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Long-Term Fallout

A summary of measurements made through June 1957 by the gummed-film network of the AEC is presented.

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Several papers have described the phenomena of long-range fallout and the methods by which it is routinely monitored (1). This paper presents estimates of strontium-90 deposition and external gamma dose which were obtained from the world-wide gummed film network of

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the U.S. Atomic Energy Commission through June 1957. Results for the continental United States and other stations are tabulated in Table 1; results for the worldwide network are mapped in Fig. 1. In addition, the estimates of strontium-90 deposition as obtained by the gummed-film method are compared with measured values obtained by sampling with open pots.

Because of their mass, it is not practical to present the detailed analytical results in this article (2). This presentation, therefore, is limited to a condensation of the cumulative fallout observations.

Sampling and Measurement

A primary technique in studying long-range fallout is the measurement of the rate of deposition and the cumulative deposit per unit area. For this purpose, three types of samples are currently used: soils, pots or funnels, and gummed film.

Soil samples represent the accumulated fallout at a given location, but these samples require tedious radiochemical analyses for the determination of specific isotopes. Moreover, soil sampling does not permit one to estimate the external gamma dose delivered by the isotopes because of difficulty in analysis and uncertainty in the time of fallout.

Open samplers, such as pots or fun-

nels, permit collection of individual rainfalls or weekly or monthly deposits, from which strontium-90 and other isotopes may be determined directly by radiochemical analyses. Gamma emitters may be evaluated by spectroscopy.

The principal advantage of the gummed-film method, in addition to its simplicity, is that it permits daily sampling. This is important for the estimation of gamma dose.

There can be no absolute sampling procedure for fallout deposition because the deposition in a given situation will be influenced by the type of surface.

However, to permit some basis of comparison, the collection performance of the gummed film has been studied in relation to the collection performance of pots.

In earlier reports, it has been shown that the gummed film, under conditions of moderate rainfall in a temperate climate, yields fallout samples with an overall efficiency of about 63 percent compared with the values from high-walled pots. In regions where much of the fallout occurs with snow, the gummed-film method may grossly underestimate the true fallout values. Despite this objec-

tion, the gummed-film technique has proved to be desirable because of the simplicity with which daily samples can be accumulated from a large number of widely scattered locations.

Since late 1954, the computation of strontium-90 from the total beta activity of the gummed-film samples has become increasingly difficult because the computed values are sensitive to the assumed age of the debris. The accumulation of long-lived fission products in the stratosphere and the greater frequency of weapon tests has greatly complicated the problem of assigning an age to the

Table 1. Strontium-90 deposition and cumulative gamma dose as estimated by gummed-film measurements through June 1957.

Station	Sr ⁹⁰ (mc/mi ²)	Ex- ternal γ dose (mrad)*	Station	Sr ⁹⁰ (mc/mi ²)	Ex- ternal γ dose (mrad)*	Station	Sr ⁹⁰ (mc/mi ²)	Ex- ternal γ dose (mrad)*
<i>Outside continental United States</i>			Iceland			Thailand		
Alaska			Keflavik	21	36	Bangkok	10	39
Anchorage	12	20	Italy			Tripoli		
Fairbanks	15	26	Milan	13	23	Libya	24	41
Juneau	16	30	Japan			Union of South Africa		
Nome	9	17	Hiroshima	19	36	Durban	4	8
Argentina			Misawa	20	39	Pretoria	10	19
Buenos Aires	9	18	Nagasaki	21	41			
Australia			Tokyo	23	43	<i>Continental United States</i>		
Sydney	6	17	Liberia			Albuquerque, N.M.	45	150
Bermuda	21	43	Monrovia	10	19	Atlanta, Ga.	20	41
Bolivia			Malaya			Billings, Mont.	26	58
La Paz	9	22	Singapore	7	23	Binghamton, N.Y.	13	25
Canada			Mexico			Boise, Idaho	27	44
Churchill, Manitoba	6	11	Mexico City	16	39	Boston, Mass.	20	69
Edmonton, Alberta	18	33	Morocco			Cape Hatteras, N.C.	14	29
Goose Bay, Labrador	13	29	Sidi Slimane	18	30	Chicago, Ill.	22	50
Moncton, New Brun- swick	13	25	New Zealand			Cleveland, Ohio	25	65
Montreal, Québec	16	33	Wellington	5	10	Concord, N.H.	11	26
Moosonee, Ontario	13	29	Nigeria			Corpus Christi, Tex.	12	25
North Bay, Ontario	17	34	Lagos	8	14	Dallas, Tex.	25	60
Ottawa, Ontario	12	25	Norway			Des Moines, Iowa	27	63
Regina, Saskatchewan	13	27	Oslo	13	23	Detroit, Mich.	22	49
Seven Islands, Quebec	12	27	Pacific Ocean			Grand Junction, Colo.	39	160
Stephenville, New- foundland	20	42	Yap, Caroline Islands	17	52	Jacksonville, Fla.	13	30
Winnipeg, Manitoba	23	45	Guam, Caroline Islands	78	160	Knoxville, Tenn.	18	45
Ceylon			Truk, Caroline Islands	33	87	Las Vegas, Nev.	23	66
Colombo	9	29	Ponape, Caroline Islands	41	140	Los Angeles, Calif.	11	20
Costa Rica			Canton Island	7	23	Louisville, Ky.	24	54
San Jose	7	17	Iwo Jima	36	170	Medford, Oreg.	13	23
Ecuador			Johnston Island	30	65	Memphis, Tenn.	24	75
Quito	5	14	Koror, Palau Island	14	44	Miami, Fla.	16	37
Ethiopia			Manila, Philippine Islands	17	48	Minneapolis, Minn.	25	51
Addis Ababa	11	21	Midway Island	19	36	New Haven, Conn.	20	43
French West Africa			Noumea, New Caledonia	8	20	New Orleans, La.	28	64
Dakar	12	22	Wake Island	22	45	New York, N.Y.	28	54
Germany			Panama Canal Zone	9	22	Philadelphia, Pa.	19	39
Rhein Main	15	27	Puerto Rico			Pittsburgh, Pa.	26	46
Greenland			San Juan	15	29	Rapid City, S.D.	18	45
Thule	9	15	Saudi Arabia			Rochester, N.Y.	19	37
Hawaii			Dhahran	15	28	Salt Lake City, Utah	54	180
French Frigate Shoals	21	42	Scotland			San Francisco, Calif.	14	23
Lihue	18	38	Prestwick	18	30	Scottsbluff, Neb.	38	73
Hilo	30	59				Seattle, Wash.	19	34
Honolulu	16	34				Tucson, Ariz.	25	49
						Washington, D.C.	18	35
						Wichita, Kan.	25	62

* The tabulated values are calculated infinity external gamma dose in millirad. The probable exposure to the population, allowing for shielding and weathering, is approximately 10 percent of this value.

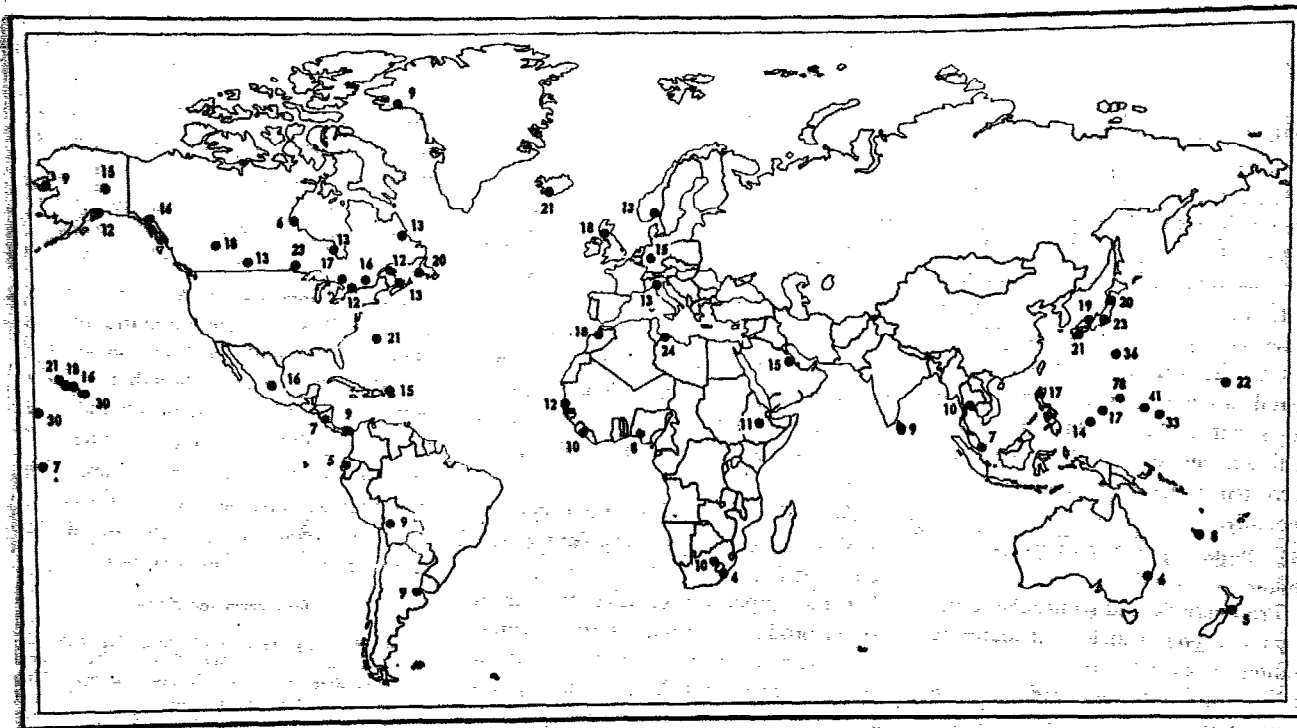


Fig. 1. Calculated cumulative strontium-90 fallout in millicuries per square mile as of June 1957.

debris. However, a method of computation has been devised by which the latter difficulty can be minimized.

Methods of Computation

The adhesive-coated films, which have been exposed for 24 hours, are shipped to the U.S. Atomic Energy Commission's Health and Safety Laboratory in New York. The total beta activity of the ashed samples is measured and corrected by the 63 percent efficiency factor. The strontium-90 component of the fallout is calculated from modified Hunter and Ballou (3) ratios. In addition, an estimate of the infinity external gamma dose in air is made from the beta activity (4).

The original calculations of strontium-90 deposition from measurements of total beta activity on ashed gummed-film samples were performed as follows:

1) The activity measured on a given sampling day was attributed to the test immediately preceding that sampling day.

2) The measured activity on the counting day was extrapolated to a fixed day by the formula

$$A = A_0 t^{-1.2}$$

3) The strontium-90 fraction of the total beta activity on this day was taken from modified Hunter and Ballou curves.

4) The strontium-90 activity values for the individual days were summed by

months, and these sums were added for the desired period of accumulation.

The assignment of activity on a given day to the most recent test was a reasonable approximation during the period of tropospheric fallout. The deviations between gummed-film estimates and radiochemical analyses became larger as the contribution from stratospheric fallout increased. To improve the estimation of strontium-90, a system was devised which takes stratospheric debris into account. Tests of this simplified model yielded values that are in good agreement with computations from more complex mod-

els. This method, which has been applied to data subsequent to May 1956, is as follows:

1) Estimates of the yields of total fission products and of strontium-90 are obtained for each weapon test.

2) The total fission-product yield for each test is added to the calculated fission-product residue from previous tests. (The $t^{-1.2}$ law is used for decaying total fission product activity.)

3) The strontium-90 activity from each test is added to the accumulated strontium-90 activity from previous tests.

4) For each sampling day, the stron-

Table 2. Comparison of strontium-90 estimates from gummed-film with radiochemical analysis of monthly pot collections.

Period of observation	Total Sr ⁹⁰ (mc/mi ²)		Ratio film/pot	Monthly ratios		Film/pot mean
	Film	Pots		Low	High	
5/56-6/57	12.3	13.7	<i>New York City</i>	0.32	2.2	1.1
			0.90			
5/56-6/57	12.1	10.6	<i>Pittsburgh</i>	0.62	2.5	1.2
			1.14			
12/56-6/57	6.3	4.6	<i>Chicago</i>	1.0	1.9	1.4
			1.37			
12/56-6/57	15.1	9.1	<i>Salt Lake City</i>	1.1	3.3	1.8
			1.66			
12/56-6/57	3.5	3.1	<i>Los Angeles</i>	0.78	2.4	1.4
			1.13			
10/56-6/57	5.6	3.7	<i>Hiroshima</i>	0.82	3.7	1.7
			1.51			
8/56-6/57	6.7	5.5	<i>Nagasaki</i>	0.64	5.5	1.6
			1.22			

strontium-90/total-fission-product-activity ratio is calculated.

5) Each day's measured beta activity is converted to strontium-90 activity by use of this factor.

This method of calculation would give high strontium values for locations near test sites on days of high fallout. This is caused by the attribution of activity to the total accumulated pool of fission products rather than to the immediate burst which caused the fallout. This can be corrected by treating these few cases individually.

The major approximations of this technique are as follows:

1) Tropospheric and stratospheric debris enter a pool which contributes to the fallout at each location.

2) The mixed fission products from each detonation decay according to the $t^{-1.2}$ law.

3) The relative tropospheric and stratospheric depletion rates are not considered at this time.

The only practical means of evaluating the new calculation technique is by comparison with radiochemical analyses of open samplers. During the period from May 1956 to June 1957, several locations had parallel sampling units for

at least part of the time. These data are shown in Table 2, in which it is shown that the gummed-film system, together with the above-mentioned method of computation, yields estimates of strontium-90 deposition which tend to be higher than the estimates derived by radiochemical analyses of pot samples. The mean ratio of strontium-90 estimated from gummed-film to pot analyses is 1.45, with a maximum ratio of 1.66 at Salt Lake City and a minimum of 0.90 in New York City.

The calculation of external gamma dose is less sensitive to variations in the source of fallout. In addition, it appears that the important gamma dose from fission products is from internal cesium-137 rather than from the external gamma radiation from distributed fission products after suitable allowance for shielding and weathering.

Conclusions

The range of values for strontium-90 deposition through June 1957 in the United States is 11 to 54 millicuries per square mile, which is somewhat higher than other large land areas of the world.

Excluding the United States, deposition in the Northern Hemisphere averages 16 millicuries per square mile, about twice the average for the somewhat fewer values reported in the Southern Hemisphere.

The calculated external gamma doses given in Table 1 are estimates of the infinity doses and have not been corrected for shielding and weathering. Our best estimate of the actual external dose to the population is approximately 10 percent of the tabulated values. The dose may actually be lower, but a factor of 10 is a conservative estimate of the effect of shielding and weathering.

References and Notes

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2. We wish to acknowledge the continued cooperation of the U.S. Weather Bureau in the collection of gummed-film samples. The computations and data handling were performed by Dr. A. E. Brandt and Dr. George D. Diehl of the Biometrics Branch of the Health and Safety Laboratory.
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