CALCINOSIS
HISTOLOGICAL AND CHEMICAL ANALYSIS

BY
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This paper is concerned with the histological appearances and the chemical analysis of calcinotic tissue. The object was to discover which tissue constituents become calcified in this condition and what modification of the tissue is associated with the calcification process.

MATERIALS AND METHODS

Tissue was obtained from seven cases of calcinosis at autopsy or biopsy. Of these (clinical data given by Ansell, Hamilton and Bywaters, 1964), four showed calcinosis associated with frank dermatomyositis; another two showed collodion patches as the only evidence of dermatomyositis; the last case showed only an isolated lesion of calcinosis post mortem. Part of each specimen was fixed in formalin for paraffin sections, while a thin slice was rapidly frozen in a mixture of solid carbon dioxide and alcohol for sectioning at -20°C. In other cases, thin slices of fresh tissue were hand-cut for incubation in enzyme, solutions, as described below, and subsequently fixed and sectioned.

Decalcification of sections or slices of tissue was usually carried out in a 10 per cent. solution of E.D.T.A., adjusted to pH 7-0, but some sections were decalcified in acid.

Sections were stained by haematoxylin and eosin, by the Picro-Mallory technique, Heidenhain's azan, and von Kossa's or the chloranilic acid method for calcium salts. Other stains used were orcein, the periodic acid—Schiff (PAS) method, Feulgen, and azure A. Some sections were stained for tyrosine by the method of Lillie (1957).

Enzymes.—Proteolytic and mucolytic fractions of elastase were kindly supplied by Dr. D. A. Hall of the School of Medicine, Leeds (Hall, 1955). The two fractions were used in combination at a ratio of five parts of the proteolytic to one of the mucolytic fraction. The enzyme was dissolved at a concentration of 1 mg. per ml. in M/15 phosphate buffer, pH 8-4. Collagenase ABC Form II was obtained from Agricultural Biologicals Corporation. It was dissolved at 1 mg. per ml. in a solution of 0-1 M NaCl, 0-1 M CaCl₂, and 0-05 M borate at pH 7-0. Crystalline trypsin (Armour Laboratories, Inc.) was used at a concentration of 1 mg. per ml. in M/15 phosphate buffer at pH 8-0.

Sections and slices of tissue were left in contact with enzyme or buffer control solution for periods of several hours, at 37°C., in a moist chamber.

Chemical Procedures.—Finely divided tissue was dried in a desiccator. Fat extraction was carried out with petroleum ether (b.p. 40-60°C). Acid hydrolysis was carried out in closed ampoules at 100°C. Nitrogen was determined by micro-Kjeldahl, hexosamine by a modification of the method of Elson and Morgan (1933), hydroxyproline by the method of Neuman and Logan (1950), and tyrosine by the method of Udenfriend and Cooper (1952). Calcium and phosphorus were estimated by standard methods described by King (1951). Two-dimensional chromatograms for amino acids were kindly carried out by Dr. R. Consden.

Results and Discussion

Morphology and Staining Characteristics

The characteristic appearance of the calcinotic lesion is shown in Figs 1, 2, and 3 (opposite). The spaces, irregular in outline and located amongst connective tissue, were filled by amorphous calcified material. In the material described in this paper we have only seen one instance in which the appearances suggested calcification of muscle fibre remnants (Fig. 4, overleaf). Cellularity of the tissue immediately surrounding the foci of calcinosis was variable. In some of our cases, the dense connective tissue was almost acellular while in others in the vicinity of foci of calcification there were considerable numbers of cells having the appearance of fibroblasts. Inflammatory cells, such as polymorphs, lymphocytes, or plasma cells, were only very rarely seen in this situation. At some distance, however, from the calcific foci, accumulations of inflammatory round cells could quite frequently be found, often surrounding small blood vessels.
Fig. 1.—Subcutaneous lesion of calcinosis. Biopsy from male aged 17 suffering from dermatomyositis. The calcified material is in the midst of relatively acellular connective tissue. Haematoxylin and eosin. × 85.

Fig. 2.—Subcutaneous calcinotic lesion (biopsy) from male aged 7 with a collodion patch as the only evidence of dermatomyositis. Section shows many fibroblasts along edge of calcified tissue. Haematoxylin and eosin. × 85.

Fig. 3.—Calcinotic lesion from deep intermuscular connective tissue of the leg. Biopsy from same patient as in Fig. 1. Connective tissue shows very few cells. Haematoxylin and eosin. × 200.
In the vicinity of some calcinotic foci, there were fibris showing the morphological characteristics of elastic fibres, although they were frequently more fragmented and bulky than the normal elastic fibres of connective tissue. These fibres could be stained by both the orcein and von Kossa methods (Figs 5 and 6, opposite).

While it must be borne in mind that abnormal collagen fibres may take the orcein stain (Loewi, Glynn, and Dorling, 1960), their situation and morphological appearances strongly suggested that these were calcified elastic fibres: calcification of these fibres may represent an early stage in the lesion of calcinosis.

Staining by the Picro-Mallory technique showed blue calcinotic material with the surrounding connective tissue similarly stained blue, with some red fibrils. Azan gave a similar blue colour to calcinotic foci. Both methods showed small red-stained fragments among the predominantly blue matrix. Collagen fibres are stained blue by these techniques, but as there were no morphologically distinct fibres in the calcinotic matrix, it was not possible to conclude the collagenous nature of the material. Staining by the PAS reaction produced intense red staining of the calcinotic matrix. Prior decalcification did not affect these staining results but, owing to the acidic conditions used with the Picro-Mallory, Azan, and PAS methods, some decalcification must have occurred during staining. Orcein stained undecalcified calcinotic matrix brown with an intensity similar to that shown by elastin. After decalcification, some areas retained this staining property, but others were coloured a light blue similar to that of surrounding collagen. With Azure A there was slight patchy metachromasia of calcinotic matrix, with orthochromasia of surrounding connective tissue. The Feulgen reaction served to demonstrate a small number of nuclei at the periphery of calcinotic areas, but failed to stain the bulk of the material. The reaction for tyrosine was similar in intensity in the calcinotic and in the surrounding connective tissue. Staining for fat by the oil-Red-O method showed a small amount of stained material irregularly distributed in calcinotic foci. Before decalcification, the Gomori technique showed degrees of staining of the calcinotic matrix as seen with collagen without showing actual fibres. After decalcification, much of this staining became less intense or changed to grey, but some small argyrophil fibrils and clumps were found amongst the amorphous material. Staining by chloranilic acid (Fig. 7, opposite) and by von Kossa's method confirmed the presence of calcium salts.

The results of staining were therefore inconclusive with regard to the nature of the calcinotic matrix, but they suggested that the material was not predominantly polysaccharide or fat, but more probably protein.

Fig. 4.—Muscle and intermuscular connective tissue from same patient as in Fig. 2. Right-hand side of picture shows calcified bundles, probably of muscular origin. Haematoxylin and eosin. × 150.
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Fig. 5.—Edge of subcutaneous calcinotic lesion. Post mortem specimen from female aged 14 suffering from dermatomyositis. Fibrils stained by von Kossa's method for calcium salts. The calcinotic areas and fibrils are blackened. × 200.

Fig. 6.—As Fig. 5, but stained by orcein. × 200.

Fig. 7.—Calcinotic lesion. Same patient as in Fig. 1. Calcium shown by chloranilic acid method. × 200.
Enzymatic Digestion.—Lesions from three patients with calcinosis were studied, giving similar results. One had dermatomyositis, another showed collodion patches as the only sign of dermatomyositis, and the third had only an isolated patch of calcinosis.

When collagenase, trypsin, or elastase was used on fresh calcinotic tissue, calcified material was not broken down, and the surrounding tissue was attacked in the expected way, i.e. collagen was broken down by collagenase and elastic fibres by elastase; trypsin affected staining and caused blurring of structures, but left the calcified tissue intact (Fig. 8). On the other hand, when incubation with enzyme was preceded by decalcification by E.D.T.A., an enzymatic effect on the matrix was seen. Both elastase and trypsin caused partial or, in some cases, complete removal of the material after such decalcification (Figs 9 and 10), unlike their effects on native collagen. Holes were left in the tissue in places where the calcinotic material had been. The surrounding tissue framework was left behind, although its staining characteristics and structure were affected by the treatment. Incubation in buffer alone did not remove the matrix. Collagenase, on the other hand, did not cause removal of decalcified matrix, although the collagen of the surrounding tissue was attacked and much of it disappeared (Fig. 11, opposite). Treatment with E.D.T.A. only or incubation with buffers after decalcification did not lead to removal of residual matrix, but only to some diminution in intensity of staining.

Two conclusions can be drawn from these results:

1. Proteolytic enzymes could only attack the tissue after decalcification. This is in agreement with the work of Fawns and Landells (1953).

2. Digestion by trypsin and elastase indicates the protein nature of the matrix.

Elastase, in spite of its name, attacks not only elastin, but also other proteins, (Fullmer and Lillie, 1958; Bagdy, 1959). Trypsin has a wide specificity for proteins, but does not digest native collagen. When collagen has been treated by acetylation,
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salicylates, or heat, however, it becomes susceptible to elastase and trypsin (Loewi and others, 1960). The resistance of calcinotic matrix to collagenase may imply either that the material is not predominantly collagen or that collagen has been changed in such a way that it has lost susceptibility to collagenase (Loewi and others, 1960). That calcification per se does not cause collagen to change its susceptibilities to these three enzymes was shown by treating bone slices similarly. The bony matrix was not attacked without prior decalcification, but after decalcification the matrix was digested by collagenase (Fig. 12, overleaf), but left intact by trypsin or elastase (Fig. 13, overleaf).

Chemical Analysis.—The analysis which is detailed here, was performed on material obtained from a patient who had an isolated patch of subcutaneous calcinosis and died from an unrelated cause. The calcified core of the tissue was dissected, free as far as possible from the surrounding connective tissue and the two collections of tissue so obtained analysed separately. The histological appearances of this lesion and results of enzymatic digestion were as described above. The results of analysis are shown in Table I, together with some data from a second case of calcinosis, in which the lesions had been associated with dermatomyositis.

### Table I

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Tissue</th>
<th>1</th>
<th>2</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>Calcinotic</td>
<td>Surrounding Connective</td>
</tr>
<tr>
<td>Water</td>
<td>45</td>
<td>84</td>
<td>50</td>
</tr>
<tr>
<td>Protein*</td>
<td>7-7</td>
<td>8-7</td>
<td></td>
</tr>
<tr>
<td>Calcium</td>
<td>15-4</td>
<td>0-4</td>
<td>13-0</td>
</tr>
<tr>
<td>Fat</td>
<td>2-2</td>
<td>3-0</td>
<td>1-8</td>
</tr>
<tr>
<td>Hexosamine</td>
<td>0-2</td>
<td>0-1</td>
<td></td>
</tr>
<tr>
<td>OH-proline N(_2) (as per cent. of Total N(_2))</td>
<td>7-35</td>
<td>5-25</td>
<td></td>
</tr>
<tr>
<td>Tyrosine N(_2) (as per cent. of Total N(_2))</td>
<td>0-97</td>
<td>0-94</td>
<td></td>
</tr>
</tbody>
</table>

Weights expressed as mg./100 mg. wet tissue.
* Obtained by multiplying figure for total N\(_2\) by 6-25.

The figures show a decreased moisture content as well as somewhat less protein in the calcinotic as
opposed to the surrounding connective tissue. Calcium, when calculated as hydroxy-apatite, the form in which it is probably present, accounts for nearly 40 per cent. of the weight of the tissue, bringing the total of five constituents in Table I (Case 1) to 100 per cent. approximately. The proportion of protein present as collagen is greater in the calcinotic material than in the surrounding tissue; that present as tyrosine is similar in the two samples, indicating a similar content of plasma proteins. Tissue hexosamine is contributed from various sources, but the figures suggest that there is no excessive amount of polysaccharide in the tissue.

Since it is possible to distinguish collagen from non-collagenous protein by its solubility in water at 100°C under pressure (Neuman and Logan, 1950), samples of tissue were so treated. The results shown in Table II confirm that most of the protein present in the calcinotic focus is a form of collagen. Amino-acid chromatograms showed a pattern con-

<table>
<thead>
<tr>
<th>Tissue</th>
<th>Calcinotic</th>
<th>Surrounding Connective</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Supernatant</td>
<td>Residue</td>
</tr>
<tr>
<td>Total N2</td>
<td>2.4</td>
<td>0.5</td>
</tr>
<tr>
<td>OH-proline N2 (as per cent. of total N2)</td>
<td>7.6</td>
<td>1.7</td>
</tr>
<tr>
<td>Ca</td>
<td>0.25</td>
<td>25.9</td>
</tr>
<tr>
<td>P</td>
<td>0.03</td>
<td>4.2</td>
</tr>
<tr>
<td>NH2-acid Chromatogram</td>
<td>Consistent with collagen</td>
<td>Not done</td>
</tr>
<tr>
<td></td>
<td>Consistent with collagen</td>
<td>Consistent with elastin + residual collagen</td>
</tr>
</tbody>
</table>

Weights given as mg./100 mg. dry tissue autoclaved.
sistent with collagen; in two other cases, less extensively studied, the chromatograms showed some hydroxyproline, thus providing evidence for collagen.

The results of chemical analysis therefore suggest that the calcified matrix in calcinosis might be altered collagen. Calcification might take place in collagen which has undergone change and also on some elastic fibres. The failure of digestion by collagenase may be compared with the findings of Loewi and others (1960), that the connective tissue of senile elastosis was refractory to the action of collagenase and that experimental acetylation of collagen similarly led to a collagenase-resistant material, thus showing that collagen may undergo changes which alter its enzyme susceptibility. That collagen may be the focus for calcification has been shown for normal bone and dentine by the work of Solomons and Irving (1958) and for collagen implants by Mergenhagen, Martin, Rizzo, Wright, and Scott (1960). In an accompanying communication, Davis and colleagues (paper in preparation) show that apatite crystals can be seen by electron microscopy to be associated with collagen in calcinosis.

Summary

Tissue from cases of calcinosis, with and without dermatomyositis, obtained at biopsy or post mortem, has been examined. It was noted that the calcified masses appeared in connective tissue, often in the neighbourhood of muscle. In some cases the tissue surrounding these masses contained small calcified fibrils giving staining reactions of elastin. Most lesions showed little cellular reaction. In an effort to elucidate the nature of the tissue undergoing calcification, histochemical, histo-enzymatic, and analytical chemical procedures were used. Trypsin and elastase were effective in digesting the calcinotic matrix only after decalcification. Collagenase was without effect. Analysis showed a high hydroxyproline content, from which it must be deduced that the protein of the calcinotic matrix consists largely of collagen, although no fibres could be seen by light microscopy. The questions why such collagen undergoes calcification and why it is not digested by collagenase have not yet been elucidated.

REFERENCES


