Matrix remodelling in dilated cardiomyopathy entails the occurrence of oncofetal fibronectin molecular variants

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Abstract

Objectives—To investigate whether disturbance of the cellular homeostasis and integrity of cardiomyocytes in dilated cardiomyopathy (DCM) is accompanied by alterations in cell-matrix relations as indicated by changes in the deposition of fibronectin (FN) isoforms.

Design—Tissue from a case series of patients with DCM was investigated by immunohistochemistry with antibodies against FN (all variants, clone IST4), ED-A+ FN (clone IST9), ED-B+ FN (clone BC1), and oncofetal glycosylated FN (clone 5C10). The sites of de novo synthesis of FN were demonstrated by means of non-radioactive RNA in situ hybridisation (ISH) with biotinylated FN cDNA fragments as the probe.

Setting—University hospital.

Patients—Samples from 10 patients with clinical criteria and histological diagnosis of DCM and from 3 individuals with normal hearts.

Interventions—Samples were obtained by right ventricular endomyocardial biopsy.

Main outcome measure—Distribution of oncofetal FN variants in DCM hearts.

Results—Immunostaining of FN (IST4, all variants) showed a coarse interstitial network in normal and diseased myocardium. ED-A+ FN was deposited in fine interstitial spots in normal myocardium and in DCM samples. Immunostaining for oncofetal glycosylated FN and ED-B+ FN was not seen in normal adult myocardium, whereas myocardium from DCM patients showed focal and delicate staining in the interstitium. RNA ISH showed that these deposits resulted from local FN synthesis.

Conclusion—The results accord with de novo expression of oncofetal FN variants in hearts with patients with DCM. The oncofetal FN variants may serve as disease markers in myocardium affected by DCM.

(Heart 1996;75:358–362)

Keywords: dilated cardiomyopathy; extracellular matrix; oncofetal fibronectin; in situ hybridisation

Dilated cardiomyopathy (DCM) is a heart muscle disease characterised by a decrease of the left ventricular ejection fraction and dilatation of both ventricles.1–3 The main histological changes are a disturbed tissue integrity and interstitial fibrosis, and both features affect outcome.4

Cardiomyocytes need cell-cell contacts and defined cell-matrix relations to maintain normal physiological cell function.5,6 Therefore, all alterations to cardiomyocytes, including cellular degeneration, can change the composition of the extracellular matrix and vice versa.7 In normal myocardium, fibronectin (FN), laminin, and different collagens are present in the intercellular spaces between myocytes and in the vicinity of blood vessels8 and are significantly increased in diseased human myocardium.9 The increase in FN is particularly prominent, and eventually thick sheaths form around the myocardial cells.10 Though the role of FN in the organisation of connective tissue in vitro and in vivo has been studied,11–14 nothing is known about occurrence and molecular variants of FN in human myocardium. Molecular variants of FN develop by alternative splicing or glycosylation. The molecular components, extra domains A and B (ED-A+, ED-B+), are skipped or included in a cell type specific manner. ED-A+ and ED-B+ are included only in cellular FN,19 in adult tissues the distribution of ED-A+ and ED-B+ FN expression is highly restricted.16 In addition the FN molecule may be subject to de novo glycosylation depending on the developmental stage of the tissue.17,18 These so-called oncofetal FN are known to occur in immature, tumorous, and reparative tissues.19–21 Are they also generated in hearts with DCM and if they are, how are they distributed?

Patients and methods

Tissue material

Endomyocard biopsy specimens from 10 patients with the clinical criteria and a histological diagnosis of DCM and three samples of normal human heart were examined. The diagnosis was confirmed by conventional histology. Fresh biopsy specimens, up to 3 mm in diameter, were snap frozen in fluid propane cooled by liquid nitrogen.

Immunohistochemistry

Cryostat sections of the frozen tissue samples were fixed in ice cooled acetone for 15 minutes and examined immunohistochemically
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Figure 1  (A) DCM myocardium showing an increase in interstitial connective tissue, diffuse pericardio-myocytic fibrosis, and small areas of fibrosis (trichrome Gomori staining (original magnification, × 200)).
(B) Coarse diffuse interstitial ED-A+ FN deposition in myocardium affected by DCM (monoclonal antibody IST9, APAAP technique (original magnification, × 200)).
(C) De novo occurrence of oncofetal glycosylated FN, demonstrated as sparse focal deposition in the interstitium of DCM affected myocardium (antibody SC10, APAAP technique (original magnification, × 200)).
(D) De novo expression of oncofetal ED-B+ FN, demonstrated as sparse focal deposits in the interstitium of DCM affected myocardium (antibody BCI, APAAP technique (original magnification, × 200)).
(E) Non-radioactive FN in situ hybridisation using cDNA as a probe. Positive signals in single stromal cells with a distribution pattern similar to the oncofetal FN variants (original magnification, × 200).
with primary antibodies to all variants of FN, including cellular and plasma FN (clone IST4, culture supernatant, diluted 1:20), ED-A FN (clone IST9, culture supernatant, diluted 1:500),22 ED-B' FN (clone BC1, culture supernatant, diluted 1:20),23 oncosetal de novo glycosylated FN (clone 5C10, culture supernatant, diluted 1:20).24 The antibody reaction was detected by the alkaline phosphatase anti-alkaline phosphatase (APAAP) method.

Sections were incubated for 30 minutes at room temperature with primary antibody. Then they were washed with Tris buffer and treated with rabbit anti-mouse immunoglobulin (IgG, Z-259, diluted 1:70, Dako, Glostrup, Denmark), and then with the mouse APAAP-complex (Dako). Both incubations lasted for 30 minutes at room temperature. When primary polyclonal rabbit antibodies were used a second mouse anti-rabbit antibody (diluted 1:400, Dako) was used.

To increase the staining intensity, the incubation with the rabbit anti-mouse immunoglobulin and with the APAAP-complex was repeated twice. Naphthol-AS-biphosphate (Sigma, 2250, St Louis, USA) and new fuchsin (Merck, 4040, Darmstadt, Germany) were used as substrate and developer, respectively. To inhibit endogenous tissue enzyme activity the developing solution was supplemented with 0.25 mmol/l levamisole (Sigma, L-9756). For evaluation of immunostaining, the primary antibody was replaced by nonimmune serum as a negative control.

**MRNA IN SITU HYBRIDISATION**

Non-radioactive fibronectin in situ hybridisation was performed on cryostat sections of specimens snap frozen immediately after human heart biopsy. Tissue sections (10 μm) were mounted on slides coated with 3% (v/v) 3-aminopropyltriethoxysilane in acetone, briefly air dried, and fixed in 4% (w/v) paraformaldehyde in phosphate buffered saline for 20 minutes at 4°C. Then slides were rinsed quickly in 70% ethanol and air dried. Sections were stored at −70°C until they were processed. Before hybridisation the slide preparations were handled as follows: 20 min at room temperature in 0.2 M HCI, rinsed in 2 × SSC, and 15 min at 37°C in 50 mM Tris-HCl, pH 7.6/proteinase K (1–5 μg/ml). Slides were rinsed twice in distilled water, dehydrated in graded ethanol (70% and 96%), and air dried.

**PREPARATION OF LABELLED cDNA PROBE**

The cDNA for human FN23-28 was purchased from Gibco BRL, UK (1.4 Kb, nucleic acids: 1993–3364) and digoxigenin labelled using the Nick Translation Kit and digoxigenin-11-dUTP from Boehringer Mannheim, Germany.

**Prehybridisation and hybridisation**—The hybridisation solution contained 4 × SSC, 50% (v/v) deionised formamide, 1 × Denhardt's solution, 1 mM EDTA, 5% (w/v) dextran sulphate, 100 μg/ml heat denatured calf thymus DNA, 100 μg/ml heat denatured transfer RNA, and 200 ng/ml digoxigenin labelled FN cDNA. Prehybridisation was carried out with hybridisation medium without cDNA for 60 min at 37°C in a moist chamber. Then 50 μl hybridisation solution containing 200 ng/ml probe were applied to each slide, RNase free coverslips were mounted and hybridisation was allowed to proceed at 37°C overnight. Slides were then washed for 2 h in 10 mM Tris-HCl, pH 7.6/2 × SSC/50% formamide/l mM EDTA followed by 1 h at 55°C in Omnisuff (JenaBioTech and WAK Chemie Medical, Bad Homburg, Germany) and rinsed in Omnisuff at room temperature. Hybridised cDNA was detected using an anti-digoxigenin antibody (Boehringer Mannheim, Germany) and the APAAP technique mentioned above.

No probe hybridisation and RNase pre-digestion were used as negative control and specificity controls, respectively.

**Results**

In normal adult human hearts immunostaining of FN (IST4, all variants) showed a delicate interstitial network. ED-A' FN (antibody IST9) deposited as fine interstitial spots. Oncosetal glycosylated FN (antibody 5C10) and ED-B' FN (antibody BC1) could not be seen.

Conventional histology of biopsies of hearts with DCM showed hypertrophied cardiomyocytes and varying degrees of interstitial fibrosis (fig 1A). A few lymphocytes and histiocytes could sometimes be seen in the interstitium. The demonstration of the total FN using the antibody IST4 showed increased immunostaining of a coarse interstitial network. Immunohistochemistry showed slight ED-A' FN deposits in hearts with DCM resembling those in the normal adult hearts (fig 1B). In contrast to the normal adult myocardium there was a focal interstitial immunoreaction indicating de novo glycosylated FN (fig 1C) and ED-B' FN (fig 1D) in failing DCM hearts. These FNs were irregularly distributed in the interstitium and were visible as small ribbons. Non-radioactive in situ hybridisation showed FN mRNA in the fibrotic interstitial areas of myocardium in DCM patients with stronger focal staining in single cells (fig 1E).

In situ hybridisation also showed an even distribution pattern of immunohistochemically demonstrated ED-B' FN and de novo glycosylated FN as well as FN synthesis.

**Discussion**

Maintenance of the structural and functional integrity of heart muscle requires a defined extracellular matrix composition.27-29 Numerous studies have described changes in extracellular matrix composition in the course of developmental and disease processes of the heart.30-32 FN, one of the major non-collagenous proteins of the extracellular matrix, is increased in diseased human myocardium33 and in the FN (all molecular variants) in DCM myocardium that we found accords with this earlier study.

The different FN isoforms are derived from a single gene by alternative processing of the
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Figure 2 Model of the domain structure of a subunit of human FN. The oncofetal domain ED-B and the oncofetal epitope recognised by the monoclonal antibody 5C10 are indicated. The figure also indicates the internal homologies, the three sites of alternative splicing (ED-B', ED-A', and IIICS), the major macromolecules interacting with the various FN domains, and the two isoforms generated by alternative splicing of the ED-B sequence with the reactivity of these isoforms with the monoclonal antibody BC1.

Figure 1b shows a model of the domain structure of a subunit of human FN. The figure also indicates the internal homologies, the three sites of alternative splicing (ED-B', ED-A', and IIICS), the major macromolecules interacting with the various FN domains, and the two isoforms generated by alternative splicing of the ED-B sequence with the reactivity of these isoforms with the monoclonal antibody BC1.

primary RNA transcript (alternative splicing) or by post-translational modifications in the C-terminal region of the molecule with an O-linked glycosylation. FN has three sites of alternative splicing, the ED-A domain, the ED-B domain, and the IIICS domain. The inclusion of the alternative type III repeats, ED-A and the ED-B, in the FN molecule as well as de novo glycosylation can be detected by specific monoclonal antibodies: IST9, BC1, and 5C10, respectively. Figure 2 shows a model of the domain structure of a subunit of human FN and the epitopes recognised by the applied antibodies. Little or no variant FN is present in normal adult tissue. In this study very little ED-A' FN and no ED-B' FN and de novo glycosylated FN was detected in normal heart muscle. ED-A' and ED-B' FN mRNA is increased in embryonic and malignant tissues. Furthermore, the de novo glycosylation of FN is associated with cellular immaturity, cancer formation, and the malignancy of breast, gastric, and oral carcinomas. Such FN variants are termed "oncofetal". Recently, Fahradian and coworkers pointed to the importance of differential splicing of FN during cardiac ontogeny and development of hypertrophy in the rat. As far as we know, ours is the first description of de novo expression of ED-B' and de novo glycosylated FN isoforms in human hearts with DCM. These FN isoforms indicate an altered extracellular matrix and may be important to the progress of myocardial damage. Nevertheless, our results do not establish that oncofetal fibronectin deposition is specific to DCM. It is possible that de novo synthesis of fibronectin isoforms is associated with the attendant fibrosis itself rather than with dilatation. Experimental studies on fetal and neonatal rat cardiomyocytes showed a high affinity to FN in contrast to adult myocytes. This corresponds with a transient physiological up regulation of collagen synthesis in utero and immediately post partum. Collagen synthesis is also increased in DCM (interstitial myocardial fibrosis).

The irregular multifocal distribution of oncofetal fibronectin variants on the one hand shows that the whole heart is involved in the disease process of DCM and on the other hand shows multiple active disease foci in the myocardium.

Though the functional role of the oncofetal FNs is not well understood our results suggest that they may modulate cellular adhesion mechanisms in cardiomyopathy. The occurrence of oncofetal molecular FN variants is likely to be associated with a decreased cellular adhesion. The simultaneous de novo expression of tenasin in DCM, which is known as adhesion modulating factor, accords with the concept that a change of cell adhesion in DCM is a significant aspect of the diagnosis.

The results of FN mRNA in situ hybridisation showed that oncofetal isoforms of FN in DCM are generated by stromal cells with a fibroelastic appearance. A more precise characterisation of the cell type responsible for synthesis of oncofetal fibronectins will be the aim of further investigations. We do not know the mechanisms that induce the synthesis of oncofetal FNs in DCM. Growth factors (released from damaged cardiomyocytes), which are known to modulate FN splicing, may be involved.

2 Kaspeh EK, Agema WRP, Hutchins GM, Deckers JW,
The tions that unmask differential splicing may generate at least 10 polypeptides from a single gene. EMBOL 1985;4:1755-9.


