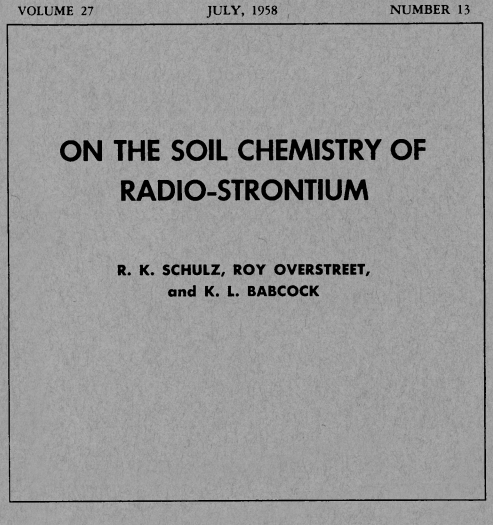
HILGARDIA

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The advent of radioactive fallout has stimulated interest in the soil chemistry of radio-strontium. In the present research, the chemical behavior of added Sr 85 in 26 California soils was studied. On the basis of the results of various extraction procedures with ammonium acetate and calcium chloride, it is concluded that added radio-strontium is retained in the water soluble and exchangeable forms. Therefore, barring long-term fixation effects, it is largely available to plants.

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ON THE SOIL CHEMISTRY OF RADIO-STRONTIUM¹ R. K. SCHULZ,² ROY OVERSTREET,³ and K. L. BABCOCK⁴

INTRODUCTION

A PHENOMENON of major concern to present-day soil chemists is the contamination of agricultural lands with strontium 90 and its subsequent introduction into food chains as a result of absorption by crops. At the present time, the general problem is all the more serious because comparatively little is known of the soil chemistry of minute traces of strontium and no means are at hand for assessing the availability of a given level of soil strontium to plants.

Perhaps the most reliable information concerning the uptake by plants has been obtained by experiments with culture solutions. The preponderance of these experiments suggests that in uptake by plants from culture solutions, Sr behaves very similarly to Ca. Actually, the evidence suggests that most plants cannot distinguish between the two elements in absorption although certain differences in behavior appear when Sr and Ca are translocated within the plant (Martin, *et al.*, 1957).⁵ Experiments with excised roots conducted in this laboratory showed that the dilution of radio-strontium in the culture solution with inactive Sr and Ca was about equally effective in reducing the uptake of the isotope (Overstreet, 1957).

It is thus reasonable to conclude that absorption of Sr from culture solutions is very closely tied to the Ca absorption. That is, when the Sr to Ca ratio in the culture solution is high, one can expect a corresponding high ratio within the plant. In line with this general idea, Comar *et al.* (1957) have introduced the term *strontium-calcium observed ratio*, OR. For the problem under consideration, as expressed below:

OR		$\frac{\mathrm{Sr}}{\mathrm{Ca}}$ in plant
On	plant-culture solution	 $\frac{\mathrm{Sr}}{\mathrm{Ca}}$ in culture solution

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⁵ See "Literature Cited" for citations referred to in the text by author and date.

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Where there is no discrimination between Sr and Ca in absorption, the OR value will be equal to unity. Middleton, as reported by Martin *et al.* (1957), has determined the OR values from culture solutions for some 10 plant species important in human diet or as sources of mineral nutrients in pastures. He found OR values ranging from 0.8 to 1.1. By and large, where gross uptake by the aerial parts of the plant is considered, it appears that satisfactory predictions concerning Sr 90 absorption from culture solutions can be made assuming an OR value of unity.

Soil series, type	pH Sat. paste	Conductivity* m.mho/cm.	Per cent lime†
Aiken c.l.	5.6	0.86	0.0
Hanford v.f.s.l.	7.7	0.46	0.0
Columbia v.f.s.l.	6.8	0.43	0.0
Fresno l.s.	6.3	3.54	0.0
Hanford s.l.	7.1	1.17	0.0
Yorkville l	5.8	0.27	0.0
Ramona s.l.	7.2	1.26	0.0
Cujunga v.f.s.l.	6.1	1.74	0.0
/ina c.l	6.2	1.56	0.0
Yolo 1	7.2	0.62	0.0
yar c	7.7	0.51	8.8
Kettleman c.l.	7.8	0.68	1.9
Meloland f.s.	8.3	2.69	4.2
Zaca c	7.4	0.74	7.7
Metz v.f.s.l.	7.5	0.75	0.7
Superstition l.f.s.	8.2	0.33	0.53
Viland s.	8.3	0.45	4.2
Cajon f.s.l.	7.7	0.63	0.7
Linne c.l.	7.4	0.87	3.6
Hovey c.	7.7	0.54	4.7
Naciemento c.l.	7.6	0.38	1.9
unnyvale c	7.8	1.22	3.0
Panoche l	7.7	1.17	1.6
Cijeras c.	7.4	114.00	7.2
Lost Hills c.	7.0	0.51	0.5
Fijeras I.	8.3	0.89	13.1

	TABLE 1									
SOIL SERIES	AND	TYPES	USED	FOR	\mathbf{Sr}	85	FIXATION	STUDY		

* Saturation extract at 25° C. † Expressed as CaCO₃.

Thus far, attempts to predict the distribution of Sr 90 between plants and soils have yielded far less satisfactory results. For example, the work of Menzel (1954) and of Romney *et al.* (1956) with soils reveals OR values of much less than unity. A number of reasons for this are conceivable. In the first place, Sr and Ca in soils may exist largely in the adsorbed or exchangeable state and the adsorption energy of Sr may be considerably different from that of Ca. For this reason, the elements will not be equally available to plants as is presumably the case in culture solutions. Moreover, it is probable that the relative adsorption energies of Sr and Ca will vary considerably from soil to soil. In the second place, the Sr and Ca extracted from soil by means of the commonly employed methods such as extraction with HC1 or NH_4Ac , electrodialysis, and fusion analysis often bear little relationship to their relative availabilities to plants. For example, calcium present in the

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soil as $CaCO_3$ may be only slightly available to crops, and yet such calcium is readily extracted from the soil with dilute acids or with NH_4Ac .

The question as to the manner in which added radio-strontium is retained in soil lies at the heart of the whole problem. Extremely insoluble compounds of strontium in which the element is unavailable to plants have been reported in some soils (Norrish, 1957). The work of Nishita *et al.* (1956) indicates that in many soils strontium is fixed in a nonexchangeable form. Kleehkovsky and Tselschcheva (1957), however, did not find such fixation in three Russian soils studied.

In the present research, Sr 85 was added to 26 soils from California. Following this, efforts were made to determine the fraction of the added Sr that was in an exchangeable form. The presumption has been that under appropriate conditions, exchangeable Sr is available to plants.

TABLE 2

ANALYSES OF NONCALCAREOUS SOILS USED IN Sr 85 FIXATION STUDY

Soil	pH 1.5 susp.	Soluble cations m.e. per 100 gms				Exchangeable cations m.e. per 100 gms				Exchange capacity m.e./
		Ca	Mg	Na	к	Ca	Mg	Na	к	100 gms (NH4Ac)
Aiken c.l.	5.8	0.00	0.00	0.00	C.00	9.12	3.24	0.08	0.70	17.40
Hanford v.f.s.l	7.9	0.37	0.09	0.07	0.05	15.3	2.90	0.38	0.60	12.2
Columbia v.f.s.l.	6.6	0.14	0.07	0.05	0.06	13.5	3.70	0.83	0.15	13.9
Fresno l.s.	7.4	0.46	0.29	0.12	0.07	4.04	1.12	0.27	0.47	3.5
Hanford s.l.	7.2	0.15	0.08	0.14	0.03	7.68	2.20	0.14	0.27	6.8
Yorkville l	6.2	0.04	0.05	0.04	0.01	7.10	6.40	0.04	0.26	13.4
Ramona s.l.	7.3	0.17	0.10	0.12	0.00	6.04	1.91	0.23	0.19	6.50
Tujunga v.f.s.l	7.3	0.41	0.17	0.11	0.05	11.9	2.74	0.36	1.15	12.6
Vina c.l.	6.5	0.15	0.17	0.04	0.00	16.2	13.2	0.25	0.24	26.12
Yolo 1	7.5	0.04	0.09	0.07	0.02	8.72	11.3	0.88	0.83	17.5

EXPERIMENTS

A number of experiments have been carried out in which soils contaminated with Sr 85 were treated and leached with neutral 1N ammonium acetate and 1N CaCl₂. Ammonium acetate of this strength is commonly used in the determination of the exchangeable cations in soils since NH_4 ion is not normally present. With calcareous soils the method leads to high results because alkaline earth carbonates are appreciably soluble in this reagent. Therefore 1N CaCl₂ was selected as an exchanging reagent because calcium carbonate is not soluble in this solution.

For the above studies, 26 soils from California were selected on the basis of their extent as mapped in soil surveys. They included representative soils over a wide range of chemical and mineralogical composition. Ten of the soils were noncalcareous; the pH of the saturated pastes varied from 5.8 to 7.9. Sixteen of the soils were calcareous and had a $CaCO_3$ content ranging from 0.5 to 13.1 per cent. Two soil samples each of the Hanford and Tijeras series were included; these were selected from widely separated locations. A list of the soil series used, together with the pH's of the saturated pastes, conductivity of the saturation extracts and lime content (expressed as per

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cent $CaCO_3$), is presented in Table 1. Additional chemical data concerning the noncalcareous soils are given in Table 2.

The leaching experiments were carried out in the following manner. Twenty-five grams of air-dry soil (pulverized to pass a 2 mm screen) was weighed into a 250 ml beaker and 50 ml of a carrier-free solution containing approximately 10 microcuries of carrier-free Sr 85 and 8×10^{-4} m.e. of HC1

TABLE 3 DATA SHOWING EFFICIENCY OF EXTRACTION OF SR 85 FROM SOILS BY LEACHING WITH 1N NH₄Ac AND 1N CaCl₂. SOILS WERE DRIED FOR ONE WEEK AT 50° C AND 110° C AFTER ADDITION OF Sr 85 AND PRIOR TO LEACH-ING. RESULTS ARE EXPRESSED AS PERCENTAGE OF ADDED Sr REMAINING IN SOIL AFTER LEACHING PROCEDURE

Soil	NH4Ac, soil dried at 50° C	NH4Ac, soil dried at 110° C	CaCl ₂ , soil dried at 110° C
Aiken c.l.	17.2	18.3	3.27
Hanford v.f.s.l.	8.68	13.2	8.87
Columbia v.f.s.l.	11.9	20.1	4.50
Fresno l.s.	3.75	5.10	2.67
Hanford s.l.	8.92	10.7	3.37
Yorkville l	5.94	7.97	2.01
Ramona s.l.	8.63	14.6	5,39
Tujunga v.f.s.l.	4.69	6.54	4.62
Vina c.l.	8.30	12.0	2.37
Yolo 1.	4.39	5.86	3.65
Ayar c	2.08	4.59	3.64
Kettleman c.l.	1.72	4.88	3.10
Meloland f.s.	1.02	4.21	3.61
Zaca c.	3.29	5.94	3.21
Metz v.f.s.l.	2.65	6.79	3.53
Superstition l.f.s.	0.83	2.94	2.48
Niland s.	1.11	4.33	4.74
Cajon f.s.l.	1.77	3.79	3.10
Linne c.l.	2.27	5.06	2.02
Hovey c	1.67	6.39	4.63
Naciemento c.l.	1.49	5.77	3.62
Sunnyvale c	5.20	5.75	4.49
Panoche l	3.09	8.30	5.26
Tijeras c	0.92	1.40	1.15
Lost Hills c	2.21	4.86	3.07
Tijeras l	1.44	3.17	4.00

were added. The resulting suspension was stirred, evaporated down, and the residue was dried at either 50° C or 110° C for one week. Following this, 100 ml of the NH_4Ac or $CaCl_2$ reagent were added and the mixture was heated on a steam bath for two hours during which time it was stirred occasionally. The suspension was then transferred to a Buchner funnel and leached with the exchanging reagent under gravity until 500 ml of leachate were collected.

Radio-assays of both the leachates and extracted soils were made from the strontium gamma radiation only, using a scintillation counter with a sodium iodide crystal. For radio-assays of the solutions, 2 ml aliquots were dried in cupped planchets of one inch diameter. The extracted soils were dried, ground, and mixed, and one gram samples were then assayed taking into account geometry and mass absorption. All experiments were performed at least in duplicate and in all cases the reproducibility was better than 5 per cent. The results of this line of study are presented in Table 3.

A second series of studies was carried out with the above soils in which a given weight of contaminated soil was repeatedly extracted with equal volumes of the NH_4Ac reagent. It was thought that this type of study would reveal any fixation of the Sr 85 in a form nonexchangeable with NH_4^+ ion.

For the repeated extraction procedure, 5 grams of air-dry soil were weighed into a 50 ml centrifuge tube and 2 ml of carrier-free solution containing approximately 10 microcuries of Sr 85 and 8×10^{-4} m.e. of HC1 were added. The soil was dried at 50° C for seven days and then suspended in 25 ml of neutral, normal ammonium acetate. After shaking for 10 minutes, the soil was centrifuged down and the supernatant liquid was decanted off. This extraction process was carried out 10 times with fresh portions of 1N ammonium acetate. The soils and extracts were assayed for Sr 85 as described above.

The percentages of added Sr 85 remaining in the various soils after 10 extractions with NH_4Ac , are presented in Table 4 and compared with the corresponding (50° C) percentages remaining after leaching with NH_4Ac .

The progress of radio-strontium removal upon successive extractions with ammonium acetate is shown graphically for nine of these soils in figures 1, 2, and 3.

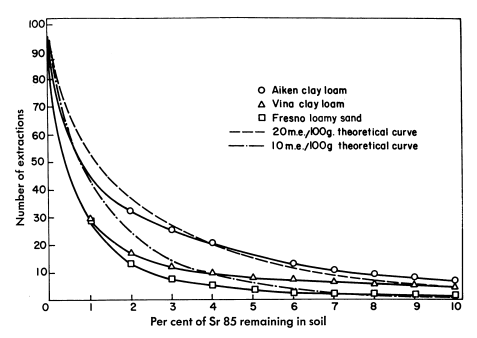


Fig. 1. Extraction of Sr 85 from Aiken, Vina, and Fresno soils with 1N NH,Ac

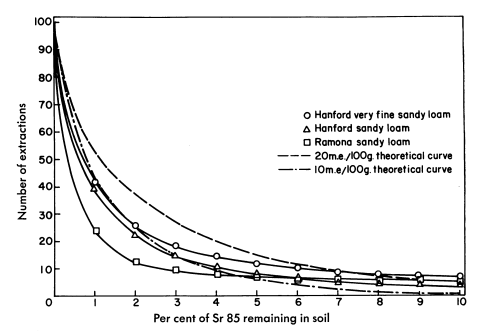


Fig. 2. Extraction of Sr 85 from Hanford and Ramona soils with 1N NH₄Ae

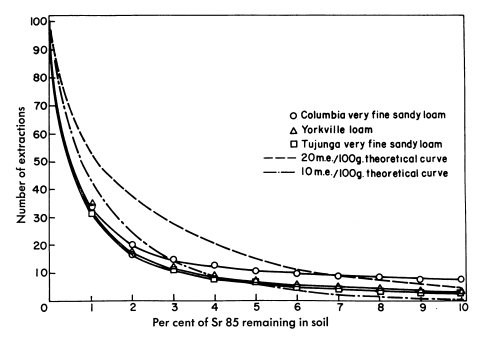


Fig. 3. Extraction of Sr 85 from Columbia, Yorkville, and Tujunga soils with 1N NH₄Ac

DISCUSSION AND CONCLUSIONS

In general, the major fraction of the Sr 85 added to the 26 Californian soils was extracted by the NH_4Ac and $CaCl_2$ leaching procedures adopted (see Table 3). This was true regardless of whether the soils had been dried at 50° C or at 110° C prior to leaching.

Certain of the noncalcareous soils dried at 50° C, namely, Aiken c.l., Han-

TABLE 4

PERCENTAGES OF ADDED Sr 85 REMAINING IN SOILS AFTER REPEATED EXTRACTION (10 EXTRACTIONS) WITH 1N NH₄Ac AND AFTER CONTINUOUS LEACHING WITH 1N NH₄Ac. ALL SOILS WERE DRIED AT 50° C FOR ONE WEEK AFTER ADDITION OF Sr 85 AND PRIOR TO TREATMENTS WITH NH₄Ac

Soil	Extracted 10 times with 1N NH4Ac	Continuously leached with 1N NH4Ac
Aiken c.l.	7.20	17.2
Hanford v.f.s.l.	6.50	8.68
Columbia v.f.s.l.	7.40	11.9
Fresno l.s.	1.80	3.75
Hanford s.l	3.30	8.92
Yorkville l	3.10	5.94
Ramona s.l.	5.40	8.63
Tujunga v.f.s.l.	2.80	4.69
Vina c.l.	5.20	8.30
Yolo l.	2.50	4.39
Ayar c.	2.08	2.08
Kettleman c.l.	2.11	1.72
Meloland f.s.	1.66	1.02
Zaca c	2.35	3.29
Metz v.f.s.l.	2.36	2.65
Superstition l.f.s.	0.99	0.83
Niland s.	1.54	1.11
Cajon f.s.l.	1.34	1.77
Linne c.l.	1.79	2.27
Hovey c	2.24	1.67
Naciemento c.l.	2.23	1.49
Sunnyvale c	3.46	5.20
Panoche l.	3.47	3.09
Tijeras c.	1.17	0.92
Lost Hills c.	1.75	2.21
Tijeras l	2.26	1.44

ford v.f.s.l., Columbia v.f.s.l., Hanford s.l., Ramona s.l., and Vina c.l., retained unexpected amounts of Sr 85 against the leaching with NH₄Ac. Also, the effect appears to have been enhanced by drying at 110° C. It is unlikely, however, that this is an actual fixation of the Sr in nonexchangeable form because the effect is not apparent when the soils are leached with CaCl₂, even after drying at 110° C. Moreover, considerably less Sr 85 is retained by these soils when repeatedly extracted with NH₄Ac (see Table 4).

The data of Table 4 indicate that the repeated NH_4Ac extraction procedure adopted is generally more efficient than the NH_4Ac leaching procedure. The graphical presentation of the repeated extraction data (figures 1, 2, and 3) Hilgardia

reveals large differences between soils in the rate at which the Sr 85 is released in exchange for NH_4^* .

According to Vanselow's equation given above, the saturation of the soil with NH_4^+ would not reach completion during the period of elevated temperature on the steam bath. Also, the subsequent leaching on a Buchner funnel possibly would not remove all of the remaining exchangeable ions owing to the existence of regions which are inaccessible, or only slowly accessible, to the leaching solution. In the repeated extraction procedure, however, the soil mass is shaken between each extraction, bringing all of the surface into contact with fresh solution.

It is reasonable to suppose that the $CaCl_2$ leaching is more efficient because the soil becomes essentially Ca++ saturated during the period of equilibration on the steam bath, and any subsequent channeling on the Buchner funnel is insignificant. There are apparently two reasons for this: in the first place, Ca++ is a much stronger replacing agent than NH_4^+ , and in the second, the soils are initially high in exchangeable Ca++.

The dotted theoretical curves in the figures emphasize that the rate of release of the Sr may depend markedly on the exchange capacity of the soil. The theoretical curves were calculated assuming that the applied Sr is exchangeable and that Vanselow's exchange equation applies (Vanselow, 1932); that is,

$$\frac{m_{{\rm Sr}^{++}} \cdot N_{{\rm NH}_{\star}^{+}}^{2}}{m_{{\rm NH}_{\star}}^{2} \cdot N_{{\rm Sr}^{++}}} = K_{v} \cdot \frac{(\gamma_{{\rm NH}_{\star}^{+}})^{2}}{\gamma_{{\rm Sr}^{++}}}$$

where m is the solution molality, N the mole fraction of the adsorbed ion, K_v the exchange constant, and γ the ionic activity coefficient in solution. For the calculations, the value of

$$K_v \cdot rac{\left(\gamma_{\mathrm{NH}_{\bullet}^+}
ight)^2}{\gamma_{\mathrm{Sr}^{++}}}$$

was taken as 10^{-4} . The curves were calculated for exchange capacities of 10 and 20 milliequivalents per 100 grams of soil.

The experimental and theoretical curves of figures 1 to 3 show further that complete removal of the added Sr 85 is approached asymptotically as the number of extractions is increased. Moreover, they point up the danger that the selection of a limited extraction procedure may lead to erroneous values for the amount of Sr held in a nonexchangeable form. For the soils under study, it is clear from the figures that in no case has the slope of the extraction curve reached a value of zero even after the tenth extraction. Thus it cannot be assumed that any of the added Sr 85 is retained in a nonexchangeable form.

The authors are led to conclude, on the basis of the evidence presented here, that essentially all of the Sr 85 added to the 26 California soils was retained in an exchangeable or water soluble form. This appears to be the case whether the soil is calcareous or noncalcareous or whether the soil is dried at an elevated temperature after addition of the Sr. This conclusion, however, does not preclude the possibility of a long-term aging effect whereby the Sr isotope is slowly converted to a nonavailable form. If such an aging effect exists, it is probable that months or even years will be required before it becomes noticeable in the soils studied in this research.

The consequent conclusion that Sr 90, added to soils by way of fallout, is probably retained in an exchangeable form and hence available to plants has important implications concerning the use of $OR_{plant-soil}$ values. That is, the generally observed inconstancy of such values must be due largely to variations in the relative adsorption energies of Sr and Ca from soil to soil. In other words, $OR_{plant-soil}$ values based on the ratio of exchangeable soil Sr to exchangeable soil Ca will not assume a particular figure and thus cannot be used to supply information concerning the uptake of Sr 90 by plants.

A solution to the problem is conceivable. Where the free electrolytic concentration of the soil is low, the $\frac{Sr}{Ca}$ ratio in a water extract of the soil is to a good approximation a measure of the activity ratio of Sr and Ca in the soil. It is also entirely possible that uptake of Sr from soil may depend on this activity ratio rather than on the ratio of exchangeable soil Sr to exchangeable soil Ca. If this should prove to be the case, the calculation of the value

$$OR_{plant-soil} = \frac{\frac{Sr 90}{Ca} \text{ in plant}}{\frac{Sr 90}{Ca} \text{ in water extract of soil}}$$

would be appropriate and could be used in assessing the Sr 90 availability in soils. Experiments bearing on this question are presently being conducted in this laboratory.

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