

calcification is imminent, and its intensity is perhaps related to the rapidity of the calcification that follows.

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Histotopochemistry of Ascorbic Acid in Tendon Fibres

FROM investigations with the electron microscope, it has been reported that the precursors of collagen fibrils are formed within the cytoplasm of fibrocytes¹. It follows that the sites of formation of the fibrils and those of the synthesis of collagen are identical. Hydroxyproline, which is a characteristic acid of collagen protein, originates from proline under the action of L-ascorbic acid².

For this reason we believe that determination of the distribution and concentration of ascorbic acid in the connective tissue, for example in the tendon, would suggest where hydroxylation of proline takes place and where the tropocollagenous cells, which are the basic elements of the fibrils, are formed.

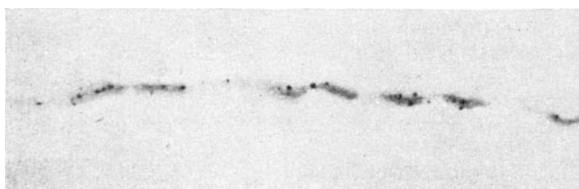


Fig. 1. Black granules of metallic silver in the tendon cells indicate L-ascorbic acid, which takes part in the conversion of proline to hydroxyproline

In this work L-ascorbic acid was determined in isolated fibres from the tail tendon of male albino Wistar rats, 4 months or 28 months old. The method of Giroud and Leblond, applying silver nitrate to metallic silver, was used. The granules occurring in the positive reaction are not granules of vitamin C but granules of metallic silver³.

It has been found that in isolated tendon fibres of both the younger and the older rats the granules of silver are deposited only in the cytoplasm of fibrocytes; the remaining fibrillar structure is negative.

The occurrence of L-ascorbic acid in the tendon cells which intensively produce collagen supports the suggestion that conversion of proline to hydroxyproline occurs within the cytoplasm of the cells, and that a collagen molecule may there be built up from the individual amino-acids.

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PATHOLOGY

An Unidentified Virus which causes the Rapid Production of Tumours in Mice

IN the course of routine passage of Moloney's leukaemogenic virus (MLV) plasma was collected from a leukaemic rat (Chester Beatty Institute outbred albino strain) which had been inoculated with MLV-containing mouse plasma when new-born. After storage at -70°C for three months the rat plasma was diluted 1 in 30 with Hanks's saline and passed through a 'Selas 02' filter, tested and found impervious to *Esch. coli*. The filtrate was injected into 15 new-born BALB/c mice, as a test of potency. Only 6 survived to weaning. On the 32nd day 5 had tumours at or near the injection site, and all had grossly enlarged spleens. Usually mice inoculated with MLV remain symptomless until they show the characteristic signs of leukaemia after 8 weeks.

Further samples of the same stored plasma filtrate were injected into new-born BALB/c mice, and Chester Beatty albino (outbred) and hooded (inbred) rats. Twenty-eight of 35 mice, and 1 of 13 rats, killed and examined *post mortem* had tumours, and all had splenomegaly.

The pathology of the disease is being examined: the result of early observations are as follows. The animals commonly die of splenic rupture, as in Friend or Rauscher disease. The histology of the spleen and the blood smears are similar to those of Friend disease. There is gross proliferation of reticulum cells in the spleen, and reticulum cells and erythroblasts appear in the blood. The tumours are of two main types, either solid and firm, or cystic and filled with blood. The former arise in the subcutaneous tissue and peritoneum, sometimes attached to the muscle of the abdominal wall, thorax, or diaphragm; their position suggests development at or near the site of injection or along the needle track. They are anaplastic sarcomata consisting of pleomorphic and spindle cells, some very large, invading and disrupting adjacent tissues. The latter may occur anywhere in the subcutaneous space or peritoneal cavity, and often in relation to lymph nodes. They are angiomatous tumours consisting of multiple dilated sinuses which are filled with blood and debris, and lined with cuboidal cells which often form ingrowths into the lumen.

Recent experiments have shown that of a number of animals injected with sarcoma virus (SV) when new-born 30/46 CB hooded rats developed gross splenomegaly, and 16 had sarcomas of the diaphragm or other sites. 4/9 cream and 5/19 golden hamsters have developed sarcomas, also mainly situated on or near the diaphragm. Many of the rats and hamsters had large thin-walled cysts at lymph node sites containing either clear or blood-stained fluid.

Two of 5 attempts to transplant sarcomas in BALB/c mice were successful, and the cell line is now in its fifth passage. All recipients have developed splenomegaly.

Tissues of tumour-bearing mice and plasma of injected rats were tested by inoculation into BALB/c mice for the presence of tumour-producing virus (Table 1).

'Selas 02' filtrates of tumour homogenates or plasma induced rapid development of tumours and splenomegaly in mice inoculated when 1-7 days old, in some cases as early as 12 days after injection. As the age of injection increased, the incidence of tumours declined, but that of splenomegaly remained high. Fifteen mice inoculated at 112 days all developed splenomegaly, but only two developed tumours.

One to 2 day-old Chester Beatty albino rats were injected with the original plasma filtrate. Their plasma was collected at 8, 30, and 54 days, and injected into 1-3 day-old BALB/c mice. Mice which received the 30- and 54-day samples developed tumours and splenomegaly in 12-15 days. Those which received the 8-day sample