, sufficiently small to discourage attempts to justify a larger limit. The margins contain no allowances for operating contingencies, e.g., double batching or a failure of analytical techniques to yield accurate values. Therefore, process specifications must incorporate margins to protect against the consequences of uncertainties in process variables and against a limit being accidentally exceeded.

The selection of limits with sufficiently, but not excessively, large margins is necessarily somewhat arbitrary, and requires the exercise of judgment, particularly, as in the present case, for mixed oxides, where pertinent experimental data are sparse. N16.1 offers the following guidance: "In the absence of directly applicable experimental measurements, the limits may be derived from calculations made by a method shown to be valid by comparison with experimental data, provided sufficient allowances are made for uncertainties in the data and in the calculations." The

- ±United Kingdom Atomic Energy Authority, Safety and Reliability Directorate, Laboratories and Transport Section, Wigshaw Lane, Culcheth, Warrington, Lancashire WA3 4NE England.
- **US Nuclear Regulatory Commission, Division of Fuel Cycle and Material Safety, Fuel Processing and Fabrication Branch, Washington, D C 20555.
- ++University of Tokyo, Department of Nuclear Engineering, 7-3-1, Hongo, Bunkyo-ku, Tokyo, Japan.
- ††Los Alamos Scientific Laboratory, PO. Box 1663, Los Alamos, New Mexico 87545.
- ‡‡General Electric Company, Nuclear Energy Systems Division, 175 Curtner Avenue, San Jose, California 95125.

^{*}E. I. du Pont de Nemours and Company, Savannah River Laboratory, Theoretical Physics Division, Aiken, South Carolina 29801.

⁺Union Carbide Corporation, Oak Ridge Gaseous Diffusion Plant, Oak Ridge, Tennessee 37830.

[†]Department of Trade and Industry, Nuclear Installations Inspectorate, NII 2 Silkhouse Court, Tithebarn Street, Liverpool, Lancashire L2 2LZ, England.

American National Standard, Validation of Calculational Methods for Nuclear Criticality Safety (N16.9-1975), offers further guidance. The material that follows gives the limits selected by the Work Group and the basis for asserting that they are indeed subcritical. Whenever numerical limits are given in a Standard, it is important that their basis be published in the open literature, where it may be examined by the user of the Standard.

SCOPE OF PROPOSED STANDARD

Although the limits may eventually be extended to lattices and other heterogeneous systems, they are presently restricted to homogeneous systems of plutonium and uranium dioxide. The mixtures may be solutions, suspended solids, precipitates, or a mechanical blend of powders, and are fully reflected by water. The following compositions were selected as having the greatest utility. The oxide mixtures contain 30, 15, 8, and 3% PuO₂ by weight. The uranium is natural. Isotopic compositions of plutonium are 100% ²³⁹Pu; 79% ²³⁹Pu, 15% ²⁴⁰Pu, 6% ²⁴¹Pu; or 60% ²³⁹Pu, 25% ²⁴⁰Pu, and 15% ²⁴¹Pu. (Plutonium-238 and -242 may be considered present, but were conservatively ignored in calculating limits.)

Subcritical limits were derived for both dry and water-moderated systems. Since a completely dry oxide system may be difficult to maintain, subcritical limits were derived also for damp $[H/(Pu + U) \le 0.45]$ oxide.² Solutions and slurries were assumed to be uniform homogeneous mixtures of UO₂ [10.96 g/cm³ (10.96 × 10³ kg/m³)] and PuO₂ (11.46 g/cm³ for ²³⁹PuO₂) in water. Critical dimensions of such systems have minima as a function of concentration, provided the oxide mixture does not contain much more than 30% PuO₂ by weight.

SUBCRITICAL LIMITS

Uniform Aqueous Mixtures

Limits for uniform aqueous mixtures³⁻⁵ of plutonium and uranium (²³⁵U ≤ 0.71 wt%) fully reflected by water are given in Table I. A margin of ~2% (see next section) in k_{eff} was considered sufficient to account for uncertainties in calculations and experimental data used in deriving the limits on mass, volume, cylinder diameter, and slab thickness. Estimated critical values are shown in Figs. 1 through 4 along with the subcritical limits for a plutonium isotopic composition of 100% ²³⁹Pu to indicate the corresponding margin in terms of mass or dimension. The margins for the subcritical limits with higher isotopes would be similar.



For sufficiently dilute mixtures, criticality is impossible regardless of mass. A margin in k approaching 5% was used in deriving the infinite sea concentration limits in Table I, because uncertainties were considered greater than for mass or dimension limits. The corresponding margin in terms of concentration approaches 10%.

The product of critical slab thickness and concentration has a minimum, leading to a limiting areal density that is useful where precipitation or evaporation is a credible possibility. As for the infinite sea concentration, uncertainties

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	3 wt% I	PuO₂ in PuC	$v_2 + UO_2$	8 wt% P	uO2 in PuO	2 + UO2	15 wt% PuO ₂ in PuO ₂ + UO ₂			30 wt% PuO ₂ in PuO ₂ + UO ₂		
	(A) ^a	(B) ^a	(C) ^a	(A)	(B)	(C)	(A)	(B)	(C)	(A)	(B)	(C)
Mass of plutonium contained in oxide (kg) (Total oxide mass,	0.73	1.35	2.00	0.61	1.06	1.53	0.54	0.94	1.28	0.50	0.87	1.16
$PuO_2 + UO_2$ (kg)	(21.3)	(51.5)	(10.0)	(0.0)	(13.1)	(21.1)	(4.1)	(1.1)	(0.1)	(1.0)	(0.0)	(1.1)
Cylinder diameter (cm) $(\times 10^{-2} = m)$	24.3	30.8	34.8	19.8	24.9	27.5	17.8	22.5	24.8	16.2	21.0	23.4
Slab thickness (cm) $(\times 10^{-2} = m)$	11.0	14.9	17.4	8.2	11.2	12.9	6.9	9.6	11.0	5.9	8.7	9.9
Volume (1)	23.5	44.8	63.4	14.0	25.9	34.4	11.0	20.4	26.6	8.5	16.8	21.6
Infinite sea limiting ^b subcritical concentra- tion of plutonium contained in oxide (g/ℓ) (×10 ⁻³ = kg/ ℓ)	6.8	8.06	9.27	6.9	8.19	9.43	7.0	8.16	9,39	7.0	8.12	9.32
(H/Pu atom ratio) ^c	(3780)	(3203)	(2780)	(3780)	(3210)	(2790)	(3780)	(3237)	(2818)	(3780)	(3253)	(2848)
(Total oxide limiting concentration, $PuO_2 + UO_2$) (g/l) (×10 ⁻³ = kg/l)	(257)	(305)	(351)	(97.8)	(116)	(134)	(52.9)	(61.7)	(71.0)	(26.5)	(30.7)	(35.2)
Areal density of plutonium contained in oxide (g/cm ²) (×10 = kg/m ²)	0.27	0.38	0.47	0.25	0.34	0.42	0.25	0.33	0.41	0.24	0.32	0.37
(Total areal density of mixed oxides, $PuO_2 + UO_2$) (g/cm ²) (×10 = kg m ²)	(10.2)	(14.4)	(17.7)	(3.5)	(4.8)	(5.9)	(1.9)	(2.5)	(3.1)	(0.9)	(1.2)	(1.4)

^a Conditions on plutonium isotopic ratios: (A) = 240 Pu > 241 Pu; (B) = 240 Pu ≥ 15 wt%, 241 Pu ≤ 6 wt%; and (C) = 240 Pu ≥ 25 wt%, 241 Pu ≤ 15 wt%. ^b For plutonium content in mixed oxides in the range below 3 wt% to 0.13 wt%, the subcritical limit of Table III is controlling. The plutonium concentration limit corresponding to 0.13 wt% is ≤ 4.9 g Pu/l. For PuO₂ content ≤ 0.13 wt%, an aqueous, homogeneous mixture of mixed oxides will remain subcritical irrespective of any H/Pu ratio or concentration of $PuO_2 + UO_2$ in the mixture. (A reduction in the subcritical concentration below the value of 6.8 g/l at 3 wt% PuO_2 is required to account for the presence of ²³⁵U in the uranium, which becomes relatively more important at lower plutonium contents. The H/Pu ratio of 3780 will ensure subcriticality, if utilized for control, but at 0.13 wt% PuO2, the corresponding plutonium concentration would be down to 4.2 g Pu/L, which is a value somewhat less than required for the subcritical limit if the limit is expressed in terms of concentration or g Pu/ℓ .) ^cLower limit. All other limits are upper limits.

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Fig. 3. Cylinder diameter limit versus plutonium content.



Fig. 4. Slab thickness limit versus plutonium content.

were considered greater than for mass or dimension limits, and a margin of $\sim 4\%$ in $k_{\rm eff}$ was used in deriving the areal density limits of Table I.

Dry and Damp Oxides

Subcritical mass limits for units of mixed oxides are given in Table II. Calculations were made for an isotopic composition of 100% ²³⁹Pu, but the limits apply to other compositions. A considerable effort (see next section) was put forth in deriving these limits. In some cases, as many as six "independent" calculations were made. The smallest limits were selected, with a margin in $k_{\rm eff}$ estimated in these calculations to be $\sim 3\%$. In terms of mass, the corresponding margin increases with mass. For dry 30% PuO₂, it is ~9%, and for damp, half-density 8% PuO₂, nearly 50%. The densities listed are the theoretical values for dry and damp mixed oxides. Subcritical limits are also included for damp mixed oxides at $\frac{1}{2}$ theoretical density.

	<u> </u>			
30 wt% PuO2	26.1 kg Pu (98.6 kg PuO ₂ + UO ₂ at density ≤ 11.11 g PuO ₂ + UO2/cm ³)	23.3 kg Pu (88.1 kg PuO ₂ + UO ₂ at density ≤ 9.52 g PuO ₂ + UO ₂ /cm ³)	67.9 kg Pu (256.6 kg PuO ₂ + UO ₂ at density ≤ 4.76 g PuO ₂ + UO ₂ /cm ³)	Just a Just a in a hom enriched enrichme in natura cality, r fration 1 given in for mixe
$15 wt\% PuO_2$	47.0 kg Pu (355 kg PuO ₂ + UO ₂ at density ≤ 11.04 g PuO ₂ + UO ₂ /cm ³)	32.9 kg Pu (249 kg PuO ₂ + UO ₂ at density ≤ 9.47 g PuO ₂ + UO ₂ /cm ³)	102 kg Pu (771 kg PuO2 + UO2 at density ≤ 4.73 g PuO2 + UO2/cm ³)	in water margin i pp s margin i jures, ~2 o s s in water margin i jures, ~2 o s in water subc Subc
8 wt% PuO2	122 kg Pu (1729 kg PuO ₂ + UO ₂ at density ≤ 11.00 g PuO ₂ + UO ₂ /cm ³)	49.4 kg Pu (700 kg PuO ₂ + UO ₂ at density ≤ 9.44 g PuO ₂ + UO ₂ /cm ³)	161 kg Pu (2282 kg PuO ₂ + UO ₂ at density ≤ 4.72 g PuO ₂ + UO ₂ /cm ³)	Market Wisk Market Wisk Marke
3 wt% PuO2	Subcritical in any amount	236 kg Pu (8919 kg PuO₂ + UO₂ at density ≤ 9.42 g PuO₂ + UO₂/cm³)	885 kg Pu (33 447 kg PuO ₂ + UO ₂ at density ≤ 4.71 g PuO ₂ + UO ₂ /cm ³)	the second secon
	<i>Dry</i> mixed oxides at theoretical density	Damb mixed oxides at theoretical density, $H/(Pu + U) \le 0.45$	Damp mixed oxides at one-half theoretical density, $H/(Pu + U) \le 0.45$	Wildation o Wildation o of Bias (Ref. For limits, perform section the resu calculat secure s
	3 wt% PuO ₂ 8 wt% PuO ₂ 15 wt% PuO ₂ 30 wt% PuO ₂	$D\gamma \text{ mixed oxides at theoretical Subcritical in any amount} \begin{bmatrix} 3 \text{ wt\% PuO}_2 \\ D\gamma \text{ mixed oxides at theoretical Subcritical in any amount} \\ \text{density} & 122 \text{ kg Pu} \\ \text{density} & 11.00 \text{ g PuO}_2 + \text{UO}_2 \text{ at } \\ \text{density} & 11.04 \text{ g PuO}_2 + \text{UO}_2 \text{ at } \\ \text{density} & 11.11 \text{ g PuO}_2 + \text{UO}_2 \text{ at } \\ \text{density} & 11.04 \text{ g PuO}_2 + \text{UO}_2 \text{ cm}^3 \\ \text{density} & -102\sqrt{\text{cm}}^3 \\ \text{maxed} & -102\sqrt{\text{cm}^3 \\ \text{maxed} & -102\sqrt{\text{cm}$	$ \begin{array}{l lllllllllllllllllllllllllllllllllll$	Dry mixed oxides at theoretical 3 wt% PuO_2 8 wt% PuO_2 15 wt% PuO_2 15 wt% PuO_2 30 wt% PuO_2 30 wt% PuO_2 Dry mixed oxides at theoreticalSubcritical in any amount $122 \text{ kg PuO_2 + UO_2}$ at 47.0 kg Pu 26.1 kg Pu $density$ Long theoreticalSubcritical in any amount $122 \text{ kg PuO_2 + UO_2}$ at 47.0 kg Pu 26.1 kg Pu $density$ Long theoreticalSubcritical in any amount $122 \text{ kg PuO_2 + UO_2}$ at 47.0 kg Pu $26.1 \text{ kg PuO_2 + UO_2}$ at $density$ $\pm UO_2/cm^3$ $\pm UO_2/cm^3$ $\pm UO_2/cm^3$ $\pm UO_2/cm^3$ $\pm UO_2/cm^3$ $Damp$ mixed oxides at theoretical $236 \text{ kg PuO_2 + UO_2}$ at 49.4 kg Pu $32.9 \text{ kg PuO_2 + UO_2}$ at $49.02 \text{ ebsisty} \leq 9.4.7 \text{ gPuO_2}$ $Damp$ mixed oxides at one-half 835 kg PuO_2 $40.02 \text{ ebsisty} \leq 9.4.4 \text{ gPuO_2}$ $24.02 \text{ ebsisty} \leq 9.4.7 \text{ gPuO_2}$ $40.02 \text{ ebsisty} \leq 9.4.7 \text{ gPuO_2}$ $Damp$ mixed oxides at one-half 835 kg Pu 100 kg PuO_2 $1002/cm^3$ $40.02 \text{ ebsisty} \leq 9.4.7 \text{ gPuO_2}$ $Damp$ mixed oxides at one-half $835 \text{ kg PuO_2$ $1002 \text{ ebsisty} \leq 9.4.7 \text{ gPuO_2}$ $1002/cm^3$ $40.02 \text{ ebsisty} \leq 9.4.7 \text{ gPuO_2}$ $Damp$ mixed oxides at one-half 835 kg PuO_2 $1002 \text{ ebsisty} \leq 9.4.7 \text{ gPuO_2}$ $40.02 \text{ ebsisty} \leq 9.4.7 \text{ gPuO_2}$ $Damp$ mixed oxides at one-half 835 kg PuO_2 $1002/cm^3$ $1002/cm^3$ $40.02 \text{ ebsisty} \leq 9.4.7 \text{ gPuO_2}$ $Damp$ mixed oxides at one-half $835 $

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Autonium Enrichment

Just as uranium oxide cannot be made critical in a homogeneous aqueous mixture without being enriched in ²³⁵U or as dry oxide without further enrichment, so there are concentrations of ²³⁹Pu in natural uranium that are necessary for criticality, regardless of mass. Subcritical concentration limits of ²³⁹Pu in natural uranium are given in Table III for dry and damp mixed oxides, for mixed nitrates in water, and for mixed oxides in water. For the dry and damp oxides, the margin in k was ~5%, and for the aqueous mixures, ~2%.

TABLE III

Subcritical Limits for ²³⁹ Pu Content in Uranium
$(^{235}U \leq 0.71 \text{ wt\%})$ Applicable to
Unrestricted Quantities

	Plutonium Content
Dry Mixed Oxides ^a	
Limiting subcritical wt $\%$ ²³⁹ Pu in Pu + U	4.4 wt%
Damp Mixed Oxides ^a	
$H/(Pu + U) \le 0.45$; limiting subcritical wt% ²³⁹ Pu in Pu + U	1.8 wt%
Wet Mixed Oxides ^a	
Limiting subcritical wt% of 239 Pu in Pu + U	0.13 wt%
Aqueous Nitrate Solutions	
Limiting subcritical wt% of ²³⁹ Pu in Pu + U in the presence of 4 nitrate ions per Pu atom [Pu(NO ₃) ₄], ^[4] and two nitrate ions per uranium atom [UO ₂ (NO ₃) ₂]	0.65 wt%

These limits are not applicable to atom mixtures of plutonium and uranium, but are restricted to the oxides of these nuclides ($PuO_2 + UO_2$).

Validation of Methods and Establishment of Bias (Ref. 6)

For the purpose of deriving the subcritical limits, several calculations were independently performed, utilizing various codes and crosssection sets. To assess the confidence with which the results of the calculations can be applied, the calculations were validated against pertinent experimental data. Unfortunately, the "areas of applicability" (quoting N16.9) defined by available data do not include many of the conditions of

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interest for mixed oxides, and "extensions" were required. Agreement between results of diverse methods with different cross-section sets' (although no guarantee) was taken to be a good indication of accuracy. Limits were not taken to be greater than the smallest critical values obtained with a reasonably good method, but in some cases were not much less where other methods gave generally larger values. Particular care was exercised to calculate limits by the various codes and cross-section sets in a manner consistent with that adopted for performing correlations.

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The MONK Monte Carlo code, a neutronics code derived from GEM, has been used extensively in the U.K. for criticality calculations.^{7,8} A sample selection of correlations with this code by two of us (Chalmers and Walker) is given in Table IV. The choice of experiments includes mixed oxides with PuO₂ contents of 30, 14.62 and 7.89 wt% and H/(Pu + U) ratios of 47.4, 30.6, and 51.8, respectively⁹; and plutonium oxides with moderation ratios of 0.04, 15, and 50 (Refs. 10, 11, and 12). The results of the calculations on the experimental systems are included in Table IV. Note that the bias is mainly to overestimate reactivity.

The computational methods selected by another of us (Clark, Savannah River Laboratory) for dry and damp mixed oxides (Table II limits) was S_n as implemented by the ANISN code.¹³ The cross sections were Hansen-Roach¹⁴ with ²³⁸U resonance cross sections modified by J. R. Knight (Oak Ridge National Laboratory). Cross sections for ²⁴¹Pu at infinite dilution were furnished by Smith (Los Alamos Scientific Laboratory). Cross sections for ²³⁵U, ²³⁸U, ²³⁹Pu, and ²⁴⁰Pu at resonance energies were selected by linear interpolation of tabulated values as a function of the logarithm of the total potential scattering cross section per absorber atom. The cross-section set for hydrogen was that obtained by fission spectrum weighting. The fission spectrum for ²³⁵U or for ²³⁹Pu was used, depending on which nuclide was predominant. For infinite systems (Table III limits; infinite sea concentration, Table I) the B_1 method was selected, as implemented by HRXN with Hansen-Roach cross sections and by GLASS (Ref. 15), with essentially HAMMER (Ref. 16) cross sections.

These methods were validated by correlation with a number of critical experiments pertinent to dry and damp mixed oxides. Except for experiments with PuO_2 (Ref. 10), which were not considered by Clark, no experiments have been performed with dry or even damp [H/(U + Pu) =0.45] oxides, but experiments have been performed at fairly low (~3) ratios of hydrogen to fissionable atoms with various ratios of fissionable to fissile atoms. Experiments have also been performed

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TABLE IV

Correlations with Experiment Using MONK Monte Carlo Code* (Refs. 7 and 8)

Experiment Number	Composition	Reflector	k _{eff}	Standard Deviation	Reference
1	$PuO_2(30)-UO_2-Polystyrene$ $H/(Pu + U) = 47.4$	Plexiglas	1.0635	0.0124	9
2	$PuO_2(14.62-UO_2-Polystyrene H/(Pu + U) = 30.6$	Plexiglas	1.0611	0.0126	9
3	PuO ₂ (7.89)-UO ₂ -Polystyrene H/(Pu + U) = 51.8	Plexiglas	1.0357	0.0116	9
4 ^a	PuO ₂ (7.89-UO ₂ -Polystyrene H/(Pu + U) = 51.8	Plexiglas	1.0624	0.0157	ń
5	PuO ₂ H/Pu = 0.04	Plexiglas	0.9926	0.0177	10
6	PuO ₂ -Polystyrene H/Pu = 15	Plexiglas	1.0298	0.0134	11
7	PuO ₂ -Polystyrene H/Pu = 50	Plexiglas	1.0400	0.0124	12

*Calculations by J. H. Chalmers and G. Walker.

^aThis calculation was made on the actual experimental array of compacts from data provided by S. R. Bierman. All other calculations were made on the homogeneous cuboid, as reported.

with dry metal. For nonspherical experiments, correlations were made with the KENO code,¹⁷ again with Hansen-Roach cross sections, since it is equivalent to S_{∞} . Some of the experiments contained ²³⁴U, ²³⁸Pu, ²⁴²Pu, and ²⁴¹Am, which are not in the Hansen-Roach tabulation. Cross sections for ²³⁸Pu were furnished by Smith. Fairly rough approximations were made for the remaining three nuclides, but they were considered adequate for the minor concentrations of these nuclides in the experiments.

There is a series of critical experiments with Plexiglas cuboids built from blocks of PuO_2 and UO_2 compacted with polystyrene in which the H/(Pu + U) ratio is 2.8, the concentration of plutonium in the U + Pu is 29.3%, and the plutonium contains 11.5% ²⁴⁰Pu (Ref. 18). Correlations with these experiments are given in Table V. The ANISN calculations, involving transverse bucklings appropriate for a bare system, are not very meaningful except as a guide for the fairly large and somewhat uncertain extrapolation to the infiinte slab.

There are PCTR experiments with UO_3 enriched to ~1% (Ref. 19) and with UO_3 enriched to 3.04% (Ref. 20) ²³⁵U at H/U ratios extending upward from ~3.5. No plutonium was present, but correlation with these experiments ought to give a good indication of the bias at high ratios of fissionable to fissile atoms. The results for the

experiments near 1% enrichment are reported as values of k_{∞} as a function of H/U ratio with an uncertainty in k_{∞} of about ± 0.005 . Since there may be some discrepancy in k_{∞} calculated by the four-factor formula of the report and k_{∞} calculated as the ratio of neutrons produced per neutron absorbed by B_1 method, reported cross sections and k_{∞} 's were used to determine the amount of boron required to make k_{∞} unity. The results, expressed as B/U ratios, are given in Table VI. The H/U ratios listed in Table VI differ somewhat from the reported values, which are inconsistent with reported fractional water contents. Also given in Table VI are values of k_{α} calculated for these compositions. These values were fitted by least-squares to linear expressions in H/U. For the Hansen-Roach correlation, k^{z} 1.0375 - 0.003427 H/U, and for GLASS, k =1.0240 - 0.002145 H/U. The data show too much scatter to display any departure from linearity over the limited range of H/U. The deviation of k from unity represents the bias of the calculational method. For the experiments at 3.04%, the values in Table VII are the values computed for the reported barns of 1/v absorber per uranium atom required to make k_{∞} unity, with the absorber taken to be boron. Correlations beyond an H/Vratio of 8.01 are not given, since these are not pertinent here. A similar analysis was carried out for nitrate experiments.²¹ The correlation is

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TABLE V

Correlations with PuO₂-UO₂ Compacts Reflected by 15 cm of Plexiglas

(Density 1.185 - 8% hydrogen, 60% carbon, 32% oxygen)*

Experimental Dimonsions ^a	k _{eff}						
(cm)	- S ₄	S ₈	S ₁₆	KENO			
$35.57 \times 35.63 \times 36.50$			0.9085	1.0163 ± 0.0062			
28.86 imes40.66 imes40.72			0.9396	1.0073 ± 0.0053			
22.10 imes50.82 imes50.90			0.9737	1.0023 ± 0.0053			
20.49 imes50.82 imes61.08			0.9832	1.0090 ± 0.0050			
19.01 imes 60.98 imes 61.08			0.9923	0.9954 ± 0.0061			
18.49 imes 61.08 imes 66.07			0.9949	1.0072 ± 0.0058			
18.04 imes 66.07 imes 71.26			1.0050	1.0129 ± 0.0061			
13.01 × ∞ × ∞	1.0286	1.0299	1.0301	1.0220 ± 0.0053			
$12.60 \times \infty \times \infty$	1.0203	1.0217	1.0219	1.0218 ± 0.0063			

ANISN or KENO with Hansen-Roach cross sections. The errors associated with the KENO calculations are one standard deviation and do not include reported experimental uncertainties in dimensions. Adjoint biasing was used in the Plexiglas reflector. Calculations were made with the reported atom densities.¹⁸ The experimental data for the 18.04-cm slab do not appear in Ref. 18; these were obtained in a later experiment by Bierman. He also revised his estimate of the infinite slab thickness to 13.01 cm (0.1301 m). The 12.60-cm (0.1260-m) thickness was obtained independently by Clark from an analysis of Bierman's data, and appears more consistent with the trend shown by the S_n calculations. The KENO results may indicate an even smaller thickness for the infinite slab. From Ref. 18.

TABLE VI

Correlations with UO₃-H₂O Experiments (Ref. 19)

		-	$k_{\infty calc}/k_{exptl}$			
% ²³⁵ U	H/U	B/U	HRXN	GLASS		
1.0059	3.834	-0.000116	1.0194	1.0120		
	5.067	-0.000113	1.0168	1.0114		
:	6.235	-0.000194	1.0104	1.0069		
	6.953	-0.000283	1.0072	1.0039		
	7.524	-0.000480	1.0104	1.0087		
1.0704	3.785	0.000095	1.0262	1.0174		
	5.841	0.000102	1.0195	1.0140		
	7.145	-0.000071	1.0157	1.0113		
1.1586	3.793	0.000472	1.0297	1.0189		
	5.996	0.000555	1.0215	1.0140		
	6.909	0.000539	1.0146	1.0071		
	7 520	0.000265	1 0166	1 0106		

(Ref. 20)							
% ²³⁵ U	H/U	k _{∞calc} /k _{exptl} HRXN					
3.04	3.58	1.0320 1.0302					

1.0296 1.0308 1.0270 1.0268

1.0336

1.0324

5.86

6.38

8.01

TABLE VII

Correlations with PCTR Experiments

given in	Table	VIII	and	is	fitted	by	k =	1.03	807	-
1 .0220	H/U	for	Han	sei	n-Roac	h,	and	by	k	=
	0.00070	JZ8 I.	1/U t	or	GLASS	3.				

The correlations with the PCTR experiments in Tables VI, VII, and VIII indicate that S_{∞} , Monte Carlo, or B_1 calculations with Hansen-Roach or HAMMER cross sections are conservative by ~2 to 3% in k_{eff} at low H/U ratios. Similarly, the correlations with the compacts in Table V indicate

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TABLE VIII

Correlation with Nitrate Experiments (Ref. 21)

		$k_{\infty calc}/k_{exptl}$	
% ²³⁵ U	H/U	HRXN	GLASS
2.14	6.36	1.0237	1.0211
	7.17	1.0179	1.0159
	8.46	1.0180	1.0182
	10.36	1.0216	1.0234
	10.40	1.0212	1.0231
2.26	8.25	1.0025	1.0022
	11.2	1.0001	1.0010

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	Critical Experiments with Spheres*					·
Experiment Number	Region	Composition	Density	Dimension	Reference	Experime Number
1	Core Reflector	Pu (5.18, 0.30, 0.02; a) H ₂ O	19.74 0.9982	4.122 ± 0.006 >30	22	. 10
2	Core	U(93.8)	18.75	8.732 ± 0.009	23	10, 11
3	Core	Pu (4.5, 0.3; a), 1.02% Ga	15.61	6.385 ± 0.013	23	
4	Core	Pu (20.1, 3,1, 0.4; a), 1.01% Ga	15.73	6.660 ± 0.017	23	10
5	Core Reflector	Pu (4.80, 0.30), 1.10% Ga U(Nat)	15.53 19.00	$\begin{array}{r} 4.533 \pm 0.008 \\ 19.609 \end{array}$	23	
6	Core Reflector	U(93.24) U(Nat)	18.62 19.00	6.116 ± 0.004 18.009	23	
7	Core Reflector	U(93.90) U(Nat)	18.69 19.00	6.326 ± 0.011 9.982	23	
8	Core Reflector	U(93.99) U(Nat)	18.67 18.67	$\begin{array}{c} 6.977 \pm 0.011 \\ 4.425 \end{array}$	23	15
9	Core Reflector	U(93.91) U(Nat)	18.70 19.00	7.755 ± 0.013 1.735	23	•First figu

TABLE IX

*Plutonium composition is % ²⁴⁰Pu, ²⁴¹Pu, ²⁴²Pu; the remainder is ²³⁹Pu. All percentages are by weight except where "a" denotes at %. Uranium composition is % ²³⁵U; except for 1.02% ²³⁴U, remainder is ²³⁸U. Natural uranium con-tains 0.71% ²³⁵U; the remainder is ²³⁸U. The dimensions are in centimetres and are the core radius and the reflector thickness. In many cases, they were derived from reported masses and densities. Densities are in g/cm³.

TABLE X

Correlations with Critical Spheres*

Euronimont		keff Cal	lculated for Exp	eriment	
Number	S4	S ₈	S 16	S ∞	KENO
1	1.0159 ± 0.0012	1.0008	0.9968	0.9954	0.9845 ± 0.0056
2	1.0103 ± 0.0008	1.0034	1.0013	1.0009	1.0094 ± 0.0053
3	1.0159 ± 0.0017	1.0060	1.0030	1.0019	0.9973 ± 0.0053
4	1.0222 ± 0.0022	1.0124	1.0095	1.0084	0.9989 ± 0.0055
5	1.0153 ± 0.0015	1.0018	0.9983	0.9971	1.0018 ± 0.0052
6	1.0108 ± 0.0005	1.0011	0.9986	0.9978	0.9933 ± 0.0043
7	1.0139 ± 0.0013	1.0040	1.0014	1.0005	1.0017 ± 0.0036
8	1.0122 ± 0.0012	1.0029	1.0003	0.9994	0.9990 ± 0.0049
9	1.0124 ± 0.0013	1.0042	1.0018	1.0014	1.0017 ± 0.0044

*The errors associated with the S4 calculations correspond to the probable errors in the experimental radii and 0 would presumably be about the same for S_{∞} .

conservatism in calculations at low H/(U + Pu). The maximum ratio to be considered for damp oxide, however, is only ~ 0.5 , which is much less than the minimum ratio in these experiments. Critical experiment data available for dry, metal spheres are listed in Table IX. For these systems, the correlations in Table X indicate good agreement between calculation and experiment and

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no apparent dependence on whether the fissile material is ²³⁵U or ²³⁹Pu.

Experimental data for bare uranium cylinders are listed in Table XI. The correlations (Table XII) appear to indicate an increasing degree of nonconservatism as the concentration of fissile material in uranium decreases.

On the basis of these various correlations,

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TABLE XI

Critical Experiments with Bare Cylinders*

Experiment Number	Composition	Density	Radius	Height	Refer- ence
10	Ave: U(0.58, 53.33) 11 pairs: 0.6076 U(Nat), 0.7978 U(93.41)	18.83	13.335	15.459 ± 0.077	23 23
¹⁴ ¹¹	Ave: U(0.40, 37.46) 13 triplets: 0.5900 U(Nat), 0.7751 U(93.43), 0.5900 U(Nat)	18.88	13.335	25.416 ± 0.127	23
12	Ave: U(0.17, 16.01) 19 pairs: 1.5262 U(Nat), 0.3021 U(93.3)	18.68	26.67	34.4 ± 0.31	23,20
13	Ave: U(0.15, 14.11) 21 pairs: 1.8098 U(Nat), 0.3062 U(93.3)	18.41	26.67	44.44 ± 0.31	23, 24
14	Ave: U(0.13, 12.32) 25 triplets: 0.6076 U(Nat), 0.3052 U(93.3) 1 5216 U(Nat)	18.64	26.67	60.86 ± 0.30	23, 24
15	Ave: U(0.11, 10.9) 22 quadruplets: 1 8041 U(Nat), 0.2975 U(93.3) 3.0077 U(Nat), 0.2975 U(93.3)	18.63	26.67	118 95 ± 3.0	23, 24

*First figure in average uranium composition is % ²³⁴U, next is ²³⁵U, remainder is ²³⁸U. Dimensions are in centimetres (×10⁻² = m). Heights were derived from reported masses, diameters, and densities, which the experimenters adjusted to compensate for warpage of plates. The dimensions (cm) (×10⁻² = m) of the stacked units were adjusted slightly from the experimental values to give an integral number of groups at critical. The average density was assumed for each component of a group.

TABLE XII

Correlations with Bare Cylinders*

	k _{eff} Calculated for Experiment				
			KENO		
Number	S4	S ₈	Homogenized	Heterogeneous	
10	0.9854 ± 0.0019	0.9840	0.9897 ± 0.0040	1.0057 ± 0.0044	
11	0.9850 ± 0.0013	0.9831	0.9864 ± 0.0045	1.0015 ± 0.0039	
12	0.9805 ± 0.0026	0.9800	0.9798 ± 0.0039	0.9840 ± 0.0038	
13	0.9777 ± 0.0015	0.9771	0.9765 ± 0.0030	$0\ 9823\pm 0.0036$	
14	0.9781 ± 0.0007	0.9774	0.9845 ± 0.0031	0.9794 ± 0.0046	
15	0.9769 ± 0.0010	0.9762	$0\ 9745 \pm 0.0028$	0.9753 ± 0.0037	

*The S_n calculations were made for the homogenized cylinders. The errors associated with the S_4 calculations correspond to the probable errors in the experimental heights. The experimental heights were used directly in the S_n calculations. The KENO calculations for the cylinders having the average compositions were for single cylindrical units. Enclosing the first three in void cuboids gave, respectively, 0.9894, 0.9854, and 0.9833. The KENO calculations for the stacks of alternating Oralloy and natural uranium plates were made for multiple units. Treating the first three as single units with all interfaces specified gave, respectively, 0.9949, 0.9890, and 0.9899.

Clark concluded that concentrations of ²³⁹Pu in $U(Nat) + {}^{239}Pu$ in dry and damp homogeneous mixtures of oxides for which B_1 calculations with Hansen-Roach cross sections give 0.95 for k_{∞} will indeed by subcritical regardless of the mass

of material; hence these concentrations may be chosen as subcritical limits (first two limits of Table III). In view of the paucity of data and of the downward trend in $k_{\rm eff}$ as a function of decreasing uranium enrichment in bare cylinders, a larger $\{k_i\}_{i=1}^{n}$

e (b

	Dry-Theoretical Density	Damp-Theoretical Density	Damp $-\frac{1}{2}$ Theoretical Density			
3 wt% PuO ₂						
D. W. Magnuson ^a	$k_{\infty} = 0.992$	688 kg Pu ($k_{eff} \approx 0.95$) (26 061 kg MO)	2389 kg Pu (<i>k</i> eff = 0.95) (90 492 kg MO)			
H. K. Clark ^b		236 kg Pu ($k_{eff} = 0.95$) (8939 kg MO)	885 kg Pu ($k_{eff} = 0.95$) (33 523 kg MO)			
		8 wt% PuO2				
D. W. Magnuson ^a	167 kg Pu ($k_{eff} = 0.95$) 312.5 kg Pu ($k_{eff} = 1.00$)	56 kg Pu $(k_{eff} = 0.95)$ 83.3 kg Pu $(k_{eff} = 1.00)$	290.9 kg Pu ($k_{\rm eff} = 1.00$)			
H. K. Clark ^b	122 kg Pu ($k_{eff} = 0.95$) 184 kg Pu ($k_{eff} = 1.00$)	49.4 kg Pu ($k_{eff} = 0.95$) 69.4 kg Pu ($k_{eff} = 1.00$)	161 kg Pu (k _{eff} = 0.95) 239.8 kg Pu (k _{eff} = 1.00)			
L. C. Davenport ^c	160.6 kg Pu ($k_{eff} = 0.95$) 265.6 kg Pu ($k_{eff} = 1.00$)	57.8 kg Pu $(k_{eff} = 0.95)$ 88.9 kg Pu $(k_{eff} = 1.00)$	203.8 kg Pu ($k_{eff} = 0.95$) 323.9 kg Pu ($k_{eff} = 1.00$)			
J. H. Chalmers ^d	337.5 kg Pu (k _{eff} = 1.00)		160.9 kg Pu ($k_{eff} = 0.95$) 268.0 kg Pu ($k_{eff} = 1.00$)			
	······	15 wt% PuO ₂				
D. W. Magnuson ^a	46 kg Pu ($k_{eff} = 0.95$) 62 kg Pu ($k_{eff} = 1.00$)	32 kg Pu $(k_{eff} = 0.95)$ 43.7 kg Pu $(k_{eff} = 1.00)$	103 kg Pu (k _{eff} = 0.95) 142.5 kg Pu (k _{eff} = 1.00)			
H. K. Clark ^b	47 kg Pu ($k_{eff} = 0.96$) 56 kg Pu ($k_{eff} = 1.00$)	32.9 kg Pu ($k_{eff} = 0.96$) 39.4 kg Pu ($k_{eff} = 1.00$)	102 kg Pu ($k_{\rm eff}$ = 0.96) 126.9 kg Pu ($k_{\rm eff}$ = 1.00)			
L. C. Davenport ^c	50.4 kg Pu ($k_{eff} = 0.95$) 66.7 kg Pu ($k_{eff} = 1 00$)	$34.4 \text{ kg Pu} (k_{eff} = 0.95)$ 46.0 kg Pu $ (k_{eff} = 1.00)$	113.4 kg Pu ($k_{eff} = 0.95$) 157.0 kg Pu ($k_{eff} = 1.00$)			
J. H. Chalmers ^d			143.0 kg Pu ($k_{\rm eff} = 1.00$)			
S. R. Bierman	$62.2 \text{ kg Pu} \ (k_{\text{eff}} = 1.00)$	48.4 kg Pu ¹ ($k_{eff} \approx 1.00$)	~			
G. Walker	$64.2 \text{ kg Pu} \ (k_{eff} = 1.00)$		141.0 kg Pu ($k_{eff} = 1.00$)			
		30 wt% PuO ₂				
D. W. Magnuson ^a	25 kg Pu ($k_{eff} = 0.95$) 30.2 kg Pu ($k_{eff} = 1.00$)	23 kg Pu $(k_{eff} = 0.95)$ 27.5 kg Pu $(k_{eff} = 1.00)$	63 kg Pu ($k_{\rm eff}$ = 0.95) 83.7 kg Pu ($k_{\rm eff}$ = 1.00)			
H. K. Clark ^b	26.1 kg Pu ($k_{eff} = 0.97$) 28.4 kg Pu ($k_{eff} = 1.00$)	23.3 kg Pu $(k_{eff} = 0.97)$ 25.5 kg Pu $(k_{eff} = 1.00)$	67.9 kg Pu ($k_{\rm eff}$ = 0.97) 76.3 kg Pu ($k_{\rm eff}$ = 1.00)			
L. C. Davenport ^c	25.4 kg Pu ($k_{eff} = 0.95$) 31.6 kg Pu ($k_{eff} = 1.00$)	23.0 kg Pu ($k_{eff} = 0.95$) 28.9 kg Pu ($k_{eff} = 1.00$)	70.4 kg Pu ($k_{eff} = 0.95$) 91.7 kg Pu ($k_{eff} = 1.00$)			
J. H. Chalmers ^d	23.5 kg Pu ($k_{eff} = 0.95$) 28.3 kg Pu ($k_{eff} = 1.00$)	21.9 kg Pu ($k_{eff} = 0.95$) 27.8 kg Pu ($k_{eff} = 1.00$)	64.2 kg Pu (k _{eff} = 0.95) 86.9 kg Pu (k _{eff} = 1.00)			
S. R. Bierman ^e	32.2 kg Pu ($k_{\rm eff} = 1.00$)	28.9 kg Pu^{f} ($k_{eff} = 1.00$)				

TABLE XIII

Summary of Calculations on Mixed Oxides-Dry and Damp* Powders

*Damp powders contain ~1.5 wt% water, H/(Pu + U) $\simeq 0.45$.

^aComputed with ENDF/B-III cross-section data and 123-group XSDRN transport code; the radii for $k_{eff} = 0.95$ were estimated as a function of k_{eff} , which was iterated during the calculational search to $k_{eff} = 1.0$.

^bComputed with ANISN, a one-dimensional discrete ordinates transport code with anisotropic scattering, utilizing Hansen-Roach cross sections with ²³⁸U resonance cross sections modified by J. R. Knight.

^cComputed with DTF-IV transport code and ENDF/B-III cross sections; ENDF/B cross sections processed by ETOG and FLANGE codes for input to GAMTEC-II code, 18 groups used in DTF-IV.

^dComputed with MONK-4 Monte Carlo code and British cross sections.

^eComputed with DTF-IV transport code and ENDF/B-III cross sections; ENDF/B cross sections processed by ETOG and FLANGE codes and averaged over 17 epithermal groups and 1 thermal group (0 to 0.683 eV) by the EGGNIT code.

^fFrom interpolation.

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TABLE XIV

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Calculated Limiting Critical Enrichment for Uranium and Limits for ²³⁹Fu Content in Urani

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Limiting Critical Enrichment Critical Pl C	
	d
Experimental Calculated Calculated	
UO ₃ -H ₂ O 1.034 ^a 1.010	
$UO_2(NO_3)_2-H_2O$ 2.104 ^b 1.970	[
$U(0.71)O_2^{-239}PuO_2^{-H_2O}$ 0.18	3
$U(0.71)O_2(NO_3)_2 - {}^{239}Pu(NO_3)_4 - H_2O \qquad 0.678 \qquad 0.81$	2

Reference 19.

BReference 21.

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TABLE XV

Calculated Values of k_{∞} for Low Enriched UO₃ and Nitrate Aqueous Mixtures

Mixture	Enrichment (wt% ²³⁵ U)	k∞ from Experimental Data	Computed k_{∞}
UO3-H2O	$\begin{array}{l} 1.006 \ (\text{H/U} = 5) \\ (\text{H/U} = 4) \\ 1.034 \ (\text{H/U} = 5) \end{array}$	$\begin{array}{c} 0.99 \pm 0.01 \\ 0.99 \pm 0.01 \\ 1.00 \end{array}$	0.97 0.97 0.98
$UO_2(NO_3)_2-H_2O$	2.104	1.00	0.989

value was not considered justified. Clark similarly concluded that mass limits for oxides mixtures in which the concentration of plutonium in (U + Pu) is 3 and 8% (Table II) should also be masses for which $k_{\rm eff}$ is calculated by S_{∞} to be 0.95. For concentrations of 15 and 30%, he judged $k_{\rm eff}$'s of 0.96 and 0.97 sufficiently far below the correlating values to provide adequate assurance of subcriticality. The limits in Table II are those calculated by Clark, since his values are the smallest. As many as six "independent" calculations were made by members of the Work Group or their fellow workers in deriving these limits. The results of the various calculations are summarized in Table XIII.

An important nuclear criticality safety parameter for the processing of mixed-oxide fuel is the amount of plutonium which can be added to an aqueous mixture, with and without nitrate, before criticality is achieved (last two limits of Table III). Since there have been no experiments performed to determine these concentrations, limits were deduced from calculations validated ^{against} the experimental limiting ²³⁵U enrichments for criticality as determined from measurements in the Physical Constants Test Reactor at Hanford. Several analyses were made utilizing different computer codes and cross-section sets. Clark's correlations have already been described. Another of us (Magnuson, ORNL) did calculations using Hansen-Roach cross sections in the ANISN transport code,¹³ but also included some calculations with the XSDRN transport code, 25 and 123-group cross sections. 26 His results are given in Table XIV.

The resultant biases in the calculations are 0.024 and 0.134, respectively, for the oxide and nitrate mixtures.

Similar type calculations were independently made by Durst (Battelle-Northwest Laboratories) utilizing ENDF/B-III cross sections in the GAMTEC-II code.²⁷ The 17 epithermal energy groups were obtained via application of the ETOG $code^{28}$ and the thermal group data via application of the FLANGE code.²⁹ The results are given in Table XV.

The bias in the computed k_{∞} is $\sim 2\%$ in the case of UO_3 and 1% for the nitrate. Correcting for the bias, the critical plutonium content in $PuO_2 + UO_2$ corresponding to a k_{∞} of unity would be 0.17 wt% plutonium in the Pu + U. The plutonium content for a $k_{\infty} = 0.98$ would be 0.136. For nitrate $\begin{bmatrix} 2^{239} Pu(NO_3)_4 - UO_2(NO_3)_2 \end{bmatrix}$ water mixtures, the corresponding plutonium content would be 0.77 for a k_{∞} of unity and 0.70 for a $k_{\infty} = 0.98$.

Clark concluded from his correlations with PCTR experiments that a 2% margin in k was sufficient to compensate for uncertainties and to ensure subcriticality. His limiting values for 239 Pu in U + Pu as oxides were 0.134 and 0.142 as calculated from Hansen-Roach cross sections and by GLASS, respectively. The corresponding limits for nitrates were 0.654 and 0.708.

The computed concentrations of ²³⁹Pu in (Pu +

Limiting Subcritical wt% as Oxides in V	of ²³⁹ Pu in Water	n Pu + U
For k_{∞} of 0	.98	
Clark		Durst
0.134 (H-R) ^a 0.142 (GLASS)		0.136
For k_{∞} of 1	.00	
Clark	Durst	Magnuson
0.166 (H-R) 0.176 (GLASS)	0.169	0.183

TABLE XVI

Limiting Subcritical wt% of ²³⁹Pa = Pi - Ja Presence of Four Nitrate Ions per Planner [Pu(NO₃)₄], and Two Nitrate Ions per Uranium Atom [UO₂(NC₂ =]

For R_{∞} of	0.98	
Clark	Turne	
0.654 (H-R) 0.708 (GLAS	1	
For k_{∞} of	1.00	
Clark	Durst	Winner A
0.772 (H-R) 0.771 (GLASS)	0.77	¥_#7

^aHansen-Roach.

TABLE	XVII
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			Minimum Critical					1.11	
wt% Pu	wt% ²⁴⁰ Pu	wt% ²⁴¹ Pu	Mass g Pu (×10 ⁻³ = kg)	Cylinder Diam, cm $(\times 10^{-2} = m)$	Slab Thickness, cm $(\times 10^{-2} = m)$	Volume, $(\times 10^{-6} = m^3)$	Areal Density, g Pu/cm ² (×10 = kg/m ²)	Concentration g Pi (×10 ⁻¹ = sig	
3	0	0	969 965	$\begin{array}{c} 26.4 \\ 26.5 \end{array}$	12.4 12.7	$30.4\\30.8$	0.308	7.22 7.25	*
	15	6	1875 1738	$\begin{array}{c} 33.9\\ 33.0\end{array}$	$\begin{array}{c} 17.2 \\ 16.9 \end{array}$	$\begin{array}{c} 60.3 \\ 56.5 \end{array}$	0.433	8.71 9.12	*
	25	15	2778 2563	$\begin{array}{c} 38.5\\ 37.4\end{array}$	20.1 19.6	85.7 79.3	0.533	10.11 10. <u>43</u>	# #
8	0	0	701 709	$\begin{array}{c} 20.8\\ 21.3 \end{array}$	8.9 9.4	$\begin{array}{c} 16.1 \\ 17.3 \end{array}$	0.285	7.15 7.3A	* **
	15	6	1275 1197	$\begin{array}{c} 26.5\\ 26.4\end{array}$	12.4 12.6	$\begin{array}{c} 30.9\\ 30.8 \end{array}$	0.393	9.27 9.11	.
	25	15	$\begin{array}{c} 1800 \\ 1657 \end{array}$	$\begin{array}{c} 29.5\\ 29.1 \end{array}$	14.314.4	$\begin{array}{c} 41.6\\ 40.4\end{array}$	0.476	10.31 10.45	**
15	0	0	616 629	18.6 19.3	$\begin{array}{c} 7.5 \\ 8.1 \end{array}$	$\begin{array}{c} 12.1\\ 13.3\end{array}$	0.276	7.3 <u>7</u> 7.5 <u>4</u>	* *
	15	6	1099 1048	$\begin{array}{c} 23.8\\ 24.0\end{array}$	$10.7\\11.1$	$\begin{array}{c} 23.3\\ 23.9 \end{array}$	0.378	9.14 9.14	*
	25	15	$\begin{array}{c} 1505\\ 1434 \end{array}$	$\begin{array}{c} 26.4\\ 26.4\end{array}$	$\begin{array}{c} 12.3\\ 12.6\end{array}$	$\begin{array}{c} 30.8\\ 30.7\end{array}$	0.453	15.29 10.39	1
30	0	0	561 552 609 ^c	16.9 17.2 17.2 ^c	6.4 6.8 6.6 ^c	9.5 9.9	0.268	7.32	ی بر اند بر اند
	15	6	975 961 926 ^c	21.9 22.3 21.0 ^c	9.5 10.0	18.7 19.6	0.364	9.1 9.1	
	25	15	1320 1285 1233 ^c	$24.1 \\ 24.5 \\ 23.0^{\circ}$	$10.8 \\ 11.4$	24.0 25.2	0.436	10.22 10.30	

Calculated Minimum Critical Conditions for Water-Reflected PuO2-UO2-H2O Mixtures*

*Calculations in this table by Clark, except as noted.

^aMGBS-TGAN normalized to plutonium solution experiments.

^bHansen-Roach cross sections, S_{\pm} . ^cCalculations by Chalmers.

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	Material	Composition $(\times 10^{-3} = \text{kg}/\ell)$	Geometry	Reflection	$\begin{array}{c} \text{Calculated} \\ k_{\text{eff}} \end{array}$
	U(5)O ₂ F ₂ -H ₂ O	915 g U/l	25.39-cm sphere	None	1.016
2	Pu(95.4)(NO3)4-H2O	58 g Pu/l	8.455-cm half-slab	None	1.025
貅	Pu(71.9)(NO ₃) ₄ -H ₂ O	202 g Pu/l	10.595-cm half-slab	None	1.013
	Pu(95.4)(NO3)4-H2O	268.7 g Pu/l	14.57-cm sphere	0.124-cm stainless steel Inf. H ₂ O	1.009
	Pu(95)(NO3)	35.1 g Pu/l 110.8 g U/l	19.30-cm sphere	0.122-cm stainless steel Inf. H2O	1.014
N. C.	$Pu(91.4)O_2-UO_2-(C_8H_8)_m$	85.0 g Pu/l 495 g U/l	5.78-cm half-slab	15-cm Plexiglas	1.029
				Ave	rage 1.018

TABLE XVIII

Calculated keff for Selected Critical Experiments Using the XSDRN-S8 Transport Code*

*Calculations by Magnuson.

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U) that provide k_{∞} = unity and 0.98, adjusting for calculational bias, are summarized in Table XVI. The computational methods selected by Clark for homogeneous aqueous mixtures (Table I limits) were S4 with Hansen-Roach cross sections (as already described) and MGBS-TGAN (Ref. 30). The latter combination of codes was applied to the three sets of mixed oxide experiments' considered by Chalmers and Walker. Extrapolations of the data to infinite slabs were made, and values of k_{eff} were calculated for the slabs. These values were ~ 0.01 larger than values similarly obtained^{6,30} for aqueous solutions of plutonium nitrate at the same $H/^{239}$ Pu ratio. Clark concluded that the bias established for nitrate solutions is appropriate for aqueous mixed-oxide mixtures. Little bias exists in the less extensive correlations of S_4 and Hansen-Roach cross sections with the solution experiments,⁶ but there appears to be a trend toward larger values of the correlating k_{eff} with larger H/239Pu ratios. Both methods were used to compute minimum critical masses, dimensions, and concentrations for PuO₂-UO₂-H₂O mixtures. The bias characteristics of plutonium nitrate solutions were used in the MGBS-TGAN calculations. No bias was assumed in the S_4 calculations. However, for the infinite sea concentrations calculated by MGBS and by B_1 from Hansen-Roach cross sections, bias was assumed for both methods and was obtained by extrapolation of trends outside the range of experimental data. Results are given in Table XVII along with a few results obtained by Chalmers. Most of the differences between MGBS-TGAN and ANISN results correspond to a difference of <1% in k_{eff} . The maxi-

mum difference is $\sim 2\%$ and occurs mainly for the slabs.

The limiting concentrations for homogeneous mixtures of plutonium and natural uranium, i.e., the "infinite sea" concentrations, were also calculated by Magnuson utilizing ENDF/B-III cross-section data and the XSDRN transport theory code.³¹ Values of k_{eff} were calculated for a number of critical experiments to provide an estimate of the bias in the calculational method. The results of these calculations and the experimental conditions are given in Table XVIII.

It is seen that there are biases that are conservative for nuclear criticality safety, i.e., critical systems are calculated to be supercritical, and calculated critical systems would then be subcritical. The biases are in general the same order of magnitude as those previously found with either ANISN and Hansen-Roach 16-group cross sections or with XSDRN and an earlier ENDF/B cross-section set.

The compositions and the k_{∞} values calculated with the transport code XSDRN are given in Table XIX. The values of the concentrations for various k_{∞} values in Table XX were obtained from Table XIX by linear interpolation and extrapolation. It is noted for the mixtures having Pu/(Pu + U) ratios of 1.0, 0.30, 0.15, and 0.08 that the addition of natural uranium decreases k_{∞} or increases the plutonium concentration for criticality ($k_{\infty} = 1$). For Pu/(Pu + U) = 0.03, a reversal in the trend is noted, and the k_{∞} values are higher than those for the 0.08 Pu/(Pu + U) weight ratio for the same plutonium concentrations.

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Weight Fraction, Pu/(Pu + U)	1.0	0.30	0.15	0.08	0.03
Pu (g/ℓ) $(\times 10^{-3} \text{ kg}/\ell)$ Calculated k_{∞} Values (XSDRN-S ₂)					
	100 wt% ²³⁹ Pu				
6.5 7.0	$0.9474 \\ 0.9855$	0.9441 0.9811	0.9421 0.9783	$0.9414 \\ 0.9766$	$0.9466 \\ 0.9809$
79 wt% ²³⁹ Pu—15 wt% ²⁴⁰ Pu—6 wt% ²⁴¹ Pu					
8.0 8.5	0.9480 0.9768	0.9433 0.9710	$\begin{array}{c} 0.9411 \\ 0.9682 \end{array}$	0.9402 0.9664	0.9472 0.9714
$\frac{1}{60 \text{ wt}\%^{239} \text{Pu}-25 \text{ wt}\%^{240} \text{Pu}-15 \text{ wt}\%^{241} \text{Pu}}$					
9.0 10.0	0.9407 0.9875	0,9354 0,9802	0.9339 0.9766	$0.9321 \\ 0.9742$	0.9397 0.9777

TABLE XIX

XSDRN Calculations of k_{∞} for PuO₂-UO₂-H₂O Mixtures

CONCLUSION

The work leading to the development of a proposed standard involves the efforts, contributions, and views of many persons. In addition to the derivation of subcritical limits, the validations undertaken in this regard have provided heretofore unpublished data on correlations against experimental data that may be used to assess the validity of the codes used, and cross sections, in certain applications.

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Plutonium Concentrations for Several Calculated k_{∞} Values

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k_{∞} 1.0		0.97	0.95			
Weight Fraction, Pu/(Pu + U)	Plutonium Concentrations (g/ℓ) (×10 ⁻³ kg/ ℓ), with H/Pu Atom Ratios in Parentheses					
	100 wt%	²³⁹ Pu				
1.0 0.30 0.15 0.08 0.03	7.19 (3676) 7.26 (3625) 7.30 (3594) 7.33 (3561) 7.28 (3533)	6.80 (3892) 6.85 (3863) 6.89 (3831) 6.91 (3805) 6.84 (3789)	6.47 (4095) 6.58 (4020) 6.61 (3996) 6.62 (3974) 6.55 (3958)			
$79 \text{ wt}\% ^{239}\text{Pu}-15 \text{ wt}\% ^{240}\text{Pu}-6 \text{ wt}\% ^{241}\text{Pu}$						
1.0 8.90 (2965) 0.30 9.02 (2905) 0.15 9.09 (2875) 0.08 9.14 (2837) 0.03 9.09 (2795)		3.38 (3169) 8.43 (3140) 8.43 (3131) 8.57 (3061) 8.47 (3042)	8.03 (3306) 8.12 (3262) 8.12 (3253) 8.19 (3210) 8.06 (3203)			
60 wt% ²³⁹ Pu-25 wt% ²⁴⁰ Pu-15 wt% ²⁴¹ Pu						
$ \begin{array}{c} 1.0\\ 0.30\\ 0.15\\ 0.08\\ 0.03 \end{array} $	10.27 (2576) 10.44 (2517) 10.54 (2479) 10.61 (2442) 10.59 (2391)	9.63 (2765) 9.77 (2715) 9.85 (2682) 9.90 (2651) 9.80 (2614)	9.20 (2892) 9.32 (2848) 9.39 (2818) 9.42 (2793) 9.27 (2780)			

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