A Material-Based Platform to Modulate Fibronectin Activity and Focal Adhesion Assembly

Frankie A. Vanterpool,1,2,* Marco Cantini,1,* F. Philipp Seib,2 and Manuel Salmerón-Sánchez1

Abstract

We present a detailed characterization of fibronectin (FN) adsorption and cell adhesion on poly(ethyl acrylate) (PEA) and poly(methyl acrylate) (PMA), two polymers with very similar physicochemical properties and chemical structure, which differ in one single methyl group in the lateral chain of the polymer. The globular solution conformation of FN was retained following adsorption onto PMA, whereas spontaneous organization of FN into protein (nano) networks occurred on PEA. This distinct distribution of FN at the material interface promoted a different availability, measured via monoclonal antibody binding, of two domains that facilitated integrin binding to FN: FNIII10 (RGD sequence) and FNIII9 (PHSRN synergy sequence). The enhanced exposure of the synergy domain on PEA compared to PMA triggered different focal adhesion assemblies: L929 fibroblasts showed a higher fraction of smaller focal plaques on PMA (40%) than on PEA (20%). Blocking experiments with monoclonal antibodies against FNIII10 (HFN7.1) and FNIII9 (mAb1937) confirmed the ability of these polymeric substrates to modulate FN conformation. Overall, we propose a simple and versatile material platform that can be used to tune the presentation of a main extracellular matrix protein (FN) to cells, for applications than span from tissue engineering to disease biology.

Key words: biomaterials; cell culture; cellular biology; extracellular matrix

Introduction

Providing cells with a reliable and robust microenvironment, able to control and understand their response, is a crucial issue for regenerative medicine and biological research in general.1 Traditional systems lack control of the intermediate layer of absorbed proteins that govern cell response and eventually direct cell fate.1,2 It is widely accepted that cell adhesion onto a synthetic surface is mediated by a layer of proteins previously adsorbed onto the material, either from the physiological fluids in vivo or from the culture medium in vitro. This layer may also be intentionally deposited, for example, through adsorption of extracellular matrix (ECM) proteins, or via chemical attachment to the substrate.1–5 These proteins include fibronectin (FN), vitronectin, and fibrinogen, known as the soluble matrix proteins of biological fluids.7 Cell adhesion is the first step in their response to a microenvironment, and plays a fundamental role in subsequent cell differentiation, growth, viability, and phenotypic response.6 Therefore, controlling the cellular response through a robust material-based strategy is applicable for a myriad of applications in biomedical research, spanning from regenerative medicine to disease biology.

FN is a high-molecular-weight glycoprotein that is found in both soluble and insoluble forms in extracellular fluids and connective tissues, respectively.7–9 It is a main component of the ECM and therefore a likely protein candidate whose state can be tailored to target cell response. The FN molecule is a dimer of ~440 kDa, whose subunits contain three types of repeating units (FN repeats I, II, and III) that mediate interactions with other ECM proteins and cell surface receptors, and regulate FN-FN interactions (Fig. 1A). The RGD sequence, located in the FN repeat III10, is of key importance in controlling cell adhesion, with further regions, such as the PHSRN sequence (synergy site), in the FN repeat III9, promoting α5β1 integrin binding to FN (Fig. 1B).9

Cells have evolved to recognize adhesive proteins through integrins, a family of trans-membrane heterodimers that serve as a link between the ECM and the cell actin cytoskeleton.11 Subsequent to binding, integrins cluster and develop into complexes, called focal adhesions, composed of structural (e.g., vinculin) and signaling molecules (e.g., focal

1Division of Biomedical Engineering, School of Engineering, University of Glasgow, Glasgow, United Kingdom.
2Strathclyde Institute of Pharmacy and Biomedical Sciences, University of Strathclyde, Glasgow, United Kingdom.
*These two authors contributed equally to this work.
adhesion kinase). These plaques anchor cells to the surface and their formation triggers cell response.\textsuperscript{12} Overall, FN adopts a fundamental role in mediating and promoting cell adhesion, and in regulating cell survival and phenotypic response.

Several studies have demonstrated that surface chemistry is a key parameter that can alter the amount and conformation of adsorbed FN.\textsuperscript{13–20} For example, using differently terminated self-assembled monolayers (SAMs), García and coworkers demonstrated that hydrophilic surfaces (OH-SAM) induced the least amount of structural change in FN or in the recombinant fragment FNIII\textsubscript{7–10}, while hydrophobic CH\textsubscript{3}-SAM provoked the largest structural changes.\textsuperscript{13,14} Surface chemistry can therefore be used to manipulate the state and activity of the protein through simple physical adsorption. We have previously demonstrated that poly(ethyl acrylate) (PEA) surfaces induce the organization of FN into physiological-like fibrillar (nano) networks upon adsorption onto the polymeric substrate.\textsuperscript{20,21} However, the removal of a methyl group from the monomer [i.e., using poly(methyl acrylate) (PMA) instead] prevented this phenomenon, leading to FN adsorption in a globular conformation. In this work, we characterized the state of the protein on these surfaces in relation to protein concentration and surface chemistry. We used L929 fibroblast, plated on engineered FN substrates, and determined the cellular response in relation to the presentation of FN at the material interface. Overall, we intended to establish a simple and robust tissue culture platform that enables us to control the state of the intermediate adsorbed protein layer, thereby tailoring the microenvironment presented to cells (Fig. 1). We propose that this material-based platform is a versatile tool to both direct and elucidate mechanisms underlying cell response for applications spanning from tissue engineering to disease biology.

### Materials and Methods

#### Samples preparation

PEA and PMA polymers were synthetized by radical polymerization of ethyl acrylate and methyl acrylate (Sigma, St. Louis, MO), initiated by benzoin at 1 wt\% and 0.35 wt\%, respectively. Samples were then dried by vacuum extraction to constant weight and solubilized in toluene 2.5% w/v for PEA and 6% w/v for PMA. PEA and PMA solutions were then spin coated onto 12 mm round glass coverslips for 30 sec at a velocity of 2000 and 3000 rpm, respectively, with an acceleration of 3000 rpm/sec. Excess toluene was removed by placing coverslips under the vacuum for 2 h at 60°C.

#### Protein adsorption

Human plasma FN (Sigma) solutions, in Dulbecco’s phosphate buffered saline (DPBS) at 2, 5, 10, and 20 µg/mL, were coated onto PEA and PMA coverslips for 1 h.

#### Water contact angle

Water contact angle analysis was carried out on PEA or PMA surfaces alone and on FN-coated surfaces. For each condition, the static contact angle (SCA), advancing contact angle (ACA), and receding contact angle (RCA) were determined ($n = 9$). SCA was determined by placing a drop of 3 µL of water on the surface using a needle and recording the images at 12 frames/sec for 30 sec while measuring the angle of the drop with the polymer surface. ACA was determined by placing the needle in the previously deposited drop and progressively adding water in order to observe an increase in the length of the baseline. RCA was determined by progressively removing the water with the needle until the drop was removed. RCA is the angle at which the baseline starts
decreasing. Measurements were carried out using a Theta optical tensiometer (Biolin Scientific, Stockholm, Sweden).

**Immunostaining of FN**

Immunostaining was done with polyclonal anti-FN (Millipore, Darmstadt, Germany) primary antibody directed against FN. PEA and PMA samples were coated with FN at different concentrations for 1 h. Freshly prepared samples were fixed with formaldehyde 3.7% v/v for 30 min at 4°C and then washed with PBS. Samples were incubated with anti-FN (1:400 dilution in PBS/BSA 1% w/v) for 1 h at room temperature and then washed twice with DPBS/Tween-20 0.5% v/v. Next, samples were incubated with Cy3 anti-mouse secondary antibody (1:200 in PBS/BSA 1% w/v) for 1 h at room temperature and then washed twice and mounted with vectashield (Vector Laboratories, Inc., Burlingame, CA) and imaged using an epifluorescence microscope (Zeiss AXIO Observer Z1, Jena, Germany).

**Atomic force microscopy**

Several areas of the surface were scanned at different area sizes and line rates: \(5 \times 5 \text{nm}^2\) (0.5 Hz); \(2 \times 2 \text{nm}^2\) (0.7 Hz); \(1 \times 1 \text{nm}^2\) (1 Hz); \(0.5 \times 0.5 \text{nm}^2\) (1 Hz). The surface height, lock-in phase, and lock-in amplitude were observed in tapping-mode atomic force microscopy (AFM; Nanowizard 3 from JPK, Berlin, Germany) using cantilevers with a force constant of 3 N/m, a resonance frequency of 75 kHz, and a pyramidal tip, with radius of curvature less than 8 nm (MPP-21120 from Bruker, Billerica, MA).

**Micro-bicinchoninic acid protein quantification**

PEA, PMA, and glass coverslips were coated with FN for 1 h at different solution concentrations and the density of adsorbed protein was determined by measuring the amount of nonadsorbed FN. A stock solution of FN was diluted at 20, 10, 5, and 2 g/mL to determine the exact concentration of the stock FN solution and the amount of FN deposited onto the coverslips. After coating for 1 h the FN solution was collected and transferred to a 96-well plate followed by the addition of the bicinchoninic acid working reagent (Thermo Fisher Scientific, Waltham, MA). The plate was then shaken for 30 sec, covered, and placed in an incubator for 2 h at 37°C and then absorbance was read at 562 nm with a Tecan NanoQuant Infinite M200 Pro plate reader (Männedorf, Switzerland).

**Enzyme-linked immunosorbent assay**

Enzyme-linked immunosorbent assay (ELISA) was performed on PEA and PMA coated with different concentrations of FN using primary monoclonal antibodies HFN7.1 (DSHB, Iowa City, IA) and mAb1937 (Millipore) combined with HRP-Goat Anti-Mouse (Invitrogen, Carlsbad, CA) secondary antibody, which is coupled with a peroxidase for absorbance reading. FN was adsorbed at 20 and 1 µg/mL as well as a control without protein on PEA and PMA for 1 h. Samples were washed with PBS, transferred to new 24-well plates, and then blocked with PBS/BSA 1% w/v for 30 min at room temperature. Subsequently, samples were incubated with the primary antibodies for 1 h at room temperature and washed with PBS/Tween-20 0.5% v/v. The secondary antibody (HRP-Goat Anti-Mouse) was then added on all samples for 1 h at room temperature in the dark and washed with PBS/Tween-20 0.5% v/v. The substrate (color reagents A and B from R&D Systems, Minneapolis, MN) was then added for 20 min at room temperature, in the dark, stopped with a stop solution (R&D Systems), and transferred to a 96-well plate and absorbance was read at 450 and 550 nm.

**Attachment assay**

This was done according to the protocol by Humphries.22 UV-sterilized PEA and PMA samples were coated with FN (20 µg/mL) for 1 h and washed twice with PBS. Samples were then blocked with heat-denatured BSA 1% w/v (Roche, Basel, Switzerland) for 30 min at room temperature. During this period, cells were harvested, trypsinized, and re-suspended in complete medium (containing 10% FBS). The cell suspension was then incubated for 10 min at 37°C, 5% CO₂.

Cells were seeded onto the surfaces at a seeding density of \(8.5 \times 10^4 \text{cells/cm}^2\) for 20 min at 37°C, 5% CO₂. Surfaces were washed twice with PBS to remove cells that were not firmly attached, fixed with formaldehyde 3.7% for 20 min at 4°C, and washed again. Samples were permeabilized for 5 min using a Triton X-100 based permeabilization buffer (0.5% v/v Triton X-100, 10.3 w/v% saccharose, 0.292% w/v NaCl, 0.06% w/v MgCl₂, and 0.476% w/v HEPES adjusted to pH 7.2) and mounted with vectashield with DAPI to stain the nuclei. Images were taken using an inverted epifluorescence microscope and the number of cells attached was determined using CellC total cell count analysis (Zeiss AXIO Observer Z1).

In the case of blocking cell binding domains on FN, the same procedures were carried out using only samples coated with 20 µg/mL of FN and blocking the RGD and PHSRN sequences with HFN7.1 and mAb1937 at a 1:1 molar ratio (14.6 µg/mL) for 1 h at room temperature.

**Early adhesion assay**

PEA- and PMA-coated coverslips were sterilized for 20 min using a UV lamp and coated with FN at different concentrations (2, 5, 10, and 20 µg/mL) for 1 h. Samples were then washed twice with PBS and seeded with cells at \(1 \times 10^5\) cells per well for 3 h. After washing with PBS, samples were fixed with formaldehyde 3.7% for 20 min at 4°C and washed again before being stored in DPBS at 4°C until ready to use. In the case of cell adhesion after blocking specific FN domains, PEA and PMA were incubated for 1 h with 150 µL of HFN7.1 and/or mAb1937 at a 1:1 molar ratio.
Antibodies were diluted in 1% BSA w/v. Samples coated with 20 μg/mL of FN only were then permeabilized (as described above) and stained for 1 h using a primary antibody against mouse vinculin hVIN-1 (Sigma-Aldrich, St. Louis, MO), which stains the focal adhesions. The secondary antibody (Cy3 anti-mouse), coupled with actin staining Phalloidin, was then added for 1 h in the dark.

Samples were then mounted with vectashield with DAPI to stain the nuclei and visualized using an epifluorescence microscope. Images were taken and channels merged using ImageJ (1.47v) to localize nuclei, actin, and focal adhesions. The latter was quantified using the focal adhesion analysis server. Data such as focal adhesion size, area, and distance from the center of the cell were obtained from the server and used to calculate the geometric moment of inertia ($I$) of cells

$$ I = \sum a_i r_i^2 $$

where $a_i$ is the area of the $i$th focal adhesion and $r_i$ is the distance between the $i$th focal adhesion and the center of the distribution of focal adhesions. Cell morphology, describing shape, was analyzed using ImageJ (1.47v).

**Results**

**FN adsorption**

Material substrates used in this study were chemically similar, consisting of a vinyl backbone chain with side group $-\text{COO}(\text{CH}_2)_x\text{H}$, with $x=1$ for PMA and $x=2$ for PEA. The difference of only one methyl group in the side group yielded substrates with similar physicochemical properties. Both surfaces are classified as hydrophobic, with a static contact angle of 75° (Fig. 2A,D), and similar behavior in dynamic contact angles, although PEA showed more hysteresis as defined by the difference between advancing and receding angles (Fig. 2D). Contact angle hysteresis has been interpreted as a measure of molecular mobility for surfaces that are chemically homogeneous and flat.

Spin-coated samples were smooth ($R_{\text{rms}}<1$ nm) and homogeneous, with a thickness of ~1 μm. When the surfaces were coated with FN from solution concentrations of 2, 5, 10, and 20 μg/mL, the static and advancing contact angles were maintained, while the hysteresis increased for both polymers due to a decrease of receding angles, reaching a plateau for concentrations higher than 2 and 5 μg/mL for PMA and PEA, respectively (Fig. 2D). Furthermore, contact angle hysteresis of FN-coated samples was higher on PEA (e.g., ~90° compared to ~65° on PMA, for concentrations greater than 5 μg/mL), suggesting a different state of the adsorbed protein layer. Immunostaining of FN adsorbed onto the polymeric substrates resulted in a different distribution of the protein at the microscale; FN forms densely packed networks on PEA, while globular aggregates appear
FIG. 3. FN adsorption and domain availability on PEA and PMA. (A) Surface density of FN on PEA and PMA coated from solutions of 1 and 20 μg/mL of FN. (B) Availability of the RGD (located within FNIII₉) cell binding domain (left) and the PHSRN (located within FNIII₁₀) synergy domain (right) on PEA and PMA coated from solutions of concentration 1 and 20 μg/mL of FN. Monoclonal antibodies against these two domains of FN HFN7.1 (FNIII₁₀) and mAb1937 (FNIII₉) were used. ***p < 0.001, **p < 0.01, *p < 0.05.

on PMA (Fig. 2B). The different FN distributions were maintained regardless the concentration of the protein solution (Supplementary Fig. S1). AFM images confirmed structural differences of FN at the nanoscale (Fig. 2C). PEA was seen to unfold the FN arms that led to interconnected nanofibrils; in contrast, PMA surfaces promoted FN adsorbed in its globular conformation, confirming our previous results.²¹,²⁷ The surface density of adsorbed protein was indirectly quantified by measuring the depletion of FN from solution. The surface density of adsorbed FN increased with solution concentration (Fig. 3A), though it remained constant between surfaces, with no statistically significant differences (e.g., ~750 ng/cm² on PEA and ~600 ng/cm² on PMA when FN is adsorbed from a 20 μg/mL solution).

To analyze the exposure and availability of integrin binding domains on FN following adsorption onto the polymeric surfaces, binding of conformation-specific monoclonal antibodies, in combination with an ELISA, was used. This is an established method to probe for structural or conformational changes of adsorbed proteins²⁸,²⁹ and can be used to measure FN activity. The selected antibodies recognized the central binding domain of FN, namely, HFN7.1. HFN7.1 is directed to the flexible linker between the 9th and 10th type III repeats of FN and has been identified as a receptor-mimetic probe for integrin binding and cell adhesion.¹³ In contrast, mAb1937 binds to the 8th type III domain,¹⁴,³⁰ therefore closer to the synergy domain (Fig. 1B). At very low concentration (1 μg/mL), FN biological activity is higher on PMA, as the binding is higher for both antibodies (Fig. 3B). However, increasing FN concentration to 20 μg/mL resulted in increased RGD accessibility on PEA that matched PMA levels, though binding of mAb1937 was significantly higher on PEA (Fig. 3B). Considering the change of FN activity with increasing FN concentration, there is a steady increase of the binding of both antibodies on PEA, while on PMA the binding of mAb1937 remains constant (Fig. 3B).

Cell attachment and adhesion

Short-term cell culture experiments were performed to investigate attachment and initial adhesion of L929 mouse fibroblasts to the surfaces. Cell attachment to the different materials was measured using a standard protocol, allowing 20 min for attachment.²² The assay showed that the presence of an FN coating is essential for short-term cell attachment, as no cells were found on both bare polymer surfaces (Fig. 4A). When the polymers were coated with FN from a 20 μg/mL solution concentration (~600 ng/cm² according to Fig. 3A), cell attachment was similar on all surfaces (~90%), even if samples were washed extensively with DPBS before cell counting.

To study how cells responded to the FN-coated surfaces, 3 h adhesion experiments were performed in serum-free conditions after coating the surfaces from solutions containing 2, 5, 10, and 20 μg/mL of FN. Cells were seeded at a low density (5,000 cells/cm²) to maximize cell–material interactions and to minimize cell–cell interactions. Fibroblasts were found to spread on all surfaces, with the actin cytoskeleton becoming progressively more developed at increasing protein concentrations; at a minimum concentration of 2 μg/mL, the actin cytoskeleton was not completely developed and only peripheral stress fibers were observed (Fig. 4C and Supplementary Fig. S2). The spreading area was similar on all surfaces for the different protein concentrations, with the exception of coatings from a 20 μg/mL FN solution, where it was significantly higher on PEA (Fig. 4B).

Focal adhesion plaques were more developed on surfaces coated with higher FN concentration, where they were located at the end of prominent stress fibers (Fig. 5 and Supplementary Fig. S2). In order to quantify the maturation level of the focal adhesions on the different surfaces, frequency distributions for their size (defined as the length of the major axis of the focal adhesion plaque) were obtained through image analysis of the vinculin images (process detailed in Supplementary Fig. S3). Focal complexes (dot-like complexes shorter than 1 μm) were discarded from the analysis. The frequency distribution was similar on all surfaces, with a monotonic decrease from a higher fraction of smaller plaques. On PEA, the fraction of smaller focal plaques diminished with increasing FN concentration, as longer focal adhesions appeared. For example, the fraction of
plaques longer than 2 \mu m increased from \sim 10\% for an FN concentration of 2 \mu g/mL to \sim 30\% for a concentration of 20 \mu g/mL (Fig. 4). For PMA polymeric substrates the change in the frequency distribution was less pronounced (from \sim 10\% to \sim 20\%).

This was confirmed by the median of the distribution; on PEA it showed a monotonic increase, while on PMA it remained constant (Fig. 6A). This trend of longer plaques became more apparent by using the focal adhesion area instead of length (Fig. 6A). By comparing the frequency distribution of the size of the focal adhesions to their respective area, it was evident that on PMA the plaques are prevalently dot-like compared to PEA, especially at lower FN concentration. As a refined measure of cell adhesion, the moment of inertia was used. This is a purely physical read out that examines the focal adhesion imprint relative to the center of mass of the distribution (Eq. 1). Overall, the moment of inertia of fibroblasts was significantly higher on 20 \mu g/mL FN-coated PEA surfaces compared to PMA (Fig. 6C).

The relevance of the RGD and the synergy domains on cell attachment and adhesion was studied by performing short-term cell culture assays on FN-coated samples after blocking these domains with HFN7.1 and mAb1937 monoclonal antibodies, respectively. For these studies, only surfaces coated with 20 \mu g/mL FN solutions were used. Blockage of the central binding domain of FN using HFN7.1 led to significantly reduced cell attachment of less than 10\% (Fig. 7A). However, blocking FN with mAb1937 did not reduce the number of cells attached to either polymer. When both antibodies were employed, cell attachment on PMA was inhibited to the same levels (10\%) as for HFN7.1 blocking, while on PEA the percentage of attached cells was significantly higher (40\%). Next, we addressed the effect of blocking these FN domains on the number, size, and distribution of focal adhesions. Blocking with either HFN7.1 or mAb1937 (and both of them) significantly
diminished the number of focal adhesions on PMA, whereas a monotonic decrease was found on PEA (from HFN7.1 to mAb1937 and then both of them; Fig. 7B).

The same trend was found for the size of the corresponding focal adhesions, which were smaller on PEA after blocking with HFN7.1, and essentially absent after blocking with mAb1937 or both antibodies (Fig. 7C). The size distribution of focal adhesions can be observed in Supplementary Figure S4. However, blocking with HFN7.1 did not alter the moment of inertia, and focal adhesions were still too far away from the center of the cell to end up in approximately the same value for this magnitude (Fig. 7D).

**FIG. 5.** Focal adhesion assembly on PEA and PMA coated with FN. (A) Representative inverted binary representation of focal adhesions of L929 on PEA and PMA coated with different concentrations of FN. (B) Images in (A) were quantified to build size distribution histograms of focal adhesions on both polymers. Scale bar: 50 μm.

**FIG. 6.** Assessment of focal adhesions on PEA and PMA. (A) Median of focal adhesion size (left) and focal adhesion area on PEA and PMA coated with FN solutions of 2, 5, 10, and 20 μg/mL (right). Note that the slope for PEA (0.0101x + 0.00203) is statistically different from 0 for the median of focal adhesion area, which supports the increasing size of focal adhesions as the concentration of FN increases as well as the difference between focal adhesion size on PEA and PMA. p < 0.05. (B) Moment of inertia for the distribution of focal adhesions calculated using Equation 1 for cells on PEA and PMA coated with different concentrations of FN. **p < 0.01.
Conversely, using mAb1937 and the combination of both antibodies drastically decreased the number and size of focal adhesions and consequently the moment of inertia as well.

**Discussion**

It is well recognized that surface chemistry modulates FN adsorption and its conformation on the material surface. Seminal studies by Garcia et al. showed that by using model surface chemistries (SAMs with well-defined chemical groups, i.e., −OH, CH₃, NH₂, COOH) the integrin-binding domain of FN (FNIII₂⁻₁₀) can be presented to cells with altered biological activity after adsorption on material surfaces. Modulation of FN activity at the material interface led to changes in integrin binding, organization of focal adhesions, and signaling that promoted cell differentiation with different efficiencies. Focal adhesions are supramolecular structures that link adsorbed FN at the material interface with the actin cytoskeleton. Focal adhesion structure, size, composition, and dynamics depend on the information received from the surrounding matrix. Availability of key FN domains can promote integrin clustering, and this in turn can determine cell signaling and fate. Moreover, the composition, function, and size of focal adhesions have been correlated to each other: focal complexes ~1 µm are involved in migration and low-tension phenotypes that contain paxillin, vinculin, and phosphorylated proteins; focal adhesions ~2–5 µm are involved in intermediate tension phenotypes; super-mature adhesions > 5 µm are involved in high-tension phenotypes.

Here we show that FN conformation and distribution can be fine-tuned by using material surfaces with very similar chemical and physical chemistries. PEA and PMA consist of a vinyl chain with a side group that differs by only one methyl group (Fig. 2A). This subtle change in the underlying chemistry does not alter significantly the hydrophilicity of the surface (Fig. 2D) and both samples are sensed as simply rigid substrates by cells. Adsorbed FN on both PEA and PMA remained constant regardless of the concentration of the adsorbing solution (Fig. 3A). However, the micro-/nanoscale distributions of FN differed significantly, with globular aggregates on PMA compared to an interconnected FN (nano) network on PEA (Fig. 2C). The different state of the adsorbed protein on the two polymers was also confirmed by dynamic contact angle measurements: contact angle hysteresis was significantly higher on FN-coated PEA due to a stronger decrease of the receding angles compared to PMA. This might suggest a higher protein surface coverage on PEA, compatible with the unfolding of the dimer arms and the formation of fibrils, compared to the maintenance of a globular conformation on PMA. Also, the extended conformation of FN on PEA might favor the molecular rearrangement of the protein in contact with water compared to the compact conformation on PMA.

The different FN presentation on the material surface has important consequences at the molecular level for the availability of the integrin binding region of FN (FNIII₀⁻₁₀). Importantly, after FN adsorption from a solution of concentration of 20 µg/mL, the availability of the RGD domain remained constant for both PEA and PMA, whereas the synergy sequence (PHSRN) located at the III₀ domain was preferentially available for cell engagement on PEA (Figs. 1C and 3). This has important consequences in terms of integrin binding and focal adhesion assembly. It has been shown that subtype binding to FN requires both the RGD sequence (FNIII₀⁻₁₀) and the synergy domain (FNIII₀). This observation also translates to cell adhesion on FN-coated PEA, where cell attachment occurred preferentially via subtype, in contrast to subtype, which was mostly used for cells to adhere to FN adsorbed onto PMA. This biological response was triggered through FN presentation, which in turn was influenced by the underlying material surface. We used vinculin as a marker of focal adhesions because it is recruited at adhesion sites where adhesion occurs via subtype or subtype receptors. In addition, vinculin is required for myosin contractility-dependent adhesion strength and the coupling of cell area with traction force.
force. The formation (including size) of focal adhesions depends on the mechanical state of the local cell microenvironment. Stiff substrates and the application of mechanical inputs (stress and strain) involve the development of large focal adhesions, whereas soft substrates and the use of inhibitors of contractility favors the formation of focal-complex-like adhesions. Moreover, nanotopography—including nanopits, nanopillars, and nanogrooves—has been shown to alter the size and orientation of focal adhesions, including integrin clustering, focal adhesion size, and cytoskeleton organization.

Here we show that conformation and distribution of FN on surfaces with very similar physicochemical properties (i.e., PEA and PMA) can be used to alter focal adhesion organization. Smaller focal adhesions were found on globular FN on PMA (focal complexes), whereas larger and better developed adhesions were quantified on PEA (Fig. 5). The size distribution of focal adhesions remained constant while the concentration of FN on the surface increased on PMA. However, on PEA, the distribution showed a higher fraction of larger focal adhesions as the concentration of FN on the surfaces increased (this can also be observed using the median—even if this is a poor representation of the whole distribution; Fig. 6). Larger focal adhesions on PEA and larger cells (Fig. 4B) resulted in higher values for the moment of inertia (Fig. 6). Larger focal adhesions on PEA and larger cells (Fig. 4B) resulted in higher values for the moment of inertia (Fig. 6C). The stress applied by cells on substrates has been estimated to be ~5.5 nN/μm². It is then suggested that cells are better positioned to exert larger forces on FN fibrils assembled on PEA than on globular FN adsorbed on PMA, even if both surfaces are sensed as rigid by cells (E above 1 MPa for both of them).

Blocking experiments with monoclonal antibodies directed against FNIII10 (HFN7.1) and FNIII9 (mAb1937) further supported the notion that PEA and PMA induced a polymer-specific biological response. Blocking FNIII10 (which prevents the availability of RGD) drastically diminished the number of cells on both surfaces, an effect not observed when blocking FNIII9 (synergy sequence) (Fig. 7A). Therefore, when the synergy sequence of FN (PHSRN) was not available for interaction (on PEA), cells were still able to adhere using only the RGD domain (and α/β) rather than αββ. Likewise, the number and size of focal adhesions was significantly diminished on PMA after blocking with either antibody, whereas a significant number of focal adhesions (but smaller) were still found on PEA after blocking each one of the domains, or both simultaneously (Fig. 7 and Supplementary Fig. S4). This result directly correlated with the extended conformation of FN molecules on PEA (that leads to the organization into nanonetworks observed by AFM; Fig. 2) and the globular-like shape of the protein on PMA. Even after blocking FNIII10, other domains were still available for integrin binding on PEA (e.g., FN1 and the variable region) but not on PMA, as a change in the conformation of the molecule (fibrillogenesis) was required for these domains to be exposed (Fig. 1D).

Overall, we present a mechanistic model for the different conformation of FN induced by the underlying material surface that accounts for the availability of the integrin binding domains (FNIII3,10) and the effect of blocking antibodies (HFN7.1, mAb1937) as discussed (Fig. 1).

Conclusions

We have shown that FN conformation can be triggered by material surfaces with very similar chemical composition (PEA and PMA differ in only one methyl group in the side chain) and physical properties (wettability and charge). FN is organized in nanonetworks on PEA, while a globular conformation was observed on PMA. As a consequence, FN molecules were adsorbed in an extended conformation on PEA, which provides better availability of the synergy sequence and other integrin-binding domains of the molecule (e.g., FN10). The distinct organization of FN on PEA led to a higher fraction of larger focal adhesions, whereas smaller focal-complex points were observed on PMA. The maintenance of focal adhesions, even after blocking FNIII10 with monoclonal antibodies on PEA, but not on PMA, confirmed the different conformation of FN on these polymeric surfaces. Our results suggest that cells are highly sensitive to the organization of FN on material surfaces regardless of the properties of the underlying polymeric coating. In summary, PEA and PMA can be used as a simple material platform to fine-tune the microenvironment presented to cells by controlling the intermediate layer of protein adsorbed onto the substrates. This material-based modulation of FN is a versatile tool to study and understand cell behavior in biomedical research, including tissue engineering and disease biology.

Acknowledgment

The support from ERC through HealInSynergy 306990 is acknowledged.

Author Disclosure Statement

The authors declare no competing financial interests exist.

References

MATERIAL-BASED MODULATION OF FIBRONECTIN


Address correspondence to:
Prof. Manuel Salmerón-Sánchez
Division of Biomedical Engineering
School of Engineering
University of Glasgow
Rankine Building, Oakfield Avenue
Glasgow G12 8LT
United Kingdom

E-mail: manuel.salmeron-sanchez@glasgow.ac.uk

Marco Cantini, PhD
Division of Biomedical Engineering
School of Engineering
University of Glasgow
Rankine Building, Oakfield Avenue
Glasgow G12 8LT
United Kingdom

E-mail: marco.cantini@glasgow.ac.uk

---

<table>
<thead>
<tr>
<th>Abbreviations Used</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACA = advancing contact angle</td>
</tr>
<tr>
<td>AFM = atomic force microscopy</td>
</tr>
<tr>
<td>DPBS = Dulbecco’s phosphate buffered saline</td>
</tr>
<tr>
<td>ECM = extracellular matrix</td>
</tr>
<tr>
<td>ELISA = enzyme-linked immunosorbent assay</td>
</tr>
<tr>
<td>FBS = fetal bovine serum</td>
</tr>
<tr>
<td>FN = fibronectin</td>
</tr>
<tr>
<td>PEA = poly(ethyl acrylate)</td>
</tr>
<tr>
<td>PMA = poly(methyl acrylate)</td>
</tr>
<tr>
<td>RCA = receding contact angle</td>
</tr>
<tr>
<td>SAMs = self-assembled monolayers</td>
</tr>
<tr>
<td>SCA = static contact angle</td>
</tr>
</tbody>
</table>