Formerly Utilized Sites Remedial Action Program (FUSRAP)

Maywood Chemical Company Superfund Site

ADMINISTRATIVE RECORD

Document Number

MISS-001.



US Army Corps of Engineers®

* & LICENSIALC

人で小田

NERGY MEASUREMENTS GROUP

M-036 EG&G Survey Report NRC-8109 April 1981

AN AERIAL RADIOLOGIC SURVEY OF THE STEPAN CHEMCIAL COMPANY

AND SURROUNDING AREA MAYWOOD, NEW JERSEY

DATE OF SURVEY: 26 JANUARY 1981

J.R. Mueller Project Director S.A. Gunn Project Scientist

APPROVED FOR DISTRIBUTION

W. Jol

W. John Tipton, Head Radiation Sciences Section

This Document is UNCLASSIFIED

G. P. Stobie Classification Officer

This work was performed by EG&G for the United States Nuclear Regulatory Commission through an EAO transfer of funds to Contract No. DE-AC08-76NV01183 with the United States Department of Energy.

COCHR13

ABSTRACT

An aerial radiologic survey to measure terrestrial gamma radiation was performed in Maywood, New Jersey over the Stepan Chemical Company and the surrounding area. This survey was conducted by EG&G for the Nuclear Regulatory Commission (NRC) 26 January 1981.

Gemma-photon data were collected over a four square mile area. Processed data indicated that detected radioisotopes and their associated gamma-photon exposure rates were consistent with those expected from normal background emitters, except directly over and immediately to the west and south of the Stepan Chemical Company. In addition, two other points demonstrated anomalous gamma-photon activity: one north of the plant and another to the southeast, both approximately one-half mile from the center of the plant.

The results are expressed as exposure rate isoradiation contours extrapolated to μ R/h at 1 meter above the ground. The background radiation, including cosmic ray contributions, generally ranged from 6 to 7.5 μ R/h. Isoradiation contours are also shown for excess radiation from the thorium chain, pinpointing the anomalous areas.

CONTENTS

ŧ

| 3 | | Abstract |
|---|-----|---------------------------------|
| ł | Sec | tions |
| 7 | 1.0 | Summary |
| 7 | 2.0 | Introduction |
| 7 | 3.0 | Natural Background Radiation |
| 8 | 4.0 | Site Description |
| 8 | 5.0 | Survey Procedures and Equipment |
| 9 | 6.0 | Results and Conclusion |
| 9 | 7.0 | Earth Sample Analysis |
| | | |

Tables

- Earth Sample Qualitative/Quantitative Spectrometric Results 0 1
- Earth Sample Exposure Rate Conversion Factors 11 2
- Radionuclide Component Microroentgens/Hour 11 3

Figures

- 13 MBB-105S Helicopter 1
- Mobile Computer Processing Laboratory 13 2
- Exposure Rate Isoradiation Contours 15 3
- Isoradiation Contours for Excess Radiation from the Thorium-232 Chain 17 4 Inferred from Measurement of the Thallium-208 Photopeak
- Net Spectrum Over Anomalous Area 19 5
- Locations of Soil Samples Taken for Analysis 21 6
- A-1 Point Source Isopieths for Energy Window 2.52 to 2.72 MeV at 61 Meter Altitude 24
- 23 Appendix
- 25 References
- 27 Distribution

000431 7

1.0 SUMMARY

An acria' radiologic survey of the Stepan Chemical Company and the surrounding area in Maywood, New Jersey, was conducted on 26 January 1981 by the Washington Aerial Measurements Department of EG&G, Inc.

Gamma radiation was detected by 20 sodium lodide (thallium activated) crystals, arranged in 2 pods of 10 crystals each mounted on either side of a Messerschmitt-Bolkow-Blohm MBB-105S helicopter. The survey altitude was 61 meters; parallel flight lines were spaced 76 meters apart. The survey covered an area of 4 square statute miles centered on the Stepan Chemical Company.

Gamma-photon data, altitude and all supporting information were recorded each second along the flight lines. An isoradiation contour map, including all sources of gamma radiation, was prepared to show variations in total gammaphoton activity over the site. A second isoradiation map was constructed which shows concentrations of thalium-208, indicating the presence of excess thorium-232.

Areas of higher than normal gamma-photon activity were observed directly over and to the west and south of the plant. Two other areas that showed an increase in thorium concentrations are located (1) near the intersection of Coles Brook and the railroad track, approximately 0.1 mile north of Essex Street; and (2) west of Passaic Street on Latham. Ground surveys at these locations are required to determine the source of these anomalies. Other areas showing radiation levels above the average background, including the Riverside Cemetery, are likely to be variations in the natural radiation levels.

Indicated activity (due to excess thorium-232) directly over the plant was in the range of 40-70 microroentgens per hour (μ R/h), compared to 6 to 7.5 μ R/h typical of background exposure rates for the area.

2.0 INTRODUCTION

The United States Department of Energy (DOE) maintains an aerial surveillance operation called the Aerial Measuring System (AMS). AMS is operated for DOE by EG&G This continuing nationwide program, started in 1958, involves surveys to monitor radiation in and around facilities producing, utilizing, or storing radioactive materials. The purpose of these surveys is to document, at a given point in time, the location of all areas containing gammaphoton emitting radionuclides (visible at the surface) and to aid DOE pesonnel in evaluating the magnitude and spatial extent of any radioactive contaminants freleased into the environment. At the request of DOE (or other federal and state agencies), AMS is deployed for various aerial survey operations.

This report is the result of a survey requested by Region 1 of the Nuclear Regulatory Commission. The measurements reported here were mode from a base of operations at the Teterboro Airport three miles south of the survey area on 26 January 1981.

Aerial radiation detection systems average the radiation levels due to gamma-photon emitting radionuclides existing over an area of several acres. The systems are capable of detecting anomalous gamma-photon count rates and determining the specific radionuclides causing the anomalies; however, because of area averaging, they tend to underestimate the magnitude of localized sources as compared with ground-based readings.

The results of the survey are reported, where possible, as radiation exposure rates in μ R/h at 1 meter above the ground surface. Approximate annual radiation dose levels, expressed as millirem per year (mrem/y), are obtained by multiplying μ R/h by 8.76. This conversion number applies only to the external radiation dose component.

3.0 NATURAL BACKGROUND RADIATION

Natural background radiation originates from radioactive elements present in the materials of the earth and cosmic rays entering from space. The terrestrial gamma-photons originate primarily from the uranium and thorium decay chains and radioactive potassium. Local concentrations of these nuclides produce radiation levels at the surface of the earth ranging from 1 to 15 μ R/h (or 9 to 130 mrem/y). Some areas with high uranium and thorium concentrations in surface minerals exhibit even higher radiation levels, especially in the western states. (For example, in the Colorado Plateau the average radiation level is above 200 mrem/y). mrem/y). One member of each of the uranium and thorium decay chains is a noble gas which can diffuse through the soil and be airborne to other locations. Therefore, the level of airborne radiation depends on the meteorological conditions, the mineral content of the soil, the soil permeability, etc. existing at each location at a particular time. The airborne radiation contributes from 1 to 10% of the natural background radiation levels.

Cosmic rays, the space component, interact in a complicated manner with the elements of the earth's atmosphere and the soil. These interactions produce an additional natural source of gamma radiation. Radiation levels due to cosmic rays vary with altitude and geomagnetic latitude and range from 3.7 to 23 μ R/h (up to 200 mrem/y).1

4.0 SITE DESCRIPTION

8

The original site of the Maywood Chemical Company, now owned by the Stepan Chemical Company, was bounded on the east by Maywood Avenue, on the south by the Sears distribution warehouse, on the north by the New York Susquehanna and Western Railroad an(extended to the west as far as West End Parkway.

The site was developed in 1895 as the Maywood Chemical Works. From about 1916 until 1957 the Maywood Chemical Company was engaged in processing thorium for use in the manufacture of gas mantles for various lighting devices. The area to the west of the plant was used as a waste dump prior to construction of Route 17 in 1932, which now bisects this area.

"Thorium is widely distributed in nature, but rich deposites are scarce. Traces of thorium have been found in over 100 minerals, a few of which contain more than 1% thorium. Monazite sand, however, is the only commercially feasible source of thorium. Monazite sand occurs as a primary mineral in granites, gneisses and pegmatites. In 1893, Welsbach patented the use of a mixture containing 98 to 99% Thoria for gas mantles and the industry grew rapidly thereafter. In the years preceding World War I, over 300 million gas mantles were consumed annually".²

In 1959 the Illinois-based Stepan Chemical Company purchased the Maywood Chemical property and in 1960 initiated a cleanup of the

area,

The affected property west of Route 17, known as the Rochelle Park side of the property, no longer belongs to Stepan Chemical. SWS Industries, a Maywood, New Jersey based firm, is now considering this site for construction of an officewarehouse.

5.0 SURVEY PROCEDURES AND EQUIPMENT

A United States Geodetic Survey (USGS) map was used to define the area to be surveyed. Parallel lines spaced at 76 meter intervals were flown at an altitude of 61 meters.

The survey vehicle, a MBB-105S helicopter (Figure 1), carried a pilot, an equipment operator and a lightweight version of a specialized data recording system. Two detector pods were mounted on the sides of the helicopter; each pod contained ten sodium iodide, NaI (TI), detectors. The crystal in each detector was 12.7 centimeters in diameter and 5.1 centimeters in height. Gamma-photon signals from twenty detectors were summed and routed through an analog-todigital converter and pulse-height analyzer. Gamma-photon counting rates and energy spectral data were accumulated in 1-second intervals and recorded on magnetic tape.

The helicopter position was established with two systems: a Microwave Ranging System (MRS) and a radar altimeter. The MRS master station, mounted in the helicopter, interrogated two remote transceivers mounted at stationary positions overlooking the survey area. By measuring the round trip propagation time between master and remote stations, the master computed the distance to each. These distances were also recorded on magnetic tape each second. In subsequent computer processing, they were converted to absolute position coordinates.

The radar altimeter determined the aircraft altitude by converting the time of a round trip pulsed signal to distance between the aircraft and the ground. These data were also recorded on magnetic tape so that any variations in gammaphoton signal strength caused by altitude fluctuation could be compensated for accurately. The detectors and electronic systems that

160431

accumulated and recorded the data are described in detail in previous reports.³⁴

ta processing was done primarily with a nputer baced analysis laboratory system in the Hemote Sensing Laboratory, located at Andrews Air Force Base, Suitland, Maryland. Frequently, such analyses are carried out with a computer moulted in a mobile van (Figure 2). An extensive collection of software routines was available for data processing. The first data reduction that was accomplished produced gross count isoradiation contours. These contours were constructed from gross count rate numbers, which refer to integral count rates in that portion of the gamma-photon energy spectrum between 0.05 and 3.0 MeV (Figure 3).

A smaller portion of the spectrum was used to separate the fraction of the total activity due to a specific nuclide and to quantify its concentration in the ground. This operation was accomplished by computer processing the data with an algorithm that examined and combined certain regions of the spectrum. A more detailed discussion of the data processing methods typical of most aerial surveys is given in the Appendix.

6.0 RESULTS AND CONCLUSIONS

- Shown in Figure 3 are exposure rate isoradiation contours (derived from gross count rates) overlaid on a USGS map. The average natural background in the area is approximately 6 to 7.5 μ R/h, which includes a cosmic radiation component of 3.7 μ R/h. The accuracy of exposure rates computed for areas of elevated thallium-208 activity may have been compromised by difficulties encountered in detarmination of proper conversion factors (see Appendix).
- Shown in Figure 4 are isoradiation levels for excess radiation from the thorium-232 chain inferred from measurement of the thallium-208 photopeak. This refers to quantities of radioactive thallium-208 (2.62 MeV photopeak) over and above that observed in the average natural background of the area. The majority of namma radiation from the thorium-232 chain is mined by thallium-208 The lechnique for determining the exposure rate at 1 meter above ground level is discussed in the Appendix.

: Both isoradiation maps show increased levels of activity centered on the Stepan plant, as well as to the west and south. Two detached areas (to the north and southeast of the plant) also show slight increases of thorium concentrations. It is not known whether these increased concentrations are a result of plant activities or natural radiationanomalies. Ground surveys are necessary for this determination. The gross count contour mat (Figure 3) shows two additional areas of increased activity not shown in Figure 4. Spectrol analysis indicated only an increase in the activity of the natural radioisotopic mix in these areas. Such variations in the natural radiation levels are not unusual. It should also be noted that the levels given in Figures 3 and 4 are normalized to 1 meter above the ground, but only as averages over a large area. Dependent on the nuclides detected and their activity and spatial extent, ground level exposure rates inferred from aerial measurements can differ by large factors from the actual value at a specific point on the ground. A portable radiation detector held 1 meter above the ground will measure activity directly below the detector and in a relatively small circle around it. At 61 meters the helicopter detector system effectively averages the activity from a much larger area. The small source limitation in discussed in the Appendix under the heading. Spatial Resolution Function.

Exposure rate isopleths in Figures 3 and 4 may not agree (after correcting for average natural background exposure rate) for the following reasons.

- 1. Conversion factors are based on sources of infinite lateral extent, whereas some high activity areas may be of small extent.
- 2. The excess thallium-208 conversion factor is calculated with the assumptions defined in the Appendix.
- 3. The gross count conversion factor applies only to typical mixes of natural emitters.

Figure 5 presents a gamma-photon energy spectrum taken over the plant site. A background spectrum has been subtracted from the data presented in Figure 5. Photopeaks characteristic of thallium-208 are prominent in this spectrum.

7.0 EARTH SAMPLE ANALYSIS

The accuracy to which the terrestrial radiologic environment can be determined from airborne

| Table 2. Earth Sample Exposure Rate Conversion Factors* | | | | | | | | |
|--|-------------------------------------|--|--|--|--|--|--|--|
| Uranium-238 | 0.62 µR/h per ppm radionuclide | | | | | | | |
| Thorium-232 | 0.31 μR/h per ppm radionuclide | | | | | | | |
| Cesium-137 | 0.13 µR/h per pCi/g radionuclide | | | | | | | |
| Potassium-40 | .179 uR/h per pCi/g radionuclide | | | | | | | |

*Assumes uniform concentrations both vertically and horizontally.

| Table 3. Radionucilde Component Microroentgens/hour | | | | | | | | | | | |
|---|-----------------|-----------------|----------------|------------------|---------------|-------|-----------------|--|--|--|--|
| Earth Sample | Uranium- 238 | Thorlum- 232 | Cesium- 137 | Potassium- 40 | Cosmic Ray | Total | Aerial Total | | | | |
| B | 1.22 | 4.52 | | 1.79 | 3.7 | 11.2 | 17-25 | | | | |
| С | 1.29 | 3.97 | .02 | 1.76 | 3.7 | 10.7 | 17-25 | | | | |
| D | .94 | 2.40 | | 1.72 | 3.7 | 8.8 | 17-25 | | | | |

060437 11





MOBILE COMPUTER PROCESSING LABORATORY



Figure 3. EXPOSURE RATE ISORADIATION CONTOURS

S



Figure 4. ISORADIATION CONTOURS FOR EXCESS RADIATION FROM THE THORIUM-232 CHAIN, INFERRED FROM MEASUREMENT OF THE THALLIUM-208 PHOTOPEAK (2.62 MeV)





060431 21



Figure 6. LOCATIONS OF SOIL SAMPLES TAKEN FOR ANALYSIS

060431 23

APPENDIX CALIBRATION PROCEDURES AND CONVERSION FACTORS

GROSS COUNTS

Gress counts refer to the sum of counts in that portion of the gamma-photon energy spectrum between 0.05 and 3.0 MeV. The detectors are calibrated by flying over a selected area of land near a body of water. The terrestrial component of the gross count rate is obtained from the land data by subtracting the water data, consisting only of those counts due to aircraft background, airborne radon daughter components and cosmic rays. The gross count conversion factor is established by comparing this land-water difference with exposure rates measured on the ground after they are similarly corrected for radon and cosmic ray contributions. The ratio of exposure rate to the land-water difference of the gross count rate is the factor that converts the measured gross count rate of terrestrial origin to the corresponding exposure rate of terrestrial origin. The conversion factor used for this survey was 1120 counts per second at 51 m altitude equals $1 \mu R/h$ at 1 m.

The terrain in the calibration area contained a typical mix of naturally occurring radionuclides, consisting of potassium-40 and members of the uranium and thorium chains. A different mix will modify the shape of the spectrum over the energy interval covered by gross counts. Since the gross count conversion factor is dependent on spectral shape, the established conversion factor will not apply precisely to areas where the mix is atypical or where extraneous radionuclides are present.

THORIUM-232 ISOPLETHS

Since spectral extractions revealed anomalous concentrations of thallium-208 and ancestors, the magnetic tape data were processed to isolate effects from this natural chain. For this purpose, an energy window centered on the prominent 2.62 MeV gamma photon from thallium-208 was monitored.

Figure 4 shows the isopleths relating to this window after naturally occurring concentrations have been suppressed by subtracting a constant

equal to the window count rate over "natural" areas in the vicinity of the anomalous areas.

These window count rates arise from "excess" thallium-208 photons that reach the detector without interacting in the air or soil. Since uncollided photons of this type behave in a mathematically predictable way, the window count rates can be related to soil concentration through the survey and detector system geometry.^{6,7} A uniform distribution with depth has been assumed in order to generate column 4 from column 2 in the conversion scale of Figure 4. In addition, the angular response of the detector has been assumed to be an average between the two extremes of isotropic and cosine. The data of Beck et al.⁶ have been used to generate column 3 from column 4 in this conversion scale.

It should be noted that the values given in the conversion scales in Figure 4 are given to two significant figures. These theoretical values are correct within the context of the assumptions; however, this does not imply that the isopleth values are accurate to these significant figures. Due to the uncertainty in determinations of many of the parameters that relate to the air-to-ground conversion factors, the exposure rate values given in Figures 3 and 4 may be uncertain by ±25% for values relating to the natural radiation level, up to about 20 μ R/h. For those areas containing elevated radiation levels, the exposure rate values are expected to be within a factor of 2 if the radiation levels at 1 m are averaged over at least several acres.

SPATIAL RESCLUTION FUNCTION

A useful way of viewing the small source limitation is through the concept of spatial resolution function. This function is the relative count rate, measured at survey altitude, versus lateral distance from a point source on the ground. Count rates that are recorded during a survey are the result of mathematically "folding" the true ground distribution with this resolution function. This folding process can be performed easily. The reverse process, the one of interest, is lengthy and inaccurate; it has not, therefore, been attempted. It is, however, often instructive to compare the resolution function contours with contours masured over certain areas to gain some insight into the lateral extent of the ground activity. If the resolution function isopleths are similar to the measured isopleths, the source is probably highly localized relative to dimensions comparable to the survey altitude.

The point source (or resolution function) contours are generated from a calculated resolution function, which depends on gammaphoton energy, depth distribution of the source in the soil, and the angular response of the detector system. The resolution function for gross counts cannot be calculated due to the lack of energy definition. The resolution function for gammaphoton energy window counts can be calculated, but the result still suffers from uncertainties in depth distribution and angular response of the detectors. In practice, two calulations are performed to bracket the true resolution function for the energy window data.

Figure A-1 shows bracketing contours for the thallium energy window counts. The radius of the D levels were chosen to match the nearly circular D levels in Figure 4. The radii of contours of lower levels surrounding the two D levels are similar to these in Figure A-1. Therefore, these sources are probably localized relative to dimensions of a few hundred feet. The tack of a D level, coupled with the large C level radius in the third active area in Figure 4, suggests that this source has some apatial extent.

060431

The width of the spatial resolution is a measure of how far equal activity point sources must be separated on the ground in order to appear as separate sources in the overflight data. This width is, therefore, a measure of the distances through which point sources can exert their influence. It is obvious that this distance will depend on the point source activity, since the vertical resolution function scale is relative. In practice, the resolution function is reduced to about half its maximum value at lateral distances from the source equal to the survey altitude.



Figure A-1. POINT SOURCE ISOPLETHS FOR ENERGY WINDOW 2.52 TO 2.72 MeV AT 61 METERS ALTITUDE

XOURI 25

REFERENCES

- Klement, A.W.; Miller, C.R.; Min, R.P.; and B. Shleren. August 1972. Estimate of Ionizing Radiation Doses in the United States 1960-2000. R.S. EPA Report ORP/CD72-1. Washington, D.C.: Environmental Protection Agency.
- 2 Gurinsky, D.H. and Dienes, G.J. 1956. "Nuclear Fuels." The Geneve Series on the Peaceful Uses of Atomic Energy. Princeton, N.J.: D. Van Nostrand Company, Inc.
- 3. Boyns, P.K. 1976. The Aerial Radiological Measuring System (ARMS): Systems, Procedures and Sensitivity. Report No. EGG-1183-1691. Las Vegas, NV: EG&G.
- 4. Jobst, J.E. 1979. "The Aerial Measuring Systems Program". Nuclear Safety, 20: 136-47.
- 5. Mohr, R.; Fritzsche, A.; and Franks L. January 1976. Ground Survey Procedures. Report No. EGG-1183-2339. Las Vegas, NV: EG&G.
- Stuart, T.P. May 1978. "Factors That Convert Aerial Measurements to Concentration of Gamma-Ray Emitters in the Soil — An Updated Calculational Procedure." Company Memorandum. Las Vegas, NV: EG&G.
- 7. Stuart, T.P. August 1977. Limiting Values for Radionuclide Concentration in the Soil from Remote Spectrometer Measurements. Report No. EGG-1183-1716. Las Vegas, NV: EG&G.
- 8. Beck, H.L.; DeCampo, J.; and Gogolak C. September 1972. In Situ Ge(Li) and Nal(TI) Gamma-Ray Spectrometry. HASL-258. U.S. Atomic Energy Commission, Health and Safety Laboratory Springfield, VA: National Technical Information Center, U.S. Department of Commerce.

DISTRIBUTION

E=09313

τ

L. K. Cohen (10)

NRC/Region 1

J. Kinneman (5)

DOE/OES

L. J. Deal (5)

DOE/NV

J. K. Magruder (1)

EG&G

Z. G. Burson, LVAO (2) J. F. Doyle, LVAO (1) S. A. Gunn, WAMD (1) H. A. Lamonds, SBO (1) R. E. Lounsbury, WAMD (4) J. R. Mueller, WAMD (2) L. G. Sasso, LVAO (1) T. P. Stuart, LVAO (1) W. J. Tipton, LVAO (1)

LIBRARIES

AMO (8) Las Vegas (1) Santa Barbara (1)