Collagen stimulating factors from lung in experimental paraquat poisoning demonstrated in vitro and in vivo

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SUMMARY Collagen stimulating factors, previously described in experimental liver injury and healing skin wounds, have now been found in experimental paraquat lung injury. These factors are active not only in culture but also in vivo. The evidence from this third example of tissue injury points to an important pathological role in chronic inflammation.

Factors which stimulate collagen synthesis and collagen prolyl hydroxylase activity in cultured fibroblasts have been found previously in the healing phase following two types of experimental tissue injury. Extracts of carbon tetrachloride damaged mouse liver and of healing skin wounds were subjected to molecular sieve column chromatography and similar patterns of stimulating activity were found in column fractions from each. This paper describes the presence of collagen stimulating factors in a third type of experimental tissue damage—paraquat lung injury. Evidence is provided that such factors are active not only in culture but also in vivo.

Experimental paraquat poisoning causes the pulmonary alveolar lining epithelium to be selectively destroyed. The subsequent severe inflammatory reaction leads to ingrowth of fibroblasts with intra-alveolar fibrosis by day 5. This histological phase of increased fibroblast activity corresponds to a biochemical phase of increased collagen synthesis, detected as collagen prolyl hydroxylase activity. The activity of this enzyme, which has a critical function in collagen synthesis, was found to reach a maximum also by day 5. Accordingly, lung tissue was taken at day 4 or 5 for detection of collagen stimulating factors.

Material and methods

Mature male Sprague-Dawley rats (average weight 300 g) were given a single toxic dose of paraquat dichloride (Sigma) in isotonic saline, 20 mg/kg body weight, by intraperitoneal injection. Test and control rats were killed at days 4 or 5 and the lungs removed promptly and placed on ice. The tissue was either used at once or stored at −20°C. Three grams of lung parenchyma taken from at least two rats in each experimental group was chopped with scissors and then homogenised at half speed for 1 min using a Silverson homogeniser in 5 volumes of 0.05 M phosphate buffer, pH 7-0, at 4°C. The homogenate was centrifuged at 12 000 g for 20 min at 4°C. The supernatant minus surface lipid was applied to a K16/90 column of G25 fine Sephadex (Pharmacia), bed height 85 cm. Fractions (3 or 4 ml) were collected at a flow rate of 30 ml/h at 4°C, and protein content was continuously monitored at 280 nM. In later experiments the supernatant was applied to Centrifiuo ultrafiltration cones (Amicon) of 50 000 dalton exclusion and the 750 g filtrate obtained.

One millilitre aliquots of column fractions or buffer were added to replicate cultures of L 929 mouse fibroblasts (Gibco-Biocult, Paisley) seeded two days previously at a density of 0.6 to 0.8 million cells per 25 cm² plastic culture flask (Nunc products). Consecutive or alternate fractions were tested depending on numbers of flasks available. The cells were grown in 5 ml of minimum essential medium with the addition of 10% fetal calf serum, 2 × 10⁻²M L-glutamine, 0.25 × 10⁻⁴M ferric nitrate, 0.25 × 10⁻³M sodium-L-ascorbate, 100 units/ml of penicillin, and 100 µg/ml of streptomycin. All media were obtained from Gibco-Biocult and chemicals were obtained from Sigma. After 4 h incubation the cells were assayed for prolyl hydroxylase activity or collagen synthesis as previously described.

The assay for prolyl hydroxylase depends on the
ability of the enzyme to convert prolyl residues in collagen to peptide bound hydroxyproline. A substrate containing L-[3,4-3H] proline labelled collagen is prepared from chick embryos in the presence of α, α1-dipyrrolid, which inhibits hydroxylation. On hydroxylation of the labelled proline by enzyme in subsequent assay material, a hydrogen atom is released and tritiated water collected by vacuum distillation for scintillation counting. The assay for collagen polypeptide synthesis depends on the exposure of material labelled with L-[5-3H] proline to bacterial collagenase free of non-specific proteolytic activity. Labelled small peptides are released and estimated by scintillation counting. Wilcoxon's sum of ranks test was used for statistical analysis.

In later experiments ultrafiltration samples were tested by intraperitoneal injection into CFLP littermate male mice using animals of similar size (20-25 g) in groups of four or five. After 2 h the mice each received a further injection of 0.1 μCi of 5-3H proline (Amersham) in 0.5 ml normal saline in the flank. The mice were killed 4 h later, and the liver and sometimes the lungs as well were removed and stored at −20°C for subsequent prolyl hydroxylase and collagen synthesis assays as previously described. As the amount of material available for assay was greater than from tissue cultures, both assays could be readily combined. The results are expressed as DPM per 50 and 200 μl of homogenate for prolyl hydroxylase and collagen synthesis respectively, being the original data. Student's t test was used for statistical analysis.

Results

With normal lung extract no increase in prolyl hydroxylase activity was seen in test compared with control (buffer only) fibroblast cultures and no consistent pattern was evident (Fig. 1). With paraquat lung extract, the continuous protein profile from the column was subtly different with greater prominence of later peaks (Fig. 2). A fairly consistent pattern of at least five zones of stimulation of activity in cultures was observed, only roughly corresponding to the major protein peaks. Comparison of test and control culture results by means of Wilcoxon’s sum of ranks test confirmed a significant overall increase (p < 0.01). Fig. 3 shows a similar experiment using paraquat lung extract following storage of tissue for one week at −20°C. The continuous protein profile shows even greater prominence of particularly the last peak. The fibroblast cultures exposed to the later fractions show the greater stimulation. Once more, the results with the test cultures were significant compared with control (buffer only) cultures (p < 0.05).

Again, with normal lung extract no increase in collagen synthesis was seen in test compared with control cultures (Fig. 4). Results with this assay, however, were more erratic than for prolyl hydroxylase. With paraquat lung extract, a less consistent pattern of stimulation was obtained, and overall, the results were not significantly different (Fig. 5). Nevertheless, individual cultures achieved high values not seen using normal lung column fractions. Paraquat lung stored frozen appeared to lose much of its activity together with a change in protein profile with further enhancement of later peaks.
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(Fig. 6). Some residual stimulatory activity appeared to persist in those lower molecular weight fractions.

Intraperitoneal injection of normal and paraquat lung extracts did not appear to distress the mice and, when killed 6 h later, there was no evidence of peritoneal inflammation. The injection of the appropriate isotope for the collagen chain synthesis assay did not adversely affect the assay for prolyl hydroxylase on the same tissue. The proline is labeled in such a position in this isotope that it is not available for release by hydroxylation.

The Table shows the effect of injection of 2 ml of either normal or paraquat lung extract on mouse liver in two separate experiments. There was considerable batch variation in the control (buffer only) values attributable to differing maturity of animal groups. Only a slight increase in prolyl hydroxylase activity and collagen synthesis was found in animals treated with normal lung extract. A significant 2-6 fold rise in prolyl hydroxylase activity was evident after addition of paraquat lung extract. A similar

Fig. 3  A further experiment similar to Fig. 2 using paraquat lung after storage at −20°C.

Fig. 4  Effect of column fractions from normal rat lung on collagen chain synthesis in cultured fibroblasts. No significant increase is seen compared with control cultures.

Fig. 5  Effect of column fractions from paraquat damaged rat lung on collagen synthesis in cultured fibroblasts.

Fig. 6  A further experiment similar to Fig. 5 using the same paraquat lung after storage at −20°C.
Hepatic collagen prolyl hydroxylase activity and collagen chain synthesis (mean ± SEM) in mice following intraperitoneal injection of lung extract

<table>
<thead>
<tr>
<th>Substance injected</th>
<th>Prolyl hydroxylase (DPM/50 μl)</th>
<th>Collagen synthesis (DPM/200 μl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Buffer</td>
<td>1167 ± 208</td>
<td>1326 ± 365</td>
</tr>
<tr>
<td>Normal lung extract</td>
<td>1306 ± 194</td>
<td>1460 ± 1245</td>
</tr>
<tr>
<td>Buffer</td>
<td>479 ± 91</td>
<td>381 ± 269</td>
</tr>
<tr>
<td>Paraquat lung extract</td>
<td>1238 ± 111*</td>
<td>814 ± 151</td>
</tr>
</tbody>
</table>

Two experiments were performed with groups of five and four mice.

*p < 0.01 compared with buffer controls (Student's t test).

Fig. 7 Effect of injection of increasing amounts of paraquat lung extract on collagen synthesis in normal mouse liver. Increase in collagen synthesis becomes significant (p < 0.05) at 1/4 and 1/2 dilution with Student's t test.

In previous work permitted statistical assessment and significance was achieved with the more sensitive and reliable assay for prolyl hydroxylase activity. Preparation of extracts after paraquat lung had been stored frozen resulted in the loss of activity from earlier column fractions and the appearance of residual activity in later fractions (Fig. 3, 6) together with enhancement of the later protein peaks, particularly the last. This could well be attributable to the effect of proteolytic activity, and indeed collagen stimulating factors from liver have been shown previously to be trypsin sensitive.1-12 The present findings suggest the possibility that the factors are not just finally degraded by proteolytic activity but are initially produced by it. Such a mechanism for increasing collagen synthesis would be consistent with the presence of large numbers of inflammatory cells in damaged tissue at this time. The presence and activity of macrophages have been shown to be of crucial importance in healing and fibrogenesis in wounds by Leibovich and Ross,13 and by us in carbon tetrachloride liver injury.10 The finding of collagen stimulating factors in this third type and site of tissue injury indicates that they are independent of the mode of injury and suggests that they are of general occurrence during recovery from injury and in all chronic inflammatory diseases.

The use of tissue culture fibroblasts as a model for detecting substances which enhance collagen production may seem appropriate, as collagen is the major synthetic product of the fibroblast.14 The rate of synthesis, however, varies considerably during different phases of culture growth, the rate of effective secretion increasing towards confluence, promoted by increasing prolyl hydroxylase activity.15,16 It is by no means clear which tissue culture conditions are analogous to cells in their natural environment. For example, platelet growth factor has been found to be ineffective in stimulating cell division when cells are cultured in flasks coated with extracellular matrix material.17 The question arises whether collagen stimulating factors, which are as yet uncharacterised, are merely a tissue culture artefact or whether they are truly operative in vivo.

Lung extracts were injected into mice intraperitoneally as this route can accept relatively large volumes and even large molecular weight material—for example, carrageenan18—gains access to the circulation from this site. Membrane pore filtration rather than column chromatography was used since an overall effect as in vivo was sought. Similarly, the filtrates used contained material < 50 000 daltons as compared with the column fractions < 5000 daltons beyond the first peak. This was because initial fibroblast culture work with carbon tetrachloride injured liver filtrates had shown the rise was shown for collagen synthesis, but the result was not significant. Significant increases with the collagen synthesis assay were shown by the dose-response experiment (Fig. 7) with paraquat lung extract, the value reaching a maximum of just over two fold.

Discussion

The pattern of five major zones of stimulating activity obtained from paraquat lung extracts resembles that previously seen with healing liver and skin material.1,12 The inclusion of more control cultures than

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greatest activity to reside in this filtrate and progressively less in filtrates of < 10 000 and < 5000 daltons (unpublished data). The term “collagen stimulating factors” has been applied previously to material < 5000 daltons but since they are uncharacterised and the term is non-specific, it seems fair to apply it to the larger molecular weight fractions also.

From the Table it is apparent that the filtrates are capable of stimulating collagen synthesis in vivo. The dose-response experiment (Fig. 7) shows that, given sufficient animals at each point, such increases can even be quantitated. The maximum of just over two fold stimulation reached as compared with up to five fold in cultures probably indicates just how non-optimal are tissue culture conditions. The fact that stimulation has been obtained in healthy mouse liver and also twice in lung (data not shown) using a rat lung extract emphasises that collagen stimulating factors are not organ or species specific. It is notable that in the paraquat treated rat itself, the lung being selectively damaged exhibits increased collagen synthesis in contrast to liver or kidney. The situation in the mouse is presumably one of unnatural systemic access of stimulating factors.

The evidence presented here is limited, but collagen stimulating factors appear to have the ability to override normal control mechanisms in vivo and this surely indicates a pathological importance.

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References


Requests for reprints to: Dr WD Thompson, Department of Pathology, University Medical Buildings, Foresterhill, Aberdeen AB9 2ZD, Scotland.
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