Juvenile hyaline fibromatosis: impaired collagen metabolism in human skin fibroblasts

F Breier, S Fang-Kircher, K Wolff, W Jurecka

Abstract

Juvenile hyaline fibromatosis (JHF) is inherited as a fatal autosomal recessive disorder characterised by multiple tumorous mucocutaneous proliferations. In this paper a 14 month old girl with JHF is described. For this condition, a malfunction of collagen synthesis is considered as the pathogenetic cause. Recently published data have revealed an absent band for type III collagen (TIIC) chain in western blot studies of clinically unaffected JHF skin. Therefore supernatants of skin fibroblast cell cultures, obtained from normal human skin, were analysed for type I collagen (TIC) and TIIC metabolites by radioimmunoassays. Besides the typical morphological connective tissue changes in the skin lesions, TIC synthesis and degradation were found increased in JHF fibroblasts compared with control fibroblasts. In contrast, TIIC overall metabolism was significantly reduced by 36% compared with controls. (Arch Dis Child 1997;77:436–440)

Keywords: juvenile hyaline fibromatosis; skin; collagen type I and type III metabolism

Juvenile hyaline fibromatosis (JHF) is a rare disease characterised by tumorous skin lesions with the onset in early infancy, joint contractures, thickening of the gums, bone lesions, and tumorous involvement of internal organs. The skin lesions may vary in number and size and represent painful fleshy cutaneous papules, nodules, or tumours mainly located at the neck, elbows, knees, shins, and ankles. The condition is believed to be inherited as an autosomal recessive trait. Consanguinity of the parents has been reported in some of the sporadic cases. Recently, Kayashima et al have suggested JHF to be a connective tissue disorder characterised by increased synthesis of glycosaminoglycans (GAG) by fibroblasts and an impaired type VI collagen metabolism. In contrast, we demonstrated recently an absent type III collagen (TIIC) chain in western blot studies of normal skin in this patient with JHF. Therefore we wanted to quantify the synthesis and degradation of type I collagen (TIC) and TIIC of human skin fibroblasts in JHF.

As all of the affected sites predominantly contain TIC and TIIC a deregulated metabolism of these collagens might be suspected. In the following paper we present findings that indicate an abnormality of these collagens in a JHF patient.

Case report

The family history on the mother’s side is remarkable: her cousin is suffering from trisomy 21, her nephew shows hyperplasia of the gingiva. In some members of the mother’s family contractures of the left hand’s fourth finger were observed as individual symptoms (see family tree, fig 1). The parents themselves show no signs of any inherited disease. Both sisters of the patient are healthy. JHF was diagnosed by tumorous plaques on the lateral and medial aspects of the elbows, wrists, knees, and ankles (fig 2), gingival hyperplasia, and papillomatous anal proliferation, and by typical dermatohistopathological and ultrastructural findings.

The patient died at the age of 14 months due to the progression of her disease complicated by pneumonia and respiratory insufficiency. A postmortem examination revealed congestion of the lung and bronchopneumonia, showing a dense peribronchial infiltrate, mostly composed of neutrophils and foamy macrophages. The visceral pleura revealed fibrous thickening.

Experimental

LIGHT AND ELECTRON MICROSCOPY

Biopsy specimens were taken from tumorous infiltrations of the left ankle under local anaesthesia. Two tissue samples were prepared for light and electron microscopy. For routine histopathology specimens were paraffin embedded and stained with haematoxylin and eosin, periodic acid Schiff (PAS) reagent, and elastica stain. For ultrastructural examination specimens were fixed in 2.5% glutaraldehyde in 0.1M phosphate buffer, pH 7.2, postfixed in osmium tetroxide, and embedded in Epon 812. Semithin sections were stained with toluidine blue. Ultrathin sections were stained with lead citrate and uranyl acetate and were analysed in a Jel 1010 transmission electron microscope.

CELL CULTURE CONDITIONS

Three further skin biopsy specimens were obtained from skin that appeared clinically normal from the trunk of the patient. Three fibroblast cell lines were grown from these specimens in minimal essential medium supplemented with 10% fetal calf serum, 1% non-essential amino acids, 1% kanamycin, and 1% glutamine (all Gibco). After five days in culture, the supernatants were analysed in order to determine collagen and extracellular matrix parameters.
For control purposes three fibroblast cell cultures from clinically normal skin taken from the trunk of three girls (age 12–15 months) undergoing orthopaedic surgery, with no genetically determined disorders, were treated in the same way. Small skin biopsy specimens were taken by informed consent of the patients’ parents. Additionally flasks with cell free medium were cultivated. All values of collagen and extracellular matrix parameters in the supernatants of the cell cultures were reduced by the concentrations found in cell free medium to exclude possible influence of fetal calf serum.

ANALYSIS OF EXTRACELLULAR MATRIX PARAMETERS AND TIC AND TIIIC METABOLITES

The following parameters were analysed in the supernatants of fibroblast cell cultures: GAG by a spectrophotometric method using dimethylmethylen blue. Total protein was measured by a method described by Lowry et al to exclude methodical interferences in determination of GAG. Hyaluronic acid was determined by radioimmunoassay (Kabi, Pharmacia, Sweden). In order to investigate collagen metabolites, carboxy terminal propeptide of collagen type I (PICP), carboxy terminal cross linked telopeptide of collagen type I (ICTP), and amino terminal propeptide of collagen type III (PIIINP) were analysed by radioimmunoassay (Orion Diagnostica, USA). PICP and ICTP can be regarded as parameters of synthesis and degradation respectively of collagen type I; PIIINP has been reported to reflect the TIIIC overall metabolism.

STATISTICAL ANALYSIS

All concentrations were correlated with the standardised numbers of cells in each culture flask and were calculated as mean (SD) µg/flask. All parameters of the cell cultures were investigated also in cell free cultivated medium. All the statistical differences were analysed by Student’s t test.

Results

LIGHT MICROSCOPY AND ELECTRON MICROSCOPY OF INVOLVED SKIN (TUMOROUS PLAQUE)

The epidermis showed mild hyperplasia but otherwise regular structures. In the dermis the extracellular component of the connective tissue was markedly increased with irregularly arranged thick collagen bundles and homogeneous eosinophilic matrix (fig 3). There was also increased cellularity, most cells representing fibroblast-like cells with bizarre shape. The skin appendages were reduced in number. The number of small vessels was increased and they were surrounded by concentrically arranged collagen bundles. Similar changes were visible in the mid-dermis and around an epithelial cyst. Elastica stain showed rarefaction of elastic fibres and PAS stain revealed fine granular PAS positive material between the collagen fibres in the homogeneous matrix around the vessels of the dermis. These findings were typical for those already described for tumorous proliferations in JHF.

In semithin sections (fig 4) and electron microscopy (fig 5) the tumorous infiltrations revealed relatively large, bizarre shaped connective tissue cells with pseudopodia-like protrusions. Nuclei were bizarre shaped with numerous invaginations, surrounded by fine granular, electron dense cytoplasm. Most of the cells contained a prominent Golgi apparatus and numerous membrane bound vesicles of varying size up to 1 µm in diameter. The vesicles were filled with a fine granular and
microfibrillar material. Similar microfibrillar material was also observed in varying amounts extracellularly between these cells. Additionally some normal mature collagen fibres and long spacing collagen fibres were seen.

**CELL CULTURES OF SKIN FIBROBLASTS**
*(CLINICALLY NORMAL SKIN OF JHF PATIENT AND CONTROLS)*

All the cell lines of fibroblasts that were obtained from the patient suffering from JHF showed different growth patterns from those of normal controls. In the confluence stage the control cell lines showed in primary and in subsequent cultures a homogeneous growth pattern with regularly shaped fibroblasts mainly arranged in parallel. However in the confluence stage of the cell cultures of the JHF patient a milky, mucoid pseudomembrane could be observed. Microscopically the cells were irregularly arranged, unevenly orientated, presenting a bizarre growth pattern. Additionally these fibroblasts were uneven in size and shape.

**CONCENTRATIONS OF GAG AND HYALURONIC ACID**

The supernatants of standardised cell cultures on day 5 showed no significant difference in GAG (69.3 (3.8) v 73.8 (9.4) µg/flask) and hyaluronic acid (137.7 (13.6) v 133.5 (22.0) µg/flask) in the child with JHF compared with controls (fig 6A).

**PINC AND ICTP IN FIBROBLAST CELL CULTURES IN JHF**

The concentrations of PINP (24.1 (8.4) v 12.6 (1.0) µg/flask, p<0.13) and ICTP (0.5 (0.1) v 0.3 (0.1) µg/flask; p<0.14) in the supernatants of the standardised patient’s cultures were increased by 91% and 62% respectively, compared with the supernatants of standardised cell cultures of the healthy individuals (fig 6A). However the ratio of both markers (PINC/ICTP = 45.4 (7.2) v 43.1 (12.3)), reflecting synthesis and degradation of TIC, was equal in both, the pathological and the control samples respectively, indicating an accelerated but yet balanced TIC metabolism (fig 6B).

**PIIINP IN FIBROBLAST CELL CULTURES IN JHF**

PIIINP was highly significantly reduced by 36% in supernatants of standardised JHF fibroblast cultures compared with those of the control cells (1.05 (0.04) v 1.65 (0.01) µg/flask; p<0.001) demonstrating a decreased TIIIC metabolism in JHF (fig 6C). In order to show the differences between the concentrations of TIC and TIIIC metabolites, the ratios of PIIINP/PINC (0.05 (0.02) patient v 0.13 (0.01) controls; p<0.05) and PIIINP/ICTP (2.02 (0.43) patient v 5.16 (1.69) controls; p<0.05) were calculated (fig 6C); both were significant.

**Discussion**

In this study we demonstrated a significantly decreased TIIIC and increased TIC synthesis and degradation in fibroblast cell cultures of normal human skin of a patient with JHF. However in JHF several pathogenetic mechanisms have been discussed (table 1). Woyke et al speculated on an impaired...
Table 1  Metabolic defects reported in JHF

<table>
<thead>
<tr>
<th>Reference</th>
<th>Metabolic defect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Woyke et al.</td>
<td>Procollagen, tropocollagen</td>
</tr>
<tr>
<td>Iwatha et al.</td>
<td>GAG, hyaluronic acid</td>
</tr>
<tr>
<td>Kitano et al.</td>
<td>Hyaluronic acid, chondroitin sulphate</td>
</tr>
<tr>
<td>Kayashima et al.</td>
<td>Type VI collagen</td>
</tr>
<tr>
<td>Luberc et al.</td>
<td>Type III and type I collagen</td>
</tr>
<tr>
<td>Glover et al.</td>
<td>Type VI collagen</td>
</tr>
<tr>
<td>Breier et al. (present study)</td>
<td>Type III and type I collagen</td>
</tr>
</tbody>
</table>

Impaired collagen type VI metabolism. However, these findings in our view would not sufficiently explain the clinical spectrum of involvement in JHF, since the skin, the mucous membranes, the joints, and the bones are the major manifestations sites of this condition mainly harbour TIC and TIIIC. Furthermore, recently we demonstrated a deregulation of TIIIC in western blot studies of clinically normal skin in JHF. The absence of the TIIIC chain accounts for a decreased TIIIC metabolism. TIIIC defects were also described in Ehlers-Danlos syndrome type IV. A TIII(α3)C defect was reported in vascu-lopathy in Ehlers-Danlos syndrome leading to a great risk of spontaneous haemorrhage from vascular rupture or bowel perforation. Recent molecular advances have identified several mutations in the gene for type III procollagen to be responsible for Ehlers-Danlos syndrome type IV. One gene has been mapped to human chromosome 2q24.3-q31 by in situ hybridisation studies. We assume that a pathological TIIIC metabolism may be induced by a similar underlying genetic defect in JHF.

Evaluation of TIC metabolites in our patient recorded a markedly increased level of PICP and ICTP but the PICP/ICTP ratio was equal in pathological and control cell lines respectively. This indicates an accelerated, but balanced collagen TIC metabolism. In contrast, concentrations of PIINP were significantly increased in supernatants of standard cell lines and/or release into culture medium. This indicates a decreased TIIIC metabolism in JHF patients and it can be speculated that the reduced enzymatic removal of the TIIIC terminal propeptides, which is necessary for regular cross linking of collagen fibres, causes a diminished stability of collagen molecules in JHF. Reduced TIIIC overall metabolism may also lead to the intracellular accumulation of mature and immature collagen macromolecules, aggregated in intracellular vacuoles and the extracellular matrix, which was demonstrated by Woyke et al. and by us.

Histopathological investigations of clinically involved skin showed increased cellularity of the dermal connective tissue cells extending from the papillary dermis to the subcutaneous fat and deposits of increased amounts of cell fibres. Ultrastructural analysis revealed, according to the findings of Woyke et al., numerous fibroblast-like cells containing a highly active Golgi apparatus. It is generally believed that vesicles of the Golgi apparatus of a fibroblast contain a solution of monomeric tropocollagen macromolecules which are precursors of synthesis of procollagen or tropocollagen, on the other hand Iwatha et al. discussed an increased or faulty synthesis of GAG by fibroblasts leading to this condition.

Further reports suspected increased chondroitin sulphate synthesis or decreased amounts of hyaluronic acid as causative agents. However, in contrast to the observations of Iwatha et al. in our patient biochemical investigations of fibroblast cultures supernatants showed no significant increases of GAG and hyaluronic acid. Our investigation was performed on cultures from morphologically unaltered skin.

More recent investigations by Kayashima et al. supposed JHF as a disease associated with an underlying genetic defect in JHF.
of stroma collagen. These macromolecules have to be secreted from the Golgi apparatus to the extracellular space where soluble tropocollagen units are quickly transformed into fibrils. Excess of such a floccular and fibrillar material can be detected between the numerous cells of the skin tumours, indicating that the basic defect is a localised metabolic disturbance in the formation of collagen. In adult human dermis TIC accounts for approximately 80% of total collagen. The synthesis of TIC accelerates during the neonatal period until the ratio of TIC to TIIC in adult human skin is 6:1. Western blot analysis of our patient showed an absent TIIC chain exclusively in the skin, but not in other organs, such as heart, liver, and kidney. This may be explained by a decreased synthesis of TIIC in skin fibroblasts.

It can be assumed that the increased TIC turnover and the excessive production of the extracellular matrix are secondary effects due to the disturbed TIIC metabolism which obviously is fundamentally involved in the pathogenesis of this disorder. The suggestion, that the presence of type III procollagen, in particular, at the surface of collagen fibrils has an important role for TIC, underlines this hypothesis. This may be an explanation for the clinical presentation of tumorous infiltrates with predominance above mechanically strained sites.

In conclusion these results reflect a decreased TIIC metabolism with a possibly secondarily increased TIC metabolism and may be regarded as the reason for clinical manifestations of JHF.