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- Derjaguin, B. V., Churaev, N. V., Fedyakin, N. N., Talaev, M. V., and Ershova, I. G., *Izv. Akad. Nauk SSSR*, (Russ.), **10**, 2178 (1967); *Bull. USSR Chem. Sci.* (Eng.), 2095 (1967).
 Henniker, J. C., *Rev. Mod. Phys.*, **21**, 322 (1949).

- Bangham, A. D., and Bangham, D. R., Nature, 219, 1151 (1968).
 Anisimova, V. I., Derjaguin, B. V., Ershova, I. G., Lychnikov, D. S., Rabinovich, Ya. I., Simonova, U. K. L., and Churaev, N. V., Russ. J. Phys. Chem., 41, 1282 (1967).
 Fedyakin, N. N., Derjaguin, B. V., Norikova, A. V., and Talaev, M. V., DAN SSSR, 165, 862 (1965).

Exchangeable Acidity in Unburnt Colliery Spoil

The more usual methods of soil extraction and chemical analysis are not entirely applicable to assessment of the potential of unburnt colliery spoil as a medium for plant growth. This is particularly true for spoil heaps in the West Riding of Yorkshire with surface spoil pH values well below 4.5. Such spoil often contains FeS2 which oxidizes to produce Fe++, SO₄-- and H+. It may also contain the carbonate mineral ankerite which is potentially capable of neutralizing the acidity. These reactions produce large quantities of soluble salts, chiefly sulphates.

Three sites in the West Riding of Yorkshire have been investigated to obtain information on the exchangeable cation status in the surface spoil. These sites were Bull-croft (SE535098), Hound Hill (SE337046) and Maltby Main (SK545925), and the surface age of the spoil was 10 yr, 50 yr and 8 yr respectively. Twelve random samples of surface spoil (0-10 cm) were taken, each yield-Twelve random ing about 5 kg of spoil, and each sample was air-dried before sieving through a 1 cm diameter riddle. determinations were carried out on separate 10 g subsamples of the ≤ 1 cm fraction.

Exchangeable acidity was estimated by the BaCl₂triethanolamine method1 and total cation exchange capa-The method described by city by NH₄+ saturation². Jackson³, in which the spoil is treated with 0.1 N HCl, was used as a measure of acid extractable cations. Where ankerite is present in the spoil, this procedure gives a measure of its potential neutralizing capacity. The pH was measured using a glass electrode in a 2:1 0·1 M CaCl2spoil suspension.

Table 1. Mean values and significant difference ($P\!=\!0.05$) for certain spoil factors for three sites

Site	Total C. E.C.	Exch. acidity	Acid extr. cation	$p\mathbf{H}$
Bullcroft	13.22	19.75	-0.33	2.76
Hound Hill	10.88	0.03	46-42	3.43
Maltby Main	10.64	6.70	99.00	3.75
lsd(P=0.05)	0.94	2.09	35.81	1.25

Table 1 shows mean values for the four sets of determinations for each site, together with the significant difference between means (P = 0.05).

Fig. 1 shows the regression of the ratio of the exchangeable acidity to total cation exchange capacity on acid extractable cations and shows a consistent relationship between the degree of saturation of the ion exchange sites

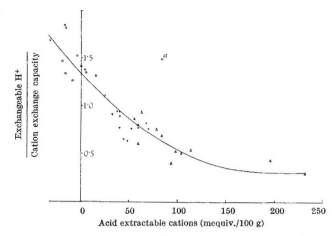


Fig. 1. The relationship between the exchangeable acidity: total cation exchange capacity ratio and acid extractable cations for unburnt spoil samples from three sites. \bigcirc , Bullcroft; +, Hound Hill; \triangle , Maltby Main. Point a has been omitted in calculating the regression: $y=1.341-0.0109x+0.0000288x^3$.

with acidic ions, and the potential neutralizing capacity of the spoil. It also indicates that some spoil (Bullcroft) has no net neutralizing capacity. Field experience indicates that this site is particularly inhospitable to plant establishment and growth. The values recorded for the exchangeable acidity of spoil may include acidic ions not adsorbed on exchange sites, hence the values of greater than unity for the saturation ratio. It has been established (unpublished work of M. E. P.) that the detrital fraction of the spoil contains significant quantities of amorphous aluminium hydroxides which may become a source of additional acidic cations in pyritic spoil.

Confirmation of a simple relationship such as this may help to identify problem sites which are unlikely to respond to conventional amelioration treatments for reclamation.

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- Chapman, H. D., and Pratt, P. F., in Methods for Analysis of Soils, Plants and Waters (Calif. Univ., 1961).
 Black, C. A., Evans, D. D., White, J. L., Ensminger, L. E., and Clark, F. E., in Methods of Soil Analysis (Amer. Soc. of Agronomy, Madison, 1965).

³ Jackson, M. L., in Soil Chemical Analysis (Constable, London, 1958).

Quenching of the Fluorescence of the Cerous Ion in Aqueous Solution by some Organic Ligands

The fluorescence of the cerous ion in aqueous solution has been the subject of a number of studies 1-3. The emission maximum occurs at about 355 nm, this being excited by absorption at about 265 and 305 nm. The fluorescence is little affected by temperature in the range 10°-40° C, or by strong acids. Nitric acid, however, oxidizes the cerium to the non-fluorescent ceric state. The fluorescence is also affected by the presence of the ceric ion1 and of some other cations2.