

ENERGY SYSTEMS GROUP
ENVIRONMENTAL MONITORING
AND
FACILITY EFFLUENT
ANNUAL REPORT
1978

BY
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APPROVED



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ABSTRACT

Environmental and facility effluent radioactivity monitoring at the Energy Systems Group (ESG) of Rockwell International (California operations) is performed by the Radiation and Nuclear Safety Group of the Health, Safety and Radiation Services Department. Soil, vegetation, and surface water are routinely sampled to a distance of 10 miles from ESG sites. Continuous ambient air sampling and also monitoring by thermoluminescent dosimetry are performed on-site for measuring airborne radioactivity concentrations and site ambient radiation levels. Radioactivity in effluents discharged to the atmosphere from ESG facilities is continuously sampled and monitored to ensure that levels released to unrestricted areas are within appropriate limits, and to identify processes which may require additional engineering safeguards to minimize radioactivity levels in such effluents. In addition, selected nonradioactive constituent concentrations in surface water discharged to unrestricted areas are determined. This report summarizes and discusses monitoring results for 1978.

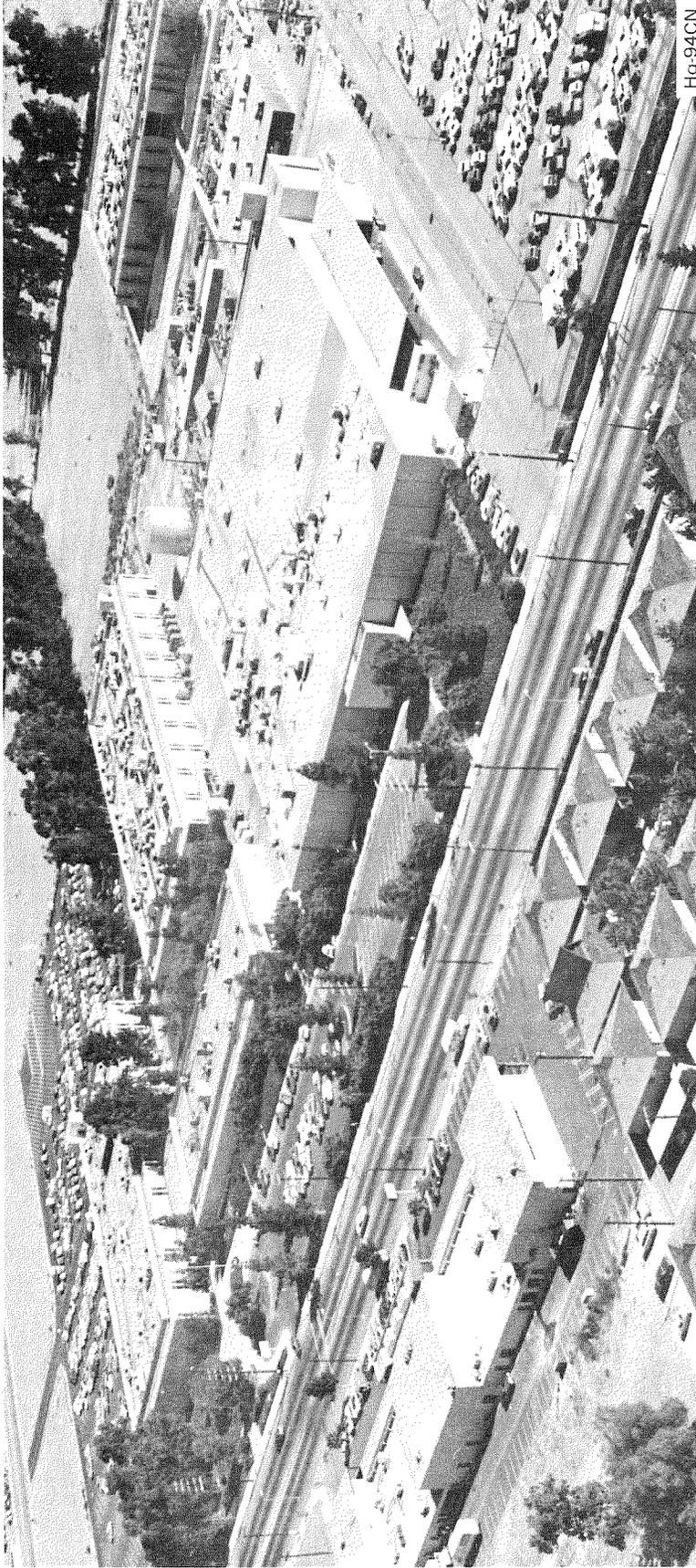
The random variations observed in the environmental monitoring data indicate that no local source of unnatural radioactive material exists in the environs. Additionally, the similarity between on-site and off-site results further indicates that the contribution to general environmental radioactivity due to operations at the ESG is essentially nonexistent.

The environmental radioactivity reported herein is attributed to natural sources and to fallout of radioactive material from foreign atmospheric testing of nuclear devices.

I. INTRODUCTION

The Energy Systems Group (ESG) of Rockwell International Corporation has been engaged in atomic energy research and development since 1946. The ESG is currently working on the design, development, fabrication, and testing of components and systems for central station power plants, on the fabrication of nuclear fuel for test and research reactors, and on the Decontamination and Disposition of Facilities (D&D) Program. Other programs include the development and fabrication of systems for stack gas SO₂ control, gasification and liquefaction of coal, and solid and liquid waste disposal.

The Group's administration, scientific research, and manufacturing facilities (Figure 1) are located in Canoga Park, California, approximately 23 miles northwest of downtown Los Angeles. The site is level, typical of the San Fernando Valley floor. Certain of the Group's nuclear programs, under licenses issued by the Nuclear Regulatory Commission (NRC) and the State of California, are conducted here. These include: (1) Building 001 containing uranium fuel production facilities, and (2) Building 004 containing analytical chemistry laboratories, and a gamma irradiation facility. The 290-acre Santa Susana Field Laboratories site (SSFL), Figure 2, is located in the Simi Hills of Ventura County, approximately 29 miles northwest of downtown Los Angeles. The SSFL site is situated in rugged terrain typical of mountain areas of recent geological age. The site may be described as an irregular plateau sprinkled with outcroppings above the more level patches and with peripheral eroded gullies. Elevations of the site vary from 1650 to 2250 ft above sea level. The surface mantle consists of sand and clay soil on sandstone. Both Department of Energy (DOE) and ESG owned facilities share this site shown in Figure 3. The SSFL also contain facilities in which nuclear operations licensed by NRC and the State, are conducted. The licensed facilities include: (1) the Rockwell International Hot Laboratory (RIHL), Building 020; (2) the Nuclear Materials Development Facility (NMDF), Building 055; (3) a neutron radiography facility containing the L-85 nuclear examination and research reactor, Building 093; and (4) several X-radiography inspection facilities. The location of these sites, in relation to nearby communities, is shown in Figure 4.



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Figure 1. Energy Systems Group – De Soto Site

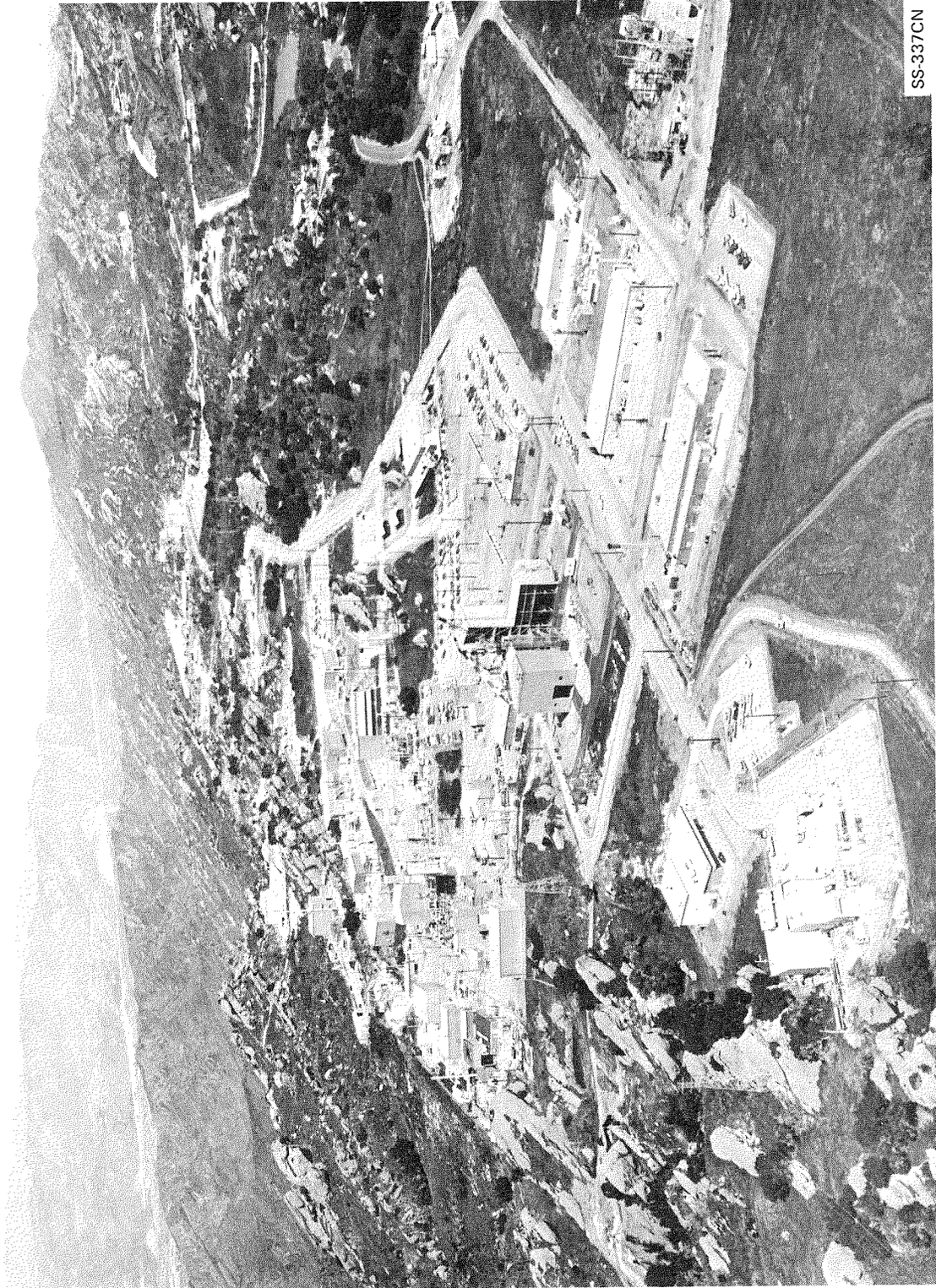
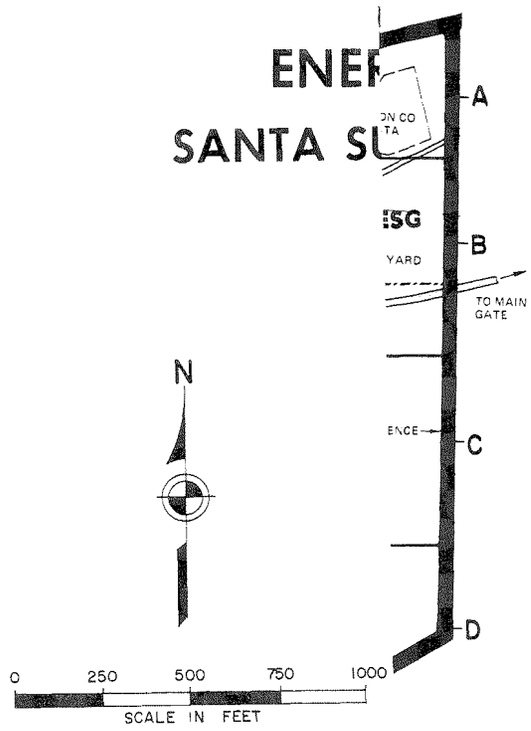


Figure 2. Energy Systems Group – Santa Susana Field Laboratories Site



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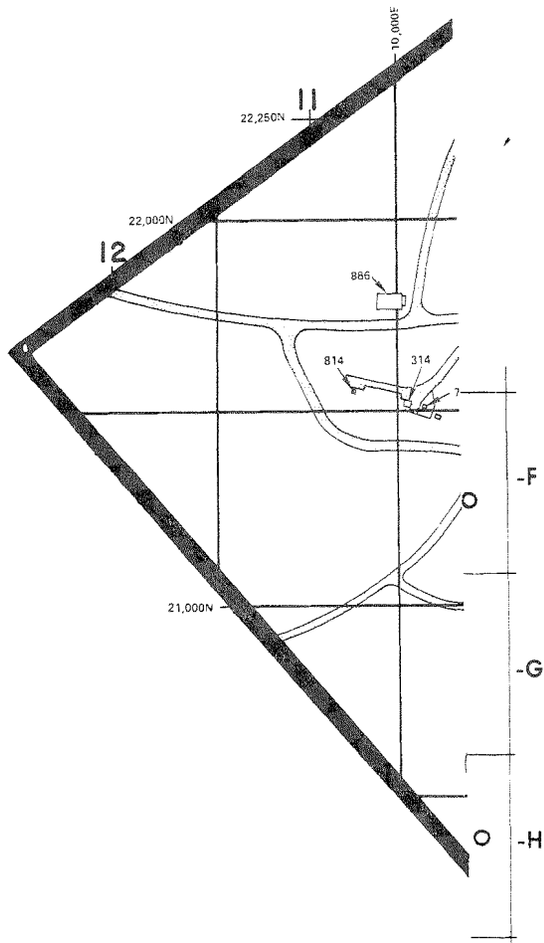


Figure 3. Map of Santa Susana
Field Laboratories Site
Facilities
(Sheet 1 of 2)

ESG-79-7

OWNED	BUILDING		DESCRIPTION	BUILDING		DESCRIPTION
	ZONE	NUMBER		ZONE	NUMBER	
ROCKWELL/GOVT.	4C	003	EXCESS EQUIPMENT STOP			
ROCKWELL	6D	005	ENVIRONMENTAL SYSTEM	9F	800	ELECTRICAL SUBSTATION
ROCKWELL	6D	006	SODIUM LABORATORY	7E	805	TIME CLOCK BUILDING
ROCKWELL	6E	007	SODIUM STORAGE	11G	814	LARGE LEAK INJECTOR DEVICE
ROCKWELL	6E	008	FLAMMABLE MATERIAL S	6E	816	RECOMBINER CANOPY
ROCKWELL	9G	009	ENGINEERING DEVELOPM	5D	836	TIME CLOCK BUILDING
GOVT.	7D	010	D&D	7H	854	TEST STRUCTURE
ROCKWELL	6F	011	MANUFACTURING SUPPOI	7H	863	HYDRAULIC TEST LOOP
GOVT.	7D	012	TOOL CRIB - ETEC OPERA	8G	873	HYDRAULIC TEST LABORATORY
GOVT.	7D	013	THERMAL TRANSIENT FA	7F	883	ELECTRICAL SUBSTATION
GOVT.	3C	014	SODIUM STORAGE BUILDI	10F	885	PISTON RANGE
ROCKWELL	7G	015	SUPPLEMENTARY STORA	11G	886	SODIUM DISPOSAL FACILITY
GOVT.	7D	019	ETEC CONSTRUCTION ST/	6D	924	ELECTRICAL SUBSTATION
ROCKWELL	8G	020	ENERGY SYSTEMS GROUP			BOWL AREA
GOVT.	6C	021	RADIOACTIVE WASTE, DE			
GOVT.	6C	022	RADIOACTIVE WASTE STC	H -13	306	PUMP HOUSE NO. 1
GOVT.	5D	023	LIQUID METALS CHEMIST	H -12	307	PUMP HOUSE NO. 2
GOVT.	6D	024	DEVELOPMENT TEST BUIL	G -12	308	CONTROL CENTER
GOVT.	6D	025	ETEC INSTRUMENTATION	G -13	320	POWER CONTROL
GOVT.	6E	026	SMALL COMPONENT TEST	H -12	334	V.T.S. - 3 OPERATIONS AND WORKSHOP
GOVT.	6D	027	ETEC QUALITY ASSURAN	H -12	391	V.T.S. - 3 WORKSHOP
GOVT.	6D	028	LMFBR FUEL SAFETY	H -12	393	PLUME STUDY BLDG.
GOVT.	3D	029	SODIUM STORAGE	G -12	401	INSTRUMENT AND WORKSHOP
GOVT.	4C	030	SITE PURCHASING OFFICE	H -12	405	STEAM PLANT BLDG.
GOVT.	5D	032	ETEC GENERAL TEST	G -13	416	ELECTRICAL BLDG.
GOVT.	5C	034	R/A WASTE OFFICE BUILD	G -13	437	PRE-TEST - BOWL
GOVT.	5D	036	ETEC OPERATIONS	F -11	964	SEWAGE TREATMENT PLANT BLDG.
GOVT.	7E	038	ETEC ADMINISTRATION			
GOVT.	7E	039	OFFICE BUILDING			
ROCKWELL	2C	040	FACILITIES AND INDUSTR			
GOVT.	5C	041	STORAGE BUILDING			
GOVT.	5D	042	LMFBR TEST			
GOVT.	6C	044	RMDF CLEAN SHOP			
ROCKWELL	4C	046	MATERIAL OFFICE ANNE			
ROCKWELL	6E	048	PDU INSTRUMENTATION E			
GOVT.	5D	049	PDV CONTROL ROOM			
ROCKWELL	8G	055	PLUTONIUM FACILITY			
GOVT.	7E	057	ETEC LABORATORY			
GOVT.	8D	059	LARGE LEAK TEST RIG			
GOVT.	7E	062	ETEC INSTRUMENTATION			
GOVT.	3D	064	SOURCE AND SPECIAL NU			
GOVT.	7E	065	ETEC CHEMISTRY LABOR			
GOVT.	7E	066	INSTRUMENTATION REPA			
ROCKWELL	4C	074	STORAGE BUILDING			
GOVT.	6C	075	CONTAMINATED EQUIPME			
ROCKWELL	4C	083	CONTROL BUILDING - NEI			
ROCKWELL	4C	093	NEUTRON RADIOGRAPHY			
ROCKWELL	9F	100	ADVANCED FUELS LABOR			
ROCKWELL	5B	114	DECON TRAILER			
ROCKWELL	5C	133	SODIUM BURN FACILITY			
ROCKWELL	4B	143	SODIUM REACTOR EXPERI			
GOVT.	8G	155	CONTROL CENTER			
ROCKWELL	4B	163	BOX SHOP			
ROCKWELL	6F	171	X-RAY BUILDING			
ROCKWELL	6F	172	X-RAY BUILDING			
ROCKWELL	8H	173	GAMMAGRAPH BUILDING			
ROCKWELL	4B	183	FIRE PUMP BUILDING - D8			
ROCKWELL	10G	314	LARGE LEAK INJECTOR DE			
GOVT.	2B	320	FUEL OIL CONTROL BUILD			
ROCKWELL	3C	333	TIME CLOCK BUILDING			
ROCKWELL	7G	343	TIME CLOCK BUILDING			
ROCKWELL	7H	353	RESEARCH AND DEVELOPM			
GOVT.	6E	354	CONTROL ELEMENT TEST			
GOVT.	7D	355	SCTI SUPPORT BUILDING			
GOVT.	7D	356	SODIUM COMPONENT TEST			
GOVT.	7D	357	ETEC PUMP BEARING TEST			
GOVT.	6E	358	SCTI SUPPORT BUILDING			
GOVT.	7E	359	COMPRESSOR BUILDING			
GOVT.	7E	360	CHEMICAL STORAGE BUIL			
ROCKWELL	8H	363	RESEARCH AND DEVELOPM			
ROCKWELL	7G	373	DEVELOPMENT TEST BUIL			
GOVT.	7G	374	TEST LOOP ENCLOSURE			
ROCKWELL	8G	375	CONTROL SHELTER BUILDI			

Figure 3. Map of Santa Susana
Field Laboratories Site
Facilities
(Sheet 2 of 2)

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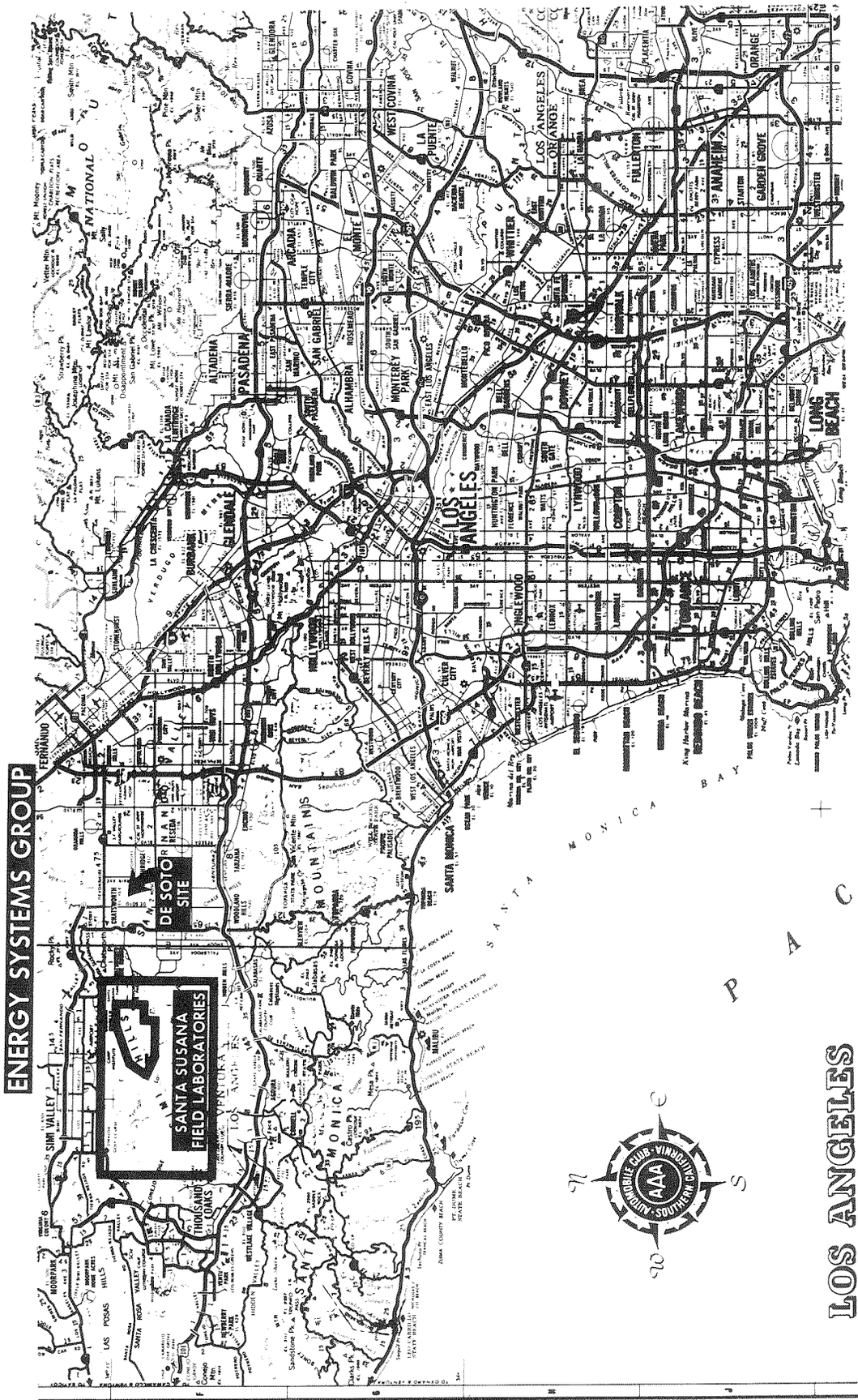


Figure 4. Map of General Los Angeles Area
 (Reproduced by permission of Automobile Club of Southern California)

Also included within the SSFL site is an 82-acre Government-optioned area where DOE-contract activities are conducted, primarily by the non-nuclear Energy Technology Engineering Center (ETEC). The major operational nuclear installation within the optioned area is the Radioactive Material Disposal Facility (RMDF), Buildings 021 and 022. This facility is used for processing wastes generated as a result of the D&D Program, begun in 1975. Several deactivated nuclear reactor and support facilities, all within the optioned area, are affected by the D&D Program. Currently involved are several facilities used for Systems for Nuclear Auxiliary Power (SNAP) reactor test operations, Buildings 010, 024, and 059, and the SRE, Building 143. There is no fissile material located at any of these facilities.

Licensed programs conducted during 1978 included: (1) commercial operation of the L-85 reactor for central station power plant operator training and for neutron radiography inspection of precision forgings, castings, and electronic and explosive devices for manufacturing defects; (2) the operation of the RIHL for nuclear reactor fuel and system component examination and the fabrication of sealed radiation sources; and (3) the operation of nuclear fuel manufacturing facilities for the production of experimental and test reactor fuel involving enriched uranium, and development of processes for fabrication of advanced fuels.

The basic policy for control of radiological and toxicological hazards at ESG requires that through engineering controls adequate containment of such materials be provided, and through rigid operational controls, that facility effluent releases and external radiation levels are reduced to a minimum. The environmental monitoring program provides a measure of the effectiveness of the Group safety procedures and of the engineering safeguards incorporated into facility designs. Specific radionuclides in facility effluent or environmental samples, although not routinely identified due to the extremely low radioactivity levels normally detected, may be identified by analytical or radiochemistry techniques if significantly increased radioactivity levels were observed.

In addition to environmental monitoring, work area air and atmospherically discharged effluents are continuously monitored or sampled, as appropriate. This provides a direct measure of the effectiveness of engineering controls and allows remedial action to be taken before a significant release of hazardous material can occur.

Environmental sampling stations that are located within the boundaries of ESG sites are referred to as "on-site" stations; those located within a 10-mile radius of the sites are referred to as "off-site" stations. The on-site environs of the De Soto and SSFL sites are sampled monthly to determine the concentration of radioactivity in typical surface soil, vegetation, and water. Soil is also sampled on site semiannually, for plutonium analysis. Similar off-site environmental samples, except for plutonium analysis, are obtained quarterly. Continuous on-site and off-site ambient air sampling provides information concerning long-lived airborne particulate radioactivity. A site ambient radiation monitoring program, utilizing thermoluminescent dosimetry (TLD), begun in 1971, was expanded during 1978.

Nonradioactive wastes released to unrestricted areas are limited to liquids released to sanitary sewage systems and to surface water drainage systems. No intentional releases of any liquid pollutants are made to unrestricted areas. Liquid wastes generated at the De Soto site are discharged into the city sewage system. This effluent is sampled for determination of radioactivity. Sanitary sewage from all DOE and ESG facilities at the SSFL site is treated at an on-site sewage plant. The plant effluent drains into a retention pond, located on the adjoining Rocketdyne Division site. The surface water drainage system of the SSFL is composed of catch ponds and open drainage ditches leading to the Rocketdyne retention pond. This pond also receives the ESG site sewage plant effluent. Water from the pond may be reclaimed as industrial process water, or it may be released off site into Bell Creek, a tributary of the Los Angeles River. The pond was also monitored at discharge for tritium, and for nonradioactive pollutants by Rocketdyne Division as required by discharge permits issued to Rocketdyne by the California Regional Water Quality Control Board.

This report summarizes environmental monitoring results for 1978. A comparison of 1978 radioactivity results with previous years appears in Appendix A.

TABLE 1-A
SOIL RADIOACTIVITY DATA — 1978

Area	Activity	No. Samples	Gross Radioactivity ($\mu\text{Ci/g}$)	
			Annual Average Value (95% Confidence Level)	Maximum Observed Value*
On Site	α	144	$(6.3 \pm 1.5) 10^{-7}$	9.9×10^{-7}
	β	144	$(2.4 \pm 0.09) 10^{-5}$	4.8×10^{-5}
Off Site	α	48	$(5.1 \pm 1.4) 10^{-7}$	10.0×10^{-7}
	β	48	$(2.4 \pm 0.09) 10^{-5}$	3.4×10^{-5}

*Maximum value observed for single sample

TABLE 1-B
SOIL PLUTONIUM RADIOACTIVITY DATA — 1978

Sample Location	June 8, 1978 Survey Results		December 13, 1978 Survey Results	
	Pu^{238} ($\mu\text{Ci/g}$)	$\text{Pu}^{239} + \text{Pu}^{240}$ ($\mu\text{Ci/g}$)	Pu^{238} ($\mu\text{Ci/g}$)	$\text{Pu}^{239} + \text{Pu}^{240}$ ($\mu\text{Ci/g}$)
S-56	$(-2.90 \pm 2.3) 10^{-9}$	$(1.04 \pm 2.3) 10^{-9}$	*	$(9.05 \pm 3.6) 10^{-9}$
S-57	$(-2.13 \pm 2.2) 10^{-9}$	$(1.14 \pm 2.2) 10^{-9}$	*	$(6.21 \pm 3.5) 10^{-9}$
S-58	$(-3.59 \pm 2.3) 10^{-9}$	$(1.46 \pm 3.0) 10^{-9}$	*	$(5.54 \pm 2.8) 10^{-9}$
S-59	$(-3.61 \pm 2.3) 10^{-9}$	$(3.10 \pm 3.4) 10^{-9}$	$(1.22 \pm 2.4) 10^{-9}$	$(5.98 \pm 2.8) 10^{-9}$
S-60	$(-2.17 \pm 3.2) 10^{-9}$	$(3.35 \pm 3.9) 10^{-9}$	$(-3.66 \pm 1.7) 10^{-9}$	$(8.28 \pm 4.4) 10^{-9}$

Note: Minus (-) indicates sample value less than reagent blank.

*Result significantly less than the minimum detection level.

II. ENVIRONMENTAL MONITORING SUMMARY RESULTS

A. RADIOACTIVE MATERIALS — 1978

The sampling and analytic methods used in the environmental monitoring program for radioactive materials are described in Section III.

The average radioactivity concentrations in local soil, vegetation, surface water, and in ambient air for 1978 are presented in Tables 1 through 5. In calculating the averaged concentration value for the tables, those individual samples having radioactivity levels less than their minimum detection levels (MDL) are assumed to have a concentration equal to the MDL. This method of data averaging, required by DOE Manual Chapter 0513, affords a significant level of conservatism in the data, as evident in the tables, in that most radioactivity concentrations are reported as "less than" (<) values. Thus, for measurements in which some apparent radioactivity concentrations are below the MDL, the true averaged value is actually somewhat less than the value reported.

The maximum level of radioactivity detected for a single sample is reported because of its significance in indicating the existence of a major episode or area-wide location of radioactive material deposition. Except for ambient air radioactivity, none of the maximum observed values, which occurred randomly during the year, show a great increase over the average values beyond natural variability. The air sample data reflect March and December atmospheric nuclear device detonations which resulted in marked but transient increases in local airborne radioactivity levels.

The results reported in Tables 1-A and 2 show no significant difference between on-site and off-site samples. Table 1-B shows no significant variations in soil plutonium concentrations for the 1978 sample sets. The detected activity is due to a variety of naturally occurring radionuclides, and to radioactive fallout resulting from dispersal of nuclear weapons materials and fission products by atmospheric testing. Naturally occurring radionuclides include Be^7 , K^{40} , Rb^{87} , Sm^{147} , and the uranium and thorium series (including the inert gas radon and its radioactive daughters). Radioactivity from fallout consists primarily of the fission products Sr^{90} - Y^{90} , Cs^{137} , and Pm^{147} , and also U^{235} and Pu^{239} .

TABLE 2
VEGETATION RADIOACTIVITY DATA — 1978

Area	Activity	No. Samples	Gross Radioactivity ($\mu\text{Ci/g}$)		
			Dry Weight Annual Average Value	Ash	
				Annual Average Value (95% Confidence Level)	Maximum Observed Value*
On Site	α	144	$(<5.8 \pm 4.1) 10^{-8}$	$(<2.4 \pm 1.7) 10^{-7}$	9.6×10^{-7}
	β	144	$(2.6 \pm 0.05) 10^{-5}$	$(1.66 \pm 0.03) 10^{-4}$	3.19×10^{-4}
Off Site	α	48	$(<6.6 \pm 4.4) 10^{-8}$	$(<2.4 \pm 1.6) 10^{-7}$	6.6×10^{-7}
	β	48	$(3.3 \pm 0.07) 10^{-5}$	$(1.43 \pm 0.03) 10^{-4}$	2.41×10^{-4}

*Maximum value observed for single sample

Domestic water used at the SSFL site is obtained from Ventura County Water District No. 17, which also supplies nearby communities, and is distributed on site by the same piping system previously used when all facility process water was obtained from on-site wells. Conversion to the domestic water supply was completed during 1969. Two on-site water wells were operated during 1978 to reduce consumption of Ventura County domestic water as a conservation measure due to local drought conditions. The well water proportion in the blend averaged about 38% for the year for a total well water consumption of approximately 6.3×10^6 gal. Pressure for the water system is provided by elevated storage tanks.

Water from the system is sampled monthly at two widely separated SSFL site locations. The average domestic water radioactivity concentration is presented in Table 3.

As discussed earlier, surface waters discharged from SSFL facilities and the sewage plant effluent drain southward into a retention pond on Rocketdyne property. When full, the pond may be drained into Bell Creek, a tributary of

TABLE 3
SSFL SITE — DOMESTIC WATER RADIOACTIVITY DATA — 1978

Area	Activity	No. Samples	Gross Radioactivity ($\mu\text{Ci}/\text{m}\ell$)	
			Average Value (95% Confidence Level)	Maximum* Observed Value
ESG-SSFL	α	24	$(<2.6 \pm 2.8) 10^{-10}$	4.4×10^{-10}
	β	24	$(3.0 \pm 0.8) 10^{-9}$	3.6×10^{-9}

*Maximum value observed for single sample

the Los Angeles River in the San Fernando Valley, Los Angeles County. Pursuant to the requirements of Los Angeles Regional Water Quality Control Board Resolution 66-49 of September 21, 1966, a sampling station for evaluating environmental radioactivity in Bell Canyon was established in 1966. It is located approximately 2.5 miles downstream from the southern Rockwell International Corporation boundary. Samples, obtained and analyzed monthly, include stream bed mud, vegetation, and water. Average radioactivity concentrations in Rocketdyne and Bell Creek samples are presented in Table 4.

Comparison of the radioactivity concentrations in water from the ponds and from Bell Creek with that of the domestic water supply shows no significant variation in either alpha or beta activity.

The SSFL site surface water and the ambient air radioactivity concentration Guide values selected for each site are the most restrictive limits for those radionuclides currently in use at ESG facilities. The identity of all such radionuclides is known, irrespective of concentration. Accordingly, for SSFL site surface water, the Guide value of $5 \times 10^{-6} \mu\text{Ci}/\text{m}\ell\alpha$ and $3 \times 10^{-7} \mu\text{Ci}/\text{m}\ell\beta$, for Pu^{239} and for Sr^{90} , respectively, is appropriate. The correspondingly most restrictive Guide value for De Soto site wastewater radioactivity discharged to the sanitary sewage system, a controlled area, is $8 \times 10^{-4} \mu\text{Ci}/\text{m}\ell\alpha$ and $1 \times 10^{-3} \mu\text{Ci}/\text{m}\ell\beta$ for U^{235} and Co^{60} , respectively. These values are established in 10 CFR 20, California Administrative Code Title 17, and DOE Manual Chapter 0524.

TABLE 4
 BELL CREEK AND ROCKETDYNE SITE RETENTION POND
 RADIOACTIVITY DATA — 1978

Area	Activity	No. Samples	Gross Radioactivity Concentration		
			Average Value (95% Confidence Level)	Maximum* Observed Value	% of Guide†
Bell Creek Mud No. 54 ($\mu\text{Ci/g}$)	α	12	$(4.2 \pm 1.3) 10^{-7}$	7.4×10^{-7}	NA
	β	12	$(2.3 \pm 0.08) 10^{-5}$	2.7×10^{-5}	NA
Pond R-2A Mud No. 55 ($\mu\text{Ci/g}$)	α	12	$(6.5 \pm 1.6) 10^{-7}$	9.0×10^{-7}	NA
	β	12	$(2.5 \pm 0.09) 10^{-5}$	2.8×10^{-5}	NA
Bell Creek Vegetation No. 54 ($\mu\text{Ci/g}$ ash)	α	12	$(<2.6 \pm 1.7) 10^{-7}$	5.1×10^{-7}	NA
	β	12	$(1.56 \pm 0.03) 10^{-4}$	2.14×10^{-4}	NA
Bell Creek Vegetation No. 54 ($\mu\text{Ci/g}$) dry weight)	α	12	$(<4.6 \pm 3.8) 10^{-8}$	1.0×10^{-7}	NA
	β	12	$(2.8 \pm 0.05) 10^{-5}$	4.4×10^{-5}	NA
Bell Creek Water No. 16 ($\mu\text{Ci/ml}$)	α	12	$(<2.4 \pm 2.8) 10^{-10}$	$<2.4 \times 10^{-10}$	<0.005
	β	12	$(2.5 \pm 0.8) 10^{-9}$	3.5×10^{-9}	0.8
Pond Water No. 6 ($\mu\text{Ci/ml}$)	α	12	$(<2.5 \pm 2.8) 10^{-10}$	3.5×10^{-10}	<0.005
	β	12	$(4.3 \pm 0.8) 10^{-9}$	7.0×10^{-9}	1.4
SSFL Pond R-2A Water No. 12 ($\mu\text{Ci/ml}$)	α	12	$(<2.5 \pm 2.8) 10^{-10}$	2.7×10^{-10}	<0.005
	β	12	$(4.6 \pm 0.8) 10^{-9}$	6.3×10^{-9}	1.5

*Maximum value observed for single sample.

†Guide: $5 \times 10^{-6} \mu\text{Ci/ml}$ α , $3 \times 10^{-7} \mu\text{Ci/ml}$ β ; 10 CFR 20 Appendix B, CAC 17, DOE Manual Chapter 0524.

NA — not applicable, no Guide value having been established.

The Guide value of 6×10^{-14} $\mu\text{Ci}/\text{m}\ell\alpha$ for SSFL site ambient air radioactivity is due to work with unencapsulated plutonium. The Guide value of 3×10^{-11} $\mu\text{Ci}/\text{m}\ell\beta$ for Sr^{90} is due to the presence of fission products in irradiated nuclear fuel at the SSFL site. The Guide value of 3×10^{-12} $\mu\text{Ci}/\text{m}\ell\alpha$ for De Soto site ambient air radioactivity is due to work with unencapsulated uranium (including depleted uranium). The Guide value of 3×10^{-10} $\mu\text{Ci}/\text{m}\ell\beta$ for Co^{60} for ambient air radioactivity is appropriate since it is the most restrictive limit for beta-emitting radionuclides present at the De Soto site. Guide value percentages are not presented for soil or vegetation data since no concentration Guide values have been established.

Ambient air sampling for long-lived particulate alpha and beta radioactivity is performed continuously with automatic sequential samplers at both the De Soto and SSFL sites. Air is drawn through Type HV-70 filter media which are analyzed for long-lived radioactivity, after a minimum 120-h decay period that eliminates the radon particulate daughters. The average concentrations of ambient air alpha and beta radioactivity are presented separately in Table 5.

Radioactivity levels observed in environmental samples for 1978, reported in Tables 1 through 5, compare closely with levels reported for recent years. Local environmental radioactivity levels, which result primarily from beta-emitting radionuclides and had shown the effect of fallout during past extensive atmospheric testing of nuclear devices, have decreased, and have been generally constant during the past several years. The effects of continuing, although infrequent, foreign atmospheric nuclear tests continue to be occasionally observable in daily ambient air radioactivity levels; however, the long-term effects on surface sample radioactivity levels are not discernible. The continued relative constancy in current levels of environmental radioactivity is due primarily to the dominance of naturally occurring radionuclides in the environment and to the longer-life fission product radioactivity from fallout.

Site ambient radiation monitoring is performed with thermoluminescent dosimeters. Each dosimeter set contains two calcium fluoride ($\text{CaF}_2:\text{Mn}$) low background, bulb-type chip dosimeters. The dosimeter sets are placed at selected locations (Figures 6 and 7) on or near the perimeters of the De Soto and SSFL sites. Each dosimeter, sealed in a light-proof energy compensation shield,

TABLE 5
 AMBIENT AIR RADIOACTIVITY DATA — 1978

Site Location	Activity	No. Samples	Average Value (95% Confidence Level)	Maximum* Observed Value (daily)	% of Guidet
De Soto On Site ($\mu\text{Ci}/\text{m}\ell$)	αs	713	$(<8.4 \pm 8.1) 10^{-15}$	9.5×10^{-14}	<0.28
	β^{**}	713	$(<9.1 \pm 1.7) 10^{-14}$	1.4×10^{-12}	<0.030
SSFL On Site ($\mu\text{Ci}/\text{m}\ell$)	αs	1724	$(<7.2 \pm 7.9) 10^{-15}$	2.1×10^{-14}	<12.0
	β^{**}	1724	$(<8.8 \pm 1.7) 10^{-14}$	1.5×10^{-12}	<0.29
SSFL Sewage Treatment Plant Off Site ($\mu\text{Ci}/\text{m}\ell$)	αs	327	$(<7.3 \pm 7.3) 10^{-15}$	4.4×10^{-14}	<12.2
	β^{**}		$(<8.4 \pm 1.6) 10^{-14}$	1.2×10^{-12}	<0.28
SSFL Control Center Off Site ($\mu\text{Ci}/\text{m}\ell$)	αs	351	$(<7.1 \pm 7.3) 10^{-15}$	3.4×10^{-14}	<11.8
	β^{**}		$(<8.9 \pm 1.6) 10^{-14}$	1.3×10^{-12}	<0.30

*Maximum value observed for single sample.

†Guide: De Soto site, $3 \times 10^{-12} \mu\text{Ci}/\text{m}\ell\alpha$, $3 \times 10^{-10} \mu\text{Ci}/\text{m}\ell\beta$; 10 CFR 20 Appendix B, SSFL site, $6 \times 10^{-14} \mu\text{Ci}/\text{m}\ell\alpha$, $3 \times 10^{-11} \mu\text{Ci}/\text{m}\ell\beta$; 10 CFR 20 Appendix B, CAC 17, and DOE Manual Chapter 0524

§MDL = $6.0 \times 10^{-15} \mu\text{Ci}/\text{m}\ell$ — Individual daily samples with activity levels of 0 to $6.0 \times 10^{-15} \mu\text{Ci}/\text{m}\ell$ are recorded and averaged as $6.0 \times 10^{-15} \mu\text{Ci}/\text{m}\ell$.

**MDL = $1.2 \times 10^{-14} \mu\text{Ci}/\text{m}\ell$ — Individual daily samples with activity levels of 0 to $1.2 \times 10^{-14} \mu\text{Ci}/\text{m}\ell$ are recorded and averaged as $1.2 \times 10^{-14} \mu\text{Ci}/\text{m}\ell$. Indicated average values are upper limits, since some data were below the minimum detection levels.

is installed in a polyethylene container which is mounted ~1 meter above ground at each location. The dosimeters are exchanged and evaluated quarterly. There were 13 on-site TLD monitoring locations used during the year. Three additional dosimeter sets, located at locations up to 10 miles from the ESG sites, are similarly evaluated to determine the local area off-site ambient radiation level, which averaged 0.014 mrem/h for 1978. The average radiation dose rate and equivalent annual dose monitored at each dosimeter location are presented in Table 6.

TABLE 6
 DE SOTO AND SSFL SITES — AMBIENT RADIATION
 DOSIMETRY DATA — 1978

TCD	Average Dose Rate (mrem/h)	Equivalent Annual Dose (mrem)
1 De Soto	0.016	140
2 De Soto	0.015	131
3 De Soto	0.014	123
4 De Soto	0.015	131
5 De Soto	0.015	131
6 De Soto	0.013*	114
7 De Soto	0.014	123
1 SSFL	0.017	149
2 SSFL	0.017	149
3 SSFL	0.017	149
4 SSFL	0.017	149
5 SSFL	0.011 [†]	96
6 SSFL	0.016	140
1 Off-Site Control	0.014	123
2 Off-Site Control	0.014	123
3 Off-Site Control	0.015	131

*Excludes second quarter data due to missing dosimeter.

[†]Dosimeter partially shielded by water tank. Relocated to 360° exposure field beginning third quarter. Unshielded annual average dose rate is 0.014 mrem/h, equivalent to an annual dose of 123 mrem.

The table shows that radiation dose rates and equivalent annual doses monitored on site are nearly identical to levels monitored at three widely separated off-site locations. These data include the natural background radiation component which exists as a consequence of cosmic radiation, radionuclides in the soil, and radon and thoron in the atmosphere, in addition to radioactive fallout from nuclear weapons tests. Locally, this is approximately 125 mrem/year. The small variability observed in the data is attributed to differences in elevation

TABLE 7
 NONRADIOACTIVE CONSTITUENTS AND TRITIUM IN WASTEWATER DISCHARGED TO UNRESTRICTED AREAS - 1978
 (Analysis Results for Wastewater Discharged from Pond R-2A to Bell Creek on Date Indicated - Sample Station W-12)

Constituents	January 5*		January 16*		February 6*		February 13*		March 1*	
	Result	% of Guide								
Total Dissolved Solids (mg/l)	225	23.7	217	22.8	272	28.6	147	15.5	107	11.3
Chloride (mg/l)	25	16.7	25	16.7	28	18.7	11	7.3	4	2.7
Sulfate (mg/l)	56	18.7	50	16.7	63	21.0	28	9.3	14	4.7
Suspended Solids [§] (mg/l)	135	90.0	40	28.7	53	35.3	58	38.7	200	133.3
Settleable Solids [§] (ml/l)	0.2	66.7	<0.1	<33.3	<0.1	<33.3	0.3	100.0	0.6	200.0
BOD (mg/l)	65	108.3	3	5.0	9	15.0	1	1.7	6	10.0
Oil and Grease (mg/l)	2	13.3	2	13.3	<2	<13.3	<1	<6.7	<1	<6.7
Turbidity (TU)	85	-	32	-	39	-	53	-	165	-
Chromium (mg/l)	0.011	110.0	0.003	30.0	0.006	60.0	0.006	60.0	0.024	240.0
Fluoride (mg/l)	0.4	40.0	0.3	30.0	0.3	30.0	0.3	30.0	0.4	40.0
Boron (mg/l)	0.1	10.0	0.1	10.0	0.1	10.0	0.2	20.0	<0.2	<20.0
Residual Chlorine (mg/l)	<0.05	<50.0	0.02	20.0	<0.04	<40.0	<0.04	<40.0	<0.04	<40.0
Fecal Coliform (MPN/100 ml)	<2.2	<9.5	<2.2	<9.5	<2.2	<9.5	<2.2	<9.5	16.0+	69.5+
Surfactants (mg/l)	<0.01	<2.0	0.006	1.2	0.01	2.0	<0.01	<2.0	<0.01	<2.0
pH		7.5		7.6		7.3		7.6		7.6
Tritium [†] (uCi/cc)	<1.1 x 10 ⁻⁵	<0.37								
Rainfall (in.)	2		4		7		7		8.75	
Estimated Rainfall Runoff (gal)	4 x 10 ⁷		1.2 x 10 ⁸		6 x 10 ⁷		2.5 x 10 ⁸		2.1 x 10 ⁸	
Release Volume (gal)	4 x 10 ⁵		6 x 10 ⁵		6 x 10 ⁵		9 x 10 ⁵		6.5 x 10 ⁵	
Constituents	March 10*		March 22*		March 30*		April 7*		April 7*	
	Result	% of Guide								
Total Dissolved Solids (mg/l)	473	49.8	326	34.3	429	45.2	384	40.4	254	26.7
Chloride (mg/l)	50	33.3	76	50.7	61	40.7	44	29.3	26	17.3
Sulfate (mg/l)	93	31.0	55	18.3	98	32.7	62	20.7	53	17.7
Suspended Solids [§] (mg/l)	25	16.7	68	45.3	13	8.7	44	29.3	100	66.7
Settleable Solids [§] (ml/l)	0.1	33.3	<0.1	<33.3	<0.1	<33.3	0.7	233.3	0.5	166.6
BOD (mg/l)	5	8.3	3	5.0	7	11.7	3.6	6.0	12	20.0
Oil and Grease (mg/l)	<1	<6.7	<1	<6.7	<1	<6.7	<1	<6.7	<1	<6.7
Turbidity (TU)	14	-	55	-	6	-	38	-	84	-
Chromium (mg/l)	0.039	390.0	0.007	70.0	0.002	20.0	0.006	60.0	0.005	50.0
Fluoride (mg/l)	4.9	490.0	1.0	100.0	1.1	110.0	0.5	50.0	0.8	80.0
Boron (mg/l)	0.2	20.0	<0.2	<20.0	0.3	30.0	0.2	20.0	<0.2	<20.0
Residual Chlorine (mg/l)	0.1	100.0	<0.04	<40.0	<0.04	<40.0	0.04	40.0	<0.04	<40.0
Fecal Coliform (MPN/100 ml)	NA	-	2.2	9.5	16	69.5	<2.2	<9.5	<2.2	<9.5
Surfactants (mg/l)	0.02	4.0	0.03	6.0	0.04	8.0	0.03	6.0	<0.01	<2.0
pH		7.8		7.9		8.0		7.8		8.0
Tritium [†] (uCi/cc)	<1.1 x 10 ⁻⁵	<0.37								
Rainfall (in.)	0.87		1.33		0.82		0.65		1.23	
Estimated Rainfall Runoff (gal)	2.1 x 10 ⁷		3.2 x 10 ⁷		2 x 10 ⁷		1.4 x 10 ⁷		3 x 10 ⁷	
Release Volume (gal)	5.5 x 10 ⁵		5 x 10 ⁵		5 x 10 ⁵		5 x 10 ⁵		7 x 10 ⁵	
Constituents	September 5*		November 13*		November 22*		December 18*			
	Result	% of Guide								
Total Dissolved Solids (mg/l)	438	46.1	548	57.7	309	32.5	403	42.4		
Chloride (mg/l)	84	56.0	87	58.0	30	20.0	44	29.3		
Sulfate (mg/l)	115	38.3	183	61.0	80	26.7	89	29.7		
Suspended Solids [§] (mg/l)	16	10.7	18	12.0	67	44.7	40	26.7		
Settleable Solids [§] (ml/l)	0.3	100.0	<0.3	<100.0	<0.1	<33.3	<0.1	<33.3		
BOD (mg/l)	5	8.3	5	8.3	14	23.3	6	10.0		
Oil and Grease (mg/l)	<1	<6.7	<1	<6.7	<1	<6.7	<1	<6.7		
Turbidity (TU)	7	-	10	-	62	-	34	-		
Chromium (mg/l)	0.003	30.0	0.005	50.0	0.009	90.0	0.008	80.0		
Fluoride (mg/l)	1.0	100.0	0.8	80.0	0.5	50.0	0.5	50.0		
Boron (mg/l)	0.2	20.0	0.3	30.0	0.1	10.0	<0.2	<20.0		
Residual Chlorine (mg/l)	<0.02	<20.0	<0.04	<40.0	<0.04	<40.0	<0.04	<40.0		
Fecal Coliform (MPN/100 ml)	<2.2	<9.5	<2.2	<9.5	<2.2	<9.5	9.2	40.0		
Surfactants (mg/l)	0.12	24.0	0.07	14.0	0.06	12.0	0.13	26.0		
pH		8.2		7.8		8.5		8.1		
Tritium [†] (uCi/cc)	<1.1 x 10 ⁻⁵	<0.37								
Rainfall (in.)	1.0		0.96		1.49		1.06			
Estimated Rainfall Runoff (gal)	3 x 10 ⁷		2.3 x 10 ⁷		3 x 10 ⁷		2.5 x 10 ⁷			
Release Volume (gal)	1.1 x 10 ⁶		2.9 x 10 ⁶		7.4 x 10 ⁵		1.6 x 10 ⁶			

NA = Not Available; analysis not requested or not performed.
 * = Rainfall related discharge.
 † = Tritium minimum detection limit: (1.1 ± 1.1) 10⁻⁵ uCi/ml.
 § = Not applicable to discharges containing rainfall runoff during or immediately after periods of rainfall.

and geologic conditions at the various dosimeter locations. Since the data for the on-site and off-site locations are nearly identical, no measurable radiation dose to the general population or to individuals in uncontrolled areas resulted from ESG operations.

B. NONRADIOACTIVE MATERIALS — 1978

Processed wastewater and most collected surface runoff discharged from the SSFL site drains to Retention Pond R-2A, operated by Rocketdyne. Water samples are taken from the pond and analyzed for various constituents, as required by the Regional Water Quality Control Board, and for tritium as required by NRC License - SNM-21, for each discharge to Bell Canyon. The discharges are normally required only as a result of excessive rainfall run-off. During such releases, the NPDES concentration limits for turbidity, and suspended and settleable solids do not apply. The results of analyses for each discharge for 1978, all of which were rainfall-related discharges, are presented in Table 7.

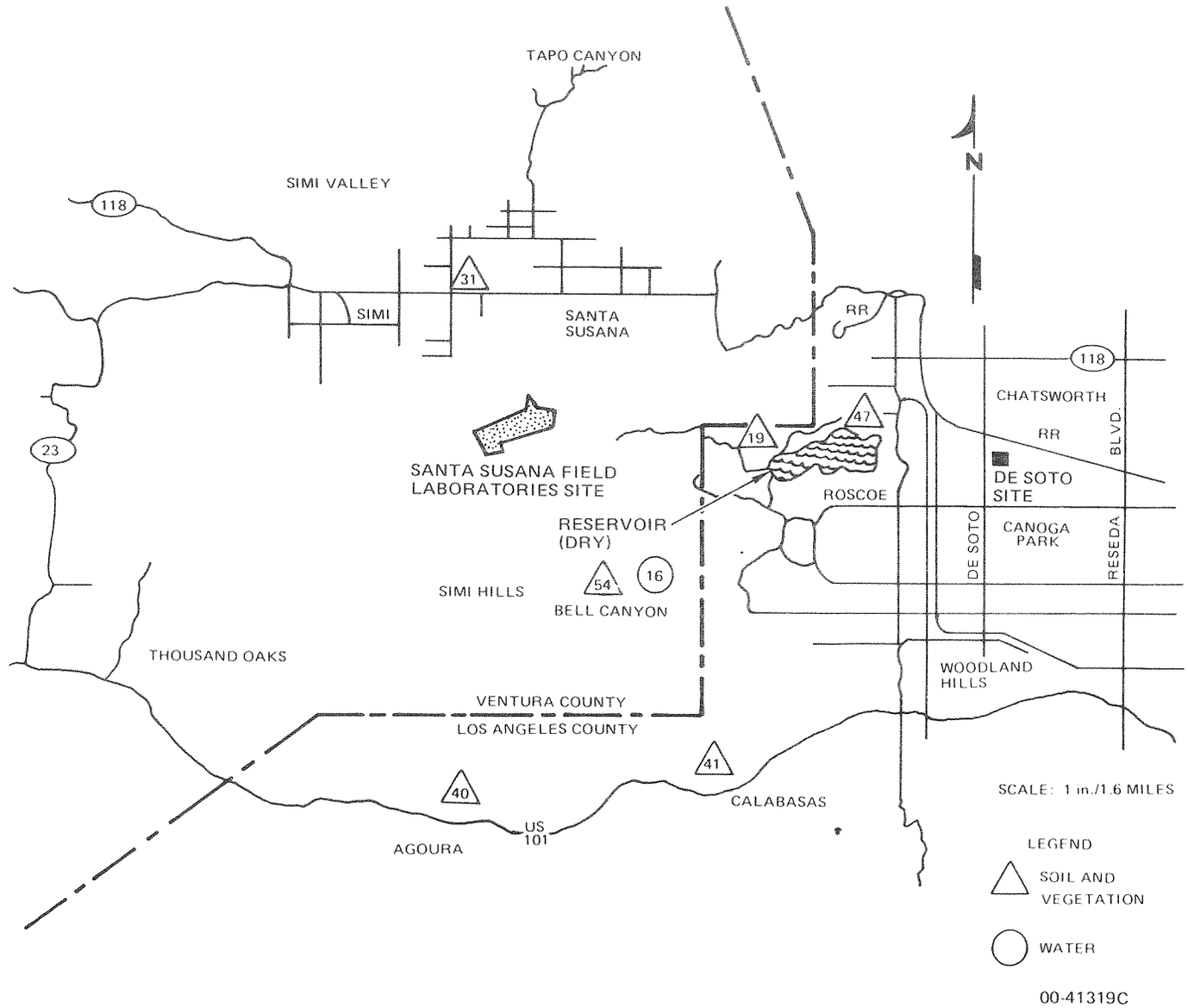


Figure 5. Map of Canoga Park, Simi Valley, Agoura and Calabasas Sampling Stations

III. ENVIRONMENTAL MONITORING PROGRAM

A. GENERAL DESCRIPTION

Soil and vegetation sample collection and analysis for radioactivity were initiated in 1952, in the Downey, California area, where the Energy Systems Group was initially located. Environmental sampling was subsequently extended to the proposed SRE site in the Simi Hills in May of 1954. In addition, sampling was begun in the Burro Flats area, southwest of SRE, where other nuclear installations were planned and are currently in operation. The Downey area survey was terminated when the Group relocated to Canoga Park in 1955. The primary purpose of the environmental monitoring program is to survey environmental radioactivity adequately to ensure that ESG operations do not contribute significantly to environmental radioactivity. The locations of sampling stations are shown in Figures 5 through 7 and listed in Table 8.

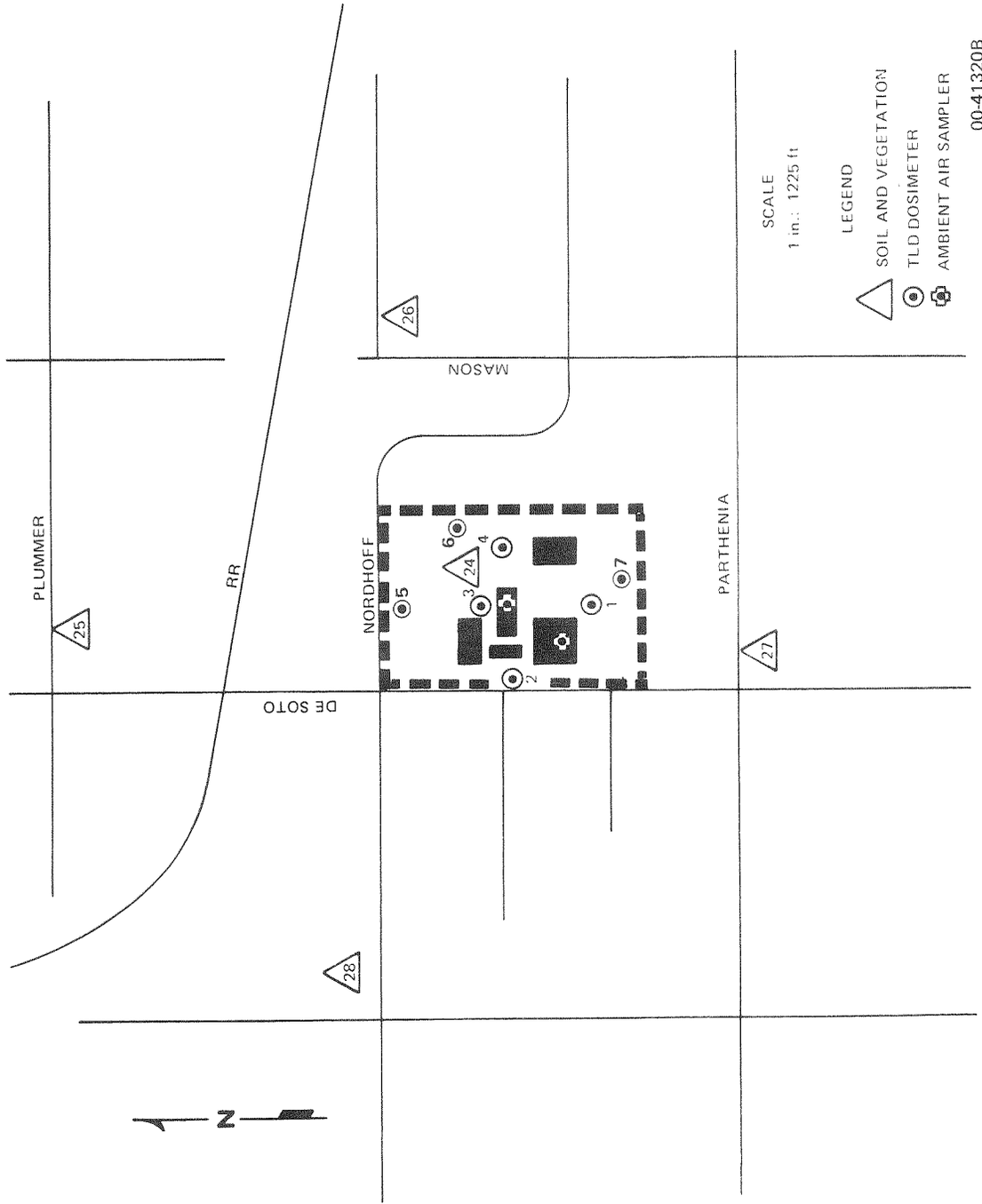
B. SAMPLING AND SAMPLE PREPARATION

1. Soil

Soil is analyzed for radioactivity to monitor for any significant increase in radioactive deposition by fallout from airborne radioactivity. Since soil is naturally radioactive and has been contaminated by atmospheric testing of nuclear weapons, a general background level of radioactivity exists. The data are monitored for increases beyond the natural variability of this background.

Surface soil types available for sampling range from decomposed granite to clay and loam. Samples are taken from the top 1/2-in. layer of undisturbed ground surface for gross radioactivity analysis and to a depth of 5 cm for plutonium analysis. The soil samples are packaged in plastic containers, and returned to the laboratory for analysis.

Sample preparation for gross radioactivity determination consists of transferring the soils to Pyrex beakers, and drying in a muffle furnace at $\sim 500^{\circ}\text{C}$ for 8 h. After cooling, the soil is sieved to obtain uniform particle size. Two-gram aliquots of the sieved soil are weighed, and transferred to copper planchets. The soil is wetted in the planchet with alcohol, evenly distributed to obtain uniform sample thickness, dried, and counted for alpha and beta radiation. Plutonium in soil analysis is performed according to the guidelines specified in



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Figure 6. Map of De Soto Site and Vicinity Sampling Stations

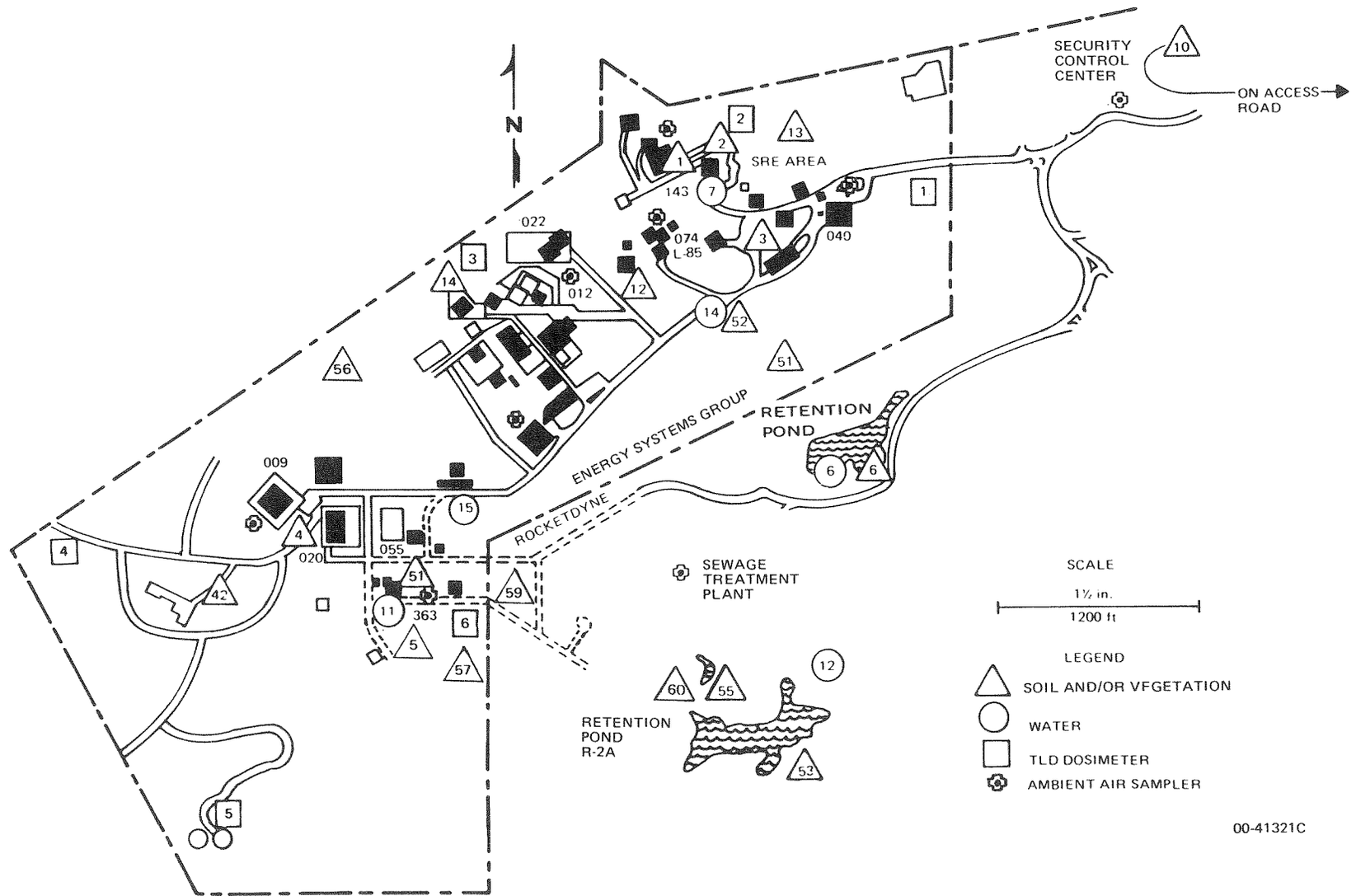


Figure 7. Map of Santa Susana Field Laboratories Site Sampling Stations

TABLE 8
SAMPLE STATION LOCATIONS
(Sheet 1 of 3)

Station	Location
SV-1	SSFL Site, Bldg. 143
SV-2	SSFL Site, Bldg. 143 Perimeter Drainage System
SV-3	SSFL Site, Bldg. 064
SV-4	SSFL Site, Bldg. 020
SV-5	SSFL Site, Bldg. 363
SV-6	Rocketdyne Site Interim Retention Pond
SV-10	SSFL Site Access Road
SV-12	SSFL Site, Bldg. 093 (L-85 Reactor)
SV-13	SSFL Site, Below Sodium Cleaning Facility at SRE Pond
SV-14	SSFL Site, Bldg. 028
SV-19	SSFL Site Entrance, Woolsey Canyon
SV-24	De Soto Site, Bldg. 004
SV-25	De Soto Avenue and Plummer Street
SV-26	Mason Avenue and Nordhoff Street
SV-27	De Soto Avenue and Parthenia Street
SV-28	Canoga Avenue and Nordhoff Street
SV-31	Simi Valley, Alamo Avenue and Sycamore Road
SV-40	Agoura — Kanan Road and Ventura Freeway
SV-41	Calabasas — Parkway Calabasas and Ventura Freeway
SV-42	SSFL Site, Bldg. 886
SV-47	Chatsworth Reservoir North Boundary
SV-51	SSFL Site, Bldg. 029
SV-52	SSFL Site, Burro Flats Drainage Control Pond, G Street and 17th Street
SV-53	Rocketdyne Site Pond R-2A Spillway, Head of Bell Canyon
SV-54	Bell Creek
S-55	Rocketdyne Site Retention Pond R-2A (Pond Bottom Mud)
S-56	SSFL Site, F Street and 24th Street

SV — Soil and Vegetation Sample Station
S — Soil Sample Station

TABLE 8
SAMPLE STATION LOCATIONS
(Sheet 2 of 3)

Station	Location
S-57	SSFL Site, J Street at Bldg. 055
S-58	SSFL Site, Bldg. 353
S-59	Rocketdyne Site Test Area CTL 4
S-60	Rocketdyne Site Retention Pond R-2A
W-6	Rocketdyne Site Interim Retention Pond (drains to Pond R-2A)
W-7	SSFL Site Domestic Water, Bldg. 003
W-11	SSFL Site Domestic Water, Bldg. 363
W-12	Rocketdyne Site Area II Final Retention Pond R-2A
W-16	Bell Creek
A-1	De Soto Site, Bldg. 001 Roof
A-2	De Soto Site, Bldg. 004 Roof
A-3	SSFL Site, Bldg. 009, West Side
A-4	SSFL Site, Bldg. 011, West Side
A-5	Rocketdyne Site, Bldg. 600, North Side
A-6	Rocketdyne Site, Bldg. 207, North Side
A-7	SSFL Site, Bldg. 074, South Side
A-8	SSFL Site, Bldg. 143, West Side
A-9	SSFL Site, Bldg. 363, West Side
TLD-1	De Soto Site, South of Bldg. 102
TLD-2	De Soto Site, West Boundary
TLD-3	De Soto Site, Guard Post No. 1, Bldg. 201
TLD-4	De Soto Site, East Fence
TLD-5	De Soto Site, North Boundary
TLD-6	De Soto Site, East Boundary
TLD-7	De Soto Site, South Boundary
TLD-1	SSFL Site, Bldg. 114

S — Soil Sample Station
W — Water Sample Station
A — Air Sampler Station
TLD — Thermoluminescent Dosimeter Location

TABLE 8
 SAMPLE STATION LOCATIONS
 (Sheet 3 of 3)

Station	Location
TLD-2	SSFL Site, SRE Water Retention Pond
TLD-3	SSFL Site, Electric Substation No. 719
TLD-4	SSFL Site, West Boundary on H Street
TLD-5	SSFL Site, Water Tank No. 701
TLD-6	SSFL Site, Bldg. 854
TLD-1	Off Site, Northridge
TLD-2	Off Site, Simi Valley
TLD-3	Off Site, Northridge

TLD – Thermoluminescent Dosimeter Location

U. S. NRC Regulatory Guide 4.5 titled "Measurements of Radionuclides in the Environment-Sampling and Analysis of Plutonium in Soil."

2. Vegetation

The analysis of vegetation is performed as an adjunct to the soil analysis and is done to determine the uptake of radioactivity by plants. These plants do not contribute to the human food chain, nor is there significant agriculture or grazing in the immediate neighborhood of either site.

Vegetation samples obtained in the field are of the same perennial plant types, wherever possible; these are usually sunflower or wild tobacco leaves. Vegetation leaves are stripped from plants, and placed in ice cream cartons for transfer to the laboratory for analysis. Ordinarily, plant root systems are not analyzed.

Vegetation samples are first washed with tap water to remove foreign matter, and then thoroughly rinsed with distilled water. Washed vegetation is dried in tared beakers at 100°C for 24 h for dry weight determination, then ashed in a muffle furnace at ~500°C for 8 h, producing a completely burned ash. One-gram aliquots of pulverized ash from each beaker are weighed, and transferred to copper planchets. The vegetation ash is wetted in the planchet with alcohol, evenly distributed to obtain uniform sample thickness, dried, and counted for alpha and beta radiation. The dry/ash weight ratio is used for the determination of the equivalent dry weight gross radioactivity concentration value.

3. Water

Surface and domestic supply water samples are obtained monthly at the SSFL site and from Bell Creek. The water is drawn into 1-liter polyethylene bottles, and transferred to the laboratory.

Five-hundred-milliliter volumes of water are evaporated to dryness in crystallizing dishes at ~90°C. The residual salts are redissolved into distilled water, transferred to copper planchets, dried under heat lamps, and counted for alpha and beta radiation.

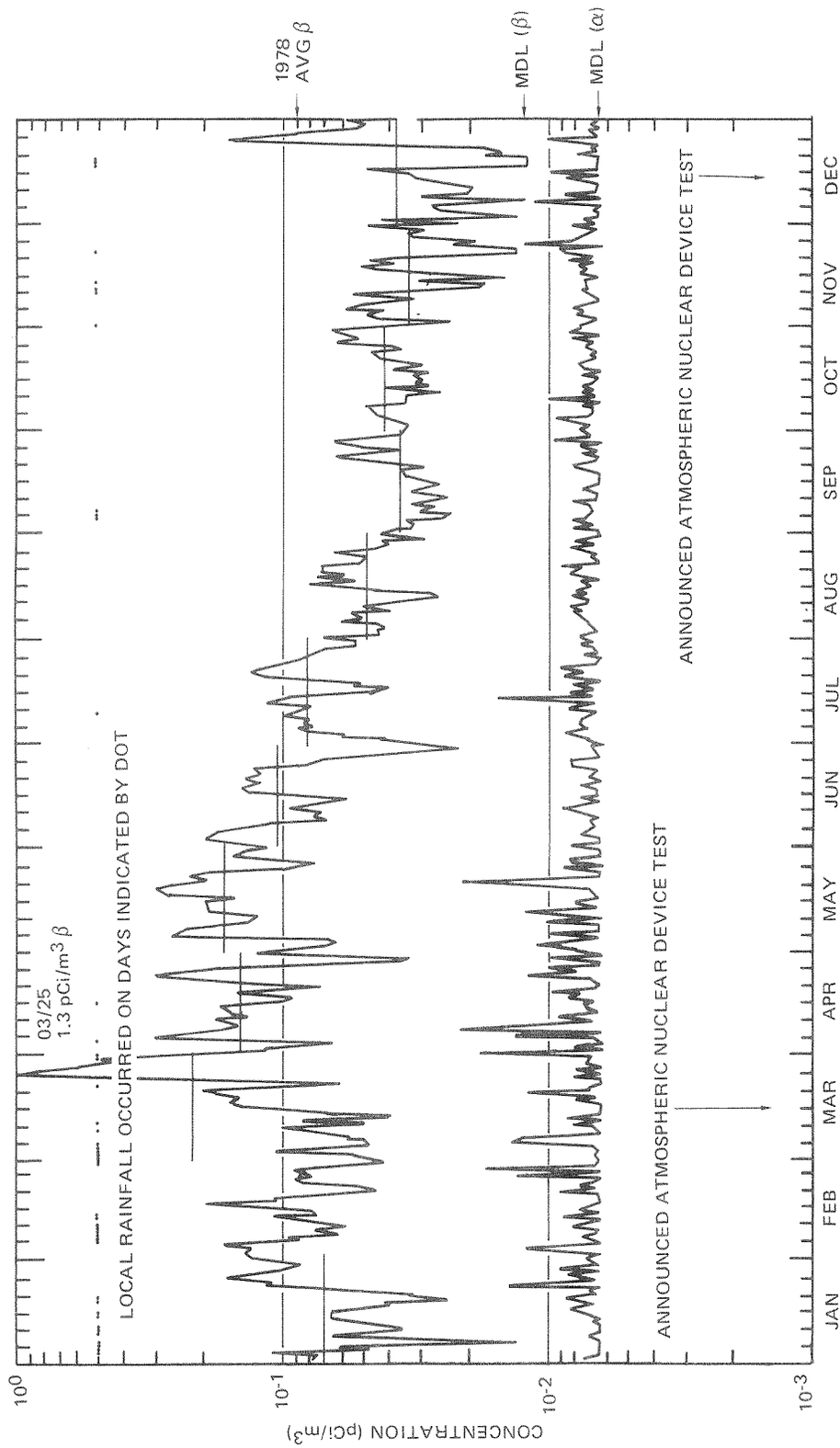


Figure 8. Daily Averaged Long-Lived Airborne Radioactivity at the De Soto and Santa Susana Field Laboratories Sites — 1978

4. Ambient Air

Air sampling is performed continuously at the De Soto and SSFL sites with automatic air samplers, operating on 24-h sampling cycles. Airborne particulate radioactivity is collected on Type HV-70 filter media, which are automatically changed daily at the end of each sampling period. The samples are counted for alpha and beta radiation following a minimum 120-h decay period to eliminate radon particulate daughters. The volume of a typical daily ambient air sample is approximately 25 m³.

Figure 8 is a graph of the daily averaged long-lived alpha and beta ambient air radioactivity concentrations for the De Soto and SSFL sites during 1978. The average beta concentration for each month is also indicated by horizontal bars. The graph shows two prominent peaks occurring during the year, a moderate spring increase in concentration and subsequent decreasing levels through the year's end.

C. COUNTING AND CALIBRATION

Environmental soil, vegetation, water, and ambient air samples are counted for alpha and beta radiation with a low-background gas flow proportional counting system, capable of the simultaneous counting of both alpha and net beta radiation. The sample-detector configuration provides a nearly 2 π geometry. The thin-window detector is continually purged with methane counting gas. A preset time mode of operation is used for all samples. The minimum detection limits shown in Table 9 were determined by using typical values for counting time, system efficiencies for detecting alpha and beta radiation, background count rates (approximately 0.05 cpm α and 1.0 cpm β) and sample size. For the table, the minimum statistically significant amount of radioactivity, irrespective of sample configuration, is taken as that amount equal in count rate to three times the standard deviation of the system background count rate.

Counting system efficiencies are determined routinely with Ra-D+E+F (with alpha absorber), Cl³⁶, Th²³⁰, U²³⁵, and Pu²³⁹ standard sources, and with K⁴⁰, in the form of standard reagent grade KCl, which is used to simulate soil and vegetation samples. Self-absorption standards are made by dividing sieved KCl into samples, increasing in mass by 200-mg increments, from 100 to 3000 mg. The samples are placed in copper planchets, of the type used for environmental samples,

TABLE 9
MINIMUM RADIOACTIVITY DETECTION LIMITS (MDL)

Sample	Activity	Minimum Detection Limits*
Soil	α	$(5.8 \pm 6.9) 10^{-8} \mu\text{Ci/g}$
	β	$(2.3 \pm 2.3) 10^{-7} \mu\text{Ci/g}$
Vegetation	α	$(1.2 \pm 1.4) 10^{-7} \mu\text{Ci/g ash}$
	β	$(3.7 \pm 3.6) 10^{-7} \mu\text{Ci/g ash}$
Water	α	$(2.3 \pm 2.8) 10^{-10} \mu\text{Ci/ml}$
	β	$(6.2 \pm 6.1) 10^{-10} \mu\text{Ci/ml}$
Air	α	$(6.0 \pm 7.2) 10^{-15} \mu\text{Ci/ml}$
	β	$(1.2 \pm 1.2) 10^{-14} \mu\text{Ci/ml}$

*95% Confidence Level

and counted. The ratio of sample activity to the observed net count rate for each sample is plotted as a function of sample weight. The correction factor (ratio) corresponding to sample weight may be obtained from the graph. The product of the correction factor and the net sample count rate yields the sample activity (dpm). This method has been proved usable by applying it to various-sized aliquots of uniformly mixed environmental samples and observing that the resultant specific activities fall within the expected statistical counting error.

Since the observed radioactivity in environmental samples results primarily from natural and weapons-testing sources, and is at such low concentrations, an effort is not made to identify individual radionuclides. The detection of significant levels of radioactivity would lead to an investigation of the radioactive material involved, the sources and possible causes.

D. NONRADIOACTIVE MATERIALS

Rockwell International Corporation, Rocketdyne Division, has filed a Report of Waste Discharge with the California Regional Water Quality Control Board, and

has been granted a National Pollutant Discharge Elimination System permit to discharge wastewater, pursuant to Section 402 of the Federal Water Pollution Control Act. The permit, NPDES No. CA0001309, became effective on September 27, 1976, and supersedes all previously held permits for wastewater discharge from the Rocketdyne Division SSFL. Discharge of overflow and storm runoff only is permitted into Bell Creek from water reclamation retention ponds. Discharge generally occurs only during and immediately after periods of heavy rainfall or during extended periods of rocket engine testing.

Only one of the retention ponds receives influent directly from the ESG SSFL site. It is identified as retention pond R-2A, Water Sample Station W-12 in Table 8. The influent includes sewage treatment plant effluent and surface runoff water. Grab-type water samples, taken at the retention pond prior to a discharge, are analyzed for non-radioactive chemical constituents by a California State certified analytical testing laboratory and for tritium. The specific constituents analyzed for, and their respective limitations in discharged wastewater, are presented in Appendix B. Wastewater originating from facilities located throughout the SSFL site is composited in the retention pond. Therefore, the point of origin of nonradioactive constituents found in wastewater is impossible to determine.

TABLE 10
 ATMOSPHERICALLY DISCHARGED EFFLUENT RELEASED
 TO UNRESTRICTED AREAS — 1978

Building	Point of Release	Approximate Effluent Volume (ft ³)	Activity Monitored	Approximate Minimum Detection Limit (μCi/ml)	Annual* Average Concentration (μCi/ml)	Sampling Period Maximum Observed Concentration (μCi/ml)	Total Radio-Activity Released (Ci)
001	Stack Exit	2.3 x 10 ¹⁰	α	1.7 x 10 ⁻¹⁶	<1.8 x 10 ⁻¹⁴	1.4 x 10 ⁻¹³	<1.6 x 10 ⁻⁵
			β	5.4 x 10 ⁻¹⁶	<5.8 x 10 ⁻¹⁵	3.8 x 10 ⁻¹⁴	<5.0 x 10 ⁻⁶
004	Stack Exit	3.4 x 10 ¹⁰	α	2.7 x 10 ⁻¹⁶	<6.8 x 10 ⁻¹⁶	4.7 x 10 ⁻¹⁵	<6.5 x 10 ⁻⁷
			β	9.2 x 10 ⁻¹⁶	<4.5 x 10 ⁻¹⁵	4.6 x 10 ⁻¹⁴	<4.3 x 10 ⁻⁶
020	Stack Exit	1.1 x 10 ¹⁰	α	0.9 x 10 ⁻¹⁶	<4.0 x 10 ⁻¹⁶	1.8 x 10 ⁻¹⁵	<1.3 x 10 ⁻⁷
			β	3.0 x 10 ⁻¹⁶	1.4 x 10 ⁻¹³	1.0 x 10 ⁻¹²	5.9 x 10 ⁻⁵
021-022	Stack Exit	1.2 x 10 ¹⁰	α	2.1 x 10 ⁻¹⁶	<3.2 x 10 ⁻¹⁶	8.3 x 10 ⁻¹⁶	<1.0 x 10 ⁻⁷
			β	7.0 x 10 ⁻¹⁶	3.5 x 10 ⁻¹⁴	2.5 x 10 ⁻¹³	<1.1 x 10 ⁻⁵
055	Stack Exit	8.0 x 10 ⁹	α	2.4 x 10 ⁻¹⁶	<3.5 x 10 ⁻¹⁶	9.7 x 10 ⁻¹⁶	<8.1 x 10 ⁻⁸
						Total	<9.6 x 10 ⁻⁵

Annual average ambient air radioactivity concentration — 1978

α	<7.5 x 10 ⁻¹⁵
β	<8.8 x 10 ⁻¹⁴

*Effluent radioactivity is generally less than ambient air radioactivity. If released concentrations were at the ambient air radioactivity concentrations, a total of 2.4 x 10⁻⁴ Ci would be released, which value is 2.5 times greater than the actual release.

IV. EFFLUENT MONITORING PROGRAM

Effluents which may contain radioactive material are generated at ESG facilities as the result of operations performed under contract to DOE, under NRC Special Nuclear Materials License SNM-21, and under State of California Radioactive Material License 0015-59. The specific facilities are identified as Buildings 001 and 004 at the De Soto site, and Buildings 020, 021, 022, and 055 at the Santa Susana site, SSFL.

A. TREATMENT AND HANDLING

Waste streams released to unrestricted areas are limited in all cases, to gaseous effluents. No contaminated liquids are discharged to unrestricted areas.

The level of radioactivity contained in all atmospherically discharged effluents is reduced to the lowest practicable values by passing the effluents through certified, high efficiency particulate air (HEPA) filters. These effluents are sampled for particulate radioactive materials by means of continuous stack exhaust samplers at the point of release. In addition, stack monitors installed at Buildings 020 and 055 provide automatic alarm capability in the event of the release of gaseous activity from Building 020 or particulate activity from Building 055. The HEPA filters used for filtering gaseous effluents are 99.97% efficient for particles of 0.3- μ m diameter. Particle filtration efficiency increases above and below this size.

The average concentration and total radioactivity in gaseous effluent released to unrestricted areas is shown in Table 10. The effectiveness of the air cleaning systems is evident from the fact that in most cases, the gaseous effluent released is less radioactive than the ambient air, which is indicative that there are not radioactivity releases during normal facility operations.

Liquid wastes released to sanitary sewage systems, a controlled area as provided for by CAC 17 and 10 CFR 20, are generated at the De Soto site only. Liquid wastes are discharged from Building 001 following analysis on a volume basis only. There is no continuous flow. Building 004 liquid chemical wastes are released to a proportional sampler installation which retains an aliquot

TABLE 11
LIQUID EFFLUENT DISCHARGED TO SANITARY SEWER — 1978

Building	Point of Release	Approximate Effluent Volume (gal)	Activity Monitored	Approximate MDL ($\mu\text{Ci}/\text{m}\ell$)	Annual Average Concentration ($\mu\text{Ci}/\text{m}\ell$)	Sample Maximum Observed Concentration ($\mu\text{Ci}/\text{m}\ell$)	Total Radioactivity Released (Ci)
001	Retention Tank	40,500	α	1.2×10^{-9}	1.5×10^{-7}	3.7×10^{-7}	2.3×10^{-5}
			β	4.1×10^{-9}	1.1×10^{-7}	3.3×10^{-7}	1.7×10^{-5}
004	Proportional Sampler	1,039,600	α	1.2×10^{-9}	$<2.1 \times 10^{-8}$	1.5×10^{-7}	$<8.0 \times 10^{-5}$
			β	4.1×10^{-9}	7.8×10^{-8}	1.4×10^{-6}	3.1×10^{-4}
020*	—	0	—	—	—	—	—
021 - 022*	—	0	—	—	—	—	—
055*	—	0	—	—	—	—	—

*All liquid radioactive wastes are solidified and land buried as dry waste.

each time a fixed volume is released to the sanitary sewage system. No radioactive liquid effluents are released from the Santa Susana Buildings 020, 021, 022, or 055, except as controlled liquid radioactive waste solidified for land burial. The average concentration and total radioactivity in liquid effluents discharged is shown in Table 11.

B. ENERGY SYSTEMS GROUP FACILITY DESCRIPTIONS

1. De Soto Site

a. Building 001 — NRC and California State Licensed Activities

Operations at Building 001 which may generate radioactive effluents consist of production operations associated with the manufacture of enriched uranium fuel elements. Only atmospherically discharged effluents are released from the building to uncontrolled areas. Following analysis for radioactivity concentration, liquid wastes are released to the sanitary sewage system, which is considered a controlled area, as provided by CAC 17 and 10 CFR 20. Nuclear fuel material handled in unencapsulated form in this facility contains the uranium isotopes U^{234} , U^{235} , U^{236} , and U^{238} . No significant quantities of these radionuclides were released.

b. Building 004 — NRC and California State Licensed Activities

Operations at Building 004 which may generate radioactive effluents consist of research studies in physics and chemistry, and the chemical analysis of small quantities of fuel materials, usually limited to a few grams. Only atmospherically discharged effluents are released from the building to uncontrolled areas. Liquid laboratory wastes are released to a proportional sampler installation which retains an aliquot of wastewater each time a fixed volume is released to the facility sanitary sewage system. The aliquots are composited and analyzed for radioactivity. Nuclear fuel material handled in unencapsulated form in this facility contains the uranium isotopes U^{234} , U^{235} , U^{236} , and U^{238} . Major quantities of other radionuclides in encapsulated form include Co^{60} and Pm^{147} . No significant quantities of these radionuclides were released. The monitoring of De Soto site sewage effluent for tritium commenced during December 1977 and continues on the basis of a monthly analysis of a daily composited sample of total facility effluent collected at the discharge point into the municipal sewerage.

2. Santa Susana Field Laboratories Site

a. Building 020 – NRC and California State Licensed Activities

Operations at Building 020 which may generate radioactive effluents consist of hot cell examination of irradiated nuclear fuels and reactor components. Only atmospherically discharged effluents are released from the building to uncontrolled areas. The effluent may contain particulate material, as well as radioactive gases, depending on the operations being performed and the history of the irradiated fuel and other material. The chemical form of such materials may be U metal, UO_2 , UC, mixed fission products, and various activation products. No radioactive liquid waste is released from the facility. Radioactive material handled in unencapsulated form in this facility includes the following radionuclides: Th^{232} , U^{233} , U^{234} , U^{235} , U^{236} , and U^{238} as constituents in the various fuel materials; and Cs^{137} , Sr^{90} , Kr^{85} , and Pm^{147} as mixed fission products. No significant quantities of these radionuclides were released.

b. Buildings 021 and 022 – DOE Contract Activities

Operations at Buildings 021 and 022 which may generate radioactive effluents consist of the processing, packaging, and temporary storage of liquid and dry radioactive waste material for disposal. Only atmospherically discharged effluents are released from the building to uncontrolled areas. No radioactive liquid waste is released from the facility. Nuclear fuel material handled in encapsulated or unencapsulated form contains the uranium isotopes U^{234} , U^{235} , U^{236} , U^{238} , plus Cs^{137} , Sr^{90} , and Pm^{147} as mixed fission products. No significant quantities of these radionuclides were released.

c. Building 055 – NRC and California State Licensed Activities

Operations at Building 055 which may generate radioactive effluents consist of fabrication of plutonium and plutonium-uranium fuel pins. Only atmospherically discharged effluents are released from the facility to uncontrolled areas. No radioactive liquid waste is released from the facility.

The various fuel materials (depleted and enriched uranium and plutonium) contain the following radionuclides: U^{234} , U^{235} , U^{236} , U^{238} , Pu^{238} , Pu^{239} , Pu^{240} , Pu^{241} , and Am^{241} . No significant quantities of these radionuclides were released.

C. ESTIMATION OF GENERAL POPULATION DOSE

Release of airborne material at the De Soto site for summer season weather conditions would generally be under a subsidence inversion into an atmosphere that is typical of slight neutral to lapse conditions. Although nocturnal cooling inversions are present they are relatively shallow in extent. During the summer season the subsidence inversion is present almost every day. The base and top of this inversion for the most part lie below the elevation of the SSFL site. Thus, any atmospheric release under this condition from the SSFL site would result in Pasquill Type D lofting diffusion conditions above the inversion and considerable atmospheric dispersion prior to diffusion (if any) through the inversion into the Simi or San Fernando Valleys. In the winter season the Pacific high pressure cell shifts to the south and the subsidence inversion for the most part is missing. The surface air flow is dominated by frontal activity moving through the area or to the east. Frontal passages through the area during this season are generally accompanied by precipitation. Diffusion characteristics are highly variable depending upon the frontal location. Generally, a light to moderate southwesterly wind precedes these frontal passages introducing strong onshore flow of marine air, and lapse rates are slight neutral to lapse. Wind speeds increase with the approach of the frontal systems, enhancing diffusion. The diffusion characteristics of the frontal passage are lapse conditions with light to moderate northerly winds. A summary of surface wind conditions for the local area is presented in Table 12.

TABLE 12
SURFACE WIND CONDITIONS

	Summer	Winter
Prevailing afternoon direction	WNW	NW
Prevailing early morning direction	ESE	ESE
Average daytime speed	8 mph	6 mph
Average nighttime speed	3 mph	3 mph

The population distributions around the De Soto and SSFL sites used to estimate population doses in this section are based on the 1970 census data projected for 1980. The projections were based on an average growth rate of 5.17%/yr for this area. For population distribution at distances >5 miles out to 50 miles, a single distribution centered on 34°14'25" north and 118°39'00" west is used. This location is between the two ESG sites which are ~6 miles apart. This population distribution is also based on the 1970 census data, with the 1980 projection based on the average growth rate of 5.17 %/yr.

The calculated downwind concentration of radioactive material discharged during 1978 from each of the four major ESG nuclear facilities is presented in Table 13. The Type B stability parameter coefficients and a mean wind speed of 2.2 m/s were used for the calculations.

TABLE 13
DOWNWIND CONCENTRATION OF GASEOUS EFFLUENTS — 1978

Facility	Q (Ci/s)	Meters to		Type B Stability (σ_y)			Type B Stability (σ_z)			χ Downwind ($\mu\text{Ci}/\text{cm}^3$)		
		Boundary	Residence	Boundary	Residence	80 km	Boundary	Residence	80 km*	Boundary	Residence	80 km
B/001	6.7×10^{-13}	110 W	171 SW	18	27	6800	12	17	2000	4.5×10^{-16}	2.1×10^{-16}	7.1×10^{-21}
B/020	1.9×10^{-12}	305 NW	1900 SE	50	290	6800	34	340	2000	1.6×10^{-16}	2.7×10^{-18}	2.0×10^{-20}
B/022	3.5×10^{-13}	350 NW	2300 SE	55	310	6800	38	500	2000	2.4×10^{-17}	3.3×10^{-19}	3.7×10^{-21}
B/055	2.6×10^{-15}	400 NW	1830 SE	68	260	6800	42	300	2000	1.3×10^{-19}	4.8×10^{-21}	2.7×10^{-23}

*Assume \bar{u} = 2.2 m/s average wind speed, constant direction, full year
* σ_z Type C stability category

The general population man-rem dose estimates calculated from demography data and the concentrations calculated for atmospherically discharged effluent data are presented in Table 14. It should be noted that these estimates assume level surrounding terrain and ignore the effect of the mountains that completely encircle the sites at distances of about 10 km. The air turbulence and changes in elevation associated with actual terrain would result in much lower concentrations than those calculated.

The off-site doses are extremely low compared to the maximum permissible exposures recommended for the general population. These values are 3 rem/year for bone, and 1.5 rem/year for the lung for an individual, and are one-third of

TABLE 14
POPULATION DOSE ESTIMATES FOR ATMOSPHERIC DISCHARGED EFFLUENTS

Sector	Dose to Receptor Population Segment (man-rem)						
	0-8 km	8-16 km	16-32 km	32-48 km	48-64 km	64-80 km	Total
N-NNE	0.0099	0.0000	0.0002	0.0000	0.0000	0.0000	0.010
NNE-NE	0.013	0.0000	0.0001	0.0000	0.0004	0.0000	0.014
NE-ENE	0.59	0.10	0.0008	0.0000	0.0000	0.0000	0.69
ENE-E	0.58	0.18	0.0016	0.0005	0.0002	0.0001	0.76
E-ESE	1.2	0.21	0.0026	0.0024	0.0014	0.0007	1.4
ESE-SE	1.4	0.11	0.0019	0.0029	0.0018	0.0008	1.5
SE-SSE	1.2	0.57	0.0008	0.0003	0.0005	0.0000	1.8
SSE-S	1.2	0.062	0.0000	0.0000	0.0000	0.0000	1.3
S-SSW	0.0016	0.0000	0.0000	0.0000	0.0000	0.0000	0.002
SSW-SW	0.005	0.0001	0.0000	0.0000	0.0000	0.0000	0.005
SW-WSW	0.0037	0.0000	0.0002	0.0000	0.0000	0.0000	0.004
WSW-W	0.031	0.0000	0.0001	0.0002	0.0002	0.0000	0.032
W-WNW	0.19	0.0008	0.0000	0.0001	0.0000	0.0000	0.19
WNW-NW	0.50	0.0001	0.0000	0.0000	0.0000	0.0000	0.5
NW-NNW	0.19	0.0000	0.0000	0.0000	0.0000	0.0000	0.19
NNW-N	0.0099	0.0000	0.0000	0.0000	0.0000	0.0000	0.010
	7.1	1.2	0.0086	0.0064	0.0045	0.0016	8.4

1. Average rem/man dose = 0.00001 rem for the 80 km segment average population.
2. Total 80 km man-rem dose estimate from naturally occurring airborne radioactivity dose to the lung of ~ 0.1 rem/year = 1,300,000 man-rem for the 80 km segment average population

these values for the general population. From Table 14, it may be seen that the highest segment dose integral is for the 0-8 km segment equivalent to an average dose/man-year of 0.48 mrem equivalent to 0.096% of the maximum permissible exposure for an individual and 0.28% of the general population recommended average exposure. Thus, atmospheric discharges from ESG nuclear facilities have resulted in estimated radiation doses which are a small fraction of the recommended limits. Doses due to internal deposition of natural radioactivity in air are ~ 50 to 100 mrem per year.

APPENDIX A
COMPARISON OF ENVIRONMENTAL RADIOACTIVITY DATA
FOR 1978 WITH PREVIOUS YEARS

This section compares environmental monitoring results for the calendar year 1978 with previous annual data.

The data presented in Tables A-1 through A-5 summarize all past annual average radioactivity concentrations. These data show the effects of both the short-lived and long-lived radioactive fallout from nuclear weapons tests superimposed on the natural radioactivity inherent in the various sample types.

Over the considerable period of time that the environmental program has been in operation, evolutionary changes have been made in order to provide more effective data. In some cases this is readily apparent in the data. For example, in Table A-1, a small but abrupt increase in the alpha activity reported for soil is seen to occur in 1971. This increase is observed in both the on-site and the off-site samples and resulted from use of an improved counting system with a thinner sample configuration. The thinner sample increases the sensitivity of the detector to alpha-emitting radionuclides in the sample, thus producing a higher measured specific activity.

Similarly, prior to 1971, gross activity in ambient air was measured, including both alpha and beta activity. In 1971, measurements were begun which allowed separate identification of these two types of activity.

The types of random variations observed in the data indicate that there is no local source of unnatural radioactivity in the environment. Also, the similarity between on-site and off-site results further indicate that the contribution to general environmental radioactivity due to operations at ESG is essentially nonexistent.

TABLE A-1
SOIL RADIOACTIVITY DATA — 1957 THROUGH 1978

Year	On Site-Average (10^{-6} $\mu\text{Ci/g}$)			Off Site - Average (10^{-6} $\mu\text{Ci/g}$)		
	Number Samples	α	β	Number Samples	α	β
1978	144	0.63	24	48	0.51	24
1977	144	0.56	24	48	0.53	23
1976	144	0.56	25	48	0.56	24
1975	144	0.60	25	48	0.58	24
1974	144	0.60	25	48	0.54	24
1973	144	0.57	25	48	0.51	24
1972	144	0.56	25	48	0.57	24
1971	144	0.55	25	48	0.53	23
1970	144	0.47	27	48	0.48	25
1969	144	0.42	27	48	0.42	25
1968	144	0.47	26	48	0.48	26
1967	144	0.42	28	48	0.39	24
1966	144	0.41	29	48	0.44	25
1965	144	0.46	36	142	0.47	29
1964	152	0.46	32	299	0.44	26
1963	156	0.43	45	455	0.42	42
1962	147	0.44	48	453	0.41	47
1961	120	0.37	34	458	0.33	23
1960	115	0.41	23	362	0.37	19
1959	107	0.43	15	377	0.32	14
1958	80	0.27	21	309	0.26	10
1957	64	0.32	11	318	0.35	10

TABLE A-2
VEGETATION RADIOACTIVITY DATA — 1957 THROUGH 1978

Year	On Site — Average (10^{-6} μ Ci/g ash)			Off-Site — Average (10^{-6} μ Ci/g ash)		
	Number Samples	α	β	Number Samples	α	β
1978	144	<0.24	166	48	<0.24	143
1977	144	<0.22	162	48	<0.21	142
1976	144	<0.19	170	48	<0.22	147
1975	144	<0.21	155	48	<0.21	141
1974	144	<0.20	152	48	<0.27	141
1973	144	<0.24	155	48	<0.24	142
1972	144	0.23	145	48	0.36	125
1971	144	0.24	165	48	0.31	132
1970	144	0.33	159	48	0.30	142
1969	144	0.40	165	48	0.36	144
1968	144	0.51	158	48	0.51	205
1967	144	0.62	286	48	0.39	413
1966	144	0.37	169	48	0.37	123
1965	144	0.56	162	142	0.61	138
1964	154	0.50	211	293	0.51	181
1963	156	0.44	465	456	0.37	388
1962	147	0.45	500	453	0.44	406
1961	120	0.35	224	459	0.29	246
1960	115	0.35	137	362	0.25	136
1959	96	0.29	212	293	0.18	168
1958	65	0.57	683	250	0.39	356
1957	58	1.1	208	304	0.89	200

TABLE A-3
 SSFL SITE DOMESTIC WATER RADIOACTIVITY DATA —
 1957 THROUGH 1978

Year	Number Samples	Average α (10^{-9} $\mu\text{Ci}/\text{m}\ell$)	Average β (10^{-9} $\mu\text{Ci}/\text{m}\ell$)
1978	24	<0.26	3.0
1977	24	<0.25	2.5
1976	24	<0.25	2.0
1975	24	<0.24	2.3
1974	24	<0.24	2.7
1973	24	<0.26	3.4
1972	24	0.22	3.7
1971	24	0.28	4.9
1970	24	0.18	5.3
1969	24	0.11	5.0
1968	24	0.16	5.0
1967	24	0.13	6.1
1966	24	0.13	4.6
1965	24	0.22	6.0
1964	23	0.18	5.3
1963	24	0.18	7.0
1962	24	0.21	12.0
1961	24	0.08	2.9
1960	22	0.08	1.9
1959	18	0.08	1.6
1958	13	0.16	4.7
1957	17	—	13.0

TABLE A-4

BELL CREEK AND ROCKETDYNE DIVISION RETENTION POND RADIOACTIVITY DATA — 1966 THROUGH 1978

Samples															
Year	Bell Creek Mud 54			Bell Creek Vegetation 54			Bell Creek Water 16			Interim Retention Pond Water 6			Final Retention Pond R-2A Water 12		
	No. Samples	Average (10^{-6} μ Ci/g)		No. Samples	Average (10^{-6} μ Ci/g ash)		No. Samples	Average (10^{-9} μ Ci/ml)		No. Samples	Average (10^{-9} μ Ci/ml)		No. Samples	Average (10^{-9} μ Ci/ml)	
		α	β		α	β		α	β		α	β		α	β
1978	12	0.42	23.	12	<0.26	156.	12	<0.24	2.5	12	<0.25	4.3	12	<0.25	4.6
1977	12	0.29	22.	12	<0.19	155.	12	<0.24	1.8	12	<0.24	4.3	12	<0.25	5.2
1976	12	0.38	23.	12	<0.17	164.	12	<0.25	2.2	12	<0.24	4.3	12	<0.28	4.4
1975	12	0.29	22.	12	<0.19	123.	12	<0.22	2.4	12	<0.24	4.2	12	<0.31	4.5
1974	12	0.32	22.	12	<0.16	142.	12	<0.21	2.5	12	<0.22	4.2	12	<0.21	4.5
1973	12	0.34	24.	12	<0.17	147.	12	<0.21	2.7	12	<0.23	4.5	12	<0.37	5.6
1972	12	0.32	22.	12	0.12	139.	12	0.20	2.5	12	0.22	5.3	12	0.22	5.5
1971	12	0.36	23.	12	0.19	128.	12	0.15	3.8	12	0.18	6.2	12	0.16	6.4
1970	12	0.44	24.	12	0.23	165.	12	0.15	3.7	12	0.15	6.9	12	0.12	7.4
1969	12	0.35	27.	12	0.28	166.	12	0.04	4.0	12	0.07	5.9	11	0.10	5.7
1968	11	0.32	24.	11	0.39	170.	8	0.05	4.6	11	0.23	8.1	12	0.33	7.7
1967	12	0.40	24.	12	0.38	180.	12	0.07	5.8	12	0.19	6.6	10	0.17	7.0
1966	3	0.39	25.	3	1.1	108.	3	0.75	2.5	9	0.11	5.8	8	1.1	6.3

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TABLE A-5
 AMBIENT AIR RADIOACTIVITY CONCENTRATION DATA —
 1957 THROUGH 1978

Year	DeSoto Site Average (10 ⁻¹² μCi/ml)			SSFL Site Average [§] (10 ⁻¹² μCi/ml)		
	Number Samples	α	β	Number Samples	α	β
1978	713	<0.0084	<0.091	2402	<0.0072	<0.088
1977	729	<0.0066	<0.17	2438	<0.0066	<0.17
1976	719	<0.0067	<0.096	2520	<0.0065	<0.11
1975	709	<0.0063	<0.076	2450	<0.0060	<0.073
1974	663	<0.0056	<0.16	2477	<0.0057	<0.16
1973	715	<0.0075	<0.041	2311	<0.0072	<0.038
1972	708	0.0085	0.14	2430	0.0086	0.14
1971*	730	0.0087	0.30	2476	0.0086	0.33
1970	668	—	0.34	2434	—	0.36
1969	687	—	0.27	2364	—	0.26
1968	650	—	0.32	2157	—	0.32
1967	712	—	0.39	2400	—	0.41
1966	706	—	0.18	2205	—	0.17
1965	483	—	0.83	1062	—	0.21
1964	355	—	2.7	—	—	†
1963	360	—	6.6	292	—	4.7
1962	343	—	7.3	314	—	5.6
1961	313	—	4.2	176	—	3.6
1960	182	—	0.24	44	—	0.44
1959	215	—	2.5	257	—	0.93
1958	366	—	4.9	164	—	2.7
1957	63	—	1.6	141	—	2.7

*Ambient air alpha radioactivity values were included in the beta values and not reported separately prior to 1971

†Insufficient data

§Includes Rocketdyne Site Air Sampler Data

APPENDIX B

CALIFORNIA REGIONAL WATER QUALITY CONTROL BOARD CRITERIA FOR
DISCHARGING NONRADIOACTIVE CONSTITUENTS FROM ROCKETDYNE DIVISION, SSFL

The discharge of an effluent in excess of the following limits given in Table B-1 is prohibited.

TABLE B-1
NPDES NO. CA00-01309, EFFECTIVE SEPTEMBER 27, 1976

Constituent	Discharge Rate (lb/day)	Concentration Limit (mg/l)	
	30-day Average	30-day Average	Maximum
Total Dissolved Solids	1,267,680	-	950
Chloride	200,160	-	150
Sulfate	400,320	-	300
Suspended Solids*	66,720	50	150
Settleable Solids*	-	0.1	0.3
BOD 20°C	26,690	20	60
Oil and Grease	13,350	10	15
Chromium	6.67	0.005	0.01
Fluoride	1,340	-	1.0
Boron	1,340	-	1.0
Residual Chlorine	-	-	0.1
Fecal Coliform (MPN/100 ml)	-	-	23.0
Surfactants (as MBAS)	667	-	0.5
pH			6.0-9.0

*Not applicable to discharges containing rainfall runoff during or immediately after periods of rainfall.

APPENDIX C
REFERENCES

1. DOE Manual Chapter 0513
2. DOE Manual Chapter 0524, Appendix
3. Code of Federal Regulations, Title 10, Part 20
4. California Radiation Control Regulations, California Administrative Code, Title 17, Public Health
5. California Regional Water Quality Control Board, Los Angeles Region, Order No. 74-379, NPDES No. CA0001309, Effective September 27, 1976
6. Meteorology and Atomic Energy – 1968, TID 24190
7. Report of Committee II on Permissible Dose for Internal Radiation (1959), ICRP Publication 2
8. Deposition and Retention Models for Internal Dosimetry of the Human Respiratory Tract, ICRP Committee II Task Group on Lung Dynamics
9. Document TI #N001TI000-046 titled "Method of Estimating General Population Radiation Dose Attributable to Atmospheric Discharge of Radioactivity from ESG Nuclear Facilities," J. D. Moore

APPENDIX D
EXTERNAL DISTRIBUTION

1. Radiologic Health Section, State Department of Public Health, California
2. Radiological Health Division, Los Angeles County Health Department California
3. Environmental Health Department, Ventura County, California
4. U.S. Department of Energy, San Francisco Operations Office
5. U.S. Nuclear Regulatory Commission, Division of Reactor Licensing
6. Gordon Facer, Division of Military Applications, DOE
7. Andrew J. Pressesky, Reactor Research and Development, DOE
8. James Miller, Division of Biomedical and Environmental Research, DOE
9. DOE-Headquarters Library, Attention: Charles Sherman

