Interpretation of Serum Calcium in Patients with Abnormal Serum Proteins

R. B. PAYNE, A. J. LITTLE, R. B. WILLIAMS, J. R. MILNER

British Medical Journal, 1973, 4, 643-646

Summary

Two hundred consecutive specimens received in this laboratory for "liver function tests" showed a wide range of abnormal protein concentrations. Calcium concentration correlated closely with albumin (r = 0.867) but less closely with total protein (r = 0.682). A simple formula for adjusting calcium concentration was derived from the regression equation of calcium on albumin. Adjusted calcium = calcium - albumin + 4.0, where calcium is in mg/100 ml and albumin in g/100 ml.

Low calcium concentrations were found in 49 (24.5%) and raised concentrations in six (3%) of the 200 blood specimens taken for liver function tests. After adjustment, the 95% limits of the observed range were identical with the 95% limits of the normal range determined in this laboratory. Unlike adjustments based on total protein or specific gravity, the adjustment on albumin in 39 specimens which showed hypergammaglobulinaemia on electrophoresis gave normal calcium concentrations.

Introduction

The total calcium of serum or plasma is nearly all accounted for as calcium bound to protein and as calcium ions in roughly equal proportions, and it is the latter which are clinically important (McLean and Hastings, 1935). Measurement of ionized or diffusible calcium is at present technically complex (Rose, 1972; Rushton et al., 1973), so that a screening procedure to detect patients with abnormal protein concentrations and to allow for the effect of changes in protein-bound calcium on the measurement of total calcium is desirable (Dent, 1962).

There are statistically significant correlations between calcium and total protein and between calcium and albumin in normal subjects, but the correlation coefficients are low (less than 0.4) and because of the limited ranges of the observed values the regression lines cannot be extrapolated with confidence to low protein concentrations (Keating et al., 1969; Williams et al., 1973). For this reason we have examined the relation between calcium and total protein and between calcium and albumin in patients who show a wide range of abnormalities of protein concentration but who are likely to have a low prevalence of disorders of calcium homeostasis.

Material and Methods

Two hundred consecutive serum specimens received in the laboratory with a request for "liver function tests" but with no request for calcium estimation were analysed for calcium in addition to total protein and albumin on a Technicon S.M.A. Plus using cresolphthalein complexone for calcium, biuret for total protein, and bromocresol green in succinic acid buffer pH 4.2 for albumin. Duplicate analyses were not done. Measurements (in mg/100 ml and g/100 ml) were made to one decimal place. The specimens included all those received from wards, outpatient departments, and general practitioners, but excluded those from the department of renal medicine in this hospital because many of their patients were receiving therapy which would affect calcium concentration. No account was taken of age, sex, or provisional diagnosis. The between-batch coefficients of variation at normal serum concentrations during the period of the investigation were 1.2% for calcium,
1.5% for total protein, and 3.5% for albumin. Cellulose acetate electrophoresis was carried out on all specimens.

**Results**

The calcium concentrations in the 200 specimens received for liver function tests ranged from 7.1 to 10.8 mg/100 ml (fig. 1). The 95% limits of the normal range determined in the laboratory shortly before the present investigation were 9.0 to 10.4 mg/100 ml. Forty-nine (24.5%) of the specimens had low calcium concentrations and six (3%) had raised concentrations. The total protein concentrations ranged from 4.6 to 8.7 g/100 ml (normal range: 6.3 to 7.7 g/100 ml), and the albumin concentrations ranged from 2.0 to 4.8 g/100 ml (normal range 3.7 to 4.7 g/100 ml). Thirty-nine (19.5%) of the specimens showed hypergammaglobulinaemia on electrophoresis.

**RELATION BETWEEN CALCIUM AND TOTAL PROTEIN**

There was a significant correlation between calcium and total protein concentrations \( r = 0.682; P < 0.001 \). The regression equation of calcium on total protein was

\[
\text{Calcium} = (0.689 \times \text{total protein}) + 4.64
\]

(S.E. of regression coefficient 0.049).

The calcium values were adjusted so that their mean was the same as the mean of the normal range (9.70 mg/100 ml) by adding to the individual intercept values the difference between the mean intercept and the mean of the normal range by using the equation

\[
\text{Adjusted calcium} = \text{calcium} - (0.689 \times \text{total protein}) + 5.06
\]

The derived 95% limits after applying this adjustment were 8.7 to 10.7 mg/100 ml (table 1). The 39 hypergammaglobulinaemic sera gave relatively low values within the overall distribution (fig. 2) and largely accounted for the unacceptable width of the range.

**TABLE 1—Calcium Concentrations in 200 Specimens on which Liver Function Tests were Requested, before and after Adjustment for Protein Concentration**

<table>
<thead>
<tr>
<th></th>
<th>Mean (mg/100 ml)</th>
<th>S.D. (mg/100 ml)</th>
<th>95% Limits (mg/100 ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Measured values</td>
<td>9.37</td>
<td>0.698</td>
<td>8.0-10.8</td>
</tr>
<tr>
<td>Values adjusted on total protein (equation 1)</td>
<td>9.70</td>
<td>0.510</td>
<td>8.7-10.7</td>
</tr>
<tr>
<td>Values adjusted on albumin (equation 2)</td>
<td>9.66</td>
<td>0.347</td>
<td>9.0-10.4</td>
</tr>
</tbody>
</table>

An adjustment suggested by Zilva and Pannall (1971) has since been slightly amended (J. F. Zilva, personal communica-

**RELATION BETWEEN CALCIUM AND ALBUMIN**

There was a highly significant correlation between calcium and albumin concentrations \( r = 0.867; P < 0.001 \) and the correlation coefficient was significantly greater than that of calcium on total protein \( P < 0.001 \). The relation is shown in fig. 3. The regression equation of calcium on albumin was

\[
\text{Calcium} = (0.989 \times \text{albumin}) + 5.70
\]

(S.E. of regression coefficient 0.0405).

The equation to adjust the calcium values so that their mean
was the same as the mean of the normal range (9.70 mg/100 ml).

Adjusted calcium = calcium - (0.989 × albumin) + 4.0

or, with an error of less than 0.04 mg/100 ml in the adjusted values at the mean of the normal range

Adjusted calcium = calcium - albumin + 4.0 . . . . . 2

where calcium is in mg/100 ml and albumin in g/100 ml.

In S.I. units the equation becomes

Adjusted calcium = calcium - (0.25 × albumin) + 10

where calcium is in mmol/l and albumin in g/l.

The distribution of adjusted calcium values after applying equation 2 is shown in fig. 4. The range of values in the hypergammaglobulinaemic sera was similar to that of the group as a whole, and the 95% limits of the overall distribution, 9.0 to 10.4 mg/100 ml, were identical with those of the normal range (table I).

All the calcium values which remained low after adjustment for albumin, 9%, of the total, were consistent with the provisional diagnoses and other biochemical measurements. The three adjusted values which were raised were all from patients with carcinoma of the breast. One of these had been at the upper limit of normal before adjustment.

Discussion

A number of factors may alter the protein content of serum, and therefore the total calcium concentration, without necessarily affecting the non-protein bound, largely ionized, fraction. These include diseases in which the synthesis of serum proteins is increased or decreased, in which there is increased protein loss; prolonged venous occlusion which may cause "ultrafiltration" of blood in the capillaries of the forearm (van Leeuwen et al., 1961); and alterations of posture, which may cause shifts of fluid into or out of the intravascular compartment (Fawcett and Wynn, 1960).

Dent and Watson (1968) emphasized that the adjustment of total calcium concentration for changes in protein by using specific gravity measurements is inaccurate and should not be applied if electrophoresis shows the protein distribution to be abnormal. We have shown that adjustment for total protein concentration is unsatisfactory in patients with hypergammaglobulinaemia, and that abnormal globulin concentrations are not uncommon in patients with diagnostic problems concerned with calcium metabolism.

The present investigation was carried out on serum specimens on which liver function tests but not calcium had been requested, because we thought there would be a high prevalence of abnormal protein concentrations but a low prevalence of disorders of calcium homoeostasis. The data support this view: the 95% limits of the calcium concentrations after adjustment for the regression of calcium on albumin were identical with the limits of the normal range in healthy persons, and the Gaussian shape of the distribution suggests that contamination with calcium values of diagnostic significance was minimal (cf. Cook et al., 1970).

Our regression coefficient of calcium on total protein (0.689) is of the same order as the regression coefficient derived from the calcium adjustment of Dent (1962) based on specific gravity (0.676). However, our regression coefficient of calcium on albumin, 0.898, is considerably higher than the coefficient of 0.707 determined by Orrell (1971) on 954 specimens received in his laboratory either for albumin or calcium determination, though our relation between calcium and albumin is of the same order as that shown graphically by Jones et al. (1967) in patients with such disorders as malabsorption syndrome, cirrhosis, and malnutrition, and by Cockel et al. (1971) in patients with rheumatoid arthritis.

There are two factors which might have contributed to this discrepancy. The first is that the series of Orrell (1971), unlike the present series, is likely to have included specimens from patients with the nephrotic syndrome. There is evidence that such patients have higher calcium levels than patients with the same degree of hypoalbuminaemia from other causes (Danowski et al., 1957; Jones et al., 1967), possibly due to the presence in nephrotic sera of other macromolecules which bind calcium more effectively than albumin (Pedersen, 1972).
The second factor is that Orrell's regression coefficient was based on specimens on which either albumin or calcium estimation had been requested, and so would have included some abnormal calcium values not due to protein abnormalities. In our experience true hypocalcaemia is more common than hypercalcaemia. Thus his data would have been weighted with low albumin values associated with relatively higher calcium values from patients with the nephrotic syndrome and with low calcium values with normal albumin values from patients with, for example, osteomalacia. These differences would account both for our higher regression coefficient and for our higher correlation coefficient (0.867 on 100 pairs compared with his of 0.720 on 954 pairs: P < 0.001). Our intercept at zero albumin concentration (5.7 mg/100 ml) is closer to reported values for normal plasma ultrafilterable calcium (Robertson, 1969; Pedersen, 1970; Roche, 1972) than the intercept calculated from Orrell's (1971) data (6.8 mg/100 ml).

Our equation for adjusting calcium concentration by using measurements of albumin

\[
\text{Adjusted calcium} = \text{calcium} - \text{albumin} + 4.0
\]

was based on correction to the mean normal calcium concentration rather than to the mean normal albumin concentration because the between-batch coefficients of variation at normal serum concentrations were 1.2% for calcium and 3.5% for albumin, so that we had greater confidence in the accuracy of the mean normal calcium concentration. The adjustment is easily remembered: the measured calcium concentration is increased by 0.1 mg/100 ml for every 0.1 g/100 ml that the albumin concentration is below 4.0 g/100 ml, and vice versa. It should be noted that the adjustment cannot be applied to calcium values on patients with the nephrotic syndrome and hypoalbuminaemia, for the reasons discussed above, nor can it be applied to data from another laboratory if the accuracy and precision of calcium and albumin measurements, and therefore the normal ranges, differ greatly from ours (Payne, 1973).

Dermatoglyphics in Children with Acute Leukaemia

S. G. PURVIS-SMITH, MARGARET A. MENSER


Summary

The dermatoglyphics of 135 children with acute leukaemia differed significantly from those of normal controls, and examination of 174 of the patients' first degree relatives indicated that familial factors were involved. The findings suggested that within the racial group studied dermatoglyphics may partly identify a population subgroup which is at increased risk of leukaemogenesis. While these observations may not have immediate clinical application, they are likely to contribute to a greater understanding of individuals who have increased constitutional susceptibility to leukaemia.

Introduction

Though there have been a number of apparently conflicting studies on the fingerprints and handprints (dermatoglyphics) of patients with leukaemia (Aleksandrowicz et al., 1966; Kobayashi et al., 1968; Carvalho, 1969; Menser and Purvis-Smith, 1969, 1972; Nora et al., 1969; Rosner, 1969, 1970; Wertelecki et al., 1969, 1973; Stowens and Sammon, 1970; Verbov, 1970; Wittwer and Giessmann, 1970; Zahálková and Břušuľová, 1970; Berka et al., 1971), careful review of the reports suggests that the dermatoglyphics of children with acute leukaemia deviate fairly consistently from normal. Such findings, if confirmed, may be valuable in identifying populations with increased susceptibility to leukaemogenesis.

Subjects and Methods

Handprints were obtained from 135 Caucasian children (77 males, 58 females) with a proved diagnosis of either acute lymphatic leukaemia or acute blast cell leukaemia. The patients ranged in age from 4 months to 15 years; the mean age at onset of the disease was 5.26 years in the males and 5.38 years in the females. The patient group included those from earlier studies.