

Acid Rain Research

REPORT 19/1989

Interpretation of Soil Data from
the RAIN Project



NIVA - REPORT

Norwegian Institute for Water Research



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Report No.:
0-82073
Sub-No.:
Serial No.:
2294
Limited distribution:

Report Title:	Date:
Interpretation of Soil Data from the RAIN Project.	November 1989
	0-82073
Author (s):	Topic group:
J.O. Reuss	Acid precipitation
	Geographical area:
	Norway
	Number of pages (incl. app.)

Contractor:	Contractors ref. (or NTNF-No)
MD, NTNF, Environment Canada, Environment Ontario, SNV (Sverige), CEGB (UK)	NTNF-No.: FK03

Abstract:

Process-oriented models for prediction of soil and water acidification at the catchment scale require soil chemical and physical data as input parameters. Choice of soil properties characteristic for an entire catchment poses a major dilemma. This report deals with the interpretation of analyses from about 200 soil samples collected 1983-86 from the 7 catchments included in the RAIN project (Reversing Acidification In Norway). Various statistical techniques are used to derive characteristic values for key soil parameters such as cation exchange capacity, base saturation, bulk density, and anion absorption.

4 keywords, Norwegian

1. sur nedbør
2. jordkjemi
3. reversibilitet
4. nedbørfelt

4 keywords, English

1. acid precipitation
2. soil chemistry
3. reversibility
4. catchment

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ISBN 82-577-1596-4

INTERPRETATION OF SOIL DATA FROM THE RAIN PROJECT

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30 October 1989

INTRODUCTION

Process-oriented models for prediction of soil and water acidification at the catchment scale require soil chemical and physical data as input parameters. Such data are generally derived from soil analyses. Because of the natural heterogeneity of soil within catchments, the choice of values for soil properties characteristic for an entire catchment poses a major dilemma. Nevertheless such characteristic values must be specified to apply predictive models.

This report deals with the interpretation of soil analyses for the 7 catchments included in the RAIN project (Reversing Acidification In Norway) (Wright et al. 1988). These interpretations generally focus on the derivation of characteristic values for those soil parameters required for modelling purposes, particularly those required by MAGIC (Model for Acidification of Groundwater In Catchments) (Cosby et al. 1985). However, in some cases material of more general interest has been included.

Samples were collected and analyzed under the supervision of Dr. Erik Lotse of the Swedish Agricultural University at Uppsala. Methods of collection and analyses and data compilations are given by Lotse and Ottabong (1985) and Lotse (1989).

PART I reports interpretation of samples from the Sogndal site, while results from the Risdalsheia site are given in PART II. Anion adsorption results from both sites are reported separately in PART III.

PART 1. SOGNDAL

1. RELATIONSHIPS WITH CARBON CONTENT

1.1 Loss on Ignition versus Percent Carbon

A very close relationship (Figure 1.1) between loss on ignition and percent carbon is apparent. One outlier has been deleted from both the graphs and regressions involving carbon. This point was also inconsistent on the graph of CEC versus carbon, suggesting an error in the carbon determination. The regression equation is,

$$y = 2.97 + 1.97x \quad r^2 = 0.950 \quad n = 83 \quad (2.1)$$

This close relationship suggests that the OM lost on ignition consistently contains about $1/1.97$ or 51% carbon, slightly below the classical 58% but well within the range expected for surface soils (Jackson, 1958). The intercept seems a little high but it is probably affected by the scatter at the high points.

1.2 CEC(KCl) versus Percent Carbon

The relationship between neutral salt (KCl) pH and percent carbon is shown in Figure 1.2. The regression equation is,

$$y = -0.526 + .555x \quad r^2 = 0.779 \quad (2.2)$$

This not only indicates a close relationship, but perhaps more importantly shows that CEC is almost exclusively of organic origin. The scatter is mostly to the upper right of the graph suggesting that the errors may not be normally distributed about the linear regression line. However, there is probably little to be gained by using a log or log-log transform. The regression indicates a CEC at soil pH of about 0.56 meq/100 g soil for every 1% carbon, very much below the 4.9 meq/100 g suggested by Jackson (1958).

This suggests that at soil pH most of the CEC (nearly 90% if we accept the 4.9 figure) is not available due to protonation of the sites. The fact that virtually all charge is organic apparently is due to a minimum of soil development, at least as far as clay formation is concerned. In graph 2.2 the points are differentiated by depth, showing the generally lower CEC (due to lower carbon) in the deeper samples.

1.3 Base Saturation versus Percent Carbon

This relationship (Figure 1.3) is highly significant with an r^2 of 0.279 and 82 degrees of freedom. However, while the relationship may have a real component, a good part of it may be spurious. The high base saturation values fall at the low carbon values where CEC is low and thus analytical error is relatively high. There seems to be a differentiation between depths in the pattern. The implications of this have not been fully explored. The base saturation as calculated here includes Ca^{2+} , Mg^{2+} , Na^+ , K^+ and NH_4^+ . The MAGIC model does not use NH_4^+ and requires separate consideration of the other four cations.

1.4 pH versus Carbon

This relationship (Figure 1.4) is also highly significant. The linear regression equation is,

$$y = 5.35 - 0.0292x \quad r^2 = 0.366 \quad (2.3)$$

Again the errors are not uniformly distributed. The pattern seems to reflect the lower base saturation at high carbon levels. These lower base saturation points in turn are associated with low pH. These high carbon and low pH points tend to be concentrated in the 0-15 cm depth. Thus the general pattern is one of greater acidity associated with high organic matter. It appears that as organic

matter accumulates there are insufficient bases to saturate the sites and the system becomes more acidic.

2.0 CATION EXCHANGE CAPACITY

In order to estimate either a single value of base saturation (BS) for the catchment or the site, or two values if broken out by depth, we must aggregate the CEC. At present no distinction is made among catchments. This assumes that there is no difference in cation exchange capacity among catchments. No such difference appears using ANOVA techniques on either the raw data or logarithmic or square root transformed data. Differences among depths are much greater than any differences among catchments.

Use of arithmetic means of CEC (meq/100 g) introduces a problem in that the effect of a few high values (where the variance is high) totally overshadows the effect of all lower values. Also the median values are well below the means, indicating skewness. Estimates of central tendencies were made using the raw data, as well as logarithmic and square root transformations (Table 2.1). The log transforms seem to over-correct a bit, i.e. while the distribution is much closer to normal than the raw data, the means tended to be somewhat lower than the medians, and chi square lack of fit test is still significant in two of the three cases. None of the lack of fits were significant for the square root transform so this estimate of central tendency, i.e. (mean of the square roots)², was selected.

As an example, normal probability plots for CEC of all samples for both the raw data and a square root transform are shown in Figure 2.1. The square root transform under corrects on the low end in this case, while there is a

tendency for the log transform (not shown) to over correct at the high end.

Aggregated values and estimates of CEC are as follows:

	<u>Aggregated CEC</u>
All	3.66 meq/100 g
0-15 cm	5.52
>15 cm	2.27

The P values in Table 2.1 are from a Chi square test using frequency histograms. This procedure was found to be highly sensitive to the number of divisions selected and was discontinued.

Another problem arises with this CEC data, in that the 1983 values are higher than those for 1984-1987, and this difference is highly significant. This is illustrated by the means plot for the square root transform (Figure 2.2). Estimates for the square root transform after dropping the 1983 data are also shown in Table 2.1.

3. BASE SATURATION

The MAGIC model tracks four base cations, i.e. Ca^{2+} , Mg^{2+} , Na^+ , and K^+ . As these are handled individually in the model, the saturation, i.e. the fraction of the total charges occupied by the ion, are evaluated separately for each of these ions.

3.1 Ca Saturation

The Ca saturation was calculated individually for all samples, and the distributions examined using frequency histograms, and especially normal probability plots. Again, distributions were examined for logarithmic and square root transformations as well as the raw data. Means, medians, standard deviations, and estimates of central tendency were calculated in each case (Table 3.1). As used here, central tendency is the mean for the raw data, the geometric mean for the log transforms, and the

square of the mean of the transformed value for the square root transform. The total population of 91 samples used for this evaluation includes 44 from 0-15 cm, 41 from 15-30 cm and 5 from 15-45 cm. While many of the parameters varied substantially with depth, this was not the case for Ca saturation. However, for consistency with the other parameters results were calculated for all samples, for the 0-15 cm depth and, for samples from greater than 15 cm. Single value estimates for the catchments were taken from the total population.

Examples of normal probability plots and frequency histograms representing the 0-15 cm depth are shown in Figures 3.1 and 3.2. The untransformed data are obviously skewed toward the low end. Some skewness can be detected at the high end on the lognormal plots, while the skewness at the low end is not entirely eliminated when the square root transform is used. There is no clear preference between the transforms in the case illustrated, and this was often the case not only for the Ca saturation data, but also where the procedure was used for other parameters. The "N P Evaluation" column in Table 3.1 is a subjective evaluation from examination of the Normal Probability plots and was used as an aid in selecting the transformation used in the final summary. The lognormal distribution may be slightly superior to the square root for Ca saturation, but in the case of Mg saturation (Section 3.2), the square root transformation was better, and this was selected for use in the summary in both cases.

The use of a single value for Ca saturation derived from all observations over a four-year sampling period assumes that Ca saturation has not changed significantly over time and that any difference in Ca saturation due to treatment (acid addition) can safely be ignored. There is a significant year effect on Ca saturation (Figure 3.3), but it is due to low values in the first year. These low values apparently arise from the high CEC values observed in 1983

as noted in the previous section, as the extractable Ca values do not differ by years. Ca saturation values are determined from the square root transforms both with and without the 1983 data (Table 3.1). No significant difference was detected in Ca saturation between catchments or to a year-by-catchment interaction.

3.2 Mg Saturation

The Mg saturation values for the transformed and untransformed parameters are given in Table 3.2. In this case there are small but highly significant differences due to depth, with higher Mg saturation at 0-15 cm than in the deeper layers. Again, there are significant differences due to years (Figure 3.4), but the pattern is difficult to interpret, as the values observed in 1983 and 1984 are similar (approximately 0.044), with significantly lower values (0.029) in 1985 and intermediate values (0.036) in 1986. Interpretation is also made more difficult by the apparently higher CEC in 1983.

The extractable Mg values in 1983 are significantly higher than those for the later years (Figure 3.5). It is possible that this could represent some export from the treated catchments; however, there is no indication of an effect due to fields ($p = 0.24$), or a year-by-catchment interaction ($p = 0.79$).

The relationship of exchangeable Mg to Ca as shown in Figure 3.6 is also of interest. The slope of the zero intercept regression is 0.295 with an r^2 value of 0.65. This seems to be below the ratios of the output fluxes as reported by Wright et al. (1988), which range from 0.66 to 0.41, suggesting that the Mg is somewhat more mobile in the system than is Ca.

3.3 K Saturation

The depth effect is relatively minor for K saturation, so only one estimate was made for this parameter. Estimates

were arrived at in two ways, i.e. regression of exchangeable K on CEC, and estimates of central tendency from raw or transformed values of K saturation.

Where a close relationship exists, the saturation may be estimated by regressing the amount of the exchangeable ion on CEC. A plot of this relationship for K is shown in Figure 3.7. As can be seen from the figure, much of the variability arises from the 1983 values, so the regressions were calculated for the total data set and again without the 1983 values. The intercepts were found to be very nearly zero, so the final regression lines were calculated with zero intercept.

Regression parameters of interest include the following.

	<u>All data</u>	<u>Without 1983</u>
r^2	0.60	0.75
Slope	0.031	0.029
s.e.(slope)	0.0016	0.0013
s.e.(est)	0.092	0.052

While the fit is substantially improved by dropping the 1983 values, the slope, i.e. the estimate of K saturation is very nearly the same, indicating a K saturation value of about 0.03. The regression line shown is for the 1984-86 data only. This procedure has the disadvantage that it may be unduly influenced by a few high values; however, from the distribution of the points this does not seem to be a problem.

When the K saturation data were examined directly, it again became apparent that the 1983 data were highly variable, so these were dropped along with one other extreme outlier. The resulting normal probability plot for the square root transform is shown in Figure 3.8. The estimate of K saturation derived from the square root mean is 0.032, with a 95% confidence interval of 0.028 to 0.036. The

untransformed data were clearly skewed, however the mean of 0.034 was not greatly different. From the above considerations a value of 0.030 was arrived at as a general estimate of K saturation.

3.4 Na saturation

The values for Na saturation are very low, with a mean of 0.030 and a median of 0.017. Because of the low values the relative error is high. The difference between mean and medians again indicates substantial skewness, with the average heavily affected by a few high values. Also the 1983 samples were quite different from those in later years, as shown by the higher values and more scatter when Na is plotted against CEC (Figure 3.9). If Na is regressed against CEC the intercept is highly significant, and the slopes are higher when the regression is forced through zero rather than if an intercept is allowed (Table 3.3).

Slopes range from 0.007 to 0.018 depending on the assumptions. Furthermore, there is a tendency towards non-linearity when the anomalous 1983 data are dropped. Due to these difficulties regression methods were discarded for calculating Na saturation.

The skewness in the Na saturation data cannot be overcome by a square root transformation. On the basis of the Normal Probability plot the log-normal distribution was selected, so the population estimate is the geometric mean. The 1983 data had less effect on this estimate than on the regression estimates, the estimate for all observations is 0.024, and without 1983 the estimate is 0.022, with 95 percent confidence interval of about 20% of the estimate. Considering the uncertainties in the various methods this estimate was rounded to a value of 0.02.

4.0 DENSITY

4.1 Estimates of Density

The aggregation of density was handled in a very similar manner to the other parameters discussed above. The 1983 sampling was again anomalous, and was discarded. Apparently it was based on wet weights while the 1984-86 determinations were on dry weights. Again, distributions were highly skewed, and means, medians, and normal probability plots were examined before deciding on the most appropriate estimate. The lognormal distribution was again selected, and estimates represent the geometric means as shown in Table 4.1.

Estimates derived from untransformed means are about 10% higher than the geometric means, while square root transforms are intermediate.

4.2 Density Related Parameters

One interesting relationship is that between density and CEC. When density is regressed against CEC using a multiplicative model of the form,

$$Y = aX^b \quad (5.1)$$

we obtain the regression relationship,

$$\text{Density} = 0.8101(\text{CEC})^{-.358}, \quad (5.2)$$

with 67 observations and an r^2 of 0.82. This relationship is shown graphically in Figure 4.1. It apparently arises from the fact that both CEC and density are controlled almost entirely by the amount of organic matter.

Another interesting relationship is that between CEC by weight and CEC by volume. CEC by volume is a function of the measured CEC by weight and the density. Because CEC by weight is closely related to density, there is also a close relationship between CEC by volume and CEC by weight, as shown in Figure 4.2. This relationship is non-linear, and can be described by,

$$\text{CEC(vol)} = 8.125\text{CEC(wt)}^{-.6375}, \quad (5.3)$$

with an r^2 of 0.935. Even though density decreases with increasing CEC(wt) this is not sufficient to compensate for the higher cation exchange capacity of the high carbon-low density organic matter. Thus, CEC by volume increases with increasing CEC(wt).

5.0 SUMMARY

A summary table (5.1) has been prepared for the Sogndal catchments showing the best estimates of the aggregated values of the parameters required by the MAGIC model. While these may change with further analysis of these or later data, the modifications are likely to be relatively small.

Table 2.1. Sogndal. Estimates of central tendency for cation exchange capacity (meq/100 g). Lack of fit by chi square.

<u>Depth</u>	<u>Mean</u>	<u>Median</u>	<u>s.d.</u>	<u>Central Tendency</u>	<u>Lack of fit (P)</u>
Raw Data:					
All	4.599	3.43	4.19	4.60 ¹	0.000****
0-15 cm	6.441	5.40	4.66	6.44	0.053*
>15 cm	2.875	1.79	2.80	2.88	0.000****
Log Transform (base 10)					
All	0.436	0.54	0.50	2.73 ²	0.066
0-15	0.617	0.73	0.20	4.44	0.189
>15	0.238	0.25	0.46	1.73	0.17
Square Root Transform					
All	1.914	1.85	0.97	3.66 ³	0.124
0-15	2.349	2.32	0.97	5.52	0.505
>15	1.507	1.34	0.75	2.27	0.12
Square Root Transform (Without 1983)					
All	1.678	1.59	0.89	2.82 ³	na
0-15	2.112	2.14	0.89	4.46	na
>15	1.281	1.00	0.69	1.64	na

¹ Mean
² Geometric mean
³ Square of square root mean

Table 3.1. Sogndal. Estimates of central tendency and associated parameters for Ca saturation. N P evaluation is a subjective evaluation of the normal probability plot. Categories are Very Good, Good, Fair, and Poor.

<u>Depth</u>	<u>Mean</u>	<u>Median</u>	<u>s.d.</u>	<u>Central Tendency</u>	<u>N P Evaluation</u>
Raw Data:					
All	0.193	0.155	0.155	0.193 ¹	P
0-15 cm	.197	.169	.169	.169	F
>15 cm	.190	.145	.147	.190	P
Log Transform (base e)					
All	-1.843	-1.863	.638	.158 ²	G
0-15	-1.777	-1.777	.570	.169	F
>15	-1.904	-1.927	.691	.149	G
Square Root Transform					
All	.418	.394	.135	.175 ³	F
0-15	.428	.412	.119	.183	G
>15	.410	.381	.149	.166	F
Square Root Transform (Without 1983)					
All	.448	.409	.135	.201 ³	na
0-15	.456	.444	.115	.208	na
>15	.440	.389	.151	.194	na

¹ Mean

² Geometric mean

³ Square of square root mean

Table 3.2. Sogndal. Estimates of central tendency and associated parameters for Mg saturation. N P evaluation is a subjective evaluation of the normal probability plot. Categories are Very Good, Good, Fair, and Poor.

<u>Depth</u>	<u>Mean</u>	<u>Median</u>	<u>s.d.</u>	<u>Central Tendency</u>	<u>N P Evaluation</u>
Raw Data:					
All	.0423	.038	.021	.042 ¹	F
0-15 cm	.051	.047	.021	.051	G
>15 cm	.033	.030	.018	.033	P
Log Transform (base e)					
All	-3.157	-3.280	.533	.043 ²	G
0-15	-3.064	-3.058	.449	.047	F
>15	-3.546	-3.546	.502	.029	G
Square Root Transform					
All	.197	.194	.051	.039 ³	VG
0-15	.221	.217	.047	.049	G
>15	.175	.172	.045	.031	G
Square Root Transform (without 1983)					
All	.193	.187	.050	.037 ³	na
0-15	.217	.215	.044	.047	na
>15	.171	.171	.046	.0296	na
¹	Mean				
²	Geometric mean				
³	Square of square root mean				

Table 3.3. Sogndal. Regression parameters for the regression of exchangeable Na on CEC.

	With Intercept		Without Intercept	
	All points	w/o 1983	All points	w/o 1983
r^2	0.443	0.707	0.406	0.333
Constant	0.026	0.026	0	0
Slope	0.015	0.007	0.018	0.011
s.e.(slope)	0.0017	0.0006	0.0012	0.0006
s.e.(est)	0.069	0.016	0.070	0.024

Table 4.1. Sogndal. Geometric means and confidence intervals for density.

<u>Depth</u>	<u>n</u>	<u>Estimated Density</u>	<u>Confidence Interval (95%)</u>
All	68	0.618 kg/l	0.585-0.653
0-15 cm	33	0.511	0.479-0.546
>15	37	0.738	0.694-0.785

Table 5.1. Sogndal. Summary table of aggregated values of soil parameters for use in the MAGIC Model.

<u>Depth</u>	<u>Density</u>	<u>CEC</u>	<u>Base Saturation</u>			
			<u>Ca</u>	<u>Mg</u>	<u>K</u>	<u>Na</u>
cm	kg/l	meq/100g	-----Equivalent Fraction-----			
All	0.618	3.66	0.175	0.039	0.03	0.02
0-15	0.511	5.52	0.183	0.049	0.03	0.02
> 15	0.738	2.25	0.168	0.031	0.03	0.02

Loss on Ignition vs Carbon

Sogndal

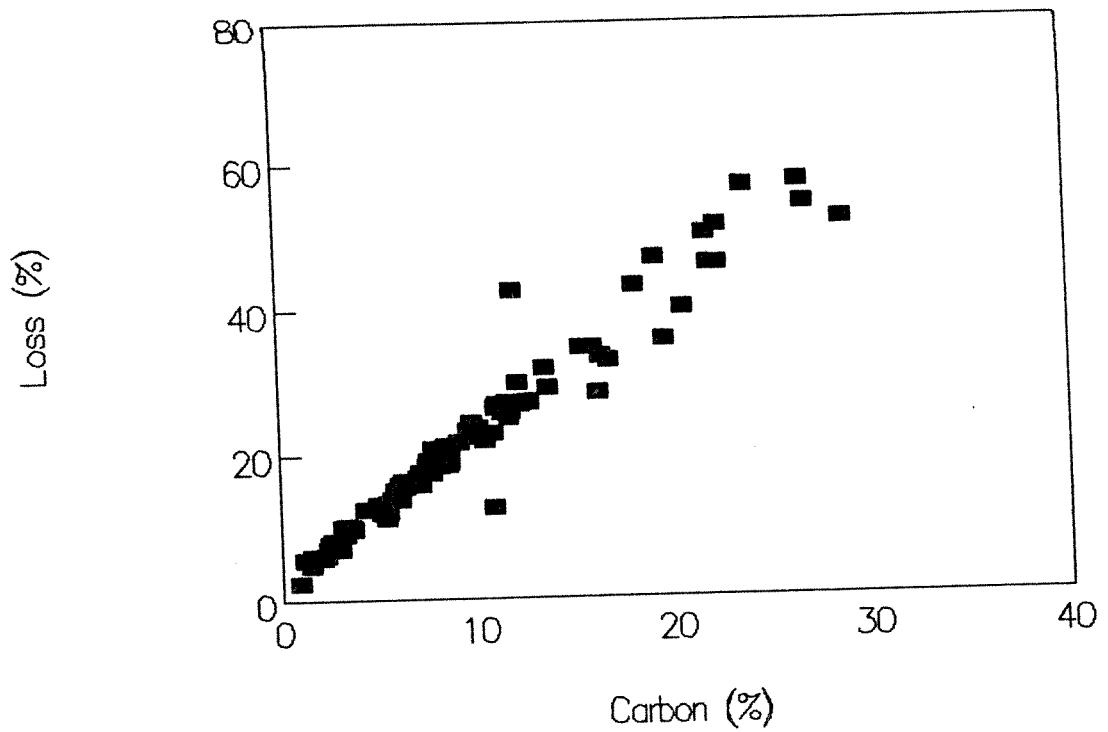


FIGURE 1.1. Sogndal. Loss on ignition as a function of percent carbon as determined by a chromic acid wet digestion.

CEC vs % Carbon

Sogndal

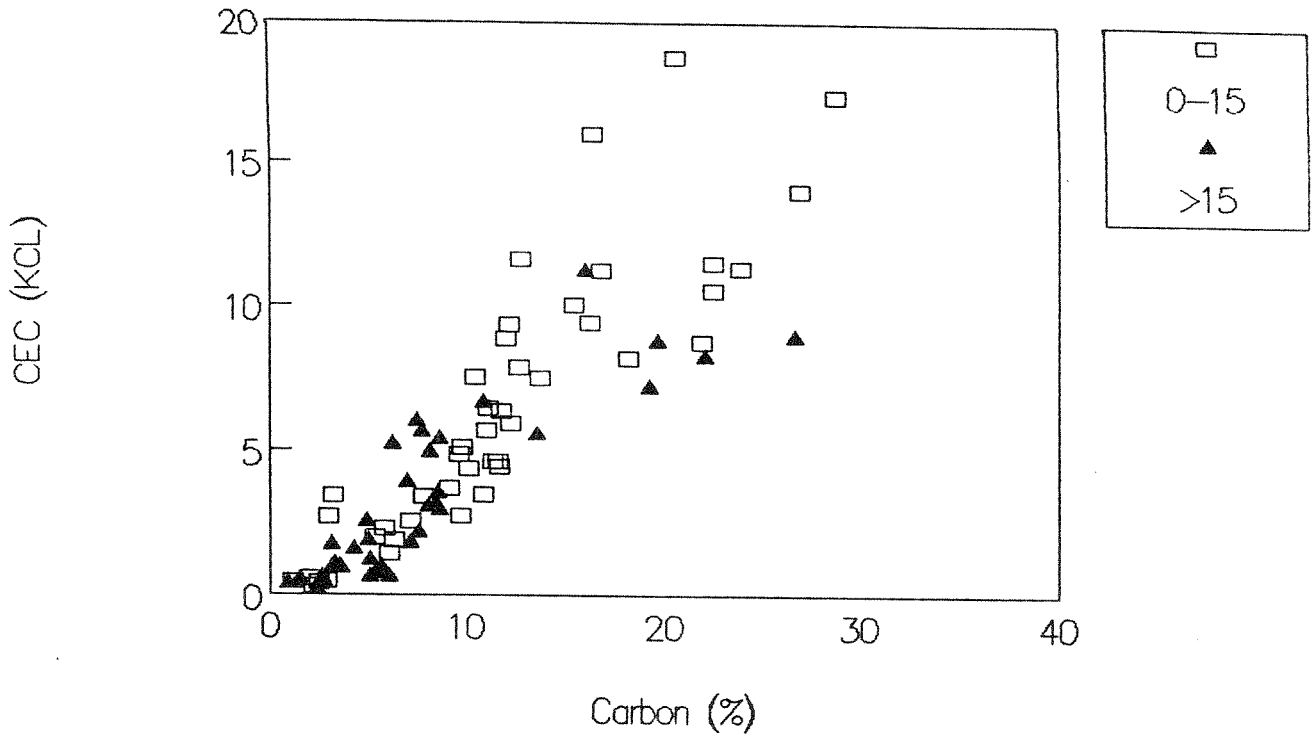


FIGURE 1.2. Sogndal. Cation exchange capacity (KCl) in meq/100g as function of percent carbon. Surface samples (0-15 cm) are shown by open boxes; deeper samples (>15 cm) are shown by triangles.

Base Saturation vs %C

Sogndal

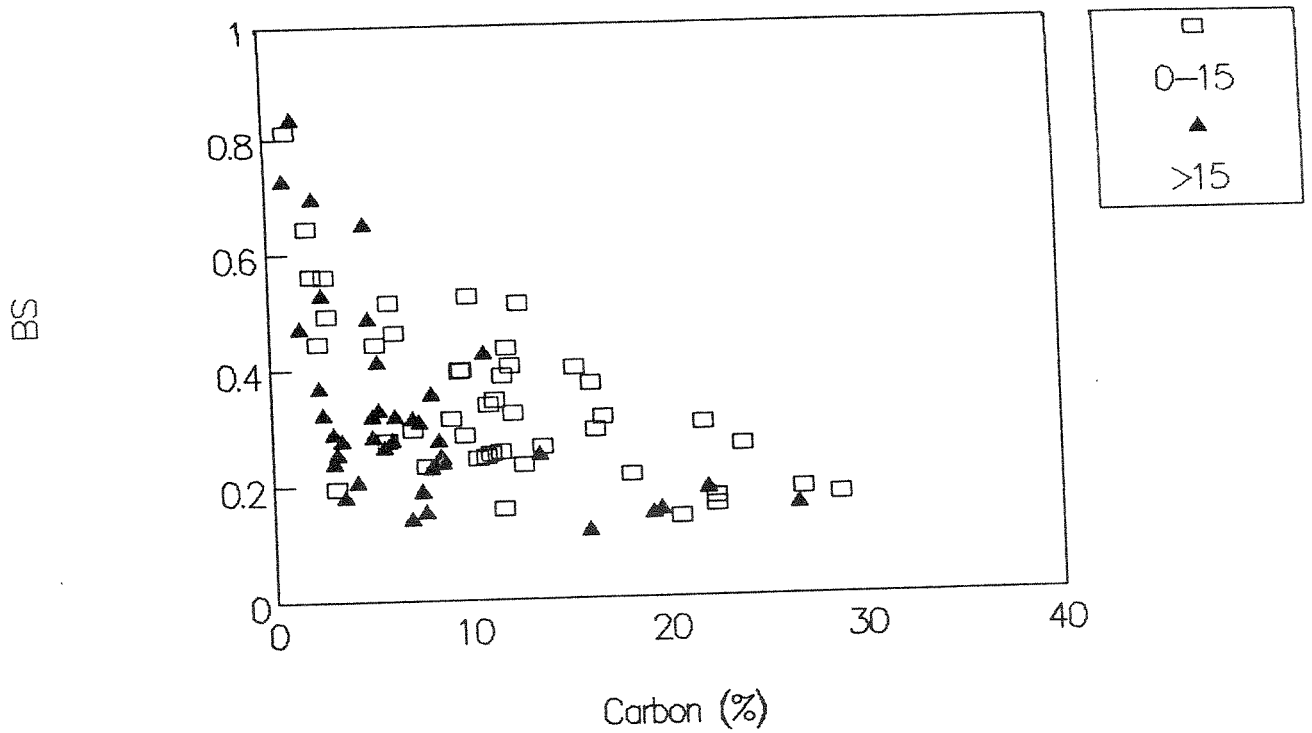


FIGURE 1.3. Sogndal. Base saturation as a function of percent carbon. Surface samples (0-15 cm) are shown by open boxes; deeper samples (>15 cm) are shown by triangles.

pH vs Percent Carbon

Sogndal

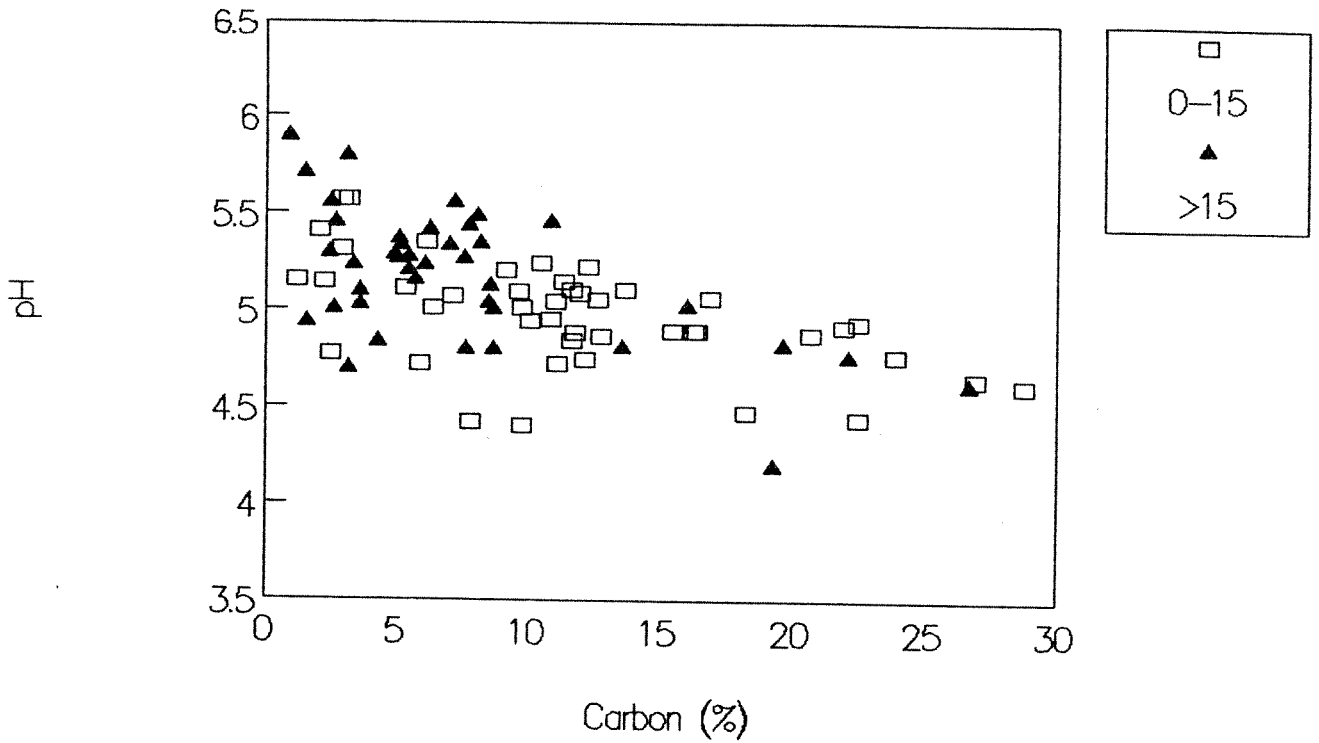
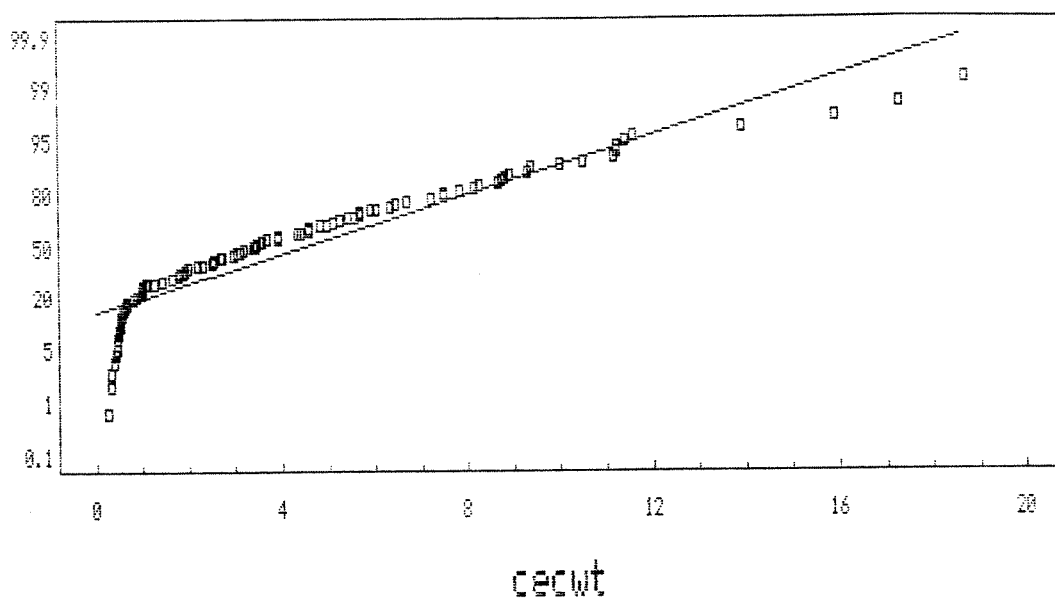


FIGURE 1.4. Sogndal. pH as a function of percent carbon. Surface samples (0-15 cm) are shown by open boxes; deeper samples (>15 cm) are shown by triangles.

Normal Probability Plot

cumulative percent



Normal Probability Plot

cumulative percent

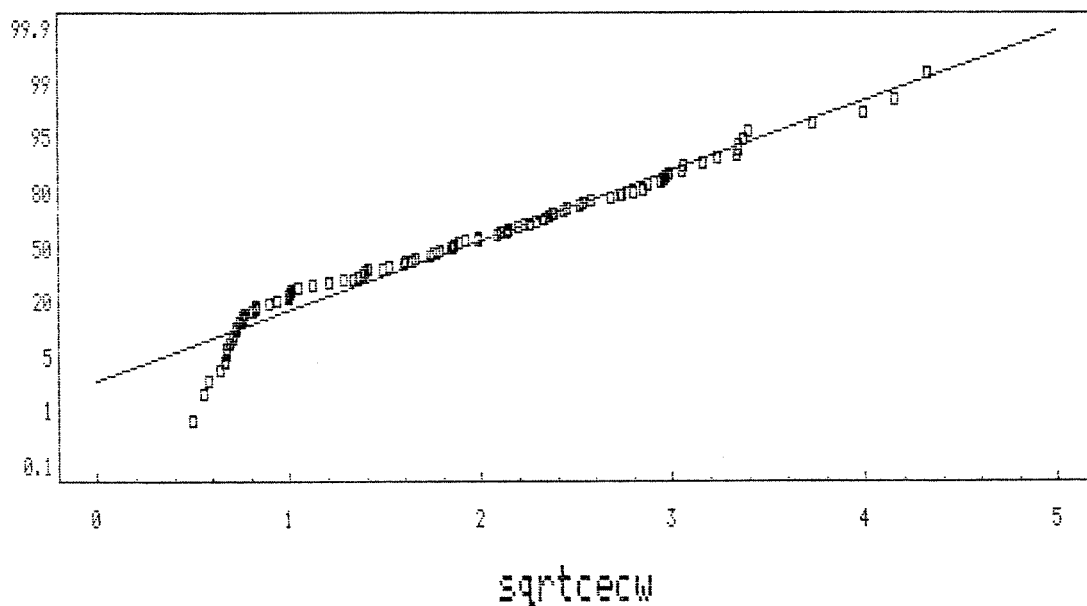


FIGURE 2.1. Sogndal. Normal probability plots for cation exchange capacity by weight, and for the square root transform of the same data.

95 Percent Confidence

Intervals for Factor Means

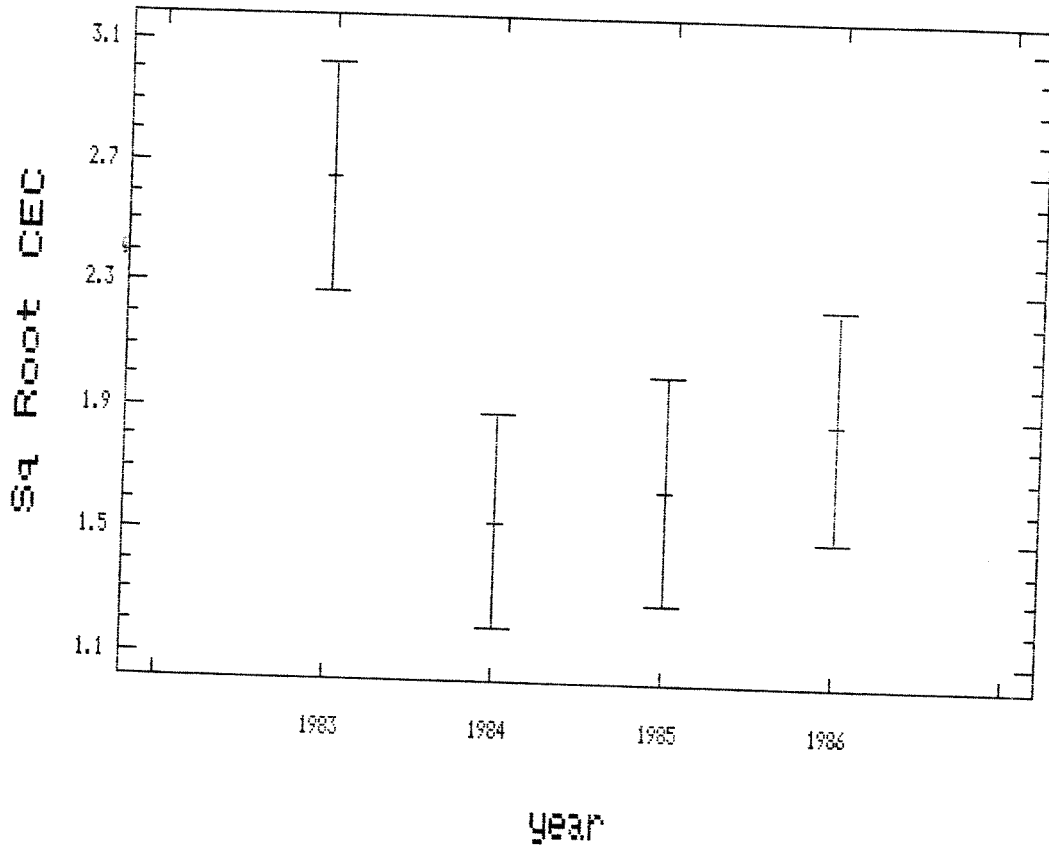
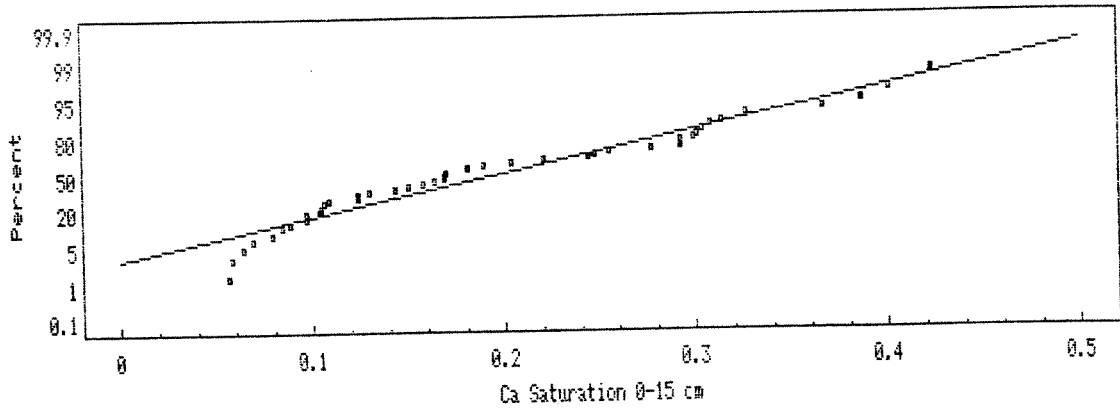
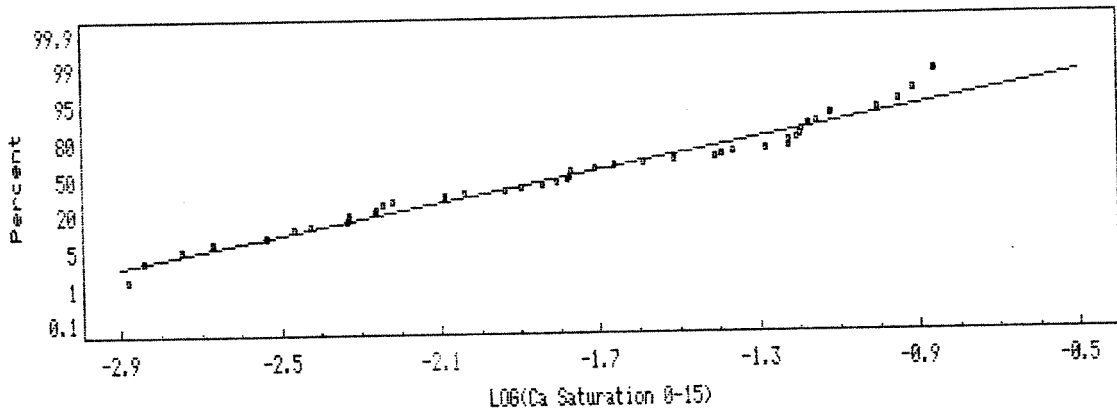


FIGURE 2.2. Sogndal. Means plot showing the 95% confidence interval for the square root transform of CEC by year.

Normal Probability Plot



Normal Probability Plot



Normal Probability Plot

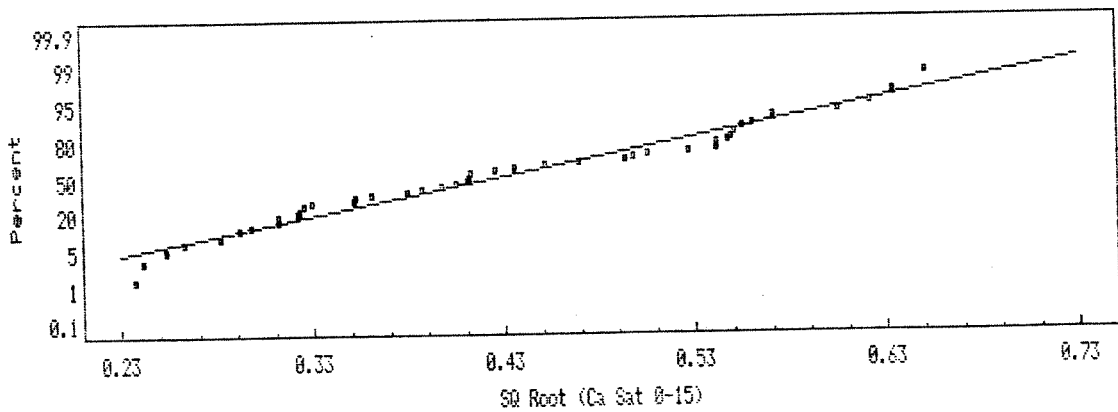


FIGURE 3.1. Sogndal. Normal Probability plots for the Ca saturation data from the 0-15 cm depth. Plots are shown for the raw data, log transform, and square root transform.

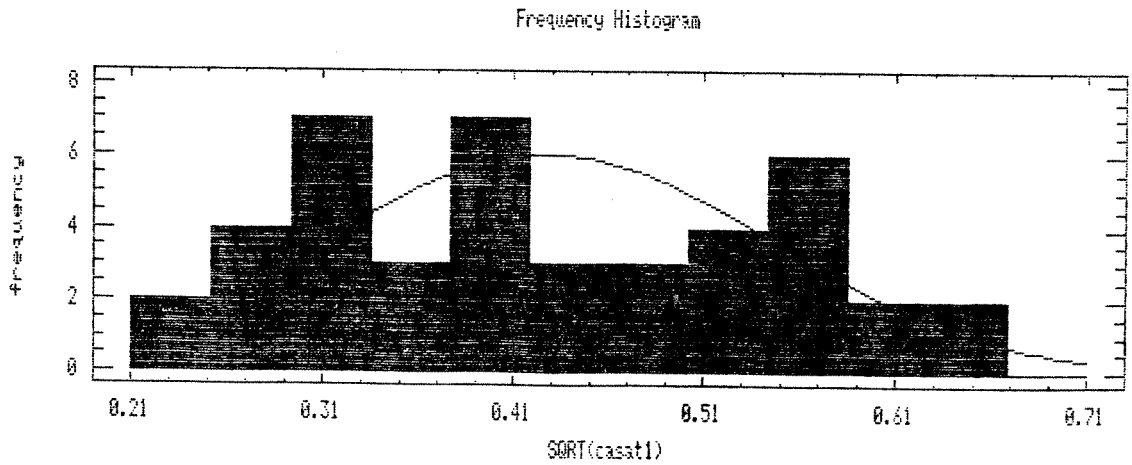
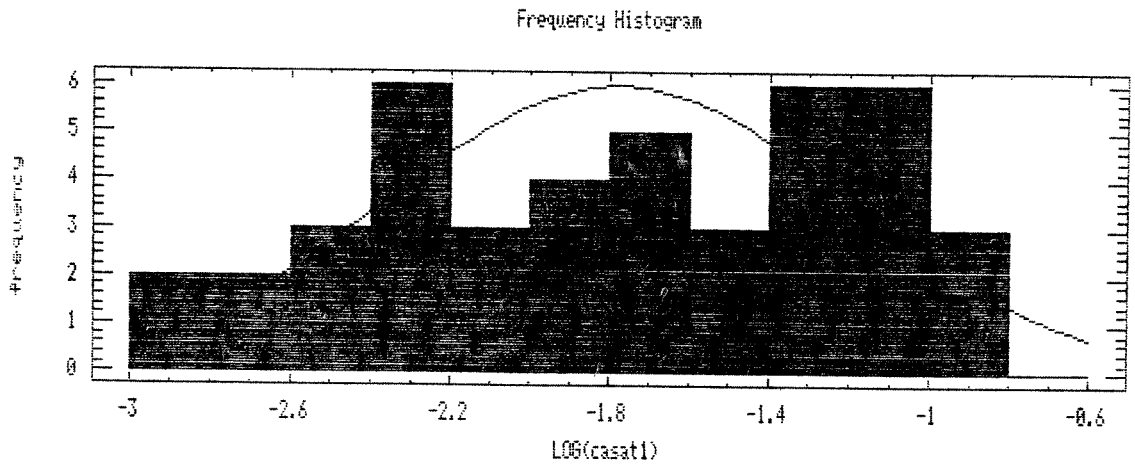
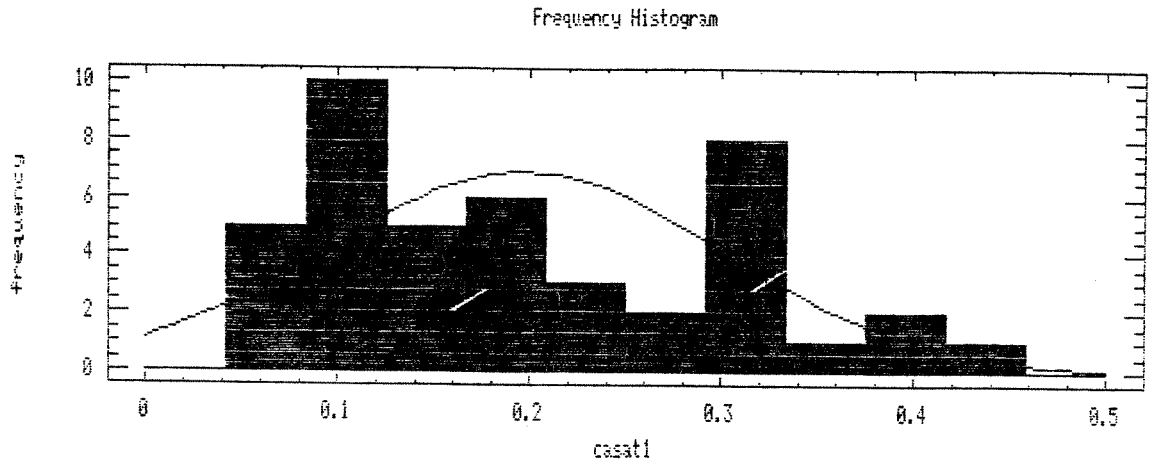


FIGURE 3.2. Sogndal. Frequency histograms for the Ca saturation data from the 0-15 cm depth. Histograms are shown for the raw data, log transform, and square root transform.

95 Percent Confidence

Intervals for Factor Means

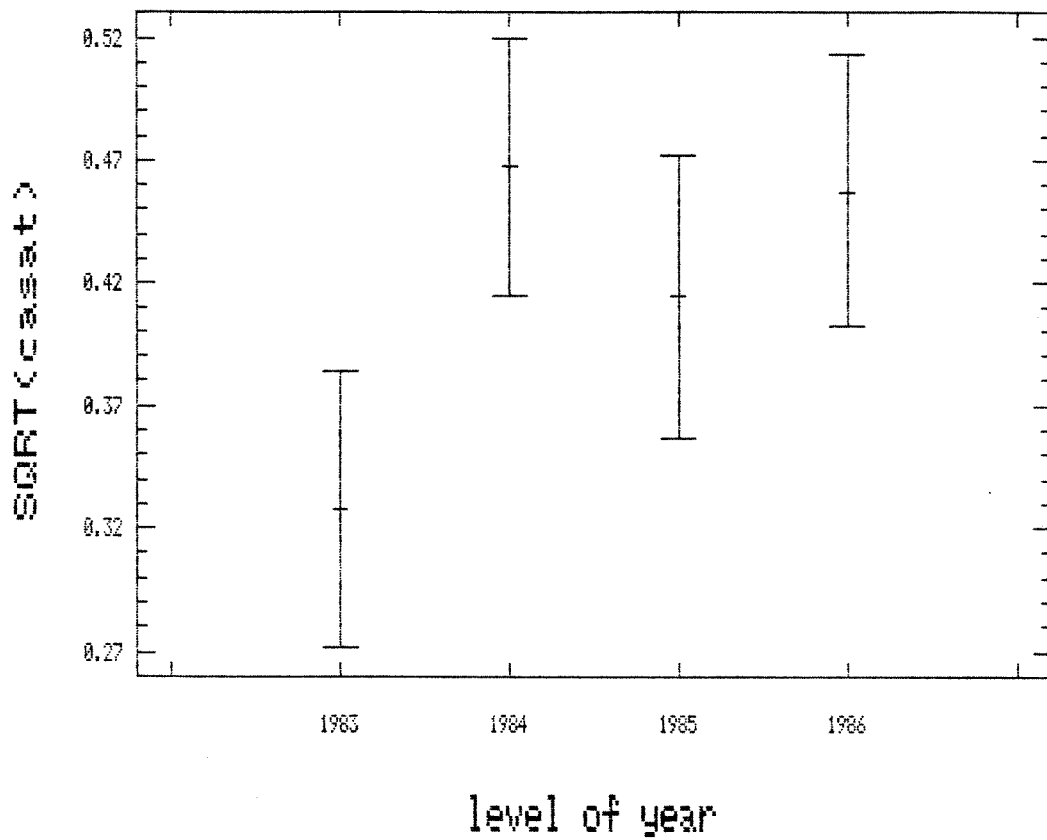


FIGURE 3.3. Sogndal. Means plot showing the 95% confidence interval for the square root transform of Ca saturation by year.

95 Percent Confidence

Intervals for Factor Means

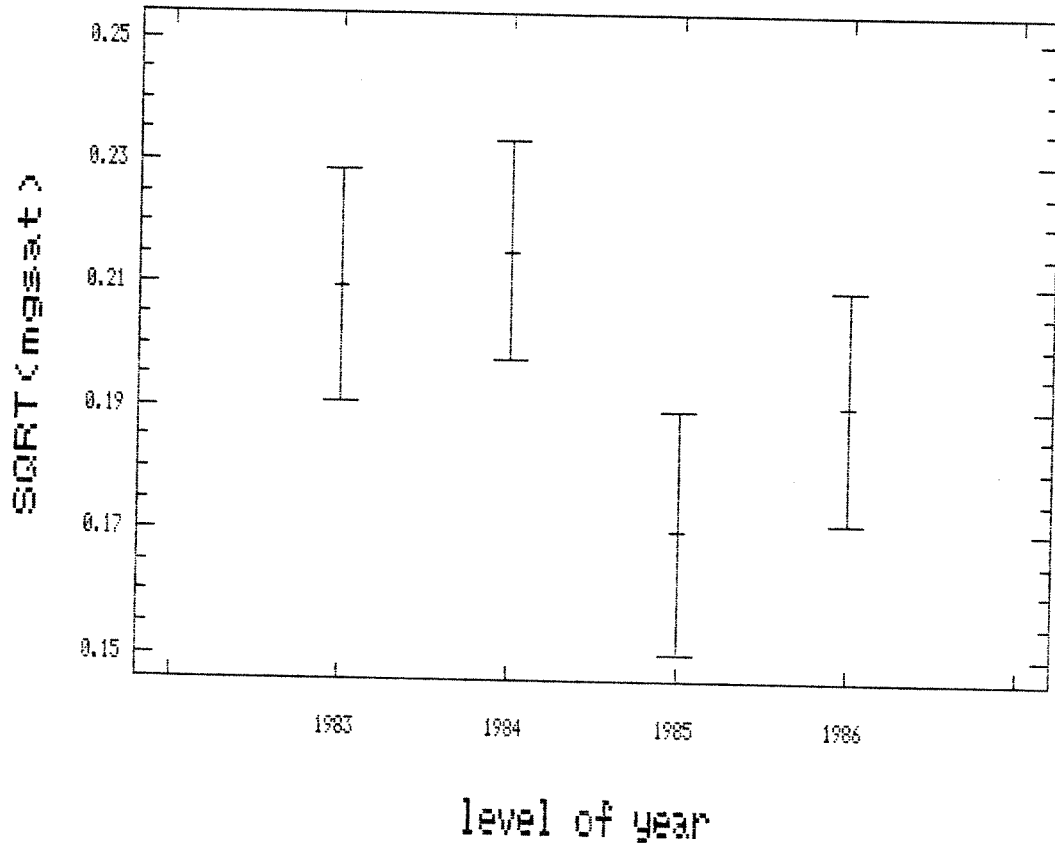


FIGURE 3.4. Sogndal. Means plot showing the 95% confidence interval for the square root transform of Mg saturation by year.

95 Percent Confidence

Intervals for Factor Means

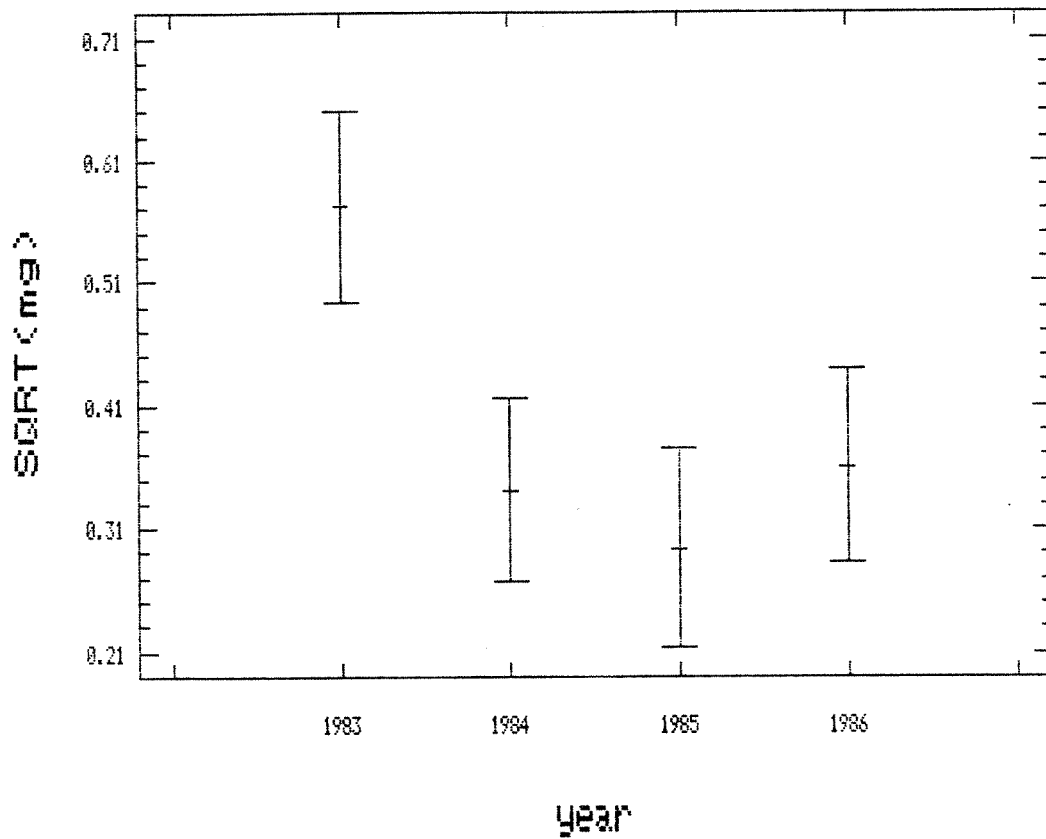


FIGURE 3.5. Sogndal. Means plot showing the 95% confidence interval for the square root transform of exchangeable Mg (meq/100g) by year.

Mg vs Ca

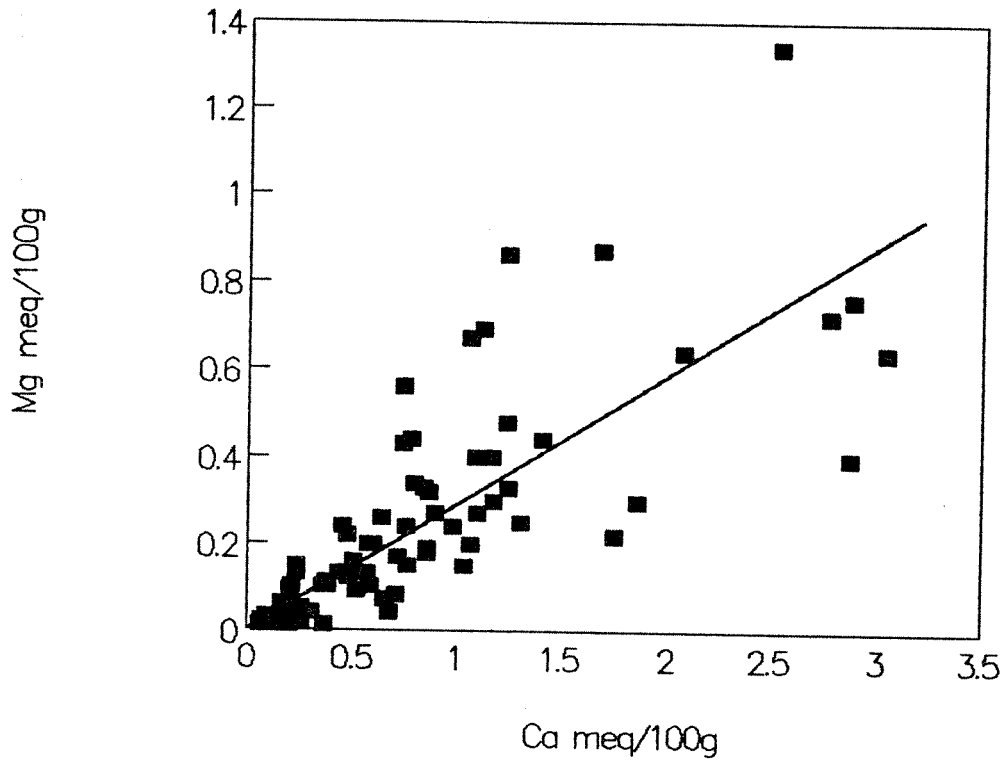


FIGURE 3.6. Sogndal. The relationship of exchangeable Mg and exchangeable Ca. The slope of the zero intercept regression equation is 0.295, with an r^2 of 0.65.

K vs CEC

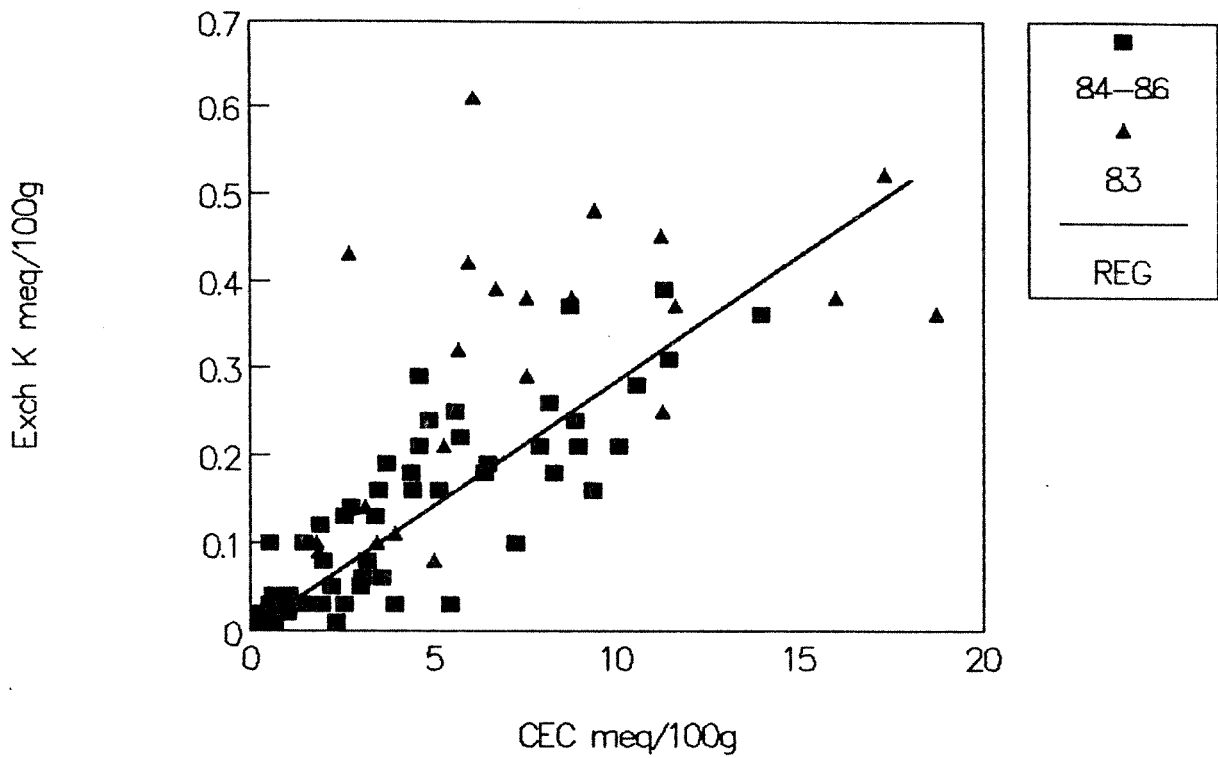


FIGURE 3.7. Sogndal. The relationship between exchangeable K and CEC, with the 1983 points differentiated. The slope of the zero intercept regression (without 1983) is 0.029 with an r^2 of 0.75.

Normal Probability Plot

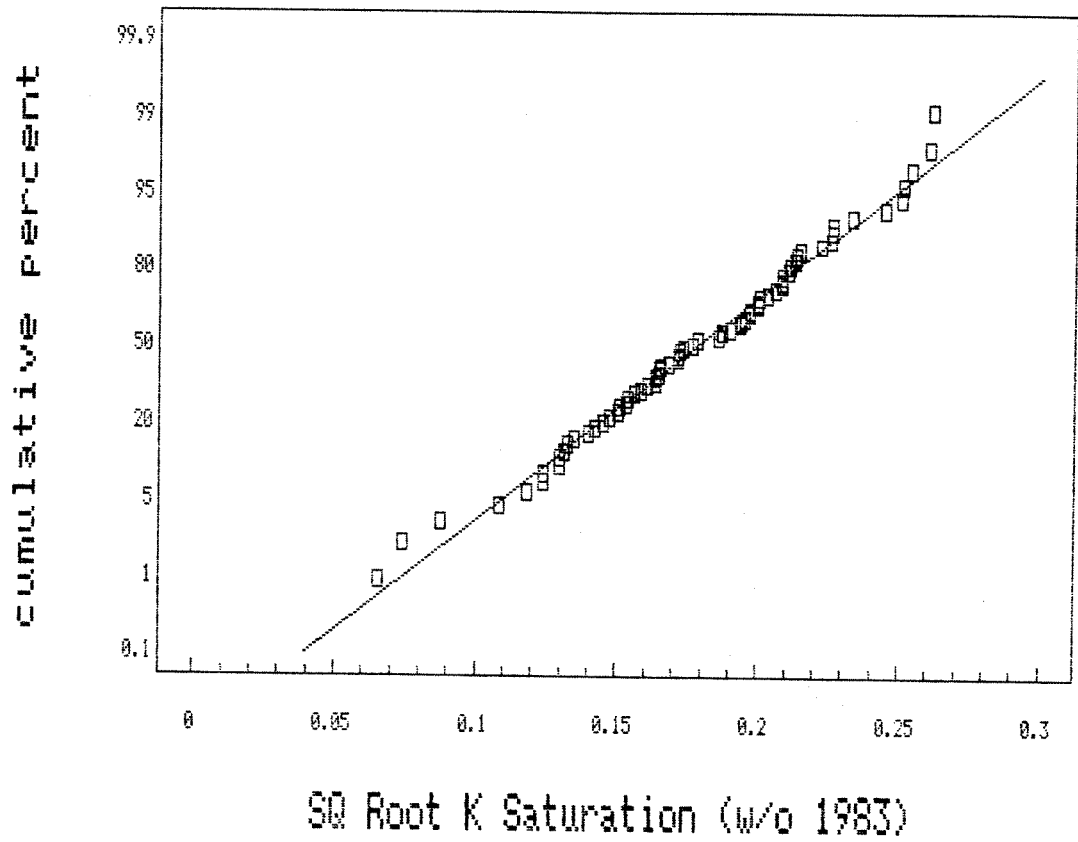


FIGURE 3.8. Sogndal. Normal probability plot for the square root transformed K saturation data.

Na vs CEC

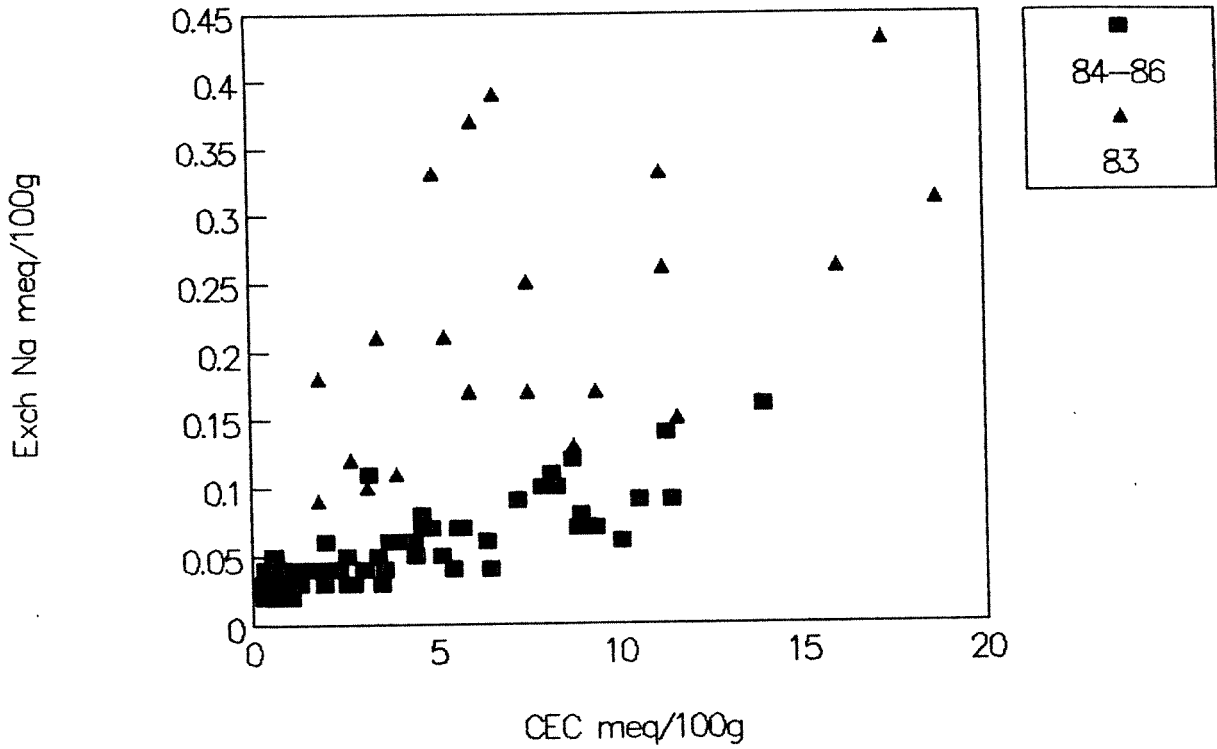


FIGURE 3.9. Sogndal. Plot of exchangeable Na as a function of CEC with the 1983 data differentiated.

$$\text{Density} = 0.810(\text{CEC}^{-.3578})$$

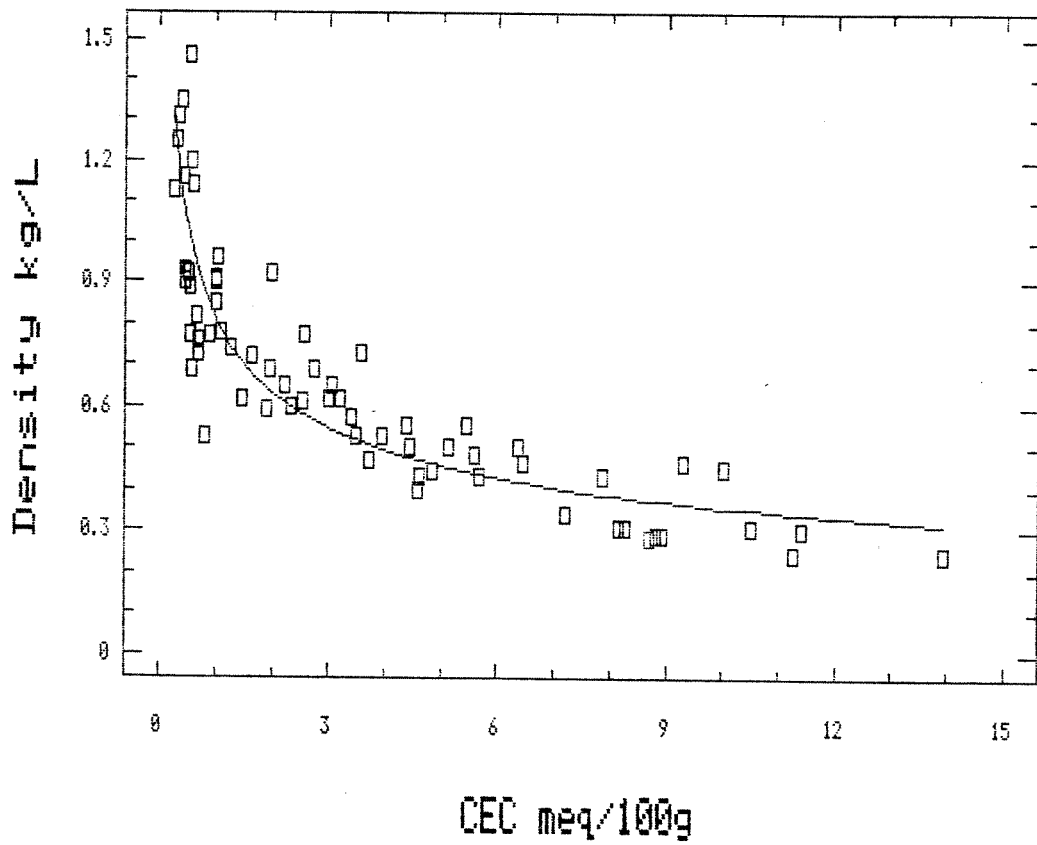


FIGURE 4.1. Sogndal. The relationship between density and cation exchange capacity.

CEC(wt) vs CEC(vol)

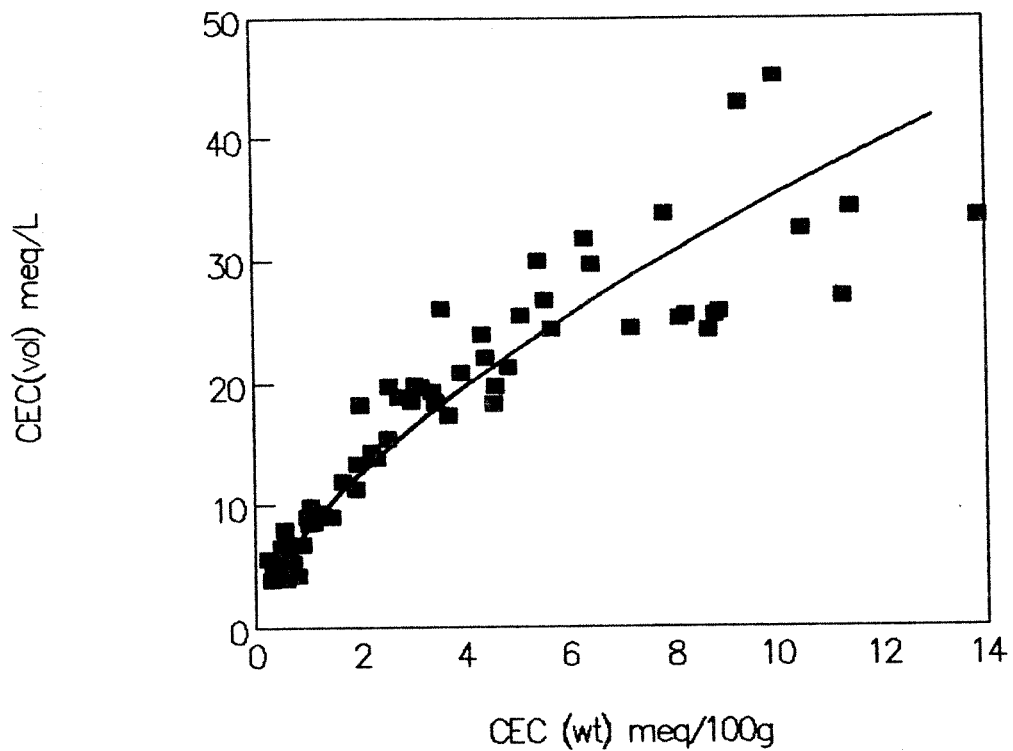


Figure 4.2. Sogndal. The relationship between CEC by weight and CEC by volume. The regression equation is $y = 8.125X^{-0.6375}$, with an r^2 of 0.935.

Part II. Risdalsheia

1. RELATIONSHIPS WITH CARBON CONTENT

1.1 Loss on Ignition versus Percent Carbon

At Risdalsheia the relationship of loss on ignition to carbon is very close as was the case at Sogndal (Figure 1.1). The regression equation in this case is:

$$y = 1.52 + 1.87x, \quad r^2 = 0.963, \quad n = 115 \quad (2.1)$$

The slope indicates that 1/1.87, or 53% of the weight loss on ignition is carbon, very near the 51% value observed for the Sogndal site.

1.2 CEC(KCl) versus Percent Carbon

Again this relationship (Figure 1.2) is very close. The regression line is:

$$y = 1.08 + 0.702x, \quad r^2 = 0.91 \quad (2.1)$$

Four points on the low side of the line are all from the 1984 sampling, and these were excluded from the regression, on the basis that the general relationship is probably better described in this manner. If these points are included the r^2 is 0.852, and the slope is 0.658.

The slope suggests a CEC at soil pH of about 0.70 meq/100g for every 1% carbon. This is above the 0.55 value found at Sogndal but still well below the 4.9 per 1% carbon suggested by Jackson (1968). Again the difference is probably largely due to use of an unbuffered method for CEC determination for these samples, as compared to methods buffered at pH 7.0 or higher which include substantial variable charge that is not available at soil pH.

2.0 CATION EXCHANGE CAPACITY (CEC)

In order to arrive at a single value for CEC, or a single value for each depth, the individual values must be aggregated. First the distributions were examined. These were highly skewed with a large influence from a few relatively high values. Normal probability plots were examined for the original data, log(base e) transformations and square root transformations. Examples for the original data and the log transforms are shown in Figure 2.1. Relevant statistics for each transform are shown in the Appendix.

While the most suitable transform varies with depth, the lognormal distribution was selected as most appropriate on an overall basis. The log transform was then used for analyses of variance and the geometric means reported. The means are summarized in Table 2.1.

With the lognormal distribution the confidence intervals are not symmetrical about the means, as these are calculated for the mean of the logs prior to taking the antilogs.

The CEC values at this site are more than double those found at Sogndal. Samples were taken from the 0-15 cm, 15-30 cm, and in some cases, from 30-45 cm. An overall estimate and a separate estimate for each depth are calculated. Because there are only about half as many samples from 30-45 cm as from the other depths, a value for > 15 cm is also given, which includes all samples from 15-30 and 30-45 cm. Depth differences are highly significant (Figure 2.2), with progressively lower values as depth increases. The largest change occurs between the 0-15 cm and 15-30 cm depths.

Again, there is a significant difference among years (Figure 2.2). The pattern differs from that observed at

Sogndal, in that while the 1983 samples tend toward high values at both sites, at Risdalsheia the 1986 values are also high, while at Sogndal the 1986 values were similar to those from 1984 and 1985. There seems to be no rational basis for rejecting any of the data sets, so values from all four years are used both for aggregation of CEC and for calculation of the base cation saturation values.

3.0 BASE SATURATION

3.1 Ca and Mg Saturation

As is evident from Figure 3.1, there is a very close relationship between exchangeable Ca and Mg at this site. The equation for the regression of Mg on Ca (zero intercept) is,

$$y = 0.53x, \quad r^2 = 0.95 \quad (4.1)$$

This close relationship was utilized in the estimation of the values for Ca and Mg saturation. First, the total saturation of (Ca+Mg) was calculated and aggregated by depth. Again, the distribution was highly skewed, and the lognormal distribution was selected. Geometric means and confidence intervals are shown in Table 3.1. The estimates of Ca and Mg saturation shown in Table 3.2 were then based on a Mg:Ca ratio of 0.53:1, as indicated by the slope of 0.53 for the regression of Mg on Ca.

The overall Ca saturation is only about one-third that observed at Sogndal, with the greatest difference at the deeper depths. Ca saturation in the 0-15 cm layer at Sogndal is estimated at 0.183, about 1.6 times the Risdalsheia value. The Sogndal value for > 15 cm is 0.168, more than five times the Risdalsheia value. Year effects are again very difficult to interpret. As noted in section 2, there was an apparent rise in CEC, with high values in 1983 and 1986. Also there were low values for exchangeable Ca+Mg in 1985 and high values in 1986 (Figure 3.2). These

effects appear to be an artifact of the measurement process, but the exact source is unknown.

3.2 K saturation

The K saturation values as estimated using the various distributions were quite similar. The log normal distribution was again selected on the basis of the linearity of the Normal Probability plots. Geometric means and confidence intervals are shown in Table 3.3.

The K saturation value for the 0-15 cm depth is 0.028, about the same as the value of 0.03 obtained for all depths at the Sogndal site. At Risdalsheia, however, the K saturation is clearly lower in the deeper layers, with a geometric mean of 0.015 for all values below 15 cm.

3.3 Na Saturation

Again a single Na saturation value is derived for all depths. This is based on the regression of exchangeable Na on CEC, as shown in Figure 3.3. In this case the intercept of the regression is not significant, and Na saturation value is taken from the slope of the regression with zero intercept. The regression equation is,

$$y = 0.0140x, \quad r^2 = 0.70 \quad (4.2)$$

The estimate of Na saturation is given by the slope, i.e. 0.014. The standard error of the slope is 0.00043, and there are 114 observations. Given the low Na saturation, no attempt was made to separate the observations by depth. However, the lack of any evidence for either an intercept or non-linearity suggests that any depth effects would be small. The points at the lower left of a graph such as Figure 3.3 tend to be from the deeper layers where cation exchange capacity is low. If the saturation decreases with depth these points tend to be below a straight line with origin at zero, resulting in a negative intercept.

Conversely an increase in saturation with depth tends to give a positive intercept.

4.0 DENSITY

There appeared to be little clear advantage for any of the distributions for estimating densities. Geometric means were selected largely on the basis of consistency with the other parameters for this site. These are shown in Table 4.1. Densities at this site are lower than at Sogndal, particularly at the 0-15 cm depth where the density at Risdalsheia is 0.221 kg/l, compared to 0.511 kg/l at Sogndal.

5.0 SUMMARY

A summary table (5.1) for the Risdalsheia site has been prepared showing the best estimates for the aggregated values of the parameters required by the MAGIC model. While these may change with further analysis of these or later data, the modifications are likely to be relatively small.

Table 2.1. Risdalsheia. Geometric means and confidence intervals for CEC (KCl).

Depth	n	Geometric mean	Confidence interval +/- 2(s.e.)
cm		----- meq/100g -----	
All	115	8.22	7.27 - 9.30
0-15	46	13.94	12.52 - 15.52
15-30	46	6.42	5.96 - 6.92
30-45	23	4.66	3.54 - 4.66
> 15	69	5.75	5.01 - 6.61

Table 3.1. Risdalsheia. Geometric means and confidence intervals for (Ca+Mg) saturation.

Depth	n	Geometric mean	Confidence interval +/- 2(s.e.)
cm		- - - - - meq/100g - - - - -	
All	115	0.082	0.069 - 0.098
0-15	46	.171	.150 - .194
15-30	46	.054	.042 - .070
30-45	23	.049	.034 - .071
> 15	69	.051	.041 - .063

Table 3.2. Risdalsheia. Estimates of Ca and Mg saturation, assuming a Mg:Ca ratio of 0.53.

Depth	Exch. Ca	Exch Mg
cm	meq/100g	meq/100g
All	0.054	.028
0-15	.112	.059
15-30	.035	.019
30-45	.032	.017
> 15	.033	.018

Table 3.3. Risdalsheia. Geometric means and confidence intervals for K saturation.

Depth	n	Geometric mean	Confidence interval +/- 2(s.e.)
cm		- - - - - meq/100g	- - - - -
All	114	0.019	0.018 - 0.021
0-15	46	.028	.026 - .030
15-30	46	.016	.014 - .018
30-45	22	.013	.013 - .016
> 15	68	.015	.013 - .016

Table 4.1. Risdalsheia. Geometric means and confidence intervals for density (kg/l).

Depth	n	Geometric mean	Confidence interval +/- 2(s.e.)
cm			kg/l
All	80	0.401	0.336 - 0.479
0-15	35	.221	.179 - .273
15-30	33	.630	.517 - .768
30-45	12	.656	.477 - .902
> 15	45	.637	.539 - .752

Table 5.1. Summary table of aggregated values of soil parameters at the Risdalsheia site for use in the MAGIC model.

Depth	Density	CEC	Base Saturation			
			Ca	Mg	K	Na
cm	kg/l	meq/100g	-----Equivalent Fraction-----			
All	0.401	8.22	0.054	0.028	0.019	0.014
0-15	.221	13.94	.112	.059	.028	.014
15-30	.630	6.42	.035	.019	.016	.014
30-45	.656	4.66	.032	.017	.013	.014
> 15	.637	5.75	.033	.018	.015	.014

Loss on Ignition vs Carbon

Risdalsheia

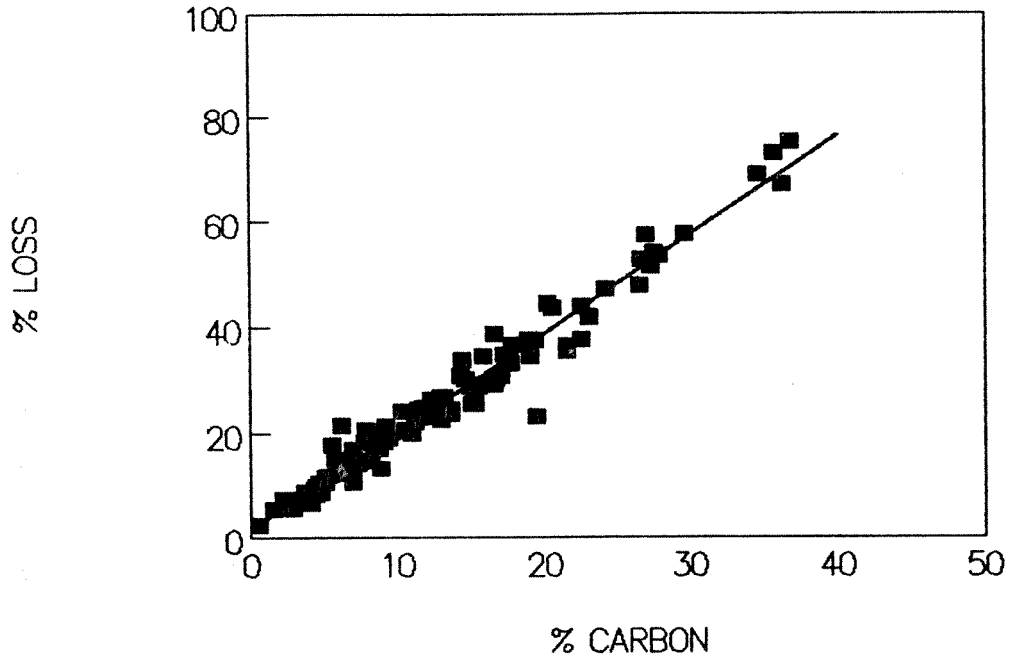


FIGURE 1.1. Risdalsheia. Loss on ignition as a function of percent carbon as determined by a chromic acid wet digestion.

CEC vs % Carbon

Risdalsheia

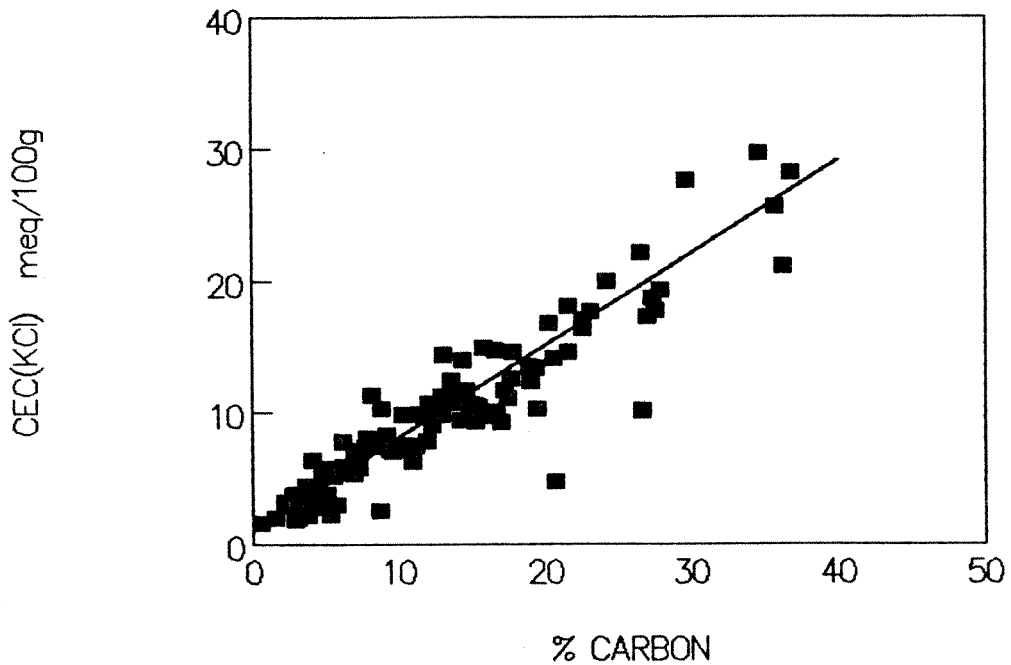
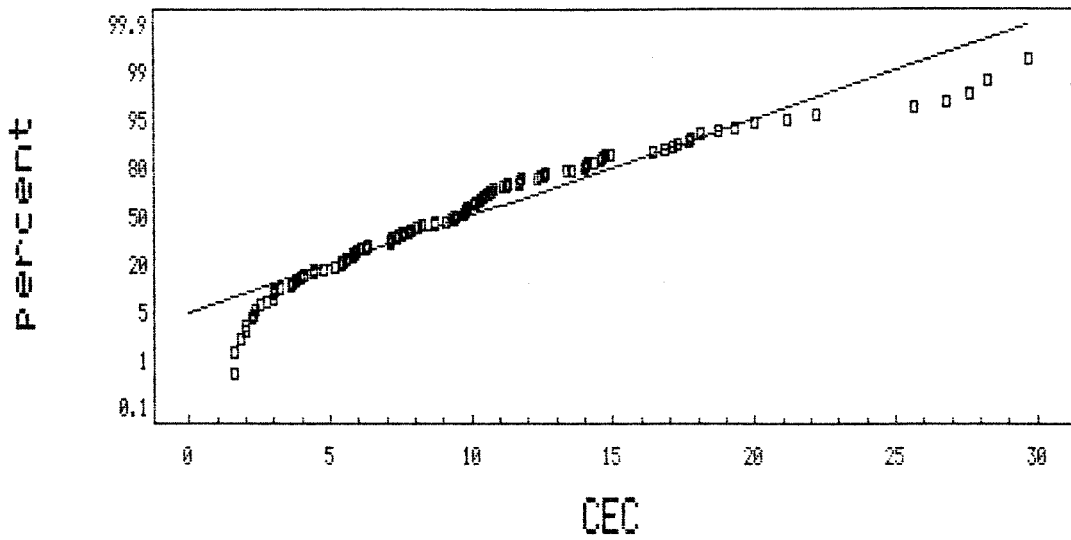


FIGURE 1.2. Risdalsheia. Cation exchange capacity (KCl) in meq/100g as function of percent carbon.

Normal Probability Plot



Normal Probability Plot

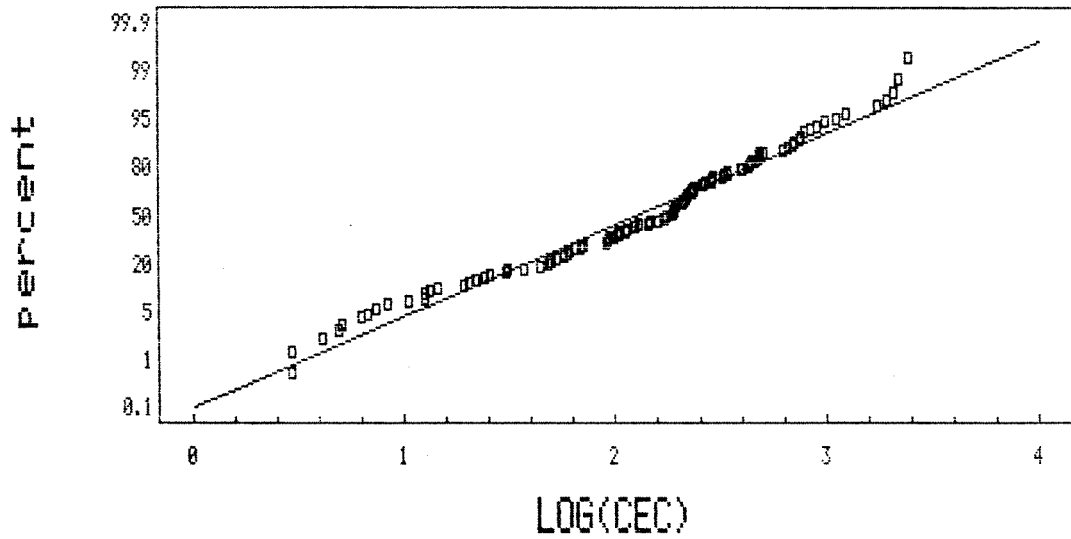
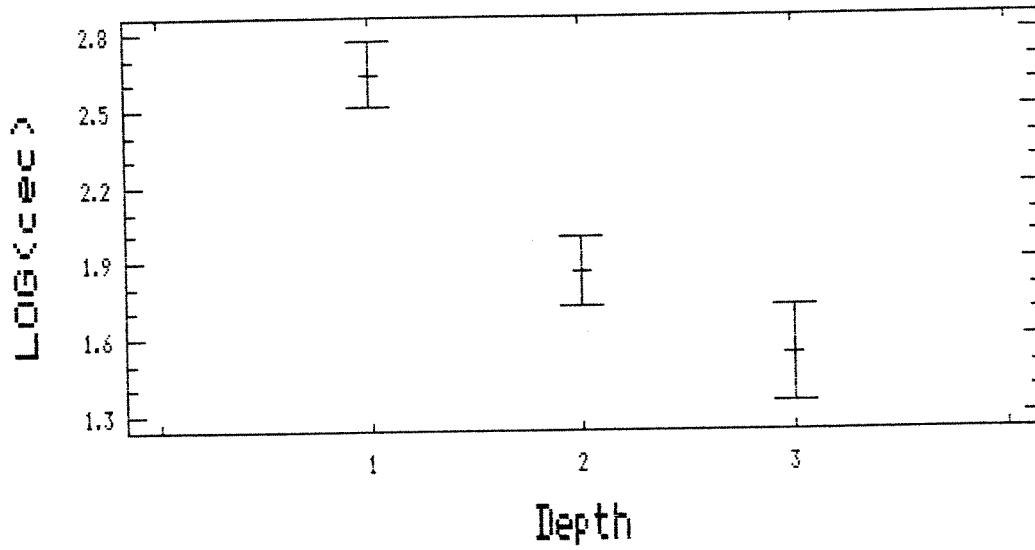


FIGURE 2.1. Risdalsheia. Normal probability plots for cation exchange capacity by weight, and for the log transform of the same data.

95 Percent Confidence Intervals



95 Percent Confidence Intervals

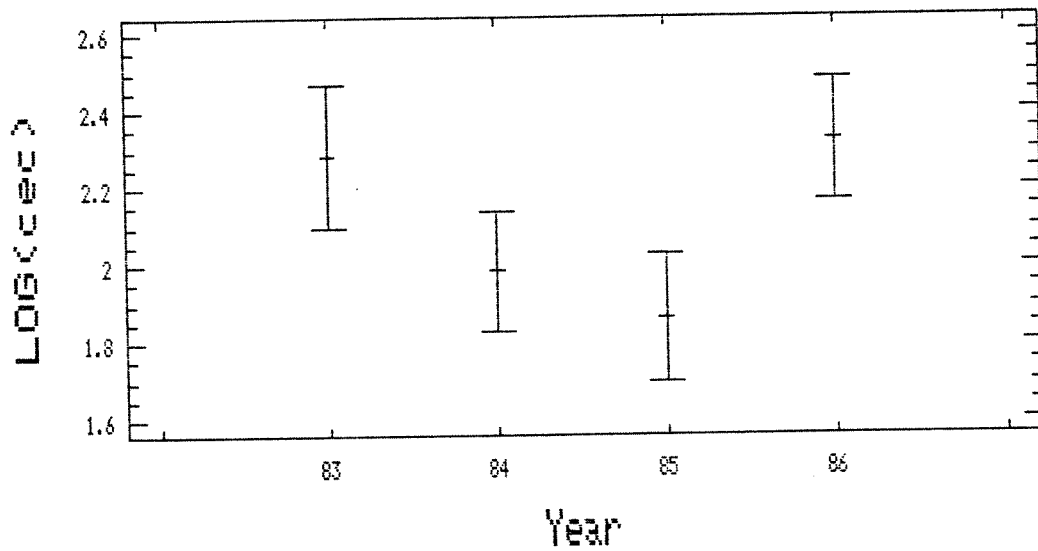


FIGURE 2.2. Risdalsheia. Means plots showing the 95% confidence interval for the logarithm (base e) of CEC by depth and year.

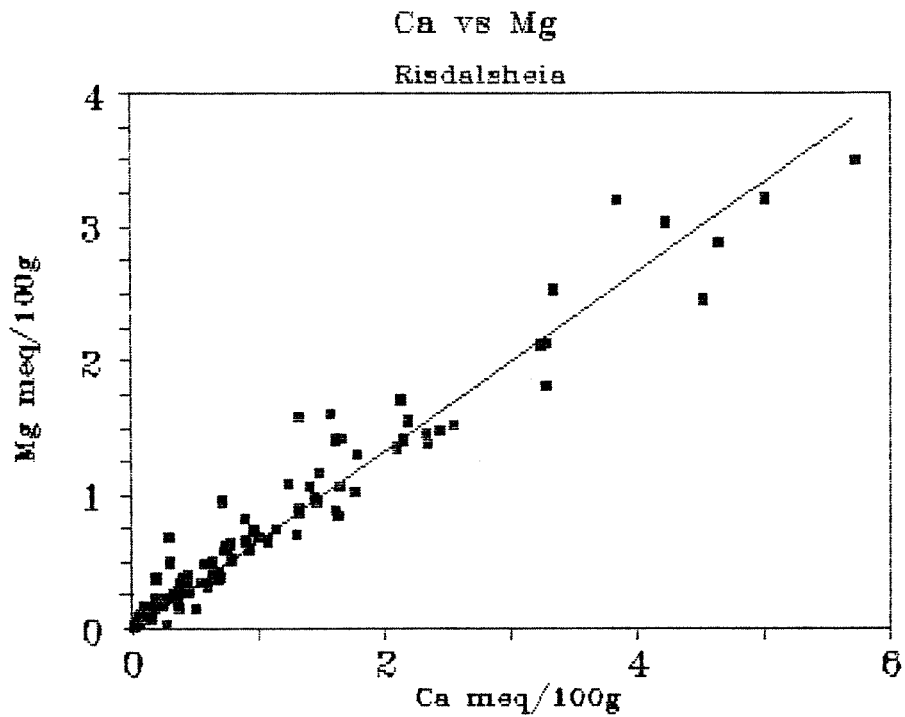


FIGURE 3.1. The relationship between exchangeable Ca and Mg at Risdalsheia. The slope (no intercept) is 0.53 and the R^2 value 0.95.

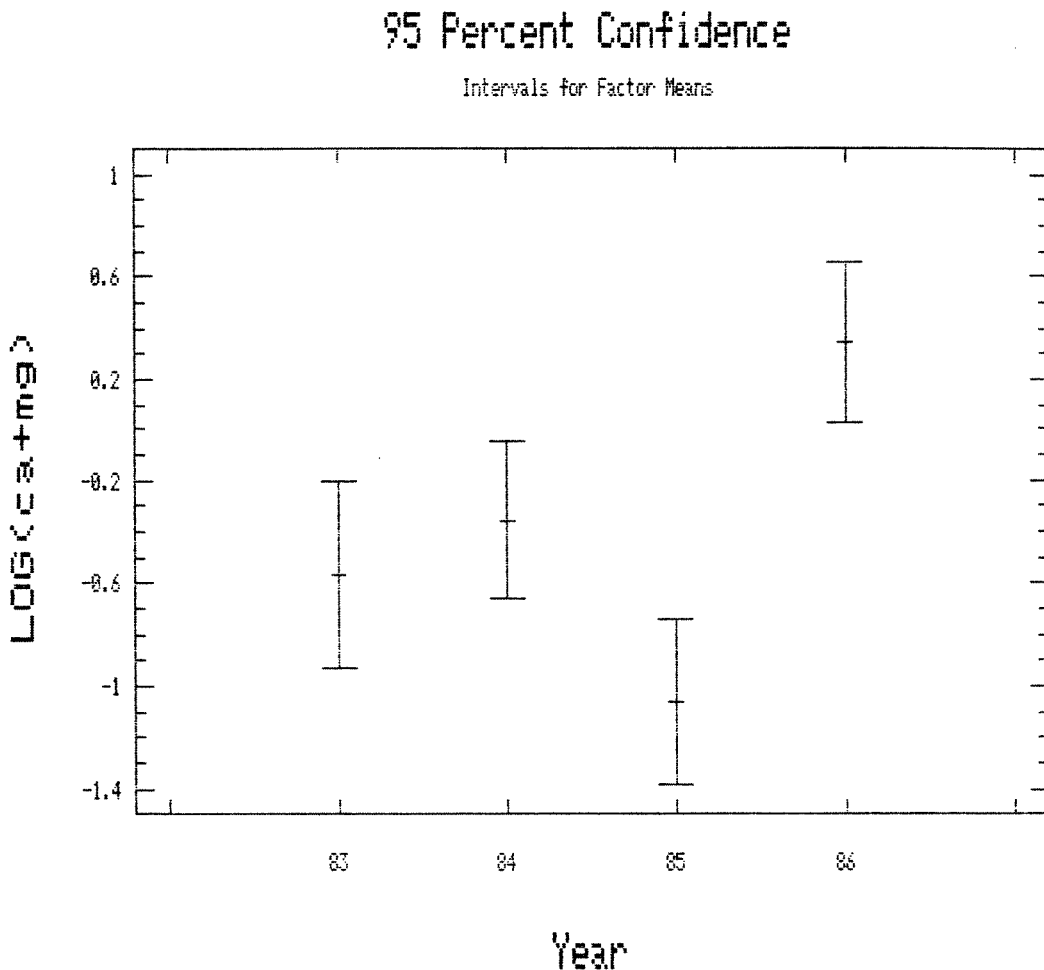


FIGURE 3.2. Risdalsheia. Means plots showing the 95% confidence interval for the logarithm (base e) of exchangeable (Ca+Mg) by years.

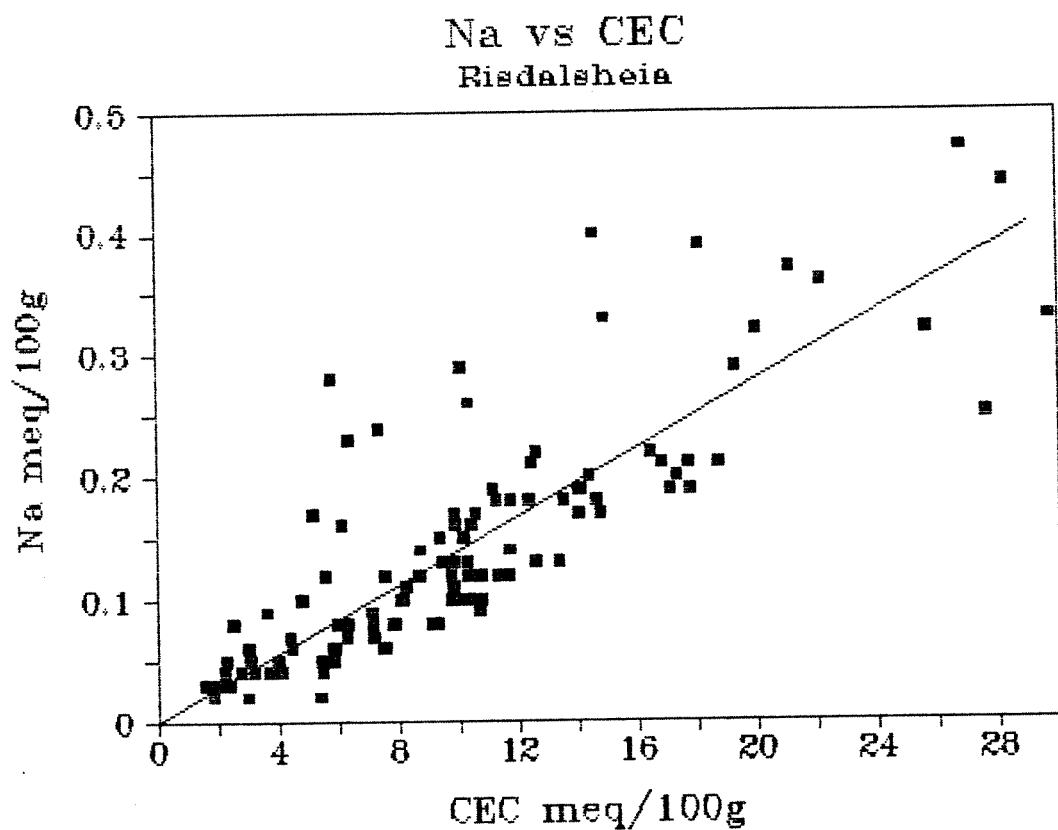


FIGURE 3.3. Risdalsheia. Regression of exchangeable Na on CEC. The slope (no intercept) of 0.014 estimates the Na saturation. R^2 is 0.70.

Part III. Anion adsorption.

1.0 INTRODUCTION

This part deals with the Cl^- and SO_4^{2-} adsorption properties of soil samples from Risdalsheia and Sogndal. These anion adsorption properties serve to buffer changes in anion concentrations of water as it passes through the soil. This buffering effect creates a lag in changes of concentration of drainage waters as compared to concentration of inputs in precipitation or dryfall. These lags, in turn, affect the response of surface waters to deposition inputs.

Anion adsorption properties are commonly described in terms of an "isotherm", i.e. a plot or mathematical function describing the amount of the adsorbed ion present as a function of the concentration of the ion in solution. While any of a number of mathematical expressions may be used to describe this relationship, the linear isotherm and the "Langmuir" isotherm will be applied here. The simplest case is the linear isotherm, for which the amount adsorbed is directly proportional to the solution concentration. For example,

$$\text{Cl}^-_{\text{ads}} = b(\text{Cl}^-), \quad (1.1)$$

where Cl^-_{ads} represents the amount adsorbed, usually expressed in mg/kg or meq/kg, and (Cl^-) represents the solution concentration, usually expressed in mg/l or meq/l. The change in the amount in the adsorbed phase per unit of concentration change is given by the slope, b .

The Langmuir isotherm is commonly used to describe SO_4^{2-} adsorption and may be described by,

$$S_{\text{ads}} = \frac{K_{\text{max}}(S)}{K_{1/2} + (S)} \quad (1.2)$$

where K_{\max} represents the maximum amount that can be adsorbed, i.e. the amount adsorbed at infinite solution concentration, while $K_{1/2}$ represents the solution concentration at which the amount adsorbed is equal to one-half K_{\max} . The use of Eq. (1.2) implies the existence of an "adsorption maximum", i.e. an adsorbed amount that will not be exceeded at any solution concentration. Eq. (1.2) is also written in terms of S rather than SO_4^{2-} , because in most cases the data used in this report are expressed in terms of mg S/kg for the adsorbed phase and mg S/l for concentration.

2.0 METHODS

The samples used to determine anion adsorption properties were collected and analyzed under the supervision of Dr. Erik Lotse at the Swedish Agricultural University in Uppsala (Lotse and Ottabong 1985, Lotse 1989). In general, the Cl^- adsorption data were obtained by equilibrating wet soil samples (approximately 2.5 gm dry soil equivalent) with 25 ml of solution with initial concentrations of 0, 8, 16, and 32 mg Cl^- per liter, and determining the Cl^- concentration after equilibration. The amount adsorbed or desorbed is then calculated from the change in concentration. For the sulfate adsorption isotherms equilibration solutions containing 0, 1, 2, 4, 8, 16, and 32 mg S/l were used.

The observed Cl^- isotherms were very nearly linear, and no attempt was made to describe them in terms of the Langmuir-type relationship. Similarly, because most of the plots for the SO_4^{2-} isotherms at the Risdalsheia site appeared linear, a linear isotherm was used and no attempt was made to describe these using the Langmuir relationship. Some curvilinearity was observed for several of the SO_4^{2-} isotherms for the Sogndal site, and in these cases the constants for the Langmuir relationship were estimated.

Concentrations determined after equilibration differ from those determined before equilibration by the amount of the ion that has been desorbed or adsorbed during the equilibration process. At low initial concentration the solution becomes more concentrated due to desorption, while at higher concentrations, adsorption occurs and the solution becomes less concentrated. The procedure used here was to convert this change in concentration (- for desorbed and + for adsorbed) to units of mg/kg and plot this change against final concentration. An example using the Cl^- data from the 0-15 cm depth at sampling site #2 at Sogndal is shown in the upper graph in Figure 2.1. The linear regression relationship for this plot is,

$$Y = -6.43 + 1.223X, r^2 = 0.988 \quad (2.1)$$

Not all the relationships are this close, and in a few cases obvious outliers were deleted before calculating the regressions. Since the relationship is linear, the intercept of -6.43 mg/kg represents the amount that would have to be desorbed in order to achieve a zero final concentration, i.e. the total amount adsorbed assuming full reversibility of adsorption. Therefore, in order to construct an isotherm showing the amount adsorbed as a function of final solution concentration, the curve must be displaced upward by 6.43 mg/kg, i.e., 6.43 mg/kg must be added to each observed value. When this is done the isotherm passes through 0,0 as required by Eq. (1.1). and shown in the lower graph in Figure 2.1.

The equilibrium concentration is given by the concentration at which no Cl^- is adsorbed or desorbed, i.e. the concentration at which the curve for the lower graph in Figure 2.1 intercepts the Y axis. From Eq. (2.1) we may calculate this equilibrium concentration for our example to be 5.26 mg/kg.

The slope is the key parameter for a linear isotherm. It represents the change in adsorbed phase (mg/kg in this case) per unit of concentration change. A steep slope means that there will be a large change in adsorbed phase for a small change in solution concentration, i.e. the system will be well buffered against changes in solution concentration. Conversely, a flatter slope is associated with less buffering due to the adsorption process.

The procedure used here is slightly different from that which was previously used to construct isotherms from these data. In the previous scheme all final concentrations were adjusted by subtracting the concentration observed after equilibration with solutions of zero initial concentration. This is equivalent to adjusting by the amount desorbed when equilibrated with the solution of initial concentration of zero rather than extrapolating to the amount that must be desorbed to equilibrate with a final concentration of zero. The differences are usually rather small, in our example the adjustment by the previous procedure would have been about 4.52 mg/kg rather than the 6.43 mg/kg arrived at by extrapolation. However, in some cases the differences may be significant and the extrapolation procedure used here appears to be more appropriate.

Two methods have been tried for fitting the Langmuir isotherm in the form shown in Eq. (1.2). It is apparent from Eq. (1.2) that at very low concentrations the function approaches linearity, as when (S) is small in relation to $K_{1/2}$, the (S) term in the denominator becomes insignificant and the function approaches,

$$S_{\text{ads}} = \frac{K_{\text{max}}(S)}{K_{1/2}} \quad (2.2)$$

Thus, if only points from the low end of the solution concentration are used, a linear extrapolation to zero concentration should be appropriate. This was done by simply visually inspecting the original plot of

adsorbed/desorbed and selecting a cutoff point below which the curve was apparently linear. An example of such a plot for the SO_4^{2-} data from sampling site #9 at Sogndal is shown in the upper graph in Figure 2.2. Linear regression was then used to extrapolate these points to zero final concentration. In the case of the example the cutoff was set at an initial concentration of 8 mg S/kg for both depths. However, in several instances the cutoff was set at an initial concentration of 4 mg S/kg to avoid the non-linear portions of the curve. For the example shown in Figure 2.2 the extrapolations yielded initial values of -20.17 and -10.64 mg S/kg for the 0-15 and 15-30 cm depths, respectively, and these values were used to adjust the adsorbed values before proceeding to the next step in the fitting procedure.

By inverting Eq. (1.2) and rearranging we obtain,

$$\frac{1}{S_{\text{ads}}} = \frac{K_{1/2}}{K_{\text{max}} (S)} + \frac{1}{K_{\text{max}}} \quad (2.3)$$

From Eq. (2.3) it is apparent that if the relationship in Eq. (1.2) applies, a plot of $1/S_{\text{ads}}$ as a function of $1/(S)$ should be linear, and the intercept is the reciprocal of K_{max} . Once K_{max} is known $K_{1/2}$ may be calculated from the slope. A reciprocal plot for the Sogndal site #9 example is shown in the lower graph in Figure 2.2. The curvilinearity of the adsorption curve is determined largely by the points at high concentrations, which are the points at the lower left of the reciprocal curve. The low concentration points, particularly those with an initial concentration of zero were sometimes inconsistent on the reciprocal plot. This inconsistency may arise simply due to measurement error, as at very low concentrations small differences will have a large effect on the reciprocal. Similarly, small differences due to errors in the extrapolation used to calculate the correction for the

native adsorbed phase would cause large differences in the reciprocals. Therefore, the fitting of Eq. (2.3) was confined to points with initial concentration (prior to equilibration) of 2.0 mg S/l or greater. The slopes and intercepts of the reciprocal curve were then determined by linear regression, and K_{max} and $K_{1/2}$ were calculated. Results for the example are shown in Table 2.1.

The adsorption plot for the 0-15 cm depth is nearly linear, and this is reflected in the relatively large values for K_{max} and $K_{1/2}$.

Again, the equilibrium concentration is the concentration at which no change in adsorption takes place, i.e. the concentration at which the uncorrected adsorption curve has a value of zero. Defining our correction values obtained from the intercepts (in our example 20.17 and 10.64 mg S/kg for the 0-15 and 15-30 cm depths respectively) as C, Eq. (1.2) for the uncorrected adsorption values may be written as,

$$S_{ads} \text{ (uncorrected)} = \frac{K_{max}(S)}{K_{1/2} + (S)} - C \quad (2.4)$$

Setting S_{ads} (uncorrected) equal to zero we may solve for the equilibrium concentration, $(S)_{eq1}$,

$$(S)_{eq1} = \frac{K_{1/2}C}{K_{max} - C} \quad (2.5)$$

The second procedure used for fitting Eq. (2.2) involves the use of a non-linear regression program for curve fitting. Equation (2.2) must go through the origin. However, if a constant C is added Eq.(2.2) takes the form,

$$S_{ads} = C + \frac{a(S)}{b + (S)} \quad (2.6)$$

The constant C represents the correction for the adsorbed phase that must be applied to bring the curve to a zero intercept. When a non-linear regression program is used to fit this curve the intercept is -C, which represents the extrapolation of the Langmuir relationship to zero concentration. When this correction is applied and the curve is refit, C is very nearly zero while a and b are direct estimates of K_{max} and $K_{1/2}$, respectively. The equilibrium concentration may then be calculated using Eq. (2.5).

An example of the fitted isotherm for the 15-30 cm depth at sampling site #9 at Sogndal is shown in Figure 2.3. In this case the correction as calculated by the non-linear regression was 11.94 mg/kg as compared to 10.64 mg/kg as calculated above by linear extrapolation of the low concentration points. In principle this procedure should estimate this correction more accurately than the linear extrapolation, particularly if the relationship is markedly curvilinear. In this case the estimated values of K_{max} and $K_{1/2}$ are 250 mg S/kg and 50.2 mg/l, respectively. These compare to respective values of 244 mgS/kg and 51.5 mg/l as calculated using the reciprocal procedure above (Table 2.1).

3.0 CHLORIDE ADSORPTION

3.1 Risdalsheia

An example of a Cl^- isotherm from Risdalsheia is shown in Figure 3.1. The "Initial" isotherm has not been corrected for the adsorbed phase, and the intercept is used to estimate the amount in the adsorbed phase in the original sample. In the "Final" isotherm this amount has been added to the observed amount in each sample, so that the isotherm starts at zero concentration and zero adsorbed Cl^- .

The isotherm characteristics for the two depths are shown in Table 3.1. The constant (intercept), slope and r^2 were

determined using linear regression. The slope represents the amount (mg/kg) that must be adsorbed/desorbed for the concentration to change by one mg/l. While all calculations were made using these units, the slope would be the same for units of meq/kg and meq/l. The amount of adsorbed Cl^- present (negative of the constant) is about three times as high in the 0-15 cm layer as in the 15-30 cm layer. The slope is also greater at 0-15 cm, indicating that on a weight basis, more Cl^- must be adsorbed or desorbed for a given change in solution concentration. The density is lower in the 0-15 cm depth, so on a volume basis these differences would largely disappear. The equilibrium concentrations are also higher at 0-15 cm than at 15-30 cm. It is not known whether this difference might be permanent, or temporary due to high sea salt input prior to sampling. However, depth effects were much less apparent at the Sogndal site.

The mean equilibrium concentrations are 17.6 mg/kg and 9.12 mg/kg for the 0-15 cm and 15-30 cm depths, respectively. If these isotherms were to accurately reflect the field system, the drainage water Cl^- concentration would be approximately the same as the equilibrium concentration. However, volume weighted mean runoff concentrations at Risdalsheia average about 105 ueq/l (Wright et al. 1988), or about 3.7 mg/l. This is less than one-half the mean equilibrium concentration as calculated from the isotherm data for the 15-30 cm depth and less than one-fourth that for the 0-15 cm depth.

3.2 Sogndal

Sample isotherms for the Sogndal site are shown in Figure 2.1. Isotherm characteristics are shown in Table 3.2. The slopes are quite similar for the two sites, but the initial amounts adsorbed are lower at Sogndal than at Risdalsheia. Also there is little or no indication of any difference due to depth at Sogndal. Mean equilibrium concentrations are 4.58 mg/l at the 0-15 cm depth and 3.70 at 15-30 cm. These

are only about half the concentration at the 15-30 cm depth at Risdalsheia. Again, this is well above the volume weighted mean of about 1.46 mg/l for runoff at Sogndal during the 1983-87 period (Wright et al. 1988). However, mean Cl^- concentrations in runoff vary by 2-3 times from year-to-year at Sogndal, and the mean for the four catchments was 2.52 mg/l during the 1984 season (Wright et al. 1988).

4.0 SULFATE ADSORPTION

4.1 Risdalsheia

The SO_4^{2-} isotherms at Risdalsheia were generally very nearly linear. There did seem to be some tendency toward non-linearity at the 0-15 cm depth, but this was not sufficient to allow fitting of the Langmuir relationship. However, because of the geometric increase in concentrations of the equilibrating solutions, in a linear regression these high points exert a substantial influence on the slope and intercept. As the accurate extrapolation of the intercept to zero final concentration is critical in establishing the correction for native adsorbed SO_4^{2-} , the points from the initial concentration of 32 mgS/l were not used for the regressions at the 0-15 cm depth. At site #12 the 8 mg/l solutions were also excluded.

Example isotherms from sampling site #10 (KIM) catchment are shown in Figure 4.1, and the isotherm characteristics are summarized in Table 4.1. As was the case with Cl^- , on a weight basis there is substantially more adsorbed SO_4^{2-} at the 0-15 cm depth than at 15-30 cm. The means are 24.02 mg S/kg at 0-15 cm and 9.43 mg S/kg at 15-30 cm. Also, the estimated equilibrium concentrations are substantially higher than those found in drainage water at the site. For example, the mean equilibrium concentration for the 15-30

cm depth is 7.06 mg S/kg or about 440 ueq/l. This exceeds weighted annual mean sulfate concentrations in runoff at EGIL catchment (roofed, acid rain) by about a factor of four (Wright et al. 1988).

It is also of interest to compare the amounts of adsorbed SO_4^{2-} , as estimated from extrapolating the isotherms, with independent measurements of "soluble" sulfate as measured by a water extraction and "adsorbed" sulfate as measured by a phosphate extraction following water extraction, as shown in Table 4.2. It is apparent that the estimates from the isotherms are similar to those from the water extraction. This is not surprising as the procedures are quite similar. However, it also appears that the additional adsorbed SO_4^{2-} that is removed by the phosphate extraction does not participate in the adsorption/desorption process, at least as measured by the isotherm.

4.2 Sogndal

The SO_4^{2-} isotherms for the Sogndal site were fit to the Langmuir isotherm using both a reciprocal plot procedure and non-linear regression. Isotherm characteristics as determined by the two methods are shown in Tables 4.3 and 4.4, respectively. In a few cases the Langmuir relationship could not be applied as the isotherm was too nearly linear. As the curvilinearity was modest in all cases, these fitting procedures were used largely to demonstrate the methods. However they do give appropriate values for the native adsorbed sulfur (i.e. the constant) and for the equilibrium concentration. No judgement can be made as to the relative merits of the methods, as the data are too nearly linear. In most cases results are very similar. However, in one case the $K_{1/2}$ and K_{\max} values as determined by the non-linear regression appear to be very large. Within the range of the data points, however, there is very little difference in the actual isotherms.

At the moment no comparison can be made between the water extractions and the phosphate extractions, as these have not been summarized. Equilibrium concentrations are generally of the order of 3.0 mg/l (48 ueq/l). This is somewhat higher than the observed values of 20-30 ueq/l in runoff from the untreated reference catchments at Sogndal (Wright et al. 1988).

REFERENCES

- Cosby, B.J., G.M. Hornberger, J.N. Galloway, and R.F. Wright. 1985. Modelling the effects of acid deposition: assessment of a lumped-parameter model of soil water and streamwater chemistry. Water Resour. Res. 21: 51-63.
- Jackson, M.L. 1958. Soil Chemical Analysis. Prentice Hall Inc. Englewood Cliffs NJ. 206p.
- Lotse, E. G. 1989. Soil chemistry 1983-86 at the RAIN project catchments. Acid Rain Research Report 18/89. Norwegian Institute for Water Research. Oslo. 24 p.
- Lotse, E., and E. Ottabong. 1985. Physiochemical properties of Soils at Risdalsheia: RAIN Project. Acid Rain Research Report 8/1985. Norwegian Institute for Water Research. Oslo. 48p.
- Wright, R.F., E. Lotse, and A. Semb. 1988. Reversibility of acidification shown by whole-catchment experiments. Nature 334: 670-675.

ACKNOWLEDGEMENTS

This work was conducted on contract to NIVA as part of the RAIN project. The RAIN project receives financial support from the Norwegian Ministry of Environment, The Royal Norwegian Council for Scientific and Industrial Research, the Ontario Ministry of the Environment, Environment Canada, the Swedish Environmental Protection Board, the Central Electricity Generating Board (UK), and from internal research funds from the participating institutes.

Table 2.1. Regression of $1/S_{ads}$ versus $1/(S)$ and calculated values of K_{max} and $K_{1/2}$ for sampling site 9 at Sogndal.

Depth cm	Intercept	Slope	r^2	K_{max} mg S/kg	$K_{1/2}$ mg S/l	Equil. mg S/l
0-15	0.0015	0.175	0.993	668	116.7	3.63
15-30	0.0041	0.211	0.954	244	51.5	2.29

Table 3.1. Characteristics of the Cl⁻ isotherms at Risdalsheia. The site numbers refer to sampling transects. Data are analyzed for KIM and EGIL catchments.

Site/ Catchment	n	Constant mg/kg	r ₂	Slope	Equilibrium mg/L
<u>0-15 cm Depth</u>					
10/KIM	8	-51.63	.950	2.74	18.8
12/KIM	8	-22.08	.927	1.76	12.5
17-19/EGIL	8	-20.05	.975	1.38	14.5
23-25/EGIL	8	-29.85	.964	1.21	24.7
mean		-30.91		1.77	17.6
s.d.		14.44		.68	5.4
<u>15-30 cm Depth</u>					
10/KIM	8	-12.46	.878	1.69	7.4
12/KIM	6	-5.49	.944	.76	7.3
17-19/EGIL	8	-12.13	.971	1.05	11.5
23-25/EGIL	7	-7.90	.998	.76	10.3
mean		-9.50		1.07	9.1
s.d.		3.38		.44	2.1

Table 3.2. Characteristics of the Cl⁻ isotherms at Sogndal.

Site	n	Constant mg/kg	r ₂	Slope	Equilibrium mg/l
<u>0-15 cm Depth</u>					
2	8	-6.44	.976	1.22	5.28
7	7	-4.00	.979	1.39	2.88
8	8	-7.04	.973	1.41	4.99
9	7	-8.83	.997	1.71	5.16
		mean	-6.57	1.44	4.58
		s.d.	2.00	.23	1.13
<u>15-30 cm Depth</u>					
2	8	-5.66	.949	1.57	3.60
7	8	-2.23	.976	1.29	1.73
8	8	-4.68	.973	.93	5.01
9	8	-7.46	.946	1.66	4.49
		mean	-5.00	1.36	3.70
		s.d.	2.18	.44	1.44

Table 4.1. Characteristics of the SO_4^{2-} isotherms at Risdalsheia.

Site/ Catchment	n	Constant mg S/kg	r_2	Slope	Equilibrium mg S/l
<u>0-15 cm Depth</u>					
10/KIM	12	-17.10	.930	2.67	6.37
12/KIM	10	-24.67	.891	1.79	13.78
17-19/EGIL	12	-30.27	.68	1.82	16.63
23-25/EGIL	12	-18.12	.953	2.24	8.08
	mean	-24.02		2.13	11.47
	s.d.	6.60		.41	4.60
<u>15-30 cm Depth</u>					
10/KIM	14	-9.02	.966	1.39	6.49
12/KIM	14	-7.18	.960	1.20	5.98
17-19/EGIL	14	-12.80	.953	1.33	9.62
23-25/EGIL	13	-8.73	.968	1.42	6.15
	mean	-9.43		1.34	7.06
	s.d.	2.39		.10	1.72

Table 4.2. Comparison of sulfate as estimated from extrapolation of the isotherms with that measured by water extraction followed by phosphate extraction.

	0 - 15 cm mg S/kg	> 15 cm mg S/kg
Water	22.4	12.62
Phosphate	9.78	8.61
Isotherm	24.02	9.43

Table 4.3. Characteristics of the SO_4^{2-} isotherm at Sogndal as fitted to the Langmuir isotherm using a reciprocal plot.

Site/Field	Constant mg S/kg	K_{max} mg S/kg	$K_{1/2}$ mg S/l	Equilibrium mg S/l
<u>0-15 cm</u>				
2/(f2)	-29.46	*	*	10.73*
7/(f2)	-14.61	140	24.5	2.85
8/(f4)	-13.73	169	51.4	4.55
9/(f1)	-20.16	668	116.7	3.63
<u>15-30 cm</u>				
2/(f2)	-26.84	318	98.6	9.09
7/(f2)	-14.20	142	25.0	2.78
8/(f4)	-11.29	814	280	3.94
9/(f1)	-10.64	244	51.5	2.35

* Parameters could not be calculated because isotherm essentially linear. Linear isotherm with slope of 2.74 used to calculate equilibrium concentration.

Table 4.4. Characteristics of the SO_4^{2-} isotherm at Sogndal as fitted to the Langmuir isotherm using non-linear regression.

Site/Field	Constant mg S/kg	K_{max} mg S/kg	$K_{1/2}$ mg S/l	Equilibrium mg S/l
<u>0-15 cm</u>				
2/(f2)	*	*	*	*
7/(f2)	-15.82	210	24.5	2.85
8/(f4)	-13.44	270	58.1	3.04
9/(f1)	*	*	*	*
<u>15-30 cm</u>				
2/(f2)	-26.55	11552	3928	9.05
7/(f2)	-14.57	316	92.0	4.45
8/(f4)	-12.27	225	66.0	3.81
9/(f1)	-11.97	254	50.0	2.47

* Parameters could not be calculated because isotherm essentially linear.

Sogndal - Cl
Site 2, 0-15 cm

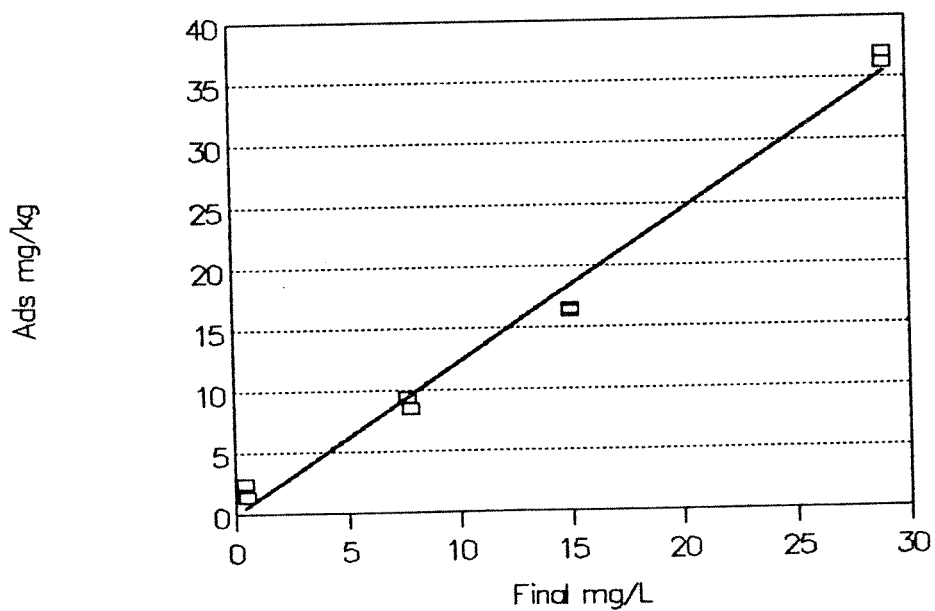
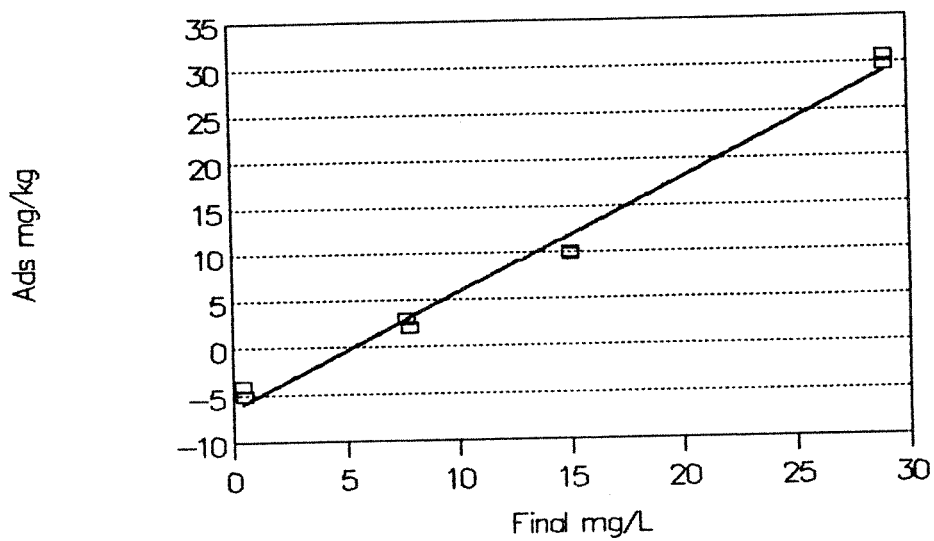
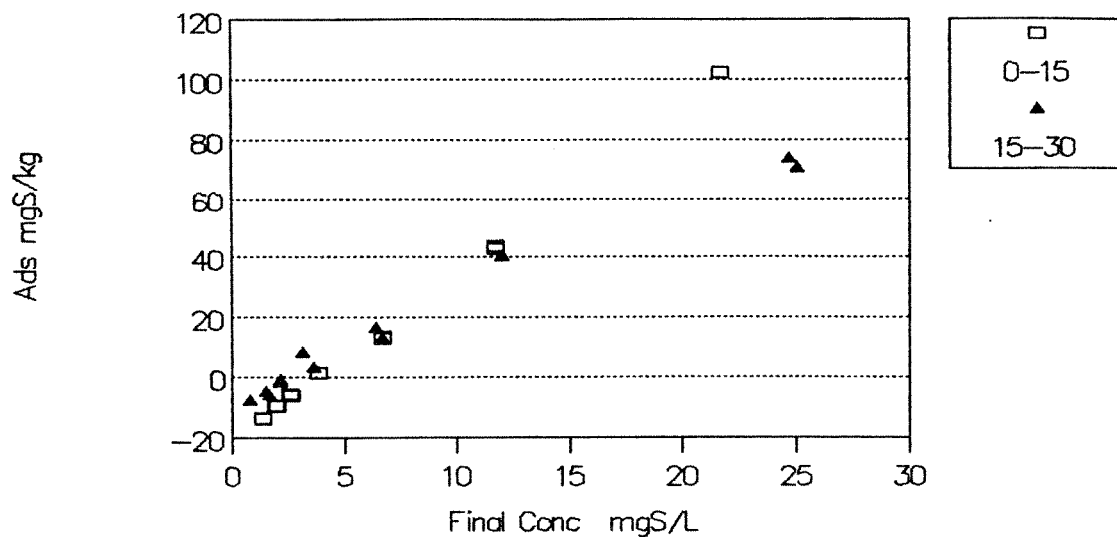


Figure 2.1. Initial uncorrected Cl⁻ isotherm (above) and final corrected isotherm (below) for site 2 at Sogndal.

Sogndal - SO4

Site 9



Site 9 - Reciprocal

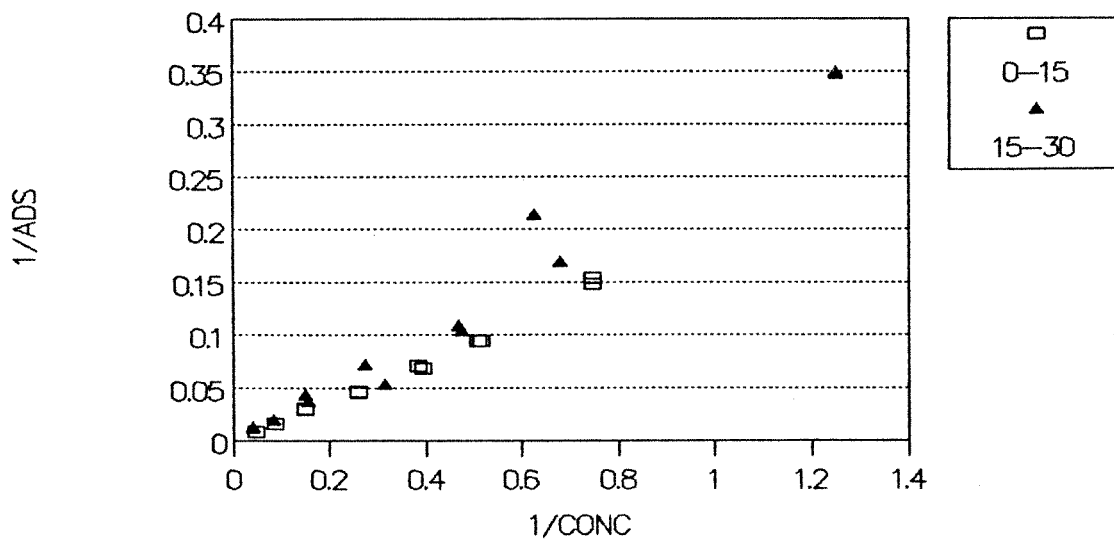


Figure 2.2. The initial uncorrected plot of SO_4^{2-} adsorbed/desorbed (above) and reciprocal plot of corrected data used in fitting the Langmuir isotherm (below) for the 15-30 cm depth of sampling site 9 at Sogndal.

Fitted Model
Site 9 - Depth 2

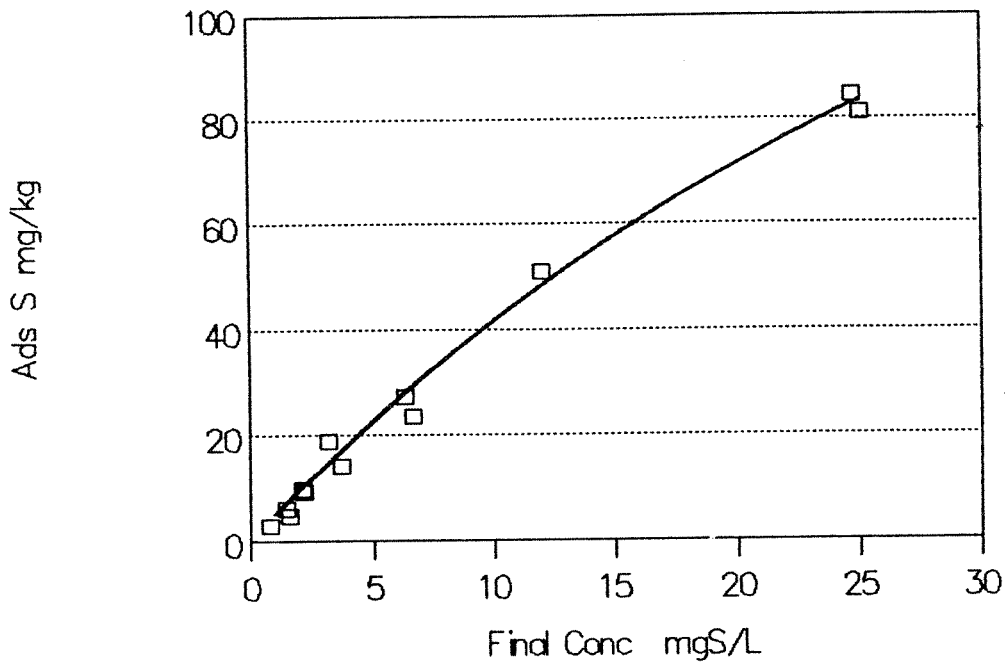
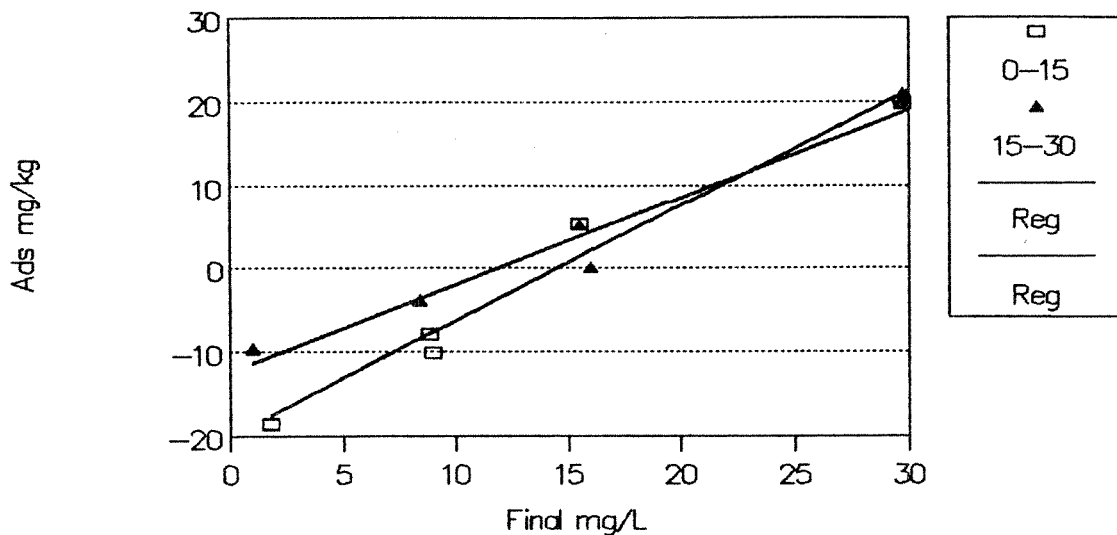


Figure 2.3. Langmuir isotherm for SO_4^{2-} adsorption for the 15-30 cm depth at sampling site 9 at Sogndal as fit using the non-linear regression procedure.

Risdalsheia - Cl Site (17-19)

Initial Plot



Final Plot

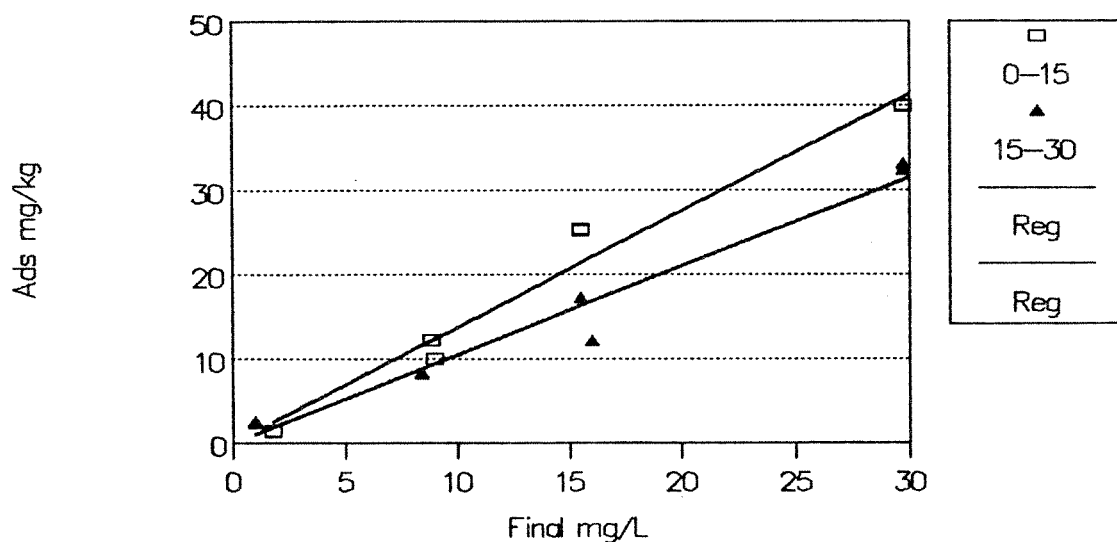


Figure 3.1. Initial uncorrected plot and final corrected plot for Cl⁻ adsorption/desorption for sampling site (17-19) at Risdalsheia.

Risdalsheia – SO₄

Site 10

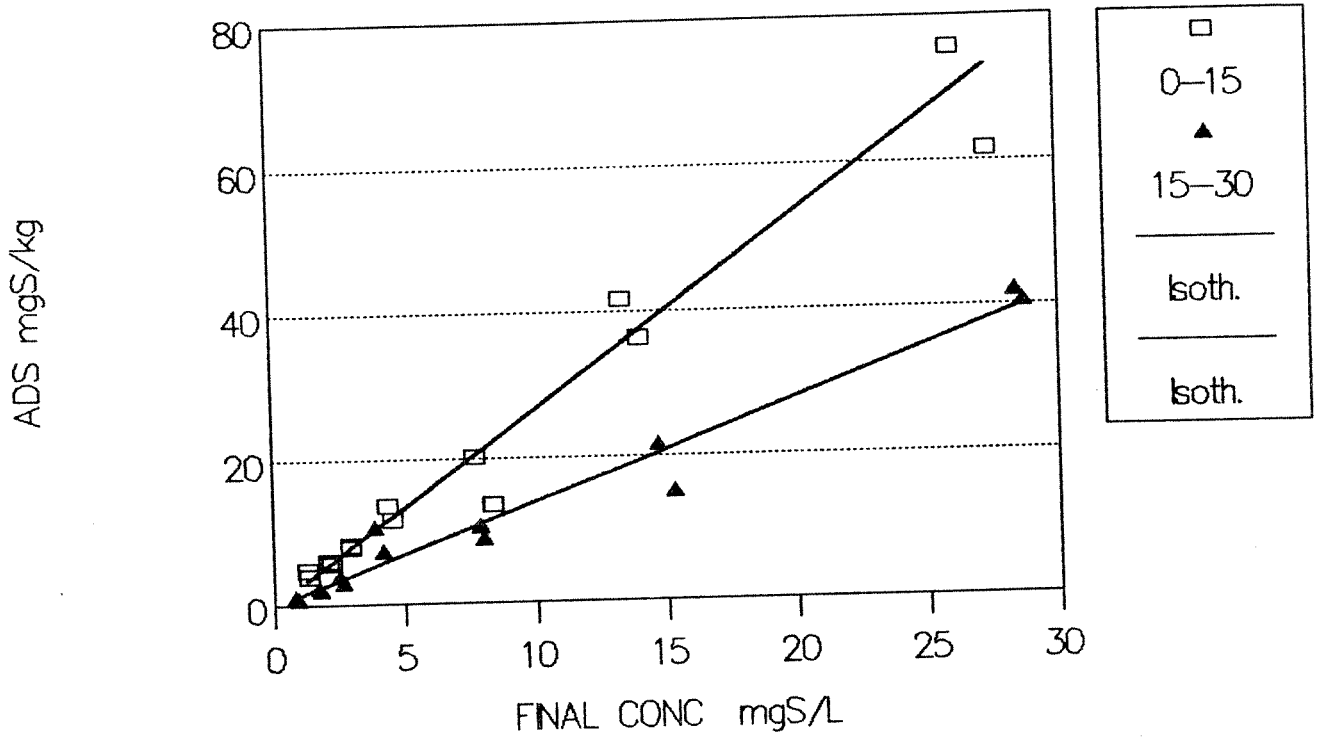


Figure 4.1. Final corrected SO₄²⁻ adsorption for sampling site 10 at Risdalsheia. Only the linear isotherms were fitted in this case.

APPENDICES

Table A 1. Risdalsheia, distribution parameters for cation exchange capacity (KCL) in meq/100 g.

Depth	n	mean	Median	s.d.	Central Tendency ^a	NP Index ^b
<u>Raw data</u>						
All	115	9.97	9.43	6.04	9.97	P
0-15	46	14.94	13.74	5.82	14.94	P
15-30	46	7.17	7.11	3.18	7.17	G
30-45	23	5.64	5.39	3.37	5.64	P
> 15	69	6.66	6.26	3.3	6.66	P
<u>Log (base e) Transform</u>						
All	115	2.11	2.24	0.66	8.22	F
0-15	46	2.635	2.621	0.365	13.94	F
15-30	46	1.859	1.962	0.252	6.42	F
30-45	23	1.538	1.685	0.662	4.66	F
> 15	69	1.752	1.834	0.576	5.77	G
<u>Square Root Transform</u>						
All	115	3.017	3.071	0.937	9.1	F
0-15	46	3.8	3.707	0.717	14.44	F
15-30	46	2.608	2.667	0.611	6.8	F
30-45	23	2.27	2.322	0.721	5.15	F
> 15	69	2.495	2.502	0.441	6.23	G

- a. Central tendency is the population mean for the raw data, the geometric mean for the log transform, and the square of the square root mean for the square root transform.
- b. The N P Index refers to a subjective evaluation of the linearity of the normal probability plots. Categories are Very Good, Good, Fair, and Poor.

Table A 2. Risdalsheia, distribution parameters for exchangeable Ca+Mg (meq/100g).

Depth	n	mean	Median	s.d.	Central Tendency ^a	NP Index ^b
<u>Raw data</u>						
All	115	1.54	0.84	1.81	1.54	VP
0-15	46	2.96	2.31	2.01	2.96	P
15-30	46	0.65	0.38	0.78	0.65	VP
30-45	23	0.48	0.19	0.65	0.48	VP
> 15	69	0.59	0.33	0.74	0.59	VP
<u>Log(base e) Transform</u>						
All	115	-0.39	-0.174	1.48	0.68	F
0-15	46	0.871	0.839	0.67	2.39	G
15-30	46	-1.06	-0.968	1.18	0.35	G
30-45	23	-1.57	-1.66	1.4	0.21	F
> 15	69	-1.23	-1.109	1.61	0.29	G
<u>Square Root Transform</u>						
All	115	1.045	0.917	0.671	1.09	VP
0-15	46	1.632	1.52	0.551	2.66	F
15-30	46	0.695	0.616	0.412	0.48	P
30-45	23	0.574	0.436	0.402	0.33	P
> 15	69	0.655	0.575	0.41	0.43	P

- a. Central tendency is the population mean for the raw data, the geometric mean for the log transform, and the square of the square root mean for the square root transform.
- b. The N P Index refers to a subjective evaluation of the linearity of the normal probability plots. Categories are Very Good, Good, Fair, and Poor.

Table A 3. Risdalsheia, distribution parameters for (Ca+Mg) saturation.

Depth	n	mean	Median	s.d.	Central Tendency ^a	NP Index ^b
<u>Raw data</u>						
All	115	0.118	0.101	0.087	0.118	VP
0-15	46	0.18	0.193	0.073	0.18	G
15-30	46	0.076	0.056	0.064	0.076	VP
30-45	23	0.065	0.031	0.057	0.065	VP
> 15	69	0.072	0.049	0.061	0.072	VP
<u>Log (base e) Transform</u>						
All	115	-2.496	-2.287	0.937	0.082	P
0-15	46	-1.766	-1.644	0.435	0.171	F
15-30	46	-2.92	-2.883	0.859	0.054	F
30-45	23	-3.106	-3.488	0.881	0.049	F
> 15	69	-2.982	-3.005	0.864	0.051	F
<u>Square Root Transform</u>						
All	115	0.317	0.319	0.131	0.1	P
0-15	46	0.423	0.44	0.087	0.178	G
15-30	46	0.254	0.237	0.109	0.064	VP
30-45	23	0.232	0.175	0.105	0.054	VP
> 15	69	0.247	0.222	0.107	0.061	P

- a. Central tendency is the population mean for the raw data, the geometric mean for the log transform, and the square of the square root mean for the square root transform.
- b. The N P Index refers to a subjective evaluation of the linearity of the normal probability plots. Categories are Very Good, Good, Fair, and Poor.

Table A 4. Risdalsheia, distribution parameters for K saturation.

Depth	n	mean	Median	s.d.	Central Tendency ^a	NP Index ^b
<u>Raw data</u>						
All	114	0.021	0.021	0.0090	0.021	P
0-15	46	0.029	0.028	0.0062	0.028	G
15-30	46	0.017	0.017	0.0074	0.017	F
30-45	23	0.014	0.014	0.0058	0.014	F
> 15	69	0.016	0.015	0.0067	0.016	F
<u>Log (base e) Transform</u>						
All	114	-3.954	-3.871	0.473	0.019	G
0-15	46	-3.572	-3.564	0.211	0.028	VG
15-30	46	-4.160	-4.098	0.420	0.016	G
30-45	22	-4.323	-4.230	0.416	0.013	G
> 15	68	-4.213	-4.177	0.423	0.015	G
<u>Square Root Transform</u>						
All	114	0.142	0.144	0.0317	0.020	F
0-15	46	0.169	0.168	0.0180	0.028	G
15-30	46	0.128	0.129	0.0267	0.016	G
30-45	22	0.118	0.121	0.0239	0.014	F
> 15	68	0.120	0.124	0.0261	0.016	G

- a. Central tendency is the population mean for the raw data, the geometric mean for the log transform, and the square of the square root mean for the square root transform.
- b. The N P Index refers to a subjective evaluation of the linearity of the normal probability plots. Categories are Very Good, Good, Fair, and Poor.

Table A 5. Risdalsheia, distribution parameters for density, (kg/l).

Depth	n	mean	Median	s.d.	Central Tendency ^a	NP Index ^b
<u>Raw data</u>						
All	80	0.524	0.45	0.356	0.524	P
0-15	35	0.263	0.24	0.149	0.263	F
15-30	33	0.72	0.62	0.334	0.72	G
30-45	12	0.746	0.67	0.361	0.746	G
> 15	45	0.727	0.62	0.338	0.727	F
<u>Log (base e) Transform</u>						
All	80	-0.9136	-0.799	0.79	0.401	P
0-15	35	-1.509	-1.427	0.63	0.221	G
15-30	33	-0.462	-0.478	0.568	0.63	F
30-45	12	-0.421	-0.406	0.553	0.656	F
> 15	45	-0.451	-0.478	0.558	0.637	P
<u>Square Root Transform</u>						
All	80	0.681	0.671	0.248	0.464	P
0-15	35	0.493	0.49	0.147	0.243	G
15-30	33	0.823	0.787	0.21	0.677	F
30-45	12	0.838	0.817	0.218	0.702	F
> 15	45	0.827	0.787	0.21	0.684	F

- a. Central tendency is the population mean for the raw data, the geometric mean for the log transform, and the square of the square root mean for the square root transform.
- b. The N P Index refers to a subjective evaluation of the linearity of the normal probability plots. Categories are Very Good, Good, Fair, and Poor.

Table A 6. Risdalsheia, distribution parameters for CEC (KCl) by volume (meq/l).

Depth	n	mean	Median	s.d.	Central Tendency ^a	NP Index ^b
<u>Raw data</u>						
All	80	36.5	35.4	16.15	36.5	P
0-15	35	32.55	29.4	13.72	32.55	P
15-30	33	43.1	39.11	18.18	43.1	G
30-45	12	29.95	29.45	10.13	29.95	G
> 15	45	39.58	36.48	17.33	39.58	F
<u>Log (base e) Transform</u>						
All	80	3.499	3.569	0.453	33.08	G
0-15	35	3.392	3.381	0.44	29.72	F
15-30	33	3.669	3.666	0.455	39.21	G
30-45	12	3.345	3.379	0.35	28.36	G
> 15	45	3.583	3.597	0.45	35.98	VG
<u>Square Root Transform</u>						
All	80	5.898	5.957	1.319	34.79	G
0-15	35	5.579	5.423	1.208	31.12	P
15-30	33	6.417	6.254	1.401	41.18	VG
30-45	12	5.4	5.421	0.93	29.16	G
> 15	45	6.146	6.04	1.361	37.77	G

- a. Central tendency is the population mean for the raw data, the geometric mean for the log transform, and the square of the square root mean for the square root transform.
- b. The N P Index refers to a subjective evaluation of the linearity of the normal probability plots. Categories are Very Good, Good, Fair, and Poor.

Acid Rain Research Reports

- 1/1982 Henriksen, A. 1982. Changes in base cation concentrations due to freshwater acidification. 50 pp. Out of print.
- 2/1982 Henriksen, A. and Andersen, S. 1982. Forsuringssituasjonen i Oslomarkas vann. 45 pp. Out of print.
- 3/1982 Henriksen, A. 1982. Preacidification pH-values in Norwegian rivers and lakes. 24 pp. Out of print.
- 4/1983 Wright, R.F. 1983. Predicting acidification of North American lakes. 165 pp.
- 5/1983 *Schoen, R., Wright, R.F. and Krieter, M.* 1983. Regional survey of freshwater acidification in West Germany (FRG). 15 pp.
- 6/1984 Wright, R.F. 1984. Changes in the chemistry of Lake Hovvatn, Norway, following liming and reacidification. 68 pp.
- 7/1985 Wright, R.F. 1985. RAIN project. Annual report for 1984. 39 pp.
- 8/1985 *Lotse, E. and Otabbong, E.* 1985. Physiochemical properties of soils at Risdalsheia and Sogndal: RAIN project. 48 pp.
- 9/1986 Wright, R.F. and Gjessing, E. 1986. RAIN project. Annual report for 1985. 33 pp.
- 10/1986 Wright, R.F., Gjessing, E., Semb, A. and Sletaune, B. 1986. RAIN project. Data report 1983-85. 62 pp.
- 11/1986 Henriksen, A., Røgeberg, E.J.S., Andersen, S. and Veidel, A. 1986. MOBILLAB-NIVA, a complete station for monitoring water quality. 44 pp.
- 12/1987 Røgeberg, E.J.S. 1987. A coulometric Gran titration method for the determination of strong and weak acids in freshwater. 28 pp.
- 13/1987 Wright, R.F. 1987. RAIN project. Annual report for 1986. 90 pp.
- 14/1988 Hauhs, M. 1988. Water and ion movement through a minicatchment at Risdalsheia, Norway. RAIN project. 74 pp.
- 15/1988 Gjessing, E., Grande, M. and Røgeberg, E.J.S. 1988. Natural organic Acids. Their Role in Freshwater Acidification and Aluminium Speciation. 28 pp.
- 16/1988 Wright, R.F. 1988. RAIN project. Annual report for 1987. 77 pp.
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