Collagen stimulating factors in hepatic fibrogenesis

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SUMMARY Four factors which stimulate collagen synthesis and prolyl hydroxylase activity in cultures of human and mouse fibroblasts have been isolated by molecular sieve chromatography from animal and human fibrotic and cirrhotic livers. These factors do not stimulate protein or DNA synthesis or total DNA in these cultures. It has also been shown that these factors, designated collagen stimulating factors F1–F4, do not owe their activity to ascorbate or glutamine. Collagen stimulating factors are heat stable, and F1 and F2 have apparent molecular weights of about 4000 and 1000 respectively. Since these factors are not present in normal animal or human liver it is suggested that they may be responsible for increased collagen production in vivo in hepatic fibrosis and cirrhosis.

The collagen content of the liver increases four to five fold in hepatic cirrhosis.1 This is at least partially due to an increase in collagen synthesis in those diseases which progress to cirrhosis. Collagen deposition in the form of fibrous septa leads to disorganisation of the microanatomy of the liver, intrahepatic shunting, and portal hypertension. Its accumulation in the space of Disse may also interfere with metabolic exchanges between the sinusoidal vascular compartment and the microvillous surface of the hepatocyte.3

In experimental hepatic fibrosis a group of low molecular weight compounds, collagen stimulating factors, have been isolated which stimulate collagen synthesis and prolyl hydroxylase (EC 1.14.11.2) activity in fibroblasts in vitro.4 Prolly hydroxylase catalyses the biosynthesis of hydroxyproline, which is necessary for the formation of a stable collagen triple helix.

We have shown that collagen stimulating factors are present in human cirrhotic liver as well as in experimental liver disease but not in normal liver. We have also shown that they stimulate collagen synthesis specifically and that they differ in the physicochemical and biological properties from other compounds which have been postulated to modulate collagen synthesis in liver disease.

Material and methods

All chemicals, of the highest purity obtainable, were purchased from British Drug Houses, Poole, Dorset, or Sigma Chemical Co Ltd, London; all radioisotopes were obtained from the Radiochemical Centre, Amersham; scintillation fluid was obtained from Nuclear Enterprises, Sighthill, Edinburgh; and all reagents for tissue culture and cells were obtained from Gibco Biocult, Paisley, Scotland.

ISOLATION OF COLLAGEN STIMULATING FACTORS

Collagen stimulating factors were isolated from murine liver by a modification of a method described earlier. Acute liver damage was produced in male albino mice (strain CFLP) weighing roughly 20 g. A single dose (0-25 ml) of a 40% solution of carbon tetrachloride in liquid paraffin was administered by oesophageal tube. Three days later six animals were killed by cervical dislocation and the livers were removed. After excision of the gall bladder, the livers were placed on ice and all subsequent operations were performed at 4°C. The tissue was chopped finely with scissors and homogenised in 2-5 volumes (wt/vol) of ice cold 0-05 M sodium phosphate buffer pH 7-0 containing 10-3 M dithiothreitol, 10-4 M edetic acid, and 0-15 M NaCl (homogenisation buffer) in a close fitting ground glass homogeniser. The homogenate was centrifuged at 10 000g for 30 min and the supernatant, minus the lipid layer, was heated at 56°C for 20 min.

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Accepted for publication 7 December 1983
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in a polypropylene vial and centrifuged for a further 30 min at 10,000 g. The supernatant was passed through a glass column (90 × 2.5 cm) of G25 Sephadex equilibrated in homogenisation buffer. The column was eluted with the same buffer and 12 ml fractions were collected at a flow rate of 30 ml/hour at 4°C. The absorbance of each fraction was monitored at 280 nM and selected fractions were assayed for their ability to stimulate collagen biosynthesis and prolyl hydroxylase activity in L-929 cultures.

Collagen stimulating factors were isolated from human liver with slight modifications of the above procedure. Liver biopsy specimens from a patient with alcoholic cirrhosis (2.5 g) and from a patient without clinical or biochemical evidence of liver disease (1.8 g) were obtained at laparotomy for these experiments. Collagen stimulating factors were extracted as above in 5 volumes (wt/vol) of homogenisation buffer, applied to a G25 Sephadex column (70 × 1.5 cm), and eluted with homogenisation buffer, and 6 ml fractions were collected. Collagen stimulating factor activity was assayed by monitoring prolyl hydroxylase activity in human lung fibroblasts (W1–38 cells).

COLLAGEN SYNTHESIS IN CELL CULTURES

Mouse skin fibroblasts (cell line L-929) were grown in 120 cm² glass flasks in 25 ml of minimal essential medium buffered with 20 mM Hepes containing Earls salts, 10% fetal calf serum, 2 × 10⁻⁵ M sodium-L-ascorbate, 4 × 10⁻⁴ M ferric nitrate, 2 × 10⁻³ M L-glutamine, and 50 units/ml of penicillin/streptomycin and incubated under air at 37°C. For assay of collagen synthesis the cells were harvested from stock bottles by addition of 0.025% trypsin; after counting they were seeded at 0.4 × 10⁶/cm² in 25 cm² plastic Falcon flasks in 5 ml of the same medium.

Cells were grown for 40 h before exposure to selected fractions of column effluents. Aliquots (1 ml) of the latter were added to the culture for 5 h and then each culture was pulsed for 1 h with 10 μCi of 3, 4 ³H-proline (SA 40 Ci/mmole) before harvesting. After removal of medium the cell layer was washed twice with 1 ml of 0.15 M NaCl and the cells scraped from the flask and resuspended to 10⁶/ml in 0.05 M Tris buffer pH 7.6. The cells were then disrupted by sonication and an aliquot (0.5 ml) digested with 20 μg/ml of RNase for 10 min at 30°C to remove any radioactive prolyl t-RNA. An equal volume of 20% trichloroacetic acid (TCA) was added to precipitate protein and the mix was allowed to stand at 4°C for 20 min. The pellet was separated by centrifugation at 3000 g for 10 min and washed a further five times in 5% TCA before being solubilised in 0.1 M NaOH. The sample was then divided into two aliquots of 0.2 ml and incubated in a reaction mix containing the following: 0.1 ml of N-ethylmaleimide, 5 μl of 0.05 M calcium chloride, 10–20 μl of purified bacterial collagenase, and water to a final volume of 0.5 ml; control tubes contained 0.2 ml of solubilised cell pellet and all other reagents but no collagenase. Samples were incubated at 37°C for 60 min and the reaction was terminated by adding 0.5 ml of 0.5% tannic acid in 10% TCA. After incubation at 4°C for 30 min the samples were centrifuged at 3000 g for 15 min. Radioactive collagen peptides in the supernatant were measured by liquid scintillation counting in NE260 scintillant. The collagenase used in this assay was repurified before use and degraded only collagenous proteins.

PROTEIN SYNTHESIS IN CELL CULTURES

An aliquot of cell sonicate prepared as described above was digested with RNase and then spotted on to a filter paper disc (Whatmann 3MM) and allowed to dry. The disc was immersed in 20% TCA for 20 min followed by sequential washings in 10% and 5% TCA and finally extracted into ether and allowed to dry. The disc was then placed in 10 ml of Bray’s solution and counted.

PROLYL HYDROXYLASE IN CELL CULTURES AND HUMAN LIVER

L-929 cells or W1–38 (human) fibroblasts, grown as described above, were seeded at 0.2 × 10⁶/cm² in 25 cm² Falcon flasks and incubated for 40 h before the addition of collagen stimulating factors. The cultures were then incubated for 6 h. After discarding the medium the cell layer was washed twice with 1 ml of saline and the cells were removed mechanically into 1 ml of saline. The cells were suspended to 10⁶/ml in 0.25 M sucrose containing 10⁻⁴ M dithiothreitol and 10⁻⁵ M edetic acid, sonicated, and an aliquot assayed for prolyl hydroxylase activity using the tritium release assay. L-929 cells and W1–38 fibroblasts were used for assay of mouse and human collagen stimulating factors respectively.

Prolyl hydroxylase activity in human liver was measured as described previously.

DNA SYNTHESIS IN CELL CULTURES

Total DNA in fibroblast cultures was measured by the method of Burton. DNA synthesis in cultured cells was measured by the incorporation of ³H-deoxyribonucleic acid (SA 7.6 Ci/mmole) into acid precipitable nucleic acid. Cultures were incubated with collagen stimulating factors for 6–48 h and then pulsed for 1 h with ³H thymidine before harvesting. After washing the cell pellet in 0.1 M NaCl the cultures were extracted with 2% perchloroacetic acid at
ASSAY OF ASCORBIC ACID IN G25 SEPHADEX COLUMN EFFLUENTS

For estimating ascorbic acid in column fractions the columns were run in buffer without dithiothreitol or edetic acid. This had no effect on the activity of collagen stimulating factors; these reagents may play some role in the temporal stabilisation of collagen stimulating factors. Ascorbic acid was detected by the method of Zannoni et al.9 This assay is based on the ability of ascorbic acid to reduce Fe3+ to Fe2+, which, when coupled to α, α'-dipyridyl, produces a colorimetric reaction. Column effluent (1 ml) was mixed with 0.5 ml of 15% TCA and the precipitate was spun out at 3000 g. The supernatant was added to a mix containing the following: 0.1 ml of 85% orthophosphoric acid, 0.8 ml of 10% aqueous α, α'-dipyridyl, and 0.1 ml of ferric chloride in a final volume of 2.5 ml. All reagents were freshly prepared for each experiment. Sodium-l-ascorbate was used as a standard. The colorimetric reaction was developed at room temperature for 30 min and absorbance was read at 430 nm.

IMMUNOHISTOCHEMICAL LOCALISATION OF COLLAGEN IN MOUSE LIVER

Sheep antiserum to acetic acid soluble rat collagen was prepared as described previously.9 The anticollagen antibody in this serum was purified by affinity chromatography on a column of rat type I collagen linked to cyanogen bromide Sepharose 6B.9 The purified antibody reacted equally well with type I and type III collagen. Blocks 3–4 μm thick were taken from mouse liver and rapidly frozen on to microscope slides. Sections from normal mouse liver and mouse liver damaged by carbon tetrachloride were cut at −20°C and placed on glass slides. The slides were washed with 0.01 M phosphate buffer pH 7–2 containing 0.15 M NaCl (PBS), 20% normal swine serum, and incubated in the same solution for 5–10 min. Slides were dried and treated with anticollagen antibody (100 μg IgG/ml) for 30 min at room temperature. Sections were washed three times for 4 min in three changes of PBS and treated with rabbit antigoat fluorescent antibody (Dakopatts) for 30 min at room temperature; the antigoat antibody, which cross reacts with sheep IgG, was diluted 1/30 in PBS. After a further three washes in PBS, the sections were mounted in 90% glycerol/PBS and viewed under ultraviolet light. Control sections were treated with normal sheep IgG (100 μg/ml) followed by fluorescein labelled rabbit antigoat serum or with the latter reagent alone.

Results

In normal liver the anticollagen antibody reacted with the collagen around central and portal veins and with sinusoidal fibres (Fig. 1a). Three days after carbon tetrachloride injury there was a considerable increase in collagen around the central zones of the liver.

Fig. 1 (a) Distribution of collagen in normal mouse liver determined by antibody which reacts with type I and III collagen. Collagen is present around the central (effluent) vein and along sinusoids. (b) Immunohistochemical localisation of collagen in mouse liver three days after treatment with carbon tetrachloride. Collagen deposition is increased in the healing centrilobular zone of liver around a central vein.
lobule where hepatocyte debris was being resorbed (Fig. 1b). Normal and carbon tetrachloride damaged liver treated with normal sheep IgG or second fluorescent antibody did not react with liver collagen.

Factors which stimulated collagen synthesis in L-929 fibroblasts were detected in extracts of mouse liver three days after carbon tetrachloride injury but not in extracts of normal mouse liver. When extracts of damaged liver were chromatographed on G25 Sephadex, four factors were isolated (Fig. 2). These factors were designated F1-F4. F1 stimulated collagen synthesis 5-7 fold, F2 6-3 fold, F3 4-7 fold, and F4 4 fold. Fig. 3 shows that these same fractions stimulated total protein synthesis to the following extent: F1 1.4 fold, F2 2.6 fold, F3 1.9 fold, and F4 2.1 fold. By comparison, chromatograms of normal liver extracts showed only one area (between fractions 30 and 45) which stimulated collagen synthesis 2-3 times (Fig. 2). The latter fractions also stimulated protein synthesis up to a maximum of 2-3 times (Fig. 3) that of controls—in other words, these fractions from normal liver have the same effect on collagen and total protein synthesis. The results shown in Figs. 2 and 3 are representative of over 200 experiments.

Fractions from G25 chromatograms of damaged and normal mouse liver were assayed for their ability to stimulate prolyl hydroxylase activity in mouse fibroblasts. As shown in Fig. 2, F1, F2, F3, and F4 increased prolyl hydroxylase activity in these cells 15-8, 10-5, 4-5, and 7 fold respectively. It should be noted that those fractions which increased hydroxylase activity also stimulated collagen synthesis (see Fig. 2). By contrast, column effluent in the F1 and F2 regions from normal liver stimulated hydroxylase activity 1.4 and 2.5 fold respectively; material eluting in the F3 and F4 region in normal liver chromatograms did not stimulate the activity of this enzyme. The data shown in Fig. 4 are representative of more than 20 experiments in which the maximum stimulation of prolyl hydroxylase activity by F1 from damaged liver was up to 20 fold; the maximum observed with F2 from normal liver being 4 fold.

None of the factors from damaged liver (and the effluent from normal liver columns) had cytotoxic activity for L-929 cells as assessed by trypan blue exclusion. The cytotoxic activity of F1 from damaged liver reported earlier² is abolished by heating liver extracts at 56°C before chromatography.

Normal human and alcoholic cirrhotic liver were examined for the presence of collagen stimulating factors by molecular sieve chromatography of crude
liver extracts. Since there was coincidence of stimulation of collagen synthesis and prolyl hydroxylase in fractions from mouse liver damaged by carbon tetrachloride, the column effluents from human material were assayed for collagen stimulating factors by examining their effect on prolyl hydroxylase activity in human fibroblasts (W138). The enzyme assay is at least five times less time consuming than that for collagen synthesis; this is important in isolating collagen stimulating factors since they are labile. As shown in Fig. 5, human cirrhotic liver contained four collagen stimulating factors; F1, F2, F3, and F4 from human cirrhotic liver stimulated prolyl hydroxylase activity 1.7, 2.6, 2.3, and 2 fold respectively whereas the same fractions from normal liver had no appreciable effect on this enzyme. Homogenates of the cirrhotic liver used in this experiment had a prolyl hydroxylase level four times that of the normal liver used, indicating that there was active fibrosis in the cirrhotic liver.\(^3\)

Prolyl hydroxylase activity in fibroblasts is increased by exposure to ascorbate and lactate.\(^1\) Ascorbate eluted from G25 columns in the F2 region (data not shown). In damaged and normal mouse liver chromatograms, ascorbate was present at a concentration of 20–70 \(\mu\)g/ml and 15–50 \(\mu\)g/ml respectively in the F2 region only. To check whether the increase in prolyl hydroxylase activity by F2 from damaged liver was due to ascorbate, the sensitivity of F2 to cuprous ion was tested; cuprous ion destroys ascorbate.\(^7\) Table 1 shows that when F2 from injured liver was treated with cuprous chloride it retained its activity whereas the ability of ascorbate to stimulate prolyl hydroxylase was inactivated by the copper. Material eluted in the F2 from normal liver, which stimulated prolyl hydroxylase, was also destroyed by cuprous chloride. Table 2 shows that ascorbate had no effect on collagen synthesis in L-929 fibroblasts while collagen stimulating factor 2 from damaged liver stimulated collagen synthesis up to 5 times that of control cultures (Fig. 2); the F2 region from normal liver did not stimulate collagen synthesis. Lactate was not present in the effluent from either normal or damaged liver extracts (data not shown). Neither ascorbate or lactate, therefore, are responsible for prolyl hydroxylase stimulation in collagen stimulating factors from diseased livers.

Recently Ronnema et al.\(^16\) isolated a factor from liver which stimulated collagen synthesis; this was identified as glutamine. The effect of glutamine on collagen synthesis and protein synthesis was therefore tested in L-929 cultures (Fig. 6). Glutamine \(10^{-3}\)M increased collagen synthesis 1.7 times and protein synthesis 1.3 times. Higher concentrations of glutamine resulted in inhibition of both collagen and protein synthesis.

![Graph](http://jcp.bmj.com/)
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The factors had no effect on total cell number or total DNA (data not shown).

Discussion

It has been shown previously that in response to acute carbon tetrachloride poisoning in mice there is an increase in collagen synthesis in the liver. This model of fibrosis has been used to study the control mechanisms of collagen synthesis. As shown in Figs. 1a and 1b collagen fibres are deposited in the liver parenchyma three days after injury in the areas surrounding the central vein and in the areas of tissue repair. Similar observations have been made in rats subjected to prolonged carbon tetrachloride poisoning.

Using this model we have shown that there is a group of low molecular weight factors in these livers which specifically stimulate collagen synthesis when added to cultured fibroblasts. Similar factors have also been found in healing wounds. Moreover, they are also present in the liver of a patient with alcoholic cirrhosis and not in normal human liver. It is therefore proposed that these factors may be the mediators of increased collagen synthesis in hepatic fibrosis in both animals and man.

A prominent feature of hepatic fibrosis is the presence in the parenchyma of mononuclear cell types such as macrophages and lymphocytes. Also notable is the increase in the number of fibroblasts. These observations have prompted workers to look for the existence of lymphokines which increase the number of collagen producing cells. One report showed evidence of a lymphokine that was released from mitogen stimulated peripheral blood lymphocytes and another report showed evidence of a T cell dependent factor that stimulated fibroblast growth. Wyler et al. also presented evidence that a lymphokine could be responsible for the increased fibroblast content in the livers of mice with experimentally induced schistosomiasis. It is therefore likely that the increase in the number of cells which make collagen, which are present in the liver in conditions of fibrosis, contributes to the total amount of collagen which is laid down. Our data, however, indicate that factors are present in cirrhotic or pre-cirrhotic livers which specifically stimulate collagen production without having any effect on cell growth. It is therefore unlikely that the lymphokines reported by other groups are the same as collagen stimulating factors. A preliminary report suggested that a factor similar to collagen stimulating factor in activity was present in lymphocytes of patients with alcoholic liver disease but the influence of this factor(s) on cell growth was not examined.

Another possible source of collagen stimulating factors is the existence of lymphokines which increase the number of fibroblasts.
factors are products derived from macrophages and so called connective tissue activating peptide. Since collagen stimulating factors are not present in normal liver it is unlikely that the latter compounds are similar to collagen stimulating factors.

Others have reported the presence of two factors which can be isolated from the livers of hypercholesterolaemic rats which stimulate the production of collagen in cultured fibroblasts. Glutamine may be responsible for the activity of one of them. Evidence presented here shows that glutamine has only a limited effect on collagen synthesis compared with collagen stimulating factors and it is improbable that glutamine is responsible for any collagen stimulating factor activity.

In summary, the properties of collagen stimulating factors may be listed as follows: (a) they are present in mouse and human liver in conditions of acute and chronic fibrosis; (b) they specifically stimulate collagen synthesis but not non-collagen protein synthesis and they stimulate the secretion of collagen; (c) they are low molecular weight compounds, heat stable, and possibly peptides as judged by their sensitivity to trypsin; (d) they are present in conditions of fibrosis in other tissues and may therefore play a fundamental role in the tissue response to injury.

The work was partially supported by a grant from the Distillers Company, UK. Dr BC Sykes supplied the anticollagen antibody. Mrs Vera Macintosh typed the manuscript.

References


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