ACCIDENTAL RADIATION EXCURSION AT THE OAK RIDGE Y-12 PLANT—III
DETERMINATION OF RADIATION DOSES

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I. INTRODUCTION
An analysis of the physical events leading to an accidental radiation excursion at the Oak Ridge Y-12 Plant has been presented in the first of this series of articles.(1) A companion article(2) is devoted to health physics problems arising from this accident. The present article describes the methods which were used to arrive at the doses received by the individuals. Although our main objective is to describe the dose analysis of this particular excursion, efforts will be made to present methods and information which may be of use in similar problems.

Very early estimates of the doses received by the individuals, based on their known positions at the initiation of the excursion and an estimate of the total number of fissions in the assembly, were unreasonably high. For example, the fast neutron component of the total dose received by employee A was estimated to have been 1500 rads. Hence one must assume that the location of personnel during a substantial fraction of their exposure time is unknown. This fact, together with the fact that the exposed employees were not wearing any type of personnel monitoring equipment, dictates that the evaluation of dose be based on the activation of body elements. Experience at Los Alamos(3) in the Pajarito accident indicates that of the body fluid methods, Na23 in blood may be the most reproducible criterion of neutron exposure and, in particular, indicates that the formation of P32 in urine is poorly understood.

In spite of the fact that the exposed individuals were wearing no dose-measuring devices,† it is believed that the final dose values are reasonably accurate (+-20 per cent). The doses were arrived at by the following procedure: (1) Blood and urine samples were collected from the exposed employees; these samples were counted for radioactive substances‡ (Na24 in the blood and P32 and Na24 in the urine) as described in Section V. (2) A burro was exposed to a mock-up of the liquid assembly to determine the relationship of Na24 activity in the blood and neutron dose. The ratio of y-dose, $D_y$, to neutron dose, $D_n$, for this particular type of critical assembly was also determined with the mock-up experiment (see Section IV). (3) A parallel program calculated these same two quantities. The calculations, described in Sections II and III, proved to be extremely helpful since they provided a basis for checking the experimental procedure at various stages.

II. SODIUM ACTIVATION OF HUMAN BLOOD

The $(n,\gamma)$ process in Na23 gives rise to Na24 which has a half-life of 14.8 hr and emits a 1.38 MeV $\gamma$-ray in cascade with a 2.76 MeV

† The personnel security badge did contain strips of indium which proved invaluable in separating the exposed from the non-exposed persons.
‡ Because of complex physiological factors involved in the interpretation of P32 activity in urine, this work will deal almost entirely with the interpretation of Na24 activity in blood. Both P32 and Na24 in urine were analyzed by L. B. FARABEE, ORNL. In the case of Na24 in urine, it appears possible to correlate activity with neutron dose, particularly if the first void after exposure is not used and if samples are collected before new sodium is given to the exposed individual. In the case of P32, the difficulties reported in Ref. 3 were encountered.
γ-ray per Na$^{24}$ decay. One expects that the cross-section for this process will follow a 1/ν behavior from the thermal region to about 300 eV. The capture contribution of the resonance at 2.94 keV has been estimated by GOLDSTEIN$^{(4)}$. He finds a very narrow resonance, with a maximum cross-section at resonance of only 1 barn. It is easy to show that the number of captures occurring in this resonance is negligible compared with those occurring at thermal energies for slowing-down spectra typical of hydrogenous media. GOLDSTEIN notes that some uncertainty exists in his estimate of the resonance cross-section and that further experimental work will be necessary to resolve the uncertainty. However, it is expected that the neglect of epithermal activation of Na$^{23}$ is not serious and this approximation will be made throughout this paper.

The human body is several mean free paths thick for fast neutrons. Consequently, the probability that a fast neutron will be captured as a thermal neutron is not very sensitive to its initial energy. Thus the body is roughly equivalent to the "long counter"$^{(5)}$ used so much in neutron flux measurements. The activation of elements in the blood should be a valid measure of the average thermal flux throughout the body and hence of the total flux incident at all energies.

This assumption is basic to the procedure described below for calculating the dose to those exposed in the nuclear excursion. The Na$^{24}$ activation (μc/cm$^2$) of blood in the human body will be calculated under the assumption that the body is equivalent to a cylindrical phantom of 15 cm radius composed of soft tissue.

Calculations have been made by SNYDER and NEUFELD$^{(6)}$ of the depth distribution of thermal neutrons in an infinite slab of tissue-equivalent material 30 cm thick due to a monoenergetic broad beam irradiation by normally incident fast neutrons. The thermal neutron flux distribution is proportional to the curves labelled $T_\nu$ in Figs. 1 through 11 of SNYDER and NEUFELD$^{(6)}$. From these curves one may obtain the fraction of those neutrons entering the phantom which are captured at thermal energy. This fraction is plotted in Fig. 1 as a function of the incident neutron energy. Also plotted are values for the fraction $\xi(E)$ captured in a right cylinder of 15 cm radius, obtained from the slab data by making approximate corrections for thermal and fast leakage through the surface of the cylinder. The fast leakage was calculated by simply averaging over the path length distribution in a cylinder irradiated normal to its axis, taking the attenuation kernels for plane slab penetration from the data given by SNYDER and NEUFELD$^{(6)}$. The thermal leakage was determined by applying one velocity diffusion theory to those neutrons which have entered the thermal region while in the cylinder. More accurate calculations of the capture fraction for a cylindrical phantom are

![Fig. 1. Thermal neutron capture probability in a phantom.](image-url)
G. S. HURST, R. H. RITCHIE and L. C. EMERSON 123

Irradiation with neutrons striking a cylindrical phantom results in the absorption of neutrons in tissue, and neutrons and γ-rays in the neighborhood of the reactor are independent of distance from the critical assembly. This is thought to be a reasonable assumption for the range of distances involved. Then one may write that the flux $\phi(E)$ at a distance $R$ from the assembly is given by $\phi(E,R) = \frac{N(E)}{4\pi R^2}$, where $N(E)$ is the number of neutrons with energy $E$ per unit energy interval leaking from the assembly.

Clearly, the replacement of the body by a cylindrical phantom for the purpose of calculation is accompanied by some uncertainties. One must assume (a) that the blood is uniformly distributed throughout the body and hence takes on a sensible average activation and (b) that the geometrical differences between the human body and the cylindrical phantom are negligible with respect to the thermal capture of incident fast neutrons. It is believed that the approximations involved will not lead to serious error in the result.

The $\text{Na}^{24}$ method of determining the fast neutron dose is most useful in cases where the number of thermal neutrons is not large compared with the number of fast neutrons. If the number of thermais comparatively large, the bulk of the activation will be due to neutrons which do relatively little damage to the body and the accuracy of the method will suffer. However, if the spectral distribution of neutrons is well known, one may still derive an accurate dose value. In most cases of interest to the health physicist the thermal flux component will be small enough that the $\text{Na}^{24}$ activation may
be related to the fast neutron dose with good accuracy.

III. CALCULATION OF $D_\gamma/D_n$ FOR LIQUID ASSEMBLIES

Following the accident it seemed desirable to estimate the leakage spectra of the neutrons and the $\gamma$-rays. The overall accuracy of the methods employed in the case of the fast neutron leakage is not easy to assess in view of the rather complicated geometry of the assembly and surrounding materials but the calculated spectrum seems to agree reasonably well with the measured spectrum. A more accurate calculation is planned, in which stochastic methods will be employed.

The neutron and gamma ray leakage spectra were calculated under the assumption that the assembly could be considered spherical and the distribution of nascent fission neutrons in the assembly could be adequately represented by the fundamental spatial mode for a spherical reactor, i.e. proportional to $1/r \sin \pi r/\rho_0$, where $r$ is the distance from the center of the assembly and $\rho_0$ is the radius of the equivalent sphere at the prompt critical stage ($24.25$ cm). The number of neutrons and $\gamma$-rays escaping the assembly was calculated by using infinite medium penetration data for point isotropic sources and by integrating these data over the volume source distribution assumed. This approximate procedure of employing infinite medium attenuation data is dictated by the lack of information relative to the bounded medium. The error involved in this procedure is believed to be small.

The details of calculation of the neutron escape spectrum will now be described. The escape of high energy neutrons ($E > 0.5$ MeV) from the assembly was carried out using infinite medium penetration data for fission neutrons in water. One fits $\psi(E,r)$, the flux of neutrons of energy $E$ at a distance $r$ in water from a unit point source of fission neutrons, by an expression of the form,

$$\psi(E,r) = \frac{1}{4\pi^2} \sum_i A_i(E) \exp(-r\mu_i(E))$$

where the constants $A_i(E)$ and $\mu_i(E)$ are determined from an empirical fit to the data.

In all cases considered, only a few terms in the series were needed to obtain a good fit to the data. Then, one needs to calculate the leakage fraction through the surface at $r_0$ for a general attenuation kernel proportional to $e^{-\mu r}/4\pi r^2$ in order to determine the leakage fraction for the normalized distribution $\mathcal{S}(r)$ inside the sphere. The leakage fraction $\mathcal{L}$ through the surface is given by

$$\mathcal{L} = 4\pi r_0^2 \int d\mathbf{v} \frac{e^{-\mu p}}{4\pi p^2} \cos \varphi \mathcal{S}(r)$$

where $\mathbf{p}$ is a vector to the volume element $d\mathbf{v}$ from the point $P$ on the surface and $\cos \varphi$ is the angle between $\mathbf{p}$ and the inward normal to the surface at $P$ (see Fig. 2). The integration is carried out over the interior of the sphere. Integrating over the volume by first varying $\mathbf{p}$ while holding $r$ constant and then finally integrating over $r$, one finds

$$\mathcal{L}(\mu r_0) = \frac{1}{4r_0^3} \int_0^{r_0} \mathcal{S}(r) \left[\mu r_0 + r \right]$$

$$\times E_2[\mu(r_0 - r)] - [r_0 - r]$$

$$\times E_2[\mu(r_0 + r)] + \frac{1}{\mu r_0}$$

$$\times [\exp(-\mu(r_0 + r)) - \exp(-\mu(r_0 - r))]$$

where

$$E_2(x) = \int_1^\infty \frac{du}{u^2} e^{-zu}.$$
supplement them by independent calculations in the low energy region.

A multigroup, multiregion reactor analysis IBM-704 code\(^{11}\) applicable to water moderated systems was available and was used for this purpose.\(^*\) Since this code is limited to a constant extrapolation distance (independent of energy group), two independent calculations were made, one with an extrapolation length found from an overall group average, and the other with a value appropriate to the high energy region. The leakage in the high energy region predicted by the second calculation was evaluated by numerical integration and is found as curve \(A\) of Fig. 3.

Then the leakage spectrum \(N(E)\) for neutrons of energy \(E\) is given by

\[
N(E) = \sum_i A_i(E) S_A[a r_0 \mu_i(E)].
\]  

The attenuation data were fitted at several energies and the values of \(A_i\) and \(\mu_i\) were substituted in this equation. The result is shown in Fig. 4. Note that the normalization in this figure refers to one fission in the core rather than to one neutron so that the data obtained from this formula have been multiplied by 2.47 neutrons/fission to obtain \(N(E)\). Since the results of Aronson et al.\(^{10}\) are valid only for relatively high energies, it was necessary to

\fig{4}{Neutron leakage spectrum.}
from the assembly per fission occurring in the interior. The total dose, \( \int_0^\infty D_0(E)N(E)\,dE \), in this spectrum is found to be \( 1.08 \times 10^{-9} \) rad-cm\(^2\)/fission, and the ratio \( D_n/A \) is \( 1.95 \times 10^5 \) rad-cm\(^3\)/\( \mu \)c [equation (3)].

As stated above, it is assumed throughout this paper that the spectrum of escaping neutrons does not depend appreciably upon distance from the assembly, so that in equation (3) for \( D_n/A \) one may substitute \( N(E) \) for \( \phi(E) \). Although this can be only approximately true, since there is clearly an appreciable amount of scattering and degradation of the spectrum in the materials of the room, one expects that this will not lead to serious error in the dose estimates. Note that the thermal component of the calculated neutron spectrum is smaller than one finds experimentally (Table 1). The higher experimental value is probably explainable in terms of room scattering and is not inconsistent with the results of Hurst, Mills and Reinhardt* on the leakage spectrum of neutrons from the Godiva reactor.

Three sources of \( \gamma \)-radiation were considered in the analysis: (a) prompt \( \gamma \)-radiation emitted directly from the fission process, (b) capture \( \gamma \)-radiation resulting from \( (n,\gamma) \) reactions taking place within the assembly, and (c) delayed \( \gamma \)-radiation from the fission products contained within the assembly. The spectral distribution of the prompt \( \gamma \)-rays, \( \Gamma(E) \), was obtained directly from the recent work of Maienschein et al.\(^{(12)}\) and is shown in Fig. 5. The spatial distribution of this source is determined by the spatial distribution of the fission events and is assumed to be proportional to \( 1/r \sin (\pi r/r_0) \). The capture \( \gamma \)-sources were obtained from the following equation:

\[
\Gamma_j = v (1 - f_\nu) \left( \frac{\sigma_{\gamma\gamma}}{\sigma_\nu} \right)_j \left( \frac{E_\nu}{E_\gamma} \right)_j \left( \frac{\Sigma_{\gamma\gamma}}{\Sigma_\nu} \right)_j
\]  

\( j \) 

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* G. S. Hurst, W. A. Mills and P. W. Reinhardt, unpublished data on neutron leakage spectrum from the Godiva reactor at LASL.
Table 2. Capture $\gamma$-ray spectra

<table>
<thead>
<tr>
<th>Element</th>
<th>$E_c$ (MeV/capture)</th>
<th>$E_\gamma$ (MeV/photon)</th>
<th>$S$ (photons/fission)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydrogen</td>
<td>2.2</td>
<td>2.2</td>
<td>0.478</td>
</tr>
<tr>
<td>Uranium</td>
<td>6.8</td>
<td>3.0</td>
<td>0.427</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1.0</td>
<td>0.171</td>
</tr>
</tbody>
</table>

where

$\Gamma_j$ = photons of energy $E_\gamma$ released from the $j$th element per fission;
$r$ = neutrons born per fission (2.47);
$f_e$ = neutron escape fraction (0.271);
$\sigma_{rc}$ = microscopic radiative capture cross-section;
$\sigma_a$ = microscopic absorption cross-section;
$F_c$ = energy release per neutron capture in form of $\gamma$-rays of energy $E_\gamma$;
$\Sigma_a$ = macroscopic absorption cross-section;
$\Sigma_t$ = total macroscopic absorption cross-section of the assembly.

Uranium and hydrogen were the only elements in the assembly which contributed appreciably to the capture $\gamma$-dose. For these elements the values for $E_c$ and $E_\gamma$ were obtained from GLASTONE[13]. In the case of uranium these values are not well known but the total neutron binding energy of 6.8 MeV is accounted for correctly by the proposed $\gamma$-spectrum. The calculated values for the capture $\gamma$-sources are shown in Table 2. As in the case of the prompt radiation, the capture $\gamma$-rays are released essentially simultaneously with fission so that the spatial distribution again follows that of the fission events.

It was not necessary to determine the complete leakage spectrum of prompt $\gamma$-rays from fission and $\gamma$-rays from capture. One is interested primarily in the $\gamma$-dose escaping the system. The calculation of the leakage dose for prompt and capture $\gamma$-rays was carried out by first computing the variation of dose $D(r)$ with distance in an infinite medium having the composition of the assembly due to a unit point isotropic source having the indicated distribution of energy. Thus,

$$[D(r)]_{\text{prompt}} = K \int_0^\infty E \Gamma(E) \mu_a^t(E) B_d(E, \mu r) \times \frac{\exp (-\mu(E)r)}{4\pi r^2} dE$$

where

$D(r) = \gamma$-dose at $r$ cm from a unit source (rad/fission);
$K = 1.6 \times 10^{-8}$ rad/MeV-gm$^{-1}$;
$\Gamma(E) = \text{number of } \gamma\text{-rays of energy } E \text{ emitted per unit interval from one fission (photon/fission MeV)}$;
$\mu_a^t(E) = \text{energy absorption coefficient for tissue dose (cm}^2/\text{g)}$;
$B_d(E, \mu r) = \text{dose buildup factor}$[14];
$\mu(E) = \text{attenuation coefficient of water for photons of energy } E \text{ (cm}^{-1})$.

The integration was performed numerically using values of $B_d$ obtained from the work of GOLDSTEIN and WILKINS[14]. $[D(r)]_{\text{capture}}$ was evaluated in the same way except that a sum over the $j$ capture photon groups was carried out. The computed curves for $[D(r)]_{\text{prompt}}$ and $[D(r)]_{\text{capture}}$ were then fitted by empirical formulas of the form used in the neutron calculation,

$$D(r) = \frac{1}{4\pi r^2} \sum_i A_i \exp (-\mu_i r)$$

and the dose, $D_{\gamma}$, escaping per fission was found from the equation

$$D_{\gamma} = \sum_i A_i \mathcal{L}_A(\mu_i, \tau_0).$$

The treatment of the delayed radiation presents somewhat more of a problem as both
the source strength and the energy distribution are strongly time dependent during the first few seconds following fission. The spectral distribution shown in Fig. 3, was obtained by normalizing the spectrum measured by MAIENSCHEN et al.\textsuperscript{[12]} at 6.2 sec to the integral over the first 15 sec of the measured decay spectrum. It was necessary to extrapolate to obtain values during the first second. The delayed $\gamma$-emitters were assumed to be homogeneously distributed throughout the assembly because of turbulence following the initial release of energy, hence it was necessary to obtain transmission factors for uniformly distributed sources of $\gamma$-rays. The leakage fraction, $\mathcal{L}_B(\mu r_0)$ for a normalized uniform distribution $\mathcal{S}(r) = \frac{3}{4\pi r_0^3}$ may be written explicitly as

$$\mathcal{L}_B(\mu r_0) = \frac{3}{8} (\mu r_0)^{-3} \times \left[ \frac{2(\mu r_0)^2}{1 + (1 + 2\mu r_0) \exp (-2\mu r_0)} \right]$$ \hspace{1cm} (12)

which may be shown by carrying out the integration indicated in equation (6). This result is shown as curve B of Fig. 3.

The movements of the exposed persons subsequent to the initial excursion influenced the amount of $\gamma$-radiation which they received. In the dose analysis it was assumed that each person exposed was exposed to a single nuclear excursion but that his individual actions in the few seconds following the excursion resulted in different $\gamma$-neutron dose ratios. This results in a delayed $\gamma$-dose slightly larger than that obtained by assuming a constant power assembly delivering the same total fission energy. The following three cases treated were based upon the actions of each of the individuals as determined from personal interviews.\textsuperscript{[15]}

Case I (employee A). This employee was exposed to the full complement of prompt and capture $\gamma$-radiation but, since it is believed that he left the area first, it was assumed that only the first 5 sec of the delayed $\gamma$-spectrum contributed to his dose.

Case II (employee E). This employee was also exposed to the prompt and capture $\gamma$-radiation source, but the $\gamma$-neutron ratio was adjusted to take into account an assumed 15 sec exposure to the delayed $\gamma$-spectrum and, further, that his exit from the area took him to within 10 ft of the assembly. This case gives the largest $\gamma$-neutron dose ratio used in analysis and reflects the fact that this employee was apparently the last to leave the area. The assumed exit path resulted in a total $\gamma$-dose which was 8 per cent more than the dose of the constant distance 15 sec exposure case. This represents what is believed to be the maximum increase that could have occurred for any of the personnel.

Case III (employees other than A and E). These employees were assumed to have received the full contribution of prompt and capture $\gamma$-radiation, but, like case I, the delayed $\gamma$-rays were received at a constant distance and, like case II, the first 15 sec of the delay spectrum was effective.

The escaping dose from delayed $\gamma$-rays which was received in the three cases considered was determined by the same method used for the prompt and capture $\gamma$-dose calculation. The dose $[D_J(r)]_{\text{delayed}}$ was found from equation (9) in which the portion of the delayed spectrum $[\Gamma(E)]_{\text{delayed}}$ seen by the $J$th case was employed. $[D_J(r)]_{\text{delayed}}$ was then fitted by an empirical expression having the form of equation (10) and the escaping dose $[D_J]_{\text{delayed}}$ seen by case $J$ was calculated from the equation:

$$[D_J]_{\text{delayed}} = \left[ \sum_i A_i \mathcal{L}_B(\mu r_0) \right]^{(J)}$$

The results for the dose multiplied by $4\pi r^2$ are shown in Table 3.

IV. DOSE MEASUREMENTS WITH A MOCK-UP OF THE CRITICAL ASSEMBLY

To establish experimentally the relationship between the total (neutron + $\gamma$) first collision dose and the blood Na activation, a mock-up of the critical assembly was constructed and operated\textsuperscript{*} as a low power reactor in two experiments. Fig. 6 shows the actual reactor tank surrounded with a large tank which rested on a concrete floor. Table 4 gives details of the mock-up assembly as compared to the actual critical assembly. The first experiment measured the ratio of $\gamma$-dose $D_{\gamma}$ to neutron dose $D_n$, i.e., $D_{\gamma}/D_n$. The assembly was operated at a power of about 6 W for 11 min. The rate at 0 ft from the center of the assembly, as in Fig. 6, was measured with an ionization

\textsuperscript{*} This phase of the work was carried out at the ORNL criticality facility.
Table 3. Results of $\gamma$-dose calculation

<table>
<thead>
<tr>
<th>Case no.</th>
<th>Prompt $\gamma$-dose (rad-cm$^2$/fission)</th>
<th>Capture $\gamma$-dose (rad-cm$^2$/fission)</th>
<th>Delayed $\gamma$-dose (rad-cm$^2$/fission)</th>
<th>Total $\gamma$-dose (rad-cm$^2$/fission)</th>
<th>$D_\gamma/D_\alpha$</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>$2.06 \times 10^{-9}$</td>
<td>$6.4 \times 10^{-10}$</td>
<td>$4.0 \times 10^{-10}$</td>
<td>$3.10 \times 10^{-9}$</td>
<td>2.87</td>
</tr>
<tr>
<td>II</td>
<td>$2.06 \times 10^{-9}$</td>
<td>$6.4 \times 10^{-10}$</td>
<td>$9.4 \times 10^{-10}$</td>
<td>$3.64 \times 10^{-9}$</td>
<td>3.37</td>
</tr>
<tr>
<td>III</td>
<td>$2.06 \times 10^{-9}$</td>
<td>$6.4 \times 10^{-10}$</td>
<td>$6.4 \times 10^{-10}$</td>
<td>$3.34 \times 10^{-9}$</td>
<td>3.09</td>
</tr>
</tbody>
</table>

Table 4. Comparison of the mock-up critical assembly with the actual critical assembly

<table>
<thead>
<tr>
<th>Actual critical assembly</th>
<th>Mock-up critical assembly</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diameter (in.)</td>
<td>22</td>
</tr>
<tr>
<td>Height (in.)</td>
<td>9.2</td>
</tr>
<tr>
<td>$^{235}$U conc. (g/l.)</td>
<td>37.4</td>
</tr>
<tr>
<td>Total $^{235}$U (kg)</td>
<td>2.1</td>
</tr>
</tbody>
</table>

that a point on his plane of symmetry coincided with the point $A$ where the neutron dose was measured in the first experiment. The burro was chosen as the experimental animal because (a) he is a large

![Fig. 6. Schematic of mock-up experiment.](image)
animal comparable in size to man, and (b) the amount of sodium per gram of blood serum is approximately the same for burro and man. Table 5 shows the concentration of sodium in blood serum for the burro and for some of the exposed individuals.* Blood samples were drawn from the burro and counted for Na\(^{24}\) as described in the next section. From these data the value of \(S/D_n\) for this particular neutron spectrum was determined to be \(6.0 \times 10^{-6}\) \(\mu\)c of Na\(^{24}\) per ml of blood and per rad of neutron dose, compared to \(5.14 \times 10^{-6}\) \(\mu\)c obtained by using the methods given in Sections II and III. Note that the results calculated in Section II apply to the case where a model of a man is the irradiated object. The close agreement of these two values is further indication that the burro is a good phantom for a man.

Also, in the second experiment a series of threshold detectors\(^1\) was used to obtain information on the fast neutron spectrum. The neutron detectors [Au (Au + Cd), S\(^{32}\) and Pu\(^{239}\), Np \(^{237}\) and U\(^{235}\) in 2.2 g/cm\(^2\) Bio] were exposed at approximately 29 in. from the center of the liquid assembly, point \(B\) in Fig. 6. The total flux of neutrons received during the 42 min period for various energy ranges is shown in Table 1. The table also shows a comparison with the calculations given in Section III. Using the flux values measured at point \(B\) the first collision neutron dose can be calculated\(^{18}\) at the burro exposure position (point \(A\)). A sulfur disc was fixed on the side of the burro facing the assembly. The measured value of the flux was corrected for distance and for flux build-up due to the burro to obtain the flux which would have existed at point \(A\) had the burro not been there. The first collision neutron dose at point \(A\) was then determined from the ratio of the neutron flux at point \(A\) to that at point \(B\) and the neutron dose calculated for point \(B\). Thus the threshold detector data when suitably corrected to refer to point \(A\) gives a first collision neutron dose at point \(A\) equal to 42 rads compared to 48 rads using the proportional counter method.

The value of the thermal neutron flux in Table 1 shows that the dose due to thermal neutrons is negligible compared to the fast neutron dose. However, the magnitude of the thermal neutron flux is of interest because of another factor. In Fig. 1 it is seen that the probability that a thermal neutron entering the human body will produce Na\(^{24}\) is about 0.4 as large as the probability that a fast neutron will become thermalized and produce a Na capture in the body. Thus it would be expected that for the spectrum shown in Table 1 only about 15 per cent of the Na\(^{24}\) activity would be produced by thermal neutrons. This fact is of considerable practical importance since it implies that it is not necessary to consider the spatial distribution of thermal neutrons in detail.

V. ASSIGNMENT OF DOSE VALUES TO THE EXPOSED INDIVIDUALS

The blood samples collected from the individuals were counted with a 2 x 4 in. NaI crystal (Fig. 7). Polyethylene bottles (4 fluid oz capacity) containing the whole blood were placed on the top of the crystal. The horizontal lines indicate liquid levels for 50 and 100 ml. Counting was done with a discriminator set to accept \(\gamma\)-rays above 0.66 MeV and 2.0 MeV. The equipment was calibrated with 50 and 100 ml solutions of Na\(^{24}\) in water. The counting geometry for Na\(^{24}\) as well as the counter background (using a 4 in. lead shield) is shown in Table 6 for the two bias values and for 50 and 100 ml samples.

A few hours after exposure, 100 ml blood samples were taken from some of the individuals, and this was counted as whole blood without the use of an anticoagulant. A second set of blood samples was

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* These measurements were made by Dr. C. H. Steffee, Oak Ridge Hospital, using the flame photometer technique. Although it was not possible to obtain whole blood sodium values at the time the experimental work was done, a subsequent evaluation of whole blood sodium for unirradiated burros showed results not inconsistent with values to be expected in normal man.

† The amount of exposure was determined with a U\(^{235}\) fission chamber which had been calibrated in terms of fast neutron dose in the first experiment.

‡ The depth dose curves taken from W. S. Snyder, Brit. J. Radiol. 28, 342 (1955), and NBS Handbook 63, November 1957, were used as a basis of correction.
Table 6. Na\textsuperscript{24} counting geometry and background for 2 by 4 in. NaI crystal at two bias levels

<table>
<thead>
<tr>
<th>Bias (MeV)</th>
<th>Geometry (50 ml) (%)</th>
<th>Geometry (100 ml) (%)</th>
<th>Background (counts/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.66</td>
<td>17.7</td>
<td>13.3</td>
<td>250</td>
</tr>
<tr>
<td>2.0</td>
<td>5.9</td>
<td>4.3</td>
<td>115</td>
</tr>
</tbody>
</table>

Table 7. Na\textsuperscript{24} activation and dose values for exposed personnel

<table>
<thead>
<tr>
<th>Exposed individuals</th>
<th>Na\textsuperscript{24} (µc/ml)</th>
<th>Experimental neutron dose (rad)</th>
<th>Theoretical neutron dose (rad)</th>
<th>Experimental (\gamma)-dose (rad)</th>
<th>Theoretical (\gamma)-dose (rad)</th>
<th>Total experimental dose (rad)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>(5.8 \times 10^{-4})</td>
<td>96</td>
<td>113</td>
<td>269</td>
<td>473</td>
<td>365</td>
</tr>
<tr>
<td>B</td>
<td>(4.3 \times 10^{-4})</td>
<td>71</td>
<td>87</td>
<td>197</td>
<td>267</td>
<td>270</td>
</tr>
<tr>
<td>C</td>
<td>(5.4 \times 10^{-4})</td>
<td>89</td>
<td>104</td>
<td>250</td>
<td>332</td>
<td>339</td>
</tr>
<tr>
<td>D</td>
<td>(5.2 \times 10^{-4})</td>
<td>86</td>
<td>100</td>
<td>241</td>
<td>308</td>
<td>327</td>
</tr>
<tr>
<td>E</td>
<td>(3.7 \times 10^{-4})</td>
<td>62</td>
<td>72</td>
<td>174</td>
<td>223</td>
<td>236</td>
</tr>
<tr>
<td>F</td>
<td>(1.1 \times 10^{-4})</td>
<td>18</td>
<td>21</td>
<td>50.5</td>
<td>64.9</td>
<td>68.5</td>
</tr>
<tr>
<td>G</td>
<td>(1.1 \times 10^{-4})</td>
<td>18</td>
<td>21</td>
<td>50.5</td>
<td>64.9</td>
<td>68.5</td>
</tr>
<tr>
<td>H</td>
<td>(0.36 \times 10^{-4})</td>
<td>6.0</td>
<td>7.0</td>
<td>16.8</td>
<td>21.6</td>
<td>22.8</td>
</tr>
<tr>
<td>Burro</td>
<td>(2.9 \times 10^{-4})</td>
<td>48</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Using \(D_\gamma/D_\alpha = 2.8\).
† Using \(D_\gamma/D_\alpha = 3.09\).

When taken about 20 hr later, but this time only 50 ml was used and heparin was added to prevent clotting. This set of data was more nearly uniform than the first and it is the basis of the final dose values reported. The blood samples were counted for several days to establish that Na\textsuperscript{24} was the primary isotope being counted. Very early counting showed some evidence of K\textsuperscript{42} in the case when the counting bias was 0.66 MeV.

Table 7 shows the final results for the Na\textsuperscript{24} activity (extrapolated to the time of exposure) for the exposed personnel as well as that for the experimental burro. Blood samples (100 ml) from the burro, receiving a neutron dose of 48 rads, were counted† in the same way as described above for the exposed individuals.

Table 7 shows the first collision neutron dose, \(\gamma\)-dose, and total dose obtained from the experimental data by direct comparison of Na\textsuperscript{24} activities in the human blood with that of the burro blood, using the known fast neutron dose given to the burro and the experimentally measured ratio, \(D_\gamma/D_\alpha\). Also shown in Table 7 are the dose values obtained by calculations, Sections II and III.§

The counting methods described above are very sensitive. For example, it is seen by using data from Tables 6 and 7 that 1 rad of neutrons give a count rate due to Na\textsuperscript{24} equal to 40 counts/min (at zero time) for 50 ml burro blood when the counting bias is 2.0 MeV. Thus, since the counter background rate is 115 counts/min, a dose of 1 rad can be detected. With minor improvements in counting techniques, it should be possible to determine doses as low as 100 mrads with the Na\textsuperscript{24} method.

§ The calculated values given in the table differ from the corresponding values in report Y-1234, *Accidental Excursion at the Y-12 Plant*, (4, August 1958). The inadvertent use of a high value of the concentration of Na\textsuperscript{24} in whole blood gave calculated values which were too low.

† For the first few hours Cl\textsuperscript{38} competed with Na\textsuperscript{24}, especially at the 0.66 MeV bias.
VI. DISCUSSION OF ERRORS

As mentioned in Section I, comparison of calculations with experiment leads to independent tests of some of the assumptions made in the experimental program. The various sources of error in the experimental approach and the extent to which comparison of theory and experiment helps to decrease the uncertainty involved in individual assumptions will now be summarized. The assumptions made in the experimental program are:

1. The actual radiation spectrum is duplicated with the mock-up reactor.
2. Na\(^{24}\) activity in blood of the exposed person is approximately proportional to the first collision neutron dose, i.e. the Na\(^{24}\) activity resulting from capture of thermal neutrons coming from the reactor constitutes a correction factor.
3. The relationship of Na\(^{24}\) activity in the blood of a burro, exposed to the radiation from the mock-up reactor, to the first collision dose is valid for a man exposed to the actual radiations.
4. The \(\gamma\)-dose for each of the exposed persons may be established by determining the ratio \(D_\gamma/D_n\) for the mock-up facility.

Assumption (1) was checked by measuring the neutron leakage spectrum from the mock-up with threshold detectors. The neutron leakage spectrum from the actual assembly was calculated. Table 1 shows comparison of these two spectra. The \(\gamma\)-spectrum for the actual assembly was calculated with what is thought to be reasonable accuracy. Errors in dose due to the difference in spectra were thought to be negligible.

Assumption (2) may lead to error for some of the cases. It was shown by calculation (see Fig. 1) that thermal neutrons are less effective than fast neutrons in producing Na\(^{24}\) in the blood. Measurements of the thermal neutron flux for the exposed burro combined with this result show that only about 20 per cent of the Na\(^{24}\) is due to thermal neutrons coming directly from the reactor. However, for those persons at large distances from the reactor, the Na\(^{24}\) method may lead to overestimates of the dose. This point deserves further exploration.

Assumption (3) was checked by comparison with calculation. The relationship of neutron dose to Na\(^{24}\) activity was calculated for a man exposed to the neutron spectrum, found from calculations on the actual assembly, and compared to the value for a burro exposed to the mock-up. The value of rad per \(\mu\)c per ml of blood was 20 per cent higher for the calculated case.

Assumption (4) has already been considered in detail in Section III. Very little error results from this assumption.

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exposed persons was determined by Dr. C. H. Steffee, Oak Ridge Hospital. We would also like to thank the entire ORNL criticality group for their work in designing and operating the mock-up reactor experiments and the University of Tennessee AEC Agricultural Research Program for the work associated with the use of the burro. We are also pleased to acknowledge continued guidance and suggestions from K. Z. Morgan, H. P. Yockey and M. J. Cook. To the many others who were called upon to render services in their fields of speciality, we also extend our thanks.

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7. National Bureau of Standards Handbook 63 Fig. 1 (1957).