The alternatively spliced domains EIIIB and EIIIA of human fibronectin affect cell adhesion and spreading

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SUMMARY

Fibronectin has a complex pattern of alternative splicing at the pre-mRNA level leading to the expression of different isoforms. The alternatively spliced domains EIIIB and EIIIA are known to be prominently expressed during development and wound healing. While the other spliced domain (CS-segment) is known to promote cell adhesion in a cell type specific manner, the biological functions of the spliced domains EIIIB and EIIIA are not well understood. In the present study, we have prepared expression proteins of specific domains of human fibronectin using a prokaryotic expression system and used the purified fragments to test their ability to support adhesion and spreading of cultured cells. Fragments from type-III domains #7 to #12 were prepared in various combinations to include or exclude the spliced domains EIIIB and EIIIA. The results indicate that cultured NIL fibroblasts adhere to many of the fragments tested. However, the cell adhesion and spreading are enhanced, especially at lower concentrations, to fragments including the domain EIIIB. The inclusion of domain EIIIA led to a decrease in the adhesion of cells and those that adhered did not spread well. When tested in a centrifugal cell adhesion assay, fragments including domain EIIIB resisted the detaching forces and stayed adhered. Fragments that included domain EIIIA were unable to resist the detaching centrifugal forces to the same extent as the fragments that included domain EIIIB alone. These results suggest that the spliced domain EIIIB may be serving important biological functions in enhancing cell adhesion and spreading. This is likely to be mediated by conformational effects because domain EIIIA alone neither exhibited any adhesive activity nor competed in inhibiting adhesion to fragments #7-10.

Key words: Fibronectin, Domain EIIIB, Domain EIIIA, Cell adhesion, Extracellular matrix

INTRODUCTION

The biological role of the extracellular matrix glycoprotein fibronectin (FN) in cell adhesion and migration, development, and wound healing has been extensively studied (Hynes, 1990). Fibronectins (FNs) are a group of related glycoproteins that arise from alternative splicing of the single FN gene transcript. FN has a relatively simple structure and consist of a dimer of two subunits joined by disulfide bonds near the COOH termini (Hynes, 1990). Three types of homologous repeating units (referred to as types I, II and III domains/repeats) form each subunit. Alternative splicing of a single primary transcript leads to the inclusion or exclusion of three domains in the central part of the molecule. Among the various functions of FN, cell adhesion is by far the most extensively studied. The short polypeptide sequence Arg-Gly-Asp (RGD), located in the 10th type-III repeating unit of FN, has been identified as the important binding site for several integrin cell surface receptors. The other cell adhesion site identified within FN is referred to as the III CS (connecting segment) and this domain is also subject to alternative splicing at the pre-mRNA level (Humphries et al., 1987, 1988). The biological function(s) of the two remaining alternatively spliced domains EIIIB and EIIIA are/is not known. It has, however, been well documented that the domains EIIIB and EIIIA of FN are prominently expressed during embryonic development and this embryonic pattern of mRNA expression reappears in healing wounds (ffrench-Constant and Hynes, 1989; ffrench-Constant et al., 1989). The expression of FN mRNA, including domains EIIIB and EIIIA, in the early chick embryo coincides at stages where there is active proliferation and migration of cells (ffrench-Constant and Hynes, 1989). Findings from a more recent study suggest that the alternatively spliced domains EIIIB and EIIIA are included in FN required for morphogenesis of ‘FN dependent’ structures during development (Peters and Hynes, 1996). Using immunohistochemistry, it has also been shown recently that the spliced domain EIIIB is included in FN around the walls of some smaller blood vessels and within cartilaginous structures (Peters et al., 1996). The possibility exists that these spliced domains of FN (EIIIB and EIIIA) may have important biological functions in the regulation of critical cellular events during embryonic development and wound healing.

Many of the biological functions of FNs have been mapped
using proteolytic fragments (reviewed by Hynes, 1990). These types of studies have several disadvantages including overlapping functional sites and the lack of information on the isoform in question. Potentially, there may be as many as 20 different isoforms of FNs possible due to alternative splicing of the primary transcript (Hynes, 1990). Hence, studies aimed at mapping the biological functions of the spliced domains EIIIB and EIIIA should focus on methods that will provide clean fragments that begin and end precisely at individual domains. In the present study, we have generated several types of recombinant polypeptide fragments of human FN by amplifying cDNA coding for the type-III domains #7-12 and cloning them into prokaryotic expression vectors. By including or excluding the spliced domains EIIIB and EIIIA, we were able to obtain ten different fragments in various combinations of the domains. These expressed domains of human FN were then purified and used to study cell adhesion. In this study, we demonstrate that the inclusion of spliced domain EIIIB alone or in combination with EIIIA, within the larger fragment of FNIII #7-12, has an enhancing effect on the adhesion and spreading of NIL fibroblasts. However, the spliced domain EIIIA alone, within the fragment FNIII #7-12, led to a reduction in the adhesion and spreading of cells. Since the domain EIIIB did not reveal any cell adhesion promoting activity when tested as a single-domain fragment, it is concluded that the domain EIIIB may be influencing cell adhesion by exerting conformational effects on the molecule.

MATERIALS AND METHODS

Materials

Human plasma FN (pFN) was provided by Dr Harold Erickson, Duke University, Durham NC. The monoclonal antibody 16G3, which has its epitope in the 10th type-III domain of human FN, was provided by Drs K. Yamada and S. Aota, NIDR, NIH. The mAB 7F9 to vinculin was provided by Dr A. Belkin. The prokaryotic expression vectors pET 11b and pET 15b were from Novagen, Inc, Madison, Wis., and the pGEM vector was from Promega Corp. Oligonucleotide primers were synthesized at the Pathology Department Oligonucleotide Laboratory at the University of North Carolina Chapel Hill using an ABI-Model 394 Machine. [35S]dATP was from Amersham Corp, Isopropyl β-D-thiogalactosidase (IPTG) was from Gibco, and all other enzymes/reagents were from New England Biolabs or Promega.

Cloning specific domains of human FN into pET expression vectors

The methods used to produce and purify bacterial expression proteins were essentially the same as described previously with some modifications (Aukhil et al., 1993). The prokaryotic expression vector pET 11b (Studier et al., 1990) and its modified version pET 15b with a histidine tag to facilitate affinity chromatography were used. The strategy we have used enables us to clone fragments that begin and end precisely at the boundaries of individual domains of FN. Briefly, oligonucleotide primers with the appropriate restriction sites were designed to amplify fragments of human FN cDNA. The cDNA template used was made by reverse transcription of total RNA extracted from cultured human periodontal ligament fibroblasts. The various recombinant human FN fragments that include or exclude the alternatively spliced domains EIIIB and EIIIA are shown in Fig. 1. Both the forward and reverse oligonucleotide primers for DNA amplification shown in Table 1 were synthesized (18-21 bases) corresponding to the 5’ and 3’ terminal sequences of the desired coding segment. The forward primer was designed to incorporate the Ndel site imme-

Protein expression and purification

Escherichia coli strain BL21(DE3) was transformed with pET 11b or pET 15b plasmid DNA with the various fragments of human FN cloned into them. FN7-10, FN7-10+B, and 7-10 were cloned into pET 15b and all other fragments were cloned into pET 11b. Overnight bacterial cultures were pelleted and the washed pellet resuspended in fresh LB medium with ampicillin (50 μg/ml). After a 3 hour period of growth in 1 liter of LB with ampicillin, IPTG was added at a final concentration of 0.2 mM and the cultures were shaken at 37°C for another 3 hours. The protocol for induction and purification of the expressed proteins has been described previously (Aukhil et al., 1993). All the expressed proteins were soluble (seen in the supernatant after cell lysis and centrifugation). Ammonium sulfate precipitation was the first step followed by ion exchange chromatography on a Mono-Q column (Pharmacia). The proteins eluted as single peaks at various salt concentrations ranging from 0.2-0.4 M NaCl. The peak fractions were pooled and dialyzed thoroughly against 20 mM Tris-HCl, pH 7.9. Analysis of the purified protein samples on SDS-PAGE revealed clean single bands. However, upon long term storage at 4°C (more than 4 weeks) some proteins showed minor bands suggesting limited proteolysis. The proteins cloned into pET 15b were purified by affinity chromatography on HIS-BIND resin columns (Novagen) that were charged with Ni2+. The bound proteins were eluted using Imidazole, dialyzed against 20 mM Tris-HCl, pH 7.4, were further purified on the Mono-Q column and the pure fractions finally dialyzed against 20 mM Tris-HCl, pH 7.9. Protein concentrations were determined by measuring the absorbance at 287 nm. The extinction coefficient for each protein was calculated and estimated as described previously (Aukhil et al., 1993). SDS-PAGE was carried out using 5-20% acrylamide gradient gels under reducing conditions and stained with Coomassie Blue.

Protein adsorption assay

The binding of pFN and the recombinant FN polypeptides was analyzed using an ELISA assay. Briefly, wells in 96-well flat bottom plastic plates (Falcon) were coated with various concentrations of each polypeptide in PBS (in triplicate) and incubated overnight at 4°C. The solution was removed and the wells were washed 3× in PBS. The
unbound sites within the wells were blocked by incubating heat-denatured BSA (2 mg/ml, denatured at 80°C for 10 minutes) for 1 hour. The wells were again washed 3× with PBS containing 0.5% Tween-20. The rat monoclonal antibody 16G3 was added (10 µg/ml, 100 µl/well) to each well and incubated for 2 hours at room temperature. The wells were then washed thoroughly 3× in PBS/Tween and then incubated with a peroxidase-conjugated goat anti-rat IgG for 2 hours. Finally, the wells were washed again in PBS/Tween and peroxidase activity was measured using o-phenylenediamine as a substrate. The reaction was stopped with 3 M H₂SO₄. The mount of polypeptides adsorbed to the well was expressed as the absorbance at 492 nm. Control wells where only heat-denatured BSA was added were used to determine background levels.

**Cell culture**

Nil.8M hamster fibroblasts were obtained from Dr Richard Hynes, MIT. Cells were cultured in DMEM-H (Dulbecco’s modified Eagle’s medium with high glucose) with 10% fetal bovine serum and antibiotics (penicillin, streptomycin and gentamycin). The evening before the cell adhesion assays, cells were detached with 0.05% trypsin, 0.5 mM EDTA and replated at about 50% confluence. The next day the cells were detached again and the effects of trypsin stopped by adding 0.5 mg/ml soy bean trypsin inhibitor. The cells were washed twice in PBS and then resuspended in DMEM-H (with no serum) and kept on ice until use.

**Gravity cell attachment assay**

The assay was carried out as described by Aukhil et al. (1993) and Joshi et al. (1993). Wells of 96-well flat bottom polystyrene plastic microtiter plates (Falcon 3912) were coated with 100 µl of various concentrations of the recombinant proteins in PBS overnight at 4°C. The wells were then washed in PBS and the unbound sites in the wells were blocked by adding 1 mg/ml heat denatured bovine serum albumin (BSA, denatured by heating to 80°C for 10 minutes, then cooled to room temperature) for 1 hour at room temperature. After washing the wells three times in PBS, each well was loaded with 100 µl DMEM containing 5-10x10⁵ cells suspended in the medium. The plates were incubated for 60 minutes at 37°C in an atmosphere of 5% CO₂. Following the incubation period, the wells were washed 3× with PBS to remove the unbound cells. The bound cells were fixed and stained with 3.7% formaldehyde and 1% Tween-20. The dishes were washed gently three times in PBS. The bound cells were fixed in 3.7% formaldehyde for 5 minutes at 4°C, washed again in PBS, and then placed into 24-well plates. After a 60 minute incubation at 37°C, the medium was removed and the wells with the glass coverslips were washed gently three times in PBS. The bound cells were fixed in 3.7% formaldehyde for 10 minutes, then washed in Tris-buffered saline (TBS): 50 mM Tris-HCl, 150 mM NaCl, 0.1% NaN₃, pH 7.6) for 5 minutes. The coverslips, with the cells attached to them, were incubated sequentially in 0.5% Triton for 7 minutes, TBS for 5 minutes, primary antibody to vinculin (mAB 7F9) for 60 minutes at 37°C, TBS for 5 minutes, secondary antibody (DAM-R, Chemicon, 1:25 in TBS), TBS for 5 minutes, dipped into water and then mounted on Mowiol (Calbiochem, Corp.). The coverslips were examined under a Zeiss fluorescence microscope and the images captured on T-MAX film (Kodak). To identify actin, coverslips were stained with phalloidin (Molecular Probes) at 1:500 dilution.

**Centrifugal cell adhesion assay**

96-well U-bottom polystyrene plastic microtiter plates (Falcon 3912) were coated with 100 µl/well of the different concentrations of the recombinant proteins in PBS by overnight adsorption at 4°C. The wells were washed and unbound sites blocked with 10 mg/ml of BSA in PBS. 100 µl of a cell suspension containing 5-10x10⁵ cells were placed in each well and the cells allowed to settle/adhere for 15 minutes at 37°C in an atmosphere of 5% CO₂. Following this incubation period, the plate was centrifuged at 250 g for 5 minutes at 4°C in a Sorval RT6000B centrifuge. The medium in the wells was gently removed and replaced with 17.5% formaldehyde with 0.005% Crystal Violet in PBS for fixation for at least one hour. The fixative was then removed and the wells washed gently in PBS. The effects of centrifugation on the adhesion of cells to the coated ligands was determined by examining the cells in the round bottom wells and scoring as follows. Wells that showed complete pelleting of the cells were scored as 0 (e.g. BSA coated wells), wells that displayed maximum resistance to centrifugation and showed adherent cells received a score of 3 (e.g. plasma fibronectin), wells with cells showing more adherent cells than pelleted cells received a score of 2, and finally cells that showed more pelleted cells than adherent cells received a score of 1. In addition, the center of each well was photographed.

**RESULTS**

**Biochemical characterization of the bacterial expression proteins**

A total of ten different expression proteins corresponding to the various type-III domains of human FN were made and their nomenclature is shown in Fig. 1. The oligonucleotide primers used to amplify the domain specific recombinant fragments of human FN are shown in Table 1. All the cDNA fragments were sequenced to verify the fidelity of DNA amplification by Taq polymerase. When cloned into the prokaryotic expression vector pET11b and induced by IPTG, all of them expressed the protein well with the yield varying between constructs. All the recombinant proteins were soluble and could be easily purified by a combination of ammonium sulfate precipitation followed by ion exchange chromatography on Mono-Q columns. When necessary, an additional step of gel filtration was carried out on Superdex-75 column. The purified expression proteins were analyzed by SDS-PAGE, and demonstrated high levels of purity (Fig. 2). Storage at 4°C (>4 weeks) in some instances led to a partial degradation, as evidenced by doublet bands that stained with the monoclonal antibody to domain #10 on western blots (data not shown). Hence experiments were carried out using fresh batches (within 2-3 weeks after purification) of recombinant FN fragments. The molecular masses

EIIB and EIIMA domains of fibronectin 2273
and extinction coefficients for each of these recombinant fragments are shown in Table 2. Fragments made in pET 15b with the histidine tag showed no difference in cell adhesion assays compared to similar fragments made in pET 11b without the histidine tag.

**Adsorption of recombinant FN fragments to plastic surface**

A modified ELISA was used to determine the binding of FN fragments to wells of plastic plates used for cell adhesion assays. Using the monoclonal antibody 16G3 to the 10th type-III repeat of FN (Nagai et al., 1994), the four large fragments (FN #7-12 with and without IIIIB and IIIA) and the four smaller fragments (FN #7-10+/--IIIIB; FN #10-12+/-IIIA) were tested for adhesion to plastic at concentrations ranging from 1 to 100 nM. The data are shown in Fig. 3 and all eight recombinant fragments showed adsorption to plastic wells. Since the monoclonal antibody was used at only one concentration, it was found to be insufficient giving a premature saturation reading for the binding of protein fragments. The apparent saturation of the protein fragments at 50 nM concentration was later found to be inaccurate since cell adhesion assays showed increasing adhesion for some fragments up to 100 nM concentration.

**Cell adhesion by gravity assays and effects on cell spreading**

The results from gravity cell attachment assays are shown in Fig. 4. Human plasma FN was used as positive control, and as expected, showed good adhesion at 50 nM concentration. The construct FN #7-10+B also showed adhesion at 50 nM that was comparable to plasma FN. The absence of spliced domain IIIIB from the FN #7-10 construct reduced the adhesion by approximately 50%, and even at concentrations of 200 nM had not reached that seen for FN #7-10+B at 50 nM (as per assays in multiwell plates). The constructs FN #10-12 with and without IIIIA showed significantly less adhesion of cells even at concentrations as high as 200 nM. In general, the presence of adjacent domains at the N-terminal end of the 10th type-III domain with the RGD sequence appeared to favor cell adhesion and this was enhanced when domain EIIIB was included. The significance of domains at the N-terminal end of the tenth domain (FN III #7,8,9) in enhancing adhesion was further confirmed by testing cell adhesion to larger fragments that contained type-III domains both at the N-terminal (FNIII # 7-9) and C-terminal ends (FNIII #11,12) of the tenth type-III domain. For example, as shown in Fig. 4, the order of cell adhesion to these fragments was (#7-12+B+A) > (#7-12+B+A) > (#7-12-B+A) > (#7-12-B+A). In both the smaller as well as the larger fragments, the inclusion of domain IIIA appeared to reduce the adhesion of cells while the inclusion of IIIB appeared to enhance it.

**Immunofluorescence staining for focal adhesions and cytoskeleton**

NIL cells plated on glass coverslips coated with the various recombinant FN fragments were stained with mAB to vinculin.
to examine the focal adhesions and with phalloidin to examine the actin network. Since the gravity adhesion assays had shown differences between the abilities of the various fragments to allow adhesion of cells at 50 nM concentrations, the coverslips were also coated at the same concentration. As shown in Fig. 5.1, the fragment FN#7-10+IIIB showed cell spreading and strong staining for focal adhesions that was comparable to native plasma FN (Fig. 5.9). The focal adhesions were also more prominent in cells plated on FN#7-10+IIIB (Fig. 5.1) than in cells plated on FN#7-10–IIIB (Fig. 5.2). The fragments FN#10-12–IIIA (Fig. 5.3) and FN#10-12+IIIA (Fig. 5.4) showed very few cells adhering and virtually no spreading. No significant focal adhesions were detectable in cells plated on these fragments. In the larger FN#7-12 fragments, the differences in focal adhesions were subtle; although the fragments FN#7-12+IIIB+IIIA (Fig. 5.5), FN#7-12–IIIB–IIIA (Fig. 5.6) and FN#7-12+IIIB–IIIA (Fig. 5.7) showed slightly more spreading of cells and focal adhesions compared to the large fragment FN#7-12–IIIB+IIIA (Fig. 5.8).

The organization of actin stress fibers was also examined in these cells by staining with phalloidin. Although the immunofluorescence was not as bright as that seen with staining for focal adhesions, the findings appeared to parallel those observed for focal adhesions and in the case of the larger fragments, the fragment FN7-12–IIIIB+IIIA showed a less intense staining for actin as compared to the other fragments (Fig. 6).

**Centrifugal cell adhesion assay**

This assay was designed as an indicator of the relative strengths of cell adhesion to the various recombinant FN fragments. Examples of the scoring criteria (0-3) are shown in Fig. 7. Cells plated on native plasma FN showed resistance to centrifugal detaching forces fifteen minutes after plating them. Other proteins that offered resistance to detaching centrifugal forces at low concentrations (50 nM) were FN#7-10+IIIB and FN#7-10–IIIB. The FN#10-12 fragments showed almost complete pelleting at 50 nM concentrations. In comparison, the larger

### Table 1. Oligonucleotide primers used to clone the fragments of human FN

<table>
<thead>
<tr>
<th>Primer Type</th>
<th>Primer Sequence and Translation</th>
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<tr>
<td>Type III domain 7</td>
<td>Fwd. ACAGGATCCATATGCGCATTTGTCCACCACAAAC</td>
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<tr>
<td>Type III domain 10</td>
<td>Fwd. GCCGGATCCATATGGTTTCTGATTTGTCAGAGG</td>
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<tr>
<td>Type III domain 10</td>
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<td>Type III domain 12</td>
<td>Rev. SINYRT* TTGCACCACACTTGAGTAAGGATCCCA</td>
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<tr>
<td>ED-B domain</td>
<td>Fwd. VTTLE* GATAGAATTCCATATGGAGGTGCCCAAGCTACT</td>
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<tr>
<td>ED-A domain</td>
<td>Fwd. TLQTQTT* CAGAGGATCCATATGACATCGCCCTAA</td>
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<tr>
<td>ED-A domain</td>
<td>Rev. MNIDRPK ATGGAAACGCAGTCCACATAAGGATCCCGCA</td>
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</table>

The reverse primer is listed as the sense strand (with translation), but the anti-strand was the sequence actually used. NdeI restriction sites are indicated by underline; BamHI sites are double underlined; EcoRI sites are indicated by -----. The asterisks show the stop codons.

### Table 2. Biochemical characterization of recombinant fragments of human FN

<table>
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<th>Peptide</th>
<th>Molecular mass</th>
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<th>Native protein</th>
<th>pET 11b</th>
<th>pET 15b</th>
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ε=(Y×1,400+W×5,600)/molecular mass. Y, Tyr; W, Trp.

OD278/ε = protein concentration (mg/ml).
FN#7-12 fragments showed mostly resistance to centrifugal detaching forces with the FN#7-12+IIIB–IIIA showing the best results followed by the FN#7-12+IIIB+IIIA fragment. These results were reproducible in three separate experiments using these recombinant fragments.

**DISCUSSION**

We have used recombinant fragments of type-III repeats of human fibronectin in the present study to examine the biological functions of the alternatively spliced domains EIIIB and EIIIA. The prokaryotic expression vectors pET15b and pET11b (with and without histidine tags respectively, for purification) yield recombinant fragments that possess biological activity in the form of promoting cell adhesion. The crystal structure of one four-domain fragment of human fibronectin fragment (FNIII 7-10), refined to 2.0 Å, prepared as described here, has been recently solved (Leahy et al., 1996), suggesting that the recombinant fragments fold properly. We encountered no technical difficulties in the production and purification of the fibronectin fragments with all the fragments being soluble in the bacterial lysate and easily purified by a combination of gel filtration, affinity (for hist-tag proteins made in pET 15b) and ion-exchange chromatography. The cloning of fragments based on amplification of cDNA to begin and terminate each protein at the boundaries of the independently folding domains successfully yields stable fragments of the FN-III domains. We had successfully used a similar approach to map the cell- and heparin-binding domains of human tenascin (Aukhil et al., 1993). In the present study, by using different combinations of type-III fragments 7-12 of human FN that include or exclude the spliced repeats EIIIB and EIIIA, we show that including the EIIIB repeat enhances the adhesion and spreading of cells. On the other hand including repeat EIIIA alone reduced the adhesion and spreading of cells, while including it along with EIIIB made no difference in terms of cell adhesion and spreading.

Recombinant fragments of human FN were made to include, in addition to the type-III repeat #10 with the putative RGD cell
binding loop, adjacent domains on both sides. Previous studies have shown that the presence of adjacent type-III domains #9 and #8 are needed to restore the ability of the putative RGD-containing domain #10 to support cell adhesion and spreading (Kimizuka et al., 1991). This along with the location of domains EIIIB (between 7 and 8) and EIIIA (between 11 and 12) dictated the design of our fragments that extended from repeats 7-12. The finding that inclusion of domain EIIIB alone in fragments FN 7-10 and 7-12 enhances the adhesion and spreading of cells is interesting and points to several possibilities. The first possibility is that the EIIIB domain itself has cell-adhesive properties via a different integrin, or uses the same alpha5 beta1 integrin. This was not found to be the case because domain EIIIB alone or fragment 7+EIIIB did not promote cell adhesion and were not successful in inhibiting adhesion to the 7-10+B fragment in competitive assays. Domain EIIIB alone showed no adhesion at concentrations of up to 10 μM. In competitive inhibition experiments (data not shown), trypsinized cells incubated with 50 μM of purified fragment EIIIB in DMEM for 15 minutes with gentle shaking at 37°C before plating in wells coated with 50 nM of the fragments (FN, 7-10, 7-10+B) showed no inhibition of cell adhesion. The second possibility is that including domain EIIIB would generate a conformational modification in the domains downstream, perhaps improving the

**Fig. 5.** Distribution of focal adhesions in cells plated on the various recombinant human FN fragments. Glass coverslips were coated with the proteins at 50 nM concentration and unbound sites blocked with BSA. Cells were plated for 1 hour following which the glass coverslips were rinsed in PBS and processed for immunofluorescence staining of focal adhesions using antibody to vinculin. The coverslips were coated as follows: 1= 7-10+B; 2= 7-10–B; 3= 10-12+A; 4= 10-12–A; 5= 7-12+B+A; 6= 7-12–A–B; 7= 7-12+B–A; 8= 7-12–B+A; 9= pFN.
Table 3. Data from centrifugal cell attachment assay

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<th>7-10+B</th>
<th>7-10−B</th>
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<td>400 nM</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>1.5</td>
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<td>3</td>
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<td>3</td>
<td>N.D.</td>
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<tr>
<td>1,000 nM</td>
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<td>N.D.</td>
<td>N.D.</td>
<td>2</td>
<td>2</td>
<td>N.D.</td>
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N.D., not determined.

Fig. 6. Distribution of actin stress fibers in cells plated on the various recombinant human FN fragments. Glass coverslips were coated with the proteins at 50 nM concentration and unbound sites blocked with BSA. Cells were plated for 1 hour following which the glass coverslips were fixed, subsequently rinsed and the actin stress fibers stained with phalloidin. The coverslips were coated as follows: 1= 7-10+B; 2= 7-10−B; 3= 10-12+A; 4= 10-12−A; 5= 7-12+B-A; 6= 7-12−A-B; 7= 7-12+B−A; 8= 7-12−B+A; 9= pFN.
access to integrin binding sites in domain 10 and 9. To this effect, a previous study by Carinemolla et al. (1992) had found that including the EIIIB repeat in their β-galactose-fibronectin fusion protein had unmasked cryptic sequences suggesting conformational effects of including EIIIB. More recently, the crystal structure of the recombinant human FN fragment 7-10 (Leahy et al., 1996) suggests some interesting structural consequences. The fragment FN 7-10, made up of four contiguous type-III modules shows relative rotations ranging from 43 to 160°. In addition, the solvent-accessible surface area buried between domains is as follows: 587 Å² between domains 7 and 8, 527 Å² between domains 8 and 9, and only 333 Å² between domains 9 and 10. It is also interesting to note that an almost twofold rotation axis relates domain 8 and 9. In summary, the small rotation and tilt angles relating domains 9 and 10 renders this segment of the 7-10 fragment almost cylindrical and contrasts with the combined tilts and rotations relating domains 7-9 giving that segment a spiral configuration (Leahy et al., 1996). While definitive information on the structural effects of including EIIIB would require crystallographic analysis, it is tempting to speculate on the structural aspects. It is possible that the large interface between domains 7 and 8 contains sequences that may have synergistic functions in promoting the adhesion and spreading of cells. While these interface sequences are buried and may be masked in the [-EIIIB] isoform, the inclusion of EIIIB may be exposing these proximal sequences. The other possibility is that the inclusion of EIIIB may affect the polarity of the domains downstream resulting in the positioning of the RGD cell-binding site in domain 10 and the other recently-discovered synergy sequences (PHSRN) in domain 9 (Aota et al., 1994) to enhance the adhesion and spreading of cells. Since domain 9 contains more basic amino acids and is located immediately before domain 10, an extra domain that may influence the proper folding and conformation of the positively charged region in domain 9 and the RGDS site could be enhancing the biological function of the synergy sequences (Kimizuka et al., 1991). The final possibility is that the NIL cells may be using another receptor that may be recognizing other sequences as ligands in addition to the α5β1 integrin used to bind to the RGD sequence. This is currently being investigated using blocking antibodies to α5 integrin in cell adhesion assays.

Recent studies by Chen and Culp (1996) and Xia and Culp (1994) have reported some cell adhesion to fragments EIIIB and EIIIA that were made as independent domains or in conjunction with their adjacent domains (7-B-8 and 11-A-12). However, the experimental conditions were very different with observation periods of 4 hours. The observation period in our experiments on cell adhesion was 1 hour for adhesion/spreading and only 15 minutes for the centrifugal assay. Our findings on the larger fragment 7-12+A differ from those reported by Xia and Culp (1994) where they report adhesion activity in domain EIIIA. The differences could be explained by the duration of adhesion assay (4 hours versus 1 hour in our study), concentration of the proteins used and cell types used. The cells that adhered to fragment 7-10+B had spread just as well as pFN at 50 nM concentrations. Similarly, the distribution of the actin cytoskeletal network was comparable. However, the 7-10–B fragment

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**Fig. 7.** Centrifugal cell adhesion assay. Round-bottom wells in 96-well plates were coated with recombinant fragments of human FN overnight. After blocking the unbound sites with heat-denatured BSA, cells were plated and allowed to adhere for 15 minutes. The entire 96 plate was then centrifuged at 250 g for 5 minutes. The medium was then gently aspirated and the cells fixed with 3.7% formaldehyde/0.1% Crystal Violet overnight. The wells were rinsed again and the center of the round-bottom wells examined and photographed. (A) Examples of scoring criteria are 0 for complete pelleting, 1 for mostly pelleting in the center with a few cells resisting detachment around the periphery, 2 for cells that are mostly resisting detachment but a few clusters at the center and, 3 for cells that have resisted detachment with no pelleting or clusters either in the center or in the periphery. (B) Examples of centrifugal assays on the various recombinant fragments of human FN. BSA is the negative control giving 0 scores and pFN is the positive control giving scores of 3.
showed slightly fewer focal adhesions and less well-organized actin network. The fact that the fragments 10-12 with and without EIIIA showed very little adhesion and virtually no spreading is not surprising given the previous findings that the RGD-containing domain alone is not sufficient to promote cell adhesion. Also, these findings suggest that domains 11 and 12 at the C-terminal end do not offer any kind of synergistic help to the putative RGD-containing domain 10. Again, structural studies (Leahy et al., 1996; Main et al., 1992) imply the need for additional specific domains on the N-terminal end of the RGD loop, perhaps acting to stabilize the integrin. One of the interesting findings in the present study is the improved spreading and distribution of focal adhesions in cells plated on the larger 7-12 fragments. Here, the inclusion of EIIIB alone or both EIIIB and EIIIA appears to help adhesion and spreading of cells, while the inclusion of EIIIA in the absence of EIIIB keeps the cells from establishing good focal adhesions. The findings on the distribution of focal adhesions and actin stress fibers complement those of the centrifugal cell adhesion assay. This assay was designed to indirectly test the adhesive strength by examining the balance between centrifugal forces and cell substrate adhesion. In non-adsorbing or weakly adhesive substrata, cells may form a pellet due to the predominance of centrifugal force. With increasing strengths of cell-substrate adhesion, cells resist centrifugal forces. Under the criteria used for scoring, a score of 3 means almost complete resistance to detachment by centrifugal forces in round-bottom wells. It should be emphasized that the cells were allowed to adhere only for 15 minutes. Cells on pFN showed scores of 3 at concentrations as low as 10 nM and fragments 7-10-B and 7-10-A were next, showing scores of 3 at about 50 nM concentrations. The larger fragments showed scores of 3 at 100 nM concentrations only when EIIIB alone was included or both EIIIB and EIIIA were excluded. Cells on the fragment 7-12-A did not show scores of 3 even at the highest concentration tested (600 nM). Similarly cells plated on the smaller fragments 10-12-A and 10-12-B never reached scores of more than 2 even at 1,000 nM concentrations. Taken together, the data on distribution of focal adhesions and the centrifugal cell attachment assay suggest that the inclusion of EIIIB enhances the adhesion and spreading of cells while including EIIIA alone in the larger fragment negatively affects the adhesion and spreading of cells. It is tempting to speculate that the inclusion of both domains EIIIB and EIIIA in the FN molecule may be a biological mechanism of achieving cell adhesion to a degree that facilitates the process of locomotion where both adhesion and loss of adhesion are needed for the advancement of cells. The EIIIB+/EIIIA+ isoformal of FN is common during embryonic development and wound healing (french-Constant and Hynes, 1989), both events where cell migration is important. More recently, it has been shown that the spliced domains EIIIB and EIIIA are included in the FN seen during the morphogenesis of ‘FN dependent’ structures such as somites, notochord and the vasculature (Peters and Hynes, 1996). Similarly, the expression of the EIIIA and EIIIB spliced variants in the adult tissues suggest that these isofoms of FN may be serving different functions (Peters et al., 1996).

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REFERENCES


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